Repair of oxidative DNA damage in *Drosophila melanogaster*: identification and characterization of dOgg1, a second DNA glycosylase activity for 8-hydroxyguanine and formamidopyrimidines

Claudine Dherin, Miral Dizdaroglu, Hélène Doerflinger, Serge Boiteux and J. Pablo Radicella*

CEA, Département de Radiobiologie et Radiopathologie, UMR217 CNRS-CEA, Radiobiologie Moléculaire et Cellulaire, 60 rue du Général Leclerc, BP6, 92265-Fontenay aux Roses, France, 1Chemical Science and Technology Laboratory, National Institute of Standards and Technology, Gaithersburg, MD 20899, USA and 2Institut Jacques-Monod, CNRS, Université Paris7-Denis Diderot, Université Paris 6-P. et M. Curie, 2 place Jussieu, 75251-Paris, France

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ABSTRACT

In *Drosophila*, the S3 ribosomal protein has been shown to act as a DNA glycosylase/AP lyase capable of releasing 8-hydroxyguanine (8-OH-Gua) in damaged DNA. Here we describe a second *Drosophila* protein (dOgg1) with 8-OH-Gua and abasic (AP) site DNA repair activities. The *Drosophila OGG1* gene codes for a protein of 327 amino acids, which shows 33 and 37% identity with the yeast and human Ogg1 proteins, respectively. The DNA glycosylase activity of purified dOgg1 was investigated using γ-irradiated DNA and gas chromatography/isotope dilution mass spectrometry (GC/IDMS). The dOgg1 protein excises 8-OH-Gua and 2,6-diamino-4-hydroxy-5-formamidopyrimidine (FapyGua) from γ-irradiated DNA. with $k_{cat}/K_m$ values of $21.0 \times 10^{-5}$ and $11.2 \times 10^{-5}$ (min$^{-1}$ nM$^{-1}$), respectively. Enzymatic assays using oligodeoxyribonucleotides containing a single lesion show that dOgg1 displays a marked preference for DNA duplexes containing 8-OH-Gua, 8-OH-Ade or an AP site placed opposite a cytosine. The cleavage of the 8-OH-Gua-containing strand results from the excision of the damaged base followed by a β-elimination reaction at the 3′-side of the resulting AP site. Cleavage of 8-OH-Gua.C duplex involves the formation of a reaction intermediate that is converted into a stable covalent adduct in the presence of sodium borohydride. dOgg1 complements the mutator phenotype of fpg mutY mutants of *Escherichia coli*. Whole-mount in situ hybridizations on tissues at different stages of *Drosophila* development reveal that the dOgg1 messenger is expressed uniformly at a low level in cells in which mitotic division occurs. Therefore, *Drosophila* possesses two DNA glycosylase activities that can excise 8-OH-Gua and formamidopyrimidines from DNA, dOgg1 and the ribosomal protein S3.

INTRODUCTION

Reactive oxygen species (ROS) induce DNA damage that has been implicated to play a role in mutagenesis, carcinogenesis and aging (1–5). ROS produce a variety of damages in DNA including base and sugar damage, sites of base loss, strand breaks and DNA–protein crosslinks (6,7). Oxidatively damaged DNA bases are thought to be primarily repaired by the base excision repair (BER) pathway in prokaryotes and eukaryotes (8–11). 8-Hydroxyguanine (8-OH-Gua) is a prevalent DNA lesion caused by oxidizing agents and ionizing radiation and is highly mutagenic (12–15). In *Escherichia coli*, two DNA glycosylases prevent mutagenesis by 8-OH-Gua: the Fpg protein which excises 8-OH-Gua in damaged DNA and the MutY protein which excises the adenine residues incorporated by DNA polymerases opposite 8-OH-Gua. Inactivation of both the fpg (mutM) and mutY (micA) genes of *E.coli* results in a strong G:C to T:A mutator phenotype (12–15). In *Saccharomyces cerevisiae*, a DNA glycosylase that is encoded by the *OGG1* gene and named yOgg1, catalyses the removal of 8-OH-Gua and formamidopyrimidines from damaged DNA (16–19). Furthermore, Ogg1-deficient strains of *S.cerevisiae* exhibit a G:C to T:A mutator phenotype (20,21). In mammalian cells, 8-OH-Gua is released by DNA glycosylases showing strong sequence homology with the yeast Ogg1 and named mOgg1 and hOgg1 for the mouse and human enzymes, respectively (22–28). To investigate the biological role of the Ogg1 protein in mammalian cells, *ogg1*−/− mice have been generated (29,30). Null animals are viable and show no marked pathological changes up to 12 months. However, Ogg1-deficient mice show an abnormal accumulation of 8-OH-Gua in their genome and exhibit a significantly higher spontaneous mutation rate in non-proliferative tissues compared to the isogenic wild-type (29,30). These results strongly suggest that excision of 8-OH-Gua by DNA glycosylases such as the bacterial Fpg and the eukaryotic Ogg1, prevent mutations induced by endogenous ROS.

*To whom correspondence should be addressed. Tel: +33 1 46 54 88 57; Fax: +33 1 46 54 88 59; Email: jpradicella@cea.fr*
Although they have similar biological functions, the Fpg and Ogg1 proteins do not exhibit significant sequence homology (19). Indeed, Ogg1 belongs to a family of DNA glycosylases/AP lyases, the signature of which is the α-helix–hairpin–α-helix–Gly/Pro-Asp motif together with a conserved catalytic lysine, which we refer to as the HhH-GPD/K family (18,31), the prototype of this family being the Nth (Endo III) protein of E.coli (18,32).

Yeast and mammalian Ogg1 proteins have very similar, if not identical, substrate specificities and catalytic properties. The Ogg1 proteins are monomers of ~40 kDa endowed with both DNA glycosylase and AP lyase activities (reviewed in 12). They excise 8-OH-Gua, 2,6-diamino-4-hydroxy-5-N-methylformamidopyrimidine (Me-FapyGua) and 2,6-diamino-4-hydroxy-5-formamidopyrimidine (FapyGua) from damaged DNA (16–19,22–28). In addition, the Ogg1 proteins efficiently remove 8-hydroxyadenine (8-OH-Ade) placed opposite a cytosine in oligodeoxyribonucleotides (33,34). Yeast and human Ogg1 proteins are also endowed with an AP lyase activity that incises DNA at AP sites placed opposite a cytosine via a β-elimination reaction (16,24). The catalytic mechanism of the Ogg1 proteins involves the formation of a transient covalent imino–enzyme–DNA intermediate presumably between the ε-NH2 group of a catalytic lysine residue and the C1′ of the abasic sugar moiety (16,24,25,31).

In Drosophila, the ribosomal protein S3 has been shown to possess an 8-OH-Gua DNA glycosylase/AP lyase activity (35–37). The glutathione S-transferase (GST) fusion of the Drosophila S3 (GST-S3) efficiently releases 8-OH-Gua and FapyGua from γ-irradiated DNA (35). These results suggested that S3 could play a role in DNA repair in addition to its involvement in protein synthesis. In the present study, we show that Drosophila also possesses an OGG1 homolog, dOgg1. The data presented show that this gene expresses a functional DNA glycosylase/AP lyase, capable of excising 8-OH-Gua and FapyGua from DNA exposed to γ-irradiation. Our results also show that dOgg1 efficiently incises 34mer DNA duplexes containing 8-OH-Gua, 8-OH-Ade and AP sites placed opposite to a cytosine. The biochemical properties of the dOgg1 protein were compared to those of S3. The expression pattern of the dOOG1 mRNA in Drosophila larvae and ovaries is also presented. Therefore, Drosophila is the first eukaryotic organism to be found having two functional 8-OH-Gua DNA glycosylase activities.

**MATERIALS AND METHODS**

**Materials**

Modified DNA bases, their stable isotope-labeled analogs and other materials for GC/IDMS were obtained as described previously (38). Calf thymus DNA and poly(dG.dC).poly(dG.dC) were purchased from Sigma and Boehringer, respectively. Restriction endonuclease, DNA polymerases, T4 DNA kinase and T4 DNA ligase were from New England Biolabs. DNA repair proteins from E.coli (Ung, Endo III, Endo IV or Fpg) and yeast Ogg1 were purified from overproducing strains of E.coli according to standard protocols (our laboratory stocks).

**Preparation of DNA substrates**

DNA samples for GC/IDMS analysis were prepared as follows. Calf thymus DNA was dissolved in phosphate buffer (pH 7.4) at a concentration of 0.3 mg/ml. An aliquot of this solution was bubbled with N2O and irradiated with γ-rays in a 60Co γ-source at a dose of 50 Gy (dose rate 35.5 Gy/min). Subsequently, unirradiated and irradiated DNA solutions were dialyzed against 10 mM phosphate buffer (pH 7.4). The [4H]-Me-FapyGua-poly(dG.dC).poly(dG.dC) substrate was prepared as previously described (39). The oligodeoxyribonucleotides used in this study are 34mers with the following sequences: 5′-GGCTTTCATCGTTGTCC[X]CAGACCTGGTGATACCG-3′ with [X] for 8-OH-Gua, 8-OH-Ade or uracil (OligoExpress, Grenoble), respectively. The complementary sequences with each of the four DNA bases placed opposite [X] in the duplex were from OligoExpress (Grenoble, France). To generate the AP sites, the 34mer DNA containing uracil was incubated in the presence of uracil DNA glycosylase as described (16).

**Cloning, expression and purification of the Drosophila dOgg1 protein**

An expressed sequence tag (EST) clone coding for dOgg1 was retrieved from the Berkeley Drosophila Genome Project database. The cDNA in a pBluescript vector (Stratagene) was sequenced using an ALFexpress automated sequencer (Amersham Pharmacia Biotech). The open reading frame identified was PCR amplified using the Pfu polymerase (Stratagene) with the following primers: 5′-GGGATCCATTGAAGCTCGTGGTCTTACTG and 5′-TAGTGAAGTACCTTTTAAAG. The resulting fragment was digested with BamHI and ligated to plasmid pGEX-4T1 (Amersham Pharmacia Biotech) digested with SmaI and BamHI. The plasmid obtained (pPR195) codes for a GST peptide with the Drosophila OGG1 cDNA fused at its C-terminus. The plasmid was introduced into the E.coli strain BH410 (pgp−). Escherichia coli BH410 harboring pPR195 was grown at 37°C in LB broth medium (2 l) containing 150 µg/ml ampicillin until the absorbance at 600 nm reached 0.3 and induced for 16 h at 20°C in the presence of IPTG (1 mM). Cells were collected and stored at −80°C. Cell pellets were resuspended in 10 ml/g of lysis buffer (20 mM Tris–HCl, pH 8.0, 1 mM Na2EDTA, 250 mM NaCl, 0.8 µg/ml antipain, 0.8 µg/ml leupeptine, 0.8 µg/ml aprotinin and 0.8 µg/ml aprotinin). After centrifugation of the cell lysate, the supernatant fraction was dialyzed against PBS buffer and applied to glutathione–Sepharose 4B (Pharmacia Biotech) equilibrated with PBS buffer. The Sepharose was washed with PBS and proteins eluted with a buffer containing 50 mM Tris–HCl pH 8.0 and 30 mM reduced glutathione. Fractions containing the enzyme activity were pooled and dialyzed overnight against a buffer containing 50 mM Tris–HCl pH 8.0 and 50 mM NaCl and subsequently 4 h against 50 mM Tris–HCl pH 8.0 with 150 mM NaCl and 2.5 mM CaCl2. Thrombin (ICN) was added (10 U/mg of GST–dOgg1 fusion protein) and the mixture incubated for 2 h at 25°C. Reactions were stopped by adding Na2EDTA to 5 mM final. Proteins were dialyzed against a buffer containing 20 mM Tris–HCl pH 8.0, 2 mM Na2EDTA, 50 mM NaCl and 2% glycerol and applied to a MonoS column (FPLC system, Pharmacia Biotech). Proteins were eluted with a linear salt gradient (50–800 mM NaCl). The active fractions were pooled and their protein concentration...
determined by the method of Bradford (40). Purification of dOgg1 was followed by SDS–PAGE and using the excision of [³H]-Me-FapyGua from [³H]-Me-FapyGua-poly(dG,dC)-poly(dG,dC) as an activity assay (39).

Enzymatic assays and GC/IDMS analysis

The determination of the dependence of excision on the enzyme amount and on the incubation time, the measurement of excision kinetics and the GC/IDMS analysis were performed as described previously (41). The amount of dOgg1 used for the measurement of excision kinetics corresponded to an enzyme concentration of 267 nM.

Assays for cleavage of 34mer DNA duplexes

The DNA strand containing the lesion (8-OH-Gua, 8-OH-Ade or AP site) was ³²P-labeled at the 5′-end and annealed with each of the four complementary sequences (16,33,34). The reaction mixtures (10 µl final volume) contained 25 mM Tris–HCl pH 7.6, 2 mM Na₂EDTA, 50 mM NaCl, 50 fmol of ³²P-labeled DNA duplex and dOgg1. Reactions were performed at 37°C for 15 min. Reactions were stopped by adding 6 µl of formamide dye and subjected to 7 M urea–20% PAGE (16,34). Gels were scanned and quantified using a PhosphorImager (Molecular Dynamics) (33).

Trapping assay

The reaction mixture (20 µl final volume) contained 25 mM Tris–HCl pH 7.6, 2 mM Na₂EDTA, 100 fmol of labeled 8-OH-Gua,C duplex, 50 mM of NaBH₄ and 50 ng of either dOgg1, yOgg1 or Fpg. The reaction was carried out at 37°C for 20 min. The reactions were stopped by addition of 10 µl of SDS–PAGE loading buffer/formamide blue dye and heating at 90°C for 3 min. The products of the reactions were separated onto 15% SDS–PAGE and analyzed as previously described (16).

Mutagenesis experiments

The complementation of the spontaneous mutator phenotype of an E.coli fpg mutY double mutant (PR195) by the expression of the different GST fusion proteins was analyzed by determining the frequency of rifampicin-resistant cells in 20 independent cultures (26).

Whole-mount in situ hybridization

Whole-mount RNA in situ hybridization on ovaries and larvae was performed with the complete dOgg1 cDNA. Digoxigenin-labeled sense and antisense RNA probes were synthesized using the RNA Genius kit (Boehringer Mannheim) according to manufacturer’s protocol. The hybridization procedures were carried out according to Tautz and Pfeifle (42) with minor modifications (43) and other modifications as follow: Dissected tissues were fixed in 4% paraformaldehyde in PBS–0.1% Tween-20 (PBT) for 20 min at room temperature (RT) and washed three times in PBT. Tissues were then incubated in boiling water for 5 min. Prehybridization was in hybridization solution (HS) for 1 h at 65°C, and hybridization was overnight at 65°C. After rinsing twice in HS for 20 min at 65°C, tissues were washed three times in PBT for 20 min at RT. After the incubation with anti-DIG antibody (1/2000 in PBT) for 1 h at RT, tissues were washed three times in PBT for 20 min at RT and brought into staining buffer. Tissues were stained for 1 h at RT, washed three times in PBT and mounted in 70% glycerol.

RESULTS

Isolation of a Drosophila cDNA coding for an Ogg1 protein homolog

A cDNA clone coding for a peptide showing a high degree of homology to the human and yeast Ogg1 proteins was identified in the EST database from the Berkeley Drosophila Genome Project. The dOgg1 cDNA is 1284 bp long and codes for a 327 amino acid peptide. Sequence alignments show that the protein coded, dOgg1, shares 33 and 37% identity with the yeast and human Ogg1, respectively. Furthermore, dOgg1 possesses all the hallmarks of the Ogg1 proteins. In particular, the highly conserved active site motif helix–hairpin–helix motif-PVD/K is present between residues 209 and 260 (Fig. 1A). Comparison of the cDNA with the complete Drosophila genome allowed the identification of the corresponding genomic sequence for the dOgg1 gene and its localization to chromosome 1. Comparison between the cDNA and the genomic sequences shows that dOgg1 has four exons separated by introns whose sizes are 58, 61 and 59 bp (Fig. 1B). The transcribed region spans 1.5 kb of genomic sequences.

Expression and purification of the Drosophila Ogg1 protein

The Drosophila OGG1 cDNA was cloned into the bacterial vector pGEX-4T1 to express dOgg1 fused to the C-terminus of GST. The GST–dOgg1 protein was purified from bacterial lysates by affinity to glutathione–Sepharose. Treatment with thrombin allowed the cleavage of the GST tag. The dOgg1 protein was separated from GST and thrombin after chromatography on a MonoS column. The purity of the dOgg1 protein was assessed by the observation of a single protein band on a SDS–PAGE with a molecular mass of ~38 kDa, which agreed well with the expected mass (Fig. 2). The specificity of the site of cleavage was confirmed by a single N-terminal sequence (data not shown). The purification steps were monitored by measuring the excision of [³H]-Me-FapyGua by the protein fractions. For the purified dOgg1 protein, the specificity constant (kₐ/Kₘ) for the excision of Me-FapyGua is 214 × 10⁻⁵ (min⁻¹ nM⁻¹). This value is very similar to that measured for the human α-hOgg1 protein (33).

Excision of modified bases by the Drosophila dOgg1 from γ-irradiated DNA: GC/IDMS analysis

We investigated the ability of the dOgg1 protein to excise modified DNA bases from γ-irradiated DNA. Using GC/IDMS, 17 modified bases were identified and quantified in DNA γ-irradiated under N₂O (44). Of these modified bases, dOgg1 protein efficiently excised FapyGua and 8-OH-Gua. No other modified base was excised significantly under the conditions used in this work. The excision of 8-OH-Gua and FapyGua was assessed by their appearance in supernatant fractions of DNA substrates incubated with active enzyme (Fig. 3). The amounts of 8-OH-Gua and FapyGua in DNA pellets after incubation of γ-irradiated DNA with active protein were significantly reduced when compared to those in DNA pellets incubated with heat-inactivated enzyme or without enzyme. The amounts found in the supernatant fractions of DNA substrates incubated with active enzyme were similar to those removed from the pellets of the same DNA substrate, demonstrating the excision of the lesions (Fig. 3). Kinetic parameters were determined by measurement of excision at six different times.
concentrations of FapyGua and 8-OH-Gua with the total amount of DNA remaining constant in each sample. The excised amounts of these products in supernatant fractions were used for the determination of the initial velocity. Excision

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**Figure 1. Drosophila OGG1 sequence analysis.** (A) Sequence alignment of the dOgg1 protein with its yeast (Sc), plant (At) and human (Hs) homologs. Black and gray boxes identify identical and similar residues respectively. Asterisks in the consensus line indicate conservation of the amino acid in all four proteins. (B) Structure of the dOGG1 gene. Boxes indicate exons.
94 67 40 14

Table 1. Kinetic constants for excision of 8-OH-Gua and FapyGua by Drosophila Ogg1 protein from DNA γ-irradiated under N2O

<table>
<thead>
<tr>
<th>Lesion</th>
<th>V_{max} (nM min^{-1})</th>
<th>K_{cat} (nM)^{a}</th>
<th>k_{cat}/V_{max} × 10^{3} (min^{-1})</th>
<th>k_{cat}/K_{cat} × 10^{5} (min^{-1} nM^{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>8-OH-Gua</td>
<td>15.0 ± 0.3</td>
<td>267 ± 16^{a}</td>
<td>56.2 ± 1.1</td>
<td>21.0 ± 0.4^{a}</td>
</tr>
<tr>
<td>FapyGua</td>
<td>12.7 ± 0.2</td>
<td>422 ± 22</td>
<td>47.5 ± 0.7</td>
<td>11.2 ± 0.2</td>
</tr>
</tbody>
</table>

{^a}Values represent the mean ± standard deviation (n = 6). k_{cat} = V_{max}/[enzyme]; ([enzyme] = 267 nM). The concentration ranges of the FapyGua and 8-OH-Gua were 0.82 – 3.8 µM, and 0.41 – 2.3 µM, respectively.

{^b}Statistically different from the value in line 2 (p < 0.05).

followed Michaelis–Menten kinetics (45). Kinetic constants and standard deviations (n = 6) were calculated using Lineweaver–Burk plots and a linear least-squares analysis of the data (data not shown). Kinetic parameters calculated from these plots are given in Table 1. The specificity constant (k_{cat}/K_{cat}) for excision of 8-OH-Gua is 1.9-fold higher than that for FapyGua (Table 1). This result suggests that, for the dOgg1 protein, 8-OH-Gua may be a better substrate than FapyGua in N_{2}O γ-irradiated DNA.

Cleavage of 34mer DNA duplexes containing 8-OH-Gua, 8-OH-Ade or an AP site placed opposite each of the four DNA bases

The specificity of the dOgg1 protein was also investigated using DNA lesions embedded in 34mer DNA duplexes as substrates. These substrates were also used to analyze the opposite base dependence of the dOgg1 activity on 8-OH-Gua, 8-OH-Ade and AP site. For this purpose, 34mer harboring one of these lesions was hybridized to complementary sequences harboring each of the four normal DNA bases opposite the lesion. The cleavage activity of the dOgg1 protein on the strand harboring the lesion was analyzed for each of the four DNA duplexes (Fig. 4). In the case of 8-OH-Gua, dOgg1 efficiently cleaves 8-OH-Gua.C and 8-OH-Gua.T duplexes whereas it cleaves 8-OH-Gua.G and 8-OH-Gua.A duplexes at a very slow rate (Fig. 4A). In the case of 8-OH-Ade and AP site, dOgg1 efficiently cleaves 8-OH-Ade.C and AP.C duplexes (Fig. 4B and C). In contrast, dOgg1 does not cleave 34mer DNA duplexes containing an 8-OH-Ade or an AP site placed opposite a thymine, a guanine or an adenine (Fig. 4B and C). This last result explains why 8-OH-Ade is not released from γ-irradiated DNA substrates which contain 8-OH-Ade.T base pairs.

Catalytic mechanism of the dOgg1 protein

To further investigate the catalytic mechanism of dOgg1, we compared the products generated by dOgg1, yOgg1 and Fpg after cleavage of an 8-OH-Gua.C duplex. Figure 5A shows that dOgg1 and yOgg1 generate the same product (P1) corresponding to the cleavage of the phosphodiester bond at the 3′-side of the AP site resulting from the excision of the 8-OH-Gua [18]. It is widely accepted that this product results from a β-elimination reaction catalyzed by the AP lyase activity of this class of enzymes (46). In contrast, the Fpg protein generates a product that migrates faster (P3) and results from successive β- and δ-elimination reactions at the AP site generated by the glycosylase activity (16,46). Control experiments show that the dOgg1 product P1 is not modified by post-treatment with Endo III (Fig. 5A). In contrast, the addition of Endo IV after cleavage by the dOgg1 protein results in a nearly quantitative conversion of the dOgg1 product P1 into another product (P2) which is probably generated by the release of the trans-4-hydroxy-2-pentenal-5-phosphate at the 3′-end of the P1 fragment (16,36,46). These results strongly suggest that the mechanism of strand cleavage by the dOgg1 protein involves the release of the 8-OH-Gua residue followed by a β-elimination reaction at the 3′-side of the resulting AP site.
The mechanism of DNA strand cleavage after excision of 8-OH-Gua by DNA glycosylases/AP lyases such as the bacterial Fpg or the yeast and human Ogg1 involves the nucleophilic attack at the C1' of the sugar moiety leading to the formation of an enzyme–DNA Schiff base intermediate (46,47). To probe for such an intermediate, the Ogg1 protein was allowed to react with a labeled 8-OH-Gua.C substrate in the presence of the reducing agent NaBH4 and the products of the reactions were analyzed by SDS–PAGE. Figure 5B shows that the presence of NaBH4 in the assay mixture results both in the inhibition of the cleavage activity of the dOgg1 protein and in the formation of a shifted band (trapped complex) with an apparent molecular weight of 55–60 kDa (Fig. 5B). The dOgg1–DNA complex can be digested with proteinase K to yield a product that co-migrates with free DNA (data not shown). Control experiments show that the dOgg1–DNA complex is not formed when NaCl is present instead of NaBH4 (Fig. 5B). Figure 5B also shows the formation of a Fpg–DNA and yeast Ogg1–DNA complexes in the presence of 32P-labeled 34mer carrying a single 8-OH-Gua residue was labeled at its 5' end and hybridized with a complementary sequence carrying a cytosine opposite the lesion. Incubation were performed with 10 ng of dOgg1, yOgg1 or Fpg protein for 15 min at 37°C. When indicated, reaction mixtures were then incubated with 10 ng of either endonuclease III (Nth) or endonuclease IV (Nfo). The products were separated by denaturing 20% PAGE. P1, P2 and P3 are the products described in the Results. (B) NaBH4-mediated trapping assay. The 34mer carrying a single 8-OH-Gua residue was labeled at its 5’-end and hybridized with a complementary sequence carrying a cytosine opposite the lesion. Fifty nanograms of either Fpg, yOgg1 or dOgg1 proteins were allowed to react for 20 min at 37°C with the 32P-labeled 34mer duplex DNA, containing a single 8-OH-Gua.C, in the presence of 50 mM of either NaCl or NaBH4. The products of the reactions were separated using 15% SDS–PAGE.
of NaBH₄. These results suggest that the Ogg1 protein forms a transient Schiff base intermediate, which is converted into a covalent protein–DNA adduct in the presence of NaBH₄.

Expression of the dOgg1 proteins in *E.coli* fpg mutY complements the mutator phenotype

Plasmid PR195 expressing GST–dOgg1 protein was transformed into *E.coli* strain PR195 in which the mutY and fpg genes are disrupted. This strain displays a strong spontaneous mutator phenotype due to its incapacity to eliminate errors induced by the presence of 8-OH-Gua in its DNA. The rates of mutation to rifampicin resistance were determined (26). Table 2 shows that the expression of the *Drosophila* protein in this strain reduces the mutation frequencies to rifampicin resistance as effectively as the human Ogg1 protein expression, partially complementing the spontaneous mutator phenotype.

Expression pattern of dOGG1 in *Drosophila* ovaries and larvae

We examined the distribution of the *dOGG1* transcripts in ovaries and larvae by whole-mount RNA *in situ* hybridization. In all the tested tissues, a specific signal has been observed after a long time of staining compared to a control probe (Fig. 6). This result suggests that the *dOGG1* mRNA might be expressed at a low level in these tissues. Ovaries are composed of about 16 ovarioles, each with a gerarium at its anterior tip and progressively older egg chambers toward the posterior. An egg chamber consists of three cell types: the oocyte connected to 15 polyplide nurse cells, both surrounded by a monolayer of somatic follicle cells. The subsequent development of the egg chamber is divided into 14 stages. From stage 10B onwards, the nurse cells expel their entire content into the oocyte. At the end of the oogenesis, the columnar follicle cells synthesize the vitelline membrane and the chorion of the egg and then both nurse cells and follicle cells degenerate. Hybridization on ovaries with an antisense *dOGG1* probe revealed a signal in the cytoplasm of the nurse cells from stage 3 (data not shown) and in the oocyte’s cytoplasm from stage 10B onwards (Fig. 6A and B). Hybridization on third instar larvae with an antisense *dOGG1* probe revealed a specific signal in the imaginal discs which are the precursors of the epidermal structures of the head, thorax and external genitalia of the adult fly (Fig. 6C and D). In late third instar larvae, they appear as sac-like epithelial organs.

**DISCUSSION**

The unavoidable but potentially mutagenic DNA base alterations resulting from endogenous ROS attack are primarily removed by the BER pathway (8–11). The first step in this ubiquitous repair pathway is the recognition and removal of the altered base by a DNA glycosylase catalyzing the cleavage of the glycosylic bond between the modified base and the sugar moiety, leaving an AP site in DNA. Subsequently, the resulting AP site is incised and the repair is completed by the successive actions of a phosphodiesterase, a DNA polymerase and a DNA ligase (8–11). In yeast three DNA glycosylases are involved in the removal of oxidatively-damaged DNA bases, namely Ntg1, Ntg2 and Ogg1 (48). Structural and/or functional homologs of these DNA glycosylases have been found in higher eukaryotes (2). The biological function of this class of enzymes is thought to prevent genetic instability due to endogenous DNA base damage such as 8-OH-Gua. Assessments of 8-OH-Gua formation and mutagenic potential are strongly supported by the G:C to T:A mutator phenotype of *E.coli*, *S.cerevisiae* and mouse.
Three classes of 8-OH-Gua DNA glycosylases have been identified in living organisms: (i) The Fpg family in Bacteria and Arabidopsis thaliana (49); (ii) the Ogg1 family in Archae (50), yeast and mammals (2); and (iii) The S3 ribosomal protein in Drosophila (35–37). In the present study, we report sequence alignments that indicate the presence of Ogg1 homologs in A. thaliana and Drosophila (Fig. 1A). The high degree of sequence homology and the conservation of critical blocks of amino acids strongly suggest that these are functional 8-OH-Gua DNA glycosylases from the Ogg1 family (2). The results from the characterization of the purified dOgg1 protein from Drosophila presented in this paper show that the dOgg1 protein is indeed a DNA glycosylase that removes Me-FapyGua, FapyGua and 8-OH-Gua from damaged DNA as do the yeast and human Ogg1 (16–19,22–28). Previous experiments had suggested that there is no Fapy glycosylase activity in human Ogg1 (49). These results lead us to conclude that Drosophila has two 8-OH-Gua DNA glycosylase activities, S3 and dOgg1, because of this apparent redundancy, it was important to analyze in detail the properties and the expression of the dOgg1 protein. Indeed, dOgg1 and S3 have very similar properties. Both dOgg1 and S3 are DNA glycosylases that excise 8-OH-Gua and FapyGua from γ-irradiated DNA. Specificity constants for the excision of 8-OH-Gua and FapyGua by dOgg1 (Table 1) and S3 (35) have been measured using the same experimental conditions. The results show that $k_{cat}/K_M$ values for excision of 8-OH-Gua and FapyGua by dOgg1 were 8.1- and 4.5-fold higher than that by S3, respectively. The better $k_{cat}/K_M$ values of dOgg1, compared to S3, are due to lower $K_M$ values.

Furthermore, both dOgg1 and S3 have a marked preference for 8-OH-Gua placed opposite a cytosine. Finally, dOgg1 and S3 are AP lyases whose reaction mechanisms involve the formation of a covalent imino enzyme–DNA intermediate. dOgg1 and S3 have then very similar, if not identical, substrate specificities and potentially the same biological function.

In the present study, we also show that dOgg1 mRNA is ubiquitously expressed in Drosophila tissues. The experiments presented suggest that the nurse cells synthesize the dOgg1 transcript and then expel it in the oocyte. The zygote is transcriptionally inactive during the first 14 division of embryogenesis. Therefore, the accumulation of the dOgg1 transcript in the oocyte might be important for the mitotic divisions that occur during early embryogenesis, in order to avoid oxidative damage-induced mutations. No dOgg1 expression is detected in follicle cells. These cells will not divide any more as they degenerated at the end of the oogenesis. Then, we could expect that they do not need an important DNA repair machinery to keep their DNA without damage. The presence at a detectable level of the dOgg1 transcript in the imaginal discs could be related to the rapid divisions of the disc’s cells during larval development. Since S3 is essential, both S3 and Ogg1 are likely to be present in most Drosophila cells. The dOgg1 possesses a cluster of basic amino acids at its C-terminal end which is probably used as a nuclear localization signal as previously reported for the human Ogg1 (24). Similarly, the S3 ribosomal protein has been suggested to possess a nuclear localization signal and to bind the nuclear matrix (37). Therefore, both dOgg1 and S3 may have access to the nuclear DNA. All the data point then to the fact both dOgg1 and S3 are functional in vivo. Although dOgg1 seems to have somehow better kinetic properties than S3 the relative importance of their activities in the Drosophila cells remains to be determined. The transcription-coupled repair of 8-OH-Gua in Ogg1-deficient rodent cells reveals the existence of an unidentified Ogg1-independent repair pathway in eukaryotes (54). If it exists in Drosophila, this could be a third repair system for 8-OH-Gua.


