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# Y-Family DNA polymerases may use two different dNTP shapes for insertion: A hypothesis and its implications

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#### ABSTRACT

Chemicals and radiation can damage DNA leading to the formation of adducts/lesions, which - if not removed by DNA repair pathways - usually block replicative DNA polymerases (DNAPs). To overcome such potentially lethal blockage, cells have lesion bypass DNAPs, which are often in the Y-Family and include several classes. One class includes human DNAP K and E. coli DNAP IV, and they insert dCTP in the non-mutagenic pathway opposite [+ta]-B[a]P-N<sup>2</sup>-dG, which is the major adduct formed by the environmental carcinogen benzo[a]pyrene. Another class includes hDNAP  $\eta$  and ecDNAP V, and they insert dATP opposite [+ta]-B[a]P-N<sup>2</sup>-dG in the dominant  $G \rightarrow T$  mutagenic pathway. Herein we develop a hypothesis for why the  $IV/\kappa$ -class preferentially does cellular dCTP insertion. On the minor groove side of the active site, Y-Family DNAPs have a cleft/hole that can be analyzed based on an analogy to a "chimney." Our models of DNAP IV show a large chimney opening from which the pyrene of [+ta]-B[a]P-N<sup>2</sup>-dG can protrude, which allows canonical adduct-dG:dCTP pairing. In contrast, our models of DNAP V have small chimney openings that forces adduct-dG downward in the active site such that canonical adduct-dG:dCTP pairing is not possible. Based on X-ray structures, sequence alignment and our modeled structures of Y-Family DNAPs, chimney opening size seems primarily controlled by one amino acid ("flue-handle"), which dictates whether nearby amino acids ("flue") plug the chimney or not. Based on this analysis, a correlation is apparent: the flue is closed in  $V/\eta$ -class DNAPs giving small chimney openings, while the flue is open for the  $IV/\kappa$ -class giving large chimney openings. Secondarily, a hypothesis is developed for why the  $V/\eta$ -class might preferentially do cellular dATP insertion opposite [+ta]-B[a]P-N<sup>2</sup>-dG: the small chimney forces adduct-dG lower in the active site, possibly leading to catalysis using a non-canonical dNTP shape that permits syn-adenine:adduct-dG base pairing. In summary, a hypothesize is developed that the pyrene moiety of [+ta]-B[a]P-N<sup>2</sup>-dG protrudes from the large chimney opening of DNAP IV, thus permitting canonical dCTP:adduct-dG pairing, while the small chimney opening of DNAP V forces [+ta]- $B[a]P-N^2$ -dG lower down in the active site, in which syn-adenine can pair with adduct-dG via a non-canonical dNTP shape.

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# 1. Introduction

Cells possess many DNA polymerases (DNAPs); e.g., human cells, yeast (*S. cerevisiae*) and *E. coli* have at least fifteen, eight and five, respectively [1–3]. Some DNAPs are present to do translesion synthesis (TLS) past DNA lesions/adducts that result from radiation and chemical insults [1–9]. Most lesion-bypass DNAPs are in the Y-

Abbreviations: B[a]P, benzo[a]pyrene; +BP, [+ta]-B[a]P-N²-dG (Fig. 1); -BP, [-ta]-B[a]P-N²-dG (Fig. 1); TLS, translesion synthesis: the insertion of a base opposite a DNA adduct as well as subsequent elongation; DNAP, DNA polymerase; S1-dNTP, dNTP active shape 1; S2-dNTP, dNTP active shape 2; MD, molecular dynamics; TT-CPD, thymine-thymine cyclopyrimidine dimer.

Family [1–9], where humans have three members (hDNAPs  $\eta$ ,  $\iota$  and  $\kappa$ ), yeast has one (scDNAP  $\eta$ ) and E. *coli* has two (ecDNAPs IV and V).

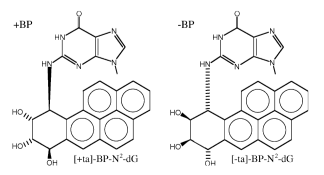
Y-Family DNAPs have a conserved  $\sim 350aa$  core, which includes the polymerase active site (representative Refs.: [10–18]). As with all DNA polymerases, Y-Family members resemble a right-hand with thumb, palm and finger domains, although their "stubby" fingers and thumb result in more solvent accessible surface around the template/dNTP-binding pocket [7]—presumably to accommodate the bypass of bulky and/or deforming DNA adducts/lesions, which protrude into these open spaces. Y-Family DNAPs grip DNA with an additional domain [7,10,11,15], usually called the "little finger". Steps in the mechanism of Y-Family DNAPs have been proposed for both protein structural changes based on a series of X-ray structures [19,20] and for chemical catalysis based on theoretical studies [21].

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Our work has focused on benzo[a]pyrene (B[a]P), which is a well-studied DNA damaging agent that is a potent mutagen/ carcinogen and an example of a polycyclic aromatic hydrocarbon (PAH), a class of ubiquitous environmental substances produced by incomplete combustion [22,23]. PAHs in general and B[a]P in particular induce the kinds of mutations thought to be relevant to carcinogenesis and may be important in human cancer (representative Ref.: [24]). B[a]P mutational spectra were established with the major metabolite that reacts with DNA, namely (+)-anti-B[a]PDE, in *E. coli* [25], yeast [26,27] and mammalian (CHO) cells ([28] and references therein). Mutagenesis has also been studied with [+ta]-B[a]P-N<sup>2</sup>-dG (+BP, Fig. 1), the major adduct of (+)-anti-B[a]PDE, and  $G \rightarrow T$  mutations predominate in most cases ([29], and references therein).

DNAPs IV and V of E. coli are both involved in translesion synthesis (TLS) with B[a]P-N<sup>2</sup>-dG adducts, although evidence suggests they play very different roles. In studies with purified proteins, DNAP IV inserted dCTP (>99%) opposite both +BP and [-ta]- $B[a]P-N^2-dG$  (-BP) in a 5'-CGA sequence, while DNAP V inserted dATP (>99%) [30], and this tendency is evident in *E. coli* where DNAP V inserts dATP opposite +BP in the  $G \rightarrow T$  pathway [31], while in the non-mutagenic pathway DNAP IV does dCTP insertion opposite +BP [32-36], -BP[34] and other  $N^2$ -dG adducts [35,36]. Regarding the non-mutagenic pathway, only DNAP IV is required for efficient TLS with -BP [34], while DNAP V is required in addition to DNAP IV with +BP [32-36]. Why are two DNAPs required for non-mutagenic TLS with +BP: certain lesions need one DNAP for insertion and a second for extension [37,38]. Thus, if DNAP IV does dCTP insertion [30,32-36], then DNAP V must do extension, which is sensible given kinetic findings with purified proteins, where DNAP V can be up to  $\sim$ 1500 times better than DNAP IV at the step directly following adduct-G:C formation (i.e., extension) in the case of +BP compared to -BP (discussed in greater detail in Ref. [34]). Finally, random mutagenesis studies with [+anti]-B[a]PDE suggest that most  $G \rightarrow T$  mutations with B[a]P-adducts require SOS-inducible Y-Family DNAPs, although a minor non-SOS-inducible  $G \rightarrow T$  pathway does exist (discussed in Ref. [34]), which has been studied with +BP in a 5'-GGA

The study of *E. coli*'s Y-Family DNAPs may provide insights about Y-Family DNAPs in general. Human DNAP  $\kappa$  was originally discovered because its sequence closely resembles *E. coli* DNAP IV [39–41], and dNTP insertion opposite a variety of adducts/lesions, including +BP, is remarkably similar for the DNAP IV/ $\kappa$  pair (Table 1), suggesting they are functional orthologs (discussed in Ref. [42]). This notion was substantiated when the identical mutation in a conserved residue (the "steric gate", which excludes rNTPs) in the active site of DNAP IV and DNAP  $\kappa$  had a similar effect on lesion bypass vs. normal replication both *in vitro* and in cells [35]. DNAPs IV and  $\kappa$  have been shown to accurately bypass a variety of N²-dG-adducts [32–36], including those formed via various cellular trioses [36], which may be the main cellular



**Fig. 1.** Structures of +BP ([+ta]-B[a]P-N<sup>2</sup>-dG) and -BP ([-ta]-B[a]P-N<sup>2</sup>-dG).

**Table 1** Dominant dNTP insertions opposite various DNA adducts/lesions by *E. coli* DNAPs IV and V and human DNAPs  $\kappa$  and  $\eta$ .

Lesion	DNAP V	DNAP η	DNAP IV	DNAP ĸ
[+ta]-BP-N <sup>2</sup> -Dg	A/C	A > G	С	С
AAF-C8-dG	C	С	C/T	C/T
AF-C8-dG	-	-	C	С
TT-CPD	AA	AA	n	n
T(6-4)T	AG	nG	n	n
AP site	Α	Α	n	A*

<sup>&</sup>lt;sup>a</sup> Dominant dNTP insertion using purified DNAPs, where "n" indicates "no" or low activity, "A\*" indicates bypass by an unusual mechanism, and "–" indicates data unavailable. Data as reviewed in Ref. [41].

rationale for the genesis of the  $IV/\kappa$ -class. *E. coli* DNAP V and human DNAP  $\eta$  are also functional orthologs, based on their similarity of dNTP insertion opposite a variety of adducts/lesions (Table 1 [42]), and a strong case has been made that the main cellular rationale for the  $V/\eta$ -class is TLS of UV-induced CPDs (discussed in Ref. [43]).

There must be structural reasons why the insertion preference opposite adducts/lesions is different for the DNAP IV/ $\kappa$  pair vs. the DNAP V/ $\eta$  pair (Table 1 [42]). However, no X-ray structures exist for UmuC (the polymerase subunit of DNAP V), DNAP IV or hDNAP  $\eta$ , which prompted us to build models taking a homology modeling approach [42]. Analysis of X-ray structures, modeled structures and sequence alignment suggests that Y-Family DNAPs lend themselves to accurate homology modeling [42,43]. An X-ray structure does exist for hDNAP  $\kappa$  with DNA [18],

Herein a molecular modeling study is presented that suggests a hypothesis for why the IV/κ-class correctly inserts dCTP opposite B[a]P-N<sup>2</sup>-dG adducts. To form an adduct-dG:dCTP base pair, the B[a]P moiety must be in the developing minor groove, since the adduction site (N<sup>2</sup>-dG) is in the minor groove in a Watson-Crick base pair. On the minor groove side, Y-Family DNAPs have an opening (or gap) next to the active site between the fingers and little finger domains. This opening looks like an elliptical hole of varying size in Dpo4 [11–14], Dbh [10], hDNAP ι [17] and in our models of DNAP IV and UmuC(V) (see below), while it looks like a slot in hDNAP κ [18]. The character of this opening can be analyzed based on analogy to a "chimney," where a cluster of nearby amino acids can be thought of as a "flue" that either plugs the chimney leaving a small opening, or does not plug the chimney leaving a large opening. Next to the flue is a single amino acid, which can be thought of as a "flue-handle" that controls whether the flue is open or closed. Three regions of the protein make up the chimney as shown in Fig. 2 for our model of DNAP IV: an upper lip (turquoise), a lower lip (dark blue) and a left lip (blue) [43]. It is not unreasonable to think that chimney opening size and shape might influence dNTP insertional mechanism given that the bulky B[a]P moiety (red, Fig. 2) must fit in it.

Herein we show that DNAP IV has a large chimney opening, which allows +BP to pair with dCTP, when dCTP adopts the canonical shape observed in all other families of DNAPs. In contrast, our molecular models suggest that UmuC(V) has a small chimney opening, which forces +BP downward in the active site into a position where catalysis is less likely to be facile.

Secondarily, we reflect on the conundrum: if the small chimney opening of DNAP V enforces a structure that seems unlikely to be active, then how might DNAP V insert dATP opposite +BP? Dpo4 is the best studied Y-Family DNAP in terms of structure (representative Refs.: [11–14]). Though the dNTPs in the active site of many Dpo4 structures adopt the canonical "chair-like" dNTP shape (Fig. 3, green), which is similar to the shape of dNTPs in the active sites of other families of DNAPs (Fig. 3, insert), a second non-canonical

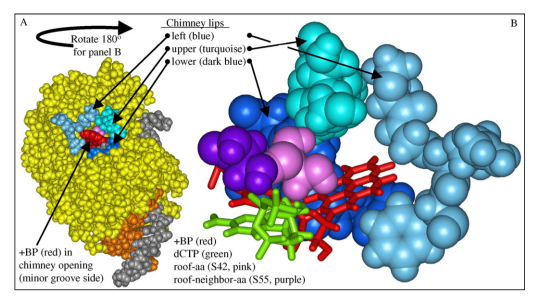
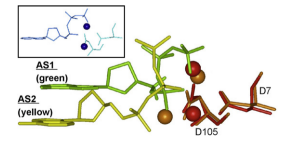


Fig. 2. DNAP IV model showing the "chimney" and "roof" regions. Panel A: View from the minor groove side of DNAP IV (yellow), showing the "chimney" opening (cleft or hole), which is encircled by the upper lip (turquoise), left lip (blue) and lower lip (dark blue), along with +BP (red) emerging from the opening. The template (gray) and primer (brown) are also shown. The view in panel A is rotated approximately 180° to give panel B, which shows the chimney (white space) between the three chimney lips. Also shown are the roof-aa (S42, pink), which is a positionally conserved residue that lies above the nucleobase of the dNTP, the "roof-neighbor-aa" (S55, purple), which is next to the roof-aa and also contacts the dNTP nucleobase (dCTP, green), and the +BP (red), whose pyrene moiety must fit in the chimney in order for the dG moiety of +BP to pair with dCTP

"goat-tail-like" shape has also been observed (Fig. 3, yellow), as noted by others [13]. (For simplicity we refer to the "chair-like" shape as S1-dNTP and the "goat-tail-like" shape as S2-dNTP.) Based on X-ray structures, modeled structures, sequence alignment and experimental results, a logical framework emerges from the hypothesis that the syn-adenine orientation of S2-dNTP pairs with adduct-dG, since S2-dNTP sits lower down in the active site and permits +BP to fit under the small chimney of UmuC(V). This hypothesis depends on the non-canonical S2-dNTP shape being active, which is considered in Section 4. The goal of the modeling work herein was to probe the insertional differences between the V/  $\eta$ -class and IV/ $\kappa$ -class of Y-family DNAPs and develop hypotheses, which we intend to test experimentally.

# 2. Methodology

Methods were identical to those described previously [43], except as noted below. Molecular dynamics (MD) runs were done with CHARMM 30 using Boston University's IBM Power4 p690 or IBM BlueGene. Starting coordinates for DNAP IV and UmuC(V) were derived from refined structures in our previous work [42],



**Fig. 3.** Side view of a dNTP in the "chair-like" shape S1-dNTP (green) vs. the "goat-tail-like" shape S2-dNTP (yellow). Key amino acids (only two are shown) from a Dpo4 structure adopting the S1-dNTP shape were superimposed on the same amino acids in a Dpo4 structure adopting the S2-dNTP. Spheres are divalent cations (S1-dNTP/red and S2-dNTP/brown). The insert shows that dNTP shape in the T7 DNAP active site resembles S1-dNTP (green). X-ray coordinates are from 1SOM-B for Dpo4/S1-dNTP, 1RYS-A for Dpo4/S2-dNTP, and 1T7P for T7 DNAP.

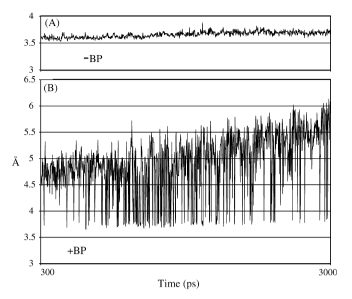
from which unadducted DNA was removed and replaced with adducted S1-dNTP or S2-dNTP DNA derived from our modeling work with Dpo4 [43]. Water was added [43]. The MD protocol (outlined here) is slightly modified from our previous work [43]. Water alone was minimized (200 steps of ABNR), after which water and protein were minimized (200 steps of steepest descent), and then, finally, water, protein and the phosphates of the dNTP were minimized (200–500 steps of steepest descent). The structures were heated to 300 K over 40 ps using the velocity Verlet integration method (time step 1 fs) with a harmonic force constraint of 5 kcal/mol per Å<sup>2</sup> on all nucleobase atoms. Thereafter, MD was conducted at 300 K for 250 ps, whereupon the constraint was reduced to 0.5 kcal/mol per Å<sup>2</sup> for 50 ps. The run was completed with an additional 1000 ps of unconstrained MD (e.g., see Fig. 4). During MD, electrostatic, van der Waals interactions and non-bonded Lennard-Jones interactions were smoothed to zero at 11 Å, and the SHAKE algorithm was used to constrain bonds with hydrogen atoms using a tolerance of 0.0005 Å. Each structure in a figure is a snapshot taken from the last 50 ps of a MD run and is representative, as all structures were more-or-less similar, except as noted in Section 3. Structures are viewed after minimization (200 steps of ABNR) to remove distortions due to the elevated temperatures.

X-ray coordinates in some figures were taken from the RSCB Protein Data Bank [44], and include Dpo4 with an S1-dNTP shape (1SOM-B), Dpo4 with an S2-dNTP shape (1RYS-A), Dpo4 with +BP (2IA1)), T7 DNAP (1T7P), hDNAP  $\kappa$  (2OH2), yDNAP  $\eta$  (1JIH).

#### 3. Results

#### 3.1. The canonical S1-dNTP "Chair-like" shape

In duplex DNA, the pyrene moieties of —BP and +BP (when paired with dC) are oriented toward the 3′- and 5′-base [45]. Herein we assume these same orientations for the pyrene moieties in UmuC(V) and DNAP IV. (Though we have not completed a thorough study, —BP in the BPmi5 conformation looks reasonably similar to +BP in BPmi5, while +BP in a BPmi3 conformation looks reasonably



**Fig. 4.** Distance between the O3'-primer and  $P\alpha$ -dCTP during the final 1 ns of the MD runs with -BP (panel A) and +BP (panel B) for DNAP IV/S1-dNTP.

similar to -BP in BPmi3. No observations from the work herein depend on the unique assignment of -BP to BPmi3 or +BP to BPmi5, and, thus, this assumption is not critical to our analysis.)

DNAP IV inserts dCTP opposite -BP (Section 1), and this possibility was modeled using the canonical S1-dNTP shape. During the final 1 ns of the MD run, the structure is stable, appeared reasonable, and showed no significant distortions (Fig. 5C) when compared to the control structure (Fig. 5A) in which unadducted-dG was paired with S1-dCTP. (In each panel of Fig. 5, the left structure shows the region around the roof-aa/roof-neighbor-aa, while a  $\sim 90~^{\circ}$ C rotation on the horizontal gives the right structure, which is of the chimney region.) For example, the distance between primer-O3′ and P $\alpha$ -dNTP was reasonably constant for the -BP structure (Fig. 4A) and averaged 3.66 Å, which was comparable to the dG no-adduct control structure that averaged 3.70 Å. These distances are close to a van der Waals contact ( $\sim 3.5~$ Å) and can be considered "reaction-ready."

DNAP IV also inserts dCTP opposite +BP (Section 1), which we modeled using the BPmi5 orientation opposite S1-dCTP. In this case, the MD run oscillated between two structures (Fig. 4B), one being reaction-ready (Fig. 5E, primer-O3'/dNTP-P $\alpha$  distance, 3.66 Å), while the second was not reaction-ready (not shown, 5.12 Å). Since the reaction-ready structure presents a significant fraction of time ( $\sim$ 20%), we only consider it. (With the exception of dCTP insertion opposite +BP by DNAP IV, all other MD runs looked stable as exemplified by Fig. 4A.)

With UmuC(V) and S1-dNTP, both -BP and +BP gave structures (Fig. 5D and F, respectively) with O3'-P $\alpha$  distances that were not reaction-ready (4.93 Å and 5.12 Å, respectively). Why did UmuC(V)/S1-dNTP not give reaction-ready distances, when DNAP IV/S1-dNTP did? DNAP IV has a large chimney opening (right side structures in Fig. 5A, C and E), while UmuC(V) has a smaller chimney opening (right side structures in Fig. 5B, D and F) into which (e.g.) the pyrene moiety of +BP does not readily fit without distortion/reorientation. During the MD run, this problem is resolved by opposing motions: the chimney opens slightly, but principally +BP is pushed downward and away from the chimney/minor groove and toward the major groove. As the adduct-dG moves so does its paired dCTP, thus forcing P $\alpha$  away from the primer-O3'. In fact, during the S1-dNTP/UmuC(V) MD run, adduct-dG:dCTP moves about halfway toward S2-dNTP/UmuC(V) posi-

tioning. Other distortions result, including the loss of co-planarity between the adduct-dG:dCTP and the base pair below it (not shown). –BP experiences similar problems, though it is the saturated ring of the B[a]P moiety that clashes with the small chimney (Fig. 5D).

## 3.2. The non-canonical S2-dNTP "Goat-tail-like" shape

A non-canonical S2-dNTP shape ("goat-tail-like") is also frequently observed in Dpo4 X-ray structures (Introduction), which we also consider. With the S2-dNTP shape in UmuC(V), both -BP and +BP show reaction-ready structures (Fig. 5] and L, respectively) with primer-O3'/dNTP-P $\alpha$  distances of 3.77 Å and 3.85 Å, respectively. Why did S2-dNTP/UmuC(V) give reactionready distances, when S1-dNTP/UmuC(V) did not? The dNTP shape in S2-dNTP is lower down compared to S1-dNTP (Fig. 3), and, thus, +BP is also lower down, allowing the pyrene to fit comfortably below the small chimney opening (Fig. 5F), with the consequence being that  $P\alpha$ -dCTP stays near the primer-O3'. With dCTP being lower down in S2-dNTP, a gap exists above the nucleobase in the initial structure; this gap diminishes during the MD run due to slight rearrangements of the bulky sidechains of the roof-aa I38 and its roof-neighbor-aa M51. The observations are similar for -BP (Fig. 5]).

With the S2-dNTP shape, both -BP and +BP gave distorted structures in DNAP IV (Fig. 5I and K, respectively) and non-reaction-ready primer-O3'/dNTP-P $\alpha$  distances of 4.52 Å and 4.85 Å, respectively. Why did DNAP IV/S2-dNTP not give reaction-ready distances and reasonable structures, when DNAP V/S2-dNTP and DNAP IV/S1-dNTP did? DNAP IV has a less bulky roof-aa (S42) and a less bulky roof-neighbor-aa (S55), neither of which has a sidechain that is big enough to rearrange and fill the gap above the nucleobase of dCTP in S2-dNTP. During the MD run, the deoxyribose of dCTP rises to try to fill in this gap (though the gap is still large in the final structure, Fig. 5I and K) and the phosphates rearrange into a "straight-chain" orientation, which is non-canonical and has not been observed to the best of our knowledge.

### 4. Discussion

We first consider a hypothesis for why chimney opening size might be different in DNAP IV vs. UmuC(V), before assessing how chimney size might affect dNTP insertion pattern. Finally, we discuss why in cells DNAP IV might insert dCTP, while DNAP V inserts dATP (Fig. 6), which are hypotheses that we intend to test experimentally, which was the goal of the modeling work described herein.

# 4.1. Structural basis for a large vs. a small chimney opening

What structural difference(s) in DNAP IV vs. UmuC(V) might result in a large vs. a small chimney opening, and is this structural difference(s) conserved in other Y-Family DNAPs in the IV/ $\kappa$ -class vs. the DNAP V/ $\eta$ -class? The chimney has three sides (Figs. 2 and 5): the upper lip (numbered aa32–35 in DNAP IV, turquoise) and left lip (aa73–76, blue) are in the fingers domain, while the lower lip (aa244–247, dark blue) is in the little finger domain. The amino acids associated with the chimney lips can be assigned reliably based on conserved amino acids. For example, the region surrounding the chimney upper lip for the IV/ $\kappa$ -class and V/ $\eta$ -class (as shown in Fig. 7) have 22 and 18 conserved amino acids, respectively, and, of these, seven are identical between the sets (boxed in gray in Fig. 7A and B). The assignment of amino acids to the left lip and lower lip are similarly anchored by the alignment of nearby conserved amino acids.

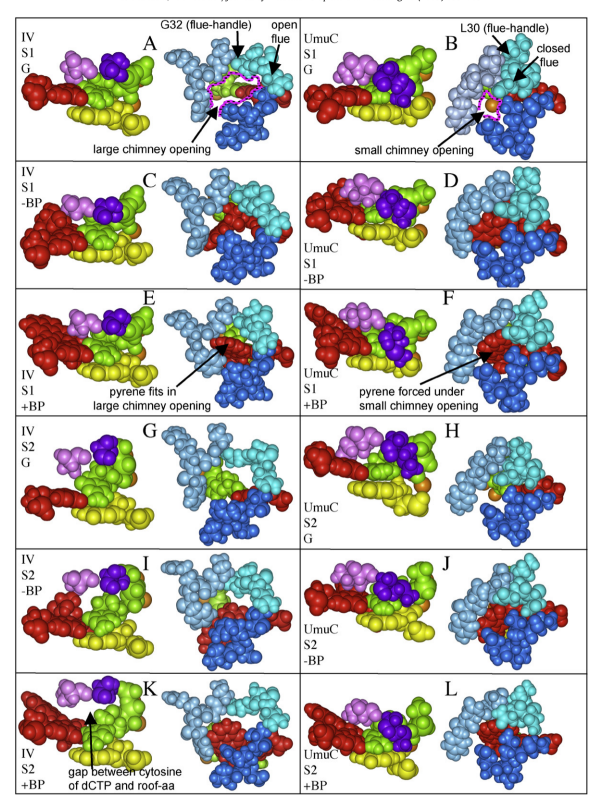
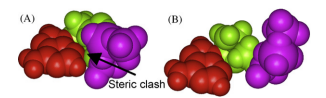


Fig. 5. (A–L) Regions of DNAP IV and UmuC(V) from our MD runs, in S1-dNTP or S2-dNTP with no adduct, –BP, or +BP. In each panel, the left structure shows the roof-aa (pink), the roof-neighbor-aa (purple), the dCTP (green), the primer (yellow), and the template adduct-dG (red), while the right structure shows the three lips of the chimney opening (turquoise/blue/dark blue) for DNAP IV (aa31-34/aa72-75/aa244-247) or UmuC(V) (aa30-33/aa71-74/aa255-258).

The chimney upper lip (turquoise, Fig. 5) is closest to the active site, and principally defines whether the chimney can accommodate the bulky B[a]P moiety. The first amino acid in the upper lip of DNAP IV is glycine (G32), and all IV/ $\kappa$ -class members from all of the species listed in Fig. 7A have a glycine at this position, often

embedded in a VGS sequence. The one X-ray structure available for this class, hDNAP  $\kappa$  [18], shows that this glycine (G131, turquoise, Fig. 8A) is followed by upward curvature of the chimney upper lip (red arrow, Fig. 8A). This glycine can be thought of as a "fluehandle" whose backbone properties allow this upward curvature



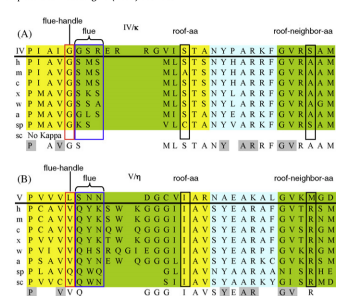
**Fig. 6.** Adenine (red) in the *syn*-orientation sterically clashes with the deoxyribose (green) and  $P\alpha$  (purple) when dATP is in the canonical "chair-like" shape S1-dNTP (panel A), while no such steric clash exists in the "goat-tail-like" shape S2-dNTP (panel B).

(discussed below), with the consequence being that the R-groups on the next several amino acids ("flue"; S132/R133, blue in Fig. 8B) are turned away from the chimney opening, which remains open. Our models of DNAP IV also have this upward curvature (Fig. 5) with an open flue, which depends on an analogous flue-handle glycine (G32).

In contrast all DNAP V/ $\eta$ -class members have bulky "flue-handles," which is usually valine in a VVQ sequence (Fig. 7B). The one X-ray structure in the V/ $\eta$ -class, scDNAP  $\eta$  [16], shows that its bulky V54 flue-handle (turquoise, Fig. 8D) is associated with downward curvature of the chimney upper lip (red arrow), which forces the "flue" (Q55/Y56, blue in Fig. 8E) to plug the chimney. In UmuC(V) the sequence is slightly different (VLSN), though the outcome is the same: the bulky L30 flue-handle causes downward curvature, and an asparagine (N32) plugs the chimney giving a closed flue (Fig. 5B).

Glycine has greater flexibility in its  $\phi/\varphi$ -angles compared to other amino acids, which appears to be the fundamental conformational reason why a glycine vs. a non-glycine flue-handle gives different curvature of the chimney upper lip. Fig. 9 is a Ramachandran plot for all valines, leucines and cysteines from Xray structures of Dpo4 [11] hDNAP κ [18] and scDNAP η [16]. Fig. 9 also shows the  $\phi/\phi$ -angles for the flue-handles in scDNAP  $\eta$  (V54, blue circle), hDNAP n (V37, red circle), UmuC(V) (L30, yellow circle) and Dpo4 (C31, pink circle), which are all clustered in the region expected for extended (β) secondary structure, with the consequence being that this combination of  $\phi/\varphi$ -angles leads to downward curvature of the chimney's upper lip, causing the flue to plug the chimney and the chimney opening to be small. In contrast, the angles for the glycine flue-handles are outside this region for hDNAP κ (G131, green square) and DNAP IV (G32, purple square), which results in upward curvature of the chimney's upper lip, thus opening the flue and giving a large chimney opening. (Given the discontinuous nature of Ramachandran plots, the  $\phi/\varphi$ -angles for DNAPs IV and κ look farther from the valine/leucine/cysteine cluster than they actually are.)

Glycines are *not* excluded from adopting the  $\phi/\varphi$ -angles observed in the valine/leucine/cysteine cluster, so in principle IV/κ-class DNAPs could have a closed flue leading to a small chimney opening as in the DNAP V/n-class. Our model for DNAP IV began with the Xray backbone coordinates of Dpo4, whose non-glycine flue-handle (C31) leads to downward curvature of the upper lip and a closed flue in X-ray structures ([11-14], as well as other Dpo4 structures). In spite of the downward curvature in the starting structure (derived from Dpo4) in our DNAP IV model, its flue-handle (G32) always rearranged during MD runs [42] to give upward curvature (e.g., Fig. 5A), resulting in a final structure with an open flue, which resembles the hDNAP K X-ray structure (Fig. 8B). This finding implies that glycine flue-handles prefer the "up" orientation, which appears to be stabilized by several interactions—notably by a R34/E25 salt bridge in our model of DNAP IV. In DNAP κ, a M133/I166 hydrophobic interaction is the major contact favoring the uporientation for the glycine flue-handle.



**Fig. 7.** Amino acid alignment of several key regions of representative Y-Family DNAPs in the IV/κ-class (panel A) and the V/η-class (panel B). The R-groups of the final three amino acids ("flue") of the chimney's upper lip either don't plug (IV/κ-class) or plug (V/η-class) the chimney, depending upon whether the first amino acid in the upper lip ("flue-handle") is glycine (flue open, IV/κ-class) or is a bulky aa (often valine, flue closed, V/η-class). The lowest line in each panel shows conserved residues, with residues conserved (≥75%) between the IV/κ-class and the V/η-class are highlighted in gray. Abbreviations: ss (Sulfolobus solfataricus) h, human (*Homo sapien*); m, mouse (*Mus musculus*); c, chicken (*Gallus gallus*); x, frog (*Xenopus laevis*); w, worm (*Caenorhabditis elegans*); a, plant (*Arabidopsis thaliana*); sp, fission yeast (*Saccharomyces pombe*); sc, budding yeast (*Saccharomyces cerevisae*). Colors: blue, α-helicies; yellow, extended (β); and green, loops.

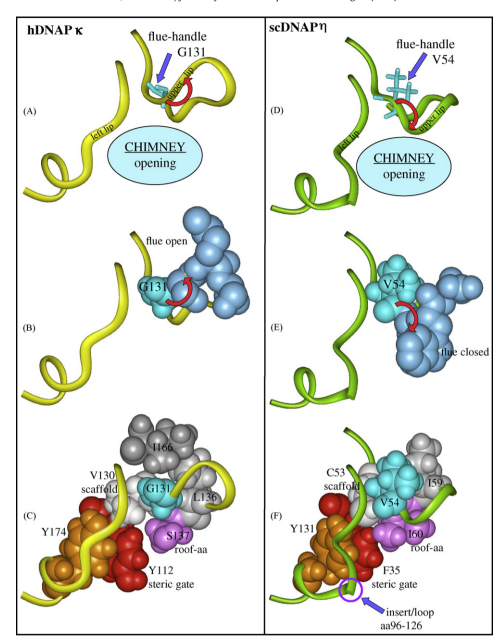
We could find no other reasonable structural rationale for the difference in chimney opening size. For example, loop-size per se does not correlate, since UmuC(V) and DNAP  $\kappa$  in most cases have relatively small loops, while DNAPs IV and DNAP  $\eta$  have relatively large loops (leftmost green region in Fig. 7).

### 4.2. Roof-aa and roof-neighbor-aa

The roof-aa (Fig. 2, pink) is a positionally conserved residue that lies above the nucleobase of the dNTP, as seen in the active site of Dpo4 [11–14], yDNAP  $\eta$  [16], hDNAP  $\iota$  [17] and hDNAP  $\kappa$  [18]. The roof-aa is next to the chimney (see Section 4.3). The "roofneighbor-aa" (Fig. 2, purple) also contacts the nucleobase of the dNTP. It is not unreasonable to imagine that the roof-aa and roofneighbor-aa could influence dNTP insertion patterns, and in fact a correlation exists. The roof-aa is isoleucine for all members of the DNAP  $V/\eta$  class listed in Fig. 7B, and the roof-neighbor-aa is always bulky (arginine or methionine). Bulky roof-aa/roof-neighbor-aa can rearrange to fill in the gap above the nucleobase in the S2-dNTP shape (Fig. 5B). The roof-aa is non-bulky (usually serine) for all IV/ $\kappa$ class members (Fig. 7A) and the roof-neighbor-aa is non-bulky (usually alanine). Non-bulky amino acids work well for S1-dNTP, but do not fill in the gap above the nucleobase in the S2-dNTP shape, making it less viable.

#### 4.3. The interconnected architecture of the chimney and roof regions

To understand the architecture of the chimney/roof region of Y-Family DNAPs, it is useful to focus on a bulky aliphatic amino acid, usually a valine (e.g., V29/UmuC(V), I30/DNAP IV, V37/scDNAP  $\eta$  and V130/hDNAP  $\kappa$ , Fig. 7), that plays a scaffolding role as revealed in X-ray structures [10–18] and in our models [42,43]. Using hDNAP  $\kappa$  as an example (Fig. 8C), this scaffolding valine (V130,

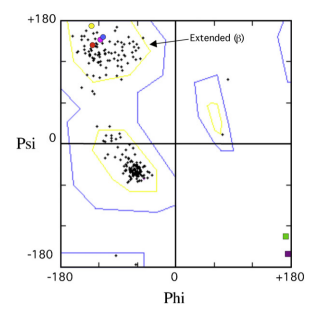


**Fig. 8.** Regions of hDNAP  $\kappa$  (panels A and B), scDNAP  $\eta$  (panels C and D) and hDNAP  $\eta$  (insert) that show why the chimney opening is large or small. Panels A–C: Y-Family DNAPs in the IV/ $\kappa$ -class have a glycine "flue-handle," such as G131 (turquoise) in hDNAP  $\kappa$ , which leads to upward curvature of the protein backbone in the chimney upper lip (red arrow, panel A) and results in the "flue" amino acids (S132/M133, blue, panel B) to point away from the chimney, giving a large chimney opening. Panel C shows V130 (white) that serves as a scaffold to organize the chimney's upper lip and left lip (yellow ribbons in panel A), along with the roof-aa (S137, purple), the steric gate (Y112, red) and a conserved tyrosine (Y174, brown), which stacks on the backbone of the left lip and orients it. V130 forms a rectangle with the G131 flue-handle, L136 (gray) and the S137 roof (pink) upon which I166 stacks (dark gray). Panels D–F: DNAP V/ $\eta$ -class DNAPs have a bulky "flue-handle," such as V54 in scDNAP  $\eta$  (turquoise, panel D), which causes downward curvature of the chimney upper lip (red arrow, panel D), and results in the "flue" amino acids (Q33/W34, blue, panel E) to plug the chimney, giving a small opening. Panel E shows scaffold C53 (white) organizing the chimney's upper lip and left lip (green ribbons in panel C), along with the roof-aa (I60, purple), the steric gate (F35, red) and a conserved tyrosine (Y131, brown), which stacks on the backbone of the left lip and orients it. C53 forms a rectangle with the V54 flue-handle (turquoise), I59 (gray) and the I60 roof (pink). scDNAP  $\eta$  has a large insert/loop (aa96–126) in the left lip, which is represented as a discontinuity. X-ray coordinates are from 20H2 for hDNAP  $\kappa$  [18] and from 1JIH for scDNAP  $\eta$  [16], where hydrogens were added using insight!l.

white) begins a loop that ends with the roof-aa, with which it forms a hydrogen bond (scaffold-C=0:HN-roof). This type of hydrogen bond is also observed in X-ray structures from Dpo4 [11–14], hDNAP  $\kappa$  [18] and scDNAP  $\eta$  [15,16]. The flue-handle (G131) is adjacent to scaffold-V130, which also contacts the bulky amino acid L136 (Fig. 8C, gray) next to roof-S137. Thus, the loop is anchored by a square of four amino acids (V130/G131/L136/S137), above which sits the R-group of I166 (dark gray, Fig. 8C) giving a rectangular pyramid of amino acids that orient the flue-handle/

flue/roof-aa region. In our model of DNAP IV, the rectangle includes I31/G32/I41/S42 with L71 sitting above it. This region in scDNAP  $\eta$  (Fig. 8F) looks similar, though only the rectangle C53/V54/I59/I60 exists, because the bulky R-group of flue-handle/V54 serves the role of the fifth amino acid (i.e., the R-group of flue-handle-V54 takes the place of I166 in hDNAP  $\kappa$ ). In our model of UmuC(V), the rectangle involves V29/L30/V37/I38.

Scaffold-V130 in hDNAP  $\kappa$  (white, Fig. 8C) also helps organize the steric gate (Y12, red), which face-stacks with Y174, a highly



**Fig. 9.** Ramachandran plot showing  $\phi/\varphi$ -angles for all valines, leucines and cysteines (crosses) from X-ray structures of Dpo4 [11] hDNAP κ [18] and scDNAP κ [15], along with the flue-handles for scDNAP η (blue circle, V54), hDNAP η (red circle, V37), UmuC(V) (yellow circle, L30) Dpo4 (pink circle, C31), which are all clustered in the region expected for extended (β) secondary structure. The glycine flue-handles for hDNAP κ (G131, green square) and DNAP IV (G32, purple square) lie outside this clustered region.

conserved tyrosine whose other aromatic face contacts the backbone of the left lip of the chimney (i.e., aa168-171 in hDNAP  $\kappa$ ), thus helping to orient it. The scDNAP  $\eta$  structure is similar (Fig. 8F) except the chimney left lip has an insert/loop (aa96-126, which is indicated by the circle). In spite of this insert/loop, the chimney left lip is similarly positioned in the vicinity of the upper lip. (Of the DNAPs shown in Fig. 7, only DNAPs  $\eta$  from yeast (both sc and sp) have an insert at this position.)

Amino acid structure in the upper lip/left lip/roof region is similar in all Y-Family DNAPs, for which X-ray structures [10–18] or modeled structures [42,43] exist.

# 4.4. dCTP insertion by DNAP IV using S1-dNTP

DNAP IV can pair dCTP with both -BP and +BP using S1-dNTP (Fig. 5C/E), importantly because DNAP IV's large chimney opening can accommodate B[a]P bulk in the developing minor groove. In contrast, DNAP V does not give a satisfactory structure when either -BP or +BP are paired with S1-dNTP (Fig. 5C/E), because DNAP V's small chimney opening forces the bulky pyrene moiety downward. In fact, UmuC(V) did not retain normal S1-dNTP-positioning in the active site during MD; rather the small chimney forced template BP-dG and its paired dCTP to move about halfway toward S2-dNTPpositioning with one consequence being that primer-O3'/dNTP-P $\alpha$ distance elongated to a non-reaction-ready distance of  $\sim$ 5 Å. These observations provide a reasonable rationale for why DNAP IV preferentially does dCTP insertion, since it can do so using the S1dNTP shape, which is presumably active, given that it is found in other DNAP families (e.g., T7 DNAP, insert Fig. 3), while DNAP V's small chimney enforces the use of S2-dNTP, which is expected to be less active (or inactive).

# 4.5. How does UmuC(V) do dATP insertion?

-BP and +BP cannot be paired using the S1-dNTP shape in UmuC(V) without significant distortions to the adduct, the dNTP,

the DNA and/or the protein, because B[a]P bulk does not fit in UmuC(V)'s small chimney opening. N31 is the main problem, as its sidechain plugs the UmuC(V) chimney leading to clashes with -BP/+BP when paired using the S1-dNTP shape. In the unadducted structure (Fig. 5B), N31 adopts its lowest energy rotational conformer with respect to the  $C\alpha-C\beta$  bond. The presence of +BP causes a rotation about the  $C\alpha-C\beta$  bond (Fig. 5D); however, no other rotations are possible to get N31 farther out of the way.

These observations lead to the question: how does UmuC(V) insert a dATP opposite +BP and -BP? One possibility involves S2-dATP pairing with adduct-dG, which allows the pyrene moiety to lie under UmuC(V)'s small chimney opening (Fig. 5J and L), since S2-dNTP lies lower down in the active site than S1-dNTP (Fig. 3). Adduct-dG can Hoogsteen pair with syn-adenine in S2-dATP. Interestingly, syn-adenine has steric clashes with atoms in the deoxyribose and the  $\alpha$ -phosphate in the S1-dATP shape (Fig. 6A), but no such steric clashes exist with the S2-dATP shape (Fig. 6B and discussed in Ref. [43]). MD runs with adduct-dG:syn-dATP pairing in UmuC(V) gave reasonable structures; e.g., the primer-O3'/dNTP-P $\alpha$  distances averaged 3.64 Å and 3.61 Å, respectively (structures not shown, though virtually identical to Fig. 5J and L, respectively).

S2-dATP Hoogsteen pairing of syn-adenine with adduct-dG is a plausible first step in the G  $\rightarrow$  T pathway. The notion that syn-dATP can be present in S2-dNTP is supported by one Dpo4 X-ray structure (1RYS-A), in which syn-dATP pairs with the 5'-T of a TT-CPD using S2-dNTP [12]. In fact, Y-Family DNAPs in the DNAP V/ $\eta$ -class may have evolved a catalytically active S2-dNTP shape to permit syn-dATP insertion opposite (e.g.) TT-CPDs, since one of the roles of the DNAP V/ $\eta$ -class is to minimize UV-light mutagenesis by just such a mechanism (discussed in Ref. [43]). Perhaps, G  $\rightarrow$  T mutations with BP-adducts are the unfortunate by-product of the occasional use of the S2-dNTP insertion pathway.

We presume that the S1-dNTP shape is optimal for catalytic efficiency given that it is observed in all families of DNAPs. But could S2-dNTP possibly be active? In previous work we analyzed thirty-four Dpo4 X-ray structures [43] and concluded that those adopting the S2-dNTP shape had accompanying protein components that should allow phosphoester bond making and breaking, notably with protein elements needed: (1) to ensure proper geometry between the nucleophilic primer-O3' and the electrophilic dNTP-P $\alpha$ ; (2) to enhance electrophilicity of P $\alpha$  of the dNTP during nucleophilic attack; (3) to dampen negative charge on the pyrophosphate leaving group; and (4) to orient the  $P\alpha-P\beta-P\gamma$ phosphate tail. In fact, S2-dNTP seemed to have all of the necessary requirements to be catalytically active. For example, in fact four out of five S2-dNTP structures had primer-O3'/dNTP-P $\alpha$  distances of less than 4.0 Å (average: 4.0 Å, range: 3.66-4.69 Å), while even the best S1-dNTP structure had a longer distance of 4.41 Å with an average of 5.9 Å (range: 4.41–7.95 Å). We note, however, that there is no evidence that S2-dNTP is active, and it is merely a hypothesis that it might be.

Other G:A mispairings are also possible. In principle, *anti*-dATP can pair with *syn*-guanine in adduct-dG, which requires the pyrene moiety to be in the major groove. *Anti*-dATP can also pair with *anti*-guanine in adduct-dG in an elongated mispair. Thus, there are scenarios other than the one involving our *syn*-dATP: *anti*-adduct-dG hypothesis.

Our modeled structures show that, in principle, either dCTP or syn-dATP can pair with adduct-dG using S2-dNTP. In a 5′-GGT sequence [30], purified DNAP V inserts both dCTP and dATP ([dATP]/[dCTP] = 0.75 and 0.18 for -BP and +BP, respectively). In a 5′-CGA sequence, however, purified DNAP V inserts dATP > 100-fold more efficiently than dCTP opposite -BP and +BP [30]. In cells we know that DNAP V does not efficiently insert dCTP in our 5′-TGC sequence, because - if it did - the non-mutagenic pathway would

not have declined to  $\sim$ 1% and  $\sim$ 3% efficiency for -BP and +BP, respectively, in *E. coli* defective in DNAP IV [34]. It will be of interest to investigate whether an S2-dNTP structure might explain these differences in dATP vs. dCTP insertion efficiency.

We note, however, that one published finding is hard to rationalize based on the mechanism proposed in this section. Of mismatched insertions, dATP is most often inserted opposite a B[a]P-dA adduct by DNAP V [30]; there is no obvious way to pair dATP with an adduct-dA, not even via Hoogsteen or wobble pairings. It is impossible to know whether this observation implies that a more nuanced variation of the mechanism proposed in this section is required, or whether it implies that the mechanism is wrong.

#### 4.6. Dpo4 compared to DNAPs IV and V

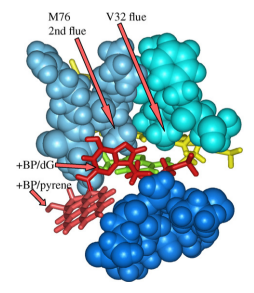
Based on biochemical and X-ray findings [14], Dpo4 insertion opposite +BP was proposed to follow a "dislocation" or "templated" pathway. Dislocation/templated insertion ([46,47] and references therein) involves DNAP stalling at an adduct, slippage to the next 5′-base along the template, and its use to direct incorporation (e.g., dATP insertion opposite the 5′-T in a 5′-TG sequence context), whereupon the newly incorporated dA slips back to form an adduct-G:A mispair, from which extension yields the mispair that ultimately gives a G  $\rightarrow$  T mutation. Dpo4 preferentially inserted dCTP, dTTP, dATP and dGTP opposite +BP in 5′-GG, 5′-AG, 5′-TG and 5′-CG sequences, respectively [14], which is consistent with a dislocation/templated mechanism.

It was suggested that the dislocation/templated mechanism might operate widely with B[a]P-N<sup>2</sup>-dG adducts [14], which-if true-would mean that the mechanism proposed herein is off-track. Though the dislocation/templated mechanism is attractive for Dpo4, considerable evidence suggests it cannot dominate in E. coli, as we have discussed previously [29,31,34]. (All findings below not specifically attributed were reviewed in Ref. [29].) (1) In five cases using various B[a]P-N<sup>2</sup>-dG isomers in 5'-TG sequences,  $G \rightarrow T$ mutations dominate, which is consistent with a dislocation/ templated mechanism. However,  $G \rightarrow T$  mutations dominate in eleven cases using various B[a]P-N<sup>2</sup>-dG isomers in various 5'-CG sequences; these findings are not consistent with a dislocation/ templated mechanism, which predicts a  $G \rightarrow C$  mutation.  $G \rightarrow A$ mutations dominated with +BP and -BP in one 5'-AG sequence, which is consistent with a dislocation/templated mechanism. However,  $G \rightarrow A$  mutations dominate in one 5'-CG sequence using three different B[a]P-N<sup>2</sup>-dG isomers, which is inconsistent with a dislocation/templated mechanism. Thus, where B[a]P-N<sup>2</sup>-dG has been studied site-specifically about twice as many examples are inconsistent than are consistent with the dislocation/templated mechanism. (Of course, even if a finding is "consistent" it does not mean it is the true mechanism.) (2) In mutagenesis studies with (+)-anti-B[a]PDE [25], where a broad range of sequence contexts were sampled,  $G \rightarrow T$  mutations dominated in 5'-TG sequence contexts, consistent with "dislocation/templated" mutagenesis. However, if this were a general mechanism, it should operate in other sequence contexts, which is not the case:  $G \rightarrow C$  and  $G \rightarrow A$ mutations did not dominate in 5'-CG and 5'-AG sequences, respectively, while 5'-GG sequences were not principally mutagenically silent [25]. In fact,  $G \rightarrow T$  mutations dominated in non-5'-TG sequences with (+)-anti-B[a]PDE, implying that it cannot be the only dATP insertion mechanism, even if the dislocation/templated pathway does occur. (3) Purified DNAP V preferentially inserted dATP opposite +BP and -BP in a 5'-CG sequence [30], which is not consistent with a dislocation/templated mechanism. DNAP V inserted dCTP opposite +BP and -BP in a 5'-GG sequence [30], which is consistent with a dislocation/templated mechanism, though dATP insertion was also significant in both cases, which is not consistent. Importantly, these findings demonstrate that DNAP V, which inserts dATP opposite +BP in the  $G \rightarrow T$  pathway in E. coli [31], has a viable means of doing so that does not depend on a dislocation/templated mechanism. (4) Purified DNAP IV correctly inserted dCTP opposite +BP and -BP in 5′-CG and 5′-GG sequences [30], where only the latter is consistent with a dislocation/templated mechanism.

Thus, while a dislocation/templated mechanism is consistent with some findings in  $E.\ coli$ , the fact that it fails to rationalize the majority of the findings implies that it is not a unifying hypothesis. Furthermore, the dislocation/templated mechanism is not compelling given that its invocation forces one to propose that the dominant  $G \to T$  mutational pathway follows one mechanism (dislocation/templated) in one sequence context (i.e., 5'-TG), but some other mechanism in other sequence contexts, when there is no obvious reason why the mechanism would change. The mechanism proposed herein (i.e., adduct-dG pairing with syn-dATP using S2-dNTP) would give  $G \to T$  mutations independent of sequence context.

Returning to Dpo4, the biochemical evidence was clear: Dpo4 appears to use a dislocation/templated mechanism with +BP [14], as summarized in the first paragraph of this subsection. Why might Dpo4 be different than DNAPs IV and V of *E. coli*? Both the roof-aa and roof-neighbor-aa for Dpo4 are non-bulky (A44/A57), while its bulky flue-handle (C31) causes downward curvature of the chimney upper lip, leading to a closed flue (V32) and a small chimney opening (based on structures from Refs. [11–14], and other Dpo4 structures). Thus, Dpo4 is a hybrid with a roof similar to the IV/ $\kappa$ -class and a chimney similar to the V/ $\eta$ -class.

In fact, M76, which is the second amino acid in Dpo4's left lip, also rests in the chimney (Fig. 10), making its chimney even more plugged than those of DNAP IV and UmuC(V), which have non-bulky G74 and S72, respectively, in the equivalent position. In fact one X-ray structure for +BP/Dpo4 reveals that the pyrene (light red, Fig. 10) is so excluded that it also drags the adduct-dG moiety (red,



**Fig. 10.** Chimney region of Dpo4 from X-ray coordinates 2IA1 [14] showing that the dG moiety (red) in addition to the pyrene (light red) of +BP are excluded from the active site, probably because Dpo4's chimney is particularly plugged, since it has both a regular flue (V32) in the upper lip (turquoise) and a second flue (M76), the latter being the second amino acid in the left lip (blue). The lower lip includes the positionally conserved four amino acids (dark blue, aa-246–249) along with aa271, the latter because the pyrene moiety stretches so far down. This is an "extension structure" with a 5'-template-dT paired with dATP (yellow) in the active site, while the +BP moiety is in the [n+1] position. Hydrogens were added using insight!l.

Fig. 10) into the chimney resulting in a gap in the template strand. (This is an "extension structure" with dATP paired with a template-5'-T in the active site, while the +BP moiety is formally in the [n+1] position, though displaced into the chimney. Extension structures, which we are also modeling (data not shown), are more similar to S2-dNTP structures, since both have the adduct-dG in the [n+1] position.) Furthermore, the V32 flue of Dpo4 is also inserted deeper into the chimney than (e.g.) the N32 flue of UmuC(V). The excessively plugged chimney of Dpo4 does not permit S2-dNTP insertion structures similar to UmuC(V) (Fig. 5J and L), which may explain why the dG moiety of +BP in the chimney is not available for pairing and why Dpo4 follows a dislocation/templated mechanism. We hasten to add that Dpo4 actually has a large cleft between the fingers and little finger domains, but it begins below the C31/M76 plugs.

This analysis suggests a reason for caution when applying conclusions from Dpo4 to other Y-Family DNAPs, especially those purely in the IV/ $\kappa$ -class or the V/ $\eta$ -class. Perhaps Dpo4 evolved its hybrid roof/chimney structure to bypass a unique set of lesions encountered by a thermophilic bacteria. Alternatively, perhaps the structure of Dpo4 at physiologically relevant elevated temperatures is different than at the temperature at which it was crystallized (r.t.) and assayed (37°C), and this affects its structure and behavior.

#### 4.7. Summary

The IV/κ-class of Y-Family DNAPs have a key glycine "fluehandle" that keeps adjacent bulky amino acids ("flue") out of an opening on the minor groove side of the active site ("chimney"). which is thus large, can accommodate the pyrene, ands allows B[a]P-dG:dCTP pairing using the canonical "chair-like" S1-dNTP shape. The  $V/\eta$ -class of DNAPs have a bulky flue-handle (usually valine) that force the flue amino acids to plug the chimney, which gives a small opening that forces +BP/-BP downward in the active site, thus preventing proper geometry for primer-O3' attack on P $\alpha$ when dCTP is in the S1-dNTP shape. This may explain why IV/κclass DNAPs do cellular dCTP insertion opposite N<sup>2</sup>-dG adducts, such as +BP and -BP. The small chimney opening of the  $V/\eta$ -class may force +BP/-BP so far down in the active site that adduct-dG can pair with the non-canonical "goat-tail-like" S2-dNTP shape, which lies lower in the active site than S1-dNTP. The S2-dNTP shape permits Hoogsteen syn-dATP pairing with B[a]P-dG and might be the first step in the  $G \rightarrow T$  mutational pathway.

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