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# A molecular view of DNA flexibility

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#### **Abstract**

DNA dynamics can only be understood by taking into account its complex mechanical behavior at different length scales. At the micrometer level, the mechanical properties of single DNA molecules have been well-characterized by polymer models and are commonly quantified by a persistence length of 50 nm (~150 bp). However, at the base pair level (~3.4 Å), the dynamics of DNA involves complex molecular mechanisms that are still being deciphered. Here, we review recent single-molecule experiments and molecular dynamics simulations that are providing novel insights into DNA mechanics from such a molecular perspective. We first discuss recent findings on sequence-dependent DNA mechanical properties, including sequences that resist mechanical stress and sequences that can accommodate strong deformations. We then comment on the intricate effects of cytosine methylation and DNA mismatches on DNA mechanics. Finally, we review recently reported differences in the mechanical properties of DNA and double-stranded RNA, the other double-helical carrier of genetic information. A thorough examination of the recent single-molecule literature permits establishing a set of general 'rules' that reasonably explain the mechanics of nucleic acids at the base pair level. These simple rules offer an improved description of certain biological systems and might serve as valuable guidelines for future design of DNA and RNA nanostructures.

#### **Table of contents**

Introduction	1
Sequence-dependent DNA mechanics	2
TpA dinucleotides are highly flexible	3
A dual role of A-tracts in DNA flexibility	4
CpG islands are rigid DNA regions depleted of nucleosomes	4
Poly(dA-dT) and poly(dG:dC) adopt A-like DNA conformations	6
Poly(dG-dC) are hotspots of Z-DNA formation	6
GC-content not always explains DNA melting	8
Chemical modifications and DNA mechanics	8
The impact of cytosine methylation on DNA physical properties	8
The effect of DNA mismatches on DNA mechanics	9
Mechanical properties of dsRNA: unexpected differences with dsDNA	10
Stretching flexibility of dsDNA and dsRNA	11
The opposite twist-stretch coupling of dsDNA and dsRNA	11
The different dynamics of plectoneme formation	13
Opposite effects of complex ions on the mechanics of dsDNA and dsRNA	13
Sequence determinants of intrinsic bending	13
Sequence-dependent mechanical properties	14
Conclusions and future perspectives	14
Sequence-dependent DNA mechanics beyond the elastic regime	14
Toward a better characterization of cytosine methylation and DNA mismatches in	
the context of DNA mechanics	15
DNA and RNA mechanics at the service of nanotechnology	15

#### Introduction

The physical properties of DNA are interrogated in virtually every process that involves storage or manipulation of the genetic information encoded in the double-helix. DNA packaging

inside eukaryotic nuclei requires bending of the DNA around the histone octamer; and gene regulation in bacteria often involves DNA looping in regulatory regions. Transcription and DNA replication require local melting of the double-helix and produce a torsional stress in the DNA that is absorbed via the formation of plectonemes. In the pursue of a quantitative characterization of these and other biological processes, intensive research has been devoted to study the mechanical properties of DNA (Bloomfield et al., 2000). In particular, in the last few decades, our understanding of DNA mechanics has been propelled by the advent of single-molecule techniques, which enable manipulating and/or measuring individual DNA molecules. Initial singlemolecule experiments on DNA stretching and supercoiling (Smith et al., 1992; Strick et al., 1996) showed that the mechanical properties of long (several kilobase pairs) DNA molecules are well described by polymer models that neglect the microscopic details of the duplex (Marko and Siggia, 1994, 1995a, 1995b). Since then, these polymer models have resulted extremely useful to quantify, for instance, the action of proteins on the DNA, and are currently employed to calibrate single-molecule biophysical instruments on a daily basis (Bouchiat et al., 1999; Bustamante et al., 2003; Madariaga-Marcos et al., 2018).

Despite the unquestionable success of polymer models, recent single-molecule assays are encouraging an alternative, i.e. microscopic description of DNA flexibility (Wiggins et al., 2006; Lipfert et al., 2010; Vafabakhsh and Ha, 2012; Lebel et al., 2014; Shon et al., 2019; Pyne et al., 2020). Ultimately, DNA is a highly sophisticated molecule with fine molecular features such as major and minor grooves, stacking interactions, or hydrogen bond donor and acceptors, that are inextricably linked to the flexibility of the double-helix. For example, dinucleotides that present weak stacking interactions are prone to induce a sharp bend in the DNA (Olson et al., 1998); sequences with a narrow minor groove resist mechanical deformations (Nelson et al., (1987); and exotic hydrogen bonds found in certain sequences can assist complex deformations of the double-helix (Dans et al., 2014; Pasi et al., 2014). Consequently, despite behaving as a homogeneous polymer at long scales, DNA presents an intricate flexibility at short distances that is strongly dependent on the nucleotide sequence (see Fig. 1). In addition, non-canonical DNA base pairs including e.g. methylated cytosines or mismatched base pairs, can substantially alter the mechanics of DNA at such short scales (see Fig. 1). Thus, given that DNA:protein interactions usually occur at the nanometer scale, it is paramount to deepen the molecular description of DNA flexibility, which will possibly provide additional insights into biological phenomena beyond the reach of classical long-range polymer approaches. Moreover, a molecular characterization of DNA mechanics may pave the way for expanding the nanotechnological applications of DNA as building material.

Here, we review recent studies that are advancing our understanding of single-molecule DNA mechanics from such a molecular perspective. We focus on two approaches that are driving major progress in this field. First, the development of new single-molecule assays that permit assessing novel mechanical properties of DNA, especially at short-length scales. Second, the synergistic combination of single-molecule experiments and computer simulations, most notably molecular dynamics (MD), that model the mechanics of DNA with atomic details. This review aims to discuss recent studies that illustrate the capability of these two approaches at providing novel relevant insights into double-stranded DNA mechanics. Thus, it is not our intention to deepen in any particular aspect of DNA mechanics or to include

non-helical DNA structures, such as G-quadruplexes or i-motifs (see Abou Assi *et al.*, 2018; Mandal *et al.*, 2019 for reviews on these topics).

This review has been divided into three sections. In Section 'Sequence-dependent DNA mechanics', we comment on how the nucleotide sequence affects the physical properties of DNA. We review sequences that facilitate or preclude DNA bending; sequences prone to adopt a double-helical structure that differs from the canonical B-DNA; and sequences where DNA melting is more favorable. In Section 'Chemical modifications and DNA mechanics', we discuss the effect of modifications of the canonical duplex, in particular, methylated cytosines and DNA mismatches. In Section 'Mechanical properties of dsRNA: unexpected differences with dsDNA', we cover recent studies that are revealing interesting differences between the mechanics of DNA and double-stranded RNA (dsRNA), the other double-helical carrier of genetic information that is found in nature. Finally, we present the conclusions of the review and suggest future lines of research.

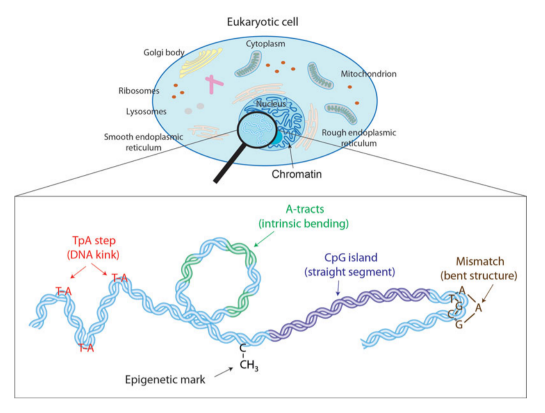
The findings on DNA mechanics hereby reviewed have often provided a new perspective on diverse biological systems, as discussed throughout the text. An exciting challenge for the near future will be to exploit our ever-growing knowledge on nucleic acids mechanics for controlled design of functional DNA and RNA nanostructures.

#### Sequence-dependent DNA mechanics

In single-molecule experiments, the mechanical properties of DNA are commonly quantified by means of its persistence length,  $P_{\rm DNA}$ . If one thinks of a long shoelace or a (cooked) spaghetti immersed in a swimming pool, one does not picture a perfectly straight line, but rather, a somewhat coiled object. The same occurs for DNA. The size of the DNA polymer in solution, as given by the distance between its two ends, is quantified by  $P_{\mathrm{DNA}}$ . An alternative – perhaps more intuitive – way of looking at  $P_{\mathrm{DNA}}$  is the threshold DNA length above which the molecule will start to bend. In other words, DNA molecules shorter than P<sub>DNA</sub> are expected to be essentially straight. Several singlemolecule experimental setups, including magnetic tweezers (MT), optical tweezers (OT) (Smith et al., 1992, 1996; Bustamante et al., 1994; Wang et al., 1997), and atomic force microscopy (AFM) imaging (Rivetti et al., 1996; Heenan and Perkins, 2019), allow extracting the value of  $P_{DNA}$  with high accuracy. These and many other single-molecule techniques yield, under standard buffer conditions, a consensual value of  $P_{\rm DNA} \sim 50$  nm, which corresponds to  $\sim 150$  bp (e.g. the length of DNA in the nucleosome core).

Note, however, that the accepted value of  $P_{\rm DNA}{\sim}150~{\rm bp}$  contrasts with several biological evidences where the DNA is severely bent at distances comparable to or shorter than 150 bp. Examples of such bends include DNA loops at regulatory regions or the wrapping of nucleosomal DNA around the histone core (reviewed in Garcia *et al.*, 2007). According to the commonly accepted worm-like chain (WLC) model, such short-scale deformations would be prohibitive for a DNA molecule with  $P{\sim}150~{\rm bp}$ .

This conundrum motivated studies on the local mechanics of single DNA molecules (Wiggins *et al.*, 2006; Vafabakhsh and Ha, 2012). Such studies revealed a striking ability of the duplex to adopt strongly bent conformations, an aspect that was not contemplated by the standard WLC model (Wiggins *et al.*, 2006; Vafabakhsh and Ha, 2012). Although the quantitative aspects of short-scale DNA bending remain a subject of debate, it is clear



**Fig. 1.** Effectors of DNA mechanical properties at short-length scales. DNA is highly compacted in the chromatin inside the nucleus of a cell. We will show along the text that multiple effectors have been described to modulate the DNA mechanical properties at short-length scales. These examples comprise kinkable TpA steps, intrinsic bending by A-tracts, rigid CGIs, mismatches that produce strong bending, and cytosine methylation that exert a versatile role in DNA physical properties.

that sharp DNA bending can proceed via concerted localized distortions of the duplex, such as DNA kinks (Vologodskii and Frank-Kamenetskii, 2013). More interestingly, the nucleotide sequence strongly affects these local deformations (Vafabakhsh and Ha, 2012), in contrast to the long-range persistence length, which shows little sequence-dependent variation (Geggier and Vologodskii, 2010). Namely, sequence effects on DNA mechanics appear amplified at short-length scales.

#### TpA dinucleotides are highly flexible

DNA kinks were predicted long ago (Crick and Klug, 1975) and have been observed in crystal structures of a number of DNA:protein complexes (Berman et al., 1992; Olson et al., 1998), including nucleosomal DNA (Olson and Zhurkin, 2011); and also in several MD simulations (Lankaš et al., 2006; Curuksu et al., 2009; Irobalieva et al., 2015; Harrison et al., 2019). The biological consequences of DNA kinks are diverse, and have been reviewed in the context of DNA wrapping in the nucleosome core (Richmond and Davey, 2003; Olson and Zhurkin, 2011) and of specific DNA:protein interactions (Rohs et al., 2010). Crystal structures of DNA:protein complexes reveal that kinks are more favorable in pyrimidine-purine steps, CpG and CpA, and particularly in the TpA step, which has shown larger flexibility (Olson et al., 1998) (see Table 1 for nomenclature). However, structural studies provide a static description of the DNA and thus, are of limited use in order to address the dynamics of kink formation.

The development of novel single-molecule assays to interrogate the flexibility of short DNA molecules is a promising tool to probe the impact of kinkable pyrimidine-purine steps on the

physical properties of DNA. Of particular relevance is the creative single-molecule assay developed by Vafabakhsh and Ha (2012). By attaching a pair of fluorescence resonance energy transfer (FRET) dyes to DNA molecules and measuring the FRET signal, the authors were able to observe single-molecule DNA cyclization in real time (see Fig. 2a). In that study, the authors showed that the cyclization dynamics, as quantified by the *J*-factor, was much faster than the WLC prediction, supporting the existence of strongly bent DNA structures, as proposed in a previous AFM study (Wiggins *et al.*, 2006).

More importantly, this single-molecule cyclization assay was employed to explore the effect of the nucleotide sequence on DNA flexibility. In a later study, the same group studied the cyclization dynamics of the two halves of the 601 Widom sequence (a strong nucleosome positioning sequence) (Ngo et al., 2015). They found that the left half, which contains four kinkable TpA steps, was highly flexible, whereas the right half, which only contains one of such steps, was relatively rigid (see Fig. 2a). Notably, when this rigid half was modified to include three additional TpA steps, it became much more flexible, albeit it was still more rigid than the left, flexible half. This experiment shows that even few TpA steps can substantially impact the flexibility of DNA molecules comprising tens of base pairs. Further single-molecule studies should aim to address whether this effect is also present in CpG and CpA steps, which have also been traditionally identified as highly flexible (Olson et al., 1998).

A deeper understanding of the physical properties of DNA sometimes offers a fresh perspective on certain biological processes. This is well illustrated in the study by Ngo *et al.* (2015). Besides the aforementioned cyclization experiments, the authors

Table 1. Nomenclature used for DNA and dsRNA sequences

Term	Definition
NpM step	A dinucleotide of sequence 5'-NM-3', e.g. TpA denotes 5'-TA-3'
Poly(dN:dM)	DNA sequences consisting of several consecutive nucleotides, e.g. poly(dA:dT) denotes sequences such as 5'-AAAAAAA-3'
Poly(dN-dM)	DNA sequences consisting of alternating N and M nucleotides, e.g. poly(dA-dT) denotes sequences of the form 5′-ATATATATA-3′ or 5′TATATAT-3′
A-tract	DNA sequences with three or more consecutive adenines and/or thymines, without the flexible TpA step. Examples are 5′-AATT-3′, 5′-TTTTTT-3′, but NOT 5′-TTAA-3′ (which contains a TpA step)
CpG island	DNA sequences with high proportion of CpG steps, e.g. 5'-ACGAGCGGCGTCG-3' or 5'-CGTCGAGCGTCGGGCG-3'
AU-tract	dsRNA sequences consisting of several (three or more) alternating adenines and uracils, e.g. 5'-rArUrArUrArUrArUrA' or 5'rUrArUrA-3'

Note: The sequences shown here refer to a canonical Watson-Crick double-helix where the reverse strand is omitted for clarity, e.g. the sequence 5'-AGTACCC-3' refers to a DNA double-helix with forward strand 5'-AGTACCC-3' and reverse strand 5'-GGGTACT-3'. Ribonucleotides are referred to as rA, rC, rU, and rG.

used a combined setup of OT and single-molecule FRET (smFRET) to observe single unwrapping events in nucleosomes. This novel assay permits visualizing which side of the nucleosomal DNA is detached from the histone core when tension is applied. They observed that reconstituted nucleosomes with the aforementioned 601 sequence showed highly asymmetric unwrapping. In particular, upon the action of an external force, the right rigid half of the nucleosomal DNA was significantly more prone to unwrap than the left, flexible half. On the contrary, nucleosomes reconstituted with the 601 sequence containing additional TpA steps in the right half showed an approximate symmetric unwrapping. Namely, when the two halves of nucleosomal DNA had a similar flexibility, nucleosome unwrapping occurred stochastically from either side. Since the publication of the study by Ngo et al. (2015), a number of experimental and computational studies have addressed the link between DNA flexibility and transient nucleosome unwrapping, or nucleosome breathing (Lequieu et al., 2016; Zhang et al., 2016; Chen et al., 2017; Culkin et al., 2017; Mauney et al., 2018; Winogradoff and Aksimentiev, 2019). These studies have revealed that, besides its well-known role in positioning nucleosomes (Widom, 2001; Segal et al., 2006; Kaplan et al., 2009), the DNA sequence can largely affect the accessibility of certain regions within nucleosomal DNA via changes in the flexibility of the duplex.

### A dual role of A-tracts in DNA flexibility

In contrast to the flexibility of TpA steps, other DNA sequence motifs are thought to be extremely rigid. The most well-known examples are A-tracts, a kind of DNA sequence that consists of three or more adenines and thymines without the flexible TpA step. Poly(dA:dT)s, a particular case of A-tracts, possess a strong nucleosome depleting character that has been associated to a presumably high rigidity of this motif, as predicted from structural studies and atomistic simulations (Nelson *et al.*, 1987; Haran and Mohanty, 2009; Segal and Widom, 2009; Dršata *et al.*, 2014).

The flexibility of A-tracts in general, and poly(dA:dT)s in particular, are controversial, because they are inextricably intertwined with the long-known intrinsic bending induced by these sequences (see, e.g. Zhang and Crothers, 2003; Thompson and Travers, 2004; Haran and Mohanty, 2009; Peters and Maher, 2010; Ortiz and de Pablo, 2011). For example, previous AFM and tether particle motion experiments indicated that A-tracts have no enhanced bending rigidity, showing a standard persistence length of ~50 nm (Rivetti et al., 1998; Brunet et al.,

2015a). However, these results contrast with coarse-grain simulations and single-molecule cyclization experiments, which suggest that A-tracts are rigid to bending (Vafabakhsh and Ha, 2012; Mitchell *et al.*, 2017).

In a recent study, these paradoxical mechanical properties of A-tracts were comprehensively studied at the single-molecule level using a wealth of techniques: MT, OT, and AFM imaging in air and in liquid (Marin-Gonzalez et al., 2020a). The authors considered a sequence from the Caenorhabditis elegans genome with several repetitions of phased A-tracts, expected to display a strongly bent character (Fire et al., 2006). Indeed, AFM measurements demonstrated that phased A-tracts induce an intrinsic bend in the DNA that could be well-described with a variant of the WLC model that includes intrinsic bending (see Fig. 2b) (Rezaei et al., 2018; Marin-Gonzalez et al., 2020a). Moreover, both AFM and OT measurements independently showed that, at long-length scales, the A-tract molecule was not particularly rigid (nor flexible) to bending, showing a persistence length of 54 nm. However, as the A-tracts were subjected to high forces (F > 10 pN), an extraordinary rigidity started to emerge, that was quantified by a large stretch modulus (see Fig. 2b). Such high forces are expected to align the intrinsic A-tracts bends, thus enabling to probe the local rigidity of the A-tract structure. It was thus proposed that, although A-tract might not seem rigid at long distances, the stiff local structure of the A-tract would resist short-scale mechanical deformations, which are likely more determinant for nucleosome formation.

We can thus conclude that A-tracts play a dual role in DNA flexibility. On the one hand, they induce a static bending in the double-helix, but on the other hand the structure of the A-tract itself appears rigid. It is tempting to state that the balance of these two effects can be regulated by the distribution of A-tracts along a given DNA region. That is, several short A-tracts in phase with the helical pitch would amplify the bending, whereas a long (i.e. >20 bp) individual A-tract would stiffen the DNA. This intriguing dual mechanism appears to be exploited *in vivo* in the context of nucleosome positioning: short, phased A-tracts are enriched in nucleosome positioning sequences Rohs *et al.* (2009), whereas long poly(dA:dT)s are characteristic of nucleosome depleting regions (Segal and Widom, 2009).

### CpG islands are rigid DNA regions depleted of nucleosomes

Another example of DNA motif whose biological functions might depend on peculiar mechanical properties are CpG islands (CGI),

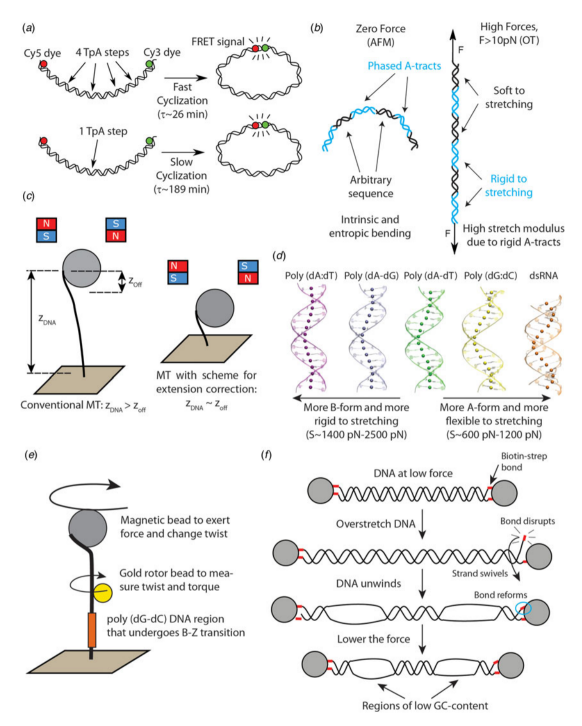


Fig. 2. Recent single-molecule and MD studies on sequence-dependent DNA mechanics. (a) The single-molecule assay developed by Vafabakhsh and Ha (2012) consists of a pair or FRET dyes attached to the extremes of a DNA molecule with cohesive ends. Cyclization results in an increase in the FRET signal, thus enabling measuring single-molecule cyclization in real time. TpA steps were found to increase DNA flexibility, resulting in faster cyclization kinetics Ngo et al. (2015). (b) Cartoon illustrating the complex mechanical properties of A-tract sequences. A-tracts located in phase with the DNA helical pitch induce a global macroscopic curvature in the molecule. When the molecule is subjected to high forces, the bends are straightened and the A-tracts are found to present a large stretching rigidity. Thus, A-tracts induce DNA bending, but the tracts themselves are rigid at a local level. Adapted from Marin-Gonzalez et al. (2020a). (c) MT are usually limited to measurements of DNA molecules with contour length longer than the bead radius. This limitation is overcome in the MT scheme for extension correction developed in Shon et al. (2019), which enables estimating the anchor point of the DNA on the bead. Using this correction scheme, accurate MT force-extension curves can be obtained for DNA molecules as short as  $\sim$ 200 bp. (d) The crookedness curvature reported in Marin-Gonzalez et al. (2019a) is responsible for sequencedependent variations of the DNA extension and distinguishes between A- and B-DNA conformations. Curved sequences, such as poly(dG:dC) are A-like and flexible to stretching; straight sequences, such as poly(dA:dT) are highly B-like and rigid to stretching. Adapted from Marin-Gonzalez et al. (2019a). (e) The gold rotor bead assay from Lebel et al. (2014) combines high-resolution torque spectroscopy and controlled stretching and supercoiling of DNA molecules. A magnetic bead permits exerting force and supercoiling DNA molecules; whereas a gold bead attached to the side of the molecule reports on its twist and torque. This system has been used to study the B-Z transition of a poly(dG-dC) sequence with high temporal resolution. (f) Assay for DNA supercoiling using OT, as reported in King et al. (2019). Upon overstretching, one of the biotin-streptavidin interactions that attach the DNA to the optical beads is disrupted. The DNA unwinds and then the bond reforms, trapping the DNA in a negatively supercoiled state. When the force is lowered, melting bubbles are observed in regions of low GC-content.

GC-rich DNA regulatory regions with a high proportion of CpG steps. In vertebrates, CGI are ubiquitous sites of transcription initiation, a feature that has been linked with a deficient assembly of CGI into nucleosomes (both *in vivo* and *in vitro*) (Ramirez-Carrozzi *et al.*, 2009; Deaton and Bird, 2011). As for poly(dA:dT), it is likely that the depletion of nucleosomes at CGI is related to a peculiar rigidity of these sequences (Marin-Gonzalez *et al.*, 2019*a*; Shon *et al.*, 2019; Pongor *et al.*, 2017)

Recent developments in single-molecule techniques have enhanced our understanding of the mechanical properties of CGI. Shon et al. (2019) developed a novel scheme for in-situ correction of the DNA extension in MT, which enables obtaining accurate force-extension curves of DNA molecules as short as 198 bp. This development extends the capability of conventional MT, which are usually limited to DNA molecules longer than ~1 kbp; and thus optimizes this technique for better exploration of sequence effects on DNA flexibility (see Fig. 2c). Using their novel correction scheme, Shon et al. found that a CGI DNA molecule possessed a large persistence length when compared with a control DNA of arbitrary sequence, indicative of a high bending rigidity of CGI. Interestingly, this and other studies have proposed that the flexibility of CGI is highly sensitive to the methylation state of these sequences (Marin-Gonzalez, 2019a; Shon et al., 2019; Pongor et al., 2017), suggesting that epigenetic marks might exert regulatory functions in CGI via changes in the mechanical properties of the DNA. This will be discussed in Section 'The impact of cytosine methylation on DNA physical properties'.

Despite these insightful studies, further inquiry is clearly needed to better understand the mechanical properties of CGI. First, there is some discrepancy among different single-molecule techniques regarding the large rigidity of CGI (Pongor *et al.*, 2017), which should be solved in future studies. Such studies should take into account that CGI not only have a high GC-content, but also a high density of CpG steps. In fact, it is expected that the frequency and distribution of CpG steps would play a more relevant role in the mechanics of DNA than the GC-content itself (Marin-Gonzalez *et al.*, 2019*a*). In parallel to single-molecule studies, computer simulations should aim at explaining the molecular mechanisms behind the proposed rigidity of CGI. This is a challenging task, given the complex deformability exhibited by the CpG step, which is highly anharmonic and dependent of its sequence context (Dans *et al.*, 2014).

#### Poly(dA-dT) and poly(dG:dC) adopt A-like DNA conformations

In addition to DNA flexibility, the nucleotide sequence can strongly impact the conformation of the double-helix. In particular, some sequences, such as poly(dG:dC), have been identified to possess structural features close to the A-form (A-philic), whereas others such as poly(dA:dT) are often classified as highly B-like (B-philic) (Lu *et al.*, 2000). Yet, another class include poly(dA-dT) sequences, which adopt a so-called TA-DNA structure, or hybrid between B- and A-forms (Lu *et al.*, 2000; Kulkarni and Mukherjee, 2017). This rich conformational landscape of the double-helix is crucial for achieving DNA sequence recognition by several proteins that read specific features of the B- or A-forms (Lu *et al.*, 2000; Rohs *et al.*, 2010) (see Table 2).

Several structural parameters have been proposed as a measure of the A-philicity of a given DNA sequence, i.e. of its propensity to assume an A-like conformation (Lu *et al.*, 2000; Waters *et al.*,

2016; Kulkarni and Mukherjee, 2017). However, calculating these parameters requires knowledge of the atomic structure of the molecule, which is often inaccessible to single-molecule experiments. Is it possible to assess the A-philicity of a DNA sequence without knowing its atomic structure?

Recently, a promising approach has been proposed. Inspired by previous studies on dsRNA (Chou et al., 2014; Lipfert et al., 2014), Marin-Gonzalez et al. (2019a) suggested a connection between the A-philicity of a DNA sequence and its stretching flexibility. The authors performed MD simulations of several DNA sequences and they observed significant variations in the extension from one sequence to another. They noticed that these changes in the extension were a consequence of a sequencedependent DNA curvature, denoted crookedness, that reasonably correlated with the A-philicity of the sequence (see Fig. 2d). For example, the poly(dG:dC), which is highly A-philic, showed a short extension and large crookedness curvature; whereas the B-philic poly(dA:dT) highly was essentially Interestingly, those DNA molecules that were more crooked, and more A-like, were more flexible to a stretching force; and vice versa, highly B-like sequences were more rigid. Among the latter are poly(dA:dT) and poly(dC-dG), whose large stretching rigidity are supported by recent experiments on A-tracts and CGI (Pongor et al., 2017; Marin-Gonzalez et al., 2020a).

It is thus tempting to state that the stretching flexibility of a given DNA sequence is a good indicator of its A-philicity. This hypothesis is supported by naïve intuition. Note that a perfectly straight textbook B-DNA helix should be extremely hard to stretch. In such an ideal B-DNA, elongations beyond the contour length should result in unwinding and unstacking of the base pairs, which would easily disrupt the double-helix. This was already realized by Bustamante and coworkers in their seminal single-molecule MT experiment, where they proposed that the stretching elasticity of DNA should be a consequence of a 'local curvature of the DNA axis' (Smith *et al.*, 1992). It is plausible that such 'local curvature' consists of sequence-dependent deviations from the straight B-form into a more curved A-form (see Fig. 2d).

However, further research is needed to better characterize the DNA A-philicity, *crookedness* and stretching flexibility, in order to establish a more solid connection between these DNA features. Along this line, a recent study by Lankas and coworkers has expanded our understanding on DNA crookedness, by assessing how this parameter depends on the temperature of the system (Dohnalová *et al.*, 2020).

#### Poly(dG-dC) are hotspots of Z-DNA formation

Besides the right-handed helical structures discussed above, DNA is able to assume, under certain conditions, a left-handed conformation with a zig-zag backbone known as Z-DNA (Rich and Zhang, 2003). This DNA structure has been observed at sequences of alternating purines and pyrimidines, most notably poly(dG–dC), and is thought to require high salt concentrations or negative torsional stress in order to form (Rich and Zhang, 2003). The structural features of Z-DNA (and also Z-RNA) are recognized *in vitro* by certain proteins (those containing the so-called  $Z\alpha$  domain, see Table 2). However, the *in vivo* relevance of Z-DNA is debated (Rich and Zhang, 2003; Herbert, 2019; Jiao *et al.*, 2020). In this regard, some studies point toward transient formation of Z-DNA as regulatory mechanism during

Quarterly Reviews of Biophysics

Table 2. Sequence-dependent DNA physical properties

Sequence motif	DNA physical properties	Reference	Biological implications	Reference
Pyrimidine-purine steps (mainly TpA)	Highly flexible	Ngo <i>et al</i> . (2015) <sup>a</sup> Olson <i>et al</i> . (1998)	Stabilize nucleosomes, DNA:protein recognition via kinks	Ngo <i>et al.</i> (2015) <sup>a</sup> Lowary and Widom (1998); Widom (2001); Rohs <i>et al.</i> (2010)
Short (~4–10 bp) phased A-tracts	Intrinsic bending B'-DNA with narrow minor groove	Rivetti et al. (1998; Moreno-Herrero et al. (2006) <sup>a</sup> Koo et al. (1986); Nelson et al. (1987); Haran and Mohanty (2009)	Stabilize nucleosomes, Promote supercoiling, DNA:protein recognition via minor groove electrostatics	Kim et al. (2018b) <sup>a</sup> Rohs et al. (2009); Rohs et al. (2010)
Long (>~10 bp) poly(dA:dT)s	Very rigid	Vafabakhsh and Ha (2012); Marin-Gonzalez <i>et al</i> . (2020 <i>a</i> ) <sup>a</sup>	Destabilize nucleosomes	Field <i>et al.</i> (2008; Kaplan <i>et al.</i> (2009); Segal and Widom (2009)
CGI (rich in CpG steps)	Rigid	Shon <i>et al.</i> (2019);Pongor <i>et al.</i> (2017) <sup>a</sup> Marin-Gonzalez <i>et al.</i> (2019 <i>a</i> ) <sup>b</sup>	Destabilize nucleosomes	Ramirez-Carrozzi <i>et al.</i> (2009); Deaton and Bird (2011)
Poly(dG:dC)	A-like conformation	(Lu <i>et al.</i> (2000); Kulkarni and Mukherjee (2017) Marin-Gonzalez <i>et al.</i> (2019 <i>a</i> ) <sup>b</sup>	DNA:protein recognition, e.g. by zinc finger proteins	Lu <i>et al.</i> (2000); Rohs <i>et al.</i> (2010)
Poly(dA–dT)	TA-DNA conformation	Lu <i>et al.</i> (2000); Kulkarni and Mukherjee (2017) Marin-Gonzalez <i>et al.</i> (2019 <i>a</i> ) <sup>b</sup>	DNA:protein recognition, e.g. by TATA-binding protein	Lu <i>et al.</i> (2000); Rohs <i>et al.</i> (2010)
Poly(dG–dC)	B-Z transition under negative torsional stress	Lee <i>et al.</i> (2010 <i>a</i> ); Lebel <i>et al.</i> (2014); Bryant <i>et al.</i> (2012); Oberstrass <i>et al.</i> (2012, 2013) <sup>a</sup>	DNA:protein recognition via $Zlpha$ domain, e.g. in ADAR	Rich and Zhang (2003)
GC-content	Explains melting in long (kbp) DNA molecules, but not in short ones	Huguet <i>et al.</i> (2010); Gross <i>et al.</i> (2011); King <i>et al.</i> (2019) <sup>a</sup> Sutthibutpong <i>et al.</i> (2016) <sup>b</sup>	DNA unwinding at the onset of replication or transcription	Gai <i>et al.</i> (2010); Larson et al. (2011)

<sup>&</sup>lt;sup>a</sup>Single-molecule experiments.

transcription, where the negative supercoils generated by RNA polymerase would stabilize this left-handed structure (Rich and Zhang, 2003).

A number of single-molecule studies have provided a quantitative description of the thermodynamics of Z-DNA formation (Lee *et al.*, 2010*a*; Lebel *et al.*, 2014; Bryant *et al.*, 2012; Oberstrass *et al.*, 2012, 2013). In a pioneer study, Lee *et al.* (2010*a*) combined MT and smFRET to simultaneously induce and measure B–Z transitions in a DNA molecule containing a (dG–dC)<sub>11</sub> sequence, prone to assume a Z-form. These single-molecule experiments indicated that, in the presence of tension, Z-DNA formation requires only a small torsional strain, much lower than predictions based on bulk experiments. The study by Lee *et al.* thus suggested that Z-DNA might occur *in vivo* more often than expected, given that tension is likely to accumulate on DNA, e.g. during transcription.

The dynamics of the B- to Z-DNA transition have also been studied using the rotor bead tracking assay (Oberstrass *et al.*, 2012, 2013; Lebel *et al.*, 2014; Lipfert *et al.*, 2015). In this experimental setup an extreme of the DNA is bound to a magnetic bead that permits stretching and supercoiling the duplex; and a second bead is attached to the side of the DNA at an intermediate position. The position and fluctuations of the latter bead inform, respectively, on the twist and torque of the system (Bryant *et al.*, 2003; Gore *et al.*, 2006a, 2006b) (see Fig. 2e). Using different versions of this rotor bead assay, Oberstrass *et al.* obtained torque-twist measurements on DNA molecules with 22 and 50 bp-long GC repeats and observed transitions consistent with the formation of a Z-DNA structure at the poly(dG-dC) regions

(Oberstrass *et al.*, 2012, 2013). These measurements were able to recapitulate relevant features of the B–Z transition, such as its high cooperativity. Recently, rotor bead assays using a gold probe have allowed a more detailed characterization of this phenomenon, unveiling the B- to Z-DNA transition with unprecedented temporal resolution (Lebel *et al.*, 2014).

Altogether, single-molecule assays have provided a rather comprehensive description of the B- to Z-DNA transition at GC repeats under torsional stress. Future single-molecule studies should aim to explore the dynamics of this transition at other repeating sequences, and the impact of Z-DNA binding proteins on this process. Recent experimental studies are starting to provide interesting insights into these questions (Kim *et al.*, 2018*a*).

In addition to single-molecule experiments, MD simulations have also explored the B- to Z-DNA transition, shedding light on the atomistic mechanisms behind it (Lee et al., 2010b; Moradi et al., 2013; Chakraborty and Wales, 2017). Of particular relevance is the study by Moradi et al. (2013), which suggested that the B- to Z-DNA transition can proceed by means of several different mechanisms, rather than via a single pathway. In that study, MD simulations revealed a rich diversity of non-canonical DNA structures - such as an overstretched-like S-DNA, extensive base flipping or unpeeling of the two DNA strands - that act as intermediate states in the B- to Z-DNA transition. Future computational efforts should further elucidate the molecular aspects of this enigmatic process. Recent refinements of DNA force-fields which focused on improving the description of the Z-DNA structure (Zgarbová et al., 2015) will offer an invaluable tool toward achieving this goal.

<sup>&</sup>lt;sup>b</sup>Computer simulations.

#### GC-content not always explains DNA melting

Negative torsional stress regulates access to the genetic information. DNA unwinding promotes disruption of the base pairing interactions, or DNA melting, resulting in larger exposure of the nucleobases. DNA is thus unwound by specialized enzymes that locally denature the double-helix in order to read its nucleotide sequence, e.g. in transcription (Larson *et al.*, 2011), replication (Gai *et al.*, 2010), or CRISPR-mediated bacterial immunity (Szczelkun *et al.*, 2014). In addition, DNA melting often occurs as a consequence of *in vivo* torsional strains, and can result in the formation of particular structures such as kinks or bubbles that are specifically recognized by proteins (Fogg *et al.*, 2012).

At a macroscopic level (several kbp), DNA melting can be well explained from the GC-content of a given sequence (Marmur and Doty, 1962; Vologodskii and Frank-Kamenetskii, 2018). Namely, higher GC-content sequences better resist melting due to the three hydrogen bonds of the G:C base pair compared to the two bonds of the A:T base pair. This idea is supported by a recent single-molecule study that employs a novel creative assay to generate negative supercoils on DNA using a conventional OT set-up (King et al., 2019) (see Fig. 2f). In typical OT experiments, the DNA molecules are attached to the optically trapped beads by means of biotin-streptavidin bonds. King et al. found that, when torsionally constrained molecules are overstretched (at forces of ~110 pN), one of the biotin-streptavidin interactions can temporarily disrupt. When this occurs, the torsional stress accumulated on the DNA during overstretching can be partially relieved, by swiveling one DNA strand around the other (see Fig. 2f). Eventually, the broken biotin-streptavidin bond forms again, locking the DNA molecule in a negatively supercoiled state. The authors exploited this discovery to controllably unwind the DNA and induce melting events that were then detected using fluorescently labeled RPA protein (which strongly binds singlestranded DNA) as a reporter. Their results indicate that the sites of DNA melting are reasonably correlated with DNA regions of low GC-content. This finding is in agreement with experiments on DNA overstretching, where the thermodynamics of sequencedependent DNA unpeeling could be predicted on the basis of the GC-content (Gross et al., 2011).

At the microscopic level (few base pairs), however, torsion-induced DNA melting remains poorly understood. For example, studies on single-molecule unzipping of DNA hairpins suggest that, in such short DNA molecules, GC-content might not be a determinant factor of DNA melting (Huguet *et al.*, 2010; Camunas-Soler *et al.*, 2016; Vologodskii and Frank-Kamenetskii, 2018). Instead, stacking interactions (which are not directly related to the GC-content; Kilchherr *et al.*, 2016) can play an even more important role in DNA melting than base pairing interactions (Huguet *et al.*, 2010; Camunas-Soler *et al.*, 2016; Vologodskii and Frank-Kamenetskii, 2018). However, DNA melting under torsion is likely to be even more complex than DNA unzipping, and might depend on additional sequence-dependent molecular features such as DNA flexibility (Vlijm *et al.*, 2015; Shepherd *et al.*, 2020).

DNA minicircles offer an attractive platform for studying supercoiling-induced DNA denaturation at the molecular level. DNA minicircles consist of small (few hundreds of base pairs) closed DNA molecules that are subjected to high bending and torsional stress. Computer simulations have revealed a rich structural diversity in DNA minicircles, including kinks, base flipping, and denaturation events (Lankaš *et al.*, 2006; Irobalieva *et al.*, 2015; Sutthibutpong *et al.*, 2016; Pyne *et al.*, 2020). In a recent study,

Sutthibutpong *et al.* combined atomistic MD, coarse-grained simulations and statistical mechanics calculations to study sequence-dependent melting in DNA minicircles (Sutthibutpong *et al.*, 2016). Coarse-grained techniques identified AT-rich regions to be more prone to undergo melting. Nevertheless, detailed atomistic MD simulations indicated that both breathing and melting events were more frequent in flexible pyrimidine-purine dinucleotides such as TpA or CpA (see Section 'TpA dinucleotides are highly flexible').

The complex interplay between base pairing, base stacking interactions and DNA flexibility in DNA unwinding remains an open question. We believe that the combination of MD simulations and high-resolution AFM imaging on DNA minicircles can be an extremely useful approach toward gaining such molecular understanding of sequence-dependent DNA melting. Ongoing research in this direction is already yielding very promising results (Pyne *et al.*, 2020).

#### **Chemical modifications and DNA mechanics**

In the previous section, we have discussed how the nucleotide sequence impacts the flexibility of canonical Watson–Crick base paired DNA molecules. Inside the cell, however, the DNA often presents changes in its chemical structure, including modified bases, mismatches, or abasic sites. These chemical modifications can occur in the form of epigenetic marks via controlled action of the cellular machinery; or in the form of DNA lesions that jeopardize the normal functioning of the cell.

In this section, we review recent findings on the effects of epigenetics and DNA mismatches on DNA mechanics. When put together, these findings reveal that such chemical modifications have a more complex effect on DNA flexibility than previously thought. Methylation marks play a versatile role in DNA flexibility attending to the sequence context on which they occur and the mechanical deformation considered. On the other hand, DNA mismatches usually enhance DNA flexibility, but confer the DNA with exotic mechanical properties that remain to be deciphered. The emerging picture is that, in both cases, DNA flexibility might act as a potent signal for downstream events. Changes in DNA flexibility resulting from epigenetic marks might affect the compaction state of chromatin; whereas DNA defects might act as flexibility antennas for the recruitment of the repair machinery.

### The impact of cytosine methylation on DNA physical properties

Epigenetic DNA marks play a myriad of roles in development (Smith and Meissner, 2013), aging (Unnikrishnan *et al.*, 2019), and the onset and progression of cancer (Esteller, 2008). The most common of these marks consists of the addition of a methyl group to the C5 carbon of cytosine, which typically occurs at CpG steps. The resulting methylcytosine is usually linked to gene silencing, but the molecular mechanism responsible for this is not completely understood (Cortini *et al.*, 2016). Notably, besides affecting the interaction of DNA with several proteins, cytosine methylation is suspected to orchestrate rearrangements of nucleosomes via changes in the mechanical properties of DNA (Dantas Machado, 2014; Cortini *et al.*, 2016).

Cytosine methylation is generally considered to reduce DNA flexibility Cortini *et al.* (2016), as supported by recent single-molecule cyclization experiments. Ngo *et al.* (2016) reported that upon methylation of one to eight (four in each strand) cytosines of an arbitrary DNA sequence, the looping time increases,

Table 3. Effect of cytosine methylation on DNA physical properties

DNA physical property	Effect of cytosine methylation	References	
DNA flexibility	Generally stiffens the DNA	Ngo <i>et al.</i> (2016) <sup>a</sup> Pérez <i>et al.</i> (2012); Portella <i>et al.</i> (2013) <sup>b</sup>	
	Could increase the flexibility of CGIs	Pongor <i>et al.</i> (2017); Shon <i>et al.</i> (2019) <sup>a</sup> Marin-Gonzalez <i>et al.</i> (2019 <i>a</i> ) <sup>b</sup>	
DNA melting	Versatile effect on DNA strand separation depending on the sequence	Severin et al. (2011) <sup>a</sup>	
DNA structure	Affects DNA structure in a complex, sequence-dependent manner	Pérez et al. (2012); Rao et al. (2018) <sup>b</sup>	
B-Z DNA transition	Facilitates B–Z transition under torsional stress	Lee et al. (2010a) <sup>a</sup>	
DNA condensation	Promotes DNA condensation	Yoo et al. (2016); Kang et al. (2018); Yang et al. (2020) <sup>a</sup>	

<sup>&</sup>lt;sup>a</sup>Single-molecule experiments.

which reflects a decrease in flexibility. This methylation-induced stiffening was well reproduced in supporting MD simulations, showing that methylcytosine dampened local fluctuations in the duplex. In addition, Ngo *et al.* (2016) combined OT and smFRET to show that methylation of the 601 Widom sequence resulted in mechanical destabilization of nucleosomes.

The idea that methylation stiffens the DNA is generally well accepted (Pérez et al., 2012; Portella et al., 2013; Cortini et al., 2016). Yet, simulation (Marin-Gonzalez et al., 2019a; Liebl and Zacharias, 2018) and experimental studies (Pongor et al., 2017; Shon et al., 2019) have indicated that, in some cases, methylation can soften the DNA. In particular, methylation could enhance the flexibility of CGIs (discussed in Section 'CpG islands are rigid DNA regions depleted of nucleosomes') (Marin-Gonzalez et al., 2019a; Shon et al., 2019; Pongor et al., 2017). Dense methylation (hypermethylation) of CGIs is a potent gene silencing mechanism, and a hallmark of many cancer types (Esteller, 2008; Deaton and Bird, 2011). A possible role of DNA flexibility has been suggested in this process (Marin-Gonzalez et al., 2019a; Shon et al., 2019; Pongor et al., 2017). In fact, CGIs appear rigid, a feature that might explain the nucleosome-binding deficient character of these sequences in vivo (see Section 'CpG islands are rigid DNA regions depleted of nucleosomes') (Pongor et al., 2017; Shon et al., 2019). However, upon hypermethylation, CGIs have been reported to become more flexible, with values of persistence length and stretch modulus closer to those of standard DNA (Pongor et al., 2017; Marin-Gonzalez, 2019a; Shon et al., 2019). This softening upon hypermethylation could increase the affinity of CGI to form nucleosomes, which may occlude the DNA to the transcription machinery, resulting in gene inactivation. Therefore, despite methylation is usually thought to stiffen the DNA, the full story might be more complex. Given the key biological function of CGIs, it would be of utmost interest to determine whether these sequences constitute an exception to the aforementioned rule.

Moreover, it is worth mentioning that cytosine methylation can have a substantial impact on DNA physical properties other than flexibility (see Table 3). For example, cytosine methylation affects force-induced DNA strand separation, as probed by single-molecule force spectroscopy measurements (Severin *et al.*, 2011). In this case, cytosine methylation plays a versatile role in DNA mechanics depending on the sequence-context: one methylated cytosine destabilized the duplex but three methylated cytosines resulted in larger mechanical stability. This versatility also appears to be present in sequence-dependent effects of cytosine

methylation on DNA structure, as predicted from computer simulations (Pérez et al., 2012; Dantas Machado et al., 2014; Rao et al., 2018). For example, in some sequence contexts, cytosine methylation would greatly affect the structure of the DNA, whereas in other sequences the structural effect of this epigenetic mark would be minimal (Rao et al., 2018). In the extreme case of poly(dG–dC) sequences, single-molecule experiments showed that cytosine methylation can even facilitate the formation of non-canonical Z-DNA structures in the presence of torsional stress (Lee et al., 2010a).

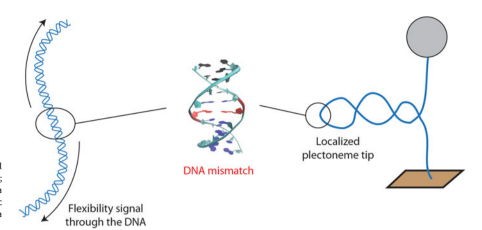
Finally, cytosine methylation has also been studied in the context of DNA condensation by polycations, the phenomenon by which highly positively charged ions mediate DNA compaction by stabilizing inter-helical interactions (Bloomfield, 1997). smFRET, MT, and MD simulations coincide that cytosine methylation enhances DNA condensation (Yoo *et al.*, 2016; Kang *et al.*, 2018; Yang *et al.*, 2020). Importantly, this effect appears consistent when using different condensing polycations (spermidine<sup>3+</sup>, CoHex<sup>3+</sup>, spermine<sup>4+</sup>, and polylysine<sup>6+</sup>), and DNA sequences.

Altogether, some general conclusions can be derived concerning the effect of cytosine methylation on DNA physical properties. Methylation usually reduces DNA flexibility and facilitates DNA condensation. Nevertheless, in many instances the role of cytosine methylation in DNA mechanics appears strongly dependent on the sequence context on which this epigenetic mark occurs. Therefore, methylation would add an additional layer of complexity to the sequence-dependent biophysical properties of DNA, instead of exerting a systematic, generalizable effect. Understanding cytosine methylation in the context of DNA mechanics would then require investigation on a case-by-case basis.

# The effect of DNA mismatches on DNA mechanics

DNA defects, such as mismatches, can induce strongly distorted DNA conformations. Inside the cell, such DNA defects are localized and repaired by the coordinated action of DNA repair proteins with an impressive efficiency. However, the mechanisms by which some of these proteins rapidly identify single DNA imperfections in a huge genome are not completely understood. Remarkably, several DNA repair proteins spanning diverse repair pathways have been reported to interact with a sharply bent DNA (Roberts and Cheng, 1998; Natrajan et al., 2003; Qi et al., 2009; Chakraborty et al., 2017; Craggs et al., 2019; Paul et al., 2019). A natural question is how (and to which extent) the deformability of damaged DNA impacts these repair processes and, in particular, the recognition

<sup>&</sup>lt;sup>b</sup>Computer simulations.



**Fig. 3.** Effects of DNA mismatches on DNA mechanical properties. A single mismatched base pair (center; PDB: 10NM; Sanchez *et al.*, 2003) can propagate a mechanical signal through the DNA via an allosteric mechanism (left); and can pinpoint the position of a plectoneme tip (right).

of the defect (Krokan and Bjørås, 2013; Marteijn *et al.*, 2014; Kunkel and Erie, 2015). For example, do DNA defects spontaneously bend the duplex to recruit repair proteins or do these defects facilitate DNA bending once the protein is attached?

The answer to this question will greatly vary attending to the particular DNA defect and repair proteins involved. Here, we will discuss the effect of mismatches, or 'pairing' between noncomplementary bases, which have recently received special attention in the context of DNA flexibility (Rossetti et al., 2015; Jeong and Kim, 2019). Mismatches arise during DNA replication or DNA exposure to damaging agents, among other processes. Although these defects are generally associated with increased DNA bendability (see e.g. Vafabakhsh and Ha, 2012; Satange et al., 2018), this might not always be the case, as suggested by recent studies (Rossetti et al., 2015; Jeong and Kim, 2019). Using MD simulations and nuclear magnetic resonance experiments, Orozco, Gonzalez, and coworkers performed a systematic study of the effect of mismatches on DNA flexibility (Rossetti et al., 2015). They observed that, in general, the magnitude of bending fluctuations was similar in mismatched duplexes and in control ones with correct base pairing, suggesting that mismatches do not enhance DNA bending flexibility. Nevertheless, DNA molecules containing mismatches were flexible at a local level, with frequent breathing events and distortions of the DNA grooves. The authors warn against simplistic interpretations and indicate that the aforementioned effects are largely dependent on the kind of mismatch considered. Nevertheless, their results suggest that, even though mismatches might not cause spontaneous DNA bending, local DNA distortions due to mismatches may aid proteins at bending the duplex.

Interestingly, Orozco, Gonzalez, and coworkers also observed that DNA distortions induced by defects not only occurred at the position of the mismatch, but also at remote locations of ~4 bp from the mismatch. This finding implies that mismatches might take advantage of a phenomenon known as DNA allostery (Kim et al., 2013) to propagate distortions along the duplex. An independent, experimental observation of such long-range effects of mismatches has been recently reported in the context of single-molecule DNA cyclization (Jeong and Kim, 2019). Jeong et al. reported that, paradoxically, mismatches destabilize DNA loops, even though these defects favor the sharply bent DNA conformations that are required for loop formation. Such destabilization of loops was attributed to allostery effects, similar to the ones described in Rossetti et al. (2015), but reaching even longer distances of ~50 bp.

Altogether, the appealing hypothesis that mismatches propagate deformations along the DNA and, thus, act as 'flexibility antennas' for DNA repair proteins will require further research (see Fig. 3). Alternative mechanisms for the cellular machinery to identify mismatches should also be explored, such as the recently reported finding that mismatches can localize plectoneme tips (Ganji et al., 2016; Dittmore et al., 2017) (see Fig. 3). Note that these two mechanisms, DNA allostery and plectoneme localization, are not necessarily mutually exclusive. Finally, it remains to be explored whether these and other new phenomena arise from other kinds of DNA defects, such as oxo-guanines, photoproducts, or interstrand crosslinks.

# Mechanical properties of dsRNA: unexpected differences with dsDNA

Single-molecule studies have also addressed the mechanical properties of dsRNA (Abels et al., 2005; Herrero-Galán et al., 2013; Lipfert et al., 2014; Fu et al., 2020). RNA is known to be predominantly single-stranded; however, dsRNA helices are also commonly found inside the cell and exert a myriad of biological functions. For example, dsRNA is the carrier of genetic information in some viruses and dsRNA helices are key elements of tertiary RNA structures, as formed e.g. by ribosomal RNA or t-RNAs (Carter et al., 2000; Nissen et al., 2001; Schimmel, 2018). The mechanical properties of dsRNA might play a role in some of these biological systems. For example, the dsRNA bending stiffness is expected to affect the energetics of genome compaction in dsRNA viruses (Zhang et al., 2015; Buzón et al., 2020). Moreover, the sequence-dependent structure and flexibility of dsRNA helices have been proposed to impact dsRNA:protein interactions or the folding of tertiary RNA structures (Perona and Hou, 2007; Yesselman et al., 2019). Finally, a quantitative understanding of dsRNA mechanics could aid the future design of RNA nanostructures (Guo, 2010).

Initial studies on the dsRNA persistence length suggested that the mechanical properties of this molecule were qualitatively similar to dsDNA (Hagerman, 1997; Abels *et al.*, 2005). In a seminal study that combined MT and AFM, Abels *et al.* (2005) obtained a value of  $P_{\rm RNA}$  ~62 nm, which is only 20% larger than  $P_{\rm DNA}$ . This slightly larger rigidity of dsRNA can be easily rationalized on the basis of the thicker and more compact structure of the A-RNA helix compared to the B-DNA helix. In parallel, MD simulations indicated some differences in the dynamics of dsDNA and dsRNA at the microscopic level (Noy *et al.*, 2004). Namely,

Table 4. Differences in the mechanical properties of dsDNA and dsRNA

Feature	dsDNA	References	dsRNA	References
Microscopic flexibility	Complex deformability, including e.g. allostery, polymorphic behavior	Kim <i>et al.</i> (2013) <sup>a</sup> Dans <i>et al.</i> (2014); Pasi <i>et al.</i> (2014) <sup>b</sup>	Simple deformability patterns	Noy <i>et al.</i> (2004); Beššeová <i>et al.</i> (2012) <sup>b</sup> Pérez <i>et al.</i> (2004)
Stretching stiffness	High (S ∼ 1200 pN)	Smith <i>et al.</i> (1996); Wang <i>et al.</i> (1997) <sup>a</sup>	Low (S ~ 400 pN)	Herrero-Galán <i>et al.</i> (2013); Lipfert <i>et al.</i> (2014) <sup>a</sup>
Twist-stretch coupling	Negative: overwinds when stretched	Gore <i>et al.</i> (2006 <i>a</i> ); Lionnet <i>et al.</i> (2006) <sup>a</sup>	Positive: unwinds when stretched	Lipfert <i>et al.</i> (2014) <sup>a</sup> Liebl <i>et al.</i> (2015); Bao <i>et al.</i> (2017); Marin-Gonzalez <i>et al.</i> (2017) <sup>b</sup>
Plectoneme formation	Fast (ms) dynamics	Crut et al. (2007); Forth et al. (2008); Brutzer et al. (2010); van Loenhout et al. (2012) <sup>a</sup> Ott et al. (2020) <sup>b</sup>	Slow (s) dynamics	Lipfert <i>et al.</i> (2014) <sup>a</sup> Ott <i>et al.</i> (2020) <sup>b</sup>
Effects of multivalent (≥2) cations	Increase bending flexibility; cause dsDNA condensation	Baumann <i>et al.</i> (1997); Wenner <i>et al.</i> (2002) <sup>a</sup>	Can decrease bending flexibility; dsRNA resists condensation	Fu et al. (2020) <sup>a</sup> Tolokh et al. (2014); Drozdetski et al. (2016) <sup>b</sup> Li et al. (2011); Katz et al. (2017)
Intrinsic bending	Occurs at A-tracts	Rivetti <i>et al.</i> (1998); Moreno-Herrero <i>et al.</i> (2006) <sup>a</sup>	Occurs at AU-tracts	Marin-Gonzalez <i>et al.</i> (2020 <i>b</i> ) <sup>a</sup>
Rigid motifs	A-tracts, CGI, ApT step	Shon et al. (2019); Marin-Gonzalez et al. (2020a) <sup>a</sup> Pasi et al. (2014) <sup>b</sup>	Homopurine regions: poly(rG:rC), poly(rA:rU) and poly(rA-rG)	Marin-Gonzalez <i>et al.</i> (2019 <i>b</i> ) <sup>b</sup>

a Single-molecule experiments.

dsRNA showed simple deformability patterns that could be well described by few essential motions, whereas dsDNA was able to explore a wider range of conformations. Nonetheless, it was not clear whether these microscopic motions would translate into changes in the global flexibility of the molecules that could be measured in single-molecule experiments.

In the last few years, a number of single-molecule and simulation studies have revealed that the mechanical properties of dsDNA and dsRNA are more different than previously thought (Table 4). The emerging picture is that previous findings on dsDNA flexibility do not necessarily apply to dsRNA. Namely, the latter has its own mechanical identity. In the following, we discuss some of those studies. First, we discuss differences in the flexibility of DNA and RNA duplexes of arbitrary sequence under standard ionic conditions (Sections 'Stretching flexibility of dsDNA and dsRNA' and 'The opposite twist-stretch coupling of dsDNA and dsRNA'). We then comment on the opposite effect of certain multivalent ions on the mechanics of DNA and RNA duplexes (Section 'The different dynamics of plectoneme formation'). Finally, we discuss sequence effects on the structure (Section 'Opposite effects of complex ions on the mechanics of dsDNA and dsRNA') and the flexibility (Section 'Sequence determinants of intrinsic bending') of dsDNA and dsRNA.

### Stretching flexibility of dsDNA and dsRNA

The first single-molecule study that reported qualitative differences in the mechanics of dsDNA and dsRNA was Herrero-Galán *et al.* (2013). The authors first performed AFM and MT experiments on dsDNA and dsRNA and found similar values of the persistence length to the ones previously published in Abels *et al.* (2005). Nonetheless, OT stretching experiments unveiled an important difference between the two nucleic acids:

dsRNA was very soft to stretching deformations, around threefold more flexible compared to dsDNA (see Fig. 4a).

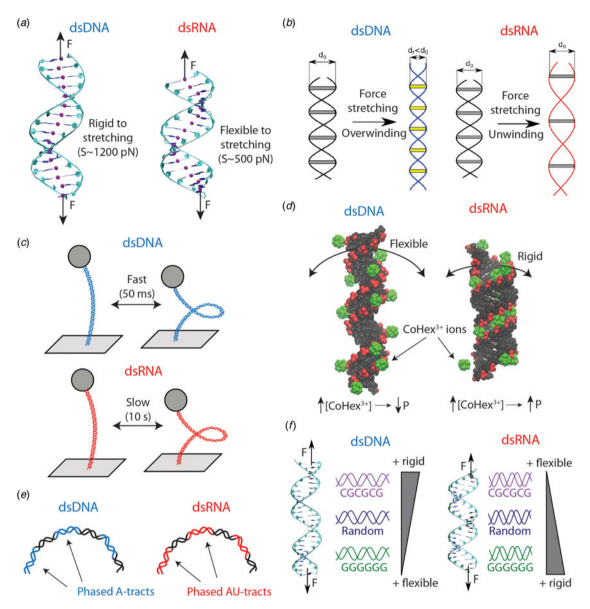
This different stretching flexibility of dsDNA and dsRNA has been reproduced in several experimental and computational studies (Chou *et al.*, 2014; Lipfert *et al.*, 2014; Bao *et al.*, 2017; Marin-Gonzalez *et al.*, 2017; Fu *et al.*, 2020). At the molecular level, the softer stretching response of dsRNA is commonly associated with the more open structure of the A-RNA helix compared to B-DNA (Chou *et al.*, 2014; Bao *et al.*, 2017; Marin-Gonzalez *et al.*, 2017). Note that this idea is similar to the one presented in Section 'Poly(dA–dT) and poly(dG:dC) adopt A-like DNA conformations', according to which DNA sequences with larger A-philicity would possess a softer stretching rigidity.

### The opposite twist-stretch coupling of dsDNA and dsRNA

Measurements of the torsional response of dsDNA and dsRNA, as performed by Lipfert *et al.* revealed more striking differences in the mechanics of these duplexes (Lipfert *et al.*, 2014). Note that when a double-helix is straightened, two parallel strands reach a longer extension than they have when coiled around each other in a helical conformation. According to this picture, stretching a double-helix should lead to unwinding. However, it was known that dsDNA had the surprising ability of overwinding when stretched (Gore *et al.*, 2006a; Lionnet *et al.*, 2006) (see Fig. 4b). Using MT, Lipfert *et al.* measured changes in the dsRNA extension as a function of twist, and they found the completely opposite behavior: contrary to dsDNA, dsRNA unwinds upon stretching Lipfert *et al.* (2014).

The opposite twist-stretch coupling of dsDNA and dsRNA resulted in a great challenge for computational models (see e.g. the review by Kriegel *et al.*, 2017*a*). Several coarse-grain models,

<sup>&</sup>lt;sup>b</sup>Computer simulations.



**Fig. 4.** Differences in the mechanical properties of dsDNA and dsRNA (see also Table 4). (a) dsRNA is around three times softer to stretching deformations than dsDNA. This difference can be explained on the basis of the more open structure of dsRNA, as evidenced from the base pair center chains of the duplexes (purple beads). (b) dsDNA overwinds when stretched, whereas dsRNA unwinds. The peculiar behavior of dsDNA can be rationalized from the shrinking of its radius upon elongation. On the contrary, the dsRNA radius is unchanged when the molecule is stretched. Adapted from Marin-Gonzalez et al. (2017). (c) When the duplexes are supercoiled at the threshold for plectoneme formation, dsDNA displays fast buckling dynamics (ms), whereas dsRNA undergoes slow (s) buckling transitions. This can be partly attributed to the larger persistence length of dsRNA; however, the precise mechanisms for this difference remain incompletely understood. (d) Increasing ionic concentrations result in larger dsDNA bending flexibility and a decrease in persistence length (P). However, some multivalent ions, such as CoHex<sup>3+</sup> or spermine<sup>4+</sup>, can have the opposite effect on dsRNA and stiffen this duplex increasing P. This phenomenon can be understood from the shape of the grooves of the duplexes. For dsRNA, the ions can bind inside the major groove, but in the dsDNA case the ions bind mostly externally. Adapted from Drozdetski et al. (2016). (e) Sequence-induced bending in dsDNA and dsRNA occur, respectively, via A-tract and AU-tract (alternating adenines and uracit) sequences. (f) The nucleotide sequence impacts in different ways the global mechanical properties of dsDNA and dsRNA. Poly(dC-dG) motifs are rigid in dsDNA, but flexible in the dsRNA case; whereas poly(dG:dC) ones are flexible in dsDNA but rigid in dsRNA. Adapted from Marin-Gonzalez et al. (2019b).

with varying levels of detail, unsuccessfully attempted to reproduce the opposite twist-stretch coupling of dsDNA and dsRNA (Kriegel *et al.*, 2017*a*). For example, a base pair level model built from crystal structures of dsDNA and dsRNA concluded that both molecules would unwind when stretched (Chou *et al.*, 2014). The oxDNA/oxRNA models, which were originally designed to reproduce the thermodynamics of duplex formation (base pairing and stacking interactions), yielded the correct twist-stretch coupling for dsRNA, but failed for dsDNA (Matek *et al.*, 2015*a*, 2015*b*).

The shortfall of coarse-grain simulations suggested that, in order to reproduce the opposite twist-stretch coupling of dsDNA and dsRNA, fine details of the dynamics of the duplexes might need to be considered. Atomistic MD simulations met this demanding condition, as was elegantly demonstrated by Liebl *et al.* (2015). In that study, the authors performed atomistic MD simulations of a dsDNA and a dsRNA molecule of analogous sequence. An analysis of the MD trajectories revealed that the correlation between elongation and twisting motions were of opposite sign for both duplexes, in semi-quantitative agreement with

the experiments. The authors further corroborated this finding by exerting a controlled torque on the nucleic acids and measuring the associated change in extension. Consistently, they found that a positive torque (overwinding) caused lengthening of dsDNA, but shortening of dsRNA; and unwinding resulted in the opposite behavior (shortening of dsDNA and lengthening of dsRNA). Liebl *et al.* elaborated a technical explanation for the opposite twist–stretch coupling of dsDNA and dsRNA based on a wealth of structural parameters of the duplexes. Another study also addressed the twist–stretch coupling of DNA and RNA duplexes and attributed their difference to the slide and inclination motions of the base pairs (Bao *et al.*, 2017).

Recently, an alternative mechanism for the twist-stretch coupling has been proposed, by directly computing the radii of the duplexes from MD simulations (Marin-Gonzalez et al., 2017) (see Fig. 4b). DNA and RNA duplexes were stretched using a novel constant force protocol for MD simulations Liebl et al. (2015). It was found that, as the force increased, dsDNA overwound, while dsRNA unwound, in agreement with the aforementioned simulations and experiments. Importantly, because the molecules were left to equilibrate under stress, it was possible to measure how their average structure changed at each value of the external force. It was thus observed that the dsDNA radius decreases with force, whereas the dsRNA radius remains approximately constant. The force-induced shrinking of the DNA radius brings together the strands of the duplex, allowing it to overwind upon stretching (see Fig. 4b). Since this capability of reducing the radius was absent in dsRNA, this molecule could only unwind when stretched. The larger flexibility of the DNA sugar was suggested to be responsible for promoting the reduction of the dsDNA radius.

#### The different dynamics of plectoneme formation

The extensive investigation of Lipfert *et al.* on the twisting dynamics of dsDNA and dsRNA yielded yet another surprising result (Lipfert *et al.*, (2014). When these double-stranded nucleic acids are twisted above a certain threshold, they undergo a buckling transition and start forming plectonemes (see Fig. 4c). Lipfert *et al.* (2014) studied the dynamics of dsDNA and dsRNA when the twist of duplexes is constrained at the threshold value for the buckling transition. At that point, pre-buckling and post-buckling states are equally populated and the nucleic acids continuously transition between states. The authors found that such buckling dynamics were much slower in dsRNA compared to dsDNA, with a difference in dwell times of at least two orders of magnitude.

This issue has been recently addressed in an insightful simulation study. Ott et al. (2020) modeled a dsDNA and a dsRNA molecule using Brownian dynamics simulations and a simple WLC model that contemplates bending and twisting deformations. They found that the difference in persistence length of dsDNA and dsRNA, although small, can result in an order of magnitude change in the buckling dynamics. This approach thus partially explains the experimental findings. However, further theoretical efforts are needed to arrive at a quantitative description of the buckling dynamics of these nucleic acids (see the recent perspective by Lankaš, 2020). We believe that such description should take into account the cross-talk between twisting and bending deformations, that is, the fact that twisting can facilitate bending. This so-called twist-stretch coupling parameter has recently been estimated for dsDNA (Nomidis et al., 2017), but a solid measurement is still lacking in the dsRNA case.

# Opposite effects of complex ions on the mechanics of dsDNA and dsRNA

The effect of salt conditions on dsDNA mechanics has been extensively studied for several decades (Harrington, 1978; Ha and Thirumalai, 2003). However, recent studies are providing very interesting insights into this topic (Brunet et al., 2015b; Kriegel et al., 2017b; Guilbaud et al., 2019). Nonetheless, in contrast to DNA, much less is known about the effect of salt on dsRNA flexibility. Initial single-molecule experiments reported a qualitatively similar decrease in  $P_{\text{RNA}}$  and  $P_{\text{DNA}}$  with increasing monovalent salt (Herrero-Galán et al., 2013; Lipfert et al., 2014). A priori, these findings could suggest a similar dependence of DNA and dsRNA flexibility on salt conditions. However, further single-molecule experiments and MD simulations have reported unexpected results when multivalent ions are present (Drozdetski et al., 2016; Fu et al., 2020) (see Fig. 4d). Motivated by the different condensation properties of dsDNA and dsRNA (Li et al., 2011; Tolokh et al., 2014), Onufriev and coworkers performed atomistic MD simulations to explore the role of the ion  $CoHex^{3+}$  in  $P_{RNA}$  and  $P_{DNA}$  (Drozdetski et al., 2016). The simulations showed that  $P_{\rm DNA}$  decreases with increasing CoHex<sup>3+</sup> concentration, in accordance with naïve intuition and with previous experiments (Baumann *et al.*, 1997). However, unexpectedly, increasing amounts of CoHex<sup>3+</sup> resulted in an increase of  $P_{\rm RNA}$ . This stiffening of dsRNA was consistent when alternative multivalent ions were used, such as spermine<sup>4+</sup> or a hypothetical Na<sup>3+</sup>. Recently, the opposite effect of CoHex<sup>3+</sup> on the flexibility of dsDNA and dsRNA has been experimentally demonstrated in a comprehensive study that combines both MT experiments and MD simulations (Fu et al., 2020). Altogether, these studies illustrate the limitations of polyelectrolyte models to describe the interaction of dsDNA and dsRNA with complex, multivalent ions such as CoHex3+. In these cases, molecular details of the double helices, such as shape of their grooves, must be taken into account in order to explain the effect of these ions on the elasticity of the duplexes.

## Sequence determinants of intrinsic bending

The phenomenon of sequence-induced DNA bending by A-tracts raises the question of whether a similar effect can occur in dsRNA. This question is more complicated than one would expect. For example, the few structures of naked RNA duplexes available from X-ray crystallography experiments often present artifacts (Šponer et al., 2018). This issue constitutes an important drawback for experimental validation of dsRNA MD simulations via X-ray crystallography data (Šponer et al., 2018). Testing MD against single-molecule experiments can therefore be a promising route for exploring sequence-dependent dsRNA features. By combining MD simulations and AFM experiments, it has been recently shown that alternating adenines and uracils - or AU-tracts - bend the RNA duplex (Marin-Gonzalez et al., 2020b) (see Fig. 4e). AU-tracts were long known to possess a peculiar structure at a local base pair level (Dock-Bregeon et al., 1989), but their effect on global dsRNA features had remained largely unexplored. The MD simulations from Marin-Gonzalez et al. (2020b) revealed that dsRNA molecules with AU-tracts were systematically more bent than sequences lacking this motif. Motivated by this finding, dsRNA molecules were fabricated with repetitions of AU-tracts spaced by 11 bp and these molecules were imaged using an AFM. The images revealed a bent character in these AU-tract molecules, which was quantified

by a value of the persistence length as low as  $\sim$ 30 nm, about half the standard value of dsRNA ( $\sim$ 60 nm). These findings argue against simplistic conceptions of dsRNA as a regular helix. Further investigation will be required to elucidate the molecular mechanisms and the possible biological consequences of this novel phenomenon.

#### Sequence-dependent mechanical properties

The aforementioned sequence-dependent dsRNA bending demonstrates that the nucleotide sequence can affect in a very distinct manner the structure of dsDNA and dsRNA. Namely, A-tracts induce an intrinsic bend in dsDNA, but not in dsRNA; and AU-tracts have the opposite effect: they bend dsRNA, but not dsDNA (see Fig. 4e). Therefore, it would be natural to expect that the nucleotide sequence would also impact the relative flexibility of dsDNA and dsRNA in different ways. A recent simulation study suggests that this might be the case (Marin-Gonzalez et al., 2019b). Using MD simulations, Marin-Gonzalez et al. measured the mechanical response of several dsRNA sequences and compared the results with dsDNA analogs of the same sequence. dsRNA molecules were always more flexible to stretching deformations than dsDNA ones, regardless of the sequence. However, the effect of the nucleotide sequence on the stretching flexibility was completely different in the two nucleic acids. For example, the poly(rG:rC) RNA duplex is relatively rigid to stretching, but a poly(dG:dC) DNA duplex is highly flexible when compared with an arbitrary sequence (see Fig. 4f). On the contrary, the poly(rG-rC) dsRNA molecule is flexible to stretching, but the poly(dG-dC) dsDNA is rigid. The molecular mechanisms behind this difference are to be examined. Furthermore, the idea that sequence effects on dsDNA and dsRNA flexibility can be substantially different still awaits experimental validation.

# **Conclusions and future perspectives**

In this review, we have revisited recent single-molecule experiments and MD simulations studies on DNA mechanical properties. These studies are collectively providing a comprehensive, molecular description of DNA mechanics by assessing how microscopic chemical features of the double-helix impact its physical properties. Importantly, as we deepen into such molecular characterization, a rich sequence-dependent conformational variability of the double-helix emerges, which is often overlooked by classical polymer approaches. For example, both experiments and simulations suggest that, besides few exceptions, the DNA persistence length is relatively insensitive to the nucleotide sequence (Geggier and Vologodskii, 2010; Mitchell et al., 2017). Even distorted DNA duplexes containing a mismatch appear to have a standard bending flexibility (Rossetti et al., 2015). On the contrary, local, strong deformations such as kinks, bubbles or Z-DNA structures are strongly sequence-dependent (Rich and Zhang, 2003; Olson and Zhurkin, 2011; Fogg et al., 2012). It is thus tempting to conclude that sequence effects on DNA mechanics are amplified under large mechanical stress, that is, when the duplex is forced to adopt a structure that substantially differs from the canonical B-DNA helix.

Furthermore, the studies here reviewed are enabling a better understanding of the mechanical impact of cytosine methylation. Several lines of evidence indicate that cytosine methylation affects many DNA physical properties in a highly complex manner that often depends on the particular sequence context (see Table 3 and reference Cortini *et al.*, 2016). The emerging view is that, rather than exerting a systematic effect on DNA mechanics, methylated cytosine acts as a 'fifth nucleotide' that expands the 'physical code' imprinted in a given DNA sequence. Similar considerations might apply to DNA mismatches, whose effects on DNA mechanics appear to be strongly dependent on the specific kind of mismatch considered.

Finally, we have revisited recent findings on dsRNA mechanics, focusing on those studies that reveal unexpected observations in the mechanical properties of dsRNA when compared to its DNA counterpart. From a molecular perspective, these studies raise an interesting consideration. Namely, that the presence of an extra –OH group in the sugar and the substitution of thymine by uracil have enormous implications in the physical properties of nucleic acids.

Despite the substantial progress made in the last few years, a number of important aspects of DNA mechanics will require further study. In the following, we comment on some of those aspects and we briefly discuss how an improved characterization of DNA physical properties can potentially impact other areas of biology, biophysics, and nanotechnology.

# Sequence-dependent DNA mechanics beyond the elastic regime

In the elastic regime, the sequence-dependent DNA deformability can be accurately described from the analysis of structural databases (Olson *et al.*, 1998), or from extensive atomistic MD simulations (Pasi *et al.*, 2014; Walther *et al.*, 2020). However, as mentioned above, the nucleotide sequence largely affects the energetics of highly distorted DNA conformations beyond the elastic regime, such as kinks or bubbles. We have outlined in a qualitative manner the main sequence determinants of such sharp DNA deformations (see Table 2). However, a quantitative characterization of DNA dynamics beyond the elastic regime is a challenge that will need to be addressed in future studies. In this respect, novel high-throughput assays based on next-generation sequencing, such as the recently developed loop-seq assay (Basu *et al.*, 2020, 2021), offer an attractive platform to interrogate such sequence-dependent DNA mechanical properties.

It is important to note that DNA kinks, bubbles, and other distorted conformations appear in a number of structures of DNA: protein complexes, including the nucleosome core (Dickerson et al., 1998; Olson and Zhurkin, 2011). Therefore, a quantitative understanding of the energetics of these conformations might shed light on several biological questions. In the paradigmatic case of nucleosome stability, DNA kinks might be more determinant than the smooth bending flexibility (Zhurkin and Olson, 2013). Together with other sequence-dependent features, such as DNA shape (Rohs et al., 2009), DNA kinks might greatly contribute to the wide sequence-dependent variability of nucleosome affinity reported in in vitro experiments (Onufriev and Schiessel, 2019).

The future challenges regarding the sequence-dependent DNA mechanics beyond the elastic regime can be summarized as:

- To quantitatively characterize the energetics of formation of highly distorted DNA conformations, most notably kinks and local denaturation events.
- To devise novel assays for systematically evaluating the sequence-dependent energetics of formation of highly distorted DNA conformations.

- To evaluate under which circumstances the *in vivo* mechanical stress, as found e.g. in the nucleosome core, is sufficient for stabilizing the formation of those highly distorted DNA structures.
- To incorporate non-elastic effects into current coarse-grain models of DNA to improve theoretical descriptions of DNA mechanics.

# Toward a better characterization of cytosine methylation and DNA mismatches in the context of DNA mechanics

As discussed in the text, cytosine methylation is usually correlated with an increased rigidity of the DNA. However, there is evidence pointing toward possible exceptions to this rule, most notably, certain sequences with high density of CpG steps (or CGIs). These (and potentially other kinds of) DNA sequences where methylation results in DNA softening clearly deserve more attention, both for experiments and simulations. The latter should aim to elucidate the molecular mechanisms behind methylation-induced changes in DNA flexibility, which still remain obscure.

In addition, future efforts are needed to better characterize the effect of cytosine methylation on DNA condensation by polycations. This question is particularly timely, in light of recent experiments and simulations that have suggested a liquid–liquid phase separation mechanism for DNA condensation (Kang *et al.*, 2018; Shakya and King, 2018). It is thus conceivable that cytosine methylation might enhance the phase separation behavior of DNA.

Understanding how DNA mismatches alter the DNA physical properties is another exciting task for the near future. For example, the hypothesis that mismatches promote allosteric effects in the DNA would benefit from more extensive experimental support at the single-molecule level. Another interesting idea to test experimentally would be whether different kinds of mismatches possess different mechanical footprints. Testing transductions (non-complementary pyrimidine:purine base pairing) against transversions (pyrimidine:pyrimidine or purine:purine base pairing) would be a promising starting point. Because the geometries of these kinds of mismatches are very different, it is expected they will possess different mechanical properties amenable to experimental observation.

Altogether, we devise the following challenges for future studies on cytosine methylation and DNA mismatches in the context of DNA mechanics:

- To further test experimentally the effect of cytosine methylation on the dynamics of CGIs and explore other potential exceptions to the rule that methylation reduces DNA flexibility.
- To further test the effect of cytosine methylation on the condensation and, potentially, liquid-liquid phase separation behavior of DNA.
- To provide additional experimental evidence on allosteric DNA effects induced by DNA mismatches.
- To explore the variability of mechanical effects among different kinds of mismatches.

# DNA and RNA mechanics at the service of nanotechnology

Quoting Richard Feynman, 'what I cannot create, I do not understand'. True comprehension of natural processes is achieved when those processes can be customized to fulfill our necessities. In the last few decades, the field of DNA nanotechnology has provided an outstanding example of this philosophy. DNA nanostructures

can be designed from molecular models that incorporate wellcharacterized biophysical properties of the DNA, including mechanical parameters such as the persistence length, the twist stiffness, or the twist-stretch coupling (Dietz et al., 2009; Ouldridge et al., 2010; Castro et al., 2011; Maffeo and Aksimentiev, 2020). Nevertheless, when conceiving DNA as nanomaterial, considerations of sequence effects are often limited to base pairing and stacking interactions (Doye et al., 2013). Only in few cases, additional sequence-dependent features of dsDNA, such as the B-Z transition or A-tract curvature, have been exploited to achieve novel functionalities of DNA nanodevices, such as molecular switches or curved DNA trajectories (Mao et al., 1999; Iric et al., 2018). Incorporating sequence-dependent biophysical properties, such as the ones delineated in Table 2, in the design of future DNA nanodevices could therefore expand the potential of DNA as nanotechnological material. Recent exciting developments in coarse-grain models of DNA mechanics hold great promise for this ambitious goal (Ouldridge et al., 2010; Edens et al., 2012; Freeman et al., 2014; Chakraborty et al., 2018; Maffeo and Aksimentiev, 2020).

In addition to DNA, the field of RNA nanotechnology is becoming increasingly popular. An improved quantitative comprehension of sequence-dependent dsRNA structure and flexibility will surely accompany a sustained and solid development of the RNA nanotechnology field (Guo, 2010; Jasinski *et al.*, 2017). Note that the rules that govern dsRNA mechanics and structure are different from those of dsDNA (see Table 4 and references therein). Namely, RNA possesses its own material properties, different from those of DNA, and might therefore offer new, unforeseen possibilities in terms of molecular design. Exploiting the unique biophysical properties of RNA for nanotechnological applications will be an exciting challenge for the years to come.

In conclusion, we foresee the following challenges for expanding the application of nucleic acids in nanotechnology:

- To exploit sequence-dependent structural and mechanical features in the design of DNA-based nanostructures.
- To advance our current knowledge on dsRNA mechanics and exploit these and other biophysical properties of RNA for improved design of RNA-based nanostructures.

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#### References

Abels JA, Moreno-Herrero F, van der Heijden T, Dekker C and Dekker N (2005) Single-molecule measurements of the persistence length of doublestranded RNA. *Biophysical Journal* 88, 2737–2744.

Abou Assi H, Garavis M, Gonzalez C and Damha MJ (2018) i-Motif DNA: structural features and significance to cell biology. *Nucleic Acids Research* 46, 8038–8056.

Bao L, Zhang Xi, Shi Y-Z, Wu Y-Y and Tan Z-J (2017) Understanding the relative flexibility of RNA and DNA duplexes: stretching and twist-stretch coupling. *Biophysical Journal* 112, 1094–1104.

- Basu A, Bobrovnikov D G, Cieza B, Qureshi Z and Ha T (2020) Deciphering the mechanical code of genome and epigenome. bioRxiv, 2020.08.22.262352.
- Basu A, Bobrovnikov DG, Qureshi Z, Kayikcioglu T, Ngo TTM, Anand R, Eustermann S, Cieza B, Morgan MT, Hejna M, Rube T, Hopfner KP, Wolberger C, Song JS and Ha T (2021) Measuring DNA mechanics on the genome scale. *Nature* 589, 462–467.
- Baumann CG, Smith SB, Bloomfield VA and Bustamante C (1997) Ionic effects on the elasticity of single DNA molecules. Proceedings of the National Academy of Sciences of the United States of America 94, 6185– 6190
- Berman HM, Olson WK, Beveridge DL, Westbrook J, Gelbin A, Demeny T, Hsieh SH, AR Srinivasan and B Schneider (1992) The nucleic acid database. A comprehensive relational database of three-dimensional structures of nucleic acids. *Biophysical Journal* 63, 751–759.
- Beššeová I, Banas P, Kuhrova P, Kosinova P, Otyepka M and Sponer J (2012) Simulations of A-RNA duplexes. The effect of sequence, solute force field, water model, and salt concentration. The Journal of Physical Chemistry B 116, 9899–9916.
- **Bloomfield VA** (1997) DNA condensation by multivalent cations. *Biopolymers* 44, 269–282.
- Bloomfield VA, Crothers D and Tinoco I (2000) Nucleic Acids: Structure, Properties, and Functions. University Science Books.
- Bouchiat C, Wang MD, Allemand J-F, Strick T, Block SM and Croquette V (1999) Estimating the persistence length of a worm-like chain molecule from force–extension measurements. *Biophysical Journal* 76, 409–413.
- Brunet A, Chevalier S, Destainville N, Manghi M, Rousseau P, Salhi M, Salome L and Tardin C (2015a) Probing a label-free local bend in DNA by single molecule tethered particle motion. *Nucleic Acids Research* 43, e72–e72.
- Brunet A, Tardin C, Salome L, Rousseau P, Destainville N and Manghi M (2015b) Dependence of DNA persistence length on ionic strength of solutions with monovalent and divalent salts: a joint theory–experiment study. *Macromolecules* 48, 3641–3652.
- Brutzer H, Luzzietti N, Klaue D and Seidel R (2010) Energetics at the DNA supercoiling transition. *Biophysical Journal* **98**, 1267–1276.
- Bryant Z, Stone MD, Gore J, Smith SB, Cozzarelli NR and Bustamante C (2003) Structural transitions and elasticity from torque measurements on DNA. *Nature* 424, 338–341.
- Bryant Z, Oberstrass FC and Basu A (2012) Recent developments in single-molecule DNA mechanics. Current Opinion in Structural Biology 22, 304–312.
- Bustamante C, Marko JF, Siggia ED and Smith S (1994) Entropic elasticity of lambda-phage DNA. Science (New York, N.Y.) 265, 1599.
- Bustamante C, Bryant Z and Smith SB (2003) Ten years of tension: single-molecule DNA mechanics. *Nature* 421, 423.
- Buzón P, Maity S and Roos WH (2020) Physical virology: from virus self-assembly to particle mechanics. WIREs Nanomedicine and Nanobiotechnology n/a, e1613.
- Camunas-Soler J, Ribezzi-Crivellari M and Ritort F (2016) Elastic properties of nucleic acids by single-molecule force spectroscopy. Annual Review of Biophysics 45, 65–84.
- Carter AP, Clemons WM, Brodersen DE, Morgan-Warren RJ, Wimberly BT and Ramakrishnan V (2000) Functional insights from the structure of the 30S ribosomal subunit and its interactions with antibiotics. *Nature* 407, 340–348.
- Castro CE, Kilchherr F, Kim D-N, Shiao EL, Wauer T, Wortmann P, Bathe M and Dietz H (2011) A primer to scaffolded DNA origami. Nature Methods 8, 221–229.
- Chakraborty D and Wales DJ (2017) Probing helical transitions in a DNA duplex. Physical Chemistry Chemical Physics 19, 878–892.
- Chakraborty D, Hori N and Thirumalai D (2018) Sequence-dependent three interaction site model for single- and double-stranded DNA. *Journal of Chemical Theory and Computation* 14, 3763–3779.
- Chakraborty S, Steinbach PJ, Paul D, Mu H, Broyde S, Min J-H and Ansari A (2017) Enhanced spontaneous DNA twisting/bending fluctuations unveiled

by fluorescence lifetime distributions promote mismatch recognition by the Rad4 nucleotide excision repair complex. *Nucleic Acids Research* **46**, 1240–1255.

- Chen Y, Tokuda JM, Topping T, Meisburger SP, Pabit SA, Gloss LM and Pollack L (2017) Asymmetric unwrapping of nucleosomal DNA propagates asymmetric opening and dissociation of the histone core. *Proceedings of the National Academy of Sciences* 114, 334.
- Chou F-C, Lipfert J and Das R (2014) Blind predictions of DNA and RNA tweezers experiments with force and torque. PLoS Computational Biology 10, e1003756.
- Cortini R, Barbi M, Caré BR, Lavelle C, Lesne A, Mozziconacci J and Victor J-M (2016) The physics of epigenetics. Reviews of Modern Physics 88, 025002.
- Craggs TD, Sustarsic M, Plochowietz A, Mosayebi M, Kaju H, Cuthbert A, Hohlbein J, Domicevica L, Biggin PC, Doye Jonathan PK and Kapanidis AN (2019) Substrate conformational dynamics facilitate structure-specific recognition of gapped DNA by DNA polymerase. *Nucleic Acids Research* 47, 10788–10800.
- Crick FHC and Klug A (1975) Kinky helix. Nature 255, 530-533.
- Crut A, Koster DA, Seidel R, Wiggins CH and Dekker NH (2007) Fast dynamics of supercoiled DNA revealed by single-molecule experiments. Proceedings of the National Academy of Sciences 104, 11957–11962.
- Culkin J, De Bruin L, Tompitak M, Phillips R & Schiessel H (2017) The role of DNA sequence in nucleosome breathing. *The European Physical Journal E* 40, 106.
- Curuksu J, Zacharias M, Lavery R and Zakrzewska K (2009) Local and global effects of strong DNA bending induced during molecular dynamics simulations. *Nucleic Acids Research* 37, 3766–3773.
- Dans PD, Faustino I, Battistini F, Zakrzewska K, Lavery R and Orozco M (2014) Unraveling the sequence-dependent polymorphic behavior of d(CpG) steps in B-DNA. *Nucleic Acids Research* **42**, 11304–11320.
- Dantas Machado AC, Zhou T, Rao S, Goel P, Rastogi C, Lazarovici A, Bussemaker HJ and Rohs R (2014) Evolving insights on how cytosine methylation affects protein–DNA binding. Briefings in Functional Genomics 14, 61–73.
- Deaton AM and Bird A (2011) CpG islands and the regulation of transcription. Genes & Development 25, 1010-1022.
- Dickerson RE (1998) DNA bending: the prevalence of kinkiness and the virtues of normality. *Nucleic Acids Research* 26, 1906–1926.
- Dietz H, Douglas SM and Shih WM (2009) Folding DNA into twisted and curved nanoscale shapes. Science (New York, N.Y.) 325, 725–730.
- Dittmore A, Brahmachari S, Takagi Y, Marko JF and Neuman KC (2017) Supercoiling DNA locates mismatches. *Physical Review Letters* **119**, 147801.
- Dock-Bregeon AC, Chevrier B, Podjarny A, Johnson J, De Bear JS, Gough GR, Gilham PT and Moras D (1989) Crystallographic structure of an RNA helix: [U(UA)6A]2. *Journal of Molecular Biology* **209**, 459–474.
- Dohnalová H, Dršata T, Šponer J, Zacharias M, Lipfert J and Lankaš F (2020) Compensatory mechanisms in temperature dependence of DNA double helical structure: bending and elongation. *Journal of Chemical Theory and Computation* 16, 2857–2863.
- Doye JPK, Ouldridge TE, Louis AA, Romano F, Šulc P, Matek C, Snodin BEK, Rovigatti L, Schreck JS, Harrison RM and Smith WPJ (2013) Coarse-graining DNA for simulations of DNA nanotechnology. *Physical Chemistry Chemical Physics* 15, 20395–20414.
- Drozdetski AV, Tolokh IS, Pollack L, Baker N and Onufriev AV (2016)
  Opposing effects of multivalent ions on the flexibility of DNA and RNA.

  Physical Review Letters 117, 028101–028101.
- Dršata T, Špačková N, Jurečka P, Zgarbová M, Šponer J and Lankaš F (2014) Mechanical properties of symmetric and asymmetric DNA A-tracts: implications for looping and nucleosome positioning. Nucleic Acids Research 42, 7383–7394.
- Edens LE, Brozik JA and Keller DJ (2012) Coarse-grained model DNA: structure, sequences, stems, circles, hairpins. *The Journal of Physical Chemistry B* 116, 14735–14743.
- Esteller M (2008) Epigenetics in cancer. New England Journal of Medicine 358, 1148–1159.
- Field Y, Kaplan N, Fondufe-Mittendorf Y, Moore IK, Sharon E, Lubling Y, Widom J and Segal E (2008) Distinct modes of regulation by chromatin

- encoded through nucleosome positioning signals. *PLoS Computational Biology* **4**, e1000216.
- Fire A, Alcazar R and Tan F (2006) Unusual DNA structures associated with germline genetic activity in *Caenorhabditis elegans*. *Genetics* 173, 1259–1273.
- Fogg JM, Randall GL, Pettitt BM, Sumners DWL, Harris SA and Zechiedrich L (2012) Bullied no more: when and how DNA shoves proteins around. *Quarterly Reviews of Biophysics* 45, 257–299.
- Forth S, Deufel C, Sheinin MY, Daniels B, Sethna JP and Wangi MD (2008) Abrupt buckling transition observed during the plectoneme formation of individual DNA molecules. *Physical Review Letters* 100, 148301.
- Freeman GS, Hinckley DM, Lequieu JP, Whitmer JK and De Pablo JJ (2014)
  Coarse-grained modeling of DNA curvature. *The Journal of Chemical Physics* 141, 165103.
- Fu H, Zhang C, Qiang X-W, Yang Y-J, Dai L, Tan Z-J and Zhang X-H (2020) Opposite effects of high-valent cations on the elasticities of DNA and RNA duplexes revealed by magnetic tweezers. *Physical Review Letters* 124, 058101.
- Gai D, Chang YP and Chen XS (2010) Origin DNA melting and unwinding in DNA replication. Current Opinion in Structural Biology 20, 756–762.
- Ganji M, Kim SH, Van Der Torre J, Abbondanzieri E and Dekker C (2016) Intercalation-based single-molecule fluorescence assay to study DNA supercoil dynamics. Nano Letters 16, 4699–4707.
- Garcia HG, Grayson P, Han L, Inamdar M, Kondev J, Nelson PC, Phillips R, Widom J and Wiggins PA (2007) Biological consequences of tightly bent DNA: the other life of a macromolecular celebrity. *Biopolymers* 85, 115–130
- Geggier S and Vologodskii A (2010) Sequence dependence of DNA bending rigidity. Proceedings of the National Academy of Sciences 107, 15421.
- Gore J, Bryant Z, Nöllmann M, Le MU, Cozzarelli NR and Bustamante C (2006a) DNA overwinds when stretched. *Nature* **442**, 836–839.
- Gore J, Bryant Z, Stone MD, Nöllmann M, Cozzarelli NR and Bustamante C (2006b) Mechanochemical analysis of DNA gyrase using rotor bead tracking. *Nature* 439, 100–104.
- Gross P, Laurens N, Oddershede LB, Bockelmann U, Peterman EJG and Wuite GJL (2011) Quantifying how DNA stretches, melts and changes twist under tension. *Nature Physics* 7, 731.
- Guilbaud S, Salomé L, Destainville N, Manghi M and Tardin C (2019)Dependence of DNA persistence length on ionic strength and ion type.Physical Review Letters 122, 028102.
- Guo P (2010) The emerging field of RNA nanotechnology. Nature Nanotechnology 5, 833–842.
- Ha B-Y and Thirumalai D (2003) Bending rigidity of stiff polyelectrolyte chains: a single chain and a bundle of multichains. Macromolecules 36, 9658–9666.
- Hagerman PJ (1997) Flexibility of RNA. Annual Review of Biophysics and Biomolecular Structure 26, 139–156.
- Haran TE and Mohanty U (2009) The unique structure of A-tracts and intrinsic DNA bending. *Quarterly Reviews of Biophysics* **42**, 41–81.
- Harrington RE (1978) Opticohydrodynamic properties of high-molecularweight DNA. III. The effects of NaCl concentration. Biopolymers 17, 919–936.
- Harrison RM, Romano F, Ouldridge TE, Louis AA and Doye JPK (2019) Identifying physical causes of apparent enhanced cyclization of short DNA molecules with a coarse-grained model. *Journal of Chemical Theory and Computation* **15**, 4660–4672.
- Heenan PR and Perkins TT (2019) Imaging DNA equilibrated onto mica in liquid using biochemically relevant deposition conditions. ACS Nano 13, 4220–4229.
- **Herbert A** (2019) A genetic instruction code based on DNA conformation. *Trends in Genetics* **35**, 887–890.
- Herrero-Galán E, Fuentes-Perez ME, Carrasco C, Valpuesta JM, Carrascosa JL, Moreno-Herrero F and Arias-Gonzalez JR (2013) Mechanical identities of RNA and DNA double helices unveiled at the single-molecule level. *Journal of the American Chemical Society* 135, 122–131.
- Huguet JM, Bizarro CV, Forns N, Smith SB, Bustamante C and Ritort F (2010) Single-molecule derivation of salt dependent base-pair free energies in DNA. *Proceedings of the National Academy of Sciences* **107**, 15431.
- Iric K, Subramanian M, Oertel J, Agarwal NP, Matthies M, Periole X, Sakmar TP, Huber T, Fahmy K and Schmidt TL (2018) DNA-encircled lipid bilayers. Nanoscale 10, 18463–18467.

- Irobalieva RN, Fogg JM, Catanese DJ, Sutthibutpong T, Chen M, Barker AK, Ludtke SJ, Harris SA, Schmid MF, Chiu W and Zechiedrich L (2015) Structural diversity of supercoiled DNA. *Nature Communications* 6, 8440
- Jasinski D, Haque F, Binzel DW and Guo P (2017) Advancement of the emerging field of RNA nanotechnology. ACS Nano 11, 1142–1164.
- Jeong J and Kim HD (2019) Base-pair mismatch can destabilize small DNA loops through cooperative kinking. *Physical Review Letters* 122, 218101.
- Jiao H, Wachsmuth L, Kumari S, Schwarzer R, Lin J, Eren RO, Fisher A, Lane R, Young GR, Kassiotis G, Kaiser WJ and Pasparakis M (2020) Z-nucleic-acid sensing triggers ZBP1-dependent necroptosis and inflammation. *Nature* 580, 391–395.
- Kang H, Yoo J, Sohn B-K, Lee S-W, Lee HS, Ma W, Kee J-M, Aksimentiev A and Kim H (2018) Sequence-dependent DNA condensation as a driving force of DNA phase separation. *Nucleic Acids Research* 46, 9401–9413.
- Kaplan N, Moore IK, Fondufe-Mittendorf Y, Gossett AJ, Tillo D, Field Y, Leproust EM, Hughes TR, Lieb JD, Widom J and Segal E (2009) The DNA-encoded nucleosome organization of a eukaryotic genome. *Nature* 458, 362–366.
- Katz AM, Tolokh IS, Pabit SA, Baker N, Onufriev AV and Pollack L (2017)
  Spermine condenses DNA, but not RNA duplexes. *Biophysical Journal* 112, 22–30
- Kilchherr F, Wachauf C, Pelz B, Rief M, Zacharias M and Dietz H (2016) Single-molecule dissection of stacking forces in DNA. Science (New York, N.Y.) 353, aaf5508.
- Kim S, Broströmer E, Xing D, Jin J, Chong S, Ge H, Wang S, Gu C, Yang L, Gao YQ, Su X-D, Sun Y and Xie XS (2013) Probing allostery through DNA. Science (New York, N.Y.) 339, 816.
- Kim SH, Ganji M, Kim, E, Van Der Torre J, Abbondanzieri E and Dekker C (2018a) Unveiling the pathway to Z-DNA in the protein-induced B–Z transition. Nucleic Acids Research 46, 4129–4137.
- Kim SH, Lim S-H, Lee A-R, Kwon Do h, Song Hyun k, Lee J-H, Cho M, Johner A, Lee N-K and Hong S-C (2018b) DNA sequence encodes the position of DNA supercoils. eLife 7, e36557.
- King GA, Burla F, Peterman EJG and Wuite GJL (2019) Supercoiling DNA optically. Proceedings of the National Academy of Sciences 116, 26534.
- Koo H-S, Wu H-M and Crothers DM (1986) DNA bending at adenine thymine tracts. Nature 320, 501.
- Kriegel F, Ermann N and Lipfert J (2017a) Probing the mechanical properties, conformational changes, and interactions of nucleic acids with magnetic tweezers. *Journal of Structural Biology* 197, 26–36.
- Kriegel F, Ermann N, Forbes R, Dulin D, Dekker NH and Lipfert J (2017b)Probing the salt dependence of the torsional stiffness of DNA by multiplexed magnetic torque tweezers. *Nucleic Acids Research* 45, 5920–5929.
- Krokan HE and Bjørås M (2013) Base excision repair. Cold Spring Harbor Perspectives in Biology 5, a012583.
- Kulkarni M and Mukherjee A (2017) Understanding B-DNA to A-DNA transition in the right-handed DNA helix: perspective from a local to global transition. *Progress in Biophysics and Molecular Biology* 128, 63–73.
- Kunkel TA and Erie DA (2015) Eukaryotic mismatch repair in relation to DNA replication. Annual Review of Genetics 49, 291–313.
- Lankaš F (2020) Simple, but not too simple: modeling the dynamics of DNA and RNA buckling. Biophysical Journal 118, 1514–1516.
- Lankaš F, Lavery R and Maddocks JH (2006) Kinking occurs during molecular dynamics simulations of small DNA minicircles. Structure (London, England: 1993) 14, 1527–1534.
- Larson MH, Landick R and Block SM (2011) Single-molecule studies of RNA polymerase: one singular sensation, every little step it takes. *Molecular Cell* 41, 249–262
- Lebel P, Basu A, Oberstrass FC, Tretter EM and Bryant Z (2014) Gold rotor bead tracking for high-speed measurements of DNA twist, torque and extension. *Nature Methods* 11, 456–462.
- Lee M, Kim SH, Kim KK and Hong S-C (2010a) Minute negative superhelicity is sufficient to induce the B-Z transition in the presence of low tension. Proceedings of the National Academy of Sciences 107, 4985.
- Lee J, Kim SH and Hong S-C (2010b) Transition between B-DNA and Z-DNA: free energy landscape for the B-Z junction propagation. *The Journal of Physical Chemistry B* 114, 9872–9881.

- Lequieu J, Córdoba A, Schwartz DC and De Pablo JJ (2016) Tension-dependent free energies of nucleosome unwrapping. ACS Central Science 2, 660–666.
- Li L, Pabit SA, Meisburger SP and Pollack L (2011) Double-stranded RNA resists condensation. *Physical Review Letters* 106, 108101–108101.
- Liebl K, Drsata T, Lankas F, Lipfert J and Zacharias M (2015) Explaining the striking difference in twist-stretch coupling between DNA and RNA: a comparative molecular dynamics analysis. *Nucleic Acids Research* 43, 10143-10156.
- Liebl K and Zacharias M (2018) How methyl-sugar interactions determine DNA structure and flexibility. Nucleic Acids Research 47, 1132–1140.
- Lionnet T, Joubaud S, Lavery R, Bensimon D and Croquette V (2006) Wringing out DNA. Physical Review Letters 96, 178102.
- Lipfert J, Kerssemakers JWJ, Jager T and Dekker NH (2010) Magnetic torque tweezers: measuring torsional stiffness in DNA and RecA-DNA filaments. *Nature Methods* 7, 977–980.
- Lipfert J, Skinner GM, Keegstra JM, Hensgens T, Jager T, Dulin D, Köber M, Yu Z, Donkers SP, Chou F-C, Das R and Dekker NH (2014) Double-stranded RNA under force and torque: similarities to and striking differences from double-stranded DNA. Proceedings of the National Academy of Sciences of the United States of America 111, 15408–15413.
- Lipfert J, Van Oene MM, Lee M, Pedaci F and Dekker NH (2015) Torque spectroscopy for the study of rotary motion in biological systems. Chemical Reviews 115, 1449–1474.
- Lowary PT and Widom J (1998) New DNA sequence rules for high affinity binding to histone octamer and sequence-directed nucleosome positioning. *Journal of Molecular Biology* 276, 19–42.
- Lu X-J, Shakked Z and Olson WK (2000) A-form conformational motifs in ligand-bound DNA structures. *Journal of Molecular Biology* 300, 819–840.
- Madariaga-Marcos J, Hormeño S, Pastrana CL, Fisher GLM, Dillingham MS and Moreno-Herrero F (2018) Force determination in lateral magnetic tweezers combined with TIRF microscopy. *Nanoscale* **10**, 4579–4590.
- Maffeo C and Aksimentiev A (2020) MrDNA: a multi-resolution model for predicting the structure and dynamics of DNA systems. *Nucleic Acids Research* 48, 5135–5146.
- Mandal S, Hoque ME and Mao H (2019) Single-molecule investigations of G-quadruplex. In Yang D and Lin C (eds), G-Quadruplex Nucleic Acids: Methods and Protocols. New York, NY: Springer New York, pp. 275–298.
- Mao C, Sun W, Shen Z and Seeman NC (1999) A nanomechanical device based on the B–Z transition of DNA. *Nature* 397: 144–146.
- Marin-Gonzalez A, Vilhena JG, Perez R and Moreno-Herrero F (2017)
  Understanding the mechanical response of double-stranded DNA and RNA under constant stretching forces using all-atom molecular dynamics.

  Proceedings of the National Academy of Sciences 114, 7049.
- Marin-Gonzalez A, Vilhena JG, Moreno-Herrero F and Perez R (2019a)DNA crookedness regulates DNA mechanical properties at short length scales. *Physical Review Letters* 122, 048102.
- Marin-Gonzalez A, Vilhena JG, Moreno-Herrero F and Perez R (2019b) Sequence-dependent mechanical properties of double-stranded RNA. Nanoscale 11, 21471–21478.
- Marin-Gonzalez A, Pastrana CL, Bocanegra R, Martín-González A, Vilhena JG, Pérez R, Ibarra B, Aicart-Ramos C and Moreno-Herrero F (2020a) Understanding the paradoxical mechanical response of in-phase A-tracts at different force regimes. *Nucleic Acids Research* 48, 5024–5036.
- Marin-Gonzalez A, Aicart-Ramos C, Marin-Baquero M, Martín-González A, Suomalainen M, Kannan A, Vilhena JG, Greber UF, Moreno-Herrero F and Pérez R (2020b) Double-stranded RNA bending by AU-tract sequences. Nucleic Acids Research 48, 12917–12928.
- Marko JF and Siggia ED (1994) Fluctuations and supercoiling of DNA. Science (New York, N.Y.) 265, 506.
- Marko JF and Siggia ED (1995a) Stretching DNA. Macromolecules 28, 8759–8770.
- Marko JF and Siggia ED (1995b) Statistical mechanics of supercoiled DNA. *Physical Review E* 52, 2912–2938.
- Marmur J and Doty P (1962) Determination of the base composition of deoxyribonucleic acid from its thermal denaturation temperature. *Journal of Molecular Biology* 5, 109–118.

- Marteijn JA, Lans H, Vermeulen W and Hoeijmakers JHJ (2014) Understanding nucleotide excision repair and its roles in cancer and ageing. Nature Reviews Molecular Cell Biology 15, 465–481.
- Matek C, Ouldridge TE, Doye JPK and Louis AA (2015a) Plectoneme tip bubbles: coupled denaturation and writhing in supercoiled DNA. Scientific Reports 5, 7655.
- Matek C, Sulc P, Randisi F, Doye JPK and Louis AA (2015b) Coarse-grained modelling of supercoiled RNA. The Journal of Chemical Physics 143, 243122.
- Mauney AW, Tokuda JM, Gloss LM, Gonzalez O and Pollack L (2018) Local DNA sequence controls asymmetry of DNA unwrapping from nucleosome core particles. *Biophysical Journal* 115, 773–781.
- Mitchell JS, Glowacki J, Grandchamp AE, Manning RS and Maddocks JH (2017) Sequence-dependent persistence lengths of DNA. Journal of Chemical Theory and Computation 13, 1539–1555.
- Moradi M, Babin V, Roland C and Sagui C (2013) Reaction path ensemble of the B–Z-DNA transition: a comprehensive atomistic study. *Nucleic Acids Research* 41, 33–43.
- Moreno-Herrero F, Seidel R, Johnson SM, Fire A and Dekker NH (2006) Structural analysis of hyperperiodic DNA from *Caenorhabditis elegans*. *Nucleic Acids Research* **34**, 3057–3066.
- Natrajan G, Lamers MH, Enzlin JH, Winterwerp HHK, Perrakis A and Sixma TK (2003) Structures of *Escherichia coli* DNA mismatch repair enzyme MutS in complex with different mismatches: a common recognition mode for diverse substrates. *Nucleic Acids Research* 31, 4814–4821.
- Nelson HCM, Finch JT, Luisi BF and Klug A (1987) The structure of an oligo (dA)·oligo(dT) tract and its biological implications. *Nature* 330, 221.
- Ngo TTM, Zhang Q, Zhou R, Yodh Jaya G and Ha T (2015) Asymmetric unwrapping of nucleosomes under tension directed by DNA local flexibility. *Cell* 160, 1135–1144.
- Ngo TTM, Yoo J, Dai Q, Zhang Q, He C, Aksimentiev A and Ha T (2016) Effects of cytosine modifications on DNA flexibility and nucleosome mechanical stability. *Nature Communications* 7, 10813.
- Nissen P, Ippolito JA, Ban N, Moore PB and Steitz TA (2001) RNA tertiary interactions in the large ribosomal subunit: the A-minor motif. *Proceedings of the National Academy of Sciences* **98**, 4899.
- Nomidis SK, Kriegel F, Vanderlinden W, Lipfert J and Carlon E (2017)
  Twist-bend coupling and the torsional response of double-stranded DNA.

  Physical Review Letters 118, 217801.
- Noy A, Pérez A, Lankas F, Javier Luque F and Orozco M (2004) Relative flexibility of DNA and RNA: a molecular dynamics study. *Journal of Molecular Biology* **343**, 627–638.
- Oberstrass FC, Fernandes LE and Bryant Z (2012) Torque measurements reveal sequence-specific cooperative transitions in supercoiled DNA. *Proceedings of the National Academy of Sciences* 109, 6106.
- Oberstrass FC, Fernandes LE, Lebel P and Bryant Z (2013) Torque spectroscopy of DNA: base-pair stability, boundary effects, backbending, and breathing dynamics. *Physical Review Letters* 110, 178103.
- Olson WK and Zhurkin VB (2011) Working the kinks out of nucleosomal DNA. Current Opinion in Structural Biology 21, 348–357.
- Olson WK, Gorin AA, Lu X-J, Hock LM and Zhurkin VB (1998) DNA sequence-dependent deformability deduced from protein–DNA crystal complexes. *Proceedings of the National Academy of Sciences* **95**, 11163.
- Onufriev AV and Schiessel H (2019) The nucleosome: from structure to function through physics. Current Opinion in Structural Biology 56, 119-130.
- Ortiz V and de Pablo JJ (2011) Molecular origins of DNA flexibility: sequence effects on conformational and mechanical properties. *Physical Review Letters* **106**, 238107–238107.
- Ott K, Martini L, Lipfert J and Gerland U (2020) Dynamics of the buckling transition in double-stranded DNA and RNA. *Biophysical Journal* 118, 1690–1701.
- Ouldridge TE, Louis AA and Doye JPK (2010) DNA nanotweezers studied with a coarse-grained model of DNA. Physical Review Letters 104, 178101.
- Pasi M, Maddocks JH, Beveridge D, Bishop TC, Case DA, Cheatham T, Iii Dans PD, Jayaram B, Lankas F, Laughton C, Mitchell J, Osman R, Orozco M, Pérez A, Petkevičiūtė D, Spackova N, Sponer J, Zakrzewska K and Lavery R (2014) μABC: a systematic microsecond molecular

- dynamics study of tetranucleotide sequence effects in B-DNA. *Nucleic Acids Research* **42**, 12272–12283.
- Paul D, Mu H, Zhao H, Ouerfelli O, Jeffrey PD, Broyde S and Min J-H (2019) Structure and mechanism of pyrimidine-pyrimidone (6-4) photoproduct recognition by the Rad4/XPC nucleotide excision repair complex. Nucleic Acids Research 47, 6015–6028.
- Pérez A, Noy A, Lankas F, Luque FJ and Orozco M (2004) The relative flexibility of B-DNA and A-RNA duplexes: database analysis. *Nucleic Acids Research* 32, 6144–6151.
- Pérez A, Castellazzi CL, Battistini F, Collinet K, Flores O, Deniz O, Ruiz ML, Torrents D, Eritja R, Soler-López M and Orozco M (2012) Impact of methylation on the physical properties of DNA. *Biophysical Journal* 102, 2140-2148.
- **Perona JJ and Hou Y-M** (2007) Indirect readout of tRNA for aminoacylation. *Biochemistry* **46**, 10419–10432.
- Peters JP 3rd and Maher LJ (2010) DNA curvature and flexibility in vitro and in vivo. Quarterly Reviews of Biophysics 43, 23–63.
- Pongor CI, Bianco P, Ferenczy G, Kellermayer R and Kellermayer M (2017)

  Optical trapping nanometry of hypermethylated CPG-island DNA.

  Biophysical Journal 112, 512–522.
- Portella G, Battistini F and Orozco M (2013) Understanding the connection between epigenetic DNA methylation and nucleosome positioning from computer simulations. PLoS Computational Biology 9, e1003354.
- Pyne ALB, Noy A, Main K, Velasco-Berrelleza V, Piperakis MM, Mitchenall LA, Cugliandolo FM, Beton JG, Stevenson CEM, Hoogenboom BW, Bates AD, Maxwell A and Harris SA (2020) Base-pair resolution analysis of the effect of supercoiling on DNA flexibility and recognition. *Nature Communications* 12, 1053.
- Qi Y, Spong MC, Nam K, Banerjee A, Jiralerspong S, Karplus M and Verdine GL (2009) Encounter and extrusion of an intrahelical lesion by a DNA repair enzyme. *Nature* **462**, 762–766.
- Ramirez-Carrozzi VR, Braas D, Bhatt DM, Cheng CS, Hong C, Doty KR, Black JC, Hoffmann A, Carey M and Smale ST (2009) A unifying model for the selective regulation of inducible transcription by CpG islands and nucleosome remodeling. Cell 138, 114–128.
- Rao S, Chiu T-P, Kribelbauer JF, Mann RS, Bussemaker HJ and Rohs R (2018) Systematic prediction of DNA shape changes due to CpG methylation explains epigenetic effects on protein–DNA binding. *Epigenetics & Chromatin* 11, 6.
- Rezaei N, Lyons A and Forde NR (2018) Environmentally controlled curvature of single collagen proteins. *Biophysical Journal* 115, 1457–1469.
- Rich A and Zhang S (2003) Z-DNA: the long road to biological function. Nature Reviews Genetics 4, 566–572.
- Richmond TJ and Davey CA (2003) The structure of DNA in the nucleosome core. *Nature* **423**, 145.
- Rivetti C, Guthold M and Bustamante C (1996) Scanning force microscopy of DNA deposited onto mica: equilibration versus kinetic trapping studied by statistical polymer chain analysis. *Journal of Molecular Biology* 264, 919–932.
- Rivetti C, Walker C and Bustamante C (1998) Polymer chain statistics and conformational analysis of DNA molecules with bends or sections of different flexibility. *Journal of Molecular Biology* **280**, 41–59.
- Roberts RJ and Cheng X (1998) Base flipping. Annual Review of Biochemistry 67, 181–198.
- Rohs R, West SM, Sosinsky A, Liu P, Mann RS and Honig B (2009) The role of DNA shape in protein–DNA recognition. *Nature* 461, 1248.
- Rohs R, Jin X, West SM, Joshi R, Honig B and Mann RS (2010) Origins of specificity in protein–DNA recognition. *Annual Review of Biochemistry* 79, 233–269
- Rossetti G, Dans PD, Gomez-Pinto I, Ivani I, Gonzalez C and Orozco M (2015) The structural impact of DNA mismatches. *Nucleic Acids Research* **43**, 4309–4321.
- Sanchez AM, Volk DE, Gorenstein DG and Lloyd RS (2003) Initiation of repair of A/G mismatches is modulated by sequence context. *DNA Repair* 2, 863–878
- Satange R, Chang C-K and Hou M-H (2018) A survey of recent unusual high-resolution DNA structures provoked by mismatches, repeats and ligand binding. *Nucleic Acids Research* 46, 6416–6434.

- Schimmel P (2018) The emerging complexity of the tRNA world: mammalian tRNAs beyond protein synthesis. *Nature Reviews Molecular Cell Biology* 19, 45–58.
- Segal E and Widom J (2009) Poly(dA:dT) tracts: major determinants of nucleosome organization. Current Opinion in Structural Biology 19, 65-71
- Segal E, Fondufe-Mittendorf Y, Chen L, Thåström A, Field Y, Moore IK, Wang J-PZ and Widom J (2006) A genomic code for nucleosome positioning. Nature 442, 772–778.
- Severin PMD, Zou X, Gaub HE and Schulten K (2011) Cytosine methylation alters DNA mechanical properties. Nucleic Acids Research 39, 8740–8751.
- Shakya A and King JT (2018) DNA local-flexibility-dependent assembly of phase-separated liquid droplets. *Biophysical Journal* 115, 1840–1847.
- Shepherd JW, Greenall RJ, Probert Matt IJ, Noy A and Leake Mark C (2020)
  The emergence of sequence-dependent structural motifs in stretched, torsionally constrained DNA. *Nucleic Acids Research* 48, 1748–1763.
- **Shon MJ, Rah S-H and Yoon T-Y** (2019) Submicrometer elasticity of double-stranded DNA revealed by precision force–extension measurements with magnetic tweezers. *Science Advances* **5**, eaav1697.
- Smith ZD and Meissner A (2013) DNA methylation: roles in mammalian development. *Nature Reviews Genetics* 14, 204–220.
- Smith SB, Finzi L and Bustamante C (1992) Direct mechanical measurements of the elasticity of single DNA molecules by using magnetic beads. Science (New York, N.Y.) 258, 1122.
- Smith SB, Cui Y and Bustamante C (1996) Overstretching B-DNA: the elastic response of individual double-stranded and single-stranded DNA molecules. Science 271, 795–799.
- Šponer J, Bussi G, Krepl M, Banáš P, Bottaro S, Cunha RA, Gil-Ley A, Pinamonti G, Poblete, S, Jurečka, P, Walter NG and Otyepka M (2018) RNA structural dynamics as captured by molecular simulations: a comprehensive overview. *Chemical Reviews* 118, 4177–4338.
- Strick TR, Allemand J-F, Bensimon D, Bensimon A and Croquette V (1996)
  The elasticity of a single supercoiled DNA molecule. Science (New York, N.Y.) 271, 1835.
- Sutthibutpong T, Matek C, Benham C, Slade GG, Noy A, Laughton C, K doye JP, Louis AA and Harris SA (2016) Long-range correlations in the mechanics of small DNA circles under topological stress revealed by multiscale simulation. *Nucleic Acids Research* 44, 9121–9130.
- Szczelkun MD, Tikhomirova MS, Sinkunas T, Gasiunas G, Karvelis T, Pschera P, Siksnys V and Seidel R (2014) Direct observation of R-loop formation by single RNA-guided Cas9 and Cascade effector complexes. Proceedings of the National Academy of Sciences 111, 9798.
- **Thompson JMT and Travers AA** (2004) The structural basis of DNA flexibility. *Philosophical Transactions of the Royal Society of London. Series A: Mathematical, Physical and Engineering Sciences* **362**, 1423–1438.
- Tolokh IS, Pabit SA, Katz AM, Chen Y, Drozdetski A, Baker N, Pollack L and Onufriev AV (2014) Why double-stranded RNA resists condensation. *Nucleic Acids Research* **42**, 10823–10831.
- Unnikrishnan A, Freeman WM, Jackson J, Wren JD, Porter H and Richardson A (2019) The role of DNA methylation in epigenetics of aging. *Pharmacology & Therapeutics* 195, 172–185.
- Vafabakhsh R and Ha T (2012) Extreme bendability of DNA less than 100 base pairs long revealed by single-molecule cyclization. Science 337, 1097–1101.
- van Loenhout MTJ, de Grunt MV and Dekker C (2012) Dynamics of DNA supercoils. Science (New York, N.Y.) 338, 94–97.
- Vlijm R, Torre JVD and Dekker C (2015) Counterintuitive DNA sequence dependence in supercoiling-induced DNA melting. PLoS ONE 10, e0141576.
- Vologodskii A and Frank-Kamenetskii MD (2013) Strong bending of the DNA double helix. Nucleic Acids Research 41, 6785–6792.
- Vologodskii A and Frank-Kamenetskii MD (2018) DNA melting and energetics of the double helix. Physics of Life Reviews 25, 1–21.
- Walther J, Dans PD, Balaceanu A, Hospital A, Bayarri G and Orozco M (2020) A multi-modal coarse grained model of DNA flexibility mappable to the atomistic level. *Nucleic Acids Research* 48, e29.
- Wang MD, Yin H, Landick R, Gelles J and Block SM (1997) Stretching DNA with optical tweezers. Biophysical Journal 72, 1335–1346.

Waters JT, Lu X-J, Galindo-Murillo R, Gumbart JC, Kim HD, Cheatham TE and Harvey SC (2016) Transitions of double-stranded DNA between the A- and B-forms. *The Journal of Physical Chemistry B* 120, 8449–8456.

20

- Wenner JR, Williams MC, Rouzina I and Bloomfield VA (2002) Salt dependence of the elasticity and overstretching transition of single DNA molecules. *Biophysical Journal* 82, 3160–3169.
- Widom J (2001) Role of DNA sequence in nucleosome stability and dynamics. Quarterly Reviews of Biophysics 34, 269–324.
- Wiggins PA, Van Der Heijden T, Moreno-Herrero F, Spakowitz A, Phillips R, Widom J, Dekker C and Nelson PC (2006) High flexibility of DNA on short length scales probed by atomic force microscopy. Nature Nanotechnology 1, 137.
- Winogradoff D and Aksimentiev A (2019) Molecular mechanism of spontaneous nucleosome unraveling. *Journal of Molecular Biology* 431, 323–335.
- Yang Y-J, Dong H-L, Qiang X-W, Fu H, Zhou E-C, Zhang C, Yin L, Chen X-F, Jia F-C, Dai L, Tan Z-J and Zhang X-H (2020) Cytosine methylation enhances DNA condensation revealed by equilibrium measurements using magnetic tweezers. *Journal of the American Chemical Society* 142, 9203–9209.
- Yesselman JD, Denny SK, Bisaria N, Herschlag D, Greenleaf WJ and Das R (2019) Sequence-dependent RNA helix conformational preferences

- predictably impact tertiary structure formation. *Proceedings of the National Academy of Sciences* **116**, 16847.
- Yoo J, Kim H, Aksimentiev A and Ha T (2016) Direct evidence for sequence-dependent attraction between double-stranded DNA controlled by methylation. *Nature Communications* 7, 11045.
- Zgarbová M, Šponer J, Otyepka M, Cheatham TE, Galindo-Murillo R and Jurečka P (2015) Refinement of the sugar-phosphate backbone torsion beta for AMBER force fields improves the description of Z- and B-DNA. *Journal of Chemical Theory and Computation* 11, 5723–5736.
- Zhang Y and Crothers DM (2003) High-throughput approach for detection of DNA bending and flexibility based on cyclization. *Proceedings of the National Academy of Sciences* 100, 3161.
- Zhang X, Ding K, Yu X, Chang W, Sun J and Hong Zhou Z (2015) In situ structures of the segmented genome and RNA polymerase complex inside a dsRNA virus. *Nature* **527**, 531–534.
- Zhang B, Zheng W, Papoian GA and Wolynes PG (2016) Exploring the free energy landscape of nucleosomes. *Journal of the American Chemical Society* 138, 8126–8133.
- Zhurkin VB and Olson WK (2013) Can nucleosomal DNA be described by an elastic model?: Comment on 'Sequence-dependent collective properties of DNAs and their role in biological systems' by Pasquale De Santis and Anita Scipioni. *Physics of Life Reviews* 10, 70–72.