A highly conserved Tyrosine residue of family B DNA polymerases contributes to dictate translesion synthesis past 8-oxo-7,8-dihydro-2'-deoxyguanosine

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ABSTRACT

The harmfulness of 8-oxo-7,8-dihydro-2'-deoxyguanosine (8oxodG) damage resides on its dual coding potential, as it can pair with the correct dCMP (dC) or the incorrect dAMP (dA). Here, we investigate the translesional synthesis ability of family B φ29 DNA polymerase on 8oxodG-containing templates. We show that this polymerase preferentially inserts dC opposite 8oxodG, its 3’–5’ exonuclease activity acting indistinctly on both dA or dC primer terminus. In addition, φ29 DNA polymerase shows a favoured extension of the 8oxodG/dA pair, but with an efficiency much lower than that of the canonical dG/dC pair. Additionally, we have analysed the role of the invariant tyrosine from motif B of family B DNA polymerases in translesional synthesis past 8oxodG, replacing the corresponding φ29 DNA polymerase Tyr390 by Phe or Ser. The lack of the aromatic portion in mutant Y390S led to a lost of discrimination against dA insertion opposite 8oxodG. On the contrary, the absence of the hydroxyl group in the Y390F mutant precluded the favoured extension of 8oxodG:dA base pair with respect to 8oxodG:dC. Based on the results obtained, we propose that this Tyr residue contributes to dictate nucleotide insertion and extension preferences during translesion synthesis past 8oxodG by family B replicases.

INTRODUCTION

The high fidelity displayed by replicases from families A, B and C outcomes from 10^4 to 10^6-fold polymerase preference for inserting correct rather than incorrect nucleotides (1). Such selectivity has traditionally been considered to rely on the base-pair shape and size, as the result of the geometric restraints imposed by the polymerase active site to tolerate equivalent Watson–Crick base pairs, ruling out those differing from this geometry (1–3). Additionally, recent studies performed with non-natural nucleotides also suggest a role for the π–π stacking interactions between the aromatic rings of the incoming dNTP and amino acid residues for an efficient polymerization at least in family B DNA polymerases (4). Polymerases contact DNA at positions in which both, topology and chemistry are identical among the four canonical base pairs. Thus, interactions occur mainly with the base, sugar and triphosphate moieties of the incoming nucleotide, with the nucleotide placed immediately 5’ with respect to the templating nucleotide and, most importantly, with the minor groove of the nascent base pair (1,3). In this sense, structural studies of DNA polymerase complexes have revealed the presence of conserved residues at their catalytic sites that make contacts with the purine N3 and pyrimidine O2 atoms that act as hydrogen-bond acceptors and are positioned at similar locations in the minor groove (1,5–8). Incorrect insertion of a nucleotide will move such H-acceptors out of position, breaking interactions with the polymerase and diminishing the DNA-binding stability at the polymerization site. This will result in an increased chance of the primer DNA to be switched to the 3’-5’ exonuclease site of the replicase to have the incorrect nucleotide removed (8–10).

Lesions in the genomes arise by their continuous exposure to the action of toxic environmental agents such as ultraviolet and ionizing radiation, genotoxic chemicals, by-products of the normal metabolism like the reactive oxygen species (ROS), in addition to spontaneous breaks of chemical bonds (11). Most organisms code for a number of enzymes to repair such damages before replication fork meeting them since replicative DNA polymerases stall when they come across one of these lesions as they cannot form a proper and catalytically competent ternary complex with the damaged nucleotide at their catalytic sites. One exception is the specially deleterious lesion

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8-oxo-7,8-dihydro-2'-deoxyguanosine (8oxodG), produced by ROS inside the cell (12) as, if it escapes the repair machinery, it can be used by the replicase as template or as incoming nucleotide. The harmfulness of this lesion resides in its dual coding potential as it can pair with both cytosine and adenine during DNA synthesis, in the latter case leading to G to T transversions (13,14). Structural studies of both 8oxodG:dC and 8oxodG:dA base pairs showed that 8oxodG adopts the classical anti conformation opposite dC, pairing in a Watson–Crick fashion, whereas its glycosidic bond adopts the syn orientation to form a Hoogsteen base pair with dA (15–19).

Structural modelling of catalytically competent complexes of RB69 and T7 DNA polymerases have suggested that the preferential insertion of dC opposite such a lesion is accomplished by a handcapped dA incorporation since the O8 atom of the 8oxodG(3yn)dA mispair sterically clashes with specific residues at the corresponding active sites (19,20). However, notwithstanding their high insertion fidelity and preferential dC insertion during the bypass of 8oxodG-containing templates, replicases can misincorporate dA with a moderately high efficiency (19–24) due to the fact that such a base pair establishes appropriate hydrogen-bond interactions with the minor groove sensing residues at the catalytic site (19).

Bacteriophage ϕ29 DNA polymerase is a protein-primed DNA-dependent replicase belonging to the eukaryotic-type family of DNA polymerases (family B). Like many other replicases, it possesses, within a single polypeptide chain, both 5'-3' polymerization and 3'-5' exonuclease activities. It displays a high intrinsic nucleotide insertion discrimination [104 to 106 (25)], which is further improved 100-fold through proofreading by the exonuclease domain (26). ϕ29 DNA polymerase displays two unique characteristics compared with most replicases. First, a DNA polymerase molecule replicates the entire genome processively without the assistance of processivity factors (27), in contrast to most replicases that require accessory proteins to clamp the enzyme to the DNA (28–31). Second, ϕ29 DNA polymerase couples processive DNA polymerization to strand displacement. This ability allows the enzyme to replicate the ϕ29 double-strand genome without the need for a helicase (27).

In this article, we study the ability of ϕ29 DNA polymerase to perform translesional synthesis past 8oxodG by assaying the nucleotide insertion opposite the lesion and further extension steps, as well as its capacity to proofread the formed pairs, this issue being of importance, as this enzyme is currently used for isothermal rolling circle amplification and whole genome amplification (32,33). Structural models mentioned above, together with multiple sequence alignments of DNA polymerases (34), as well as the availability of the crystallographic structure of ϕ29 DNA polymerase (35), have led us to analyse the role in translesional synthesis past 8oxodG of the invariant Tyr residue of the highly conserved B motif of family B DNA polymerases by means of substitutions at the corresponding ϕ29 DNA polymerase residue Tyr390. The results obtained allow us to propose a principal and dual role for this residue as one of the key determinants that dictate nucleotide insertion and extension preferences during translesion synthesis past 8oxodG by family B replicases.

MATERIALS AND METHODS

Nucleotides and DNAs

Unlabelled nucleotides were purchased from Amersham Pharmacia Biochemicals. [γ-32P]ATP (3000 Ci/mmol) was obtained from Amersham Pharmacia. Oligonucleotides Pber (5’CTGCAGCTGATGC), Pber-2 (5’CTGCA GCTGATGC), PberA (5’CTGCAGCTGATGCAGCA), PberC (5’CTGCA GCTGATGCAGC) and sp1 (5’GATC GACGTGAGTAC) were 5’-labelled with [γ-32P]ATP and phage T4 polynucleotide kinase and purified electrophoretically on 8 M urea–20% polyacrylamide gels. Labelled Pber, Pber-2, PberA and PberC were hybridized to the 34-mer 8oxodG containing oligonucleotide 5’GATACCCGGGATCCGTAC (8oxodG) GGCATCA GCTGCA, and labelled sp1 to sp1c+5 (5’TCTATCG TACCTCAGTGTAC) and to sp1cA+5 (5’TCTATAG TACCTCAGTGTAC). Hybridizations were performed in the presence of 0.2 M NaCl and 50 mM Tris-HCl (pH 7.5), resulting in primer/template structures. Oligonucleotides were obtained from Invitrogen.

Proteins

Phage T4 polynucleotide kinase was obtained from New England Biolabs. ϕ29 DNA polymerase variants at Tyr390 residue Y390F and Y390S were constructed by J. Saturno in an exonuclease deficient background (D12A/D66A) (hereafter PolY390FExo and PolY390SExo, respectively (unpublished data) and further purified from Escherichia coli BL21(DE3) cells harbouring the corresponding recombinant plasmid (36).

Translesion synthesis past 8oxodG by ϕ29 DNA polymerase

The hybrid molecules Pber-2/T4 and Pber-2/8oxodG, containing the unmodified dG and an 8oxodG lesion at the +3 position of the template, respectively, can be used both as substrate for the 3’-5’ exonuclease activity and for DNA-dependent DNA polymerization. The incubation mixture contained, in a final volume of 12.5 μl, 50 mM Tris-HCl (pH 7.5), 1 mM dithiothreitol, 4% glycerol, 0.1 mg/ml bovine serum albumin (BSA), 10 mM MgCl2, 1.2 nM of either 5’-labelled Pber-2/T4 or Pber-2/8oxodG, 24 nM of either wild-type or mutant D12A/D66A (Exo-) DNA polymerase, and the indicated concentration of the four dNTPs. After incubation for 5 min at 25°C, the reaction was stopped by adding EDTA up to 10 mM. Samples were analysed by 8 M urea–20% PAGE and autoradiography. Polymerization or 3’-5’ exonuclease activity are detected as an increase or decrease, respectively, in the size (13-mer) of the 5’-labelled Pber-2 primer.

3’-5’ exonuclease activity of wild-type ϕ29 DNA polymerase

The incubation mixture contained, in 12.5 μl, 50 mM Tris-HCl (pH 7.5), 10 mM MgCl2, 1 mM dithiothreitol, 4% glycerol, 0.1 mg/ml BSA, 6 nM of wild-type ϕ29 DNA

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polymerase and 1.2 nM of either 5′-labelled Pber/C/T4, PberA/8oxodG or PberC/8oxodG double-stranded DNA (dsDNA) substrate. Samples were incubated at 25°C for the indicated times and quenched by adding EDTA up to 10 mM. Reactions were analysed by 8 M urea–20% PAGE and autoradiography.

Insertion of dC and dA opposite 8oxoG

The incubation mixture contained, in a final volume of 12.5 µl, 50 mM Tris-HCl (pH 7.5), 1 mM dithiothreitol, 4% glycerol, 0.1 mg/ml BSA and 10 mM MgCl₂. As substrate, 1.2 nM of the 5′-labelled hybrid molecule Pber/8oxoG dsDNA was used. The amount of DNA polymerase added (12, 30 and 18 nM of Pol wtExo⁻ and Pol Y390FExo⁻ mutants, respectively) was adjusted to obtain linear conditions. Samples were incubated for 15 s at 25°C, in the presence of the indicated concentrations of either dCTP or dATP, and quenched by adding 3 µl of gel loading buffer. Reactions were analysed by electrophoresis in 8 M urea–20% PAGE and quantified using a Molecular Dynamics PhosphorImager. Formation of dCTP:dA and dATP:dC base pairs, respectively. Assay was carried out under the same conditions without 8oxoGTP, using 8oxoGTP labelled hybrid molecules sp1/sp1cC+5 or sp1/sp1cA+5 was used. The incubation mixture contained, in a final volume of 12.5 µl, 50 mM Tris-HCl (pH 7.5), 1 mM dithiothreitol, 4% glycerol, 0.1 mg/ml BSA, 10 mM MgCl₂ and 12 nM of Pol Y390FExo⁻. As substrate, 1.2 nM of the 5′-labelled hybrid molecule sp1/sp1cC+5 or sp1/sp1cA+5 was used. Samples were incubated at 25°C for 15 s in the presence of the indicated concentrations of 8oxoGTP and quenched by adding 3 µl of gel loading buffer. Reactions were analysed and quantified as described above.

Extension of dC:8oxoG and dA:8oxoG base pairs

The incubation mixture contained, in a final volume of 12.5 µl, 50 mM Tris-HCl (pH 7.5), 1 mM dithiothreitol, 4% glycerol, 0.1 mg/ml BSA, 10 mM MgCl₂ and 12 nM of Pol Y390FExo⁻. As substrate, 1.2 nM of the 5′-labelled hybrid molecule sp1/sp1cC+5 or sp1/sp1cA+5 was used. Samples were incubated at 25°C, in the presence of either 100 nM dGTP or 64 µM 8oxoGTP to obtain the dC:dG, dC:8oxoG and dA:8oxoG base pairs poised for extension by the indicated dATP concentrations. Samples were incubated at 25°C for 15 s and quenched by adding 3 µl of gel loading buffer. Reactions were analysed by electrophoresis in 8 M urea–20% PAGE.

RESULTS AND DISCUSSION

Translesion synthesis past 8oxoG by φ29 DNA polymerase

φ29 DNA polymerase is the most processive replicase known, being able to incorporate, without the assistance of processivity factors, more than 70 000 nt during a single encounter with an undamaged DNA template (27). Considering the presence of ROS derived from normal cellular metabolism that react with and modify DNA, is predictable that φ29 DNA polymerase has to deal with such DNA lesions during replication. Here, we have analysed the efficiency of translesion synthesis past one of the most abundant DNA lesions caused by ROS, the 8oxoG. To test φ29 DNA polymerase ability to carry out nucleotide insertion and further extension opposite such a lesion, primer extension reactions were conducted in the presence of increasing concentrations of the four dNTPs DNA polymerase molecules. After the delay times indicated, 100 µM dGTP was added. Reactions were quenched by adding 3 µl of gel loading buffer 15 s after addition of dGTP. For 0 min delay time point, dGTP was included in the solution with the trapping DNA. Control reaction of the effectiveness of the trap was performed by incubating the DNA polymerase with the labelled and trapping DNA simultaneously. Reactions were analysed by electrophoresis in 8 M urea–20% PAGE and quantified using a Molecular Dynamics PhosphorImager. The fraction of primer molecules extended (y) by the saturating concentration of dGTP was plotted against the delay time (t). These data were fit to the single exponential relationship

\[ y = Ae^{-kt} + B \]

where A and B are the lower and upper limits, respectively, and k is the rate constant for dissociation. Assays were performed at least three times to guarantee reproducibility.

8oxoG incorporation opposite dC and dA

The incubation mixture contained, in a final volume of 12.5 µl, 50 mM Tris-HCl (pH 7.5), 1 mM dithiothreitol, 4% glycerol, 0.1 mg/ml BSA, 10 mM MgCl₂ and 12 nM of Pol Y390FExo⁻. As substrate, 1.2 nM of the 5′-labelled hybrid molecule sp1/sp1cC+5 or sp1/sp1cA+5 was used. Samples were incubated at 25°C for 15 s in the presence of the indicated concentrations of 8oxoGTP and quenched by adding 3 µl of gel loading buffer. Reactions were analysed and quantified as described above.

Measurement of DNA polymerase dissociation rates

Reactions were carried out essentially as described (37,38). The incubation mixture contained, in a final volume of 12.5 µl, 50 mM Tris-HCl (pH 7.5), 1 mM dithiothreitol, 4% glycerol, 0.1 mg/ml BSA, 10 mM MgCl₂, 1.2 nM of the 5′-labelled hybrid molecule PberA/8oxoG or PberC/8oxoG and 25 nM of either Pol Y390FExo⁻ or 75 nM of Pol Y390SExo⁻ mutant. After 5 min at 25°C to reach the binding equilibrium, samples were mixed with a 1000-fold excess of unlabelled substrate to trap dissociated DNA polymerase molecules. After the delay times indicated, 100 µM dGTP was added. Reactions were quenched by adding 3 µl of gel loading buffer 15 s after addition of dGTP. For 0 min delay time point, dGTP was included in the solution with the trapping DNA. Control reaction of the effectiveness of the trap was performed by incubating the DNA polymerase with the labelled and trapping DNA simultaneously. Reactions were analysed by electrophoresis in 8 M urea–20% PAGE and quantified using a Molecular Dynamics PhosphorImager. The fraction of primer molecules extended (y) by the saturating concentration of dGTP was plotted against the delay time (t). These data were fit to the single exponential relationship

\[ y = Ae^{-kt} + B \]

where A and B are the lower and upper limits, respectively, and k is the rate constant for dissociation. Assays were performed at least three times to guarantee reproducibility.
(see the Materials and Methods section). The 5'-labelled primer molecule was annealed to templates containing at the +3 position with respect to the 3'-end of primer terminus either the natural dG (X=G) or 8oxodG (X=8oxodG). As shown in Figure 1, in the absence of nucleotides, the wild-type enzyme digests the primer strand by means of its 3'-5' exonuclease activity and requires 50–100 nM dNTPs to get a net elongation of the primer strand with the unmodified template, yielding full-length (34-mer) products. In contrast, at those high dNTPs concentrations (100 nM), the polymerase stalls at the +1 and +2 positions, that is, prior to the 8oxodG lesion, showing nucleotide insertion opposite damage as a limiting step. As dNTP concentration increases, a band at the +3 position appears just opposite 8oxodG, indicating that extension of the primer terminus paired to 8oxodG is also impaired, needing an 800 nM dNTP concentration to come across the damage. In parallel, we tested the behaviour of the exonuclease deficient φ29 DNA polymerase mutant D12A/D66A on these molecules to study the effect of the 3'-5' exonuclease activity during translesion synthesis past 8oxodG (Figure 1). The absence of competition by the 3'-5' exonuclease activity allows mutant polymerase to insert and further elongate nucleotides at dNTPs concentrations lower than those of the wild-type enzyme with the unmodified template. Interestingly, the mutant polymerase was able to insert nucleotides opposite 8oxodG at 100 nM dNTPs, as manifested by the presence of a band at the +3 position. To get full-length products with the damaged DNA, both mutant and wild-type DNA polymerases required a dNTP concentration 16-fold higher than that needed with the undamaged template. From this, it could be inferred that the 3'-5' exonuclease activity of the wild-type enzyme exerts a similar pressure with both substrates, suggesting that during translesional synthesis DNA polymerase could proofread the primer terminus paired with 8oxodG.

To carry out a comparative analysis of the capacity of the wild-type DNA polymerase to excise 3' terminal dA or dC paired to 8oxodG versus dC opposite the unmodified dG, we performed a 3'-5' exonuclease assay using as substrates the hybrid molecules depicted on top of Figure 2 (see also Materials and Methods section). As shown in Figure 2, the wild-type DNA polymerase was able to degrade the three substrates with the same efficiency, irrespective of their terminus base pair. The inherent potential deleteriousness of the 8oxodG(dyn):dA mispair is a consequence of not being identified as aberrant by the polymerase since, in such orientation, the O8 group mimics the O2 group of a thymidine, establishing a proper hydrogen-bond contact with the minor groove sensing residues at the catalytic site (19). This fact allows stable mispair formation at the polymerization site, avoiding its exonucleolytic correction. In addition, DNA polymerases contain residues that make interactions through the minor groove of the DNA at post-replication positions that contribute to binding affinity and to the ability to sense base-pair geometry (3). The misinsertion and further extension of nucleotides provoke changes in the geometry of the DNA.
with the subsequent lost of several of these contacts, causing DNA polymerase to stall and proofread the misinserted nucleotides (9). Protrusion of the Hoogsteen base pair 8oxodG(syn):dA into the major groove does not alter the overall structure of the DNA, contributing to avoid its exonucleolytic proofread.

### Error-free synthesis opposite 8oxodG depends on the invariant Tyrosine residue of motif B

As most DNA polymerases assayed in their capacity to insert nucleotides opposite 8oxodG [see (39) and references therein], 829 DNA polymerase is able to catalyse mainly the insertion of both dC and dA (data not shown). Crystallization of bacteriophage T7 (family A) (19) and RB69 (family B) (20) DNA polymerases complexed with a nascent 8oxodG(anti):dCTP base pair have given light about the structural rationale for the efficient bypass of 8oxodG, in contrast to other lesions as abasic sites and thymine dimers that completely stall the replication machinery until the lesion is either repaired or bypassed. In both structures, the unfavourable steric interaction between the O8 group and O4 oxygen of the 8oxodG in its non-mutagenic (anti) conformation is alleviated by a kinking of the phosphodiester backbone of the template strand at the catalytic site. In T7 DNA polymerase, there is an additional contact between the ε amino group of residue Lys536 and the O8 group of 8oxodG steadying the anti conformation for base pairing with dCTP (19,24). On the other hand, modelling of a nascent 8oxodG(syn):dATP base pair in the active site of both DNA polymerases allowed Brieba et al. (19) and Freisinger et al. (20) to predict steric and electrostatic clashes between the O5 group and the side chain of residues located at the fingers subdomain, as responsible for the discrimination against the 8oxodG in its syn (mutagenic) conformation. In spite of the presence of polymerase-dependent specific residues, in both DNA polymerases the Tyr residue of the highly conserved motif B (Tyr567 and Tyr530 in RB69 and T7 DNA polymerases, respectively) (40) is proposed to play a role in discrimination during nucleotide insertion opposite an 8oxodG site.

To study the role of such Tyr residue in 8oxodG translesion synthesis, we have analysed the insertion and elongation capacities opposite 8oxodG-containing templates of 829 DNA polymerase variants at the homologous Tyr390 residue, in an exonucleic deficient background (D12A/D66A). PolY390FExo− and PolY390SExo− mutants, allowed us to evaluate the importance of the hydroxyl and aromatic groups, respectively, in both translesional phases by comparing their activities with the wild-type enzyme (PolwtExo−).

To analyse the ability of 829 DNA polymerase mutants to carry out nucleotide insertion opposite 8oxodG, primer extension reactions were conducted in the presence of increasing concentrations of the corresponding dNTP (see Materials and Methods section). A 5'-labelled primer molecule (15-mer) was annealed to a template (34-mer) containing 8oxodG at the nascent position (see Materials and Methods section). As it can be observed in Figure 3 and Table 1, mutant PolwtExo− displayed a clear preference to incorporate dC with respect to dA, mainly due to the lower Km for dC, showing a discrimination factor (fins in Table 1) of about 5, a moderately high insertion fidelity in comparison with other polymerases belonging to families A, B, X, Y and reverse transcriptase (RT) (13,20,21,23,24,41–50), in which the dC:dA insertion ratio ranges from 0.075 for HIV-1 RT (21) to 91 for Dpo4 DNA polymerase (51) [compiled in (39)]. The incorporation of dA cannot be attributed to a decreased insertion fidelity of PolwtExo−, as it discriminates 3 × 105-fold against dA insertion opposite the undamaged dG (data not shown). Under those conditions, and consistent with their reduced efficiency in incorporating nucleotides opposite undamaged templates due to a reduced dNTP-binding capacity (52,53), PolY390FExo− and PolY390SExo− mutants were also less active than PolwtExo− in inserting both dA and dC at 8oxodG (Figure 3 and Table 1). Thus, correct dC insertion by PolY390FExo− and PolY390SExo− mutants is reduced to two and one orders of magnitude relative to the wild-type polymerase, respectively. These data are in good agreement with the comparative efficiencies displayed by these two mutants during incorporation of dNTPs on undamaged substrates (53), precluding any additional incorporation defect due to the presence of 8oxodG at the templating position. However, their discrimination factors clearly changed. Whereas PolY390Exo− mutant discriminates against dA insertion in a similar fashion as PolwtExo− (fins~4), PolY390SExo− mutant displays nearly identical insertion efficiency for both nucleotides (fins~1). Such behaviour is not a direct consequence of the diminished insertion fidelity showed by this mutant, as it has a discrimination factor against erroneous nucleotide insertion opposite a non-damaged nucleotide of 3 × 105 (52,53). This result clearly reflects
the importance of the aromatic moiety of φ29 DNA polymerase residue Tyr390 in discriminating against the syn orientation (error-prone) of the purine base of the 8oxodG. Freisinger et al. modelled a nascent 8oxodG:dATP base pair at the catalytic active site of RB69 DNA polymerase (20). Such a model led them to propose that the O₈ group of 8oxodG(syn) would clash with the molecular surface formed by residues Tyr567 and Gly568, allowing the canonical and non-mutagenic anti conformation to be preferentially placed (20). The DNA and incoming nucleotide from the structure of the RB69 ternary complex (8) can be homology modelled onto the structure of the apo φ29 DNA polymerase by aligning the palm subdomains of both φ29 and RB69 DNA.

Table 1. Insertion of dCTP and dATP opposite 8oxodG

<table>
<thead>
<tr>
<th>Mutant polymerase</th>
<th>dA insertion</th>
<th>dC insertion</th>
<th>dA insertion</th>
<th>dC insertion</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Kₘ (µM)</td>
<td>K_{cat} (min⁻¹)</td>
<td>(K_{cat}/Kₘ)(µM⁻¹min⁻¹)</td>
<td>Kₘ (µM)</td>
</tr>
<tr>
<td>PolwtExo⁻</td>
<td>28.05 ± 6.2</td>
<td>0.124</td>
<td>4.42 × 10⁻³</td>
<td>5.8 ± 1.2</td>
</tr>
<tr>
<td>PolY390FExo⁻</td>
<td>457.6 ± 122.1</td>
<td>0.027</td>
<td>5.9 × 10⁻³</td>
<td>168.2 ± 33.3</td>
</tr>
<tr>
<td>PolY390SExo⁻</td>
<td>116.7 ± 16.4</td>
<td>0.098</td>
<td>8.4 × 10⁻⁴</td>
<td>104.1 ± 18</td>
</tr>
</tbody>
</table>

(K_{cat}/Kₘ) stands for the catalytic efficiency displayed by mutant polymerases for the insertion of the indicated nucleotide. 
f_{ins} stands for insertion discrimination factor obtained by dividing catalytic efficiency for the insertion of dC by that for the insertion of dA.
polymerases, the position of the modelled DNA and incoming nucleotide being consistent with the placement of the φ29 DNA polymerase catalytic carboxylates, steric gate residues that distinguish ribo- from deoxyribonucleotides, and residues involved in binding DNA (35). This fact allowed us also to model a nascent 8oxodG:dATP in the corresponding active site of φ29 DNA polymerase (Figure 4) in which φ29 DNA polymerase residues Tyr390 and Gly391 are placed at the same location and orientation that the homologous RB69 DNA polymerase residues Tyr567 and Gly568, mentioned above. Based on that, the absence of discrimination showed by PolY390Exo− mutant could reflect that the lack of the aromatic group avoids the steric contacts with the O8 of the 8oxodG(syn) either by removing a direct interaction with Tyr390 or by tolerating a subtle shift of Gly391 backbone to accommodate the O8 atom, or both. Interestingly, recent studies have involved the homologous Tyr766 residue of E. coli DNA polymerase I in preventing incorrect nucleotide insertion across from the dG:N-2-aminofluorene adduct, as mutant Y766S was significantly less selective for correct nucleotide incorporation than the wild-type enzyme (54).

Tyrosine residue of motif B dictates the extension preferences of nucleotides base-paired to 8oxodG lesion

To study the next step after insertion opposite 8oxodG, 16/34-mer hybrid molecules, with the primer 3'-end containing either dC or dA paired with 8oxodG at the template (see Materials and Methods section), were used as substrate to analyse single nucleotide incorporation of dG, the next correct nucleotide following the 8oxodG:dC/8oxodG:dA base-pair formation. As seen in Table 2 (see also Supplementary Figure S1), PolwtExo− mutant displayed a 14-fold favoured extension of the 8oxodG:dA base pair (see in Table 2 discrimination factor, fex). Interestingly, PolY390Exo− mutant lost discrimination ability to elongate the base pairs under study, as it showed a 5-fold preference to extend the 8oxodG:dA pair. In contrast, PolY390FExo− mutant recovered a nearly wild-type phenotype, exhibiting a 12-fold preference in the extension of such a base pair (see Table 2). These results were pointing to the importance of the hydroxyl group of φ29 DNA polymerase residue Tyr390 in the establishment of contacts with the syn conformation of 8oxodG that allow the pair 8oxodG:dA to be preferentially extended.

The structural basis for the poor extension efficiency of the 8oxodG:dC base pair have been highlighted by modelling this pair, poised for extension, at the polymerization site of T7 and RB69 DNA polymerases (19,20). Thus, the anti conformation of 8oxodG would promote local perturbations of the DNA backbone to alleviate steric clashes between the O8 group of 8oxodG and its 5' phosphate group, leading to a subtle shift in the sugar moiety of 8oxodG which would decrease the catalytic efficiency of the elongation of this base pair (19). In addition, avoidance of those steric clashes would be accounted by a slight movement of the newly replicated duplex towards the DNA major groove with the direct consequence of a diminished binding stability at the polymerase active site (20). To elucidate differences in the proper stabilization at the polymerase active site of both DNA substrates, koff assays with φ29 PolwtExo− were carried out by measuring the DNA-binding stability of the enzyme to DNA hybrid molecules containing either a dA- or dC- 3'-end primer terminus paired to 8oxodG (see Materials and Methods section and Supplementary Figure S2). Interestingly, the 8oxodG:dA base pair (koff = 0.23 ± 0.01 min−1) was bound 3-fold more stably to the PolwtExo− polymerization site than the 8oxodG:dC pair (koff = 0.70 ± 0.02 min−1). Thus, our results support the original proposal of a diminished binding stability

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Figure 4. Modelling of nascent 8oxodG(anti):dC and 8oxodG(syn):dA base pairs in the polymerase active site. Modelling an 8oxodG(syn):dA base pair shows the steric hindrance between the O8 atom and the surface formed by φ29 DNA polymerase residues Tyr390 and Gly391, according to (20). van der Waals surfaces are shown for base pairs (light brown) and for protein side chains (coloured according to their electrostaticity). 8oxodG(anti):dC [coordinates obtained from PDB1 Q9Y (20), and 8oxodG(syn):dA [PDB1 TK8 (19)] base pairs were first modelled into RB69 DNA polymerase active site (PDB1 IG9). Superposition of RB69 and φ29 (PDB1 XHX) DNA polymerase active sites allowed us to model both base pairs at the active site of the latter. Green dotted lines represent the potential hydrogen bonds formed between the nucleotides.
of the template 8oxodG\(_{\text{syn}}\) when is forming a base pair poised for extension (20).

RB69 DNA polymerase Tyr567 forms part of the minor groove mismatch sensing mechanism (8) by means of making, through a water molecule, hydrogen bonds to the sugar moiety and the acceptors located at the N\(^3\) of a purine or O\(^2\) of a pyrimidine of the ultimate template base (8). Modelling of 8oxodG\(_{\text{syn}}\):dA pair ready for extension predicted that the O\(^8\) group could substitute this water molecule and make a hydrogen bond with the hydroxyl group of Tyr567, in an analogous manner as with the N\(^3\) of a non-modified dG (20) (Figure 5), not detecting this mispair as being aberrant. In fact, the minor groove surface of this mispair is identical to the normal dT:dA base pair (19). On the contrary, the steric clashes between the O\(^8\) of 8oxodG\(_{\text{anti}}\) and sugar moiety could difficult the proper formation of the water-mediated hydrogen bond between the Tyr residue and the N\(^3\) of 8oxodG in the 8oxodG:dC pair. This elegant model proposed by Freisinger et al. (20) to explain the structural determinants for insertion and further extension opposite 8oxodG is substantiated by the biochemical results presented here. Thus, the absence of the hydroxyl group in Pol\(^{Y390F}\)Exo\(^{-}\) mutant would preclude stabilization of the 8oxodG\(_{\text{syn}}\) through such hydrogen bond, diminishing the discrimination for extending both base pairs. In this sense, relative stabilization of the 8oxodG:dA base pair \((k_{\text{on}} = 0.32 \pm 0.02 \text{ min}^{-1})\) with respect to 8oxodG:dC \((k_{\text{off}} = 0.58 \pm 0.03 \text{ min}^{-1})\) at the polymerization site of Pol\(^{Y390F}\)Exo\(^{-}\) mutant decreased in comparison with Pol\(^{Y1}\)Exo\(^{-}\) (Supplementary Figure S2), while Pol\(^{Y390F}\)Exo\(^{-}\) variant displayed a wild-type-like phenotype (data not shown). The smaller size of serine could allow one of its rotameric conformations to locate its hydroxyl group in a position suitable to contact the O\(^8\) group of 8oxodG\(_{\text{syn}}\). Additionally, the absence of the bulky aromatic ring in mutant Pol\(^{Y390F}\)Exo\(^{-}\) could favour a better accommodation of the nitrogen base of 8oxodG\(_{\text{syn}}\) that would permit the establishment of the above-mentioned interaction. Thus, allowing this mutant to discriminate against 8oxodG:dC elongation at a wild-type level. In this extension step, the aromatic group of Tyr would be dispensable, their role being restricted to confer specificity during the insertion step.

Our biochemical results, together with the structural models, let to predict the dual role of the conserved

<table>
<thead>
<tr>
<th>Mutant polymerase</th>
<th>8oxodG:dA</th>
<th>8oxodG:dC</th>
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<tr>
<td></td>
<td>(K_m)</td>
<td>(k_{\text{cat}})</td>
</tr>
<tr>
<td>Pol(^{Y1})Exo(^{-})</td>
<td>0.09 ± 0.01</td>
<td>0.15</td>
</tr>
<tr>
<td>Pol(^{Y390F})Exo(^{-})</td>
<td>2.86 ± 0.26</td>
<td>0.07</td>
</tr>
<tr>
<td>Pol(^{Y390F})Exo(^{-})</td>
<td>0.56 ± 0.05</td>
<td>0.13</td>
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</table>

\(K_{\text{cat}}/K_m\) stands for the catalytic efficiency displayed by mutant polymerases for extension of either 8oxodG:dA or 8oxodG:dA base pair. \(f_{\text{ext}}\) stands for extension discrimination factor obtained by dividing catalytic efficiency for the extension of 8oxodG:dA base pair by that for the extension of 8oxodG:dC base pair.

Figure 5. Modelling of 8oxodG\(_{\text{anti}}\):dC and 8oxodG\(_{\text{syn}}\):dA base pairs poised for extension in \(\phi29\) DNA polymerase active site. Modelling an 8oxodG\(_{\text{syn}}\):dA base pair shows the potential contact through a H-bond (green dotted line) between the O\(^8\) atom of 8oxodG and the OH group of Tyr390, in accordance to (20). van der Waals surfaces are shown for base pairs (light brown) and for Tyr390 side chain (coloured according to its electrostaticity). 8oxodG\(_{\text{anti}}\):dC [coordinates obtained from PDB1 Q9Y (20)], and 8oxodG\(_{\text{syn}}\):dA [PDB1 TK8 (19)] base pairs were first modelled into RB69 DNA polymerase ternary complex. Superposition of RB69 and \(\phi29\) (PDB1 XHX) DNA polymerase active sites allowed us to model both base pairs at the active site of the latter.
Tyr residue of motif B of family B DNA polymerases during the bypass of the 8oxodG lesions of the DNA by means of its aromatic and hydroxyl groups. The former will dictate the preferential 8oxodG(anti) conformation during the nucleotide insertion step, and as a consequence will favour the non-mutagenic dC incorporation, while the latter will do it during further base-pair extension, stabilizing the 8oxodG(syn) form promoting extension of the mispair 8oxodG:dA. Although the role of this Tyr as a key residue for controlling nucleotide misincorporation on natural DNA has been extensively described in family B DNA polymerases (55–57), this is the first time that the specific contribution of its chemical substituents during the two steps of 8oxodG replication is biochemically demonstrated. On the other hand, contrarily to thermophilic A-family DNA polymerases, currently used for PCR, as Taq polymerase from Thermus aquaticus that incorporates indistinguishably dC and dA opposite 8oxodG (58), preferential incorporation of dC opposite 8oxodG makes α29 DNA polymerase one of the most suitable enzymes to amplify faithfully DNA substrates that, as ancient DNAs, could be widely oxidized.

8oxodGTP incorporation and extension by α29 DNA polymerase

In addition to promote lesions in the DNA, the presence inside cells of ROS also generates oxidatively altered purines and pyrimidines, the most abundant being 8oxodGTP (14). Although organisms synthesize 8oxodGTPases to sanitize the dNTP pool, the incorporation of such modified nucleotide could have adverse consequences as pairing opposite adenine of the template strand would result in A:T to C:G transversions (59).

Primer extension analyses similar to those described above were performed to analyse the templating nucleotide preference (dA versus dC) during 8oxodGMP incorporation by α29 DNA polymerase, using the 15/21-mer hybrid molecules spl/sp1C+5 and spl/sp1aA+5 described in the Materials and Methods section (see also Supplementary Figure S3). As it can be seen in Table 3, α29 DNA polymerase inserts 8oxodG preferentially opposite dC, although 2000-fold less efficiently than insertion of the unmodified dGTP. Additionally, qualitative analysis of the extension of both dA:8oxodG and dC:8oxodG base pairs obtained after insertion of 8oxodG in these molecules (see Materials and Methods section), show the preferential extension of the correct dC:8oxodG base pair (Figure 6), with an efficiency similar to that of the normal dC:dG base pair (the appearance of the 18- and 20-mer bands is due to the pairing of 8oxodG with the templating dA and dC at these positions, respectively). These are very unusual results as most of DNA polymerases assayed displayed a favoured 8oxodG insertion opposite dA, as T7 and γ DNA polymerases from family A (22, 60), β and λ DNA polymerases from family X (39, 45, 47) and Dpo4 and Dbh DNA polymerases belonging to family Y (61). Solely HIV-RT, Pol Ikk (family A), E. coli DNA pol II (family B) and DNA polymerase from African Swine Fever Virus (family X) showed an 8oxodG insertion pattern similar to that described for α29 DNA polymerase (22, 45). Conversely, PolIk and α29 DNA polymerases are the only enzymes studied so far exhibiting a prone extension of the correct dC:8oxodG base pair (22) (this study).

At present, DNA polymerase ternary complexes with 8oxodGTP as incoming nucleotide have not been solved. This fact precludes the analysis of the structural rationale for 8oxodG insertion preferences. In Figure 7, we have modelled dC:8oxodGTP(anti) and dA:8oxodGTP(syn) base pairs into the RB69 DNA polymerase active site. As it happens with a non-modified incoming nucleotide, Lys560 residue, placed at the fingers subdomain, will make a hydrogen bond to one of the two negatively charged equatorial oxygens of the α phosphate of the incoming 8oxodGTP(anti), enhancing catalysis by allowing the proper stabilization of the catalytic state (8). Nevertheless, protrusion into the major groove of N1, N2 and O6 atoms of the incoming 8oxodGTP(syn) could lead to a sterical and/or electrostatic clash between the N2 atom and the edge of the side chain of Lys560 (red patch in Figure 7), affecting the proper interaction between this residue and the α phosphate of the 8oxodGTP(syn) and, as a consequence, its insertion. Structural alignments of both RB69 and α29 DNA polymerases have allowed us to identify α29 DNA polymerase Lys383 as the homologous lysine residue (62). By the contrary, we have not found a structural rationale for the preferential extension of the dC:8oxodG base pair, as once inserted, the 8oxodGMP(syn) should be translocated backwards to the primer terminus site. At this location, the only contacts between the protein and the 3′-end nucleotide are through the phosphodiester bond. The impaired elongation of the incorrect base pair could suggest that the syn conformation of the 3′ terminus either promotes a subtle distortion in the proper orientation of the attacking 3′-OH group to form the phosphodiester bond with the next incoming nucleotide, or a slightly variation in the angle of the glycosidic bond that could make difficult the establishment of the proper stacking interactions between the nitrogen bases of the incoming nucleotide and primer terminus. Difficulties in the translocation step due to the presence of 8oxodGMP(syn) at the 3′ primer terminus cannot be ruled out.

As mentioned above, several DNA polymerases from eukaryotes and prokaryotes can use 8oxodGTP as incoming nucleotide, leading to transversions. Surveillance mechanisms have evolved to maintain a low

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<tr>
<td>dG insertion</td>
<td>1.71 × 10⁻³</td>
<td>3.68</td>
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<tr>
<td>opposite dC</td>
<td>± 0.16 × 10⁻³</td>
<td>2.15 × 10⁻³</td>
</tr>
<tr>
<td>8oxodG insertion</td>
<td>3.27 ± 0.58</td>
<td>3.48</td>
</tr>
<tr>
<td>opposite dC</td>
<td>1.07</td>
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<tr>
<td>8oxodG insertion</td>
<td>9.32 ± 1.25</td>
<td>3.3</td>
</tr>
<tr>
<td>opposite dA</td>
<td>0.35</td>
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\((K_{cat}/K_{m})\) stands for the catalytic efficiency displayed by mutant polymerase for the formation of the indicated base pair.
frequency of these types of mutations, owing to the action of enzymes able to degrade such mutagenic substrates (14). Thus, *E. coli* synthesizes MutT protein that hydrolyses 8oxodGTP to 8oxodGMP (59), this latter form being unable to be rephosphorylated, preventing insertion of 8oxodGTP during DNA synthesis. Analogously, mammalian cells code for MTH1, a protein showing a similar enzymatic activity (63–65). However, proteins as *E. coli* MutY, which excises a dA mispaired with 8-oxodG as part of the process to restore the original G:C base (66) could have a deleterious effect if 8oxodG had been the erroneously incorporated nucleotide.

**Fig. 7.** Modelling of nascent dC:8oxodGTP(anti) and dA:8oxodGTP(syn) base pairs in the RB69 DNA polymerase active site. Modelling a dA:8oxodGTP(syn) base pair shows the potential steric hindrance between the N2 atom of 8oxodGTP and the edge of the side chain of Lys560 (red patch). van der Waals surfaces are shown for base pairs (light brown) and for Lys560 side chain (coloured according to its electrostaticity). Green dotted line represents the hydrogen bond between Lys560 and the γ phosphate of the incoming 8oxodG(anti). Modelling was performed essentially as described above.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

**ACKNOWLEDGEMENTS**

We are grateful to Dr Javier Saturno for the preparation of the φ29 DNA polymerase mutants PolY<sup>390F</sup>Exo<sup>-</sup> and PolY<sup>390S</sup>Exo<sup>-</sup>, to José M. Lázaro for the purification of the proteins and to Dr Luis Blanco for critical reading of the manuscript and for helpful discussions.
REFERENCES