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Interaction of the Family-B DNA Polymerase from the Archaeon Pyrococcus furiosus with Deaminated Bases

ARTICLE in JOURNAL OF MOLECULAR BIOLOGY · OCTOBER 2007

Impact Factor: 4.33 · DOI: 10.1016/j.jmb.2007.07.015 · Source: PubMed

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Probing the Interaction of Archaeal DNA Polymerases with Deaminated Bases Using X-ray Crystallography and Non-Hydrogen Bonding Isosteric Base Analogues^{†,||}

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Received March 19, 2010; Revised Manuscript Received May 5, 2010

ABSTRACT: Archaeal family-B DNA polymerases stall replication on encountering the pro-mutagenic bases uracil and hypoxanthine. This publication describes an X-ray crystal structure of Thermococcus gorgonarius polymerase in complex with a DNA containing hypoxanthine in the single-stranded region of the template, two bases ahead of the primer-template junction. Full details of the specific recognition of hypoxanthine are revealed, allowing a comparison with published data that describe uracil binding. The two bases are recognized by the same pocket, in the N-terminal domain, and make very similar protein—DNA interactions. Specificity for hypoxanthine (and uracil) arises from a combination of polymerase—base hydrogen bonds and shape fit between the deaminated bases and the pocket. The structure with hypoxanthine at position 2 explains the stimulation of the polymerase 3'-5' proofreading exonuclease, observed with deaminated bases at this location. A β -hairpin element, involved in partitioning the primer strand between the polymerase and exonuclease active sites, inserts between the two template bases at the extreme end of the double-stranded DNA. This denatures the two complementary primer bases and directs the resulting 3' single-stranded extension toward the exonuclease active site. Finally, the relative importance of hydrogen bonding and shape fit in determining selectivity for deaminated bases has been examined using nonpolar isosteres. Affinity for both 2,4-difluorobenzene and fluorobenzimidazole, non-hydrogen bonding shape mimics of uracil and hypoxanthine, respectively, is strongly diminished, suggesting polar protein—base contacts are important. However, residual interaction with 2,4-difluorobenzene is seen, confirming a role for shape recognition.

The family-B DNA polymerases from the archaea are unusual in specifically recognizing uracil and hypoxanthine, the deamination products of cytidine and adenine, respectively. During replication, these polymerases scan the template strand ahead of the replication fork and tightly bind such deaminated bases, should they be encountered four positions ahead of the primertemplate junction (1-5). Following the capture of uracil, replication is aborted, thereby preventing the incorporation of adenine and the conversion of a C·G to a T·A base pair. Similarly, cessation of polymerization in response to hypoxanthine results in the avoidance of an $A \cdot T$ to $G \cdot C$ transition mutation. Thus, read-ahead recognition appears to be the first step in a DNA repair pathway that prevents mutations arising as a consequence of the deamination of cytidine or adenine. Additional surveillance is provided by strong stimulation of the 3'-5' proofreading exonuclease activity seen when the polymerase approaches within four bases of uracil or hypoxanthine, e.g., when these bases are at position 2. This activity trims back the extending primer, resetting the stalling position to four bases (6). Interaction with deaminated bases seems to be confined to the family-B DNA polymerases from the archaea. Family-B enzymes from

Read-ahead recognition requires exquisite selectivity for uracil and hypoxanthine. Any interaction with canonical DNA bases would result in aberrant termination of replication and premature cell death. Recently, a crystal structure of the family-B polymerase from *Thermococcus gorgonarius* (Tgo-Pol), in complex with a primer-template containing uracil at the optimal 4 position in the template, has elucidated the features responsible for specificity (4). Uracil was flipped into a specific binding pocket, located within the polymerase N-terminal domain, with the formation of two protein-base hydrogen bonds. Both interactions involve the peptide backbone with the amide nitrogens of Ile-114 and Tyr-37 interacting with the O2 and O4 atoms of uracil, respectively. The flipped uracil fits extremely snugly into the recognition pocket, with the side chains of Val-93 and Pro-36 packing above the base and Ile-114 and Arg-119 below the base. Val-93 stacks on top of the uracil ring, with the isopropyl segment of the valine side chain lying flat and in the same plane as the uracil ring, mimicking the hydrophobic stacking seen in duplex DNA. The polymerase makes strong interactions with the phosphates flanking uracil, Tyr-7

bacteriophages (I) and eukaryotes (7), which are strongly similar in sequence and structure to the archaeal enzymes, are unable to recognize these bases.

[†]B.A.C. was supported by grants from the European Commission (MRTN-CT-2005-019566) and the UK BBSRC (BB/F00687X/1). T.K. is a UK BBSRC-supported Ph.D. student. E.T.K. acknowledges support from the National Institutes of Health (GM072705).

[&]quot;Protein Data Bank accession code: Coordinates have been deposited under the accession code 2xhb.

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¹Abbreviations: Fpg, formamidopyrimidine (fapy)-DNA glycosylase (also known as 8-oxoguanine DNA glycosylase); diFBz, difluorobenzene; FBzIm, fluorobenzimidazole; U, uracil; H, hypoxanthine; Hex, hexachlorofluorescein; Cy-5, cyanine-5; Pfu-Pol, family-B DNA polymerase from *Pyrococcus furiosus*; Tgo-Pol, family-B DNA polymerase from *Thermococcus gorgonarius*; PCNA, proliferating cell nuclear antigen.

forming a hydrogen bond with the 5'-phosphate and Arg-97 a salt bridge with the 3'-phosphate. Until now, less information has been available for the mechanism responsible for specific recognition of hypoxanthine, which appears to interact with the polymerase $\sim 1.5-4.5$ -fold less strongly than uracil does (5, 8). Nor has it been obvious how the polymerase is able to interact with both uracil (a pyrimidine) and hypoxanthine (a purine), while simultaneously rejecting the four standard DNA bases. It has been proposed that hypoxanthine may bind in the less favored syn conformation (5), in contrast to uracil which clearly binds as the preferred anti form (4). This work presents a structure of Tgo-Pol with an oligodeoxynucleotide containing hypoxanthine two bases ahead of the primer-template junction, revealing fully how the deaminated purine is specifically recognized. Further, the ability of the polymerase to tightly bind both a purine and a pyrimidine is elucidated, and clues about why locating these bases in position 2 stimulates exonuclease activity are provided.

The X-ray structure data published previously for uracil (4), along with that revealed for hypoxanthine in this paper, show binding of the two bases in an exquisitely tailored pocket, with formation of several protein-base hydrogen bonds. However, the role of these hydrogen bonds in the selective binding of deaminated bases has yet to be tested, for example, using base analogues lacking hydrogen bonding capability. Nonpolar nucleoside isosteres were designed to be the same size and shape as natural bases (with benzenes and indoles/benzimidazoles replacing pyrimidines and purines, respectively) but to completely lack polar carbonyl, amino, and imino functions. These bases have found extensive use as probes of the importance of polar functional groups and hydrogen bonding in protein-nucleic acid interactions (9-12). Perhaps the most surprising results were seen with A- and B-family DNA polymerases, which accepted isosteres (e.g., difluorotoluene, a deoxythymidine mimic), during replication with efficiency and specificity approaching those of natural bases (12-15). It was concluded that steric factors played a critical role during polymerase-catalyzed replication, and the formation of base pairs with the appropriate size, shape, and geometry was more important than the generation of Watson-Crick hydrogen bonds. However, polar interactions are important in other biological systems. For example, family-Y polymerases, involved in translesion synthesis and DNA repair, replicate poorly with difluorotoluene (15-18). The 3'-5' proofreading exonuclease activity of family-A DNA polymerases also depends on correct Watson-Crick base pairing (19). With DNA repair proteins such as Fpg and MutY, isosteric analogues have shown that both Watson-Crick interactions between the two DNA strands and hydrogen bonds between the protein and the damaged base play important roles in the repair process (20). In this work, the nonpolar isosteres difluorobenzene (diFBz) and fluorobenzimidazole (FBzIm), uracil and hypoxanthine analogues, respectively (Figure 1), have been used to shed light on the importance of hydrogen bonds in the selection of deaminated bases by archaeal DNA polymerase.

METHODS

Polymerase Purification. The purification of the family-B DNA polymerases from Pyrococcus furiosus (Pfu-Pol) has been described previously (21). The polymerase from T. gorgonarius (Tgo-Pol) was produced in an identical manner. All polymerases used in this publication lacked the 3'-5' proofreading exonuclease

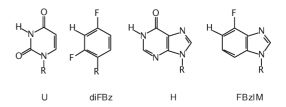


Figure 1: Deaminated bases and isosteric base analogues used in this study. $R=2^{\prime}$ -deoxyribose.

activity, by the incorporation of the D215A mutation (21) which abrogates exonucleolytic degradation of DNA but has no effect on the binding of the polymerase to deaminated bases (1).

Crystallization of Tgo Polymerase. Complexes between Tgo-Pol and a number of oligodeoxynucleotides containing hypoxanthine were prepared for crystallization by mixing at a molar ratio of 1:1.2. Crystals with 5'-AAHGGAGACAC-GGCTTTTGCCGTGTCTC-3' were obtained in 0.1 M Bicine (pH 9.0) and 20% PEG 6000 from a robot screen in which 100 nL of the complex (protein at 10 mg/mL) was mixed with 100 nL of well solution. Crystals were flash-frozen in liquid nitrogen using Paratone-N as a cryoprotectant. Diffraction data were collected at Diamond light source on beamline I04. Data were processed and scaled using MOSFLM (22) and SCALA (23) from the CCP4 suite (24), respectively.

Structure Determination and Refinement. The cocrystal structure is in space group P2₁2₁2₁ with one Tgo-pol-DNA complex per asymmetric unit and the following unit cell dimensions: a = 79.5 Å, b = 98.5 Å, and c = 116.9 Å. The structure was determined by molecular replacement using MOLREP (25) with the protein component of Protein Data Bank (PDB) entry 2VWJ (4) as the search model. After the domains had undergone rigid body fitting, clear density for a base was visible in the pocket. In addition, positive density, consistent with base stacking of DNA, was visible adjacent to the thumb, and the sequence of the oligodeoxynucleotide used during crystallization fitted the density. In addition, density indicative of DNA was observed on the exterior surface of the polymerase, bridging the space between symmetry-related molecules of the crystal. The positioning of the oligodeoxynucleotide used within this density was not possible, and hence, only a sugar-phosphate backbone has been modeled; all nucleotides are labeled as deoxyadenosines within the PDB entry. Manual rebuilding was conducted in COOT (26) interspersed with cycles of refinement using REFMAC (27) until convergence. Superimpositions were performed with LSQMAN (28).

Preparation of Oligodeoxynucleotides Containing Difluorobenzene and Fluorobenzimidazole. The diFBz and FBzIM nucleosides and their phosphoramidite derivatives were prepared as previously reported (29, 30). Both modified bases were incorporated into three oligodeoxynucleotides using standard phosphoramidite DNA synthesis chemistry and characterized by MALDI-TOF mass spectroscopy. Two sequences were prepared: 5'-GGAGACAAGCXTGCTTGCCTGCAGGTC-GACTCTAGAGGATCCCC-3' (X = diFBz, calcd 13523, found 13527; X = FBzIm, calcd 13544, found 13538) and 5'-GGAGACAAGCTTGCXTGCCTGCAGGTCGACTCTAGAGGATC-CCC-3' (X = diFBz, calcd 13522, found 13523; X = FBzIm, calcd 13544, found 13537).

Primer-Template Extensions. The ability of Pfu-Pol to copy beyond modified bases was measured using primer-template extension assays (1–5). A Cy-5-labeled primer 24 bases in length (5'-Cy-5-GGGGATCCTCTAGAGTCGACCTGC-3')

was used with two related 44-mer templates (5'-GGAGACA-AGCX₁TGCX₂TGCCTGCAGGTCGACTCTAGAGGATCC-CC-3'). When $X_1 = T$ and $X_2 = uracil$, hypoxanthine, a stable abasic site (tetrahydrofuran; sold as dSpacer phosphoramidite, Glenn Research, Sterling, VA), diFBz, or FBzIm, the modified bases are positioned six bases ahead of the primer-template junction. When X_1 = uracil, hypoxanthine, a stable abasic site, diFBz, or FBzIm and $X_2 = T$, the modified bases are positioned 10 bases ahead of the primer-template junction. Reactions were conducted at 37 °C in 150 μL volumes containing 20 mM Tris (pH 8.5), 10 mM (NH₄)₂SO₄, 2 mM MgSO₄, 15 μ g of acetylated bovine serum albumin, the four dNTPs (0.1 mM each), 20 nM primer-template, and 100 nM Pfu-Pol. After 2, 7, 15, and 30 min, a 25 μ L aliquot was withdrawn and the reaction quenched by addition of an equal volume of 80% formamide, 10% glycerol, 0.1 M EDTA, and orange G. An excess of "competitor" DNA, corresponding to the sequence of a fully extended primer (but lacking Cy-5), was added, and the samples were heated at 90 °C for 10 min and then rapidly cooled on ice (the competitor sequesters the template and so prevents any reannealing of the extended Cy-5 primer, giving cleaner electrophoresis data) (6). Extension products were detected by denaturing polyacrylamide (15%) gel electrophoresis followed by fluorescence detection using a Typhoon scanner (GE Healthcare).

Binding of Pfu-Pol to DNA Containing Modified Bases. The interaction of Pfu-Pol with DNA containing modified bases was assessed using fluorescence anisotropy with hexachlorofluorescein (Hex)-labeled DNA (3, 4). A 5'-Hex-labeled primer (5'-Hex-GGGGATCCTCTAGAGTCGACCTGCAG-3'), 26 bases in length, was annealed to a 44-mer template (5'-GGAGACA-AGCTTGCXTGCCTGCAGGTCGACTCTAGAGGATCCCC-3', X = T, uracil, hypoxanthine, diFBz, or FBzIm). This combination locates the modified base four positions ahead of the primer-template junction, resulting in the tightest possible binding (3). Binding was assessed in volumes of 1 mL containing 25 mM Hepes (pH 7.5), 100 mM NaCl, 1 mM EDTA, and 5 nM primer-template. Pfu-Pol was added in aliquots and the anisotropy measured after each addition. Data were fitted to binding isotherms, assuming a 1:1 stoichiomety, using Grafit (Erithacus Software, London, U.K.).

Incorporation of a Single dNTP into Primer-Template-Containing Modified Bases. Single-turnover incorporations (8) used essentially the same volumes, buffer conditions, and temperatures described for primer-template extensions, save the primer-template concentration was 10 nM. The primertemplate consisted of 5'-Cy-5-GGGGATCCTCTAGAGTC-GACCTGCAG-3' (primer) and 5'-GGAGACAAGCTTGCX-TGCCTGCAGGTCGACTCTAGAGGATCCCC-3' (X = T, diFBz, or FBzIm). This combination places the modified bases four positions ahead of the primer-template junction. In addition Pfu-PCNA (300 nM) was added to ensure complete binding of the polymerase (100 nM) to the DNA and, hence, single-turnover conditions (8). dGTP, complementary to the first single-stranded base in the template, was the only dNTP added. The reaction was quenched and analyzed by gel electrophoresis as described above. Data obtained at each dGTP concentration were fitted to single exponentials to obtain k_{obs} values, which were replotted versus dGTP concentration using the relationship $k_{obs} = k_{pol}[dGTP]/(K_D +$ [dGTP]). Grafit (Erithacus Software) was used for both fitting procedures.

RESULTS

Structure of Tgo-Pol with a Hypoxanthine-Containing *Oligodeoxynucleotide*. The crystal structure of Tgo-Pol with a primer-template containing uracil at position 4 has been described previously (4). Comparison with bacteriophage RB69 Pol (a family-B member) containing DNA bound in either the polymerization (31) or editing (32) mode indicated that the overall localization of the U + 4 primer-template junction was reminiscent of that observed for the DNA in the RB69 editing structure. However, no bases at the 3'-end of the U + 4 primer were unwound to give single strands, and primer-templates with uracil at this position are only slowly subject to proofreading exonuclease activity (6). In light of results showing that deaminated bases at positions 0, 1, 2, and 3 lead to more rapid exonucleolytic degradation of the primer than in the case when a deaminated base is at position 4 (6), we were curious to investigate such a complex. Additionally, the absence of data for the binding of hypoxanthine precluded a full understanding of the selectivity of the pocket, which is able to recognize both a deaminated pyrimidine and a purine (5). We thus determined the structure of a DNA-polymerase complex, using AAHGGA-GACACGGCTTTTGCCGTGTCTC, an oligodeoxynucleotide that folds to a stem-loop structure to produce a primer-template mimic, which places hypoxanthine at position 2 (Figure 2a). An analogous oligodeoxynucleotide was used in our previous study of the binding of U + 4 (4). Data collection and refinement statistics for the final model are summarized in Table 1.

The protein-DNA interactions seen with H + 2 are summarized in Figure 2b, and the DNA largely makes the same contacts with the thumb, palm, and amino-terminal domains that are observed in the uracil structure (4). However, with H + 2, the thumb domain of the polymerase is moved even further, in comparison to U + 4, from the position it occupies in the apoenzyme form (33). The two bases immediately 3' of the hypoxanthine (G4 and G5) are found stacking within the single-stranded template channel T of the polymerase between Pro-115 and Met-244 (Figure 2c,d). The majority of the remaining DNA is in the approximately double-stranded B form within the main polymerase cavity, although six bases (T15–C20) at the hairpin bend and the two extreme 5'-bases (A1 and A2) are not visible in the electron density maps. The sequence should allow G5 and A6 to base pair with T27 and C28, and indeed, with the U + 4 structure, all bases in the double-stranded region adjacent to the primer-template junction are paired. However, in the H+2structure, T27 and C28 have peeled away from their complementary bases in the template strand to give a two-base singlestranded extension, with T27 positioned approximately 6 Å from A6. This denaturation appears to be driven by the β -hairpin motif, two antiparallel β -strands joined by a loop and comprising amino acids 240-251. Arg-247, found in the loop region of the β -hairpin, inserts between G5 and A6, stacking against A6 and acting as a wedge to force these two bases apart (Figure 2c,d). The separation between the C1' atoms of G5 and A6 is 8.4 Å; for the N9 atoms, a distance of 9.8 Å is observed. Additionally, D246, also present in the loop region, appears to form a hydrogen bond with A6 and may contribute to strand separation. As a consequence, T27 and C28 are displaced into the editing channel (Figure 2c,d), and additional hydrogen bonds between the phosphates of T27 and C28 and the backbone amide nitrogens of Y273 and L275 (amino acids in the exonuclease domain) are observed (not shown). Y261 helps maintain the primer strand in

FIGURE 2: Structure of the Tgo-Pol-DNA complex. (a) Hypoxanthine-containing DNA used to obtain structural data. The oligodeoxynucleotide folds to produce a primer-template mimic that places hypoxanthine two bases (+2) ahead of the primer-template junction. (b) Summary of the interactions seen between Tgo-Pol and the H + 2 primer-template mimic. Although all the bases are shown, A1, A2, and C14—G19 are not visible in the structure. The extreme 3'-bases (T27 and C28) become single-stranded when bound to the polymerase. The amino acids contacting the DNA are color-coded according to polymerase subdomain: thumb, green; palm, pink; amino-terminal, yellow; exonuclease, blue. The interactions made by the β -hairpin with G5 and A6 are not shown. (c and d) Two views illustrating key features of the interactions of the H + 2 primer-template mimic with Tgo-Pol. Hypoxanthine (Hyp 3) is bound in the *anti* conformation. G4 and G5 are sandwiched between Pro-115 and M244; R247 stacks against A6, and D246 appears to form a hydrogen bond with this base. As a consequence, DNA denaturation occurs, with primer bases T27 and C28 becoming single-stranded and well-separated from their complementary template bases, G5 and A6. T27 and C28 are located in the editing channel, and Y261, which is positioned between T27 and A6, hinders reannealing. Polymerase amino acids are color-coded by subdomain as in panel b. (e) Contacts between Tgo-Pol and hypoxanthine. Three protein—base hydrogen bonds are seen: Tyr-37 (backbone)—O6, Ile-116 (backbone)—N3, and Glu-111/R119—N1 (water-mediated). The side chains of E111 and Arg-119 form a salt bridge. The interactions of Tyr-7 and Arg-97 with the flanking phosphates (see also Figure 3a) and the stacking of Val-93 with hypoxanthine are also shown.

the editing channel by holding T27 away from its template partner, A6 (Figure 2c,d). Although T27 and C28 are directed toward the 3'-5' exonuclease site, the extreme 3' base, C28, does not actually enter the active site and so is not correctly positioned for excision. Previously (6), it was predicted that the location of a

deaminated base at position 2 resulted in the formation of a single-stranded region in the primer, with the 3'-base inserting into the 3'-5' exonuclease active site for rapid removal. While the H+2 structure confirms the formation of a single-stranded region, entry into the exonuclease site is not seen. The latter may

Table 1: X-ray Diffraction Data Collection and Refinement Statistics

Data Collection

$P2_{1}2_{1}2_{1}$
a = 79.45, b = 98.37, c = 116.63
47.01-2.72 (2.87-2.72)
23740 (3473)
95.1 (97.0)
3.6 (3.7)
10.9 (25.0)
8.8 (4.2)

Refinement

47.01-2.72 (2.79-2.72)
23.3 (31.5)
29.4 (39.4)
6484
5949
512
22
34
0
97.82
0.006
0.944

^aValues in parentheses refer to values in the highest resolution shell. ^bRoot-mean-square deviation.

arise through the use of D215A (an exo mutant) and the absence of divalent metal ions essential for exonuclease activity (34). It is presumed that in the complete system a further conformational change places the 3'-base in a suitable position for catalysis.

Hypoxanthine is observed in the same binding pocket used to accommodate uracil and is clearly bound in the anti conformation (Figure 2c), contrary to an earlier prediction that the purine may be recognized in the *syn* conformation (5). Thus, both uracil and hypoxanthine interact with the polymerase as anti conformers. Comparison of the structure of the uracil complex (4) with the apoenzyme (33) revealed a preformed binding pocket, with no conformational change taking place following interaction with the deaminated base. Similarly, the binding of hypoxanthine does not alter the structure of the pocket, and the amino acids involved in selective binding of uracil are again observed to be critical for recognition of hypoxanthine (Figure 2e). The same two backbone amide nitrogens position the hypoxanthine within the pocket; Tyr-37 and Ile-114 hydrogen bond with the exocyclic O6 group and the ring N3, respectively (Figure 2e) (with uracil, these amino acids recognize the spatially near-equivalent O4 and O2 functions). It is clear that hypoxanthine alone is not sufficient to account for all the observed density in the pocket, as an additional feature is visible $\sim 2.8 \text{ Å}$ from N1 of the purine. This density, which is situated within hydrogen bonding distance of NH1 of Arg-119, and the main chain O of Glu-111, has been modeled as a water molecule (Figure 2e). A water-mediated hydrogen bond network linking Arg-119 and Glu-111 with N1 of hypoxanthine would further increase selectivity for this deaminated base. In addition, the side chains of Arg-119 and Glu-111 form a salt bridge (Figure 2e), as they do in the U + 4 (4) and apoenzyme (33) structures. While a corresponding water molecule is not visible in the model of the U + 4 structure, the modest resolution does not allow the presence of such a water to be unequivocally excluded. Indeed, it is possible to model a water molecule in the pocket in a manner that makes chemical sense, albeit at a slightly shifted position because of the differences

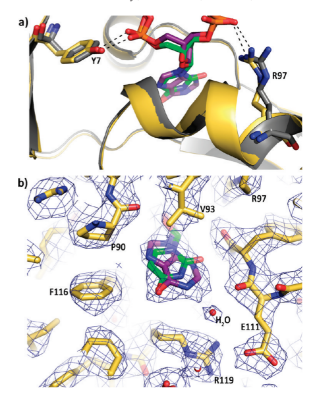


FIGURE 3: Comparison of hypoxanthine and uracil recognition. (a) Overlay of the hypoxanthine (purple) (this publication) and uracil (green) (4) structures with Tgo-Pol. The base, sugar, and phosphates of the two deaminated bases are nearly superimposable, as are the amino-terminal domains (yellow and gray for the hypoxanthine and uracil structures, respectively). The degree of side chain overlap is also very high, as shown for Tyr-7 and Arg-97, which bind the phosphates flanking the deaminated bases. (b) Superimposition of uracil (green) onto the structure of hypoxanthine (purple) bound to Tgo-Pol. The C5-C6 edge of uracil is near Pro-90 and Phe-116 (and also Pro-36, which is not visible in this view), and C2 of hypoxanthine is close to the main chain carbonyl oxygen of Glu-111. Such tight packing of the deaminated bases results in steric exclusion of the larger thymine (5-CH₃) and guanine (2-NH₂) bases. The water molecule involved in a water-mediated hydrogen bond with hypoxanthine is also shown.

between the purine and pyrimidine rings. The hydrophobic contributions seen with uracil are preserved with hypoxanthine. Again of particular note is Val-93, whose isopropyl group effectively stacks with the plane of the base. The N7 side of the imidazole ring is flanked by Pro-90, Pro-36, and Phe-116, and the side chains of Arg-119 and Ile-114 also contribute to the shape of the pocket (Figure 2e). An overlay of the U + 4 and H + 2structures (Figure 3a,b) demonstrates that the base, sugar, and flanking phosphates of the two deaminated bases occupy virtually identical positions, as do the side chains of Tyr-7 and Arg-97, which tether the flanking phosphates at the mouth of the pocket. The three amino acids, Pro-90, Pro-36, and Phe-116, on one side of the pocket (the left side of Figure 3b) are close to the C5-C6 double bond of uracil, such that binding of thymine would give rise to a severe steric clash with the 5-CH₃ group (4). The main chain oxygen of Glu-111, on the other side of the pocket (right-hand side in Figure 3b), plays a similar role with purines. This backbone atom is located near C2 of hypoxanthine, such that the exocyclic amino group present in guanine would lead to a steric clash ensuring its exclusion from the pocket (Figure 3b).

Replication of Modified Base-Containing Primer-Templates. The inability of Pfu-Pol to copy beyond deaminated

bases is most simply observed using primer extension assays, where the base is located at a defined position in the template strand, and its ability to impede polymerization noted (1-8). This method has been applied as an initial test of the interaction of both diFBz and FBzIm with Pfu-Pol. Although different Pols were used for crystallization and biochemical experiments, Tgo-Pol and Pfu-Pol are ~80% identical in amino acid sequence, and their crystal structures are essentially identical (33, 35). The two polymerases show indistinguishable behavior with deaminated bases (4). The results observed for difluorobenzene (diFBz), after incubation for 15 min, are shown in Figure 4a. The control template, comprising only the four natural DNA bases, was completely extended (lane 2). Templates containing uracil (U), either 6 or 10 steps ahead of the primer-template junction, gave truncated products, consistent with stalling of replication three to four bases prior to its encounter (2, 3, 6) (lanes 3 and 6, products resulting from stalling marked with an arrow). With a stable abasic site (Ab), shortened products were also observed (lanes 4 and 7) as the polymerase cannot effectively add a base opposite the noncoding site. Note that the polymerase pauses at the abasic site itself, rather than stalling three to four bases upstream, and therefore, any truncated products seen with abasic sites are longer than those observed with uracil. With diFBz, some evidence of stalling is seen for position 10, with an extended product, corresponding to that found with uracil, being produced (lane 8, key product marked with an arrow). However, a band equivalent to that seen with the abasic site (arising as the polymerase is poor at inserting a base opposite the template strand isostere) along with full-length product is also seen. Further, there is no evidence of stalling with diFBz at position 6, the observed band running with that produced by an abasic site (lane 5). A shorter time (2 min) was used to better visualize any transient stalling caused by diFBz (Figure 4b). With diFBz at position 10, most of the product now corresponds to the stalled band seen for U + 10 (lane 8, key band indicated with an arrow). Even with diFBz + 6, traces of pocket-mediated stalling are now visible as a minor band (lane 5, critical band arrowed) running just below the main truncated product, which comigrates at the Ab + 6 location. Overall, it is clear that any pausing of replication due to the specific capture of diFBz in the deaminated base binding pocket is significantly weaker than seen with uracil. Figure 4c shows the gel patterns observed with fluorobenzimidazole (FBzIm) after 15 min. This gel additionally contains lanes corresponding to hypoxanthine (H) at positions 6 and 10, which give stalled products similar to those seen with uracil at the same location (lanes 3, 4, 7, and 8, bands arising from stalling marked with arrows). With FBzIm, the gel shows no evidence of pocketmediated capture (lanes 6 and 10); rather, all extended products correspond to those seen with an abasic site, consistent with the enzyme's active site being inefficient at using FBzIm as a coding base. Shorter incubation times, comparable to the period of 2 min used with uracil, gave results identical to those seen after 15 min (data not shown). Thus, FBzIm does not interact with the deaminated base binding pocket, resulting in stalling of replication.

Binding of Pfu-Pol to Primer-Templates Containing Isosteric Bases. Results obtained using primer-template extension assays only give a qualitative indication of recognition by the deaminated base binding pocket. To obtain a more quantitative measure of any affinity for diFBz and FBzIm, binding titrations using fluorescence anisotropy have been performed (2–5). These studies have been conducted with primer-templates that position

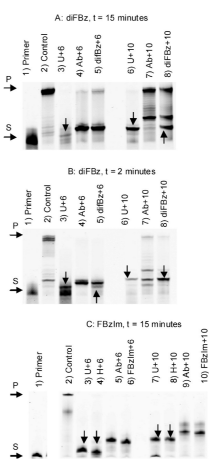


FIGURE 4: Primer extension reactions. Pfu-Pol was used to extend a Cy-5-labeled primer (24 bases in length) annealed to different templates (all 44 bases long) that place uracil (U), hypoxanthine (H), a stable abasic site (Ab), difuorobenzene (diFBz), and fluorobenzimidazole (FBzIm) either 6 (+6) or 10 (+10) bases ahead of the primertemplate junction. Controls use a template that contains only canonical DNA bases, resulting in full extension of the primer. U and H stall replication three to four bases prior to encounter, resulting in truncated products (such stalling products are highlighted with a vertical arrow). The abasic site also gives shortened products, but these are longer than those seen with U and H as, here, the polymerase stops directly at the abasic site. The positions of the starting primer and fully extended primer are shown with the horizontal arrows at the sides of the gel, labeled S and P, respectively. (A) Results seen with diFBz after 15 min. A stalled product (arrow) is seen with diFBz + 10but not with diFBz + 6. (B) Results found with diFBz after 2 min. Here a stalled product (arrow) can be seen with diFBz + 6, although the main product runs with that produced by the abasic site. (C) Results seen with FBzIm after 15 min. No evidence of stalling is visible; rather, extended products seen with this isostere correspond to those produced by the abasic site. Similar results were found at shorter times.

the modified base at position 4, a location that results in the tightest binding (3). As shown in Figure 5 and summarized in Table 2, a primer-template with uracil at position 4 bound strongly to the polymerase, giving a K_D value of ~ 5 nM. The binding of diFBz and FBzIm, located in an identical sequence, is clearly much weaker. In fact, no differences in affinity between FbzIm and a control, containing T at position 4, could be discerned; in both cases, a K_D of ~ 175 nM was found. Slightly better binding was seen with diFbz ($K_D \sim 50$ nM), but the U + 4 primer-template still interacts some 10-fold more strongly with the polymerase. The K_D values summarized in Table 2 agree reasonably well with those obtained previously for primer-templates using competitive fluorescence titrations (3), also

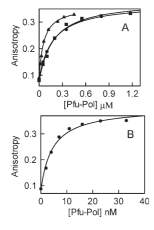


FIGURE 5: Binding of Pfu-Pol to Hex-labeled primer-templates (sequences given in Methods) containing varying bases at position 4 in the template strand: (A) diFBz + $4(\triangle)$, FBzIM + $4(\square)$, and T + 4(control) ($\hat{\bullet}$) and (B) U + 4 (\bullet). The K_D values found for these titrations are summarized in Table 2.

Table 2: Binding of Primer-Templates to P. furiosus DNA Polymerase

primer-template	$K_{\mathrm{D}}\left(\mathrm{nM}\right)^{a}$
control (T)	$182 \pm 23 (270)$
U diFBz	$4.4 \pm 1.3 (1.5)$ 52 ± 8
FBzIm	176 ± 19

^aThe binding constants for the interaction of primer-templates (sequences given in Methods) containing the bases indicated at position 4 were measured by direct titration (Figure 5). The values (±standard deviation) are the averages found from four determinations. The values given in parentheses for T and U were measured previously for analogous primer-templates using competitive titration (3).

mentioned in Table 2. Unfortunately, it proved impossible to apply the competitive approach, where a nonlabeled primertemplate is used to displace a hexachlorofluorescein-labeled uracil-containing oligodeoxynucleotide from the polymerase. The poor affinities of both diFBz and FBzIm necessitate the use of high concentrations to produce full or nearly full displacement, and the amount of material required was not available.

Single dNTP Incorporation into Primer-Templates. A final approach to determine the barrier to polymerization that arises from the presence of a deaminated base involves measuring the kinetic constants for the incorporation of one dNTP, under single-turnover conditions (8). To ensure complete binding of Pfu-Pol to the primer-templates, particularly those lacking deaminated bases, it is necessary to include an excess of PCNA. As found previously, PCNA increases the affinity of the polymerase for DNA, allowing saturation of the DNA (measured by attainment of the ceiling reaction velocity), at achievable concentrations of the enzyme (8). It is not necessary to add the PCNA loader (RF-C in archaea) or "block" the ends of the DNA, as used in studies of other polymerases, to realize this outcome (36). Values of k_{pol} (reaction rate under single-turnover conditions) and K_D (binding affinity) have been determined for primer-templates containing thymine (control), diFBz, and FBzIm at position 4, as shown in Figure 6 and summarized in Table 3. Very similar values of k_{pol} and K_D were observed with T and FBzIm, showing that the purine-based isostere does not result in any significant block to dNTP incorporation and the progression of the polymerase. With diFBz, the reaction was

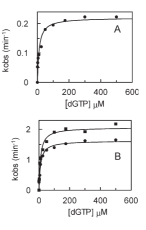


FIGURE 6: Incorporation of a single dGTP by Pfu-Pol into primertemplates containing varying bases in the template strand at position 4, under single-turnover conditions. Initially, k_{obs} was determined at various dGTP concentrations (data not shown) and used to generate secondary plots of $k_{\rm obs}$ vs [dGTP] for determination of kinetic parameters: (A) diFBz + 4 (\bullet) and (B) T + 4 (control) (\blacksquare) and

slowed by ~1 order of magnitude, as a result of a 10-fold decrease in k_{pol} and, consequently, a similar decrease in the specificity constant $k_{\text{pol}}/K_{\text{D}}$. Therefore, diFBz shows some degree of interaction with the deaminated base binding pocket, resulting in a reduced level of incorporation of incoming dNTPs and abatement in the rate of polymerization. With primer-templates containing U or H at position 4, incorporation of the single dNTP was too slow ($\sim 10^3 - 10^4$ -fold reduced when compared to that with T) to allow the determination of kinetic constants. However, Table 3 also gives results obtained in an earlier investigation with T, U, and H (8). The two studies used different temperatures (37 °C here and 50 °C previously), accounting for the lower k_{pol} observed for T controls in this investigation. Additionally, the changes in both primer-templates and incoming dNTP (dGTP here and dATP earlier) can account for the small changes in K_D . Nevertheless, it is abundantly clear that the decreases of $\sim 4 \times 10^3$ in $k_{\rm pol}$ and $k_{\rm pol}/K_{\rm D}$ seen with U and H, relative to the values seen for the T control, are much greater than the 10-fold reduction observed with diFBz. The results presented in this section are in agreement with those found above, confirming diFBz interacts with the deaminated base binding pocket slightly better than controls, but much more weakly than either U or H. With FBzIm, kinetic parameters identical to that observed with the control primer-template, containing T + 4, were found. Thus, in agreement with the primer-template extension and binding assays, FBzIm does not appear to interact with the deaminated base binding pocket.

DISCUSSION

Comparison of Uracil and Hypoxanthine Recognition. The X-ray structure of Tgo-Pol bound to a primer-template containing hypoxanthine elucidates how a deaminated purine is recognized and reveals the mechanism for specific binding of both a purine and a pyrimidine deaminated base. Comparison of the structure of apo Tgo-Pol (33) with a uracil-bound form (4) showed no conformational changes in the amino-terminal domain, which houses the deaminated base binding pocket. The hypoxanthine structure confirms the rigidity of this domain, with a preformed pocket poised to accept deaminated bases as they are encountered during replication. Comparing the uracil and

Table 3: Kinetic Parameters for the Interaction of dNTP with P. furiosus Polymerase-Primer-Template Complexes

primer-template	this study ^a		from ref 8^b			
	$k_{\text{pol}}^{c} (s^{-1})$	$K_{\mathrm{D}}{}^{c}\left(\mu\mathrm{M}\right)$	$k_{\rm pol}/K_{\rm D}^{c} ({\rm M}^{-1} {\rm s}^{-1})$	$k_{\text{pol}}^{c}(s^{-1})$	$K_{\mathrm{D}}{}^{c}\left(\mu\mathrm{M}\right)$	$k_{\rm pol}/K_{\rm D}^{c} ({\rm M}^{-1} {\rm s}^{-1})$
thymidine	2.1	9.7	1.7×10^{5}	7.7	45	1.5×10^{5}
diFBz	0.2	10.5	2.0×10^{4}	not determined		
FBzIm	1.6	10	1.6×10^{5}		not determine	ed
uracil	too slow for determination of $k_{\rm pol}/K_{\rm D}$			1.8×10^{-3}	52	35
hypoxanthine	too slow for determination of $k_{\rm pol}/K_{\rm D}$			2.5×10^{-3}	51	50

^aExperiments conducted at 37 °C with insertion of a single dGTP. ^bExperiments conducted at 50 °C with insertion of a single dATP. ^cThe kinetic parameters for the incorporation of a single dNTP into primer-templates (sequences given in Methods) containing the base indicated at position 4 were measured under single-turnover conditions (Figure 6). Values are the averages of three determinations and are accurate to ±25%.

hypoxanthine structures reveals remarkable superposition between atoms involved in both polar and hydrophobic protein-DNA contacts, with profound overlap of the two bases (Figure 3a). Uracil and hypoxanthine are discriminated from the four canonical DNA bases by a combination of shape match to the pocket and protein-base hydrogen bonds. The backbone amide nitrogen of Tyr-37 is critical for identifying an oxygen at the deaminated position (O4 in uracil and O6 in hypoxanthine), with the backbone amide of Ile-114 stabilizing binding by selecting a hydrogen bond acceptor (O2 in uracil and N3 in hypoxanthine) on the opposite side of the ring. With hypoxanthine, an additional, water-mediated hydrogen bond is observed between the ring NH1 group and Glu-111/Arg-119. As mentioned in Results, higher-resolution structures will be needed to unequivocally confirm if such an interaction is also formed to the equivalent NH3 position of uracil. Forming two hydrogen bonds to the C(6)O-N(1)H system of hypoxanthine [and possibly to the analogous C(4)O-N(3)H group of uracil provides excellent discrimination against adenine (and potentially cytosine), as in the latter two bases the hydrogen bonding pattern is reversed. This recognition pattern is analogous to that used by uracil DNA glycosylases to distinguish between uracil and cytosine (37, 38). The importance of Glu-111 and Arg-119 is confirmed by their high degree of conservation within the euryarchaeal family-B DNA polymerases and the observation that the R119A mutation no longer interacts with deaminated bases (4). While hydrogen bonding accounts for preferential binding of uracil and hypoxanthine, relative to their precursors, cytosine and adenine, rejection of thymine and guanine relies on shape matching and steric features. The additional bulk of thymine (5-CH₃ group) and guanine (2-NH₂ group) would not occupy the same spatial positions, should these two bases interact with the pocket in the same manner as uracil and hypoxanthine (Figure 3b). Therefore, different groups in the pocket are used for rejecting the thymine 5-methyl and guanine 2-amino functions. As described previously, the 5-methyl group is excluded by the side chains of Pro-36, Pro-90, and Phe-116 (4). By contrast, an exocyclic amino group at C2 of guanines would result in a steric clash with the main chain oxygen of Glu-111, thus precluding binding. This steric clash also explains the lack of binding seen with xanthine (the deamination product of guanine) (5), which contains an exocyclic oxygen at this position.

Coupling Deaminated Base Recognition to 3'-5' Exonuclease Activity. Polymerase stalling in response to uracil and hypoxanthine is a DNA damage avoidance mechanism, designed to avoid copying of these pro-mutagenic bases. With the U + 4 structure (4), the DNA is bound in an editing mode away from the polymerization active site, and additionally, the lack of

unwinding of the primer strand inhibits proofreading exonuclease activity (6). Thus, the nucleic acid is stably and inertly bound, presumably awaiting the recruitment of, as yet, unidentified repair proteins. However, stalling does not confer an absolute block on DNA replication, and prolonged incubation leads to read-through of deaminated bases (6). As the polymerase inches toward the lesion, uracil and hypoxanthine become progressively located three, two, and one base ahead of the primer-template junction, and this progression results in a profound stimulation of proofreading exonuclease activity (6). Thereby, the slowly, and inappropriately, extending primer is cut back, to reset the deaminated base to position 4. Such "idling" cycles of extension and degradation provide an additional barrier to replication beyond the damaged bases, as shown by the increased propensity of exo mutants to extend beyond uracil and hypoxanthine (6). The structure with hypoxanthine at position 2 rationalizes the link between deaminated base binding and activation of the polymerase proofreading exonuclease. The U + 4 and H + 2 structures are broadly similar with regard to base recognition, the disposition of the polymerase domains, and the localization of the bound nucleic acid. There exists, however, one key difference; with U + 4, the primer-template is fully basepaired, whereas with H + 2, the last two primer bases are singlestranded and located in the editing channel. The denaturing seen with H + 2 is mediated by the β -hairpin motif, a critical region in family B polymerases for the partition of primer strands between the polymerase and exonuclease active sites (39, 40). In an editing complex seen with the family-B polymerase from bacteriophage RB69, Arg-260, an amino acid found in the β -hairpin, interacts with the primer-template and appears to contribute to the melting of three primer bases and positioning of the resulting singlestranded primer region in the editing channel (32). It has been proposed that the equivalent amino acid in the archaeal family-B polymerase from Pyrococcus kodakaraensis KOD1, Arg-247, fulfills a similar role (41). With Tgo-Pol bound to hypoxanthine at position 2, Arg-247 pushes into the template strand, between the two bases in the double-stranded region immediately adjacent to the primer-template junction (G5 and A6). This wedgelike insertion pries G5 and A6 apart and unwinds the complementary primer bases, C28 and T27. Comparing the H + 2 and U + 4structures shows almost perfect superimposition of the β -hairpin (Figure 7), and the entire amino-terminal and exonuclease domains overlap almost completely. In particular, the separations among the deaminated base binding pocket, the β -hairpin, and the nearest double-stranded base pair are largely unchanged between the two structures (Figure 7). In the case of U + 4, this locates the hairpin near single-stranded template bases and both Arg-247 and the proximal bases appear disordered. With H + 2,

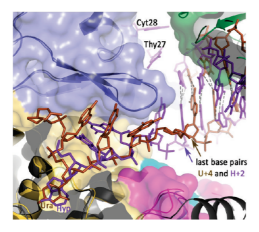


FIGURE 7: Relative spatial positions of uracil or hypoxanthine, the β -hairpin, and the nearest double-stranded base for the U + 4 and H + 2 structures. The two structures have been superimposed, and the space filling model for H + 2 is shown (thumb domain, green; exonuclease domain, blue; amino-terminal domain, yellow). The uracil-containing DNA is colored brown and the hypoxanthinecontaining DNA blue, with both the deaminated bases labeled. The β -hairpin (β -strand-loop- β -strand) is illustrated for both structures, with that of uracil colored slightly darker blue. The positions of the base pairs (last base pairs) nearest the deaminated bases and the β hairpin are marked with arrows, showing the relative disposition of these three elements is unchanged between the two structures. However, with U + 4, the last base pair contains the 3'-base of the primer. This is not the case with H + 2, as the two bases at the 3'-end of the primer (T27 and C28) become single-stranded and positioned in the editing channel of the polymerase.

the hairpin is close to the double-stranded region of the primertemplate and Arg-247 behaves differently, becoming ordered and acting as a wedge. Thus, the two polymerase domains differentially manipulate and distort the conformation of bound DNA, depending on the relative positions of the deaminated base and the nearest base pair, thereby activating exonuclease activity with U/H at position 2. With euryarchaeal polymerases from *Thermo*coccus and Pyrococcus species, sequence alignment shows that amino acid 247 is Arg (21), Met (8, with Pfu-Pol having this residue), or Ser (9) (numbers in parentheses refer to the frequency of occurrence). While the long side chain of Met could act like Arg inserting between bases, it is unclear if the much shorter Ser could fulfill this role. The same alignment shows that the amino acid at position 246, which forms a hydrogen bond with base A6, is overwhelmingly Asp, which occurs 36 times. There are single instances of Glu and Gln at this location, but all three bases are compatible with the function proposed and, in principle, able to hydrogen bond to any standard DNA base at this position. However, other species of euryarchaeal family-B polymerases show very little amino acid conservation at these two locations. This may be related to variability in the overall size of the β -hairpin observed between polymerases, e.g., from phage RB69 (31) and herpes simplex virus (42), and the tendency for structural rather than amino acid sequence conservation (39). In contrast, amino acid 261, which prevents rehybridization by blocking access of T27 to A6, is functionally well conserved in all euryarchaeal family-B polymerases, being either an aromatic or bulky hydrophobic residue: Y(51), F(12), W(12), or L(21).

Hydrogen Bonding and Shape Recognition. To assess the relative contributions of hydrogen bonds and shape recognition to specific recognition of deaminated bases, we made use of the uracil and hypoxanthine isosteric analogues illustrated in Figure 1. With difluorobenzene, some interaction with the

polymerase is seen using primer extensions, binding assays, and single-turnover dNTP incorporation. However, the preference over controls is only 4- and 10-fold (binding and single turnovers, respectively), significantly lower than that observed with uracil. In the case of fluorobenzimidazole, none of the assays reveal any indication of positive binding to the polymerase. The stronger binding of diFBz relative to FBzIm mimics the behavior seen with their parent bases, uracil and hypoxanthine, respectively (5, 8). diFBz and FBzIm are shape-matched to uracil and hypoxanthine but cannot participate in polar interactions; therefore, we can conclude that the hydrogen bonds seen between the polymerase and the deaminated bases play a key role in selectivity. The ability to specifically bind the damaged bases uracil and hypoxanthine, which occur at very low levels in DNA, while simultaneously rejecting the four canonical bases, present in large excess, is demanding. Therefore, it appears likely that the polymerase would make use of every distinguishing feature of the two deaminated bases. It is possible to differentiate between the uracil/thymine and hypoxanthine/guanine pairs using size criteria, as the canonical bases are larger. Indeed, rejection of thymine and guanine relies largely on the pocket being shaped to prevent binding of the additional 5-CH₃ and 2-NH₂ groups on steric grounds. However, it is almost impossible to distinguish the near isosteric bases uracil/cytosine and hypoxanthine/adenine in this manner, and for this purpose, the polymerase makes use of hydrogen bonding. The critical nature of these interactions accounts for the loss of tight binding when diFBz and FBzIm are used.

DNA polymerases are able to accept nonpolar isosteric base analogues with relatively high efficiency during replication, inferring a preference for the production of a correctly shaped base pair, rather than Watson-Crick hydrogen bonds (12-15). However, in this work, isosteres have been used to probe specific recognition of a damaged base by the polymerase, a DNA repair response, rather than DNA synthesis itself. Examples of the use of isosteric bases for investigation of DNA repair include Fpg (20) and AlkA (43), which target 8-oxoguanine and hypoxanthine, respectively. Locating an isostere opposite the damaged base, in double-stranded DNA, usually results in faster excision, as abolition of Watson-Crick base pairs facilitates flipping of the damaged base into the enzyme active site. However, in both cases, the isostere does not directly probe recognition of the damaged base by the protein but rather the influence of base pairing on overall repair. MutY removes a canonical DNA base, adenine, when paired opposite the damaged base 8-oxoguanine. In general, replacing adenine with an isosteric analogue results in better binding by promoting base flipping (18, 44). However, catalysis is severely compromised, and a study with adenine isosteres in which the ring nitrogens were gradually replaced uncovered a critical role for N3 and N7 (20). Recognition of uracil and hypoxanthine by DNA polymerases differs, somewhat, from the repair systems previously analyzed. No catalytic activity is manifested, and the polymerase interacts with the deaminated bases in single-stranded DNA, removing any influence of complementary bases. Nevertheless, the results found with the polymerase are reminiscent of those seen with MutY; isosteric bases greatly weaken binding in the case of the polymerase and drastically slow catalysis with MutY. It is clear that base pairs can be recognized largely by shape and steric factors, as seen with DNA polymerases during replication. However, it remains to be demonstrated whether such a mechanism is adequate for highly specific interaction of a single isolated

base with a protein. With archaeal polymerases, faithful recognition of deaminated bases is clearly dependent on hydrogen bonding. However, as the enzyme must recognize both uracil and hypoxanthine, extreme steric tailoring, needed to distinguish near isosteric base combinations such as uracil/cytidine and hypoxanthine/adenine, is almost certainly precluded. Therefore, the question of shape recognition of single bases by other proteins remains an open one.

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