



Structure of a DNA glycosylase that unhooks interstrand cross-links

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DNA glycosylases are important editing enzymes that protect genomic stability by excising chemically modified nucleobases that alter normal DNA metabolism. These enzymes have been known only to initiate base excision repair of small adducts by extrusion from the DNA helix. However, recent reports have described both vertebrate and microbial DNA glycosylases capable of unhooking highly toxic interstrand cross-links (ICLs) and bulky minor groove adducts normally recognized by Fanconi anemia and nucleotide excision repair machinery, although the mechanisms of these activities are unknown. Here we report the crystal structure of *Streptomyces sahachiroi* AlkZ (previously Orf1), a bacterial DNA glycosylase that protects its host by excising ICLs derived from azinomycin B (AZB), a potent antimicrobial and antitumor genotoxin. AlkZ adopts a unique fold in which three tandem winged helix-turn-helix motifs scaffold a positively charged concave surface perfectly shaped for duplex DNA. Through mutational analysis, we identified two glutamine residues and a β -hairpin within this putative DNA-binding cleft that are essential for catalytic activity. Additionally, we present a molecular docking model for how this active site can unhook either or both sides of an AZB ICL, providing a basis for understanding the mechanisms of base excision repair of ICLs. Given the prevalence of this protein fold in pathogenic bacteria, this work also lays the foundation for an emerging role of DNA repair in bacteria-host pathogenesis.

DNA repair | DNA glycosylase | interstrand cross-link | azinomycin B | winged-helix

The chemical integrity of DNA is constantly challenged by cellular and environmental genotoxic agents that produce a diverse array of covalent nucleobase adducts. If left unrepaired, these DNA lesions impair important cellular processes, including replication, transcription, and cell cycle regulation, leading to mutations, chromosomal rearrangements, and genomic instability that threaten the livelihood of the organism and lead to human diseases (1). Small adducts are typically removed by base excision repair (BER), in which a lesion-specific DNA glycosylase catalyzes hydrolysis of the *N*-glycosidic bond of the aberrant nucleobase, liberating it from the phosphoribose backbone (2). The resulting abasic site is nicked at the 5' side by an apurinic/aprimidinic (AP) endonuclease to create a free 3'-hydroxyl group necessary for the synthesis of new DNA. In contrast, bulky and helix distorting lesions are removed by nucleotide excision repair (NER), which involves removal of an oligonucleotide segment containing the lesion through the action of dual nuclease incisions, followed by helicase removal of the damaged segment.

Interstrand cross-links (ICLs) from various endogenous and environmental sources covalently tether the opposite strands of DNA and pose a major obstacle to normal DNA metabolism (3, 4). The cytotoxicity of cross-linking agents makes them particularly effective as antitumor drugs (5). Because ICLs damage both strands of DNA, repair of ICLs is more elaborate than repair of monoadducts and involves multiple pathways (4, 6). In ICL repair, nuclease-dependent dual incisions on one strand flanking the lesion produce a gapped intermediate that must be filled by translesion synthesis (TLS) or homologous recombination (HR)

before NER-associated repair of the remaining unhooked monoadduct can occur.

Azinomycin B (AZB) (Fig. 1A) is a cytotoxic, nonribosomal peptide-polyketide secondary metabolite of *Streptomyces sahachiroi* and *Streptomyces griseofuscus* (7, 8). This bifunctional alkylating agent forms ICLs in vitro and in vivo (9) and displays potent antibiotic and antitumor activities comparable to the chemotherapeutic mitomycin C (8, 10, 11). AZB ICLs are formed between the electrophilic aziridine and epoxide functional groups of AZB and the N7 nitrogens of guanosine and adenosine within d(GNC) and d(GNT) sequences (Fig. 1B) (12, 13). Computational modeling of the AZB ICL structure is consistent with the entire AZB molecule residing in the major groove of the DNA without nucleobase intercalation by the naphthalate moiety (14, 15).

In contrast to NER-associated repair of other ICLs, AZB ICLs were recently identified as the target of repair by a DNA glycosylase (16). The *azi36* (*orf1*) gene sits adjacent to the AZB synthesis cluster in *S. sahachiroi* (17) and provides self-resistance to cells from AZB toxicity. The *orf1* gene product unhooks AZB ICLs by cleaving the *N*-glycosidic bond on at least one side of the lesion to produce an abasic site that is recognized by the bacterial AP endonuclease EndoIV (Fig. 1C) (16). Orf1 belongs to the HTH_42 (Pfam 06224) superfamily of uncharacterized winged helix (WH)-containing proteins that exist in various pathogenic and antimicrobial-producing bacteria. However, there are no known structures of any HTH_42 protein, and thus the molecular rationale for glycosylase repair of AZB ICLs is unclear.

The significance of this alternative, glycosylase-mediated ICL repair pathway is underscored by the recent discovery that the unrelated eukaryotic NEIL3 glycosylase unhooks psoralen and

Significance

DNA glycosylases are important repair enzymes that safeguard the integrity of the genome by excising chemically damaged DNA bases from the phosphoribose backbone. Recently, these enzymes were found to repair DNA interstrand cross-links (ICLs). ICLs are highly toxic DNA lesions formed by various bifunctional metabolites, environmental toxins, and chemotherapeutic agents that block normal DNA metabolism. This work describes the crystal structure of a newly discovered bacterial DNA glycosylase that repairs ICLs formed by azinomycin B, a potent antimicrobial and antitumor agent. The protein belongs to a structural superfamily prevalent in pathogenic bacteria and may serve as an important therapeutic target.

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The authors declare no conflict of interest.

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Data deposition: The atomic coordinates and structure factors have been deposited in the Protein Data Bank, www.pdb.org (PDB ID code 5UUJ).

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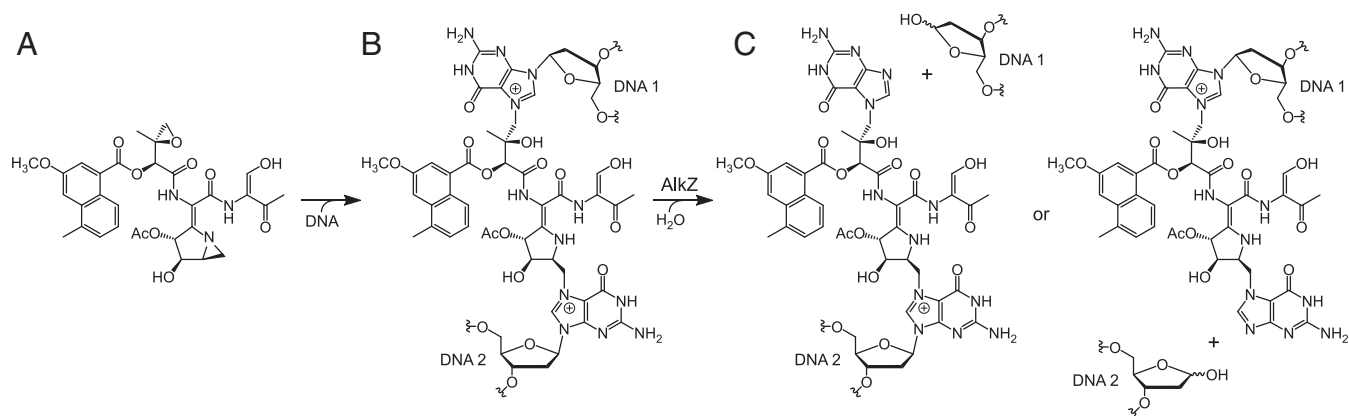


Fig. 1. Unhooking of AZB ICLs by AlkZ. (A) AZB. (B) AZB cross-links opposing guanines at their N7 nitrogens via reactive aziridine and epoxide groups. (C) AlkZ glycosylase catalyzes hydrolysis of the *N*-glycosidic bond of either cross-linked dG to produce two possible dG-AZB-Gua monoadducts.

abasic site-adenine ICLs during DNA replication (18), and that human NEIL1 can excise unhooked psoralen monoadducts and ICLs (19, 20). Thus, glycosylase-mediated ICL repair activity has been identified in both prokaryotes and eukaryotes by two distinct proteins. The molecular details of how these enzymes can excise adducts covalently tethered to both DNA strands is unknown, however. To better understand the mechanisms for DNA glycosylase-mediated ICL repair, we determined the crystal structure of the *S. sahachiroi azi36* gene product and identified the active site by mutational analysis. The structure is the defining member of the HTH_42 superfamily and an eighth structural class of DNA glycosylase. Our analysis shows how the HTH_42 architecture can support unhooking of either side of an AZB ICL or excision of an AZB-dG monoadduct. To better reflect the activity of the enzyme for base excision of alkylated DNA, we propose renaming the *Azi36/Orf1* protein “AlkZ.”

Results

AlkZ Excises N7-Methylguanine from DNA. Recently, the AlkZ protein was shown to act on AZB ICLs and monoadducts through hydrolysis of one or more of the *N*-glycosidic bonds (16), but whether the enzyme would act on adducts other than AZB was unclear. DNA glycosylases that excise alkylated DNA lesions often exhibit activity for a number of N3- and N7-methylated purines in addition to their preferred substrates (21, 22). For example, *Bacillus cereus* AlkD displays robust activity for bulky minor groove N3-yatakemycinyldeoxyadenosine lesions, but also cleaves N3-methyldeoxyadenosine and N7-methyldeoxyguanosine (d7mG) (23, 24). Therefore, we tested the ability of AlkZ to catalyze excision of N7-methylguanine (7mGua) from an oligonucleotide containing d7mG (Fig. 2A).

Using a standard *in vitro* assay that measures glycosylase activity by alkaline hydrolysis of the resulting abasic site product, we found that AlkZ hydrolyzed d7mG with an observed rate (k_{obs}) of $(8.5 \pm 0.4) \times 10^{-5} \text{ s}^{-1}$ (Fig. 2B and C). This d7mG activity is modest compared with that of other alkylpurine DNA glycosylases under similar conditions (21, 25, 26), but demonstrates that an AZB adduct is not necessary for AlkZ activity and corroborates the previous data showing activity toward monoadducts (16). In addition, we found that treatment with NaOH or EndoIV is required to nick the abasic DNA produced by AlkZ, indicating that AlkZ is a monofunctional DNA glycosylase that does not have lyase activity (Fig. 2D).

AlkZ Adopts a Unique DNA-Binding Fold. HTH_42 superfamily proteins are predicted to contain two WH motifs, which are common helix-turn-helix (HTH)-related DNA-binding elements found in transcription factors and DNA processing enzymes (27).

The canonical WH fold contains a three-helix bundle at the N terminus and a three-stranded β -sheet (wing) at the C terminus, and is related to the HTH sequence-specific DNA-binding motif in that $\alpha 3$ serves as a “recognition helix” by inserting into the DNA major groove (28). However, because there is no known structure of an HTH_42 protein, how or if the putative WH motifs bind DNA remains unclear. Moreover, there are no WH motifs in any of the seven known DNA glycosylase protein folds (29), so the molecular details of how AlkZ acts as a DNA repair enzyme are also unclear.

To understand the structural basis for DNA glycosylase-mediated ICL repair by AlkZ, we determined the crystal structure of the *S. sahachiroi* enzyme. Experimental X-ray phases were determined from single-wavelength anomalous dispersion using selenomethionyl-substituted protein (Fig. S1). The resulting crystallographic model, which contains all 371 residues of the protein, was refined against X-ray diffraction data extending to 2.3 Å, resulting in an R factor of 15.7% and an R_{free} value of 20.1% (Table S1). The protein adopts a cluster of three tandem WH motifs (WH1, WH2, and WH3) that pack against a C-terminal β -barrel subdomain to form a C-shaped molecule (Fig. 3). WH1 is centrally located and bridges the WH2 and WH3 motifs. WH1 packs against WH2 with twofold rotational symmetry and their β -wings at the interface, and against WH3 in a 90°-offset head-to-tail arrangement. This

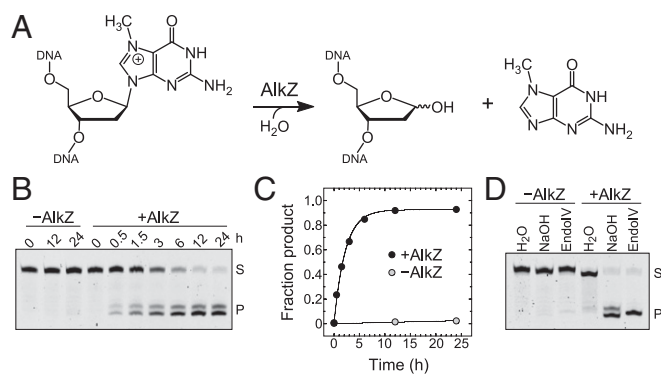


Fig. 2. Excision of N7-methylguanine by AlkZ. (A) Base excision of d7mG by AlkZ. (B) Denaturing PAGE of d7mG-DNA substrate (S) and nicked abasic-DNA product (P) after treatment without (-) or with (+) AlkZ for the specified time, followed by alkaline hydrolysis. (C) Quantitation of the gel shown in B, averaged across three independent experiments. Error bars represent SD. (D) Denaturing PAGE of d7mG-DNA substrate (S) and nicked abasic-DNA product (P) after treatment without (-) or with (+) AlkZ for 24 h, followed by treatment with water, NaOH, or EndoIV.

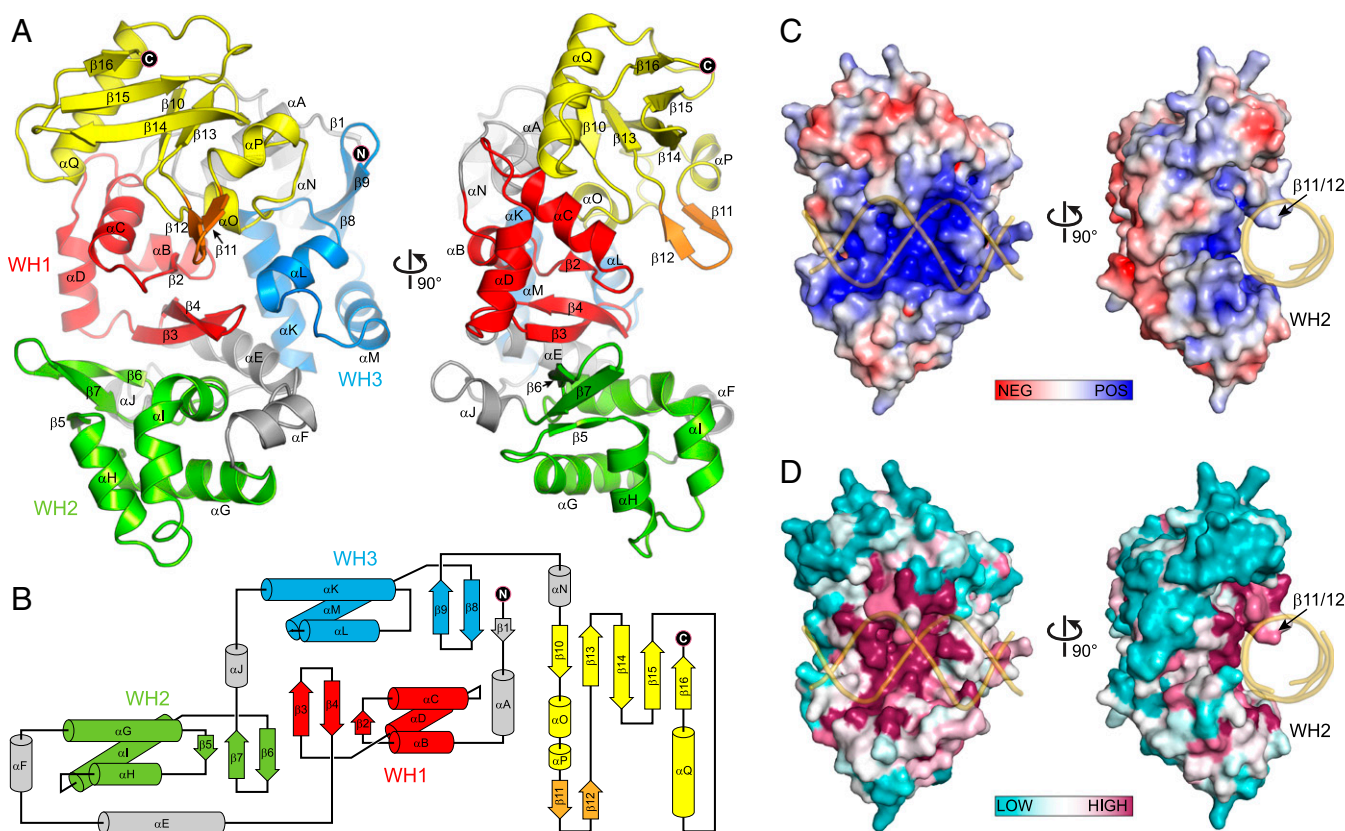


Fig. 3. Crystal structure of AlkZ. (A) Orthogonal views of the enzyme colored by subdomain. WH motifs are in red, green, and blue; the C-terminal domain is in yellow; the putative DNA-binding $\beta 11/12$ loop is in orange; and connecting helices are in gray. (B) Topology diagram color-coded as in A. (C) Solvent-accessible surface of AlkZ colored by electrostatic potential (-7 to $+7 k_B T/e_c$). (D) Solvent-accessible surface of AlkZ colored by sequence conservation. In C and D, B-DNA modeled against the crystal structure is shown as a transparent gold backbone trace.

cluster of WH motifs appears to be a unique protein architecture, as a search for structural homologs using the DALI server (30) returned no results aside from those of each individual WH (Fig. S2). Consistent with the established structural diversity among WH motifs, the three motifs in AlkZ show slight variations from one another (Fig. S2), including a highly kinked $\alpha 3$ “recognition helix” in WH2 (helix αI). However, with the exception of WH3, which lacks the short $\beta 1$ strand, each WH adopts the canonical $\alpha 1$ - $\beta 1$ - $\alpha 2$ - $\alpha 3$ - $\beta 2$ - $\beta 3$ topology.

The most notable feature of the C-shaped structure is the concave channel that spans the width of the protein. The channel is defined by all three WH motifs and a short β -hairpin ($\beta 11/12$) that protrudes from the C-terminal domain. The surface of this channel is lined with positively charged, highly conserved residues and is appropriately sized to accommodate duplex DNA (Fig. 3 C and D). DNA bound along this channel would be clamped between the $\beta 11/12$ hairpin on one side of the duplex and helix αI —the putative “recognition helix” of WH2—on the other side, indicating that WH2 may play a role in DNA recognition similar to that of other WH motifs. In contrast, the corresponding helices from WH1 (αD) and WH3 (αM) reside outside the central channel in positions not likely to bind DNA. Indeed, docking DNA from other WH-DNA and HTH-DNA complexes onto WH1 or WH3 positioned the DNA well outside of the putative DNA-binding channel.

Active Site Residues Reside Within the Putative DNA-Binding Channel.

To gain insight into how AlkZ properly orients an AZB lesion for catalysis, we docked a B-DNA model containing a computationally derived AZB ICL (14) as a rigid body into the putative DNA-

binding channel (Fig. 4 A–D). Two nearly equivalent DNA orientations, related by the symmetric nature of the AZB ICL relative to the DNA dyad axis, optimized van der Waals and electrostatic interactions without altering the structure of the protein or the DNA. Both orientations placed the $\beta 11/12$ hairpin into the minor groove directly across from the AZB ICL and placed WH2 against the adduct in the major groove.

We used these models to identify candidate catalytic residues among the highly conserved polar, cationic, aromatic, and hydrophobic side chains lining the concave surface (Fig. S3A), and thus likely in the vicinity of a bound DNA lesion. Monofunctional glycosylases generally initiate nucleophilic attack from the minor groove using carboxylate (Asp, Glu) or carboxamide (Asn, Gln) side chains (31, 32). In AlkZ, side chains from conserved glutamines Q37 and Q39 (Fig. S4) point into the minor groove at either point of ICL attachment in both orientations of our docking model (Fig. 4 A–D). Whereas Q37 is recessed at the rear of the channel, Q39 protrudes far enough into the minor groove to orient a water nucleophile for catalysis.

We tested involvement of the helix-penetrating QxQ motif and $\beta 11/12$ hairpin in enzyme activity by substituting with alanine or glycine and measuring d7mG excision activity from purified proteins (Fig. 4E). We verified by circular dichroism that all mutant proteins were folded under the conditions of our assay (Fig. S3 B and C). Both Q37A and Q39A mutants completely abrogated d7mG excision, even after 24 h. In contrast, alanine substitution of the nearest glutamate side chains, E45 and E221, which reside just outside the putative active site on helices αC (WH1) and αL (WH3), respectively, had no effect on base excision activity. In addition to catalysis by polar residues, DNA

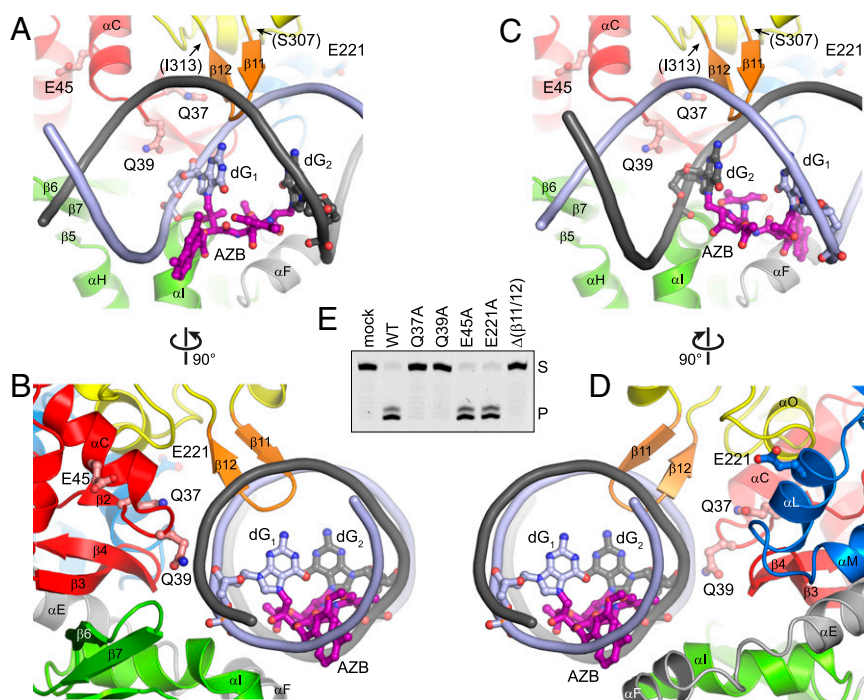


Fig. 4. Identification of the AlkZ active site. (A and B) Orthogonal views of the AlkZ active site with AZB-ICL-DNA docked in the putative DNA-binding channel. The protein is color-coded as in Fig. 3, and AlkZ residues tested for their roles in base excision activity are labeled. The AZB-ICL-DNA model is shown as a slate/charcoal backbone trace with adducted guanines (slate/charcoal) and AZB (purple) depicted in ball-and-stick representation. (C and D) Orthogonal views of AlkZ bound to AZB-ICL-DNA in a hypothetical alternate binding orientation. (E) Base excision activity of wild-type (WT) and mutant AlkZ proteins as measured by the formation of nicked abasic-DNA product (P) after incubation of protein or buffer (mock) with d7mG-DNA substrate (S) for 24 h.

glycosylases often stabilize a particular DNA conformation in the enzyme-substrate complex by inserting a steric wedge—either bulky side chains or a loop—into the DNA helix. In AlkZ, this role is likely played by the $\beta 11/12$ hairpin, which protrudes into the minor groove in our docking model (Fig. 4A–D). Consistent with this hypothesis, replacing residues 307–313 that form the $\beta 11/12$ loop with two glycine residues ($\Delta\beta 11/12$) abrogated d7mG excision activity (Fig. 4E). These results identify this central region of the protein as the active site, with important roles for Q37, Q39, and the $\beta 11/12$ hairpin in base excision.

Discussion

This work defines the architecture of the HTH₄₂ family of DNA-binding proteins (33) and an eighth class of DNA glycosylase (29, 34). A unique cluster of three tandem WH motifs scaffold a putative DNA-binding channel, with only one WH (WH2) positioned to participate in the canonical helix-major groove recognition used by other WH and HTH motifs (28). The active site contains a QxQ motif and a β -hairpin essential for base excision activity and conserved among AlkZ homologs and the HTH₄₂ superfamily (Fig. S4). Our docking models predict insertion of the $\beta 11/12$ hairpin into the DNA minor groove at the lesion site. This motif is likely involved in stabilizing a particular DNA conformation in the enzyme-substrate complex, similar to that observed for β -hairpins and loops in other glycosylases (29). The N-terminal glutamine (Q37) in the QxQ motif is recessed in the active site cleft and thus predicted to bind the DNA backbone, consistent with histidine occupying this position in some AlkZ homologs. The C-terminal glutamine (Q39), on the other hand, likely plays a catalytic role. Our docking models predict that Q39 protrudes into the minor groove so that its carboxamide side chain is positioned to orient a water nucleophile in the manner observed for N140 in human thymine DNA glycosylase (32) and for aspartate or glutamate side chains in other glycosylases (31). Consistent with such a catalytic role for Q39, approximately one-half of all HTH₄₂ proteins have an aspartate at this position, suggesting that at least some other members of this previously uncharacterized superfamily also have DNA glycosylase activity.

AlkZ is one of two DNA glycosylases that unhook an ICL (16, 18). The mechanism by which a DNA glycosylase unhooks an ICL is not immediately obvious, because these enzymes typically capture the modified nucleoside inside the active site by extruding it from the DNA helix, and such a base-flipping mechanism would be inhibited by an ICL. However, we recently discovered a non-base-flipping mechanism that enables glycosylase excision of bulky minor groove adducts and conceivably could enable ICL excision. The bacterial AlkD glycosylase is able to recognize and cleave deoxyadenosine adducts of the bulky natural product yatagemycin (YTM) without rotating the lesion from the duplex (24). Like AlkZ, AlkD adopts a C-shaped fold that engages DNA along the concave surface (22). The AlkD active site positions a catalytic aspartate and two catalytic tryptophan side chains (25) against the deoxyribose of the alkylated adenosine, which is displaced only slightly into the minor groove of the DNA. Similarly, in our AlkZ docking models, the QxQ motif is positioned sufficiently far into the minor groove to access the *N*-glycosidic bonds of the ICL without rotating the cross-linked nucleosides (Figs. 4 and 5). The eukaryotic DNA glycosylase with ICL unhooking activity, NEIL3, also has an active site (35) that conceivably could access an unflipped lesion.

Previous work has established that AlkZ can unhook an AZB ICL, but whether the enzyme cleaves both sides is less clear (16). Although full ICL excision by a DNA glycosylase would create opposing abasic sites susceptible to a deleterious double-strand break, it also would simplify repair by eliminating the need for NER or TLS/HR pathways to remove the remaining monoadduct. Pol I and II bypass of abasic sites and major groove monoadducts (36, 37) suggests that, in principle, BER could fully process opposing abasic sites in bacteria. Several lines of evidence suggest that AlkZ can excise the entire cross-link by one of two mechanisms, either by sequentially unhooking the ICL and excising the remaining monoadduct or by simultaneously cleaving the glycosidic bonds of both modified dG residues. First, AlkZ has both unhooking activity for ICLs and excision activity for monoadducts (Fig. 2 and ref. 16). Second, our DNA docking models illustrate that AlkZ can bind to either side of the AZB ICL (Fig. 5A and B). Both binding orientations position the putative active site against the minor groove on either side of the ICL

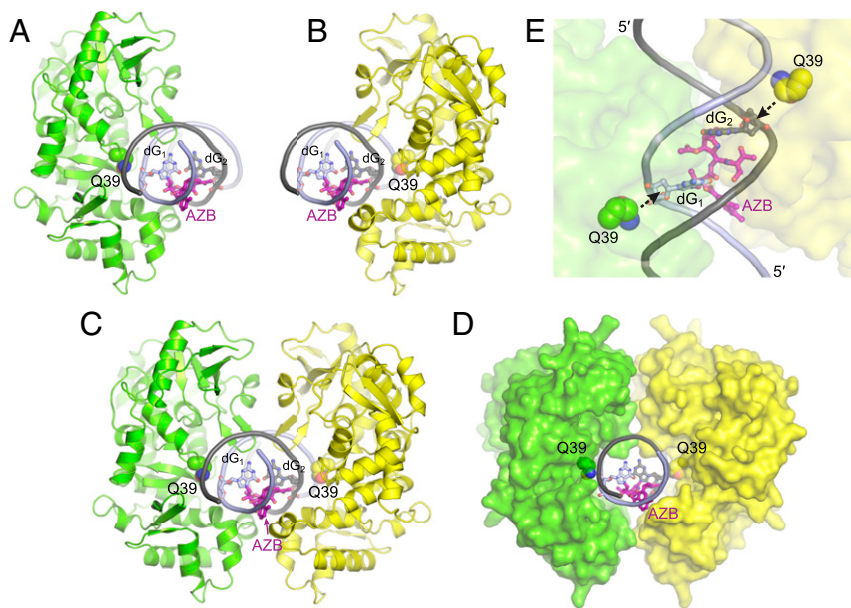


Fig. 5. Possible mechanisms of ICL repair by AlkZ. (A and B) Two binding orientations of AlkZ against each side of an AZB ICL. The catalytic Q39 side chain is shown in green or yellow, and the AZB ICL is shown in purple. (C and D) Dimeric model of two AlkZ subunits bound to one AZB ICL. The proteins are shown as green and yellow cartoons (C) or solvent-accessible surfaces (D). (E) Top view of the hypothetical AlkZ dimer interface, showing simultaneous engagement of both cross-linked dG residues by the catalytic Q39 side chain.

without obstruction from AZB. Third, our docking models show that the two binding orientations can coexist at the same ICL. Modeling two AlkZ molecules on opposite sides of the DNA with their respective active sites against the two cross-linked dG residues reveals a remarkable complementarity between protein surfaces (Fig. 5 C–E). Although we have no evidence of dimerization of AlkZ in the presence of d7mG-DNA, dimerization may be dependent on the AZB ICL. The recent discovery of a dimeric DNA glycosylase acting as part of a restriction-modification system in thermophilic bacteria (34) raises the possibility that AlkZ could use a similar strategy to simultaneously excise the entire cross-link. However, at the lower temperatures at which AlkZ functions, spontaneous formation of a double-strand break from opposing abasic sites would be less likely. Moreover, the bound AlkZ dimer may transiently shelter both abasic sites from hydrolysis until further repair by enzymes in the BER pathway occurs.

Pathogenic and antibiotic-producing microbes have resistance mechanisms against their own toxins that often are genetically clustered with the antibiotic synthesis operon (38–40). AlkZ is the second DNA glycosylase discovered to act as a toxin resistance mechanism (16). The AlkD homolog YtkR2 is produced in *Streptomyces* sp. TP-A0356 as a means to protect against its natural product YTM, a bulky genotoxin of the spirocyclopropylcyclohexadienone family (41, 42). Similar to the AlkZ gene (*azi36/orf1*), *ytkR2* and putative BER genes are embedded within the YTM synthesis gene cluster. Although AlkZ and YtkR2 likely originated in AZB- and YTM-producing organisms, homologous proteins are evident in diverse bacteria, including human pathogens (16, 43). Thus, the two bacterial DNA glycosylases known to repair bulky and cross-linked DNA damage arising from potent bacterial toxins have transferred among bacterial populations. This implicates DNA repair and alternative mechanisms thereof, in possible treatments against antibiotic resistance, and opens the possibility that other DNA repair mechanisms and genotoxic metabolites await discovery in pathogenic and antibiotic-producing bacteria.

Methods

Protein Purification. The *azi36* gene was amplified by PCR from genomic DNA isolated from *S. sahachiroi* (American Type Culture Collection; 33158) and

cloned into pBG102 (Vanderbilt University Center for Structural Biology). His₆-SUMO-AlkZ was overexpressed in *Escherichia coli* Tuner(DE3) cells at 16 °C for 18 h in LB medium supplemented with 100 µg/mL kanamycin and 50 µM isopropyl β-D-1-thiogalactopyranoside (IPTG). Cells were lysed with gentle sonication, and cell debris was removed by centrifugation at 45,000 × g at 4 °C for 30 min. Clarified lysate was passed over 5 mL of Ni-NTA agarose resin equilibrated in buffer A [50 mM Tris-HCl pH 8.5, 500 mM NaCl, and 10% (vol/vol) glycerol]. The column was washed with 100 mL of 20 mM imidazole/buffer A, and protein was eluted in 50 mL of 250 mM imidazole/buffer A. Protein fractions were pooled and supplemented with 0.1 mM EDTA and 1 mM tris(2-carboxyethyl)phosphine (TCEP) before incubation with ~0.5 mg of Rhinovirus 3C protease (PreScission) at 4 °C overnight. Cleaved protein was diluted 10-fold in buffer B [50 mM Tris-HCl pH 8.5, 10% (vol/vol) glycerol, 1 mM TCEP, and 0.1 mM EDTA]. Diluted protein was loaded onto a 5-mL heparin Sepharose column, washed with 25 mL of 50 mM NaCl/buffer B, and eluted with 50 mL of a 50–1,000 mM NaCl/buffer B linear gradient. Fractions were pooled and re-passed over Ni-NTA agarose resin equilibrated in buffer A. Flow-through and protein that eluted with 50 mL of 20 mM imidazole/buffer A were pooled, concentrated and filtered, and passed over Superdex 200 size exclusion resin equilibrated in buffer C [20 mM Tris-HCl pH 8.5, 100 mM NaCl, 5% (vol/vol) glycerol, 1 mM TCEP, and 0.1 mM EDTA]. Protein was eluted with 1.5 CV of buffer C, concentrated to 6 mg/mL, flash-frozen in liquid nitrogen, and stored at –80 °C.

AlkZ mutants were generated using the Q5 Mutagenesis Kit (New England BioLabs). The Δ(β11/12) mutant was generated by replacing residues 307–SRAAGWI-313 with two glycine residues. Mutant proteins were overexpressed and purified the same as wild-type (WT). The structural integrity of mutant proteins was verified by thermal denaturation using far-UV circular dichroism spectroscopy, in which molar ellipticity at 222 nm was monitored using a Jasco J-810 spectropolarimeter equipped with a Peltier temperature controller as a solution containing 5 µM enzyme, 50 mM Hepes pH 8.5, 100 mM KCl, and 10% (vol/vol) glycerol was heated at 1 °C/min in a 0.1-cm cell (Fig. S3 B and C). Selenomethionyl-substituted AlkZ (SeMet-AlkZ) was overexpressed and purified the same as the WT protein, except that protein was overexpressed in M9 medium supplemented with 0.4% (wt/vol) dextrose, 1 mM MgSO₄, 0.1 mM CaCl₂, 1 mg/L thiamine, 70 mg/L selenomethionine, 50 mg/L leucine, 50 mg/L isoleucine, 50 mg/L valine, 100 mg/L phenylalanine, 100 mg/L lysine, 100 mg/L threonine, 100 µg/mL kanamycin, and 50 µM IPTG.

Crystallization, X-Ray Data Collection, and Structure Refinement. SeMet-AlkZ was crystallized by sitting drop vapor diffusion at 21 °C by mixing equal volumes of 2 mg/mL protein and reservoir solution containing 18% (wt/vol) PEG 8000, 10 mM Tris-HCl pH 7.5, and 10 mM MgCl₂. Crystals were harvested

4 d after setting the drops, cryoprotected in reservoir solution supplemented with 15% (vol/vol) glycerol, and flash-frozen in liquid nitrogen. X-ray diffraction data from a single frozen crystal were collected at a wavelength of 0.97857 Å at the Advanced Photon Source Beamline 21-ID-G at Argonne National Laboratory and processed with HKL2000 (44). Phasing and refinement were carried out using the PHENIX suite of programs (45). Phases were calculated from the positions of eight Se atoms and modified by solvent flattening using PHENIX AutoSol. A partial model consisting of 347 of 371 SeMet-AlkZ residues was built automatically by PHENIX AutoBuild; the remaining residues were built manually in Coot (46). After adding riding hydrogens with PHENIX ReadySet, atomic coordinates and TLS-derived anisotropic B-factors were refined using PHENIX Refine. The final model was validated with MolProbity (47) and contained no residues in the disallowed regions of the Ramachandran plot. Refinement and validation statistics are presented in Table S1. Atomic coordinates and structure factors were deposited in the Protein Data Bank (PDB) under accession code 5UUJ. The AlkZ-DNA model was prepared by extending the previously determined AZB-ICL-DNA model (14) with three B-form base pairs on each end, followed by manual docking onto the AlkZ structure using PyMOL. All structural biology software was curated by SBGrid (48).

Base Excision Assay. DNA substrates containing a single d7mG lesion and a 5'-6-carboxyfluorescein (FAM) label within the sequence FAM-d(CACCACTAC-ACC(7mG)ATTCCTTACAAC)/d(GTTGTAAGGAATCGGTGTAGTGGTG) were prepared as described previously (23). In 60- μ L reactions, 10 μ M AlkZ was incubated

with 100 nM FAM-DNA in glycosylase buffer [50 mM Hepes pH 8.5, 100 mM KCl, 10 mM EDTA, and 10% (vol/vol) glycerol] at 25 °C. At various times, 8- μ L aliquots were added to 2 μ L of 1 M NaOH and heated at 70 °C for 2 min. To test for AP lyase activity, the 8- μ L aliquots were added to 2 μ L of glycosylase buffer, 1 M NaOH, or *E. coli* EndoIV (New England Biolabs). Samples were denatured at 70 °C for 5 min in 5 mM EDTA pH 8.0, 40% (wt/vol) formamide, 0.5 mg/mL bromophenol blue, and 0.5 mg/mL xylene cyanol, and electrophoresed on a 20% (wt/vol) acrylamide/8 M urea sequencing gel at 20 W for 2 h in 0.5 \times TBE buffer (45 mM Tris, 45 mM borate, and 1 mM EDTA pH 8.0). Gels were imaged on a Typhoon Trio variable mode imager (GE Healthcare) using 532-nm excitation and 526-nm emission fluorescence, and bands were quantified with ImageQuant (GE Healthcare). Reactions were performed in triplicate.

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