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Understanding the coupling between DNA damage detection and UvrA’s ATPase using bulk and single molecule kinetics

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ABSTRACT: Nucleotide excision repair (NER) protects cells against diverse types of DNA damage, principally UV irradiation. In *Escherichia coli*, damage is recognized by 2 key enzymes: UvrA and UvrB. Despite extensive investigation, the role of UvrA’s 2 ATP domains in NER remains elusive. Combining single-molecule fluorescence microscopy and classic biochemical methods, we have investigated the role of nucleotide binding in UvrA’s kinetic cycle. Measurement of UvrA’s steady-state ATPase activity shows it is stimulated upon binding DNA ($k_{cat}$, 0.71–1.07/s). Despite UvrA’s ability to discriminate damage, we find UV-damaged DNA does not alter the steady-state ATPase. To understand how damage affects UvrA, we studied its binding to DNA under various nucleotide conditions at the single molecule level. We have found that both UV damage and nucleotide cofactors affect the attached lifetime of UvrA. In the presence of ATP and UV damage, the lifetime is significantly greater compared with undamaged DNA. To reconcile these observations, we suggest that UvrA uses negative cooperativity between its ATPase sites that is gated by damage recognition. Only in the presence of damage is the second site activated, most likely in a sequential manner.—Barnett, J. T., Kad, N. M. Understanding the coupling between DNA damage detection and UvrA’s ATPase using bulk and single molecule kinetics. FASEB J. 33, 000–000 (2019). www.fasebj.org

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Genomic DNA is constantly damaged by both exogenous and endogenous sources and must be repaired efficiently to maintain genome integrity. Nucleotide excision repair (NER) is an evolutionarily conserved DNA repair mechanism across all kingdoms of life. NER primarily repairs bulky lesions, including UV-induced damage, such as cyclobutane pyrimidine dimers and 6–4 photoproducts. However, NER also acts promiscuously to recognize and repair myriad lesion types (1). The classic description of NER involves the processing of the lesion by multiple enzymes and requires detection, incision, and repair resynthesis (2). Damage verification requires UvrA (a homodimer that is referred to as UvrA here for clarity) to locate damage with UvrB. Upon location of damage, UvrA is ejected from the UvrAB “preincision” complex, leaving UvrB alone on DNA. UvrC is then recruited to perform 2 DNA incisions on either side of the lesion on the same strand. Subsequently, UvrD and DNA polymerase I carry out downstream damaged oligonucleotide removal and repair resynthesis (3, 4). Finally, DNA ligase seals the remaining nick in the DNA backbone.

Successful lesion excision in *Escherichia coli* requires ATP (3, 5–7). Analysis of UvrA’s sequence reveals 2 type A Walker motif sites, one at residues 31–45 (N-terminal site) and one at residues 640–654 (C-terminal site) (8). In the 3-dimensional structure of dimeric UvrA, the N-terminal ATPase domain of 1 UvrA is close to the C-terminal ATPase site of the other. Furthermore, all 4 are positioned beneath the DNA binding cleft that runs across the face of the protein (9, 10). The role of UvrA’s ATPase activity is uncertain but has been linked to dimer formation and lesion searching (5, 11, 12). Previous studies that mutated catalytic lysine residues in each ATPase site observed a drastic loss of overall ATPase activity (13). Such mutational studies have indicated a role for ATPase in loading UvrB onto DNA (13, 14). Because many of these studies conflict in their outcomes, a clearly defined ATPase mechanism and definition of its role in UvrA’s function is needed.

UvrA’s ATPase sites are thought to have distinct roles (13, 15, 16); the C-terminal site discriminates DNA damage, and the N-terminal site is implicated in UvrA dimer formation and UvrB binding (17, 18). Using the *in vitro*
DNA tightrope assay in conjunction with mScarlet-labeled *E. coli* UvrA (19), we have investigated, at the single-molecule level, how different nucleotide cofactors affect UvrA’s interaction with DNA. UvrA was observed to perform a 3-dimensional search on DNA, with a lifetime altered by the nucleotide condition, and the presence of UV damage. The rate-limiting step for ATP turnover occurs on DNA, and the presence of damage alters UvrA’s DNA bound lifetime but not its steady-state ATPase rate. This paradox is resolved if UvrA sequentially hydrolyzes ATP in its 2 sites, suggesting that UvrA has a negatively cooperative ATPase that is tightly coupled to damage recognition.

**MATERIALS AND METHODS**

Unless stated otherwise, all chemicals were purchased from Thermo Fisher Scientific (Waltham, MA, USA), and DNA oligonucleotides were from MilliporeSigma (Burlington, MA, USA). All *in vitro* experiments were carried out at room temperature in ABC buffer [50 mM Tris (pH 7.5), 500 mM KCl, 0.1 mM EDTA, 10 mM MgCl₂, and 10 mM DTT].

**Protein expression and purification**

The gene for *E. coli* UvrA, obtained from National Bioresource Project (NIG, Kyoto, Japan), was engineered onto a C-terminal mScarlet fluorescent protein separated via a flexible linker (20). Further C-terminal to mScarlet, we placed a His tag for purification and induced expression of this pET21a construct overnight at 18°C using isopropyl β-d-thiogalactopyranoside. Purification was performed using nickel affinity followed by heparin chromatography (GE Healthcare, Chicago, IL, USA). To avoid protein precipitation, we kept KCl concentrations >200 mM. Once purified, the UvrA concentration was determined using mScarlet’s extinction coefficient (M⁻¹ cm⁻¹) at 569 nm and stored in 50% glycerol 50 mM Tris (pH 7.5), 500 mM KCl, 0.1 mM EDTA, and 10 mM DTT at −20°C (18).

**Complementation assays**

To investigate whether our UvrA-mScarlet construct can rescue UvrA wild-type (UvrA<sup>WT</sup>) knockout cells, we performed *in vitro* UV complementation assays. UvrA<sup>WT</sup> knockout (Keto) cells were obtained from the National Bioresource Project (NIG) and transformed with UvrA-mScarlet or empty mScarlet vector. Previously, we have shown UvrA<sup>WT</sup> knockout cells complemented with ecotropically expressed UvrA<sup>WT</sup> survive UV irradiation at 5 J/m² (21). Transformed cells were grown in Luria-Bertani medium with 25 µg/ml chloramphenicol to OD600 of 0.5. Five microliters of undiluted and three 10-fold serial dilutions were plated on Luria-Bertani agar and subjected to either no UV or 5 J/m² UV (254 nm) irradiation and incubated overnight at 37°C in the dark. UvrA<sup>WT</sup> and UvrA-mScarlet fully restored UV survival compared with UvrA<sup>WT</sup> knockout and UvrA<sup>WT</sup> knockout cells ecotopically expressing mScarlet alone.

**NADH-linked ATPase assay**

ABC buffer supplemented with 0.5 mM phosphoenol pyruvate solution was stored at −20°C; 1 mM DTT was added upon thawing. The phosphoenol pyruvate solution with DTT was blanked at 340 nm in a spectrophotometer, and then 10 µl of pyruvate kinase (600–1000 U/ml) and lactate dehydrogenase (900–1400 U/ml, premixed stock from MilliporeSigma) per 500 µl reaction were added to a cuvette with 210 µM NADH. The change in OD340 was fitted linearly to calculate loss of NADH (6220 M⁻¹ cm⁻¹ at 340 nm), enabling calculation of k<sub>cat</sub>. Reactions were repeated 3 times, and the error represents the SD.

**DNA substrates**

DNA tightropes were constructed as previously described (21). Nucleotides [ATP, ADP, adenosine 5'-O-(3-thio)triphosphate (ATP<sub>s</sub>)] were used at 1 mM final concentration. For experiments with P<sub>1</sub>, 1 mM AMP with 50 mM Tris-HCl (pH 7.5), 10 mM MgCl₂, and 10 mM DTT was supplemented with 19 mM sodium phosphate (freshly autoclaved to remove pyrophosphate), and ionic strength was balanced by the removal of 50 mM KCl (23). For experiments with damaged λ DNA, the DNA was exposed to 1000 J/m² of 254 nm light immediately before creating tightropes.

**Single molecule imaging and analysis**

Imaging was performed using a custom-built fluorescence microscope capable of oblique-angle fluorescence excitation and multichannel emission (21). mScarlet was excited using a 561 nm diode OBIS LS laser (Coherent, Santa Clara, CA, USA) at 5 mW. To determine if photobleaching affected our observed attached lifetimes, we measured durations of attachment for the longest associating nucleotide condition, UvrA-ATP<sub>s</sub>, at different laser powers. The attached lifetime was reduced only above 10 mW, indicating that, at the laser powers used in our experiments, photobleaching is not defining the lifetime. Videos were collected with a 300 ms exposure time for 90 s using 2 x 2 binning and were transformed into kymographs using ImageJ.

Attached lifetimes in each condition were plotted as cumulative frequency histograms to remove any bin size dependence. Cumulative frequencies were fitted to exponentials, and F-tests were used to determine the requirement for single vs. double exponential fits.

**RESULTS**

**DNA accelerates the steady-state ATP hydrolysis rate of UvrA**

Although it is known that *E. coli* nucleotide excision repair requires ATP (3, 6, 7), its role in UvrA’s function has not been fully determined. Using an NADH-linked assay, the steady-state UvrA-mScarlet (UvrA hereafter) ATP turnover rate was measured with and without DNA. Absorbance at 340 nm was used to linearly quantify ADP release because
1 NADH is oxidized to NAD<sup>+</sup> for each ADP released. Linear fits gave the rates of ATP hydrolysis shown in Fig. 1.

This assay was repeated for various damaged and undamaged DNA substrates (Fig. 2A). Undamaged DNA substrates stimulate the ATPase activity of UvrA significantly (k<sub>cat</sub> UvrA alone, 0.71 ± 0.05 ATP/UvrA/second vs. UvrA + undamaged DNA, 1.07 ± 0.08 ATP/UvrA/second; P < 0.0001). The ATPase rate was also stimulated by a fluorescein-containing oligonucleotide F26.50 (1.06 ± 0.142 ATP/UvrA/second), UV damaged pUC18 (1.13 ± 0.05 ATP/UvrA/second), or λ phage DNA (1.03 ± 0.12 ATP/UvrA/second) compared with ATP alone (P < 0.0001). However, damaged DNA does not affect the maximal ATPase activity relative to undamaged equivalents (P = 0.9594) (Fig. 2A, combined data). To investigate if damage discrimination affects UvrA’s ATPase, we titrated ATP (Fig. 2B) in the presence of undamaged DNA or UV-irradiated DNA. The K<sub>m</sub> for ATP decreased from 195 μM in the presence of undamaged DNA to 60 μM with damaged DNA. The ATP titration independently confirmed that the k<sub>cat</sub> (per UvrA monomer) was unchanged, remaining at 1.4/s. Because ATP hydrolysis is accelerated by DNA binding, this indicates that the rate-limiting kinetic process occurs while bound to DNA. If the rate-limiting step did not occur on DNA, there would be no change in steady-state ATPase upon addition of DNA.

Determining the kinetic state affected by DNA binding

To identify which nucleotide state is affected by DNA binding, we measured the DNA-bound lifetime of UvrA using single-molecule fluorescence imaging. To do this, we used an in vitro DNA tightrope assay, which comprises individual DNA molecules suspended between silica beads in a microfluidic chamber (24, 25). Oblique angle illumination is used to limit the background from molecules in solution, leading to clear images of mScarlet-labeled protein binding to the DNA tightropes. The effects of damage were studied using DNA exposed to UV light (254 nm) immediately before the formation of tightropes. Videos of UvrA molecules binding to DNA were transformed into kymographs (Fig. 3A) for dwell-time analysis and determination of the search mechanism. One-dimensional sliding on DNA will appear as movement on the Y axis (position) over time, whereas a 3-dimensional search will appear as horizontal streaks (24) (Fig. 3A). The length of any continuous interaction corresponds to the dwell time. It was clear from the data that UvrA uses a 3-dimensional search mechanism because no positional movement was seen. The individual attached lifetimes were plotted as cumulative frequency histograms and fit to exponentials, which are appropriate for stochastic interactions (Fig. 3B). These fits across various nucleotide and DNA damage conditions provided dissociation rate constants (Fig. 3C and Table 1).

The bulk-phase steady-state ATPase k<sub>cat</sub> was determined to be 1.07 ATP/UvrA per second in the presence of damaged or undamaged DNA (Fig. 2). This is in excellent agreement with the single-molecule detachment rate constant of 1.24 ± 0.038/s for UvrA on undamaged DNA with ATP. These data fit better to a double exponential (F-test = 204) than a single, suggesting 2 populations of attachments, the vast majority of which (amp1 = 94%) reflect molecules binding to undamaged DNA. The remainder most likely represents molecules that bind to nonspecific damage on λ DNA, as previously shown (24). In the presence of UV-damaged DNA and ATP, UvrA shows a dramatic decrease in the detachment rate constant (0.411 ± 0.023/s; i.e., a 3-fold increase in attached lifetime). The UV damage cumulative frequency histogram also fits better to a double exponential (F-test = 154; amp1 96%, k1 0.411 ± 0.023/s; amp2 4%, k2 0.098 ± 0.030/s; Fig. 3B), consistent with 2 populations of attached proteins. However, the small contribution (amp2 = 4%) of k2 means very few complexes are long lived on DNA. We did not investigate the origin of this species further.

To determine the contribution of the ATP binding to the attached lifetime of UvrA, we used ATP<sub>γ</sub>S, a non-hydrolyzable analog of ATP. ATP<sub>γ</sub>S has been shown to permit formation of UvrA dimers and stimulate binding of UvrA to DNA (5). Figure 3C shows that ATP<sub>γ</sub>S slows the release of UvrA from DNA (0.155 ± 0.002/s), equivalent to a 6.5-fold increase in attached lifetime. DNA damage does not affect the dissociation kinetics of UvrA in the presence of ATP<sub>γ</sub>S (0.127 ± 0.001/s), suggesting that ATP-bound UvrA is not involved in damage recognition (5, 12). The slow rate constant for ATP-bound detachment implies that hydrolysis and formation of subsequent nucleotide states occur on DNA; consequently, one of these states will be sensitive to damage.

A possible candidate for controlling the release of UvrA from DNA is the ADP-bound state. ADP facilitates UvrA...
dimerization and binding to DNA (4, 11, 12). Surprisingly, UvrA in the presence of ADP did not substantially bind to DNA tightly compared with identical concentrations of UvrA with ATP or ATPγS. Nonetheless, from the attachments observed it was possible to determine the lifetime as 4.9 s (0.20 \pm 0.006/s), which was 5-fold longer than in the presence of ATP. The low level of attachment suggests that few UvrA dimers exist in the presence of ADP (5), conflicting with previous observations (12). With UV-damaged DNA, the attached lifetime remains unchanged at 4.6 s (equivalent to 0.22 ± 0.016/s). It is likely that UvrA does not dimerize with only ADP bound and that the few dimers that form are strongly bound to the DNA. Given the low abundance of this state, it is likely that detachment occurs from the ADP state. Furthermore, the similar rates of release in the presence and absence of DNA damage suggest that UvrA-ADP is not a damage sensing state.

The step between ATP binding and formation of the UvrA ADP-bound state is therefore the most likely to detect damage. To investigate this, we attempted to create the ADP+P_i state of UvrA by adding a large excess of free phosphate to UvrA and ADP. This required careful consideration of ionic strength changes that were balanced by a reduction in the buffer salt (26). On undamaged DNA, the addition of phosphate to the ADP state increases the detachment rate constant slightly from ADP alone (0.202 ± 0.006 to 0.234 ± 0.007/s) but does not resemble the ATP condition (1.07 ± 0.019/s), nor does phosphate restore UvrA’s affinity for DNA; the number of binding events was also low, similar to the ADP-bound state. However, in the presence of UV-damaged DNA, UvrA’s detachment rate constant significantly increases (from 0.217 ± 0.016 to 0.364 ± 0.011/s; \( P < 0.0001 \)), resembling the rate constant for release in the presence of UV damage and ATP (0.411 ± 0.023/s; \( P = 0.501 \)). This change in lifetime suggests that UvrA with P_i and ADP is populating the damage-sensitive state, which is rate limiting for UvrA detachment from DNA.

**DISCUSSION**

The role of ATP in the mechanism of NER is not clearly understood. Here, we compare the kinetics of ATP turnover by UvrA at the bulk- and single-molecule level. We
have found that the bulk-phase ATPase $k_{cat}$ of UvrA was stimulated by the presence of DNA; however, there was no further change with damaged DNA. This strongly contrasted with the attached lifetimes of UvrA on DNA. The presence of damage increased the attached lifetime 2- to 3-fold in the presence of ATP. After determining the attached lifetimes of UvrA molecules on either damaged or undamaged DNA under various nucleotide conditions, we infer that UvrA most likely functions as a negatively cooperative ATPase, using its 2 ATP sites in sequence.

**UvrA’s second ATPase site is activated by DNA damage**

The steady-state ATPase rate for UvrA increased ~50% upon addition of DNA, demonstrating that UvrA is a DNA-stimulated ATPase. However, no further change in the DNA-stimulated ATPase was observed with damaged DNA. Similar changes in bulk-phase ATPase using the linked assay have been reported (17). We used damage substrates at various damage densities and different types of substrate (see Supplemental Information); however, in all conditions the type or density of damage had no effect on the observed results. We also ensured the DNA damage concentration was higher than the UvrA concentration used. Furthermore, linear and plasmid DNA substrates stimulated the ATPase of UvrA equally, indicating that the DNA ends play no part in the observed activation. This contrasts with previous studies, where undamaged DNA was seen to inhibit the ATPase and damage returned the activity to that seen without DNA (16). This difference may be due to ADP’s strong affinity for UvrA (14), which may inhibit the ATPase; our use of the linked assay ensures nucleotide is recycled, abrogating any possible issues arising from such product inhibition.

By directly imaging single UvrA molecules binding to DNA, we were able to correlate attached lifetimes with the biochemically measured ATPase rates. With ATPγS, ample binding to DNA was observed; however, with ADP this was virtually abolished, suggesting that UvrA binds DNA
in the ATP-bound state and releases from DNA in the ADP-bound state. In both cases the attached lifetimes are too long to correlate with the steady-state ATPase rate and were not affected by DNA damage; therefore, these states are unlikely to be substantially populated, and their role in the damage-modulated ATPase of UvrA is irrelevant. The most interesting observation derives from a comparison of the bulk $k_{\text{cat}}$ of 1/s vs. the lifetime in the presence of ATP (~1/s); these values are identical. The $k_{\text{cat}}$ was calculated per UvrA monomer; therefore, we infer that on DNA where UvrA is a dimer (4, 12, 24, 27) 2 ATP molecules are hydrolyzed and that attachment to DNA is tightly coupled to ATP turnover. However, in the presence of damage, although $k_{\text{cat}}$ is unchanged, the attached lifetime increases substantially. The ~3-fold increased affinity for ATP on damaged DNA corresponds well to the 3-fold increase in lifetime on damaged DNA, demonstrating that, in the presence of damage, there remains tight coupling between ATP turnover and UvrA release from DNA. Therefore, to reconcile the unchanged $k_{\text{cat}}$ with the increased lifetime, we suggest that when damage is detected during turnover of the first ATP, rather than detaching from DNA, the second site is activated. This second site has a similar ATP hydrolysis rate, thus keeping the total number of ATPs hydrolyzed per second the same but increasing the attached lifetime. This sequential action is indicative of negative cooperativity between the sites that is tightly coupled to DNA damage recognition. Similar mechanisms of negative cooperativity have been noted for other ABC ATPase superfamily members, including the bacterial mismatch repair protein MutS (28) and its eukaryotic homolog (29, 30).

**UvrA’s rate-limiting step is affected by the presence of DNA damage**

Because the rate-limiting step is not ADP release, we sought to determine whether hydrolysis and subsequent release of Pi controls the turnover rate. To push the pathway backward, P_i was added to ADP and UvrA. If ATP hydrolysis and release of ADP + P_i is the rate-limiting step, the addition of P_i to ADP should restore the lifetime to the ATP condition. With undamaged DNA, the detachment rate constant did not resemble that of UvrA in the presence of ATP (0.23 vs. 1.2/s). Therefore, the addition of phosphate does not reverse hydrolysis of the first site. However, with damaged DNA the detachment rate constant was very similar to that of UvrA with ATP and UV damage (0.36 vs. 0.41/s). We surmised that the 0.41/s detachment rate constant was derived from sequential ATP hydrolysis at both sites. This suggests that, in the presence of damage, both nucleotide binding sites can reload P_i and that the rate-limiting steps are recapitulated, possibly through reversal of hydrolysis. The identity of the 2 sites was revealed using ATPase mutants (14) and in vivo (13). Based on these previous observations that nucleotide needs to be bound to the C-terminal (or distal) ATPase site for damage recognition, we suggest this is the first ATPase site described here. Therefore, the second ATPase site (N-terminal or proximal) is activated after damage recognition at the first site. This confirms the 2-site hypothesis, which suggests the N-terminal site is necessary for recruitment of UvrB (13, 14, 17).

Communication between the ATPase sites is likely facilitated by their close proximity. The N-terminal site of 1 ATPase site is adjacent to the C-terminal site. Interestingly, these sites are connected by a tunnel (Fig. 4); it is plausible that P_i release occurs through this tunnel and gates the recruitment of nucleotide or hydrolysis at the second site. Future structural investigations of the transition state are imperative to reveal how P_i is released, whether through the tunnel or by monomerization.

![Crystal structure of Thermotoga maritima UvrA showing the phosphate tunnel.](image-url)
CONCLUSIONS

In summary, we have combined data from single-molecule and bulk-phase studies of the UvrA ATPase and DNA interaction. We find excellent correlation between the kinetics using both methods and that UvrA's interaction with DNA is tightly coupled to its ATPase rate-limiting step. Two ATPs are hydrolyzed per dimer per second when bound to undamaged DNA. However, in the presence of damage, although the steady-state ATPase is unaltered, the attached lifetime increases. We suggest this occurs through a sequential mechanism whereby if damage is located, the second ATPase site is activated, resulting in another 2 ATPs per dimer per second being hydrolyzed. This accounts for the observed kinetics and suggests a tight coupling model due to strong negative cooperativity between the UvrA ATPase sites. Therefore, UvrA couples its ATPase activity to its damage-sensing function by activating the second ATPase site.

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AUTHOR CONTRIBUTIONS

J. T. Barnett collected data; and J. T. Barnett and N. M. Kad designed the experiments, analyzed the data, and wrote the manuscript.

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