

DNA damage recognition and repair by 3-methyladenine DNA glycosylase I (TAG)

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DNA glycosylases help maintain the genome by excising chemically modified bases from DNA. *Escherichia coli* 3-methyladenine DNA glycosylase I (TAG) specifically catalyzes the removal of the cytotoxic lesion 3-methyladenine (3mA). The molecular basis for the enzymatic recognition and removal of 3mA from DNA is currently a matter of speculation, in part owing to the lack of a structure of a 3mA-specific glycosylase bound to damaged DNA. Here, high-resolution crystal structures of *Salmonella typhi* TAG in the unliganded form and in a ternary product complex with abasic DNA and 3mA nucleobase are presented. Despite its structural similarity to the helix–hairpin–helix superfamily of DNA glycosylases, TAG has evolved a modified strategy for engaging damaged DNA. In contrast to other glycosylase–DNA structures, the abasic ribose is not flipped into the TAG active site. This is the first structural demonstration that conformational relaxation must occur in the DNA upon base hydrolysis. Together with mutational studies of TAG enzymatic activity, these data provide a model for the specific recognition and hydrolysis of 3mA from DNA.

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Introduction

The information encoded within the sequence and structure of DNA is vital to the survival of any organism. The integrity of the genome is constantly threatened by the chemical reactivity of the nucleobases, which are modified by a variety of alkylation, oxidation or radiative processes (reviewed in Friedberg *et al*, 2006). DNA alkylation by cellular metabolites, environmental toxins, or chemotherapeutic agents produces a wide spectrum of aberrant nucleotides that are cytotoxic or mutagenic, and hence can lead to cell death and heritable disease. A large number of alkylated purines,

including cytotoxic 3-methyladenine (3mA), 7-methylguanine (7mG), and the highly mutagenic lesion 1,N⁶-etheno-adenine (ϵ A), have been detected in humans after exposure to various carcinogens (Shuker *et al*, 1987). As a safeguard against alkylation damage, cells have devised a number of DNA repair strategies to remove these modifications and restore the DNA to an undamaged state. The base excision repair pathway is the principal mechanism by which alkyl-purines are eliminated from the genome. DNA glycosylases initiate this pathway by locating and removing a specific type of modified base from DNA through cleavage of the C1'–N glycosylic bond.

Alkylpurine DNA glycosylases have been shown to be essential for the survival of both eukaryotic and prokaryotic organisms (Clarke *et al*, 1984; Chen *et al*, 1989), and have been identified in humans, yeast, and bacteria. Among these are *Escherichia coli* 3mA DNA glycosylase I (TAG) and II (AlkA), *Thermotoga maritima* methylpurine DNA glycosylase II (MpgII), *Helicobacter pylori* 3mA DNA glycosylase (MagIII), yeast methyladenine DNA glycosylase (MAG), and human alkyladenine DNA glycosylase (AAG) (Riazuddin and Lindahl, 1978; Thomas *et al*, 1982; Chen *et al*, 1990; O'Connor and Laval, 1991; Begley *et al*, 1999; O'Rourke *et al*, 2000). Although structurally unrelated, the human and bacterial alkylpurine glycosylases have evolved a common base-flipping mechanism for gaining access to damaged nucleobases in DNA (reviewed in Roberts and Cheng, 1998; Hollis *et al*, 2000b). The bacterial enzymes TAG, AlkA, and MagIII belong to the helix–hairpin–helix (HhH) superfamily of DNA glycosylases (Labahn *et al*, 1996; Nash *et al*, 1996; Drohat *et al*, 2002; Eichman *et al*, 2003). The HhH motif is used by hundreds of repair proteins for binding DNA in a sequence-independent manner (Doherty *et al*, 1996). Crystal structures of HhH glycosylases AlkA, hOgg1, EndoIII, and MutY in complex with DNA illustrate how the HhH motif is used as a platform for base flipping to expose damaged bases in DNA (Bruner *et al*, 2000; Hollis *et al*, 2000a; Fromme and Verdine, 2003b; Fromme *et al*, 2004).

Alkylpurine DNA glycosylases from bacteria have widely varying substrate specificities despite their structural similarity. TAG and MagIII are highly specific for 3mA (Bjelland *et al*, 1993; O'Rourke *et al*, 2000), whereas AlkA is able to excise 3mA, 7mG, and other alkylated or oxidized bases from DNA (McCarthy *et al*, 1984; Bjelland *et al*, 1994; Saparbaev *et al*, 1995). The importance of specificity during base excision is underscored by the fact that glycosylases must identify subtle alterations in base structure amidst a vast excess of normal DNA. Recognition of the substrate base must occur at two steps—interrogation of the DNA duplex during a processive search and direct read-out of the target base that has been flipped into the active site of the enzyme (Stivers and Jiang, 2003; Banerjee *et al*, 2006). Our structural understanding of 3mA processing by bacterial alkylpurine DNA glycosylases is currently limited to structures of TAG and MagIII bound to alkylated bases in the absence of DNA. Crystal structures of

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MagIII bound to 3mA and ϵ A revealed that direct contacts to nucleobase substituent atoms are not necessary for binding alkylpurines in the binding pocket (Eichman *et al*, 2003). NMR studies of *E. coli* TAG bound to 3mA demonstrated that TAG makes specific contacts to the base, and that the enzyme lacks the hallmark catalytic aspartic acid present in all other HhH glycosylases (Nash *et al*, 1996; Drohat *et al*, 2002; Cao *et al*, 2003). Given the lack of DNA in these structures, the mechanism by which specific 3mA glycosylases locate and excise their target bases from DNA is currently a matter of speculation.

Presented here are the crystal structures of *Salmonella typhi* TAG alone and in complex with abasic DNA and 3mA, together with mutational studies of TAG enzymatic activity. TAG binds damaged DNA in a manner similar to other HhH glycosylases, but uses a different strategy to intercalate the DNA in order to gain access to the damage site. Surprisingly, the abasic ribose adopts two specific conformations, neither of which is fully flipped into the active site pocket as has been observed in all other glycosylase product complexes. Extensive interactions with the bases on

both DNA strands provide a structural rationale for how TAG detects 3mA lesions within DNA. Inside the base binding pocket, a conserved glutamic acid has been identified to play a significant role in catalysis of base excision. A comparison of structures of HhH alkylpurine DNA glycosylases provides a basis for understanding the unique mechanisms by which 3mA is selected and removed from DNA.

Results and discussion

TAG from the bacterium *S. typhi* is 82% identical and 91% conserved overall to the *E. coli* enzyme. *S. typhi* TAG was crystallized alone and in complex with 3mA base and DNA containing a tetrahydrofuran (THF) abasic site analog. The THF moiety is isosteric with enzymatically generated apurinic sites, but is not prone to ring opening owing to the lack of a C1' hydroxyl group (Takeshita *et al*, 1987). The crystal structures of TAG and the TAG/THF-DNA/3mA complex were determined using experimental phases from multi- and single-wavelength anomalous dispersion (MAD, SAD) experiments, respectively (Table I). A crystallographic model of the

Table I Data collection, phasing, and refinement statistics

	Unliganded TAG				TAG/THF-DNA/3mA
	Native	Se peak	Se remote	Se inflection	Se peak
<i>Cell parameters</i>					
<i>a</i>	57.5 Å	57.5 Å			102.0 Å
<i>b</i>	63.7 Å	63.9 Å			102.0 Å
<i>c</i>	62.1 Å	62.1 Å			55.5 Å
α	90°	90°			90°
β	106.9°	107.0°			90°
γ	90°	90°			120°
Space group	P2 ₁	P2 ₁			P6 ₃
Molecules/asu	2	2			1
<i>Data collection^a</i>					
Wavelength (Å)	1.000	0.9793	0.9718	0.9795	0.9793
Resolution (Å)	50–1.5	50–1.5	50–1.5	50–1.5	50–1.85
Unique reflections	64 266 (4195)	121 469 (7253)	125 767 (8960)	122 660 (7290)	54 963 (5415)
Completeness (%)	93.6 (61.1)	89.7 (53.4)	92.8 (65.8)	90.4 (53.5)	99.6 (97.9)
Redundancy	7.0 (4.6)	3.7 (2.6)	3.8 (2.9)	3.7 (2.6)	6.4 (5.3)
R_{sym}^b	0.064 (0.247)	0.089 (0.185)	0.097 (0.176)	0.095 (0.192)	0.053 (0.433)
$\langle I \rangle / \langle \sigma(I) \rangle$	23.8 (4.8)	17.6 (4.1)	17.2 (4.6)	17.5 (3.8)	27.2 (3.4)
<i>Phasing</i>					
Resolution (Å)		50–1.5	50–1.5	50–1.5	50–1.85
Phasing power ^c		1.41	1.55	—	—
R_{cullis} (c/a) ^d		0.63/0.69	0.59/0.65	—	—
R_{cullis} (anom)		0.60	0.73	0.65	0.77
<i>Refinement</i>					
Resolution	50–1.5				50–1.85
No. of reflections	60 990				26 778
R_{cryst}^e	0.161				0.175
R_{free}^e	0.196				0.198
Number of atoms					
Protein	2903				1450
DNA	0				518
Solvent	399				160
Average <i>B</i> -factor (Å ²)	31.6				34.3
R.m.s.d. bonds (Å)	0.014				0.018
R.m.s.d. angles (deg)	1.427				1.868

^aValues in parentheses refer to data in the highest resolution shell.

^b $R_{\text{sym}} = \sum_{hkl} \sum_j |I_j - \langle I \rangle| / \sum_{hkl} \sum_j I_j$ where $\langle I \rangle$ is the mean intensity of *j* observations of reflection *hkl* and its symmetry equivalents.

^cPhasing power = $\sum_{hkl} F_H / \sum_{hkl} |F_{PH} - F_{PH,calc}|$. Value shown is the weighted average of centric and acentric data.

^d $R_{\text{cullis}} = (\sum_{hkl} |F_{PH} \pm F_P| - F_{H,calc}) / \sum_{hkl} |F_{PH} - F_P|$. *c/a*, centric/acentric data.

^e $R_{\text{cryst}} = \sum_{hkl} |F_{\text{obs}} - kF_{\text{calc}}| / \sum_{hkl} |F_{\text{obs}}|$. $R_{\text{free}} = R_{\text{cryst}}$ for 5% of reflections that were not used in refinement (Brünger, 1992).

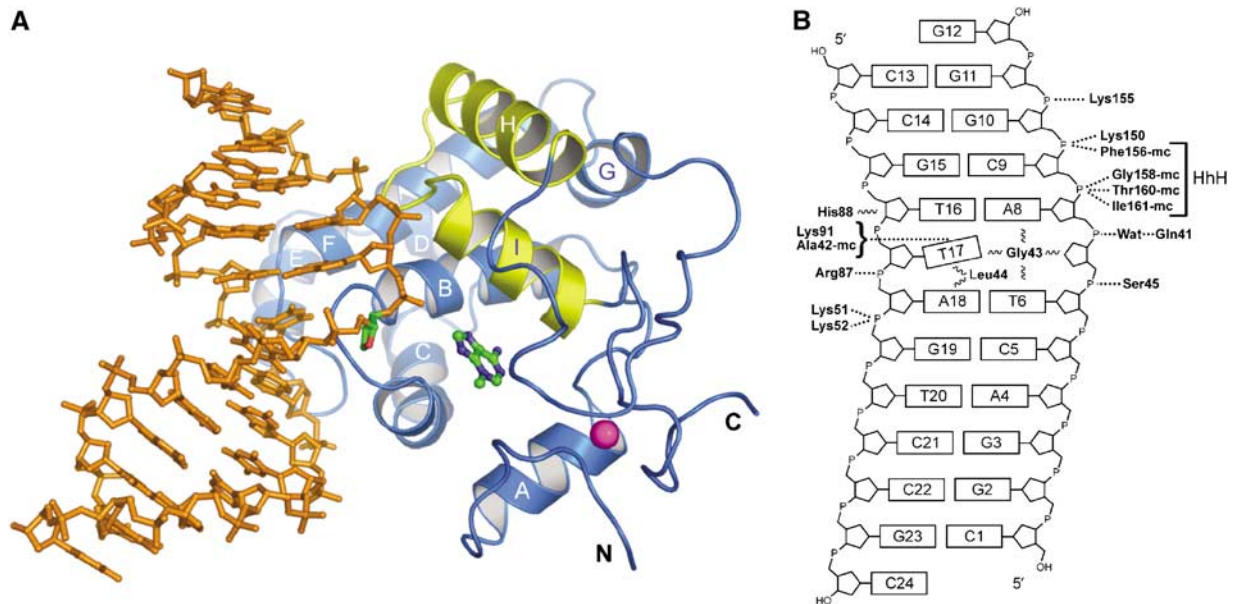


Figure 1 The structure of the TAG–DNA complex. **(A)** The crystallographic model of TAG (blue ribbons; yellow HhH motif) bound to DNA (orange sticks) and 3mA (ball-and-stick). The abasic site in the DNA is highlighted in green and the coordinated Zn^{2+} ion is shown as a magenta sphere. **(B)** Schematic representation showing the electrostatic (dashed lines) and van der Waals (wavy lines) interactions between protein side-chain and main-chain (mc) atoms with the DNA.

free protein, which consists of two TAG molecules in the asymmetric unit, was built into 1.5-Å MAD electron density (Supplementary Figure S1) and refined to a crystallographic residual of 0.161 ($R_{free} = 0.196$). Likewise, the model of the TAG/THF–DNA/3mA product complex (Figure 1) was built into 1.85-Å SAD experimental electron density (Supplementary Figure S1) and refined to a crystallographic residual of 0.175 ($R_{free} = 0.198$).

The crystal structures of *S. typhi* TAG are consistent with NMR structures of the *E. coli* enzyme that identified TAG as a member of the HhH superfamily of DNA glycosylases (Drohat *et al*, 2002). TAG adopts a globular fold consisting of an α -helical domain that contains the HhH motif (helices H and I) and a second, unique Zn^{2+} -binding domain that tethers the N- and C-termini (Figure 1A) (Kwon *et al*, 2003). The 3mA binding pocket is located at the interface between the two domains (Figure 1A) (Cao *et al*, 2003). Superposition of the *S. typhi* (crystal) and *E. coli* (NMR) structures shows that the protein backbones and positions of bound 3mA are virtually identical (with an r.m.s. deviation of 1.8 Å for all main-chain atoms; Supplementary Figure S2). Surprisingly, the largest differences between the two structures occur in the positions of two conserved tryptophan side chains in the 3mA binding pocket. Each of the indole rings of Trp 6 and Trp 21 are rotated $\sim 120^\circ$ between the two models (Supplementary Figure S2). Based on the high degree of sequence and structural conservation between *S. typhi* and *E. coli* TAG, these differences are likely an artifact of structure determination and not inherent differences between the two orthologs.

DNA binding by TAG

The HhH glycosylases use a common mechanism for binding DNA. These proteins anchor both strands of the DNA duplex from the minor groove side through van der Waals and polar interactions with the bases and the phosphate backbone. Main-chain atoms from the HhH hairpin form hydrogen

bonds with two phosphate groups immediately 3' to the lesion, whereas positively charged side chains from a conserved protein loop engage the non-lesioned strand. An intercalating side chain occupies (or 'plugs') the gap in the DNA left by the flipped-out nucleotide, and a second side chain wedges into the non-lesioned DNA opposite the flipped-out nucleotide. Collectively, these interactions stabilize a 60–70° bend in the duplex and help the protein gain access to the modified base.

TAG binds DNA similarly to other HhH glycosylases (Bruner *et al*, 2000; Hollis *et al*, 2000a; Fromme and Verdine, 2003b; Fromme *et al*, 2004), with subtle unique differences that categorize TAG as a divergent member of the superfamily and that likely result in its high specificity for positively charged 3mA bases. The DNA is anchored to the protein by three hairpin loops formed from helices B/C, E/F, and the HhH motif (Figure 1A). Basic side-chain and main-chain atoms from the HhH motif bind the phosphate groups 3' to the abasic site, whereas basic residues from the E/F loop contact the DNA backbone on the non-lesioned strand (Figure 1B). The loop between helices B and C inserts into the abasic gap in the DNA duplex, and the details will be discussed below. The DNA is kinked at the THF site by $\sim 62^\circ$, with the two duplex arms on either side of the bend primarily B-form DNA. Interestingly, there are no protein–DNA contacts with the 5 base pairs upstream of the lesion (C1·G23–C5·G19), and the B-factors for the DNA are significantly higher at that end. The structures of TAG in the free state and when bound to product DNA are essentially identical, with r.m.s. deviations of 0.6 Å (backbone atoms only) and 1.0 Å (all atoms) (Supplementary Figure S3). Thus, no significant protein movement is necessary to engage the DNA.

TAG contains a unique HhH motif that accounts for about half of the polar interactions with the DNA backbone. Amide nitrogens from Phe156, Gly158, Thr160, and Ile161 form hydrogen bonds to the phosphate groups 3' to the THF site

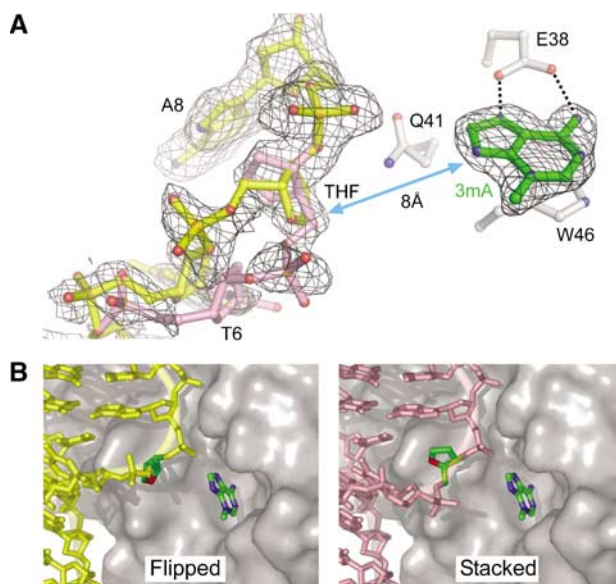


Figure 2 Two conformations of the abasic DNA backbone. (A) Refined models showing the flipped (yellow) and stacked (pink) conformations of the THF abasic site are superimposed on an unbiased composite omit electron density map (contoured at 2σ). Only density corresponding to nucleic acid is shown for clarity. The proximity of the DNA to the 3mA (green sticks) is highlighted by a double-headed arrow. (B) The flipped and stacked DNA conformations are shown separately against a van der Waals surface representation of the protein. THF and 3mA carbon atoms are highlighted in green.

(cytosine C9 and guanine G10) (Figure 1B). In contrast to DNA complexes of AlkA, hOgg1, and EndoIII, TAG does not coordinate a cation (Na^+ , Mg^{2+} , or Ca^{2+}) at the hairpin (Bruner *et al*, 2000; Hollis *et al*, 2000a; Fromme and Verdine, 2003b). Instead, a water molecule links the hairpin with the DNA backbone by coordinating in a tetrahedral arrangement only four ligands: the main-chain nitrogen of Val157, the amino N_ϵ nitrogen of Lys150, the O1P phosphate oxygen of guanine G10, and a water molecule (Supplementary Figure S4). Despite its structural divergence from other HhH glycosylases, TAG's HhH motif serves the same functional role of anchoring the protein to the DNA.

The abasic site in two conformations

One surprising aspect of the TAG/DNA complex structure is the conformational flexibility of the THF abasic site. This residue exists in two discrete orientations in the crystal (Figure 2). Both experimental MAD and unbiased composite omit electron density maps clearly show two equally occupied trajectories for the DNA backbone at residues T6 and THF7 (Figure 2A). In one orientation, the THF ribose ring is partially rotated $\sim 90^\circ$ out of the helical base stack and into the minor groove toward the protein ('flipped' conformation; Figure 2B). In the second orientation, the THF ribose remains stacked into the helix in its normal position in B-DNA. The shift in the position of the THF moiety is accompanied by a concomitant rotation of the DNA backbone that forces the THF 5'-phosphate to point either away from (flipped conformation) or toward (stacked conformation) the protein (Figure 2B). The largest deviations in the DNA backbone occur predominantly as rotations around the C3'-O3' bonds (ϵ torsion angle) of nucleotides T6 and THF7 and around the

O3'-P (ζ) bond, although the entire backbone of nucleotides C5, T6, and THF7 significantly deviates from that of B-DNA (Figure 2A). In addition to torsional rotation, the two DNA conformations differ by a 2 \AA translation around thymine T6, a movement that affects the positions of both the backbone and thymine base. The slight positional disorder in thymine T6 is reflected in the discontinuous electron density and high *B*-factors of this residue.

The multiple conformations of the phosphate backbone are likely a consequence of the sharp kink in the DNA and the lack of specific protein–DNA contacts at the abasic site and in the duplex 5' to the lesion. Surprisingly, both flipped and stacked orientations of the ribose ring make only nonspecific van der Waals contacts with TAG. Even in the flipped conformation, the abasic ribose is only partially rotated out of the DNA duplex and is located $\sim 8 \text{ \AA}$ away from the 3mA base bound in the active site pocket (Figure 2A). This unflipped ribose is in stark contrast to the structures of all other HhH glycosylases bound to abasic DNA. In these structures, the ribose is rotated a full 180° around the backbone and forms specific polar interactions inside the active site (Hollis *et al*, 2000a; Norman *et al*, 2001; Fromme and Verdine, 2003b). The structure of hOgg1 bound to THF-DNA shows the THF moiety in the same position as the ribose ring in the hOgg1/8-oxoG-DNA substrate complex (Norman *et al*, 2001), indicating that the protein–DNA interactions necessary to stabilize the flipped nucleotide in the hOgg1 active site need not involve the 8-oxoG base itself. In contrast, the TAG/THF-DNA/3mA structure suggests that the intact glycosylic bond is necessary for TAG to hold 3mA DNA substrate in a specific extrahelical orientation, and that the bound abasic DNA product relaxes its conformation after 3mA excision.

Interrogation of a DNA lesion

The HhH glycosylases use a common strategy for probing the DNA bases within the double helix. A bulky, intercalating side chain (typically, Leu, Asn, Gln, or Arg) plugs the gap in the DNA left by the flipped-out nucleotide, and a second side chain (Phe, Tyr, Leu, or Pro) wedges between the bases opposite the flipped-out nucleotide. Both plug and wedge residues are important for stabilizing the conformation of the DNA necessary to accommodate an extrahelical nucleotide. It has recently been suggested that the wedge residue is important for locating damaged DNA during the search process (Banerjee *et al*, 2006).

TAG interacts with the DNA bases in a manner different from the other HhH glycosylases. Most notable is the intercalation of Gly43 at the tip of the B/C loop into the abasic gap (Figure 3). To our knowledge, this is the first reported case of a base-flipping enzyme that intercalates backbone atoms, as opposed to a bulky side chain, into the DNA base stack. Second, the side chain of Leu44 serves as the wedge residue and intercalates between thymine T17 and adenine A18 bases on the non-lesioned strand. Interestingly, both plug and wedge residues are located on the same secondary structure element (B/C loop), and not on both the B/C-(plug) and E/F-(wedge) loops, as is observed in all other HhH glycosylase structures (Supplementary Figure S5). Thus, TAG uses a modified strategy to form the plug and wedge interactions present in all DNA glycosylases (Barrett *et al*, 1998; Parikh *et al*, 1998; Bruner *et al*, 2000; Hollis *et al*, 2000a; Fromme *et al*, 2004; Banerjee *et al*, 2006). The conservation of this

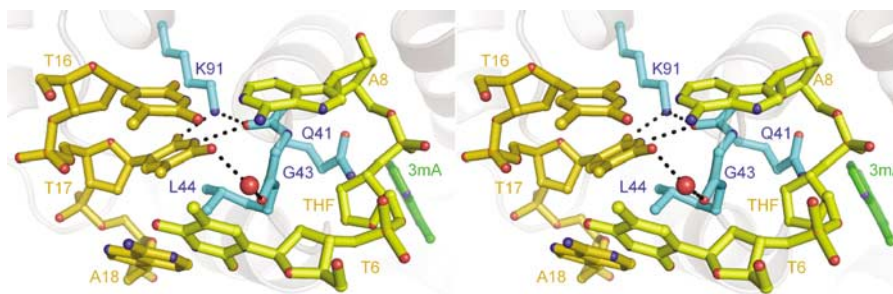


Figure 3 TAG interrogation of the DNA base stack. TAG binds the DNA damage site by intercalating side chains (blue) into both the lesioned (yellow) and non-lesioned (gold) DNA strands. Hydrogen bonds to the estranged thymine are shown as dashed lines and the 3mA base is colored green.

Table II Glycosylase activities for wild-type and active site mutants of TAG

Mutant	Conservation ^a	3mA excision rate ^b ($10^{-3} \text{ min}^{-1} \mu\text{M}^{-1}$)	Relative activity (mutant/WT)
WT	—	233.6 ± 16.4	(1.0)
L44A	Invariant	6.5 ± 4.4	2.8×10^{-2}
G43L	Invariant	2.5 ± 0.8	1.1×10^{-2}
K91A	Invariant	26.9 ± 1.5	1.1×10^{-1}
Q41A	Invariant	36.2 ± 3.1	1.5×10^{-1}
E38A	High	0.7 ± 0.1	3.0×10^{-3}
Y16F	Invariant	19.0 ± 4.6	8.1×10^{-2}
W46A	Invariant	20.7 ± 7.4	8.9×10^{-2}
W6A	Moderate	29.2 ± 16.9	1.2×10^{-1}
Y13F	Low	54.8 ± 39.3	2.3×10^{-1}
T160V	Moderate	273.8 ± 68.3	1.2
S45A	Moderate	230.0 ± 7.8	1.0
Q167A	Moderate	107.2 ± 55.7	4.6×10^{-1}

^aHigh, moderate, and low conservation refers to 1, 2–20, and >20 amino-acid substitutions, respectively, out of 250 total aligned TAG sequences (Altschul *et al*, 1997).

^bSecond-order rate constants for [³H]3mA release from *N*-[³H]methyl-*N*-nitrosourea-treated genomic DNA.

base intercalation mechanism in divergent protein architectures highlights the importance of this interaction in DNA glycosylase function.

The functional significance of the Gly43 plug and Leu44 wedge identified in the TAG/DNA crystal structure was tested by measuring the glycosylase activity of TAG site-directed mutants. The rate of 3mA excision was measured using genomic DNA treated with the alkylating agent *N*-methyl-*N*-nitrosourea (Bjelland *et al*, 1994). This agent primarily produces 7mG and 3mA lesions in DNA, and TAG selectively excises 3mA but not 7mG (Bjelland *et al*, 1993). Substituting Gly43 with a leucine residue decreased the glycosylase activity by two orders of magnitude (Table II). This decrease may partially be a result of reduced stability of the Gly43Leu protein, which is ~50% denatured under the conditions of our assay (Table II and Supplementary Figure S6). It is likely that the remaining 50-fold decrease in 3mA excision activity, which is measured by necessity under subsaturating conditions (see Materials and methods), is a result of compromised DNA binding activity of Gly43Leu. The reciprocal experiment using the closely related enzyme MagIII showed that removal of the bulky asparagine plug (Asn42Ala) enhanced DNA binding (Eichman *et al*, 2003). It is interesting to note that TAG and MagIII, both highly specific for 3mA, show greater base excision or DNA binding activity in the absence of a bulky side-chain plug.

Substitution of Leu44 with alanine decreased the glycosylase activity 36-fold in comparison to wild-type TAG

(Table II). A comparable effect of the wedge residue on DNA binding and glycosylase activity has been observed for MagIII (Eichman *et al*, 2003) and MutY (Chmiel *et al*, 2003). The predominance of phenylalanine or tyrosine wedge residues in DNA glycosylases MutY, hOgg1, and MutM suggests that aromatic stacking is important for intercalation of the bases opposite the lesion. However, the presence of leucine wedges in TAG and EndoIII (Fromme and Verdine, 2003b) and the observation that an *E. coli* MutY Tyr82Leu wedge mutant has similar activity compared to wild-type MutY (Livingston *et al*, 2005) demonstrate that van der Waals contacts are sufficient in this capacity.

As a result of the Leu44 wedge interaction, the estranged thymine T17 is highly distorted opposite the abasic site (Figure 3). This distortion is manifest as a large tilt ($\tau = -22.5^\circ$) and twist ($\omega = -8.4^\circ$) for the T16/T17 base step as compared to B-DNA ($\tau_{\text{B-DNA}} = -0.1^\circ$; $\omega_{\text{B-DNA}} = 36.0^\circ$). Such a large distortion in the estranged base has been observed in the structures of MutY and MutM bound to DNA (Fromme and Verdine, 2003a; Fromme *et al*, 2004; Banerjee *et al*, 2006). The estranged thymine is held in this distorted conformation in the TAG/DNA complex through an extensive hydrogen bond network involving lysine 91 at the N-terminal end of helix F and the B/C loop backbone (Figure 3). The N_ϵ amino group of Lys91 donates hydrogen bonds to the O2 keto oxygen of thymine T17 and to the backbone carbonyl oxygen of Ala42. The Ala42 backbone oxygen also accepts a hydrogen bond from the N3 nitrogen of

thymine T17 to form a closed T17-Lys91-Ala42 network. Substituting Lys91 with alanine reduced the rate of 3mA excision eight-fold (Table II). In addition to the T17-Lys91-Ala42 network, a water-mediated hydrogen bonding interaction links the Gly43 carbonyl oxygen from the B/C loop to the estranged thymine T17 at its O4 keto oxygen. Thus, TAG makes intimate and specific contacts with the estranged thymine base in addition to the van der Waals interactions from the intercalating residues.

The extensive interactions between TAG and the estranged base help explain the specificity of this enzyme for 3mA and 3mG residues. The same hydrogen bonds between TAG and thymine observed in the crystal structure can be formed with a cytosine but not a purine base. A model constructed with a cytosine in place of the thymine shows that a cytosine would be slightly rotated toward the minor groove of the DNA to make favorable van der Waals contacts with the surface of the protein. Alternatively, purine bases are clearly sterically excluded from this position. Specific interactions between the protein and the estranged nucleobase commonly account for HhH glycosylase substrate specificity. For example, the specificity of hOgg1 for 8oxoG·C base pairs can be rationalized by the extensive contacts between the estranged cytosine and Asn149, Arg154, and Arg204 (Bruner *et al*, 2000). AlkA, on the other hand, does not form hydrogen bonds with the estranged base, which partially accounts for its broad specificity (Hollis *et al*, 2000a).

The effect of Leu44 on the estranged base and on TAG glycosylase activity contributes to the growing body of evidence suggesting that this wedge interaction helps the enzyme find damaged base pairs among a vast excess of unmodified DNA. It has been shown that DNA glycosylases search for damage by a processive mechanism of sliding along DNA (Francis and David, 2003; Blainey *et al*, 2006). Recently, a series of crystal structures of MutM in complex with undamaged DNA demonstrate that a phenylalanine wedge (Phe114) intercalates into the base stack and severely buckles the surrounding base pairs (Banerjee *et al*, 2006). These structures suggest that such a probe in the nucleobase stack might serve as an early test of base-pair stability and thus allow the enzyme to flip into the active site only those bases whose Watson-Crick pairing has been destabilized by the presence of a modification. The distortion to the estranged thymine imposed by the TAG Leu44 wedge is consistent with the idea that TAG uses this residue to probe for DNA damage. The network of hydrogen bonds to the

estranged base would help lock the protein in place to facilitate base flipping into the active site.

3mA selection and hydrolysis in the TAG active site

The active site clefts of the HhH glycosylases have distinct chemical and physical characteristics that are suited for a particular nucleobase substrate and are located adjacent to the DNA-binding elements described above. The location of the active site with respect to the DNA lesion is important when considering how glycosylases couple damage recognition, nucleotide flipping, substrate specificity in the binding pocket, and base excision. The proximity of the TAG base binding cleft to the DNA lesion was identified by co-crystallization of all three components in the TAG/THF-DNA/3mA ternary product complex. The 3mA base was clearly observed in the experimental electron density to reside deep in the active site pocket (Figures 2A and 4). The addition of free 3mA to the crystallization experiment increased the size and quality of the crystals, suggesting that the ternary complex with bound 3mA is more stable than a binary TAG/THF-DNA complex.

The TAG active site is perfectly shaped to accommodate 3mA. An unbiased composite omit electron density map clearly distinguishes the exocyclic 3-methyl and 6-amino substituents, indicating that the base binds in one orientation (Figure 2A). The nucleobase ring nitrogen N9 that is linked to the ribose before catalysis points toward the bound DNA, suggesting that the crystal structure reflects a catalytically competent orientation of 3mA. The 3mA is constrained by hydrogen bonding and aromatic stacking interactions with active site residues (Figure 4). As observed in the NMR structure of *E. coli* TAG bound to 3mA (Cao *et al*, 2003), the side chains of Glu38 and Tyr16 line the back of the active site pocket and form hydrogen bonds to the Hoogsteen and Watson-Crick faces of 3mA, respectively. The side chains of Trp46 and Trp6 pack against one face and edge of the nucleobase ring, whereas the opposite face is contacted by water molecules held in place by hydrogen bonds from peripheral active site residues.

Despite the 8 Å distance and lack of direct contacts between the THF moiety and 3mA, the DNA damage/abasic site is linked to the base binding pocket through a series of interactions that provide insight into the base-flipping step. A water-mediated hydrogen bond network extends from Glu38 in the active site to the phosphate 3' to the THF moiety (Figure 4). Importantly, an invariant glutamine (Gln41)

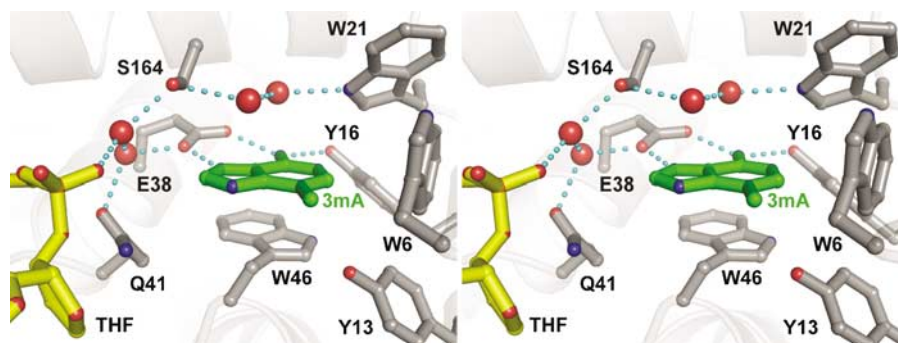


Figure 4 The nucleobase binding pocket of TAG. Protein residues are shown in gray, 3mA in green, DNA in yellow, and waters as red spheres. Hydrogen bonds are depicted as blue dashed lines.

residue is positioned directly between 3mA and THF (Figures 2A and 4), and is located on the B/C loop that plugs the abasic gap (Figure 3). Substitution of this residue with alanine reduces the rate of base excision ~6-fold with respect to wild-type TAG (Table II). On the basis of its location at the active site/THF interface and its effect on TAG activity, it is intriguing to speculate that Gln41 is involved in guiding 3mA into the base binding pocket during base flipping. Independent of whether 3mA rotates around the phosphate backbone through major or minor grooves, the modified nucleobase will likely make its first contact with Gln41. Interestingly, this is the only side chain in the base binding pocket that shifts position upon DNA binding.

The aromatic character and shape of TAG's nucleobase binding pocket is particularly well suited for interactions with alkylated purines. Electron-rich aromatic active sites that stack against electron-deficient, ring-substituted purines are common among the bacterial and human 3mA DNA glycosylases, and this feature has been shown to be important for 3mA specificity (Labahn *et al*, 1996; Hollis *et al*, 2000a; Lau *et al*, 2000; Eichman *et al*, 2003). In TAG, substitution of Trp46 with alanine had a 10-fold effect on base excision activity (Table II). A Trp6Ala mutant, on the other hand, was severely destabilized with respect to wild-type TAG

(Supplementary Figure S6), suggesting that Trp6 is important for the structural integrity of the active site.

Despite the similarities in aromaticity among 3mA base binding pockets, TAG's active site differs significantly from other glycosylases in two aspects. First, TAG lacks the conserved aspartic acid that is located 8–9 residues C-terminal to the HhH motif (Figure 5) and that is essential to the base excision activity in other HhH glycosylases (Thayer *et al*, 1995; Labahn *et al*, 1996; Guan *et al*, 1998; Mol *et al*, 2002; Norman *et al*, 2003). The lack of this catalytic residue has led to the suggestion that excision of a destabilized 3mA lesion does not require the same catalytic assistance as other more stable alkylpurines (Drohat *et al*, 2002; Stivers and Jiang, 2003), and that TAG must therefore use a unique mechanism of 3mA excision (Cao *et al*, 2003). Second, specific hydrogen bonds between 3mA and active site residues analogous to Glu38 and Tyr16 in TAG were not observed in a MagIII/3mA complex (Eichman *et al*, 2003), nor were they predicted from structures of AlkA or AAG (Hollis *et al*, 2000a; Lau *et al*, 2000). It seems likely, therefore, that the 3mA-specific contacts from Glu38 and Tyr16 contribute to TAG's narrow substrate specificity (Drohat *et al*, 2002). Indeed, the Glu38 side chain has been shown to sterically exclude N7-substituted methylpurine bases from *E. coli* TAG (Figure 4 and Cao *et al*, 2003).

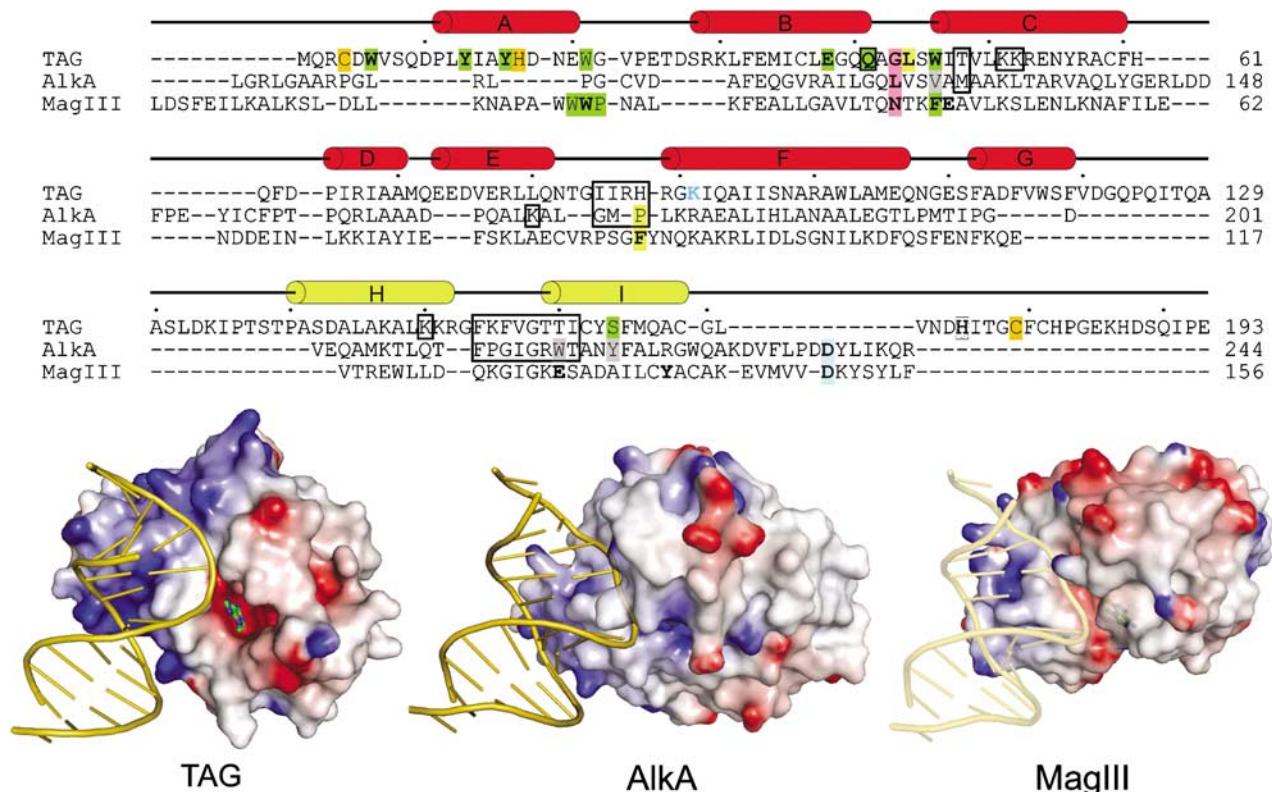


Figure 5 Comparison of 3-methyladenine DNA glycosylases. Top: structure-based sequence alignment of TAG, AlkA, and MagIII shows the relative positions of residues important for DNA binding and base excision. TAG secondary structure elements are shown schematically, with the HhH motif colored yellow. Residues contacting the DNA backbone are boxed, intercalating plug (pink) and wedge (yellow) residues are highlighted, and side chains contacting the estranged base are labeled blue. Side chains confirmed (green) or postulated (gray) to contact 3mA in the base binding pocket are highlighted. Residues verified biochemically to affect substrate binding or catalysis are shown in boldface and the catalytic aspartates in AlkA and MagIII are shaded blue. TAG residues that coordinate Zn²⁺ are shaded orange. Bottom: crystal structures of TAG/DNA/3mA, AlkA/DNA, and MagIII/3mA (with modeled DNA) are shown. Protein solvent-accessible surfaces are colored according to the electrostatic potential (blue, positive; red, negative). An alternate version of this figure showing all HhH glycosylase/DNA complexes is available as Supplementary data.

To gain insight into nucleobase specificity and excision by TAG, the enzymatic contributions of Glu38 and Tyr16 were examined by mutational studies. Steric exclusion of 7mG by Glu38 was tested by measuring the ability of a Glu38Ala mutant to excise 7mG from a defined oligonucleotide substrate. Wild-type TAG is unable to cleave 7mG from this substrate (Supplementary Figure S7). Removal of the glutamate side chain in the Glu38Ala mutant did not result in any measurable 7mG release even after 24 h (Supplementary Figure S7). Thus, TAG's inability to excise 7mG is not merely a consequence of steric exclusion by Glu38. Using the genomic DNA substrate, we found that substitution of Glu38 with alanine reduced the rate of 3mA excision 333-fold with respect to wild-type TAG, whereas a Tyr16Phe mutant had 12-fold reduced activity (Table II). Although the observed rate of 3mA excision in our assay reflects both binding and catalysis (see Materials and methods), it is unlikely that substrate binding fully accounts for the reduced activity of the Glu38Ala mutant. We do not expect the large difference in observed rate constants between Glu38 and Tyr16 mutants to be a consequence of the one additional hydrogen bond that Glu38 contributes to 3mA, as Glu38Ala and Tyr16Ala mutations each reduced the 3mA binding affinity for *E. coli* TAG by ~15-fold (Cao *et al*, 2003). Likewise, the observation that wild-type *E. coli* TAG binds weaker to positively charged 3,9-dimethyladenine base than to neutral 3mA (Cao *et al*, 2003) suggests that the reduction in base excision by the Glu38Ala mutant is not due to a loss of the electrostatic interaction between Glu38⁽⁻⁾ and DNA-3mA⁽⁺⁾. Although the precise mechanism for 3mA excision remains to be determined, these data clearly demonstrate that Glu38 has a significant effect on base excision, and are consistent with the idea that TAG is specific for destabilized 3mA lesions simply because it lacks the catalytic power to remove the more stable alkylpurine adducts from DNA (Stivers and Jiang, 2003).

Comparison of alkylpurine DNA glycosylases

The structures of TAG and AlkA bound to DNA (Figure 5) highlight important features that provide a physical basis for substrate selectivity by alkylpurine glycosylases. First, the TAG–DNA contact surface is more extensive than that of AlkA. TAG forms additional van der Waals and electrostatic interactions with the non-lesioned strand that are not present in AlkA. Additionally, DNA bound to TAG shows less backbone distortion and a closer resemblance to canonical B-DNA than in any of the other DNA complexes of HhH superfamily members (Supplementary Figure S5). This difference is not likely an artifact of the abasic THF moiety as DNA containing this analog was observed in structures of EndoIII and hOgg1 to be highly distorted as a consequence of being pulled into the active site (Norman *et al*, 2001; Fromme and Verdine, 2003b). The base binding pockets of TAG and MagIII are highly electronegative and provide a snug fit for 3mA, in contrast to AlkA's electropositive, shallow active site surface (Figure 5). This difference helps to explain the exquisite specificity of TAG and MagIII for positively charged 3mA, and suggests that the most important requirement for 3mA excision is a high-affinity binding pocket.

Based on the structures of TAG and AlkA bound to DNA, we constructed a model for TAG in complex with a 3mA-DNA substrate that illustrates a likely mechanism for 3mA excision (Figure 6). The model was constructed by superposition of

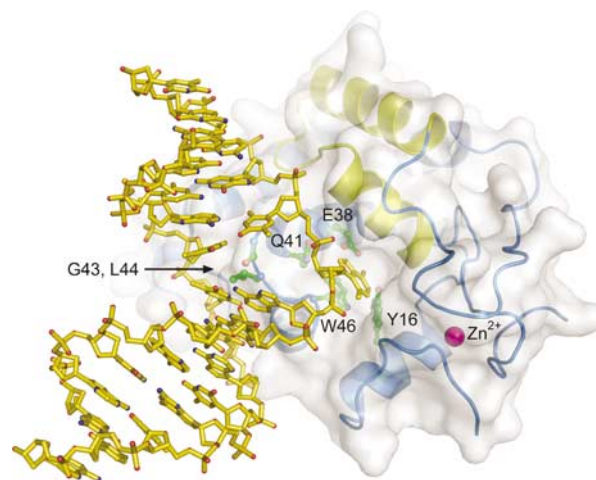


Figure 6 Model of a TAG/3mA-DNA substrate complex. TAG side chains important for DNA intercalation (Gly43, Leu44) and 3mA binding (Glu38, Trp46, Tyr16, Gln41) are shown as green ball-and-sticks. See text for details.

the DNA from the AlkA-DNA complex (Hollis *et al*, 2000a) onto the TAG/DNA/3mA structure, while retaining the position of the estranged thymine, flanking base pairs, and 3mA base from the TAG structure. This model confirms that the positions of 3mA and abasic DNA in the TAG crystal structure are aligned in biologically relevant orientations with respect to one another. The redirection of the phosphate backbone necessary to link the damage site to the 3mA base illustrates that the structure of the DNA in the TAG/THF-DNA/3mA product complex is relaxed relative to the substrate complex before hydrolysis of the glycosylic bond. This supports a previously described ground-state destabilization mechanism for catalysis of base excision (Cao *et al*, 2003; Stivers and Jiang, 2003). Collectively, TAG's enhanced interactions with both the non-lesioned strand and the 3mA base, together with the large distance between the abasic moiety and TAG's active site in the product complex argue that the 3mA glycosylic bond is strained in the substrate complex. This strain would be relieved upon cleavage of the glycosylic bond, allowing the DNA to relax to the position observed in the crystal structure.

Conclusions

The crystal structures of *S. typhi* TAG alone and bound to abasic-DNA and 3mA base provide the first structural information for how a highly specific alkylpurine DNA glycosylase engages damaged DNA. In contrast to other glycosylase-DNA structures, the abasic ribose in the TAG complex is not fully rotated into the active site, suggesting that a conformational relaxation in the DNA takes place after base hydrolysis. TAG stabilizes damaged DNA differently than other HhH glycosylases by inserting a single hairpin loop into both strands of the DNA duplex. Specific interactions with the DNA bases help to explain the enzyme's preference for 3mA bases. Results from mutagenesis studies illustrate the functional significance of key residues identified in the crystal structure, and reveal an important catalytic dependence on a highly conserved glutamate residue (Glu38) in the base binding pocket. The crystal structures and mutational data are consistent with a model in which conformational strain in the

3mA DNA substrate drives base excision by destabilizing the ground state of the reaction.

Materials and methods

TAG purification and crystallization

S. typhi was expressed as an N-terminal His₁₀-fusion protein from a pET-19b plasmid (Novagen). *E. coli* C41 cells transformed with the TAG/pET-19b plasmid were propagated in LB media supplemented with 5 μ M ZnSO₄, and protein was overexpressed for 4 h at 25°C upon addition of 0.5 mM IPTG. Cells were harvested in 50 mM Tris buffer (pH 7.5), 500 mM NaCl, and 10% glycerol and lysed with an Avestin Emulsifier C3 homogenizer operating at ~20 000 psi. TAG protein was purified using Ni-NTA (Qiagen) affinity chromatography. After cleavage of the His₁₀ tag, TAG was further purified by heparin affinity and gel filtration chromatography to >99% homogeneity as estimated by Coomassie staining. Protein was concentrated to 8 mg/ml and stored in 20 mM Tris (pH 8.5), 5% glycerol, 100 mM NaCl, 2 mM DTT, and 0.1 mM EDTA.

Selenomethionyl-substituted (SeMet) TAG was prepared similar to wild-type protein, except that the protein was overexpressed under conditions that suppress normal methionine biosynthesis (Van Duyne *et al*, 1993). Briefly, SeMet TAG was overexpressed for 16 h at 25°C in C41 cells grown in minimal media supplemented with 70 mg/ml selenomethionine (Acros Organics). After the Ni-NTA step, 5 mM methionine and 20 mM DTT were added to all buffers for the remainder of the purification.

Crystals of unliganded TAG were grown at 21°C by vapor diffusion, in which drops containing equal volumes of protein (8 mg/ml) and reservoir (30% PEG 200, 5% PEG 3000, 100 mM MES pH 6.0) were equilibrated against the reservoir. Crystals grew as single blocks (25 \times 25 \times 25 μ m³) and were used as microseeds for a second crystallization experiment using a reservoir solution containing 16% PEG 200, 5% PEG 3000, and 100 mM MES pH 6.0. Crystals grown from seeds appeared as larger single blocks (100 \times 100 \times 100 μ m³) after 1–2 days, and were flash-frozen in liquid nitrogen for X-ray data collection. To crystallize the TAG/THF-DNA/3mA complex, 0.23 mM TAG was preincubated for 15 min at 4°C with 0.27 mM DNA (d(CGGACTXACGGG)/d(CCGTTAGTCCGC), where X is a THF abasic analog) and 2 mM 3mA. Crystals were grown at 21°C by vapor diffusion using equal volumes of protein/DNA/3mA and reservoir (2 M (NH₄)₂SO₄, 2% PEG 400, 100 mM HEPES pH 7.5) solutions. The crystals grew as hexagonal rods (75 \times 75 \times 500 μ m³) in 1–2 days, and were soaked in 2 M sodium malonate (pH 7.5) before flash-freezing.

X-ray data collection, phasing, and structure refinement

X-ray diffraction data (Table I) on flash-frozen TAG and TAG/THF-DNA/3mA crystals were collected at beamline 22-ID at the Advanced Photon Source (Argonne, IL) and processed using the HKL 2000 package (Otwinowski and Minor, 1997). Data collection statistics are summarized in Table I.

Experimental X-ray phases for unliganded and DNA-bound TAG structures were obtained from MAD and SAD experiments, respectively, using crystals grown with SeMet-substituted protein. Diffraction data (Table I) were collected at energies corresponding to the selenium peak, inflection point, and high-energy remote settings (TAG) and at the peak energy only (TAG/DNA). Selenium positions (nine in TAG and five in TAG/DNA) in the asymmetric unit were located and refined using the program SOLVE (Terwilliger and Berendzen, 1999). Density modification and phase calculation were carried out using RESOLVE (Terwilliger, 2000). The protein chain was built *de novo* into 1.5-Å electron density from the TAG-only crystals. This model was docked into experimental SAD density for the TAG/DNA complex, followed by manual building of the DNA and 3mA portions of the model.

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The models were refined using experimental phases and amplitudes from native (TAG, 50–1.5 Å) and SeMet (TAG/DNA, 50–1.85 Å) diffraction data against a maximum likelihood target as implemented in REFMAC 5.1 (Murshudov *et al*, 1997). Improvements to the models were guided by manual inspection of σ_A -weighted $2mF_o - DF_c$ and $mF_o - DF_c$ electron density maps, and were judged successful by a decrease in R_{free} during refinement. Anisotropic *B*-factors were refined explicitly for each atom in the TAG structure, and translation/libration/screw-rotation (TLS) refinement in REFMAC was used to model anisotropic motion of four protein/DNA domains in the complex. Individual anisotropic *B*-factors were derived from the refined TLS parameters and held fixed during subsequent rounds of refinement, which resulted in a decrease in both *R* and R_{free} and a noticeable improvement in the electron density maps. Protein and DNA models were validated using PROCHECK (Laskowski *et al*, 1993), and DNA parameters were analyzed using CURVES 5.2 (Lavery and Sklenar, 1988).

The model coordinates, experimental phases, and structure factor amplitudes for TAG and TAG/THF-DNA/3mA structures have been deposited in the Protein Data Bank under accession numbers 2OFK and 2OFI, respectively.

Mutagenesis and enzyme activity assays

Mutations were made in the TAG expression plasmid using the Quik Change Site-Directed Mutagenesis kit (Stratagene), and they were verified by DNA sequencing. Mutant proteins were expressed and purified in the same manner as the wild-type enzyme, but without the final gel filtration step.

DNA glycosylase activity assays for 3mA excision were performed similar to the method described previously (O'Rourke *et al*, 2000; Eichman *et al*, 2003). The reaction mixture (50 μ l) contained 6 μ M enzyme and 3000 c.p.m. of *N*-[³H]methyl-*N*-nitrosourea-treated calf thymus DNA in activity buffer (50 mM HEPES pH 7.5, 100 mM KCl, 10 mM DTT, 1 mM EDTA). Reactions were incubated at 37°C and terminated at various time points by ethanol precipitation of the DNA. The release of radioactive bases into the soluble fraction was quantitated by liquid scintillation counting. Rate constants were determined from single-exponential fits to data from three different experiments for each mutant and then corrected for the concentration of each enzyme. For this assay, the enzyme concentration was subsaturating with respect to substrate at the highest concentration of enzyme tested (41 μ M). The failure to saturate might be caused by nonspecific binding of TAG to the vast excess of unmodified bases in the genomic DNA substrate. The observed second-order rate constants were shown to be linearly dependent on enzyme concentration up to at least 40 μ M, and therefore reflect both binding and catalysis under these conditions.

Supplementary data

Supplementary data are available at *The EMBO Journal* Online (<http://www.embojournal.org>).

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