Helicase on DNA: A Phase coexistence based mechanism

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We propose a phase coexistence based mechanism for activity of helicases, ubiquitous enzymes that unwind double stranded DNA. The helicase-DNA complex constitutes a fixed-stretch ensemble that entails a coexistence of domains of zipped and unzipped phases of DNA, separated by a domain wall. The motor action of the helicase leads to a change in the position of the fixed constraint thereby shifting the domain wall on dsDNA. We associate this off-equilibrium domain wall motion with the unzipping activity of helicase. We show that this proposal gives a clear and consistent explanation of the main observed features of helicases.

Nucleic acid helicases are defined as enzymes that translocate directionally through double stranded nucleic acid substrates to catalyze the separation of the complementary strands. They facilitate various biological processes such as DNA replication, recombination and repair, RNA transcription, editing and splicing. There are several structural varieties of helicases like monomeric (e.g. PcrA), dimeric (e.g. Rep), trimeric (e.g. RecBCD), tetrameric (e.g. RNA polymerase) or closed hexameric (e.g. DnaB), but all use the hydrolysis of ATP to ADP as the preferred source of energy.

Bulk behaviour in solutions like average unwinding rates, step size, average number of base-pairs opened per helicase, etc., are known for a few helicases like Hepatitis-C virus helicase, PcrA, DnaB and others. Much attention has recently been devoted toward a quantitative characterization of RecBCD enzymes, a multifunctional trimeric protein complex (the products of the recB, recC and recD genes) that participates in the repair of chromosomal DNA through homologous recombination. In bacteria, like Escherichia coli, RecBCD is involved, e.g., in protection against damages by UV or gamma irradiation, and infection by bacteriophages. In all cases, the full functionality of RecBCD relies on the helicase and the nuclease actions of its subunits. The use of single-molecule, micromanipulation tools allowed for monitoring in detail translocation, unwinding and processivity (rate of dissociation) of individual RecBCD enzyme molecules on dsDNA. Such experiments elucidated several new aspects of helicase behaviour and showed that many properties could be more related to general principles than on specific chemical details. In particular it has been observed that (a) RecBCD unwinds dsDNA at a uniform rate, over a wide range of ATP concentrations, as it moves on one strand, (b) the nuclease activity does not affect unzipping, and c) the helicase can work in presence of DNA gaps upto certain lengths. More recently, winding-rewinding for E.Coli Rep helicase-DNA complex has been observed at a single molecule level.

Despite these varieties of experimental findings no clear mechanism coupling the motor action and the helicase activity is known yet. To fill this gap, in this paper we present a simple, but powerful argument, based on the principle of phase coexistence, that provides clear and robust explanations to the gross observed features. There are a few biological operational models built on how a helicase presumably might work. The analysis reported here gives a thermodynamic basis to a model called the “wedge model” according to which the motion of the helicase “provides enough force to enable the helicase to destabilize the base pairs at the junction by a process resembling the action of a wedge”. In our proposed mechanism, energy (from ATP) is required for translocation activity or the motor action of the helicase and not directly for base pair breaking and therefore, according to the classification scheme of Ref. this corresponds to passive helicases. Additional features required for active helicases are ignored in this first study. Our proposal is supported by computer simulation of an exactly tractable model. To our knowledge, this is the first theoretical study of the dynamics of a DNA-helicase complex.

We study the joint dynamics of the helicase and the DNA in a two dimensional fork model (Y-Model). The two strands of DNA are represented by two directed and mutually avoiding walks formed by N bases each. In two dimensions, on the square lattice (see Fig. this implies that the two walks follow the positive direction of the diagonal axis (z); in other words the coordinate along such direction always increases. The perpendicular direction x measures, in unit of the elementary square

[8] In general, some nucleoside triphosphate (NTP) is required, e.g. T7 gp4 can use GTP, SV40 large T antigen uses non-ATP nucleotides for unwinding RNA.
diagonal, the distance between the two complementary monomers belonging to the two strands. When this distance is equal to 1 they are considered in contact: a binding energy is gained which is uniform ($\epsilon = 1$) for a homogeneous model of DNA (homo-DNA) but chosen randomly from two different values ($\epsilon_1, \epsilon_2$) for a heterogeneous DNA (hetero-DNA). Notice that due to the geometrical properties of the lattice the two complementary monomers are labeled by the same $z$-coordinate, as one would require for base pairing in DNA. In the Y-model unzipping can occur only processively, e.g. bubbles are suppressed along the chain: the only conformations considered have the first $N - m$ monomers bounded, whereas the remaining $m$ are separated in a Y-like configuration. The fact that the Y-model does not allow rejoining of the unzipped portion of the dsDNA is similar to the geometry observed in the experiment of Ref. [13] (see also Fig. 4 of Ref. [22]). Also, bubbles are suppressed for DNA at temperatures much below its melting temperature $T_m$, temperature below which the two strands are zipped. In the case of homogeneous interaction the exact phase diagram and other static and dynamical quantities can be exactly determined (also in the presence of a stretching force) [14, 17].

The coarse-grained nature of the model[21] needs to be stressed here. Monomers are to be thought of as groups of bases, and ignoring helicity or restricting to two dimensions are more for simplification of the calculation than artifacts[8]. Such coarse-grained models, or even simpler ones, are used in various DNA-related problems [19, 24]

\[ g_c(T) = \frac{T}{2} \cosh^{-1}(1^{1/T} - 1), \quad \langle x \rangle = \tanh\left(\frac{g}{2T}\right), \]

where temperature $T$ is measured in units of $k_B/\epsilon$, $k_B$ being the Boltzmann constant. From these equations the phase coexistence curve ($x)/N$-vs-$T$ can be determined exactly and it is schematically shown in Fig. 1b. Under a fixed-distance constraint, represented by the vertical line in Fig. 1b, the DNA chain splits into domains of zipped and unzipped phases. The length of the unzipped strand can be read off from the upper line of the coexistence curve. This fact can be checked independently and we have verified it not only for the Y-model but also for models that allow bubbles.

We now make our hypothesis. If we think that the helicase, by virtue of its size larger than the separation of the two DNA strands and the excluded volume interaction, acts as a geometrical separator, the helicase-DNA complex constitutes exactly a fixed-stretch ensemble. The unwinding activity can then be simply associated with the motion of the domain wall which necessarily forms and follows the motor action of the helicase. In other
words, the helicase plays a double role: firstly, its presence thermodynamically implies the existence of a domain wall, and secondly, its translocation induces a motion of the domain wall to reach its equilibrium position. Such an approach puts the primary role on the translocation motion. The thermodynamic force, that drives the domain wall toward its equilibrium position, provides the mechanism for base pair opening.

To verify our hypothesis we have numerically studied the DNA model described above mimicking the effects of the helicase with a rod of length \( l \) (see Fig. 1). Dynamics is introduced by a Monte-Carlo procedure. For the DNA, one among the \( 2N \) monomers is randomly chosen and an attempt is made to modify its position with respect to all the others (which remain fixed). The move can in principle increase or decrease by one unit the distance between the strands. The move is accepted according to standard Metropolis rules. A Monte-Carlo unit time is defined as \( 2N \) single-monomer attempted moves. The scaling properties of this DNA dynamics, also in the presence of a stretching force, have already been determined \[16\]. The helicase moves forward (motor action) by unit step, along the \(-z\) direction, on the DNA, if it is not hindered by the chain configuration (excluded volume interaction). The motion is kept unidirectional as found for RecBCD in Ref. \[12\] (see below also). The motion of the helicase is attempted at every Monte Carlo step. In our simulation we start with a configuration where the helicase is attached to one end of a dsDNA (as in experiments of Refs \[12, 13, 14\]). The temperature is maintained constant for the DNA to be in the zipped phase (below \( T_m \)). The overall dynamics is off-equilibrium. Notice that there is no specific interaction but both DNA and helicase dynamics are constrained by the excluded volume.

Throughout the simulation we monitored the position of the last zipped base pair (fork position see Fig. 1a), the zipping probability \( p(s, t) \) that a base pair at site \( s \) at time \( t \) is zipped, and the average position of the helicase at time \( t \). In most simulations the length \( l \) of the helicase was taken equal to 6 (though we studied up to \( l = 14 \)) whereas 1000 thermal averages were necessary. The length \( N \) of the strands were varied according to the different experiments ranging up to \( N = 1000 \). For an analysis of the domain wall, the zipping probability \( p(s, t) \) can be fitted by a function:

\[
p(s, t) = \frac{1}{2} \left[ 1 - \tanh \left( (s - s_0(t))/w(t) \right) \right]
\]

where \( s_0(t) \) and \( w(t) \) are the position and the width, respectively, of the domain wall.

In Fig 2 we plot for two different times the zipping probability \( p(s, t) \) for the homogeneous and for an heterogeneous case. In all cases the domain wall behaviour predicted is well fitted by a tanh profile (Eq. 2). We point out that such domain walls are not found if the DNA strands are noninteracting, \( i.e. \epsilon = 0 \) (see inset of Fig. 2). In Fig. 3 we show the instantaneous positions of the helicase and the domain wall (computed through Eq. 2) as a function of time. Movement starts at \( t = 0 \) when the helicase is released. The two quantities proceed uniformly and cooperatively through the DNA unwinding it. The domain wall evolves toward the equilibrium position whenever the helicase motion is stopped and this position turns out to be not very far away from the instantaneous position. This indicates an adiabatic adjustment of the domain wall to the instantaneous position of the mobile helicase. We stress that there was no interaction between the DNA and the helicase, the latter, as a phantom motor, would have moved with the assigned speed with the DNA remaining bound (since we are below \( T_m \)). It is important to compare this motion with the unzipping dynamics in a fixed force ensemble which shows a characteristic scaling behavior \[10\], namely a nonlinear evolution \( m(t) \sim t^{1/3} \), where \( m(t) \) is the number of unzipped bases at time \( t \). In contrast, and this is the central point of our work, we find that the combined dynamics involving the excluded volume interaction between the helicase and the DNA (but no external force) leads to the uniform motion of both the helicase and the domain wall. Our results of Fig. 3 should be compared with Fig. 2 of Ref \[13\]. The effective velocity is smaller than the unhindered one and depends on the size \( l \) of the helicase, the temperature and the sequence. The \( l \) and \( T \) dependence can be estimated if we think that to move the helicase we have to unzip a base pair at the fork position, and to allow the formed kink to reach the position of the helicase. The base pair is broken with a probability \( \exp (-\frac{m}{l}) \) and the kink needs a time of order \( l \) to reach the helicase. Therefore the velocity is proportional to \( \exp (-\frac{m}{l})/l \) (confirmed by the fit of our data, see the inset of Fig 3). A strong dependence on \( T \) has also been found in Ref. \[13\]. In our simulation no sequence dependent nonuniformity in motion was ever discernible so long as the heterogeneity was uncorrelated. The velocity we observed, originating solely from the motor action, is a lower bound because any periodic conformational change of the helicase during its motion (ignored here mainly to illustrate the role of the domain wall) would assist the motion of the domain wall itself.

Our proposal lends itself to several predictions. We discuss a few here. Since the position \( s_0(t) \) of the wall is determined by the location of the stretching constraint put up by the helicase, the unzipped part beyond the helicase should not affect the action of the helicase. We verified this explicitly in our simulation: a part of the unzipped DNA beyond the helicase is chopped off (which mimics RecD activity \[12\]) at arbitrarily chosen times and there is neither any change in the nature of the wall (e.g. \( p(s, t) \)) nor in the rate of unzipping. This agrees with the observations in Refs. \[12, 13\] that the nuclease activity does not modify the unwinding action of RecBCD. Next, the width of the wall, as defined by Eq. 2 gives a length-scale for the helicase activity. The sequence randomness in a heterogeneous chain does not matter if there is no correlation beyond this scale, as found both in simulation and real experiments. We have seen a periodic mod-
Fixed at a value $P(\text{ity 1})$ when this is sterically acceptable) but also a probability affecting the forward motion of the helicase could be a phenomenological characterization of the gradual wobbliness of the helicase on the track.

In conclusion, we have shown that associating the helicase activity with the domain wall motion in a fixed-stretch ensemble accounts for several observed features, as e.g., the uniformity of unzipping, no sequence-dependent nonuniformity, and the insensitivity to the nuclease action, without any requirement of extra specific bond-cutting chemical processes. A domain wall also gives a quantitative meaning to the Y-fork in the terminology of DNA replication. The underlying thermodynamic basis gives a robustness to the mechanism that it could be at work for hexameric helicases also.

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