Human DNA polymerase θ possesses 5’-dRP lyase activity and functions in single-nucleotide base excision repair in vitro

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ABSTRACT

DNA polymerase θ (Pol θ) is a low-fidelity DNA polymerase that belongs to the family A polymerases and has been proposed to play a role in somatic hypermutation. Pol θ has the ability to conduct translesion DNA synthesis opposite an AP site or thymine glycol, and it was recently proposed to be involved in base excision repair (BER) of DNA damage. Here, we show that Pol θ has intrinsic 5’-deoxyribose phosphate (5’-dRP) lyase activity that is involved in single-nucleotide base excision DNA repair (SN-BER). Full-length human Pol θ is a ~300-kDa polypeptide, but we show here that the 98-kDa C-terminal region of Pol θ possesses both DNA polymerase activity and dRP lyase activity and is sufficient to carry out base excision repair in vitro. The 5’-dRP lyase activity is independent of the polymerase activity, in that a polymerase inactive mutant retained full 5’-dRP lyase activity. Domain mapping of the 98-kDa enzyme by limited proteolysis and NaBH4 cross-linking with a BER intermediate revealed that the dRP lyase active site resides in a 24-kDa domain of Pol θ. These results are consistent with a role of Pol θ in BER.

INTRODUCTION

DNA sequences homologous to the mus308 gene of Drosophila melanogaster were identified in human genome database, and the corresponding cDNA was cloned and mapped to chromosome 3q13.31. As the partial human cDNA encoded a 1762-amino-acid DNA polymerase with a molecular mass of 198 kDa, this new polymerase was designated DNA polymerase θ (Pol θ), and the corresponding vertebrate locus was named POLQ (1). Subsequently, a full-length cDNA for POLQ was cloned and shown to encode a 2592-amino-acid polypeptide with an amino-terminal helicase domain, a carboxy-terminal polymerase domain and an intervening spacer region (2). The mammalian POLQ and Drosophila mus308 gene products have similar domain arrangements. In addition to these orthologs, vertebrates have paralogs of the POLQ gene, HEL308 and POLN. The HEL308 gene product has homology to the helicase domain of POLQ but does not have a polymerase domain, whereas the POLN gene product has homology to the polymerase domain of POLQ, but does not contain a helicase domain (3,4).

DNA polymerase θ belongs to the family A polymerases (2,5) typified by Escherichia coli Pol I (6). Biochemical studies showed that purified full-length human Pol θ exhibited template-directed DNA polymerase activity on nicked double-stranded DNA and on a single-primed DNA template. Consistent with family A DNA polymerases, this activity was resistant to aphidicolin and inhibited by dideoxynucleotides. In addition, Pol θ exhibited a single-stranded DNA-dependent ATPase activity (2). Fidelity measurements of human Pol θ revealed that the polymerase generates single base pair substitutions at a rate 10- to 100-fold higher than other characterized family A DNA polymerases, making it one of the least faithful member of the family A DNA polymerases (7).

Pol θ is capable of conducting translesion DNA synthesis by inserting bases opposite an AP site or thymine glycol residue in the template strand. It also can extend an unpaired primer base opposite these lesions (8). Thus, Pol θ is sufficient for both the insertion and extension steps required for bypass of an abasic site. In contrast, Pol θ cannot insert bases opposite a cyclobutane pyrimidine dimer or a (6–4) photoproduct, but is able to extend primers after DNA polymerase ι had inserted a base opposite these UV-induced photoproducts (9). A role for Pol θ in somatic hypermutation (SHM) of immunoglobulin genes...
has been proposed where the excision of uracil by uracil-DNA glycosylase (UDG) during SHM generates an apurinic/apyrimidinic (AP) site that is repaired by either trans-lesion bypass or a base excision repair (BER) process involving several DNA polymerases and accessory proteins (10–14). Recently, Takeda and associates (15) suggested that Pol θ and Pol β are involved in BER of DNA damage inDT40 cells. They reported that POLQ−/− cells exhibited a significant increase in sensitivity to methyl methanesulfonate (MMS) or HmdUrd, but deletion of the genes for both POLQ and Pol β resulted in cells more sensitive to these base-damaging agents. Furthermore, subcellular localization studies demonstrated that Pol θ is rapidly recruited to sites of base damage that are induced by laser-delivered irradiation (15). These data along with extract-based in vitro repair assays suggested that Pol θ participates in BER in mammalian cells under certain conditions.

To understand the potential involvement of Pol θ in BER in mammalian cells, we cloned, overexpressed and purified to near homogeneity the C-terminal 98-kDa region of Pol θ corresponding to the core polymerase domain. We also prepared a polymerase-deficient variant of the enzyme by site-directed mutagenesis, and assessed activities associated with BER. Our results reveal that the Pol θ polymerase domain possesses both 5′-deoxyribose phosphate (dRP) lyase and DNA polymerase activities that function together during single-nucleotide BER in vitro. The 5′-dRP lyase activity was also fully active in the polymerase-deficient mutant. Our observation that Pol θ exhibits DNA polymerase and 5′-dRP lyase activities in vitro lends credence to the argument that Pol θ may have a role in BER in vivo.

MATERIALS AND METHODS

Materials

Synthetic oligodeoxyribonucleotides were from Oligos Etc, Inc. (Wilsonville, OR, USA) and The Midland Certified Reagent Co. (Midland, TX, USA). [γ-32P]dATP and [γ-32P]ddATP (3000 Ci/mmol), and [γ-32P]ATP (7000 Ci/mmol) were from GE Healthcare (Piscataway, NJ, USA) and Biomedicals (Irvine, CA, USA), respectively. Optikinase and terminal deoxynucleotidyl transferase were from USB Corp. (Cleveland, OH, USA) and Fermentas Inc. (Hanover, MD, USA), respectively. Protease inhibitor complete (EDTA-free) was from Roche Molecular Diagnostics (Pleasanton, CA, USA). Leupeptin, aprotinin and phenylmethylsulfonyl fluoride were from Calbiochem (La Jolla, CA, USA). Recombinant human Pol β was overexpressed and purified as described previously (16). Human AP endonuclease (APE), uracil-DNA glycosylase (UDG) with 84 amino acids deleted from the amino terminus and DNA ligase I were purified as described previously (17–19).

Overproduction and purification of the Pol θ polymerase domain

Based on the cDNA sequence of human Pol θ (1,2), a 2.7-kb cDNA clone was amplified for overexpression and production of a 98-kDa C-terminal fragment of human Pol θ using a 5′-BamH1 primer: 5′-TGC CAA TCA TGA TGG ATC CTC ATC CCT CTT ACC-3′ and a 3′-Sal1 primer: 5′-CTC TGT TCT TTG CAG TCG ACT GCA TCT GCA C-3′. The amplified DNA was digested with BamH1 and Sal1 restriction enzymes, and the correct DNA fragment was gel-purified and ligated into BamH1–Sal1 digested pQE80L (Qiagen). The resulting plasmid pQE80L-02.7 was transformed into E. coli BL21-CodonPlus-RP cells (Stratagene) for protein production.

Pol θ was purified from 1 to 21 of E. coli BL21-CodonPlus-RP cells transformed with pQE80L-02.7 following induction with 1.0 mM IPTG at a temperature of 30°C for 3.5 h. After harvesting by centrifugation and storage at –80°C, cells were resuspended in buffer (35 ml) containing 50 mM Tris–HCl, pH 7.5, 150 mM NaCl, 0.5 mM 2-mercaptoethanol, 5% glycerol and a cocktail of protease inhibitors. Cells were disrupted in a French Press chilled to 4°C with a constant pressure of 25 000 psi. The lysate was clarified by centrifugation at 12 000 g for 20 min, and the supernatant was saved for further purification. Imidazole was added to the cleared lysate to a final concentration of 20 mM before being applied to a 2-ml Ni–NTA column (Qiagen). The resin was washed with 50 column volumes of buffer containing 500 mM NaCl, 1% Triton X-100, 50 mM Tris–HCl, pH 7.5 and 20 mM imidazole, followed by several column volumes of a second wash buffer containing 500 mM NaCl, 0.005% NP-40, 50 mM Tris–HCl, pH 7.5 and 20 mM imidazole. Bound proteins were step-eluted with the second wash buffer also containing 0.25 M imidazole. The buffer was exchanged on a Fast Desalt HR 10/10 column (GE Healthcare) equilibrated in 25 mM potassium phosphate, pH 7.5, 10% (v/v) glycerol, 1 mM EDTA, 1 mM 2-mercaptoethanol, 0.005% (v/v) NP-40 and 75 mM KCl. The desalted protein fraction was applied to a 1.0 ml HR 5/5 Mono S column equilibrated in 25 mM potassium phosphate, pH 7.5, 10% (v/v) glycerol, 1 mM EDTA, 1 mM 2-mercaptoethanol and 75 mM KCl. Protein was eluted with a linear KCl gradient (0.075–0.5 M) in this buffer, and fractions containing Pol θ were flash frozen in liquid nitrogen in small aliquots.

Generation of polymerase variant

The DNA polymerase activity of Pol θ was inactivated by disrupting the ability of two essential carboxylate side chains to bind Mg2+ within motif C of the polymerase active site. Polymerase domain residues Asp829 and Glu830, corresponding to residues 2540 and 2541 in full-length Pol θ, were changed to Asn and Gln, respectively, using the QuikChange Site-Directed Mutagenesis Kit (Stratagene) and the following oligonucleotides: forward Pol′: 5′-CAT CCT TCA ACT CCA TAA TCA ACT CCT ATA TGA AGT G-3′ and reverse Pol′: 5′-CAC TTC ATA TAG GAG TTG ATT ATG GAG TGG AAG GAT G-3′. The resulting plasmid bearing the polymerase activity mutations was sequenced in its entirety to confirm the D829N and E830Q substitutions. E. coli BL21-CodonPlus-RP cells were transformed with this...
plasmid, pQE80L-θPol\(^{-}\), followed by induction and purification of the protein as described above.

**DNA polymerase assay**

Pol θ activity was measured with a gel-based oligonucleotide extension assay as described previously (20–23). An 18-mer primer (5'-TGA CCA TGT AAC AGA GAG-3') was gel purified, labeled at the 5'-end with \(\gamma\)-\(32\)P]ATP and annealed in a 1:1.4 ratio to a 36-mer template (3'-ACT GGT ACA TTG TCT CTC GCA CTC ACT CTC TCT TCT-5') in 10mM Tris-HCl, pH 7.5 by heating to 90°C for 5 min followed by slow cooling to room temperature. One picomole primer/template (\(32\)P-end labeled primer) was incubated with 5nM Pol θ in a reaction mixture (10\(\mu\)l) containing 25mM HEPES, pH 7.5, 2mM 2-mercaptoethanol, 0.1mM EDTA, 5mM MgCl\(_2\), 50\(\mu\)g/ml acetylated bovine serum albumin and 100\(\mu\)M each dNTP, as indicated. Following incubation at 37°C for 10 min, reactions were stopped on ice by addition of 10\(\mu\)l formamide gel-loading dye. Reaction products were separated by electrophoresis in a 15% polyacrylamide gel containing 8mM urea in 89mM Tris–HCl, 89mM boric acid, and 2mM EDTA, pH 8.8. Imaging and data analysis were performed with a Typhoon PhosphorImager and the ImageQuant software (GE HealthCare).

**Substrate preparation and dRP lyase activity assay**

Preparation of the dRP lyase substrate and the dRP lyase assay conditions were essentially as described previously (24). Briefly, the reaction mixture (20\(\mu\)l) contained 50mM HEPES, pH 7.5, 20mM KCl, 1mM EDTA and 50nM preincised \(32\)P-labeled AP site containing DNA. The reaction was initiated by adding the indicated amounts of Pol θ, mutant Pol θ or Pol β, followed by incubation at 37°C. Aliquots (9\(\mu\)l each) were transferred at the indicated time intervals into the tubes that contained 1\(\mu\)l of freshly prepared 200mM NaBH\(_4\). Reaction mixtures were shifted to 0–1°C (on ice) and incubation was continued for 30min. After heating to 75°C for 2 min, the reaction products were separated by electrophoresis in a 15% polyacrylamide gel containing 8M urea as described above. Imaging and data analysis were performed by PhosphorImager and ImageQuant software.

**5'-End labeling and substrate preparation for dRP lyase and NaBH\(_4\) cross-linking reactions**

Dephosphorylated 17-mer oligodeoxyribonucleotide (5'-UGTS-SGGATCCCGGGGTAC Biotin-3') containing a uracil residue at the 5'-end, a disulfide bond (S-S) 3 nt from the 5'-end, and biotin at the 3'-end was phosphorylated with Optiokinase and \(\gamma\)-\(32\)P]ATP. A 34-mer (5'-GTAC CCGGGATCCGTACCGCGCATCAGCTGCG-3') template was then annealed with 15-mer (5'-CTGCCAGCTGATGCCG-3') and 17-mer \(32\)P-labeled oligonucleotides by heating the solution at 90°C for 3 min and allowing the solution to slowly cool to 25°C. The \(32\)P-labeled duplex DNA was treated with human UDG to generate the \(32\)P-labeled deoxyribose sugar phosphate containing single nucleotide gapped substrate. The S-S bond was included in the substrate molecule to enable future studies on cross-linking within the dRP lyase active site.

**Covalent cross-linking of DNA to Pol θ**

A NaBH\(_4\) trapping technique was utilized to covalently cross-link Pol θ to DNA (25). The reaction mixture (10\(\mu\)l) contained 50mM HEPES, pH 7.4, 20mM KCl, 1mM EDTA, 200nM \(32\)P-labeled UDG-treated duplex DNA, 150nM wild-type Pol θ, mutant Pol θ or 20nM Pol β and 1mM NaBH\(_4\). Reactions were incubated first on ice for 60 min and then for 10 min at room temperature. After these incubations, 10\(\mu\)l SDS-PAGE gel-loading dye was added to each reaction, boiled for 5 min and protein–DNA cross-linked complexes were separated by electrophoresis in a 10% NuPAGE Bis-Tris gel with a MOPS running buffer system. Radioactive bands were visualized with a Typhoon PhosphorImager.

**Kinetic measurements of dRP lyase activity of Pol θ**

Kinetic analysis of the dRP lyase activity of Pol θ was performed with a \(32\)P-labeled 34-bp substrate that had been pretreated with UDG to generate a \(32\)P-labeled dRP flap within a single-nucleotide gap in the duplex DNA. The reaction mixture contained 50mM HEPES, pH 7.4, 20mM KCl, 1mM EDTA and 100nM \(32\)P-labeled UDG-treated duplex DNA. For the time course experiment, the reaction mixture (50\(\mu\)l) was assembled at 0–1°C in the above buffer. Reactions were initiated by adding 400nM Pol θ or the dilution buffer (control), as indicated in figure legends, and incubated at 37°C. Aliquots (9\(\mu\)l each) were withdrawn at different time intervals and transferred to 0–1°C to stop the reaction. DNA products were stabilized by addition of 20mM NaBH\(_4\) and incubated for 30 min on ice. Then, an equal volume of gel-loading buffer was added, and the reaction mixture was incubated at 75°C for 2 min. The reaction products were separated by electrophoresis in a 15% polyacrylamide TBE-Urea gel (Invitrogen, Pre-cast gel) for 30 min at constant voltage (200 V). To quantify the reaction products, gels were scanned on a PhosphorImager and the data were analyzed as above. Reaction rates were determined by plotting the amount of substrate released as a function of time, and data were fitted to the appropriate equation by nonlinear least squares methods. To examine the influence of protein concentrations on dRP removal, the reactions were assembled on ice, as above, and initiated by adding appropriate dilutions of Pol θ, as indicated in the figure legends. Reaction mixtures were incubated at 37°C for 10 min and processed as above.

**In vitro SN-BER**

Repair assay was performed in a final reaction volume of 30\(\mu\)l. A 35-bp oligonucleotide duplex DNA (250 nM) containing uracil at position 15 was incubated in a BER reaction mixture that contained 50mM HEPES, pH 7.5, 0.5mM EDTA, 2mM DTT, 20mM KCl, 4mM ATP, 5mM phosphocreatine, 100\(\mu\)g/ml phosphocreatine kinase, 0.5mM NAD, 15nM UDG, 15nM APE, 200nM DNA ligase I and 400nM or 800nM purified Pol θ, as indicated. Repair reactions were initiated...
by the addition of 10 mM MgCl$_2$ and 2.2 mM $[^{32}\text{P}]$dCTP (specific activity, $1 \times 10^6$ dpm/pmol), followed by incubation at 37°C. Aliquots (9 μl) were withdrawn at the indicated time intervals. Reactions were terminated by the addition of an equal volume (9 μl) of DNA gel-loading buffer. After incubation at 75°C for 2 min, the reaction products were separated by electrophoresis in a 15% polyacrylamide gel containing 8 M urea, and the data were analyzed as above.

**Limited proteolysis and amino-terminal sequencing of Pol 0**

The purified human Pol 0 98-kDa polymerase domain (66 μg) was subjected to limited proteolysis by mixing with trypsin (0.66 μg) at a 1:100 weight ratio (trypsin:Pol 0) in 100 mM Tris–HCl, pH 8.0 and then incubating the solution at 25°C. The final reaction mixture volume was 90 μl. Aliquots (20 μl each) were withdrawn at 5-, 15-, 30- and 60-min time intervals. A portion of each sample (18 μl) was mixed immediately with 10 μl SDS–PAGE gel-loading buffer, boiled for 5 min and separated by electrophoresis in a 12% NuPAGE Bis–Tris gel with a MOPS running buffer system. Proteins were electrophoretically transferred onto an Immun-Blot PVDF membrane (7 × 8.4 cm) (Bio-Rad) using a transfer buffer that contained 10 mM CAPS [3-(cyclohexylamino)-1-propanesulfonic acid] and 10% methanol, pH 11.0 at 50 V (7 V/cm) for 1 h. The membrane was stained briefly with 0.2% (w/v) Coomassie blue R-250 in 45% methanol and 10% acetic acid and destained with 90% methanol and 10% acetic acid. The membrane was air-dried, and protein bands were cut with a scalpel. Amino-terminal sequencing was performed using a Model 492 Procise sequencing system (Applied Biosystems) at Wake Forest University, Winston-Salem, NC. The remaining portion of the trypsin digested sample (2 μl) was subjected to NaBH$_4$ cross-linking to a 5'-end labeled dRP lyase DNA substrate, as described above. The resulting gel was subjected to phosphorimaging, and then the same gel was silver-stained for protein detection.

**RESULTS**

**Production of Pol 0 polymerase domain**

To facilitate the high-yield production of recombinant human Pol 0 in *E. coli*, we expressed several nested POL0 cDNA clones and screened them to find the largest fragment that retained DNA polymerase activity. From the full-length human Pol 0 cDNA (1,2, Copeland, unpublished data), we amplified 2.7-, 2.1-, 1.8- and 1.3-kbp cDNA fragments containing the DNA polymerase domain. Expression of the 2.7-kbp cDNA fragment, corresponding to the 98-kDa polymerase domain at the C-terminus of Pol 0 (amino acids 1712–2590), produced soluble protein with the highest yield (data not shown). A (His)$_6$ affinity tag was added to the N-terminus by transferring this fragment to the pQE80L expression plasmid, and protein was overproduced in the *E. coli* host BL21-CodonPlus-RP cells. As described in ‘Materials and Methods’ section, recombinant protein was purified to apparent homogeneity by sequential passage over a Ni–NTA affinity column, a buffer exchange column, and a Mono S cation exchange column (Figure 1A, lane 2). As a control for analysis of the polymerase activity, a mutant variant of the Pol 0 polymerase domain was constructed that abolished DNA polymerase activity. This was achieved by altering critical carboxylate residues that bind active site Mg$^{2+}$ ions. Similar alterations in other DNA polymerases are known to abolish DNA polymerase catalytic activity without dramatically affecting the overall structure of the protein (26). Asp829 and Glu830 (corresponding to residues 2540 and 2541 in the full-length Pol

![Figure 1](image-url)
(98-kDa peptide), mutant Pol β or Pol β. The results demonstrated that both wild-type Pol θ and the mutant Pol θ could release the dRP group from the substrate (Figure 2B), and the activities appeared to be similar. The apparent rates of dRP removal of Pol θ and Pol β were 0.016/min and 0.64/min, respectively. Under the same reaction conditions, the rate of dRP lyase activity of Pol θ appeared to be about 40-fold lower than that of Pol β (Figure 2B).

### Kinetics and mechanism of dRP lyase

To determine kinetic parameters of 5′-dRP group removal by Pol θ, we prepared a 34-bp duplex DNA that contained uracil at position 16 and a nick between positions 15 and 16, by annealing both a 15-mer and a 5′-end 32P-labeled uracil-containing 17-mer to the 34-mer complementary DNA strand. The resulting 32P-labeled nicked DNA was pretreated with UDG to generate the 5′-dRP-containing single-nucleotide gapped substrate; the DNA substrate, thus prepared, contained a 32P-labeled dRP flap in a single-nucleotide gap (Figure 3A, 16-mer + 32P-dRP). The rate of dRP removal by Pol θ was determined under single turnover conditions (i.e. enzyme/ DNA = 4). The

#### Table 1. Steady-state kinetic parameters for polymerase activities of full-length and 98-kDa fragment of Pol θ

| Enzyme                  | \(K_{\text{m}}\) μM | \(k_{\text{cat}}\) min\(^{-1}\) | \(k_{\text{cat}}/K_{\text{m}}\) μM\(^{-1}\) min\(^{-1}\) | References |
|-------------------------|---------------------|----------------------|----------------------|------------|
| Full-length 290-kDa Pol θ | 7.0                 | 0.67                 | 0.096                | (8)        |
| Full-length 290-kDa Pol θ | 9.0                 | 0.42                 | 0.047                |            |
| 98-kDa fragment Pol θ   | 9.7                 | 0.7                  | 0.072                | This study |

These measurements were made by a single nucleotide-primer extension assay.

![Figure 2](image) dRP lyase activity of wild-type and mutant Pol θ. (A) A schematic representation of the dRP lyase substrate (18-mer + 32P-dRP) generated by pretreatment of 32P-labeled 34-bp DNA with UDG and APE and the expected products 32P-labeled 18-mer product formed as a result of Pol θ (sample) or Pol β (positive control) catalyzed excision of 5′-terminal dRP. (B) Preincised DNA substrate (50 nM) was incubated either with no enzyme (lane 1), 40 nM wild-type Pol θ (lane 2), 40 nM mutant Pol θ (lane 3), or 5 nM Pol β (lane 4). Aliquots were withdrawn at the indicated time intervals, and the DNA products were stabilized and analyzed as described under Materials and Methods section. A photograph of the phosphorimage, illustrating the reaction products, is shown. Reaction products were quantified using ImageQuant software, and the fraction of the substrate converted into product was indicated as a percentage under each lane. The apparent rates for the dRP lyase activity of Pol θ and Pol β were 0.016 and 0.64/min, respectively. The positions of substrate and the product are indicated.
results revealed that Pol θ cleaved 5’ dRP in a time-dependent manner as monitored by disappearance of the substrate (Figure 3B), with an observed rate of dRP removal of ~0.14/min (Figure 3C). In other experiments where substrate was in excess, removal of the 5’-dRP group from the substrate was dependent on the enzyme concentration (Figure 4A and B).

To confirm that the 5’-dRP lyase activity of Pol θ was intrinsic to the 98-kDa peptide, we utilized the NaBH₄ cross-linking technique (25). This cross-linking technique relies on the ability of the C1’-aldehyde group of the 5’-deoxyribose in the substrate to react with a lysine γ-amine of a protein to form a β-elimination intermediate (Schiff base) protein–DNA complex; this Schiff base is recovered as a covalent bond by NaBH₄ trapping. We had previously shown that DNA polymerases, such as Pol β, Pol γ and Pol ι, catalyze dRP removal from the BER intermediate via β-elimination, and the Schiff base intermediate between these polymerases and DNA can be trapped by NaBH₄ reduction (27). As demonstrated in Figure 5, both wild-type Pol θ and the polymerase-deficient mutant Pol θ could form covalent protein–DNA products with the BER intermediate substrate upon treatment with NaBH₄. This indicates that the Pol θ polymerase domain possesses functional 5’-dRP lyase activity.

**Domain mapping of the 98-kDa peptide of Pol θ**

To probe domain organization of the Pol θ 98-kDa polymerase domain and to localize the 5’-dRP lyase active site of Pol θ, we subjected the purified protein to controlled proteolysis with trypsin, and then we identified the digestion products by amino-terminal sequencing. Samples from the incubation with trypsin were withdrawn at different time intervals. A portion of each sample was separated by SDS–PAGE, transferred onto a PVDF membrane and subjected to amino-terminal sequencing as described under ‘Materials and Methods’ section (Figure 6). Digestion for 5 min completely degraded the 98-kDa enzyme and produced two major fragments with molecular masses of about 33 and 24 kDa; several minor polypeptides of higher and lower molecular masses also were produced. With longer incubations of 30 and 60 min, four fragments persisted and were designated as fragments II, III, IV and V, respectively (Figure 6A). The 24-kDa
fragment appeared to be further digested to a smaller molecular mass polypeptide resulting in a doublet in the gel, whereas the 20-kDa polypeptide and some high-molecular-mass polypeptides almost disappeared (Figure 6A). Amino-terminal sequence analysis of several of these protease-resistant fragments (denoted as I–VI in Figure 6A) revealed that the 33-, 24-, 23-, 14- and 9-kDa polypeptides (II to VI) begin with Phe124, Cys 545, Gly 557, Leu423 and Asn 41, respectively (Figure 6B). These results indicated that the 23-kDa (IV) polypeptide resulted from removal of 12 amino acids from the amino terminus of the 24-kDa (III) polypeptide. Fragment I, with a molecular mass of 42 kDa, failed to exhibit reasonable sequence data due to low abundance. The domains of the 98-kDa Pol θ were aligned (Figure 6C) based on the molecular masses of these protease-resistant fragments and their amino-terminal sequences.

Localization of the dRP lyase active site

To localize the dRP lyase active site within the 98-kDa Pol θ, portions of the tryptic digests of Pol θ (see above) were incubated with the [32P]-labeled dRP lyase substrate and subjected to NaBH4 cross-linking as in Figure 5. This protocol was used to assess the capacity of the tryptic fragments for cross-linking and if successful to improve the yield of cross-linked material for sequencing. Reaction mixtures that were digested with trypsin for 5 min or 15 min produced major 33-kDa and 24-kDa fragments (Figure 6A), but only the 24-kDa fragment was radiolabeled following cross-linking (Figure 6D). Furthermore, longer incubations of 30 min and 60 min revealed that the radiolabeled 23- and 24-kDa peptides persisted and appeared as a doublet in the gel (Figure 6D), similar to the doublet observed upon staining with Coomassie blue or silver (Figure 6A and E, respectively). These observations suggested that even prolonged incubation with trypsin only partially converted the 24-kDa peptide to the 23-kDa peptide, and that both fragments contain the Schiff base dRP lyase nucleophile, i.e. dRP lyase active site. Taken together, these results indicate that the 24-kDa tryptic fragment is a domain of Pol θ that contains the dRP lyase active site and retains the ability to form a covalent DNA–protein complex with a BER intermediate (Figure 6D).

In vitro base excision repair activity

Since the 98-kDa Pol θ was able to cleave the 5′-dRP flap from a BER intermediate and also retained DNA polymerase activity, we asked whether the 98-kDa enzyme was capable of supporting BER in vitro. We assembled repair reaction mixtures containing a 35-bp DNA substrate with uracil opposite guanine and purified human enzymes, including UDG, AP endonuclease, DNA ligase I and the MRE11-RAD50-NSB1 nuclease complex rather than by the incorporation of [α-32P]dCMP in place of dUMP into a ligated 35-bp DNA product as a function of incubation time and concentration of Pol θ (Figure 7). In vitro repair activity was dependent on both enzyme concentration and incubation time (Figure 7A and B), confirming that human Pol θ catalyzed sufficient dRP removal and 1-nt gap filling to support single-nucleotide BER.

DISCUSSION

Abasic sites in DNA are common intermediates in several important DNA transactions within eukaryotic cells, including SHM, translesion DNA synthesis and BER. For example, SHM of immunoglobulin variable region genes is triggered by the action of activation-induced cytidine deaminase (28), which deaminates deoxycytosine to deoxyuracil. The abasic sites could be processed by AP endonuclease, and Pol β in an error-free pathway, but such a repair system would not be beneficial for SHM. Several lines of evidence support the idea that faithful repair mediated by Pol β is suppressed during SHM. First, using a mouse model Rajeswky and his associates (29) demonstrated that Pol β is not required in SHM. Second, we recently showed that the expression of Pol β in the SHM-proficient human BL2 cell line was strongly down regulated (30). Third, Maizels and associates (31,32) reported that during immunoglobulin gene diversification in B cells, AP sites were preferentially processed by the MER11-RAD50-NSB1 nuclease complex rather than by AP endonuclease.
Although Pol  is a family A DNA polymerase, it exhibits unusual properties that permit efficient bypass of abasic sites. In fact, Pol  is able to synthesize through these damaged sites with an efficiency that is comparable to that displayed on a corresponding undamaged template (8). The unique ability of Pol  to insert and extend mispaired nucleotides opposite AP sites suggests an important role for Pol  in SHM consistent with the observation that

Figure 6. Limited proteolysis of Pol  with trypsin. (A) Purified 98-kDa Pol  (66 μg) was digested at 25°C with trypsin (0.66 μg) at a weight ratio (trypsin:Pol ) of 1:100 in 100 mM Tris–HCl, pH 8.0. Aliquots were withdrawn at 5-, 15-, 30- and 60-min time intervals, as indicated at the ‘top’ of the photograph. A portion of each digested sample was mixed with 10 μl SDS–PAGE gel-loading buffer and resolved on a 12% NuPAGE Bis–Tris gel. The proteins were transferred onto a PVDF membrane for amino-terminal sequencing, as described under Materials and Methods section. The positions of Pol , tryptic peptides and protein markers are indicated. (B) Amino-terminal sequencing was performed using the Procise sequencing system, Model 492 (Applied Biosystems). The amino-terminal sequences of peptides I–VI are shown. (C) The domain organization of 98-kDa Pol  is depicted. (D) Portions of the trypsin digested samples from (A) were reacted with 5'-end-labeled dRP lyase DNA substrate and subjected to NaBH₄ cross-linking as in Figure 5. The gel was scanned on a PhosphorImager. The positions of cross-linked Pol , 23- and 24-kDa peptides are indicated. (E) The gel in (D) was stained with silver to detect protein fragments. The positions of Pol , 23- and 24-kDa peptides are indicated.
nucleophile for the dRP lyase activity. In investigating roles for the N-terminal 2/3 of the enzyme, it will be of interest to determine how the full-length enzyme influences overall BER efficiency in vivo. Similarly, Pol 0 is highly expressed in germinal center B cells, where SHM and class switch recombination occur (33). Identification of protein partners or cofactors that interact with full-length Pol 0 may help to assign roles for Pol 0 in these processes.

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