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## N-Glycosyl Bond Formation Catalyzed by Human Alkyladenine DNA Glycosylase<sup>†</sup>

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ABSTRACT: The removal of damaged bases by DNA glycosylases is thought to be effectively irreversible, because of an overall equilibrium that favors hydrolysis over synthesis of the N-glycosyl bond. Surprisingly, human alkyladenine DNA glycosylase (AAG) can make damaged DNA by catalyzing formation of an N-glycosyl bond between  $1,N^6$ -ethenoadenine ( $\varepsilon$ A) and abasic DNA. We attribute the ready reversibility of this glycosylase reaction to the exceptionally tight binding and slow subsequent hydrolysis of DNA containing an  $\varepsilon$ A lesion. In principle, reversibility could provide a mechanism for direct reversal of base damage by a DNA glycosylase, allowing the glycosylase to bypass the rest of the base excision repair pathway.

The equilibria for hydrolysis of the N-glycosyl bonds of nucleotides and nucleosides favor their nucleobase and ribose products (I, 2). Indeed, de novo and salvage pathways for nucleotide biosynthesis typically use 5-phosphoribosyl- $\alpha$ -pyrophosphate (PRPP) as a substrate for N-glycosyl bond formation, as its utilization is driven by the subsequent hydrolysis of released PP<sub>i</sub>. Base excision repair DNA glycosylases hydrolyze the analogous N-glycosyl bond between a damaged base and C1 of a deoxyribosyl moiety, giving rise to a free base and an abasic site within DNA (3). Although the overall equilibrium for this reaction favors hydrolysis, human alkyladenine DNA glycosylase (AAG) unexpectedly synthesizes  $1,N^6$ -ethenodeoxyadenosine  $(\varepsilon A)$  within DNA when supplied with  $1,N^6$ -ethenoadenine  $(\varepsilon A)$  and abasic DNA. The internal equilibrium constant  $(K_{int})$  for this enzymatic transformation is near unity.

Mutagenic  $\varepsilon$ dA lesions form when adenine bases in DNA react with certain lipid peroxidation products or pollutants (4, 5). AAG is the only human glycosylase known to excise  $\varepsilon$ A from  $\varepsilon$ dA-containing DNA, and its mechanism has been proposed to include fast two-step DNA binding, rate-limiting *N*-glycosyl bond hydrolysis, and rapid dissociation of  $\varepsilon$ A (6). We explored possible inhibition of the AAG reaction by its  $\varepsilon$ A product by adding the free base to AAG- $\varepsilon$ dA-DNA, where  $\varepsilon$ dA-DNA is a 25 bp oligonucleotide that contains a centrally located  $\varepsilon$ dA (Scheme 1). The single-turnover rate constant of 0.04 min<sup>-1</sup> obtained for the AAG-catalyzed hydrolysis of  $\varepsilon$ dA-DNA to ab-DNA in the absence of added  $\varepsilon$ A matched the value of  $k_{hyd}$  measured previously [Figure 1A ( $\blacksquare$ )] (6). Addition of 0.6 mM  $\varepsilon$ A to an otherwise identical reaction resulted in faster, but only partial

Scheme 1: Duplex DNA Oligonucleotides Used in This Study

5 ' - CGATAGCATCCT**x**CCTTCTCTCCAT
3 ' -GCTATCGTAGGATGGAAGAGAGGTA

£dA-DNA: **x** = £dA

ab-DNA: **x** = deoxyribose

( $\sim$ 70%), conversion of  $\varepsilon$ dA-DNA to ab-DNA [Figure 1A ( $\square$ )], whereas a slower reaction that proceeded to completion was expected for simple product inhibition of AAG by  $\varepsilon$ A.

We tested the possibility that  $\varepsilon A$  reacts reversibly with ab-DNA to regenerate  $\varepsilon dA$ -DNA by adding free  $\varepsilon A$  to AAG·ab-DNA. Reactions containing 0.6 mM εA that were initiated with either AAG·ab-DNA or AAG·\varepsilon dA-DNA proceeded to the same end point of  $\sim$ 0.3 [Figure 1A (open symbols)], which shows the same equilibrium between AAG-bound species being approached from opposite directions. To verify that the plateau reached in the AAG·ab-DNA reaction represented the equilibrium between AAG·ab-DNA and newly synthesized AAG·\varepsilon dA-DNA, we tried to perturb it predictably. We first increased the AAG. ab-DNA concentration while maintaining the  $\varepsilon$ A concentration at 0.6 mM and followed the conversion of ab-DNA to \varepsilondA-DNA for 100 min, until the expected end point of  $\sim 0.3$  had been attained. The reaction mixture was then diluted 25-fold, resulting in a mixture containing only 24  $\mu$ M  $\varepsilon$ A, and the decay of the putative  $\varepsilon dA$ -DNA was followed to its new equilibrium position (Figure 1B). The rates of decay of the putative εdA-DNA and authentic εdA-DNA were the same within error (Figure 1B, inset), consistent with the model in which AAG converts ab-DNA and  $\varepsilon A$  into  $\varepsilon dA$ -DNA via direct reversal of the glycosylase hydrolysis step (Figure 2,  $k_{\rm syn}$ ).

We next investigated the dependence of the synthesis reaction on the concentration of  $\varepsilon A$ . As shown in Figure 3A, reactions of AAG·ab-DNA with εA proceeded with faster initial rates to higher end points as the  $\varepsilon A$  concentration increased. The reaction end points are plotted as a function of  $\varepsilon A$  concentration in Figure 3B, and saturation with  $\varepsilon A$  is observed with a  $K_{1/2}$  of  $0.9 \,\mathrm{mM}$ . At saturating concentrations of  $\varepsilon A$ , the fractional concentration of AAG $\cdot$   $\varepsilon$ dA-DNA is 0.7 (the maximum end point), and the fractional concentration of AAG·ab-DNA· $\varepsilon$ A is correspondingly 0.3, indicating that \( \varepsilon dA-DNA \) is more stable than its hydrolysis products when bound to AAG. The internal equilibrium constant for these AAG-bound species is the ratio of their fractional concentrations ( $K_{\text{int}} = 0.3/0.7 = 0.4$ ).  $K_{\text{int}}$  is also equivalent to  $k_{\rm hyd}/k_{\rm syn}$ , the ratio of the respective rate constants for formation of AAG·ab-DNA·εA and AAG·εdA-DNA from one another, so a value of 0.1  $min^{-1}$  for  $k_{syn}$  can be calculated from  $K_{\text{int}}$  and the reported value of 0.04 min<sup>-1</sup> for

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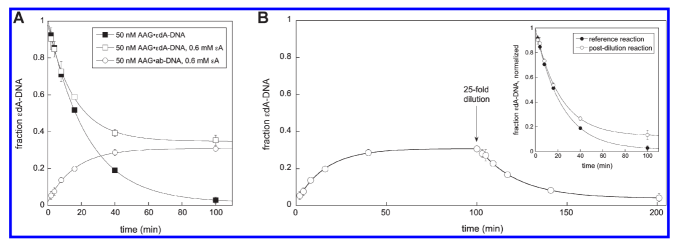


FIGURE 1: AAG synthesizes  $\varepsilon$ dA-DNA from  $\varepsilon$ A and ab-DNA. (A) The fraction of total DNA present as  $\varepsilon$ dA-DNA in quenched samples from single-turnover reactions of AAG with  $\varepsilon$ dA-DNA or ab-DNA was determined using a gel-based glycosylase assay (see the Supporting Information). Error bars indicate the standard deviation from the mean for at least two independent determinations, and lines are the best fits to single-exponential equations. (B) A single-turnover reaction containing 1.25  $\mu$ M AAG·ab-DNA and 0.6 mM  $\varepsilon$ A was followed to completion and then diluted 25-fold with buffer, resulting in a postdilution reaction of 50 nM AAG·DNA and 24  $\mu$ M  $\varepsilon$ A that was followed to its new equilibrium position. The inset shows the normalized postdilution reaction alongside a single-turnover reference reaction of 50 nM AAG· $\varepsilon$ dA-DNA. Although best fits of both inset reactions to single-exponential equations give  $k_{\rm obs}$  values of 0.04 min<sup>-1</sup>, representing the rate constant for single-turnover excision of  $\varepsilon$ A from  $\varepsilon$ dA-DNA ( $k_{\rm hyd}$ ), the postdilution reaction does not reach an end point of zero because of the residual  $\varepsilon$ A that it contains. All reactions were performed at 25 °C and pH 6.5, and  $\varepsilon$ dA-DNA reactions were corrected to account for the small percentage of ring-opened substrate that is refractory to AAG reaction (see the Supporting Information).

$$\begin{array}{c} \text{DNA} \\ \text{AAG} \bullet \text{EdA-DNA} \\ \text{AAG} \bullet \text{ab-DNA} \bullet \text{EAJ/[AAG} \bullet \text{EdA-DNA}] \\ = k_{\text{hyd}}/k_{\text{syn}} \end{array}$$

FIGURE 2: Model for reversible excision of  $\varepsilon A$  from  $\varepsilon dA$ -DNA by AAG under single-turnover conditions, where all the DNA present is bound to AAG.

 $k_{\rm hyd}$  [Figure 1A (6)]. The best global fit of the data in Figure 3A to the model in Figure 2 gives slightly different values of 0.2 for  $K_{\rm int}$  and 0.2 min<sup>-1</sup> for  $k_{\rm syn}$  (see the Supporting Information), so we report an average value of 0.3  $\pm$  0.1 for  $K_{\rm int}$ .

To the best of our knowledge, the conversion of ab-DNA and  $\varepsilon A$  to  $\varepsilon dA$ -DNA by AAG is the first example of N-glycosyl bond synthesis by a DNA glycosylase. We assessed the generality of this reverse reaction by investigating several related reactions. AAG did not insert either adenine or hypoxanthine into ab-DNA, despite extended incubation times with millimolar concentrations of each free base, nor was any reaction observed when a different DNA glycosylase, 3-methyladenine DNA glycosylase II (AlkA), was incubated with ab-DNA and εA (data not shown). Thus, the ability of AAG to stably synthesize  $\varepsilon dA$ -DNA appears to be unusual. A related enzyme activity that inserts purines into depurinated DNA has been reported previously (7, 8), and kinetic isotope effects for the DNA glycosylase MutY are consistent with a dissociative mechanism in which reversible C-N bond cleavage precedes irreversible nucleophilic attack by water (9).

Interestingly, the structure of AAG crystallized in the presence of an \(\epsilon\)dA-containing DNA substrate revealed an intact N-glycosyl bond (10). Although the authors proposed that the high concentration of MgCl<sub>2</sub> present during crystallization may have inhibited the glycosylase activity of AAG, our results raise the possibility that the substrate may have instead persisted because of the internal equilibrium that favors synthesis of \( \varepsilon d A \)-containing DNA over its hydrolysis by AAG. The crystal structure shows εdA flipped into the AAG active site, securely held by aromatic stacking interactions and a hydrogen bond between the backbone amide of His-136 and N<sup>6</sup> of the nucleobase. This snug fit helps to account for the observed tight binding of \(\epsilon\)dA-containing DNA by AAG (6, 11, 12) and the highly favorable equilibrium for flipping  $\varepsilon dA$  into the active site [ $K_{\rm flip} \sim 1300~(6)$ ]. The unusual stability of EdA-DNA when bound to AAG, which enabled us to detect its synthesis and to measure the internal equilibrium constant for its hydrolysis, presumably arises from its tight binding to the active site coupled with slow subsequent hydrolysis of its N-glycosyl bond. In contrast, human uracil DNA glycosylase (UDG) has been crystallized bound to its uracil and abasic DNA products (13, 14), consistent with an internal equilibrium for UDG that favors hydrolysis of uracil-containing DNA (15).

As described above, results from reactions at saturating  $\varepsilon A$  concentrations allowed determination of a value of 0.3 for  $K_{\rm int}$ , the equilibrium constant for hydrolysis of the N-glycosyl bond of AAG-bound  $\varepsilon dA$ -DNA.  $K_{\rm int}$  values near unity are considered to be optimal for enzymes that operate reversibly (16). While it stands to reason that certain metabolic enzymes that act at the N-glycosyl bond may benefit from reversibility (e.g., nucleosidases and nucleoside phosphorylases), synthesis of  $\varepsilon dA$ -containing DNA by AAG is expected to be negligible under typical conditions, because of low cellular concentrations of abasic DNA and  $\varepsilon A$ . In principle, however, ready reversibility provides a potential mechanism for direct repair by DNA glycosylases, in which excision and dissociation of an aberrant base from DNA are

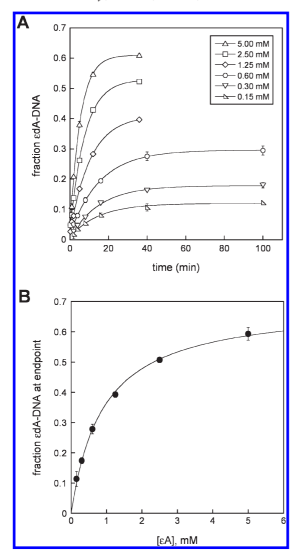


FIGURE 3: Dependence of AAG-catalyzed synthesis of \( \varepsilon AA-DNA \) on the concentration of  $\varepsilon A$ . (A) Reactions of 50 nM AAG·ab-DNA at varying EA concentrations, as indicated in the legend. Error bars indicate the standard deviation from the mean for at least two independent determinations, and lines are the best fits to single-exponential equations. All reactions were performed at 25 °C and pH 6.5. (B) Reaction end points from the curves in panel A plotted as a function of  $\varepsilon A$  concentration. A maximum end point value of 0.7 and a  $K_{1/2}$  of 0.9 mM were obtained from the best fit to a saturation equation: end point =  $[(\text{end point}_{\text{max}})[\varepsilon A]]/(K_{1/2} + [\varepsilon A])$ .

followed by binding and stable incorporation of the appropriate base. Such a repair mechanism, if present in nature, would be reminiscent of the overall base exchange processes catalyzed by tRNA guanine transglycosylases, nucleotide deoxyribosyltransferases, and nucleoside ribosyltransferases (17, 18).

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## SUPPORTING INFORMATION AVAILABLE

Detailed experimental procedures and supplementary figures. This material is available free of charge via the Internet at http:// pubs.acs.org.

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