ABSTRACT
Type II DNA topoisomerases (topos) catalyse changes in DNA topology by passing one double-stranded DNA segment through another. This reaction is essential to processes such as replication and transcription, but carries with it the inherent danger of permanent double-strand break (DSB) formation. All type II topos hydrolyse ATP during their reactions; however, only DNA gyrase is able to harness the free energy of hydrolysis to drive DNA supercoiling, an energetically unfavourable process. A long-standing puzzle has been to understand why the majority of type II enzymes consume ATP to support reactions that do not require a net energy input. While certain type II topos are known to ‘simplify’ distributions of DNA topoisomers below thermodynamic equilibrium levels, the energy required for this process is very low, suggesting that this behaviour is not the principal reason for ATP hydrolysis. Instead, we propose that the energy of ATP hydrolysis is needed to control the separation of protein–protein interfaces and prevent the accidental formation of potentially mutagenic or cytotoxic DSBs. This interpretation has parallels with the actions of a variety of molecular machines that catalyse the conformational re-arrangement of biological macromolecules.

INTRODUCTION
Type II DNA topoisomerases (topos) perform the remarkable feat of passing one double-stranded segment of DNA through a transient break in another (1). This reaction allows these enzymes to manipulate topological properties of DNA such as supercoiling, unknotting and decatenation (unlinking of DNA circles or loops) (2). DNA topoisomerases are classified into two types, I and II, depending on whether they catalyse reactions involving the breakage of one or both strands of the DNA (3,4). All known cellular organisms have at least one type II topo, whose prototypical reaction is generally considered to be the decatenation of daughter chromosomes after DNA replication. Successful replication and partitioning of chromosomes requires all the double-helical turns linking the parental strands to be removed. This process largely occurs by relaxing positive supercoils ahead of replication forks, which can in principle be carried out by either type I or type II topos (5,6). However, any rotation of the forks or incomplete relaxation also results, after replication is complete, in links and intertwinings between the daughter chromosomes that can only normally be removed by the double-strand passage reaction of type II topos (6). Recent work has also discussed the possible physiological importance of unknotting reactions (7).

Type II topos are divided into two classes on the basis of structural and evolutionary considerations (4). Type IIA enzymes are ubiquitous in eubacteria and eukaryotes and include bacterial DNA gyrase and topo IV, as well as eukaryotic topo IIs. The IIB topos (all designated topo VI) are distant relatives with some domains in common with the IIAs (8–11); they are found in archaea, plants and a few bacteria (12,13).

All type II topoisomerases are ATPases (as part of their catalytic reaction cycle. This energetic requirement appeared natural and obvious when the first type II enzyme, DNA gyrase, was discovered in Escherichia coli (14). Gyrase uses nucleotide turnover to introduce negative supercoils into DNA, exploiting the free energy of ATP hydrolysis for the formation of thermodynamically unfavourable reaction products. In contrast, all other type II enzymes (including the type IIBs) catalyse reactions that do not have an
obvious energetic cost, such as supercoil relaxation, knotting/unknotting and catenation/decatenation. Indeed, gyrase itself can catalyse the efficient relaxation of positively supercoiled DNA in an ATP-dependent reaction (15). Why type II topoisomerases have evolved to rely on a chemical energy source to catalyse otherwise thermodynamically favourable reactions remained a puzzle for many years.

**Structure and mechanism of type II topoisomerases**

Structural and biochemical studies, particularly on gyrase and yeast topo II, have led to the formulation of a general mechanistic model for type IIA topoisomerases (Figure 1A) (3). The enzymes operate as symmetrical dimers; the eukaryotic proteins are homodimers, while the bacterial homologues divide the polypeptide into two distinct gene products and are A2B2 tetramers. Both classes of type IIA topoisomerases interact with two DNA segments. The G- (or ‘Gate’-) segment first binds to and is strongly bent by the enzyme [by as much as 150° (16–18)]. Each strand of this DNA is then cleaved by one of a pair of tyrosines, at sites 4nt apart, forming two covalent, 5'-phosphotyrosine intermediates (19). ATP binding to each monomer results in dimerization of the N-terminal domains to form a new protein–protein interface (termed the N-gate), enclosing a second DNA (the T- or ‘Transported’-segment), which is passed through the G-segment; this process requires not only DNA cleavage, but also the separation of the DNA ends by disruption of an existing protein dimer interface (the DNA gate). The T-segment subsequently leaves the complex through a third protein interface (the C- or exit-gate) (20,21), having passed through the G-segment and across the entire dimer interface of the enzyme (Figure 1A). ATP hydrolysis and product release allows the N-gate to open and resets the enzyme for further rounds of reaction, although hydrolysis of one ATP and release of phosphate also appears to stimulate strand passage (22,23).

Figure 1. Structure and mechanism of type II topoisomerases. (A) Type IIA topo core mechanism—see text for details. Different domains of the protein are indicated: yellow, ATPase domain (B subunit N-terminus in prokaryotic enzymes); orange, B subunit C-terminus (or homologous region in homodimeric eukaryotic enzymes); blue, A protein N-terminal breakage-reunion domain (or homologue). The A protein C-terminal domain is not shown. The G-segment (green), T-segment (red/pink) and DNA, N- and C-gates are indicated. Movement of the T-segment is shown from the pink to the red position. (B) Type IIB (Topo VI) mechanism. The B subunit N-terminal ATPase domain and C-terminal domain are in yellow and orange, respectively; the A subunit is in blue. The N- and DNA gates are indicated. (C) DNA gyrase structure, showing DNA wrapping around the C-terminal domains of the A subunit (cyan) to deliver a contiguous T-segment (red) to the enzyme, with a right-handed crossing over the G-segment (green). Only the wrapped segment captured by the enzyme is shown for clarity.
The specific reaction carried out by the type II topos depends on the topological relationship between the G- and the T-segment. Intra-molecular strand passage (G- and T-segments on the same circular molecule) leads either to supercoiling or relaxation, with a linking number change of ±2 (24), or to a knotting/unknotting reaction. If the G- and T-segments are on separate molecules, the result is catenation or decatenation.

This basic idea underlying this mechanism, in which the T-segment is captured by the closure of the N-terminal dimer interface, then exits through the cleaved G-segment and its associated protein interface, is known as the ‘two-gate’ model (25,26). However, since subsequent structural studies have shown that the IIA enzymes have three protein dimer interfaces in all (Figure 1A (27,28)), the second gate of the ‘two-gate’ mechanism (the ultimate exit route for the T-segment) is generally taken to be the C-gate. Roca and Wang (29) have described a theoretical alternative ‘one-gate’ scheme (1), in which the enzyme bridges the site of DNA cleavage (30,31) and strand passage of the T-segment through the G-segment into an enclosed protein cavity is followed by partial dissociation of the G-segment to allow the T-segment to leave by the same route without reversing strand passage. In Figure 1A, this mechanism would correspond to a situation where the C-gate remained closed during the reaction. Using a covalently cross-linked dimer of yeast topo II, Lindsley (32) showed that the enzyme could still carry out strand passage. Although this experiment is suggestive of a one-gate mechanism, these experiments do not rule out the two-gate scheme. To date, biochemical and structural evidence largely supports the two-gate mechanism (16,21,29,33,34) and it is generally accepted that an efficient, processive type II reaction requires an enzyme that passes a T-segment between its protein subunits as well as through a DNA gate.

Although type IIB enzymes (topo VI) share the basic structural and mechanistic features of the type IIA enzymes, they differ in some important respects (3). Topo VI is an A₂B₂ heterotrimer believed to catalyse topo reactions via a two-gate mechanism involving transient double-strand cleavage, albeit with 2 nt rather than 4 nt overhangs (35). However, structural work has shown that the enzyme possesses only two dimer interfaces (36), lacking the separate C-gate found in type IIA enzymes (Figure 1B). This configuration more closely mirrors the original two-gate model than that of the type IIA topos, but raises interesting questions as to how enzyme integrity is maintained during strand passage (see below). In this context, it may be significant that DNA cleavage by topo VI seems to be strictly dependent on ATP binding (35).

DNA gyrase—a special case

With respect to the core type IIA topo mechanism, we can understand in general terms how gyrase operates specifically to reduce linking number, both relaxing positive supercoils and maintaining a steady-state level of negative supercoiling in the presence of ATP. Gyrase is known to wrap ~130 bp of DNA around itself (Figure 1C) (37–43), presenting a T-segment to the enzyme that is immediately contiguous with the G-segment (44). The selection of a T-segment by this juxtaposition facilitates an intra-molecular reaction with the appropriate orientation for unidirectional strand passage to result in a reduction in linking number (24,44). The ability of gyrase to perform directional strand passage, i.e. to reduce linking number, is crucially dependent on the C-terminal domain of the GyrA protein, the so-called DNA-wrapping domain (Figure 1). Deletion of this domain essentially converts gyrase into a conventional (DNA-relaxing) type II topo, hence implicating wrapping in T-segment selection (45). Topo IV possesses a degenerate version of this domain (the C-terminal domain of ParC) that is unable to stabilize DNA wraps (46) and this enzyme does not supercoil DNA, but is a very efficient decatenase (47).

G-segment cleavage and DNA gate opening

Two essential and separate features of the strand-passage mechanism of type II topos are the transient, double-stranded cleavage of the G-segment and the physical opening of the DNA gate to allow the T-segment through. The cleaved-DNA species is generally believed to represent only a small fraction of the topo complexes, although many agents are known that stimulate cleavage by stabilizing this intermediate [e.g. quinolone antibiotics and antitumour agents such as etoposide (48,49), as well as ATP itself and its non-hydrolysable analogue, 5'-adenyl-β,γ-imidodiphosphate (ADPNP) (50–53)]. Experiments with several eukaryotic topo IIs have shown that the normal equilibrium level of DNA cleavage is generally less than ~1% (54,55), although Chlorella virus topo IIs appear to be exceptions (55,56). Work with E. coli gyrase suggests that the level of cleavage is also low, but measurable (57). The fraction of complexes that additionally have the DNA gate interface in a physically open state is likely to be even lower than these values, since, for almost all type IIA enzymes, strand-passage reactions do not occur in the absence of ATP (gyrase being a notable exception) (58).

Recent single-molecule studies have begun to investigate DNA gate opening further. Fluorescence resonance energy transfer experiments using Drosophila topo II and a labelled 28 bp double-stranded DNA oligo have monitored the interconversion of the topo II–DNA complex between open and closed states during ATP hydrolysis and found that the DNA gate may be open ~50% of the time (59) in the presence of ATP. In contrast, single-molecule experiments using Bacillus subtilis gyrase in which either the enzyme or DNA substrate (60 bp linear or supercoiled plasmid DNA) was labelled, found that the enzyme predominantly resides in the DNA gate-closed conformation (60) and could not detect a sizable gate-open population in the presence of ATP or ADPNP. At present, it is not clear why the Drosophila and B. subtilis studies produced different results, although it is possible that these differences could reflect either a genuine mechanistic difference between enzymes from different sources or distinctions between the
The role of ATP in strand passage

It has been assumed that the central DNA strand-passage mechanism is common to gyrase and non-supercoiling type IIA enzymes (Figure 1). Consequently, attention has focused on the role of ATP binding and hydrolysis—in driving unidirectional transport through the DNA gate and the enzyme subunits. This directionality is a clear requirement of the gyrase supercoiling reaction. It has long been known that substitution of ATP with ADPNP can result in a single-turnover (non-catalytic) strand-passage reaction (64,65). Studies with yeast topo II and ADPNP in particular have suggested that the ATP-dependent closure of the N-terminal domains may physically drive DNA cleavage (61), opening of the DNA gate and transport of the T-segment (58,63). This latter result has been taken to suggest that the cavity between the dimerized N-terminal domains may be too small to stably retain a T-segment (66,67). A recent structure of the dimerized C-terminal domain of the Mycobacterium tuberculosis gyrase B subunit has also been interpreted to suggest a possible role for the T-segment in the opening of the DNA gate, via a steric-based mechanism (62). Interestingly, the situation is somewhat different for E. coli gyrase, where there is good evidence that ATPase activity and strand passage can be uncoupled (15,68). Structural data for gyrase also suggest that a T-segment may be retained between the dimerized N-terminal domains (69), while limited amounts of DNA relaxation can occur when the enzyme is pre-incubated with ADPNP (70), suggesting that strand passage can be reversed even when the N-gate is closed.

Pre-steady-state kinetics and mutagenesis experiments with yeast topo II have indicated that the hydrolysis of one ATP and concomitant release of product (phosphate) may precede and accelerate transport of the T-segment (3,23). Structural studies of gyrase (71) and human topo IIα (67), as well as the type IIB topo, topo VI (36,72), have revealed that nucleotide binding, hydrolysis and phosphate release may transmit conformational changes to the DNA gate region through a link between the ATP binding domain and an adjacent ‘transducer’ domain, thus providing a potential physical context for the kinetic results. However, the precise interaction between these components is unclear, as no crystal structure is available for an intact gyrase B subunit or the homologous region of a non-supercoiling type IIA enzyme.

The role of ATP in topology simplification by type II topoisomerases

In 1997, Rybenkov et al. (73) described a process of topology ‘simplification’ that is carried out by non-supercoiling type IIA topoisomerases. In this process, the enzymes use the free energy of ATP hydrolysis to reduce the steady-state levels of supercoiling, knotting or catenation below those seen at thermodynamic equilibrium (73). This finding suggested that standard type IIA topoisomers might employ an energy transduction function analogous to that used in the supercoiling reaction of gyrase. The simplification process has been proposed to be physiologically important, since complete decatenation of replication products is absolutely required for chromosome partitioning during cell division. There is likewise some evidence that the removal of knots may also be an important topo function (74,75).

There has been considerable interest in the detail of the mechanism behind topology simplification, which requires both unidirectional strand passage and a selection step amongst potential T-segments to directionally perturb the equilibria (a situation again analogous to that of gyrase) (76). Topology simplification probably arises, at least in part, from bending of the G-segment by the enzyme, which helps to bias the enzyme to select T-segments from within the curved contour of the DNA (18,77–79). However, it has recently been calculated that the energy required for topology simplification is very low. Specifically, the excess free energy of the steady-state distribution of topoisomers (in the presence of a topo II and ATP) over that of the equilibrium state is <1kcal mol⁻¹ (79,80), substantially less than kT. Thus, type II topoisomerase activity is at least a thousand times less efficient at shifting chemical energy into the state of their DNA products than gyrase, which can maintain a steady-state supercoiling free energy of >500kcal mol⁻¹ (79). Moreover, some data suggest that topoisomerase activity may be less crucial physiologically (79). For example, in chromosome segregation, topo II has been shown to be responsible for decatenation (81), but partitioning forces tend to pull daughter replicons apart, perturbing the equilibrium towards complete decatenation, suggesting that there would not be a need for topology simplification (81,82). Supercoiling is thought to promote decatenation (47,83,84) and unknotting (85) and a recent report has suggested that yeast mitotic chromosomes become positively supercoiled to drive efficient decatenation (86), suggesting again that chromosome separation may be more efficient in vivo than suggested by the equilibration levels determined for dilute solutions of nicked DNAs in vitro. Furthermore, recent experiments (36,79) show that Methanosarcina mazei topo VI does not carry out topology simplification, suggesting that archaeal species can adequately unknot and partition their DNA without active simplification on the part of a topo. In summary, it is clear that while non-supercoiling type IIA topoisomerases can simplify DNA topology in an ATP-dependent manner, the energy requirements of the effect are extremely small; thus, topology simplification, if it is physiologically important, is very poorly optimized in energy transduction terms. Given that there is considerable circumstantial evidence to suggest that other factors may promote decatenation and unknotting beyond levels seen in vitro, we have considered an alternative, more fundamental role for ATP.
An ancestral role for ATP hydrolysis?

In common with other enzymes that push a complex mechanical reaction forward (87,88), it has often been stated that in type II topos ATP hydrolysis serves to ‘drive conformational changes’ (23,89,90). However, while one can imagine how nucleotide binding and turnover promote structural rearrangements to facilitate unidirectional strand passage, it is not clear whether or how the free energy of hydrolysis is being channelled to perform some type of otherwise unfavourable molecular ‘work’ to increase the free energy of the DNA product (gyrase being the exception).

Indeed, a re-consideration of the type II topo reaction cycle suggests that this line of reasoning derived from gyrase, where the enzyme must transduce energy into the DNA substrate, may obscure the real reason why evolution selected for a nucleotide-regulated enzyme. Rather than an energy transduction issue per se, we suggest that the primary ancestral requirement for ATP by the type II enzymes may derive from an evolutionary pressure to avoid aberrant double-strand breaks (DSBs). Type II topos play an essential but potentially dangerous role in the manipulation of DNA (91). The enzymes must produce a transient break in both strands of the DNA, perhaps as many as a million times per human cell cycle (92) and break re-sealing must occur reliably to prevent the accumulation of mutagenic or cytotoxic lesions. Implicit in the notion of a two-gate mechanism is the idea of protection against DSBs: a simple one-gate type II enzyme, such as that shown as a cartoon in Figure 2A, would clearly dissociate to give permanent DSBs at every strand-passage event (this is distinct from the C-shaped, one-gate model discussed above) (29,58). This issue has been alluded to recently (3,63,92), but the role of ATP hydrolysis in ensuring the prevention of DSBs has not been explicitly elaborated. We can explore these issues by imagining the hypothetical evolution of a non-ATP-dependent type II topo.

Type II topos clearly require more than one protein gate so that a protomer–protomer interaction can be maintained throughout strand passage (Figure 2B) (25,29). In the simplest such case, two gates (one of which acts as a DNA gate) would operate independently, with the passive bidirectional transfer of a T-segment through each gate in turn (Figure 2B). However, independent operation of these two gates implies the possibility of having both gates open at the same time (Figure 2B, 4), a situation that would result in a permanent DSB. To minimize the probability of this outcome, the association of the two dimer interfaces must remain very tightly associated or the G-segment must remain uncleaved. In type IIA topos, resistant to DSB formation, either the DNA gate protein interface must remain very tightly associated or the G-segment must remain uncleaved. In type IIA topos, the evidence suggests that DNA cleavage is independent of the opening of the DNA gate (94) and that the distribution between intact and broken DNA is in equilibrium in the absence of ATP, with the fraction of broken DNA being low but measurable (57,95), suggesting that the enzyme dimer is crucial to DSB resistance. On the other hand, in topo VI, there is evidence that the DNA cleavage
reaction is tightly coupled to the binding of ATP (35), indicating that the integrity of the G-segment itself may contribute to the DSB resistance of the initial complex.

For strand passage to take place, the initially tight DNA gate/G-segment interface must be made sufficiently weak for its dissociation to be frequent, while at the same time a new, strong dimer interface must be formed to maintain the stability of the complex and avoid DSB formation. This situation is essentially equivalent (in Figure 2B) to going from State 1 (where the T-segment is free to enter the enzyme cavity) to State 3 (where strand passage can take place) in a concerted and largely irreversible step, to avoid the ‘both gates open’ (DSB) and ‘both gates closed’ (no reaction) traps. Such a conformational change requires an energy input to render it unidirectional and hence must have an external driver; i.e. the new dimer interface should be essentially as tight as the initial one was. A possible conformational change is shown in Figure 3, with the process driven by the binding of a small molecule X (dyadic symmetry probably requires two such agents). Binding switches the enzyme from a form with a stable DNA gate (Figure 3, 1), to one with a labile DNA gate, with a large negative free energy change (Figure 3, 2–4; the gate does not have to be permanently open).

If the conformational change on binding molecule X happens with a T-segment in the cavity, then passive strand passage out of the complex can take place safely; in fact in the X-bound form, the T-segment can potentially pass through the G-segment in either direction (Figure 3, 2–4). However, the enzyme is now trapped in the low free energy X-bound state and the conformational change must be reversed to ensure a catalytic reaction. Thus, there must be a second step that drives the re-opening of the top gate. This process could only happen through a thermodynamic cycle (Figure 3, 4→5→1), with free energy input from a chemical change in X (to Y), with the overall free energy of the protein conformational changes being zero over the whole catalytic cycle. Thus, Figure 3 describes a model for the putative simplest viable type II topo.

The binding of X, the (overall) energetically favourable transformation of X to Y and the dissociation of Y now drive the cycle round, dissipating energy. Only in the X-bound form is the DNA gate labile enough to allow strand passage at a significant rate (Figure 3, 2–4) while the new tight dimer interface maintains the complex. Without a change in energy states from a chemical cycle, the alternation of two very strong dimer interfaces could not be accomplished. In principle, the driver for the reaction could be any chemical cycle (e.g. phosphorylation-dephosphorylation, redox change, etc.); however, nucleoside triphosphates are of course a common source of free energy in coupled reactions. We argue that the free energy of ATP hydrolysis in type II topoisomerases is driving the sequential breakage and formation of two strong dimer interfaces—the greater the energy available per cycle, the stronger the dimer interfaces can be and the safer the enzyme is from the formation of accidental DSBs. The free energy needed to modulate protein–protein interactions at the DNA gate and ‘stabilization gate’ interfaces should, according to this model, be a significant fraction of the free energy available from the hydrolysis of two ATP molecules. Indeed, a preliminary estimate of the interface free energies (△G) for the DNA gates of various type II topoisomerases using the program PISA (96) yields values of around –20 to

![Figure 3](https://academic.oup.com/nar/article-abstract/39/15/6327/1018983/6332)
−70 kJ mol⁻¹ (Lawson, D.M. and A.M., unpublished data), compared with ∼100–150 kJ mol⁻¹ available from the hydrolysis of two ATPs (79), consistent with a potential role for ATP hydrolysis in gate opening. However, it is important to point out that simple dissociation of the DNA gate is unlikely to be the only structural alteration involved. In addition, these calculations are preliminary and based on a number of assumptions and should hence be treated with caution.

It is important to note that in this simplest model case, the T-segment can be transported passively in either direction. For example, in Figure 3, the T-segment could enter at 3 and be carried through 4 and 5 to be released at 1, in a ‘bottom-up’ reaction. This possibility means the enzyme would then behave as a true ‘phantom-chain’ device (97,98), whilst still requiring ATP hydrolysis. In this hypothetical ancestral scheme, the ATP cycle serves (and is required) only to protect against DSBs. However, if we allow the presence of a T-segment to differentially affect the rate of binding of X (ATP) or the rate of dissociation of Y (ADP), then directionality can be introduced into the strand-passage reaction, even though the enzyme would still only equilibrate topoisomers. Further elaborations might include the conformational changes dependent on ATP hydrolysis that have been hypothesized to drive the T-segment unidirectionally through the protein complex (see above) (3). A gyrase or a topology-simplifying enzyme requires some mechanism for T-segment selection in addition to preferential strand passage in one direction.

Overall, the hypothetical scheme in Figure 3 is essentially a representation of the canonical two-gate reaction mechanism proposed for type II topoisomerases (Figure 1) and closely matches the proposed mechanism for topo VI (Figure 1B) (36). Type IIA enzymes have an additional protein gate, the C-gate, which may function to increase the security of the DNA gate and further reduce the possibility of DSB formation (63,92). In the presence of the non-hydrolysable analogue ADPNP, only single-turnover strand passage occurs (65); this process is effectively equivalent to the reaction scheme shown in Figure 3, 1–4 only.

As noted above, the addition of ATP or ADPNP can actually promote DNA cleavage, apparently counter to our argument that ATP hydrolysis prevents DSBs. However, this effect is only revealed in experiments involving the destructive denaturation of the whole complex, either in the presence or absence of a drug (53,99,100). This simply means that the addition of nucleotide changes the cleavage–religation equilibrium to produce more cleaved DNA, as would be expected with this model. Our main point is that this inevitable and necessary cleavage of the G-segment and weakening of the DNA gate do not translate into permanent DSBs because the newly closed N-gate maintains the integrity of the overall complex. It is interesting to note that the binding of nucleotide really does ‘form a new tight protein interface’. The ADPNP molecules in structures of the dimerized N-terminal domain actually bridge the gap between the subunits, making substantial contacts with both monomer proteins (101).

Our argument represents a shift of perspective from the idea that the closing of the ATP-operated clamp simply ‘captures the T-segment’, to the concept that it forms a tight new interface, while concomitantly weakening the DNA gate, to allow strand passage while preventing complete dissociation. ‘Capture’ of a T-segment is not specifically required; in the hypothetical ‘bottom-up’ reaction described above (see Figure 3), the binding of X does not capture anything, but sets up the enzyme to allow the T-segment to pass through the G-segment from below and escape from the top, after dissociation of Y. Indeed, it remains a formal possibility that topo VI might be able to work via a ‘bottom-up’ reaction; this idea has not been specifically tested.

The evolution of type II topoisomerases

This rationale also suggests that the simplest successful type II topoisomers in evolutionary terms might have looked something like topo VI, a two-gate type IIB. This enzyme would have been required as soon as genomes evolved to be long enough for tangling or formal linking of the daughter helices to become a problem, but probably before any requirement for active supercoiling or topology simplification. The additional C-gate of the type IIA topoisomerase could have evolved later, possibly for additional resistance to DSBs [although a recent discussion of topo phylogenomics has suggested a more complex picture, in which the enzymes may have evolved in an ancient virosphere, with their modern distribution dependent on a number of horizontal gene transfer events (13)]. The ATP-operated clamp domain of the type II topoisomerases is not unique, but is a member of the GHKL ATPase family (102) (see below), so its ultimate evolutionary origin may lie in another system. It remains an outstanding question as to whether gyrases or non-superciling type IIA enzymes are more primitive. There is evidence that topo IVs are derived from gyrase by loss of the DNA wrap through modification of the C-terminal wrapping domain (13,46,103); however, whether gyrase is also ancestral to the eukaryotic type IIA topoisomerases, or vice versa, is not clear (103). A circumstantial argument can be made either way. If gyrase is the ancestral form, then the G-segment bend and unidirectional strand passage that leads to topology simplification by the non-superciling enzymes could be a relic of the directional gyrase reaction (79). On the other hand, Roca (92) has recently suggested that the G-segment bend might have a role as a sensor of tension in DNA and that an inability to correctly bend DNA could prevent the cleavage reaction under conditions where DNA tension might increase the probability of DSB formation during enzyme action. In the latter instance, the gyrase wrap could have developed from the pre-existing G-segment tension sensor. Interestingly, recent single-molecule experiments support the idea of DNA wrapping by gyrase being sensitive to tension in the DNA (104); however, in some cases, notably at the onset of chromosome partition in the presence of residual catenates, DNA tension might be thought, a priori, to be a potentially useful trigger of topo action leading to decatenation, rather than an inhibitor of it.
The ATP-independent relaxation reaction of gyrase

Although almost all of the reactions catalysed by type II topois are dependent upon ATP hydrolysis, one activity of gyrase is a notable exception. In bacterial cells, the primary in vivo roles of this enzyme are thought to be the ATP-dependent relaxation of positive supercoils and introduction of negative supercoils (105). However, it has been established that gyrase can also carry out the ATP-independent relaxation of negative supercoils (99,100). This reaction (for E. coli gyrase) is ~20-fold slower than ATP-dependent supercoiling and ATP-free gyrase does not relax supercoils processively. Whether ATP-independent relaxation is physiologically important is not known, although this seems unlikely; recent estimates of ATP concentrations in E. coli are sufficiently high to largely saturate the enzyme (106). Nevertheless, the reaction represents a significant distinction between gyrase and non-superciling type II enzymes. It has been reported that phage T4 topo II can also carry out non-catalytic strand-passage events in the absence of nucleotide, but this result has not been investigated further (107).

The gyrase relaxation reaction is an obvious counter argument to the idea that ATP hydrolysis is required to avoid DSBs, as it demonstrates that the gyrase DNA gate must be able to open in the absence of a new dimer interface created by ATP binding. However, gyrase is unique amongst type II topois and this exception might be consistent with our hypothesis. In particular, the enzyme can transduce a significant fraction of the free energy of ATP hydrolysis into DNA supercoiling free energy (108–111), suggesting that less free energy may be available to drive the conformational changes necessary for DSB prevention in the enzyme. This, in turn, suggests that the gyrase DNA- and C-gates should be intrinsically more labile, thus allowing slow relaxation, as observed. However, our preliminary consideration of interface binding energies does not do so far support this; we did not find any consistent differences in the free energies associated with DNA gyrase and those for other type II topois. These considerations imply that gyrase ought to be more susceptible to forming DSBs than other type II topois, although the unique DNA wrap around the gyrase complex may serve to stabilize the complex and mitigate this effect.

The role of type II topoisomerases in illegitimate recombination

Illegitimate recombination (IR) is a rearrangement of DNA that occurs between nucleic acid segments that bear short regions of homology (<10 bp) or no apparent homology at all (112). It has been appreciated for many years that gyrase can participate in IR reactions (113) and that this reaction is stimulated (2–3 orders of magnitude) by the quinolone drug oxolinic acid (114), which binds at the DNA gate. In contrast, this stimulation is blocked by the coumarin drug coumermycin A1, which prevents ATP binding (115). Gyrase-mediated IR can occur in vitro and requires the presence of an E. coli extract, but is independent of RecA (114). One proposed mechanism for this reaction is that IR can take place when a pair of covalently bound gyrases exchange GyrA subunits (114). Stimulation by oxolinic acid is thought to occur because the drug increases the lifetime of the DNA-cleaved state, enhancing the likelihood that subunit swapping can occur. Support for this model comes from the identification of temperature-sensitive gyrA mutants in E. coli that confer spontaneous illegitimate recombination (116). Of these substitutions, one mutant in particular (Leu<sup>492</sup>→Pro) displayed normal supercoiling activity, but generated linear DNA during the reaction, suggesting that it has a defect at the DNA rejoining step or in its subunit interactions (116).

IR has also been shown to occur in other type II topois, including T4 and eukaryotic topo II. In the case of the T4 enzyme, it appears that recombination can occur with the purified enzyme alone [i.e. without an added E. coli extract (117–119)]. Purified calf thymus topo II can also mediate IR in a reaction that can be inhibited by the coumarin drug novobiocin (120), whereas in yeast, IR is stimulated by the topo II poison etoposide (VP-16) (121). Subunit exchange is also the suggested mechanism for IR with these type II enzymes, implying that subunit–subunit interactions can be disrupted under certain conditions. Taken together, several lines of data indicate that the aberrant disruption of type II topo subunit interfaces may be a source of illegitimate recombination. If true, these reactions highlight the detrimental consequences to the cell when the security normally afforded by the various topo gates break down.

Spo11 and meiotic recombination

Spo11 is a homologue of the topo VIA subunit (9,122), the dimer of which forms the DNA gate of the type IIB enzymes (11,36) (Figure 1B). Spo11 is present in all eukaryotes, where it is responsible for the initiation of DSBs in meiotic recombination (122,123). Roca (92) has proposed that the relative insecurity of topo VI as a true two-gate (rather than three-gate) enzyme makes the A subunit an appropriate candidate for co-option into a system evolved to produce DSBs. On the other hand, Forterre and Gadelle (13) have suggested that the opposite may be true and Spo11 may have been recruited by the topo VIB subunit to make a functional topo.

To date, there has been no identification of a topo VIB ATPase homologue (or analogue) in the meiotic recombination apparatus. At first glance, our hypothesis suggests that such a component should be necessary to open the protein interface responsible for DSB formation. However, studies of meiotic DSB formation (124) suggest that free 5'-ends are produced after Spo11 binding by nucleolytic cleavage of the covalently bound strands at each side of the Spo11–DNA complex. These data can easily be accommodated by a variation of our model in which separation of Spo11 monomers is not required to generate free DNA ends, but rather binding of Spo11 marks the DNA for homologous recombination repair proteins by generating a covalent protein–DNA link, although it is not yet clear what induces Spo11 to cleave the DNA. Consistent with this line of reasoning, it is known that many DSB repair factors, such as Rad50 and Mre11, are recruited to sites of Spo11 action (125).
Alternatively, it may be possible that some as yet undiscovered ‘subunit separation’ component acts on Spo11 in meiotic recombination.

The role of nucleotide hydrolysis in disrupting protein interfaces in other systems

If the primary role of ATP in type II topois is to promote a particular type of mechanical manipulation, DNA gate opening, through the disruption of protein interfaces, then we might expect to see evidence of a similar role for nucleotides in other systems. Although many examples could be chosen, we limit a brief discussion here to other DNA-remodelling systems.

One parallel may be with the Structural Maintenance of Chromosomes (SMC) ATPase superfamily, which are key mediators of chromosome cohesion, condensation and repair in many organisms (126,127). Most bacteria possess a complex involving a dimer of SMC subunits, and a pair of accessory proteins known as ScpA and ScpB (128); γ-proteobacteria possess a diverged form of this assembly comprising the Smc homologue, MukB and two interacting subunits, MukE and MukF (129,130). MukE and MukF form an elongated, dimeric particle, which allows MukF’s C-terminal domains to engage the nucleotide binding ‘head’ regions of MukB and generate either closed ring-like structures or repetitive arrays (131,132); this general organization between SMCs and their accessory subunits is thought to be preserved across different SMC systems (126,127). Nucleotide binding further stabilizes the association of the MukB head domains (a feature shared with other Smc proteins and the ABC ATPase superfamily in general (133–135)) and it has been proposed that disruption of the MukB–MukF interaction depends upon ATP, as well as DNA (131,136). Some studies have even suggested that the ‘hinge’ region of Smc proteins, which lies distal to the ATPase domains, may function as a second regulatable gate to create transient openings within the molecule (137,138).

Another example of ATP regulation of protein–protein interactions occurs in DNA mismatch repair (MMR) (139). In E. coli, three proteins are responsible for the initiation of MMR-directed events: MutS, MutL and MutH. In the case of MutS, an ABC ATPase like the SMCs, it has been shown that DNA-binding domains N- and C-terminal to the nucleotide binding site become predisposed to associate tightly with DNA in the presence of ATP and to loosen their grip following ATP hydrolysis (140,141). For its part, MutL is a GHKL ATPase (102), a protein superfamily that includes the ATPase region of type II topois. The MutL dimer has been proposed to undergo ATP-dependent association of its ATPase domains (142,143), an event that serves to control the endonuclease action of MutH. Interestingly, in many bacteria apart from E. coli, as well as in eukaryotes, the domain C-terminal to the MutL ATPase region is in fact an endonuclease whose activity is regulated by nucleotide turnover (144,145). This arrangement and conformational cycle has striking parallels to that seen for the catalytic modules of type II topois, topo VI in particular.

A third example of the role of ATP hydrolysis in DNA remodelling through the formation and resolution of protein–protein interactions occurs with certain enzymes belonging to the AAA+ (ATPases associated with various cellular activities) superfamily, specifically, bacterial enhancer binding proteins (bEBPs) and replication initiation factors such as DnaA and the Origin Recognition Complex (ORC) (146,147)). The bEBPs such as NtrC and PspF initially bind DNA as dimers in an ‘off’ state, but upon phosphorylation via a two-component response regulator system, rearrange into ring-shaped oligomers that loop DNA and engage RNA polymerase/σ54 complexes at adjacent promoters (148,149). Assembly is mediated by the AAA+ ATPase subunits, which further remodel σ54 in a nucleotide-dependent manner to allow the polymerase to clear the promoter. For its part, DnaA binds to replication origins as a monomer, but at elevated concentrations undergoes an ATP-dependent transition to assemble into a helical nucleoprotein complex that melts DNA and assists replisome formation (150–153).

CONCLUDING REMARKS

The type II topois are well-studied molecular machines and many details of their DNA double-strand-passage mechanisms have been elucidated using a combination of structural and biochemical studies. However, their absolute requirement for ATP hydrolysis for their primary reactions has been puzzling. Although it seems that gyrase can transduce much of the free energy of ATP hydrolysis into DNA supercoiling, the non-supercoiling enzymes channel only a tiny fraction of this energy into their substrates. The hypothesis proposed here—that free energy is required to manipulate protein–protein interactions to maintain a high degree of enzyme dimer stability during the strand-passage reaction—explains how these enzymes are able to avoid the formation of permanent DSBs during their reactions. The proposal also provides a framework for explaining a number of diverse features of these enzymes, including the nucleotide-independent relaxation reaction of gyrase, the participation of type II enzymes in illegitimate recombination and the relationship between the meiotic recombination protein Spo11 and the type IIB topois. Further experiments are needed to directly investigate this proposed relationship between ATP hydrolysis and complex stability and the extent to which these ATP-dependent control mechanisms are shared by other DNA-remodelling systems, as well as non-DNA systems such as ATP-dependent transporters and heat-shock proteins.

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REFERENCES

1. Wang,J.C. (1998) Moving one DNA double helix through another by a type II DNA topoisomerase: the story of a simple molecular machine. Q. Rev. Biophys., 31, 107–144.

2. Bates,A.D. and Maxwell,A. (2005) DNA Topology. Oxford University Press, Oxford.

3. Schöfferl,A.J. and Berger,J.M. (2008) DNA topoisomerases: harnessing and constraining energy to govern chromosome topology. Q. Rev. Biophys., 41, 41–101.

4. Wang,J.C. (1996) DNA topoisomerases. Annu. Rev. Biochem., 65, 635–692.

5. Postov,L., Crisona,N.J., Peter,B.J., Hardy,C.D. and Cozzarelli,N.R. (2001) Topological challenges to DNA replication: conformations at the fork. Proc. Natl Acad. Sci. USA, 98, 8219–8226.

6. Wang,J.C. (2002) Cellular roles of DNA topoisomerases: a molecular perspective. Nat. Rev. Mol. Cell Biol., 3, 430–440.

7. Liu,Z., Deibler,R.W., Chan,H.S. and Zechiedrich,L. (2009) The phylogenomics of type II DNA topoisomerases. Nucleic Acids Res., 37, 661–671.

8. Aravind,L., Deininger,P., Hardy,C.D. and Tewari,A. (2002) Topoisomerase VI-B subunit: implications for type II topoisomerases. Nucleic Acids Res., 30, 4205–4213.

9. Bergerat,A., de Massy,B., Gadelle,D., Varoutas,P.C., Nicolas,A. and Forterre,P. (1997) An atypical topoisomerase II from Archaea with implications for meiotic recombination. Nature, 386, 414–417.

10. Corbett,K.D. and Berger,J.M. (2003) Structure of the topoisomerase VI-B subunit: implications for type II topoisomerase mechanism and evolution. EMBO J., 22, 151–163.

11. Nichols,M.D., DeAngelis,K., Keck,J.L. and Berger,J.M. (1999) Structure and function of an archaean topoisomerase VI subunit with homology to the meiotic recombination factor Spo11. EMBO J., 18, 6177–6188.

12. Gadelle,D., Filee,J., Buhtler,C. and Forterre,P. (2003) Phylogenomics of type II DNA topoisomerases. Bioessays, 25, 232–242.

13. Forterre,P. and Gadelle,D. (2009) Phylogenomics of DNA topoisomerases: their origin and putative roles in the emergence of modern organisms. Nucleic Acids Res., 37, 679–692.

14. Gellert,M., Mizuuchi,K., O'Dea,M.H. and Nash,H.A. (1976) DNA gyrase: an enzyme that introduces superhelical turns into DNA. Proc. Natl Acad. Sci. USA, 73, 3872–3876.

15. Bates,A.D., O'Dea,M.H. and Gellert,M. (1996) Energy coupling in Escherichia coli DNA gyrase: the relationship between nucleotide binding, strand passage, and DNA supercoiling. Biochemistry, 35, 1408–1416.

16. Dong,K.C. and Berger,J.M. (2007) Structural basis for gate-DNA recognition and bending by type IIA topoisomerases. Nature, 450, 1201–1205.

17. Lapontogov,I., Sohi,M.K., Veselkov,D.A., Pan,X.-S., Sawhney,R., Thompson,A.W., McAuley,K.E., Fisher,L.M. and Sanderson,M.R. (2009) Structural insight into the quinolone-DNA cleavage complex of type IIA topoisomerases. Nat. Struct. Mol. Biol., 16, 667–669.

18. Vologodskii,A.V., Zhang,W., Rybenkov,V.V., Podtelezhnikov,A.A., Subramanian,D., Griffith,J.D. and Cozzarelli,N.R. (2001) Mechanism of topology simplification by type II DNA topoisomerases. Proc. Natl Acad. Sci. USA, 98, 3045–3049.

19. Sander,M. and Hsieh,T. (1983) Double strand DNA cleavage by type II DNA topoisomerase from Drosophila melanogaster. J. Biol. Chem., 258, 8421–8428.

20. Roca,J., Berger,J.M., Harrison,S.C. and Wang,J.C. (1996) DNA transport by a type II topoisomerase: direct evidence for a two-gate mechanism. Proc. Natl Acad. Sci. USA, 93, 4057–4062.

21. Williams,N.L. and Maxwell,A. (1999) Probing the two-gate mechanism of DNA gyrase using cysteine cross-linking. Biochemistry, 38, 13502–13511.

22. Baird,C.L., Gordon,M.S., Andreason,D.M., Marecek,J.F. and Lindsley,J.E. (2001) The ATPase reaction cycle of yeast DNA topoisomerase II. Slow rates of ATP resynthesis and Pi release. J. Biol. Chem., 276, 27893–27898.

23. Baird,C.L., Harkins,T.T., Morris,S.K. and Lindsley,J.E. (1999) Topoisomerase II drives DNA transport by hydrolyzing one ATP. Proc. Natl Acad. Sci. USA, 96, 13685–13690.

24. Brown,P.O. and Cozzarelli,N.R. (1979) A sign inversion mechanism for enzymatic supercoiling of DNA. Science, 206, 1081–1083.

25. Mizuuchi,K., Fisher,M., O'Dea,M. and Gellert,M. (1980) DNA gyrase action involves the introduction of transient double-strand breaks into DNA. Proc. Natl Acad. Sci. USA, 77, 1847–1851.

26. Wang,J.C., Gumpert,R.L., Javaherian,K., Kirkegaard,K., Klevan,L., Kotezlu,M.L. and Tse,Y.-C. (2005) In Alberts,B. and Fox,C.F. (eds), Mechanistic Studies of DNA Replication and Genetic Recombination. Academic Press, NY, pp. 769–784.

27. Berger,J.M., Gamblin,S.J., Harrison,S.C. and Wang,J.C. (1996) Structure and mechanism of DNA topoisomerase II. Nature, 379, 225–232.

28. Morais Cabral,J.H., Jackson,A.P., Smith,C.V., Shikotra,N., Maxwell,A. and Liddington,R.C. (1997) Crystal structure of the breakage-reunion domain of DNA gyrase. Nature, 388, 903–906.

29. Roca,J. and Wang,J.C. (1994) DNA transport by a type II topoisomerase: evidence in favor of a two-gate mechanism. Cell, 77, 609–616.

30. Brown,P.O. and Cozzarelli,N.R. (1981) Catenation and knotting of duplex DNA by type I topoisomerases: A mechanistic parallel with type II topoisomerases. Proc. Natl Acad. Sci. USA, 78, 843–847.

31. Tse,Y.-C. and Wang,J.C. (1980) E. coli and M. luteus DNA topoisomerase I can catalyse catenation or decatenation of double-stranded DNA rings. Cell, 22, 269–276.

32. Sanderson,M.R. (1996) Intradianly tethered DNA topoisomerase II is catalytically active in DNA transport. Proc. Natl Acad. Sci. USA, 93, 2975–2980.

33. Roca,J., Berger,J.M., Harrison,S.C. and Wang,J.C. (1996) DNA transport by a type II topoisomerase: direct evidence for a two-gate mechanism. Proc. Natl Acad. Sci. USA, 93, 4057–4062.

34. Roca,J. and Wang,J.C. (1992) The capture of a DNA double helix by an ATP-dependent protein clamp: a key step in DNA transport by type II DNA topoisomerases. Cell, 71, 833–840.

35. Buhtler,C., Lebbink,J.H., Bosc,C., Ladenstein,R. and Forterre,P. (2001) DNA topoisomerase VI generates ATP-dependent double-strand breaks with two-nucleotide overhangs. J. Biol. Chem., 276, 37215–37222.

36. Corbett,K.D., Benedetti,P. and Berger,J.M. (2007) Hollenzyme assembly and ATP-mediated conformational dynamics of topoisomerase VI. Nat. Struct. Mol. Biol., 14, 611–619.

37. Fisher,L.M., Mizuuchi,K., O'Dea,M.H., Ohmori,H. and Gellert,M. (1981) Site-specific interaction of DNA gyrase with DNA. Proc. Natl Acad. Sci. USA, 78, 4165–4169.

38. Kirkegaard,K. and Wang,J.C. (1981) Mapping the topography of DNA wrapped around gyrase by nucleolytic and chemical probing of complexes of unique DNA sequences. Cell, 23, 721–729.

39. Liu,L. and Wang,J. (1978) Microccoccus luteus DNA gyrase: Active components and a model for its supercoiling of DNA. Proc. Natl Acad. Sci. USA, 75, 2098–2102.

40. Liu,L. and Wang,J.C. (1978) DNA-DNA gyrase complex: the wrapping of the DNA duplex outside the enzyme. Cell, 15, 979–984.

41. Morrison,A. and Cozzarelli,N.R. (1981) Contacts between DNA gyrase and its binding site on DNA: features of symmetry and
55. Fortune, J.M., Lavrukhin, O.V., Gurnon, J.R., Van Etten, J.L.,
54. Fortune, J.M., Dickey, J.S., Lavrukhin, O.V., Van Etten, J.L.,
53. Sander, M. and Hsieh, T.R. (1983) Double strand DNA cleavage by
51. Kampranis, S.C., Bates, A.D. and Maxwell, A. (1999) A model for
52. Morrison, A., Higgins, N.P. and Cozzarelli, N.R. (1980) Interaction
48. Deweese, J.E. and Osheroff, N. (2009) The DNA cleavage reaction
45. Kampranis, S.C. and Maxwell, A. (1996) Conversion of DNA
46. Corbett, K.D., Schoeffler, A.J., Thomsen, N.D. and Berger, J.M.
44. Heddle, J.G., Mitelheiser, S., Maxwell, A. and Thomson, N.H. (2004)
43. Wang, J.C. (1998) Moving one DNA double helix through
42. Orphanides, G. and Maxwell, A. (1994) Evidence for a
41. Gubaev, A., Hilbert, M. and Klostermeier, D. (2009) The DNA-gate
62. Fu, G., Wu, J., Liu, W., Zhu, D., Hu, Y., Deng, J., Zhang, X.E., Bi, L.
61. Corbett, A.H., Zechiedrich, E.L. and Osheroff, N. (1992) A role for
59. Wang, J.C. (1998) Moving one DNA double helix through
58. Wang, J.C. (1998) Moving one DNA double helix through
57. Sander, M. and Hsieh, T.R. (1983) Double strand DNA cleavage by
56. Classen, S., Olland, S. and Berger, J.M. (2003) Structure of the
55. Fortune, J.M., Lavrukhin, O.V., Gurnon, J.R., Van Etten, J.L.,
54. Fortune, J.M., Dickey, J.S., Lavrukhin, O.V., Van Etten, J.L.,
53. Sander, M. and Hsieh, T.R. (1983) Double strand DNA cleavage by
6338 Nucleic Acids Research, 2011, Vol. 39, No. 15

86. Baxter, J., Sen, N., Martinez, V.L., De Carandini, M.E., Schwartzman, J.B., Difley, J.F. and Aragon, J. (2011) Positive supercoiling of motic DNA drives decatenation by topoisomerase II in eukaryotes. Science, 331, 1328–1332.

87. Delagoutte, E. and von Hippel, P.H. (2003) Helicase mechanisms and the coupling of helicases within macromolecular machines. Part II: Integration of helicases into cellular processes. Q. Rev. Biophys., 36, 1–69.

88. Muller, U.F. (2006) Re-creating an RNA world. Science, 311, 1861–1866.

89. Wang, J.C. (2002) Cellular roles of DNA topoisomerases: a double-edged sword? Cell, 62, 403–406.

90. Roca, J. (2009) Topoisomerase II: a fitted mechanism for the chromatin landscape. Nucleic Acids Res., 37, 721–730.

91. Kouzine, F., Sanford, S., Elisha-Feil, Z. and Levens, D. (2008) The role of DNA topoisomerases in recombination and genome stability: a double-edged sword? Cell, 62, 403–406.

92. Westerhoff, H.V., O’Dea, M.H., Maxwell, A. and Gellert, M. (1988) DNA supercoiling by DNA gyrase. A static head analysis. Cell Biophys., 12, 15–18.

93. Wilstermann, A.M. and Osheroff, N. (2003) Stabilization of eukaryotic topoisomerase II-DNA cleavage complexes. Curr. Top. Med. Chem., 3, 321–338.

94. Wilstermann, A.M. and Osheroff, N. (2003) Stabilization of eukaryotic topoisomerase II-DNA cleavage complexes. Curr. Top. Med. Chem., 3, 321–338.

95. Kouzine, F., Sanford, S., Elisha-Feil, Z. and Levens, D. (2008) The role of DNA topoisomerases in recombination and genome stability: a double-edged sword? Cell, 62, 403–406.
architecture and functional relationship to kleisins. *Embo J.*, 24, 1921–1930.

133. Hopfner,K.P., Karcher,A., Shin,D.S., Craig,L., Arthur,L.M., Carney,J.P. and Tainer,J.A. (2000) Structural biology of Rad50 ATPase: ATP-driven conformational control in DNA double-strand break repair and the ABC-ATPase superfamily. *Cell*, 101, 789–800.

134. Thomsen,N.D. and Berger,J.M. (2008) Structural frameworks for considering microbial protein- and nucleic acid-dependent motor ATPases. *Mol. Microbiol.*, 69, 1071–1090.

135. Haering,C.H., Schoffnegger,D., Nishino,T., Helmhart,W., Nasmyth,K. and Lowe,J. (2004) Structure and stability of cohesin’s Smc1-kleisin interaction. *Mol. Cell*, 15, 951–964.

136. She,W., Wang,Q., Mordukhova,E.A. and Rybenkov,V.V. (2007) MukEF Is required for stable association of MukB with the chromosome. *J. Bacteriol.*, 189, 7062–7068.

137. Gruber,S., Arumugam,P., Katou,Y., Kuglitsch,D., Helmhart,W., Shirahige,K. and Nasmyth,K. (2006) Evidence that loading of cohesin onto chromosomes involves opening of its SMC hinge. *Cell*, 127, 523–537.

138. Moreno-Herrero,F., de Jager,M., Dekker,N.H., Kanaar,R., Wyman,C. and Dekker,C. (2005) Mesoscale conformational changes in the DNA-repair complex Rad50/Mre11/Nbs1 upon binding DNA. *Nature*, 437, 440–443.

139. Kunkel,T.A. and Erie,D.A. (2005) DNA mismatch repair. *Annu. Rev. Biochem.*, 74, 681–710.

140. Lamers,M.H., Perrakis,A., Enzlin,J.H., Winterwerp,H.H., de Wind,N. and Sixma,T.K. (2000) The crystal structure of DNA mismatch repair protein MutS binding to a G x T mismatch. *Nature*, 407, 711–717.

141. Obmolova,G., Ban,C., Hsieh,P. and Yang,W. (2000) Crystal structures of mismatch repair protein MutS and its complex with a substrate DNA. *Nature*, 407, 703–710.

142. Ban,C. and Yang,W. (1999) Transformation of MutL by ATP binding and hydrolysis: a switch in DNA mismatch repair. *Cell*, 97, 85–97.

143. Ban,C. and Yang,W. (1998) Crystal structure and ATPase activity of MutL: implications for DNA repair and mutagenesis. *Cell*, 95, 541–552.

144. Kadyrov,F.A., Holmes,S.F., Arana,M.E., Lukianova,O.A., O’Donnell,M., Kunel,T.A. and Modrich,P. (2007) *Saccharomyces cerevisiae* MutLalpha is a mismatch repair endonuclease. *J. Biol. Chem.*, 282, 37181–37190.

145. Kadyrov,F.A., Dzantiev,L., Constantin,N. and Modrich,P. (2006) Endonucleolytic function of MutLalpha in human mismatch repair. *Cell*, 126, 297–308.

146. Bose,D., Joly,N., Pape,T., Rappas,M., Schumacher,J., Buck,M. and Zhang,X. (2008) Dissecting the ATP hydrolysis pathway of bacterial enhancer-binding proteins. *Biochem. Soc. Trans.*, 36, 83–88.

147. Erzberger,J.P. and Berger,J.M. (2006) Evolutionary relationships and structural mechanisms of AAA+ proteins. *Annu. Rev. Biophys. Biomol. Struct.*, 35, 93–114.

148. Schumacher,J., Joly,N., Rappas,M., Zhang,X. and Buck,M. (2006) Structures and organisation of AAA+ enhancer binding proteins in transcriptional activation. *J. Struct. Biol.*, 156, 190–199.

149. Chen,B., Sysoeva,T.A., Chowdhury,S. and Nixon,B.T. (2008) Regulation and action of the bacterial enhancer-binding protein AAA+ domains. *Biochem. Soc. Trans.*, 36, 89–93.

150. Bramhill,D. and Kornberg,A. (1988) A model for initiation at origins of DNA replication. *Cell*, 54, 915–918.

151. Erzberger,J.P., Mott,M.L. and Berger,J.M. (2006) Structural basis for ATP-dependent DnaA assembly and replication-origin remodeling. *Nat. Struct. Mol. Biol.*, 13, 676–683.

152. Margulies,C. and Kaguni,J.M. (1996) Ordered and sequential binding of DnaA protein to oriC, the chromosomal origin of *Escherichia coli*. *J. Biol. Chem.*, 271, 17035–17040.

153. Leonard,A.C. and Grimwade,J.E. (2005) Building a bacterial orisome: emergence of new regulatory features for replication origin unwinding. *Mol. Microbiol.*, 55, 978–985.