Nucleotide-dependent Domain Movement in the ATPase Domain of a Human Type IIA DNA Topoisomerase

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Type IIA DNA topoisomerases (Topo IIs) are complex multifunctional enzymes that use the energy of ATP hydrolysis to resolve topological problems encountered in the genome during cell growth and division (reviewed in Refs. 1–3). The functions served by Topo II enzymes are essential for cell viability; hence these evolutionarily conserved enzymes have been widely exploited as targets for clinically important antibiotics and anti-tumor drugs (4). A broad range of biochemical and structural studies has provided a detailed understanding of the molecular mechanism employed by Topo II enzymes, including those of eukaryotic origin (1–3). Eukaryotic Topo II is a homodimer having two functionally distinct domains per subunit, an N-terminal ATPase domain, and a C-terminal DNA cleavage-religation domain. Structures of these individual functional domains from Saccharomyces cerevisiae have been solved (5–7), leading to a composite picture of Topo II as a theta-shaped dimer having three major dimer interfaces, one at the N terminus, one in the middle, and one at the C terminus. Each of the three interfaces constitutes a gate allowing capture and releasing passage of duplex DNA. In the absence of DNA and ATP, the N-terminal gate (N-gate) is open, and the middle (M-gate) and C-terminal (C-gate) gates are closed. According to the two-gate model (8), the Topo II catalytic cycle is initiated by the binding of the G-segment to the middle gate. Binding of ATP then triggers the closing of the N-gate via establishment of a dimer interface in the ATPase domain and captures the T-segment when present. Closing of the N-gate promotes the cleavage of the G segment and opening of the M-gate. The T segment then passes through the M-gate, after which the M-gate closes and the C-gate opens to allow release of the passed T-segment. Finally, the C-gate closes, and the N-gate opens to prepare for another round of strand passage (9). Following multiple rounds of T-segment passage, the G-segment is released from Topo II.

One of the central mysteries surrounding Topo II structure and function concerns the mechanisms through which nucleotide binding, hydrolysis, and release are transduced into gate opening and closure. To gain insight into this question, we solved the crystal structure of the ATPase domain of human Topo IIa (HT2ATPase, residues 29–428) bound to either AMP-PNP (a non-hydrolyzable ATP analog) or ADP to 1.86 and 2.51 Å, respectively. Comparison of these two structures revealed rigid-body motions of structural modules within the ATPase domain consistent with the opening and closing of a molecular gate.

### EXPERIMENTAL PROCEDURES

**Plasmid Construction**—The gene of HT2ATPase (residues 29–428) was amplified from YEpWOB6 (10) by PCR using two primers, 5’-CCGGCGCGCCGCGCGACGTCGAGAACCTGACCTCTCAGTTCGTGTGAAAGATCTATCATAAGAAAAAC-3’ and 5’-GCCGCGCGCGCGCGCTCAGTTTTATGAAACACTCTTGTATTACCTGAGC-3’. The DNA sequence corresponding to tobacco etch virus protease recognition sequence is italic. To make a plasmid that has both His and MBP tags, the MBP encoding sequence was excised from pIADL 16 vector with NcoI (New England Biolabs) and XhoI (New England Biolabs) restriction enzymes and ligated with pET 30a vector (Novagen) treated with the same restriction enzymes to produce the pSB vector. Both the PCR product and the pSB vector were then digested with SacI and Xhol and ligated to each other to produce the pSB23 vector, which encodes a fusion protein that has an N-terminal His tag followed by MBP and HT2ATPase with a tobacco etch virus protease cleavage site in between.

**Expression and Purification**—Rosetta(DE3)pLysS (Novagen) was transformed with pSB23. Cultures were grown in LB media at 37 °C. When $A_{600}$ of 0.6–0.8 was achieved, isopropyl-1-thio-β-D-galactopyranoside was added to a final concentration of 1 mM. After further incubation for 10 h at 30 °C, cells were harvested by centrifugation and lysed by sonication. The fusion protein (HisMBP-HT2ATPase) was purified by a nickel-nitrirotriacetic acid column followed by a heparin column. It was then cleaved by tobacco etch virus protease (made from...
pRK793, Scientific Reagents) with no scar into two pieces, His-MBP and HT2ATPase. His-MBP was removed when the cleavage product was run through QFF column (Amersham Biosciences). HT2ATPase was further purified by Hilo16/60 Superdex 75 size exclusion column (Amersham Biosciences).

Crystallization and Structure Determination—HT2ATPase was concentrated to 8 mg ml⁻¹ and dialyzed against 20 mM Tris, pH 7.4, 5 mM MgCl₂, 1 mM diethiothreitol. To get crystals of HT2ATPase bound to AMPPNP, AMPPNP (Sigma) was first added to the dialyzed protein solution to 1 mM. Crystals were grown at 4 °C using hanging drop vapor diffusion (NEXTAL crystallization tool) by mixing 2 μl of protein solution with 2 μl of well solution (4–8% polyethylene glycol 6,000, 10 mM β-mercaptoethanol). Crystals grew to a size of 150 × 150 × 1,000 μm within 2 days. The crystals belonged to the orthorhombic space group P2₁2₁2₁ (a = 70.3 Å, b = 90.5 Å, c = 148.3 Å, α = γ = 90.0°, β = 90.0°). Crystals were transferred via several steps to a solution of 21% polyethylene glycol 1,500, pH 8.0). Crystals reached the size of 120 μm × 120 μm × 120 μm.

The two protomers, although not strictly identical, are sufficiently similar that the dimer possesses a distinctly C₂-symmetric overall appearance (Figs. 1 and 2A). The polypeptide chain folds into two discrete structural modules, with the N-terminal module (residues 29–264, blue and cyan in Fig. 1A) bearing the characteristic nucleotide-binding fold of the GHKL superfamily (16), so named for its founding members, DNA gyrase, Hsp90, histidine kinase, and MutL. In this case, the fold consists of an eight-stranded, mixed β-sheet backed by four α-helices.

The C-terminal module (residues 265–428, magenta and peach in Fig. 1A) is known as the transducer for the important role it serves in communicating conformational information between the ATPase domain and the cleavage-religation domain (17–20). The core of this C-terminal module is also an α-β structure consisting of a four-stranded, mixed β-sheet with one α-helix on one face and two on the other. The electron density corresponding to the dozen residues at the C terminus of this module was strong enough to reveal an overall helical architecture but too weak to position each residue unambiguously; hence these residues (412–428 in protomer A, 406–428 in protomer B) were not included in the final model.

The heart-shaped dimer has dimensions of roughly 75 Å wide and 79 Å long when viewed from the perspective of Fig. 1A. The primary dimerization interface involves the nucleotide-binding modules of the two monomers and buries roughly 1,800 Å² of total solvent-accessible surface area. The N-terminal 20 residues from each monomer, comprising two short helices connected by an extended strand, interact exclusively with the nucleotide-binding module of the C₆ symmetry-related monomer. The contacts between the two transducer modules are much less extensive, burying only about 400 Å² of the total solvent-accessible surface area. Contacts are also established between the nucleotide-binding module of one monomer and the transducer module of the other monomer, and these bury ~500 Å² of the total solvent-accessible surface area. The entire dimer interface thus encompasses some 2,700 Å² of protein surface area, with half being contributed by each protomer. The HT2ATPase contains a small cavity some 17 Å wide and 14 Å high, bounded at the front and back by a 22-amino acid β-hairpin from the nucleotide-binding module of each monomer and at the sides and bottom by the transducer module. This cavity is more readily apparent in Fig. 1, B and C, in which the surface of the β-hairpin has been omitted to show the cavity more clearly. This hairpin structure is unique to eukaryotic Topo II enzymes and contains many residues that are conserved from yeast to human. The overall structure of the HT2ATPase·AMPPNP complex superimposes well with the S. cerevisiae Topo II ATPase·AMPPNP complex (5) (supplementary Fig. 1) with C-α root mean square deviations of 3.4 Å for the dimer and 1.8 Å for the protomer, respectively.

The nucleotide-binding mode observed here in the HT2ATPase·AMPPNP complex is quite similar to that reported for the corresponding complexes of other type II Topo II ATPases (5, 21–23). Briefly, each of the monomers contains one nucleotide-binding pocket, formed mainly by residues of the nucleotide-binding module (Figs. 1A and 3A, A and B). Three α-helices and one loop of the nucleotide-binding module form four sides of a box-like cleft; the β-sheet element contributes the bottom, and an extended loop termed the ATP lid (16) forms the top. The adenine ring is held in place through direct hydrogen bonding to the Asn-120 side-chain carbonyl plus water-mediated contacts to the side-chain groups of Asn-120, Thr-215, and Tyr-34’ (from the other protomer). The ribose ring of the nucleotide analog appears to adopt the C₁′exo sugar pucker (supplementary Fig. 2A), stabilized through hydrogen bonding of the 2′- and 3′-hydroxyls to the side chains of Ser-149 and Asn-150, respectively. Backbone amide N-H groups from six consecutive residues of the ATP-lid (residues 162–167) contact the α- and γ-phosphates of AMPPNP via their non-bridging oxyanions. The backbone N-H groups of residues 162 and 164 are also positioned...
within hydrogen-bonding distance of the bridging \( \beta, \gamma \)-imino nitrogen of AMPPNP; these contacts are likely to be more favorable with the oxygen atom of the actual substrate ATP. Ser-148 donates a hydrogen bond to the \( \beta \)-phosphate, and Lys-168 and the \( \delta \) amide nitrogen of Asn-91 interact with the \( \alpha \)-phosphate. An active site Mg\(^{2+}\) ion contacts all three phosphates of AMPPNP, with the \( \delta \) carbonyl oxygen of Asn-91 and two water molecules completing the distorted octahedral metal-ion coordination shell (Fig. 3, A and B). The floor of the nucleotide-binding pocket is lined with apolar side chains that contribute van der Waals contacts to the adenine ring and ribose moiety of the nucleotide (Fig. 3E). Only one contact to the nucleotide is contributed by a residue outside of the nucleotide-binding module. The highly conserved Lys-378 from the QTK-loop of the transducer domain forms a salt bridge with the \( \gamma \)-phosphate of AMPPNP (Fig. 3, A and B, indicated by electron density map). This salt bridge is believed to stabilize the transition state of the hydrolysis reaction because mutation of the corresponding Lys residue in DNA gyrase (K337Q) results in dramatically decreased ATPase activity while preserving the ability of the enzyme to perform DNA cleavage (24).

### Structure of HT2ATPase-ADP Complex

Like the HT2ATPase-AMPPNP complex, the HT2ATPase-ADP complex is a pseudosymmetric dimer (Fig. 1D), with two dimers per asymmetric unit. The dimers are not identical crystallographically but are similar enough that either one is representative of both. The individual protomers in the ADP complex show greater variation than in the AMPPNP complex (see Fig. 2, B and A, respectively), suggesting a less constrained overall conformation in the ADP complex. Viewed from the perspective in Fig. 1D, the dimer is heart-shaped and elongated relative to the AMPPNP complex (75 Å wide and 90 Å high). This elongation results primarily from the ordering of a C-terminal extension on the transducer domains that is too disordered to model in the AMPPNP complex; the extensions on each monomer contact each other at the base of the dimer to encircle completely an irregular cavity some 26 Å wide and 18 Å high at its upper end and 40 Å wide and 23 Å high at the lower end (more readily apparent in Fig. 1, E and F).

Each ligand-binding pocket (Fig. 3, C and D) contains one bound ADP molecule. The resolution of the ADP complex is not as high as that of the AMPPNP complex; thus the ADP-binding mode cannot be described in the same detail as that for AMPPNP, but the salient features of recognition are clear nonetheless. Based on proximity to side chains

### Data collection and model statistics

| Data collection model statistics | HT2ATPase-AMPPNP complex | HT2ATPase-ADP complex |
|---------------------------------|--------------------------|----------------------|
| Data collection                 | X-25 of the National Synchrotron Light Source | Northeastern Collaborative Access Team of the Advanced Photon Source |
| Space group                     | \( P_{2_1} \) \( P_{2_1} \) | \( P_{2_1} \) |
| Unit cell dimensions            | \( a, b, c (\text{Å}) \) \( 70.8, 75.3, 164.9 \) \( 71.0, 90.5, 148.3 \) | \( 90.0, 90.0, 90.0 \) \( 90.0, 90.0, 90.0 \) |
| Resolution (Å) \( ^{\text{a}} \) | \( 50–1.86 (1.93–1.86) \) | \( 50–2.51 (2.60–2.51) \) |
| Unique reflections \( ^{\text{a}} \) | \( 71.651 (6.371) \) | \( 61.096 (5.268) \) |
| Rmerge \( ^{\text{a,b}} \) | \( 0.078 (0.495) \) | \( 0.053 (0.492) \) |
| Rfree \( ^{\text{a}} \) | \( 15.7 (2.5) \) | \( 23.0 (2.9) \) |
| Refinement and model statistics | | |
| Limiting resolution (Å) | 1.86 | 2.51 |
| Rcryst (%) | 22.0 | 23.5 |
| Rfree (%) | 24.3 | 28.1 |
| Mean B-value, all atoms | 37.9 (31.1) | 57.9 (71.0) |
| (Å\(^2\)) | | |
| Bond lengths (Å) | 0.006 | 0.007 |
| Bond angles (°) | 1.22 | 1.28 |
| Dihedral angles (°) | 23.4 | 23.9 |
| Ramachandran plot (%) | 87.6 | 83.1 |
| Most favored | 87.6 | 83.1 |
| Additionaly allowed | 12.1 | 15.9 |
| Generously allowed | 0.3 | 1.0 |
| Non-water atoms | 5,993 | 11,408 |
| Water molecules | 528 | 238 |

\( ^{\text{a}} \) Values in parentheses refer to the highest resolution bin.

\( ^{\text{b}} \) \( R_{\text{merge}} = \sum_{hkl} |F_o(hkl)| - |F_c(hkl)|/\sum_{hkl} |F_o(hkl)| \).

\( ^{\text{c}} \) \( R_{\text{free}} = \sum_{hkl} |F_o(hkl)| - |F_c(hkl)|/\sum_{hkl} |F_o(hkl)| \).

\( ^{\text{d}} \) \( R_{\text{free}} \) was calculated based on 5% (HT2ATPase-AMPPNP complex) or 7.5% (HT2ATPase-ADP complex) of the total data omitted during structure refinement.

\( ^{\text{e}} \) Values in parentheses were obtained by Wilson plot.

\( ^{\text{f}} \) Values provided by PROCHECK.
Motion of the Human Topo IIα ATPase Domain

![Figure 1](http://www.jbc.org/)

**FIGURE 1.** Overall structure of the 46-kDa ATPase domain of human topoisomerase IIα (HT2ATPase) (residues 29 – 428) bound to AMPPNP (A–C) and ADP (D–F). A and D, ribbon drawings emphasizing the modularity of the protein structure. Each of the four modules in the HT2ATPase homodimer is colored individually: blue and cyan, ATPase modules; magenta and peach, transducer modules. One protomer of HT2ATPase is blue and magenta; the other is cyan and peach. Asterisks denote the β-hairpin motif, unique to eukaryotic Topo II enzymes, which forms the upper end of the DNA channel. B and E, surface representation in the same orientation as A and D, with the β-hairpin shown in yellow. Red in underlying ribbon trace indicates C-terminal portions of the structure that become ordered in the ADP-bound state. C and F, surface drawing rotated by 90° about the x-axis with respect to B and E, looking inward from the C terminus–proximal end of the cavity.

![Figure 2](http://www.jbc.org/)

**FIGURE 2.** Analysis of nucleotide-dependent domain movement in HT2ATPase. A, least-squares superposition of the two protomers in the asymmetric unit of the AMPPNP structure. B, superposition of the four protomers in the asymmetric unit of the ADP structure. C, superposition of the dimers of the AMPPNP (blue) and ADP (yellow) structures. The magnitude of domain rotation and translation is indicated. Red X indicates the approximate pivot point for rigid-body rotation. D and E, superposition of the individual (D) nucleotide-binding and (E) transducer modules in the AMPPNP structure and ADP structure (coloring as in C).

on the protein, the adenine ring in the ADP complex appears to make the same set of direct and water-mediated contacts as in the AMPPNP complex; however, the bridging waters are not as evident in the former. Although the sugar pucker in the ADP complex cannot be deduced solely from the density of the furanose ring, the density for the substituents on the ring are incompatible with the C₄-endo pucker seen in the AMPPNP complex and are most consistent with a C₃-exo pucker (supplementary Fig. 2B). The contacts to both the α- and β-phosphate are the same in the ADP complex as in the AMPPNP complex, with the exception that the ADP complex has an additional hydrogen bond between the β-phosphate and the Asn-150 side chain; this is made possible by the disengagement of Asn-150 from the 2'-OH upon flipping from the 2'-exo to 2'-endo pucker. The details of Mg²⁺ ion coordination are also nearly identical in the two complexes, except that a water molecule fills the coordination site vacated by the γ-phosphate oxygen. Although the ATP-lid has lost three hydrogen-bonding contacts to the γ-phosphate, the lid remains in place because of the preservation of multiple contacts with the α- and β-phosphates.

**Nucleotide-dependent Structural Reorganization**—Superposition of the structures of the two complexes reveals significant changes in overall conformation (Fig. 2C). Although the nucleotide-binding modules superimpose very well, the transducer domain of each monomer of the ADP complex swings out by about 10°, increasing the width of the channel at the most narrow point from 6 to 17 Å. To assess whether this conformational change results from rigid-body movement of the nucleotide-binding modules with respect to the transducer modules, individual modules of the AMPPNP complex were superposed individually with their counterparts in the ADP complex (Fig. 2, D and E). This superimposition shows clearly that the individual modules possess nearly identical overall structures; the average root mean square deviations of C-α atoms between the nucleotide-binding domains is 0.82 Å, and that between the transducer domains is 1.84 Å.

As mentioned above, the nucleotide-dependent domain movement is also associated with an apparent difference in domain mobility. Whereas the two monomers in the asymmetric unit of the AMPPNP complex overlay with C-α root mean square deviations of 0.72 Å, the four monomers of the ADP complex overlay with root mean square deviations of 1.22 Å (see Fig. 2, A and B, respectively).

To gain a better sense of the subtle structural changes in the nucleotide-binding site that give rise to rigid-body domain motions in the transducer module, we superimposed the nucleotide-binding sites of both complexes (Fig. 4). The architectures of the active sites are very similar. The nucleotide, the Mg²⁺ ion, and many key residues of the active site (Glu-87, Asn-91, Asn-120, Ser-148, Lys-168) are in essentially the same position in both structures. The difference in sugar pucker is associated with some local adjustments in contacts to the sugar and β-phosphate but has remarkably little effect on the relative positioning of the key recognition elements in the nucleotide, namely its adenine and phosphate anhydride moieties (Fig. 4B). The γ-phosphate of AMPPNP is replaced by what appears to be a water molecule in the HT2ATPase-ADP complex. The only nucleotide contact residue that
undergoes a major change in contact status and position is precisely the single residue from the transducer module that makes a nucleotide contact, Lys-378. This residue, which is absolutely conserved in all type II topoisomerases, forms a salt bridge to the \( \text{H}9280 \)-phosphate in the AMPPNP structure. The rigid-body movement of the transducer module relative to the nucleotide-binding module in the ADP structure retracts Lys-378 away from the nucleotide. In one of the two dimers in the asymmetric unit of the ADP complex, both Lys-378 side chains are disordered; in the other dimer, each side chain is ordered and pointing toward the nucleotide but is retracted by 2.6 Å relative to its position in the AMPPNP complex (Fig. 4). In the AMPPNP complex, the distance between the Lys-378 ε-amino nitrogen and the nearest \( \gamma \)-phosphate oxygen atom is 2.6 Å, consistent with a hydrogen bond between the two. If the transducer module were to adopt the conformation observed in the ADP complex but instead bound an ATP nucleotide the distance between the Lys-378 ε-N and the nearest \( \gamma \)-phosphate oxygen atom would be 4.2 Å (Fig. 4B), as determined by modeling on the basis of the present structures. Because this region of the structure is exposed to solvent, a separation of 4.2 Å between the Lys-378 ε-N and the ATP \( \gamma \)-phosphate in the retracted state would be unlikely to produce a significant hydrogen-bonding interaction. We thus propose that the Lys-378 ε-N acts as both a sensor and a relay for the nucleotide-bound status of the protein, drawing the transducer domains inward upon interaction with the \( \gamma \)-phosphate on ATP and allowing outward movement in the absence of such a stabilizing electrostatic interaction.

**DISCUSSION**

Topo II performs a complex multistep reaction sequence on the genome involving introduction of a double strand break in one segment (G-segment) of the bound DNA substrate, passage of another duplex segment (T-segment) through the break, and religation of the break to restore the continuity of the DNA backbone (Fig. 5). The enzyme is believed to accomplish this difficult operation by opening and closing three different gates consisting of dimer interfaces at distinct points in the protein structure (9). The management of these gates, and hence the precise orchestration of strand scission/religation and passage, is controlled by the hydrolysis of ATP to ADP and inorganic phosphate. By analogy to other nucleotide-dependent switching proteins, it has been suspected for some time that the nucleotide hydrolysis cycle drives conformational changes in the ATPase domain of Topo II, but the nature of
these conformational changes has remained unelucidated, as has the mechanism by which these changes enable regulated strand passage. Here we have determined the structure of the human Topo IIα ATPase domain with two different bound nucleotides, the ATP analog AMPPNP and the hydrolysis product ADP. The former structure is very similar to that determined previously for the AMPPNP-bound state of yeast Topo II ATPase (5), whereas the latter structure represents the first observation of the product state of the ATPase domain of the type IIA DNA topoisomerases. Comparison of these structures reveals a nucleotide-dependent rigid-body movement of the transducer module relative to the nucleotide-binding module; in the ADP state, the cavity is relatively open and conformationally mobile, and in the AMPPNP (ATP-like) state the cavity is relatively closed and conformationally fixed. Similar module motion has been found in the monomeric form of *Sulfolobus shibatae* Topo VI (a type IIB Topo) B subunit (22, 27) and *Thermus thermophilus* GyrB in complex with novobiocin (28). The structures thus suggest that hydrolysis of ATP to ADP or phosphate release is associated with the opening of the cavity between the N-gate and M-gate in Topo II. Without the structure of the full-length protein dimer with DNA in a variety of ATP hydrolysis states, it is impossible to conclude how ATP hydrolysis is conformationally transduced to the cleavage-reunion core. However, the differences between the two structures presented here are clearly suggestive of and are readily incorporated into the current two-gate model.

How does the cavity opening observed in the present structures fit into the larger picture of Topo II function? A model for the Topo II catalytic cycle that incorporates our findings together with those of previous studies (1–3, 5, 7–9, 18, 20, 25, 29–43) is illustrated in Fig. 5.

The present structures indicate that expansion is likely to occur by rigid-body displacement of the transducer module. In the ATP-bound state, rigid-body displacement of the transducer module would result in retraction of the Lys-378 ε-amino group from the γ-phosphate. There is little doubt that such movement of Lys-378, a residue that is essential for ATPase activity, would affect catalysis, but it is impossible at this stage to conclude whether the movement would decelerate ATP hydrolysis or accelerate it. Whatever the case, the structures establish the notion that transducer movement can influence ATP hydrolysis and that ATP hydrolysis in turn allows relaxation of the transducer domains to an open conformation.

### FIGURE 4. Nucleotide-dependent changes in the active site of HT2ATPase

A, superposition of the nucleotide-binding sites of the AMPPNP structure (blue) and ADP structure (yellow). In both structures, the nucleotide is in gray. B, superposition showing only the nucleotide and region around Lys-378. Distance measurements are made using the ε-amino of Lys-378 and the nearest γ-phosphate oxygen.

### FIGURE 5. Model for nucleotide-dependent DNA translocation by Topo II, incorporating the present findings

The ATPase domain is depicted schematically in the context of the full protein dimer (nucleotide binding module in green, transducer module in red). The remainder of the protein, known as the cleavage-reunion core, comprises the module directly C-terminal to the ATPase (B′ cleavage-reunion module, yellow) and a domain that bears homology to gyrase A(A′ cleavage-reunion module, blue). Step 1, G-segment DNA (gray) binds to the topoisomerase II dimer at the second gate of the cleavage-reunion core. Step 2, an incoming T-segment of DNA (pink) is captured in the upper cavity between the upper and middle gates by ATP (red star) binding-promoted dimerization. Step 3, hydrolysis and release of one of the two bound ATP molecules to ADP (black star) and P, lead to an asymmetric retraction of the catalytic Lys-378, opening of the enzyme-bridged middle gate of the protein-DNA complex, and movement of the T-segment from the upper cavity formed in the ATPase domain into the lower cavity formed by the cleavage-reunion core. Step 4, hydrolysis of the remaining ATP and opening of the bottom gate allow for T-segment egress from the complex and resetting the system.
The present structures suggest the need for transducer movement to accommodate a DNA duplex, but they provide no insight into whether one or both transducers move. The fact that ATP hydrolysis occurs sequentially (30–33) raises the intriguing possibility that only one of the two transducer modules swings outward, and it is this structural asymmetry that gives rise to catalytic asymmetry in the ATPase activity.

Whatever the origins of sequential hydrolysis, the present structures predict that the transducer module of the ADP-bound protomer (and for similar reasons, the nucleotide–free protomer) is the one that swings out to expand the cavity. Even though the ADP/ADP-bound state captured here structurally is not believed to represent an actual intermediate in the overall reaction scheme, the nucleotide–dependent movement of the transducer module is likely to occur similarly in the ADP-bound protomer of the ATP/ADP state. We assert this because the nucleotide-bound status (ATP versus ADP) has a negligible effect on the structure of the nucleotide binding/dimerization module, and its effects are almost entirely localized to a specific pivot point in the transducer module. There is good reason therefore to expect that the ability of Lys-378 ε-N to act as both a sensor and a relay for nucleotide status operates independently in the two protomers.

After the expansion of the upper cavity mediated by the motion of the transducer domain, inorganic phosphate is released, the central gate dissociates from the enzyme, but the upper gate remains closed because the lower chamber opens and releases the T-segment from the complex; the remaining ATP undergoes hydrolysis, and the protein-DNA complex is poised to receive another T-segment.

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