Clustered DNA Lesions Containing 5-Formyluracil and AP Site: Repair via the BER System

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Abstract

Lesions in the DNA arise under ionizing irradiation conditions or various chemical oxidants as a single damage or as part of a multiply damaged site within 1–2 helical turns (clustered lesion). Here, we explored the repair opportunity of the apurinic/apyrimidinic site (AP site) composed of the clustered lesion with 5-formyluracil (5-foU) by the base excision repair (BER) proteins. We found, that if the AP site is shifted relative to the 5-foU of the opposite strand, it could be repaired primarily via the short-patch BER pathway. In this case, the cleavage efficiency of the AP site-containing DNA strand catalyzed by human apurinic/apyrimidinic endonuclease 1 (hAPE1) decreased under AP site excursion to the 3’-side relative to the lesion in the thymine methyl group under UVA- or ionizing irradiation is of lesions that can be removed from the DNA by different repair systems potentially enhance the risk of the appearance of mutations, since cluster may include a few types of lesions (5-foU) that are directly repaired by different repair systems depending on their nature. One of the major lesions arising in the cell as a product of the thymine methyl group under UVA- or ionizing irradiation is 5-formyluracil (5-foU) [2]. The presence of 5-foU in the DNA does not lead to replication block on both DNA strands; however high-fidelity DNA polymerases such as the Klenow fragment of DNA polymerase I, DNA polymerase alpha and gamma are able to incorporate any of the 4 dNTPs opposite 5-foU allowing transition and transversions [3]. The 5-foU residue is weakly mutagenic, with mutation frequencies in double-stranded vectors in the range of 0.01–0.04% [4]. One of the troublesome outcomes of the existence of 5-foU is the formyl group. First, it can interact with its surroundings, for example with the amino group of DNA binding proteins, and form covalent adducts via Schiff bases [5]. Second, the N-glycosyl bond of 5-formyluridine is less stable than that of 2-deoxyuridine even in physiological conditions, which results in the generation of an apurinic/apyrimidinic site (AP site) in the DNA chain that has no coding base, and goes to spontaneous or enzymatic cleaving single-strand break [6]. Additionally, the presence of AP site in DNA strand force the linear form of DNA to kink which may lead to changing the recognition of the damaged site by repair proteins [7]. Thus, if 5-foU in the DNA is located close to the AP site or its precursor thereby forming a cluster, it can lead to additional danger relative to the formation of single- or double-strand breaks or involving an alternative repair system.

Data exist concerning 5-foU repair in reconstituted systems and also in cell-free extracts from HeLa and MRC5 cells [8,9]. It was suggested that basically 5-foU is repaired by the base excision repair system (BER). Several DNA glycosylases from Escherichia coli and human cells are known that can remove this lesion from DNA with different efficiencies [10,11]. However, some facts demonstrate that 5-foU can be removed by the nucleotide excision repair system (NER). The group of Hanaoka observed that one of the main initiators of the NER process, XPC-HR23B, interacts with DNA duplexes containing all four base pairs coupled to 5-foU [12]. Furthermore, the region of the DNA with 5-foU is effectively excised from such DNA duplexes by cell-free NER extracts, whose efficiency depends on the instability of the base pair but in this case

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Table 1. DNA substrate constructs used in the present study.

| name      | sequence                                                                 | primer length (nts)* |
|-----------|---------------------------------------------------------------------------|----------------------|
| DNA1      | 5'-GACTACATTTCATCGTGGGCTTCATGTTGCGTcUfCAGACCTGGTTGATACCAG                | 17                   |
|           | 3'-CTGATGTAANAGTACGAACCAGAACCGGAAATGGCAGAGGAAGCAAG C QCTCGACCACCTATTGCG*|                      |
| DNA2      | 5'-GACTACATTTCATCGTGGGCTTCATGTTGCGTcUfCAGACCTGGTTGATACCAG                | 18                   |
|           | 3'-CTGATGTAANAGTACGAACCAGAACCGGAAATGGCAGAGGAAGCAAG O QCTCGACCACCTATTGCG*|                      |
| DNA3      | 5'-GACTACATTTCATCGTGGGCTTCATGTTGCGTcUfCAGACCTGGTTGATACCAG                | 19                   |
|           | 3'-CTGATGTAANAGTACGAACCAGAACCGGAAATGGCAGAGGAAGCAAG O QCTCGACCACCTATTGCG*|                      |
| DNA4      | 5'-GACTACATTTCATCGTGGGCTTCATGTTGCGTcUfCAGACCTGGTTGATACCAG                | 18                   |
|           | 3'-CTGATGTAANAGTACGAACCAGAACCGGAAATGGCAGAGGAAGCAAG O QCTCGACCACCTATTGCG*|                      |

* primer length – the length of the primed oligonucleotide created under hAPE1 activity, cUf – 5-formyl-2'-deoxyuridine, O – AP site, * – 32P-radioactive label.

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the point mutation is fixed. Another NER factor RPA interacts with ssDNA containing 5-foU [13].

Thus, in the situation when 5-foU and the AP site compose one cluster, the cell must coordinate the choosing of the correct repair system (BER or NER) as well as the order of lesion repair. In any case, the appropriate DNA polymerase can provide DNA synthesis using the damaged template. If this synthesis is proceeds in a frame of the BER process, the role of such DNA polymerase can play DNA polymerase beta or lambda [14].

In this study, we explored the repair opportunity of the AP site composed of the clustered lesion by the BER proteins. As DNA substrates, DNA duplexes were used containing 5-foU and uracil as AP site precursor in the different strands displaced from each other towards the 3’- or 5’-side, i.e. in the positive or negative orientation (Table 1). The lesions are said to be in the positive orientation when the opposing strand lesion is 3’- of the base opposite the reference lesion, and to be in the negative orientation when the lesion in the opposite strand is 5’- of the base opposite the reference lesion. The main repair activities investigated were the activity of human apurinic/apyrimidinic endonuclease 1 (hAPE1), DNA polymerase activity of DNA polymerases beta and lambda and their cooperation with endonuclease activity of hPCNA on the BER process was studied.

Figure 1. Efficiency (kcat/Km, 1/mM sec) of the DNA endonuclease cleavage of the AP-DNA strand by hAPE1. Empty and shaded columns designate the values obtained in the presence of Mg2+ and Mn2+ ions, respectively. The results are presented as the average value of four independent experiments. Standard error was estimated as 10%.

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Materials and Methods

Materials

Synthetic oligodeoxyribonucleotides were obtained from GenSet (Switzerland). Reagents for electrophoresis and the basic components of buffers were from Sigma (USA). (γ-32P)ATP (specific activity 5000 Ci/mmol) was from the Laboratory of Biotechnology (Institute of Chemical Biology and Fundamental Medicine, Novosibirsk, Russia). T4 polynucleotide kinase and T4 DNA ligase were from Novobio (Novosibirsk, Russia). Ultra pure dNTPs were from Promega (USA).

Proteins

Human recombinant DNA pol lambda and DNA pol beta were purified from E. coli BL21(DE3) RP cells as described in [15] and [16], respectively. hPCNA was purified according to [17]. Uracil-DNA glycosylase (UDG) was from E. coli, and hAPE1 and hFEN1 were purified and kindly provided by Svetlana Khodyreva (Institute of Chemical Biology and Fundamental Medicine Siberian Branch of the RAS, Novosibirsk, Russia).

XRCC1 purification

Human full-length XRCC1 with an N-terminal histidine tag was expressed in E.coli from the plasmid pET16BHX kindly supplied by J.P.Radicella (from Dr Keith W. Caldecott, University of Sussex, Brighton, UK) and purified as described in [18] with some modifications. In particular, E. coli BL21(DE3) Rosetta cells containing the plasmid were grown in LB medium to OD600 0.4 and induced with 0.5 mM IPTG for 3 h. After centrifugation, the cells were quick-frozen at -40°C. The pellets were thawed on ice and resuspended in 20 ml ice-cold sonication buffer containing 50 mM Na phosphate, pH 7.8, 300 mM NaCl, 1% NP-40, 1 mM β-mercaptoethanol (β-Me), 10% glycerol, 1 mM PMSF and 1 mM benzamidine (Bz), and sonicated on ice (10×15 s with 100 s cooling intervals). Cellular debris was removed by centrifugation (18000g, 45 min, 4°C). The protein was further purified on a Ni-sepharose fast flow His-tag affinity column (GE Healthcare, Sweden). Imidazole, pH 7.2 was added to the supernatant (23 ml) to 3 mM final concentration, and the mixture was loaded onto 3 ml Ni- resin and stirred on ice for 1 h. The flow-through was then collected. The column was washed with 3 column volumes (CV) of each buffer A (50 mM Na phosphate, pH 7.2, 300 mM NaCl, 0.1% NP-40, 3.5 mM β-Me, 1 mM PMSF and 1 mM Bz), containing 5, 40 and 50 mM imidazole and 10 ml fractions collected. The column was then washed with 3 CV of each buffer A, containing 200 and 300 mM imidazole, with 1.5 ml fractions
Figure 2. Influence of hXRCC1 on the endonuclease activity of hAPE1 (A), and on the incorporation of dNMP catalyzed by DNA polymerases beta (B) and lambda (C). The data are presented in logarithmic scale. Solid lines designate the yield of the reaction products obtained in the presence of Mg²⁺ ions; dotted lines designate the yield of the reaction products obtained in the presence of Mn²⁺ ions. The yield of the reaction products is the ratio of the reaction products over the initial amount of substrate expressed as a percentage. dGTP was used for the reactions with DNA1 and DNA3, and dATP – for the reactions with DNA2 and DNA4. The results are presented as the average value of four independent experiments. Standard error was estimated as 10%.

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centrifugal concentrators with a PES membrane 10 kDa. The final purity was $\approx 80\%$, as estimated by acrylamide gels stained with Coomassie blue R250. To test the hXRCC1 activity, its ability to form complexes with hAPE1 and DNA polymerase beta was assayed according to [18] with minor changes. In particular, 350 pmol hXRCC1, 114 pmol hAPE1 and 130 pmol DNA polymerase beta in 40 µl reaction mixture were used for the analysis. Buffer A contained 1 mM β-Me instead of DTT, and incubation time was increased to 1 h. Before electrophoresis all mixtures were incubated with an extra 5 mM DTT on ice overnight. It was observed that the recombinant protein is able to perform protein-protein interactions with hAPE1 and DNA polymerase beta (data presented in Fig. S1).

Oligonucleotide substrates
All modified oligonucleotides were synthesized by the phosphoramidite method on an automatic ABI 3400 DNA synthesizer (Applied Biosystems, USA) using the conditions recommended by the manufacturer, and double purified a) by electrophoresis in a 20% polyacrylamide/7 M urea gel and b) by RP-HPLC with an acetonitrile gradient in 0.1 M ammonium acetate (pH 7). HPLC purification was carried out on AKTA Purifier equipped with Jupiter C18 or C5 column (Phenomenex, size 4.6×250 mm, 5 µm), at 45°C, 1 ml/min with a UV-Vis detector. T, d(NzBz), dG(iBu), dC(Ac), 5-(1,2-diaceoxyethyl)-2′-deoxyuridine phosphoramidites and Universal UnyLinker CPG Support 500 Å were purchased from ChemGenes (USA). To generate the formyl group, 300 pmol of oligonucleotide were incubated with 500 nmol fresh NaIO₄ and 11.5 µmol NaOAc, pH 4.5, in 50 µl the reaction mixture for 2 h at room temperature. The reaction was stopped by adding 8 µl of 10% Na₂SO₃ at pH 7. All modified oligonucleotides were then purified by RP-HPLC using a Jupiter C5 column.

Table 2. Efficiency ($k_{cat}/K_m$, µM⁻¹·sec⁻¹) of dNMP incorporation by DNA polymerases beta and lambda via the short-patch BER pathway.

|        | pol β | pol λ |
|--------|-------|-------|
|        | Mg²⁺ | Mn²⁺ | Mg²⁺ | Mn²⁺ |
| DNA1/dGMP | 0.717 | 3.62 | 3.92 | 32.6 |
| DNA2/dAMP | 0.036 | 0.44 | 0.81 | 8.20 |
| DNA3/dGMP | 0.020 | 2.3 | 2.36 | 9.35 |
| DNA4/dAMP | 0.006 | 0.23 | 1.14 | 96.0 |

Note: the results are presented as the average value of three independent experiments. Standard error was estimated as 10%.

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Figure 3. Specificity of dNMP incorporation catalyzed by DNA polymerases beta and lambda in the presence of Mg²⁺ (A) or Mn²⁺ (B) ions using DNA1 (lanes 1–10), DNA2 (lanes 11–20) and DNA3 (lanes 21–30). Lane C, position of the 5’-(32P)-labeled 55 and 18 nt-long DNA; lanes 1, 11 and 21, initial DNA substrate after UDG treatment; lanes 2, 12 and 22, initial DNA substrate after sequential treatment by UDG and hAPE1; 18, 19 and 20 nt, lengths of the reaction products. The autoradiographs present the results from one of three independent experiments.
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mixture at room temperature for 1 h [19]. The reaction was terminated by adding 70 ml of 3 M LiClO4 and pure acetone to 1.5 ml. The oligonucleotide precipitate was generated at 240°C for 2 h, centrifuged, washed twice with 1 ml cooled acetone (4°C), air dried, dissolved in water to the required concentration and used for annealing.

Preparation of 5'-[32P]labeled primers

Radioactive label was incorporated into the 5'-end of the U-containing oligonucleotide using phage T4 polynucleotide kinase as described [20]. The reaction mixture (10 ml) contained 0.5 mM primer, 10 MBq [γ-32P]ATP, and 5 U T4 polynucleotide kinase. The reaction was performed at 37°C for 30 min, and then further overnight at 4°C. Nucleotide material was resolved by 20% denatured PAGE, visualized by autoradiography, and isolated by electroelution onto DE81-paper (Whatmann) in 50 mM Tris-borate buffer, pH 8.3. The product was eluted from DE81-paper by five portions (20 ml per portion) of 3 M LiClO4, and then 1.2 ml cooled acetone (4°C) was added to the eluate. The probe was incubated at –40°C for 2 h. The precipitate was collected by centrifugation, air dried, dissolved in water to the required concentration and used for annealing.

Annealing of DNA oligonucleotides

Two oligonucleotides in a 1:1.3 molar ratio where annealed in water in which 1 is the 5'-[32P]phosphorylated U-containing oligonucleotide and 1.3 is the 5-fO/T-containing oligonucleotide. The solution was heated at 97°C for 5 min and slowly cooled to 73°C, then incubated at 73°C for 15 min and slowly cooled to room temperature. The amount of duplex DNA substrate was controlled by native 12% polyacrylamide gel electrophoresis (PAGE) analysis PAAG. The oligonucleotide substrates used are presented in the Table 1.

AP-site was obtained using the glycosylase activity of uracil-DNA glycosylase directly before the following reactions. Mixtures contained 1 pmol of 5'-[32P]phosphorylated DNA substrate and 0.1 U of UDG in 10 μl water were incubated at 37°C for 30 min and used for the following reactions. Complete excision of uracil was confirmed by PAGE analysis in each case (specified on the figures).

**Table 3.** Efficiency (kcat/Km, μM⁻¹·sec⁻¹) of dNMPs incorporation by DNA polymerases beta and lambda via the long-patch BER pathway.

| polβ | polλ |
|------|------|
| DNA1 | 0.003 | 0.0084 | 0.0084 | 0.012 |
| DNA2 | 0.01 | 0.0003 | 0.0034 | 0.037 |
| DNA3 | 0.022 | 0.0164 | 0.0131 | 0.235 |
| DNA4 | 0.008 | 0.0072 | 0.0074 | 0.118 |

Note: the results are presented as the average value of three independent experiments. Standard error was estimated as 10%.

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Figure 4. Strand-displacement activity of DNA polymerases beta (A, B) and lambda (C, D) in the presence of Mg2+ (A, C) or Mn2+ (B, D) ions using DNA1 (lanes 1–6), DNA2 (lanes 7–12), DNA3 (lanes 13–18) and DNA4 (lanes 19–24). Lane C, position of the 5'-[32P]labeled 55 and 18 nt-long DNA; lanes 1, 7, 13 and 19 (upper panel), initial DNA substrate after UDG treatment; lanes 2, 8, 14 and 20 (upper panel), and 1, 7, 13 and 19 (bottom panel), initial DNA substrate after sequential treatment by UDG and hAPE1; 18, 19 and 20 nt, lengths of the reaction products. The final concentrations of dNTPs are indicated below the gels. The autoradiographs present the results from one of three independent experiments.

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All the following reactions were carried out in the presence of 1 mM MgCl₂ or MnCl₂. This concentration of bivalent ions was shown to be optimal for the TLS activity (i.e. translesion synthesis) of DNA polymerases beta and lambda in the range of 0.1–10 mM for MgCl₂ and 0.05–1.2 mM for MnCl₂ (data not shown).

#### Endonuclease activity of hAPE

Endonuclease activity of hAPE was investigated in reaction buffer (TDB buffer) containing 50 mM Tris-HCl (pH 7.5), 0.5 mM dithiothreitol, 0.25 mg/mL bovine serum albumin, and 1 mM MgCl₂ or MnCl₂ as follows. First, the kinetics of the endonuclease reaction by APE1 was studied. For this, the reaction mixtures (final volume 80 μl) including 10 nM 5’-[32P]phosphorylated DNA substrate, 10 nM hAPE1 and 1 mM MgCl₂ or MnCl₂ in TDB buffer were incubated at 37°C for 20 sec, 40 sec, 1, 1.5, 2, 3, 5, 10 and 15 min. The aliquots (10 μl) were collected at the indicated times, and the reaction products were stabilized at 4°C for 30 min after addition of NaBH₄ to a concentration of 20 mM. The reactions were terminated by adding the gel loading solution (90% formamide, 0.1% xylene cyanol and 0.1% bromophenol blue) and heated at 97°C for 5 min. The mixtures were resolved on a 20% polyacrylamide gel containing 7 M urea in TBE buffer [21]. The gels were dried and subjected to autoradiography and/or phosphorimaging for quantitation using the Molecular Imager Pro FX+ and “Quantity One” software (Bio-Rad, USA), and analyzed using OriginPro7.5 (Microlab Software, USA). Based on the results obtained, a 1.5 min interval was chosen as the reaction time. Second, the DNA cleavage efficiency by APE1 was estimated. All mixtures (final volume 10 μl) included 10 nM 5’-[32P]phosphorylated DNA substrate, 0.1–10 nM hAPE1 and 1 mM MgCl₂ or MnCl₂ in TDB buffer. All the reaction mixtures were incubated at 37°C for 1.5 min. The reaction products were stabilized by addition of NaBH₄ to 20 mM at 4°C for 30 min. The reactions were terminated and analyzed as outlined above. To estimate the kinetic parameters Kₘ and Vₘₐₓ, where the Vₘₐₓ value was measured as the amount of cleavage product in pmol generated per sec, and the Kₘ value was the DNA concentration needed to achieve a rate equal to Vₘₐₓ/2, the hyperbolic equation was used. The influence of hXRCC1 on the endonuclease activity of hAPE was studied as follows. All mixtures (final volume 10 μl) included 10 nM 5’-[32P]phosphorylated DNA substrate, 1 nM hAPE1 and 0.1–500 nM hXRCC1 and were incubated at 37°C for 1.5 min. The reaction products were stabilized by addition of NaBH₄ to 20 mM at 4°C for 30 min. The reactions were terminated and analyzed as outlined above. All experiments were repeated several times.

#### DNA pol assay

DNA synthesis catalyzed by DNA polymerases beta and lambda was performed in TDB buffer. First, all DNA substrates were treated by UDG (as mentioned above). Then all mixtures (final volume 10 μl) included 10 nM 5’-[32P]phosphorylated DNA substrate and 10 nM DNA polymerase (beta or lambda). The reactions were incubated at 37°C for 10 min. The reaction products were resolved on a 10% polyacrylamide gel containing 7 M urea in TBE buffer. The gels were dried and subjected to autoradiography.

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**Figure 5. Influence of hPCNA on long-patch BER activity of DNA polymerases beta and lambda using DNA1 (A), DNA2 (B), DNA3 (C) and DNA 4 (D).** Lane C, position of the 5’-[32P]labeled 55 and 18 nt-long DNA; lane 1, initial DNA substrate after UDG treatment; lanes 2 and 6, initial DNA substrate after sequential treatment by UDG and hAPE1; 18, 19 and 20 nt, lengths of the reaction products. The final concentrations of hPCNA are indicated below the gels. The autoradiographs present the results from one of three independent experiments.

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DNA substrate treated by UDG, 2 nM hAPE1 and 1 mM MgCl2 or MnCl2 in TDB buffer were incubated at 37°C for 5 min to complete cleavage of the AP site-containing DNA strand which was confirmed by PAGE analysis in all cases (specified in the figures). DNA polymerase beta or lambda (10 nM), and 50 μM dNTP (dGMP using DNA1, 3 and dAMP using DNA2, 4) at the reconstitution of the short-patch BER pathway, or 50 μM dNTPs and 50 nM hFEN1 at the reconstitution of the long-patch BER pathway were then added. All reactions were additionally incubated at 37°C for 15 min, then terminated and analyzed as outlined above. All experiments were reproduced several times.

Michaelis constants, Km, and maximum velocities, Vmax, of DNA synthesis catalyzed by DNA pols beta or lambda. Michaelis constants, Km, and maximum velocities, Vmax, of DNA synthesis catalyzed by DNA pols beta or lambda

Figure 6. AP site repair via short-patch (A, B and E) and long-patch (C, D and E) BER pathways. Step-by-step reconstruction using DNA1 (lanes 1–6 of A, B, C and D), DNA2 (lanes 7–12 of A, B, C and D), DNA3 (lanes 13–18 of A, B, C and D), and DNA4 (E). The experiments were performed in the presence of Mg2+ (A, C, and lanes 1–6 and 12–15 of E) or Mn2+ (B, D, and lanes 7–11 and 16–19 of E) ions. Lane C, position of the 5’-(32P)labeled 55 and 18 nt-long DNA; lanes 1, 7 and 13 (A, B, C and D) and 1 (E), initial DNA substrate after UDG treatment; lanes 2, 8 and 14 (A, B, C and D), and 2 and 7 (E), initial DNA substrate after sequential treatment with UDG and hAPE1; 18, 19, 20 and 55 nts, lengths of the reaction products. The autoradiographs present the results from one of three independent experiments.

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The influence of hXRCC1 on the incorporation of dNMP was studied as follows. All mixtures (final volume 10 μl) included 10 nM 5’-[32P]phosphorylated DNA substrate treated with UDG and APE1, 10 nM DNA polymerase beta or lambda, 1 mM MgCl2 or MnCl2, 50 μM dNTP (dGMP using DNA1, 3 and dAMP using DNA2, 4), and 0.1–500 nM hXRCC1. All reactions were incubated at 37°C for 15 min, then terminated and analyzed as outlined above. All experiments were reproduced several times.
were determined in two steps. In the first step, the kinetics of dNMP incorporation into different DNA substrates by the DNA polymerases beta and lambda were estimated. The reaction was performed in 90 \( \mu \)l of a mixture containing 10 nM DNA polymerases beta or lambda, 50 \( \mu \)M dNTP, 10 nM 5'-\[32P\]phosphorylated DNA substrate treated with UDG and hAPE1 (as mentioned above), and 1 mM MgCl\(_2\) or MnCl\(_2\) in TDB buffer at 37°C. Aliquots (10 \( \mu \)l) were removed after selected times in the range of 0–40 min. The reaction was terminated by adding the gel loading solution. The reaction products were separated as described above. In the next step the Michaelis constants, \( K_m \), and the maximum velocities, \( V_{max} \), of DNA synthesis catalyzed by DNA polymerase beta or lambda were determined by the variation of dNTP concentration from 0.5 to 50 \( \mu \)M of each dNTP, and the reaction products were analyzed as described above. The data were fitted according to the Michaelis–Menten kinetic equation [22]. The \( k_{cat} \) values were calculated as \( V_{max}/e_0 \), where \( e_0 \) is the total concentration of DNA polymerase in the reaction mixture.

The effect of hPCNA on the strand-displacement activity of DNA polys beta and lambda. The effect of hPCNA on the strand-displacement activity of DNA polys beta and lambda was studied as follows. The reaction mixtures (final volume 10 \( \mu \)l) contained 10 nM 5'-\[^{32}P\] labeled DNA substrates treated with UDG and hAPE1 (as mentioned above), 10 nM DNA polymerase beta or lambda, 5 \( \mu \)M each dNTP, 50 \( \mu \)M hFEN1, 50 or 100 nM hPCNA, and 1 mM MgCl\(_2\) or MnCl\(_2\) in TDB buffer. All reactions were incubated at 37°C for 15 min. The reaction products were analyzed as outlined above. All experiments were reproduced several times.

AP site repair via the short- and long-patch BER pathways. AP site repair via the short- and long-patch BER pathways was carried out as defined for the study of the DNA polymerase assay. Aliquots were removed before adding each protein. At the final step, 1 U of T4 DNA ligase was added to 10 \( \mu \)l reaction mixture and incubated at 37°C for 15 min. The reaction products were analyzed as outlined above. All experiments were reproduced several times.

Results

Endonuclease activity of hAPE1 on DNA substrates containing cluster lesions

The major enzyme that initiates the repair of AP sites during the BER process, is hAPE1 [23]. Here, we investigated the influence of the position of 5-foU relative to the AP site in the complementary DNA strand on the endonuclease activity of hAPE1 in the presence of Mg\(^{2+}\) or Mn\(^{2+}\) ions. To this end the kinetic parameters \( K_m \) and \( V_{max} \) were determined, and the efficiency of endonuclease cleavage of the AP-DNA strand catalyzed by hAPE1 calculated as the ratio of \( k_{cat} \) to \( K_m \) (Materials and Methods; Fig. 1). In the presence of Mg\(^{2+}\) ions the cleavage efficiency of the AP-containing DNA strand did not depend on the position of the 5-foU on the other strand. However, in the presence of Mn\(^{2+}\) ions the cleavage efficiency of AP-DNA strand decreased upon AP site excursion to the 3'-side relative to the lesion in the other DNA strand. It is noteworthy that in the last case, cleavage efficiency by hAPE1 was almost 2-fold higher using damaged substrate DNA1 rather than control DNA, DNA4. These results can be explained by the formation of a more productive intermediate complex including hAPE1, DNA and Me\(^{2+}\) and the simultaneous existence of the appropriate reciprocal orientation of the AP site and 5-foU:A pair in DNA1 in the presence of Mn\(^{2+}\) [24,25].

Influence of hXRCC1 on the endonuclease cleavage of AP-DNAs

Many studies have suggested that XRCC1 plays a central role due to its action as coordinator of the BER process [24]. It is known, that hXRCC1 physically interacts with hAPE1 and DNA polymerase beta, and modulates their endonuclease and polymerase activity [26,27]. Here, hXRCC1 was expressed in E.coli, and

Figure 7. BER pathways for repair of AP site introduced into the cluster damage with 5-foU in the opposite strand. Uf, 5-foU, O, AP site. doi:10.1371/journal.pone.0068576.g007
complex formation of the purified protein with recombinant hAPE1 and DNA polymerase beta was demonstrated (Fig S1). The influence of hXRCC1 on the endonuclease activity of hAPE1 was investigated using AP-DNAs containing 5-formyl-2'-deoxuryridine (Fig. 2, A); addition of hXRCC1 up to 500 nM did not affect AP-DNA cleavage.

dNMP incorporation catalyzed by DNA polymerases beta and lambda

To simulate the short-patch BER pathway, dNMP incorporation catalyzed by DNA polymerase beta or lambda into 5-foU-containing DNAs exactly after AP-DNA cleavage by hAPE1 was examined (Fig. 3). Clearly, DNA polymerase lambda is the more accurate enzyme during DNA synthesis across 5-foU. Independently from the reaction conditions, this enzyme could incorporate correct dNMP: dGMP as a complement to template C using DNA1 and DNA3, and dAMP as complement to the initial template T using DNA2 (Fig. 3). dGMP incorporation into the DNA strand from DNA2 in the presence of Mg²⁺ (∼10%) and Mn²⁺ (∼25%) ions could be explained, on one hand, by a template slippage mechanism as described for DNA polymerase lambda [28,29]. The most effective and accurate DNA synthesis catalyzed by DNA polymerase lambda was observed using DNA1 and DNA3 (Table 2, and Table S1).

DNA polymerase beta is an extremely error-prone enzyme that can catalyze the elongation of the 3'-end of the primer opposite 5-foU and also close to the lesion using any dNMP (Fig. 3). Most likely the incorporation of dCMP and dTMP is associated with the template slippage mechanism as described for DNA polymerase lambda [28,29]. The most effective and accurate DNA synthesis catalyzed by DNA polymerase lambda was observed using DNA1 as substrate (Table 2, and Table S1).

Influence of hXRCC1 on dNMP incorporation by DNA polymerases beta and lambda

Since hXRCC1 has been suggested to be a scaffold protein for the BER process, its effect on DNA synthesis catalyzed by DNA polymerase beta or lambda during the short-patch pathway was examined. To this end, the incorporation level of dNMP into DNA1 and DNA3, and dAMP to the reaction mixture. To confirm the existence of protein-protein and nucleic acid-protein interactions in replication and repair complexes. Several studies have reported physical interactions between DNA polymerases beta and lambda and hPCNA [33,34]. Evidence also exists of the influence of hPCNA on the efficiency and fidelity of translesion DNA synthesis catalyzed by DNA polymerases beta and lambda [35,36].

Influence of hPCNA on the efficiency of the long-patch BER pathway

One of the inherent components of the BER system is hPCNA. First, hPCNA is a processivity factor for replicative DNA polymerases. Additionally, it participates in the coordination of protein-protein and nucleic acid-protein interactions in replication and repair complexes. Several studies have reported physical interactions between DNA polymerases beta and lambda and hPCNA [33,34]. Evidence also exists of the influence of hPCNA on the efficiency and fidelity of translesion DNA synthesis catalyzed by DNA polymerases beta and lambda [35,36].

Here, the effect of hPCNA on the ability of DNA polymerases beta and lambda to perform DNA synthesis using 5-foU-containing templates was investigated during AP site repair via the long-patch BER pathway. The effect of the protein factor was estimated by the changes in quality and amount of reaction products obtained in the presence or absence of hPCNA (Fig. 5). As can be seen from the results presented, hPCNA did influence the yield of reaction products; however it increased DNA polymerase processivity, if the first dNMP was incorporated opposite the template 5-foU (Fig. 5B).

AP site repair via short- and long-patch BER pathways

A reconstructed system was used for repair of an AP site inside the cluster lesion. Upon modeling the short-patch BER pathway, we sequentially added hAPE1, DNA polymerase beta or lambda, and dNTP to the reaction mixture. To confirm the existence of the possible reaction product, i.e. nicked DNA with 3'-OH and 5'-phosphate groups, and to confirm its ligation, T4 DNA ligase was used (Fig. 6A, B). Reaction products whose lengths corresponded to the initial full-length DNA substrate were obtained in all cases. It should be noted that after ligation the amount of the DNA products elongated by the DNA polymerase was increased.

Similar experiments were performed to study possible repair of model DNA substrates via the long-patch BER pathway. In this case, hAPE1, DNA polymerase beta or lambda combined with hFEN1, dNTP and T4 DNA ligase were sequentially added to the reaction (Fig. 6C, D). The ligated product was obtained in all cases although in smaller amounts relative to the repair via the short-patch BER pathway. However, even in this case the products of DNA synthesis were presumably shifted to the full-length product. Additional evidence about the preferred BER pathway could be derived from analysis of the kinetics of dNTP incorporation and the influence of auxiliary factors specific for individual pathway (discussed later). It should be also noted that BER is workable via both repair pathways using a control DNA substrate containing the AP site only (Fig. 6E).
Discussion

Radiation, radiomimetic anticancer drugs or internal active oxygen species induce high levels of clustered DNA damage, in which several individual lesions are located within one or two turns of the DNA helix. Clustered DNA damage could lead to deleterious biological consequences such as point mutations or even cell death, if the repair process becomes erroneous or incomplete. If the AP site is one of the individual lesions, it can lead to very perilous damage resulting in point mutations or single-strand breaks. In this study we analyzed step-by-step the repair of AP sites consisting of double-stranded clustered DNA damage with 5-formyl-2'-deoxyuridine which is another “hot point” for the DNA repair system. We investigated the AP-DNA repair process via the short- and long-patch BER pathways.

The capacity of the endonuclease activity of hAPE1 to cleave the AP site located in one DNA strand close to the 5-foU of the other strand was first investigated. Cleavage efficiency depends on the position of the 5-foU in the opposite DNA strand: its efficiency decreased in the row from negative to positive orientation of the lesions. Current data are in agreement with the results obtained using clustered DNA damages with the other set of individual lesions [37]. However, in terms of the next step of the BER process such hAPE1’s “preferences”, these are much more severe for the cell since following DNA synthesis they could occur using damaged DNA templates. In this case, the TLS activity of the DNA polymerases beta and lambda seems promising. DNA synthesis during short-patch BER pathway catalyzed by DNA polymerase lambda was clearly more accurate than the process carried out by DNA polymerase beta. Upon reconstruction of the long-patch BER pathway, DNA polymerase beta demonstrated more processive DNA synthesis in comparison to DNA polymerase lambda. Obtained efficiencies of the dNTPs incorporation at the 3’-end of different 5-foU-containing DNA substrates lead to suggestions that the choice for DNA polymerase depends on the relative positions of 5-foU and initial AP site. Under the reaction conditions, hPCNA, which normally acts as co-factor of long-patch BER pathway, stimulates strand-displacement activity of both enzymes using DNA2 as substrate, in which the 5-foU is located opposite the first incorporated nucleotide. It should be noted that no effect of hXRCC1 on DNA polymerase and hAPE1 activities was detected. These observations were probably due to the incomplete reaction conditions required for protein-protein interaction within the complicated BER machine. Since during the process XRCC1 is strongly associated with Ligase III [18], the lack of hXRCC1 effect might be explained by the absence of this enzyme.

Thus, based on the results obtained we propose the following summary (Fig. 7). The AP site being part of the cluster damage with 5-foU can be repaired through the BER process. If the AP site is shifted relative to the 5-foU of the opposite strand it could be repaired primarily via the short-patch BER pathway. In this case, DNA synthesis catalyzed by DNA polymerase lambda could be more accurate, and, therefore, point mutations would not accumulate. If the AP site is located exactly opposite 5-foU it is more likely to switch the repair to the long-patch BER pathway. In this situation, hPCNA stimulates the strand-displacement activity of both DNA polymerases.

Supporting Information

Figure S1 Complex formation between recombinant hXRCC1 and either hAPE1 (A) or DNA polymerase beta (B). Initially, hXRCC1 was incubated with hAPE1 or DNA polymerase beta, then Ni-NTA agarose (Qiagen) was added to the mixture. After incubation Ni-NTA agarose was gently pelleted, the supernatant containing non-adsorbed material was removed (lanes indicated as FT, flow-through sample), and the Ni-NTA agarose beads washed several times with buffer containing 25 mM imidazole (lanes indicated as 25). Proteins were finally eluted from the Ni-NTA agarose beads by washing several times with buffer containing 250 mM imidazole (lanes indicated as 250). All samples were examined by Coomassie blue R250 staining of SDS-PAGE. Lanes M – molecular weight markers.

(TIF)

Table S1 Kinetic parameters ($K_m$, $V_{max}$) of dNMP incorporation by DNA polymerases beta and lambda via the short-patch BER pathway.

(DOC)

Table S2 Kinetic parameters ($K_m$, $V_{max}$) of dNMPs incorporation by DNA polymerases beta and lambda via the long-patch BER pathway.

(DOC)

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Author Contributions

Conceived and designed the experiments: EAB OIL. Performed the experiments: EAB. Analyzed the data: EAB TSO OIL. Contributed reagents/materials/analysis tools: IAV NAM TSZ TSO. Wrote the paper: EAB.

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