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Novel thermostable Y-family polymerases: applications for the PCR amplification of damaged or ancient DNAs

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For many years, Taq polymerase has served as the stalwart enzyme in the PCR amplification of DNA. However, a major limitation of Taq is its inability to amplify damaged DNA, thereby restricting its usefulness in forensic applications. In contrast, Y-family DNA polymerases, such as Dpo4 from Sulfolobus solfataricus, can traverse a wide variety of DNA lesions. Here, we report the identification and characterization of five novel thermostable Dpo4-like enzymes from Acidianus infernus, Sulfolobus shibatae, Sulfolobus tengchongensis, Stygiolobus azoricus and Sulfurisphaera ohwakuensis, as well as two recombinant chimeras that have enhanced enzymatic properties compared with the naturally occurring polymerases. The Dpo4-like polymerases are moderately processive, can substitute for Taq in PCR and can bypass DNA lesions that normally block Taq. Such properties make the Dpo4-like enzymes ideally suited for the PCR amplification of damaged DNA samples. Indeed, by using a blend of Taq and Dpo4-like enzymes, we obtained a PCR amplicon from ultraviolet-irradiated DNA that was largely unamplifiable with Taq alone. The inclusion of thermostable Dpo4-like polymerases in PCRs, therefore, augments the recovery and analysis of lesion-containing DNA samples, such as those commonly found in forensic or ancient DNA molecular applications.

INTRODUCTION

The Y-family of DNA polymerases consists of phylogenetically separated UmuC, DinB, Rev1 and Rad30 subfamilies (1). UmuC-like proteins are only found in prokaryotes, while Rev1 and Rad30 (polη and poli) orthologs exist exclusively in eukaryotes. DinB-like proteins are, however, found in all three kingdoms of life. In prokaryotes, DinB-like proteins are typified by Escherichia coli polIV; in eukaryotes, by polk; and in archaea, by Dbh and Dpo4. The Dbh (DinB homolog) protein was originally identified from Sulfolobus acidocaldarius using a degenerate PCR approach designed to obtain UmuC homologs (2), while Dpo4 (DNA polymerase IV) was identified by searching the completed genome of Sulfolobus solfataricus (3). Both Dbh and Dpo4 have been characterized biochemically and have been shown to be thermostable Y-family DNA polymerases. While Dbh exhibits poor processivity and poor lesion bypass activity (4,5), Dpo4 is significantly more robust on both undamaged and damaged templates (3,6). The crystal structures of both Dbh and Dpo4 have been solved, revealing that they are topologically similar to classical polymerases in that they resemble a right hand and possess ‘fingers’, ‘palm’ and ‘thumb’ sub-domains (7–9). However, they also possess an additional domain that has been termed the ‘little finger’ (LF) (9). The

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LF domain holds the polymerase on to the primer-template and studies of Dpo4/Dbh chimeras in which the LF domains were interchanged, revealed that the LF domain is the major factor determining the overall processivity of the enzyme (6).

Similar to many Y-family polymerases, Dpo4 is capable of replicating past several types of DNA lesions, including abasic sites (3,10), a cis–syn cyclobutane pyrimidine dimer (CPD) (3,11), and to a lesser degree pyrimidine 6-4 pyrimidone (6-4PP) at a dithymine site (3), cis-Platin (cis-Pt) guanine adducts (3) and N-acetyl-2-aminofluorene (AAF) guanine adducts (3). Indeed, crystal structures of Dpo4 in the process of bypassing an abasic site (12), a CPD (13) and a Benzo[a]pyrene-adenine lesion (14) have served as a paradigm for lesion bypass by the Y-family polymerases and have provided insights into the molecular mechanisms of translesion replication.

In contrast to Y-family polymerases, polymerases that belong to the A- or B-family possess a smaller and constricted active site that cannot accommodate bulky or non-coding adducts. As a consequence, such lesions are often kinetic blocks to continuing replication. Indeed, a major limitation in forensic science or the analysis of ancient DNAs is the ability of conventional PCR enzymes, such as Taq, to amplify damaged DNA (15,16). One way to potentially circumvent this problem would be to utilize a Y-family polymerase to replicate the damaged DNA template, so as to generate one or more lesion-free copies of the template that could be subsequently amplified by a higher fidelity PCR enzyme. In the first steps toward achieving this goal, we report here a quick and simple strategy for the identification of thermostable Y-family polymerases. We have overproduced, purified and characterized five novel Y-family polymerases, as well as two recombinant chimeras with enhanced enzymatic properties, and we provide ‘proof of principle’ that the inclusion of a thermostable Y-family DNA polymerase in the PCR amplification of damaged DNA facilitates the recovery of amplicons previously unobtainable with Taq polymerase alone.

**MATERIALS AND METHODS**

**Identification and cloning of novel dpo4-like genes**

Based on protein alignments of *S. solfataricus* Dpo4, *S. acidocaldarius* Dbh and a hypothetical Dpo4-like protein identified within the genome of *Sulfolobus tokodaii*, the degenerate PCR primers, dpo4_FRIDegen1 and dpo4_RXhDegen1, were designed and used to amplify an ~300 bp region of dpo4-like genes from the genomes of *Acidaminis infernus* (Ain) (DSM 3191), *Sulfolobus shibatae* (Ssh) (DSM 5389), *Sulfolobus tengchongensis* (Ste) (17), *Stygiolobus azoricus* (Saz) (DSM 6296) and *Sulfurisphaera ohwakuensis* (Soh) (DSM 6296) and *S. tengchongensis* in RW382, a *D. acidorubescens* strain (18). The sequence of dpo4_FRIDegen1 is 5’-GCC GAA TTC AAT AAT GTA CGG AAT ACT CTC T-3’ and is based on the conserved amino acid sequence (FH)(VI)D/1(FY)A. The sequence of dpo4_RXhDegen1 is 5’-GCC GCT CGA GTC TAD RWA IGC YTC RTT IAY ISW IRY-3’ and is based on the conserved amino acid sequence (AVI)(IV)S/DEA(YF)(LY)(ID)(DE). The PCR products were ‘TA’ cloned into pGEM-T (Promega) and sequenced. All subsequent PCR products were also cloned into pGEM-T except where noted. Additional regions of the dpo4-like genes were amplified using the degenerate PCR primers, dpo4_RXhDegen5 (5’-GCC GCT CGA GYT TRT WRT AIS HYT CIT YDA TIG C-3’ [AI(NED)E(A5)Y(NY)K]) and dpo4_RXhDegen6 (5’-GCC GCT CGA GRT AIG CIB KIT CIW TIS WIA TIC C-3’ [GIS(K)(DE )(NTR)AY]), in conjunction with gene-specific primers (A1_FRI276 (5’-CCG GAA TTC AAT AAT GTA CGG AAT ACT CTC T-3’), Sa_FRI267 (5’-CCG GAA TTC TTC TAA TAG TAT GAA CGG A-3’), So_FRI283 (5’-CCG GAA TTC CAT AAT CAG TAG CTA TTA GG-3’) and Sa_FRI235 (5’-CCG GAA TTC GCC CAT GAG AAA GAA AGT GT-3’). The sequence of the 5’ and 3’ ends of the *A. infernus* and *S. shibatae* genes were determined by inverse PCR using the primer pairs Sa_FRI142 (5’-CCG GAA TTC GCT ACA GGC AAT TAC GAC G-3’) and Sa_RXh74 (5’-GCC GCT CGA GTC AAT GCA GAC AAT TCG C-3’) and the primer pairs Sa_RXh138 (5’-CCG GAA TTC GTG ACT GCC AAG TAC TAC-3’) and Sa_RXh96 (5’-GCC GCT CGA GTC CTA TCC TAA CTA CCA-3’). The sequence of the 5’ and 3’ ends of *A. infernus*, *S. ohwakensis* and *S. shibatae* genes were isolated by performing PCR utilizing degenerate primers designed to the conserved upstream ribokinase gene and conserved downstream hypothetical gene; ribo_FRIDegen1 (5’-CCG GAA TTC GAY ACI ACI GCI GCI GGI GAY-3’ [DTTAGGD]) and hypo_RXhDegen1a (5’-GCC GCT CGA GTA YGA YGT IGA RGG IGG3’ [YEDVEGG]) in combination with gene-specific primers. The gene-specific primers used were Ai_FRI1772 (5’-CCG GAA TTC ACA TCA GGA GAT ATT AAA GAG A-3’), Ai_RXh43 (5’-GCC GCT CGA GAG TTC TGG GTT AAG GAC TCC-3’), Sa_FRI762 (5’-CGG GAA TTC ATT ACC GTA TAA TAC GAG AAA G-3’), So_FRI783 (5’-CCG GAA TTC AAA GTG ATA GAA CCT TAT CTG-3’), So_RXh76 (5’-GCC GCT CGA GCA GAA TAA ACA CAG ACT ATA AG-3’) and Sa_FRI696 (5’-CCG GAA TTC AGA GAT GAG TAT AAC GAA CCT-3’). Isolation of the 5’ and 3’ ends of the *S. tengchongensis* dpo4-like genes was accomplished by performing PCR using the degenerate PCR primer pairs ribo_FRIDegen1/dpo4_RXhDegen1 and dpo4_FRIDegen4 (5’-CCG GAA TTC RYI WSI RTI GAY GAV GCI TWY HTI GA-3’ [AVI](IV)DEA(YF)(LY)(IDL)(DE)) and hypo_RXhDegen1a (5’-GCC GCT CGA GTA YGA YGT IGA RGG IGG3’ [YEDVEGG]) and the degenerate PCR primers dpo4_FRIDegen1 and dpo4_RXhDegen4 (5’-CCG GAA TTC RYI WSI RTI GAY GAV GCI TWY HTI GA-3’ [AVI](IV)DEA(YF)(LY)(IDL)(DE)). The full-length nucleotide sequence of the polIV-like genes can be located in the following GenBank accession numbers: DQ124670 (A. infernus); DQ124673 (S. shibatae); DQ124669 (S. tengchongensis); DQ124671 (S. azoricus); and DQ124672 (S. ohwakensis).

The full-length *dpo4*-like genes were then PCR amplified using gene-specific primers and cloned or sub-cloned into pET22b+ (Novagen) from Ndel or FauI to BamHI for expression in *E. coli* strain BL21 (DE3) (Novagen) (18). The primers sets are Ai_FNde10 (5’-TTA ATG TTA AAT GCA TAT GAT TGT AC-3’) (the initiation ATG is italicized)/Ai_RBam1088 (5’-GTA CGG TGA TAT GCA GCC TCC TCC-3’), Sa_FNde2 (5’-TTA CTA CCC TAC TCC ACG CCG CCG TCT GAT ATG ATT G-3’) and Sa_RBam1078 (5’-TAT CGA AGT ATG GAT CCC TAA ATT TC-3’), So_FNde5 (5’-AGC CTA AGG TAA AAT GAA TCT CTA ATT TAC-3’) and Sa_RBam1078 (5’-CGA AGA ATG CCT ACT AAA GTG-3’), Sa_FNde5 (5’-GAA ACC AAA TGT TAT ATG CAT ATG AT-3’) and Ss_RBam1089 (5’-CTA TTT GTG GGA GAA ACC AAA TGT TAT ATG CAT ATG AT-3’).
TCC TAA GGA AAT AG-3′) and Ste_FNde5 (5′-AAA AAC CAA AAG TTA TAT GCA TAT GAT-3′)/Ste_RBam1089 (5′-TTA CCT CAA GGA TCC TAA GGA AAT TG-3′). These clonings created plasmids overexpressing Ain dpo4-like pol (pJM534), Saz dpo4-like pol (pJM520), Soh dpo4-like (pJM521) Ssh dpo4-like pol (pJM522), and Ste dpo4-like pol (pJM551) respectively.

**Construction of chimeric enzymes**

Chimeric dpo4-like genes were constructed by introducing a novel MscI restriction site into the Ste dpo4-like and the Ain dpo4-like genes in pJM551 and pJM534, just prior to the region encoding the little finger (LF) domain, using the QuickChange Mutagenesis kit (Stratagene, La Jolla, CA) to create pJM563 and pJM564 respectively. The Ain dpo4-like LF domain was replaced in pJM564 by sub-cloning the Ste dpo4-like LF domain out of pJM563 from MscI to BamHI to create the Ain/Ste dpo4-like chimera in pJM580. In a similar way, the Sso LF domain from the previously described plasmid p1947 (6) was sub-cloned from MscI to BamHI into pJM564 to create the Ain/Sso dpo4-like chimera in pJM590. The full-length nucleotide sequence of the pol IV numbers: DQ124674 (A. infernus with the LF from the HY50) can be located in the following GenBank accession numbers: DQ124674 (A. infernus with the LF from Soh Dpo4) and DQ124675 (A. infernus with the LF from Ste Dpo4).

**Purification of Dpo4-like enzymes**

Purification of the new Dpo4-like and chimeric enzymes was accomplished essentially as previously reported (3) except the protocol was simplified as outlined below. One or two liter LB-ampicillin cultures of the RW382 strain harboring the various Dpo4-like expression plasmids were grown at 37°C overnight. In the case of Ain/Sso dpo4-like pol, the undamaged (HTU50) and the abasic site-containing (HTX50) synthetic oligonucleotides used in the in vitro primer extension assays were synthesized by Lofstrand Laboratories (Gaithersburg, MD) using standard techniques and were PAGE purified prior to use. The synthetic abasic site (dSpacer) was purchased from Glen Research (Sterling, VA) and incorporated into the oligonucleotide template using standard techniques by Lofstrand Laboratories. The cis-syn CPD-containing oligonucleotide (HMTT50) was synthesized by Phoenix Biotechnologies (Huntsville, AL) and has been described previously (6). The 5-hydroxy-5-methylhydantoin-containing oligonucleotide (ODN) was synthesized as described previously (19). Single-stranded M13mp18 DNA was purchased from New England Biolabs (Beverly, MA). The sequence of each primer-template pair is given in the legend of the respective figures. Primers were labeled with [γ-32P]ATP using T4 polynucleotide kinase by Lofstrand Laboratories. Radiolabeled primers were annealed to the unlabeled templates at a ratio of 1:2 or 1:1.4, in the case of M13mp18, by heating the primer and template together at 95°C in annealing buffer (0.4 M Tris-HCl, pH 8.0, 50 mM MgCl2, 5 μg/ml BSA and 14.2 mM β-mercaptoethanol) and allowing the mixture to slowly cool to room temperature. Standard 10 μl reactions contained 40 mM Tris-HCl, pH 8.0, 5 mM MgCl2, 100 μM each dNTP (or a single dNTP, where noted), 10 mM DTT, 250 μg/ml BSA, 2.5% glycerol and 10 nM primer-template DNA. Reactions were carried out at 60 or 37°C, in the case of the hydantoin-containing template, with varying concentrations of enzyme and for varying lengths of time which are specified in the legend of the respective figures. Reactions were terminated by the addition of 10 μl of 95% formamide, 10 mM EDTA, heated to 95°C for 5 min and briefly chilled on ice. Aliquots containing 5 μl of the samples were subjected to electrophoresis through polyacrylamide, 8 M urea gels and replication products were visualized by PhosphorImager analysis.

**PCRs on undamaged DNA**

For the undamaged plasmid DNA template, 50 μl PCRs were set up containing standard ThermoPol reaction buffer [20 mM Tris-HCl, pH 8.8, 10 mM KCl, 10 mM (NH4)2SO4, 2 mM MgSO4, 0.1% Triton X-100] (New England Biolabs), 10 ng of circular pJM548 (Ste dpo4-like in pGEM-T), 200 μM of each ultrapure dNTP and 1 μM of each primer; Ste_FNde5 and Ste_RBam1089 whose sequences are provided above. Aliquots containing 2.5 μl of Taq DNA polymerase (New England Biolabs) were used as a positive control. Each of the Dpo4 enzymes was added to a concentration of 200 nM, except the Ssh Dpo4-like enzyme, which was added to a concentration of 400 nM. Reactions were heated to 85°C for 3 min and then treated with 35 cycles of 85°C for 30 s, 56°C for 1 min and 60°C for 4 min. Products were separated on a 0.9% agarose gel stained with ethidium bromide.

**Genomic DNA sample preparation**

K562 human genomic DNA (Promega Corporation, Madison, WI) was diluted in sterile water to a concentration of 100 ng/ml in a 1.5 ml microcentrifuge tube (Fisher Scientific, Norcross, GA) and exposed to UVC in a Stratalinker 1800 (Stratagene, La Jolla, CA) at a flux rate of 0.197 J/cm²/min.
PCR amplification and detection of Alu sequences

PCR amplification of the Alu insert was carried out with 2 ng of genomic DNA. The 25 μl reaction contained 1× Reaction Buffer (40 mM Tris–HCl, pH 8.0, 10 mM DTT, 60 mM KCl and 2.5% glycerol), 2.5 μM dNTPs, 2.5 U Taq Gold DNA Polymerase (Applied Biosystems, Foster City, CA), 3 mM MgCl₂, 20 pmol each of the forward and reverse primers, and 100 nM Dpo4, if applicable. Primer sequences were as follows: forward, 5’-GGT GTG GCC TTC GTC 3’ and reverse, 5’-GGA TGC TCG CTC TGT CG 3’ (20). The forward primer was labeled at its 5’ end with the reactive fluorescent dye 6-carboxyfluorescein (6-FAM). The forward primer binds a site near the Alu transcription start site. This sequence is identical in both the Major and the Precise Alu subfamily consensus (21), which comprise ~99% of Alu repeats in the human genome (20). The cycling conditions were: initial denaturation at 95°C for 11 min; 17 cycles: 94°C for 10 s, 56°C for 30 s, 72°C for 30 s; final extension 72°C for 5 min. For some experiments a smaller Alu fragment was amplified: the 25 μl reaction was carried out with 2 ng genomic DNA, 2.5 μM dNTPs, 3.25 mM MgCl₂, 10 μg non-acetylated BSA, 2.5 U AmpliTaq DNA Polymerase (Applied Biosystems), and 20 pmol each of the forward and reverse primers in 1× Buffer D3 (40 mM Tris–HCl, pH 8.0, 10 mM DTT, 6 mM KCl, 2.5% glycerol), and 100 nM Ste or Ain, if applicable. Cycling conditions were as follows: 85°C for 1 min; 22 cycles: 85°C for 30 s, 56°C for 1 min, 60°C for 5 min; 60°C for 5 min.

Amplified fragments were detected using the ABI Prism 310 capillary electrophoresis (CE) system (Applied Biosystems). A 1.5 μl aliquot of each amplified sample was added to 24 μl Hi-Di formamide (Applied Biosystems) and 1 μl of GeneScan 500 TAMRA internal lane standard (Applied Biosystems). Tubes were heated at 95°C for 3 min and snap cooled on ice for at least 3 min. Samples were injected onto the CE using the GS STR module (5 s injection, 15 kV, 60°C, run time 28 min, Filter Set C) module. Samples were subject to laser-induced fluorescence and analyzed with GeneScan 3.1.2 software (Applied Biosystems).

RESULTS AND DISCUSSION

Identification of novel Dpo4-like genes

We utilized the amino acid sequences of *S.acidocaldarius* Dbh (Sac) and *S.solfataricus* Dpo4 (Sso), as well as a hypothetical Dpo4-like protein encoded within the *S.tokodaii* genome (GenBank Accession number NC 003106) to design degenerate PCR primers to identify additional genes encoding polIV-like enzymes from other members of the *Sulfolobaceae* family. Five new polIV homologs were identified from *A.infernum* (Ain), *S.shibatae* (Ssh), *S tengchongensis* (Ste), *S.sozoricus* (Saz) and *S.ohwakuensis* (Soh). Once a fragment of a polIV-like gene was amplified, cloned and sequenced, we took advantage of the conserved genomic architecture surrounding the known polIV-like genes to develop a strategy to quickly obtain full sequence of the new genes (Figure 1A). Degenerate PCR primers to the upstream ribokinase gene and the downstream hypothetical gene were designed and in combination with either *dpo4*-like gene-specific or degenerate PCR primers, the ends of each of the new *dpo4*-like genes were cloned and sequenced. Subsequently, PCR primers to the 5′ and 3′ ends of each of the new genes were used to amplify the full-length *dpo4*-like gene.

Dbh and Dpo4 are from closely related organisms, but have very different biochemical properties (6). We therefore reasoned that determining the phylogenetic relationship of the new polIV-like polymerases to Dpo4 or Dbh might be predictive of their general enzymatic activities. Figure 1B shows an unrooted phylogenetic tree of *S.acidocaldarius* Dbh, *S.solfataricus* Dpo4, *S.tokodaii* Dpo4-like and the five
Figure 2. Purification and processivity of Dpo4-like proteins. (A) Scheme for the purification of Dpo4-like proteins. Whole-cell extracts (WC) were clarified and then heat treated at 70°C for 10 min. Denatured *E. coli* proteins were removed from the extracts by centrifugation (70). Subsequently, the soluble extracts were passed over a Hydroxyapatite Bio-Gel HTP Gel column (HAP) and then a SP Sepharose HP column (Sph). The Saz and Soh Dpo4-like proteins were purified over one additional column, a HiLoad 26/60 Superdex 75 column (data not shown). M designates the Benchmark prestained protein size marker (Invitrogen). (B) M13mp18 primer extension reactions using the previously identified *S. solfataricus* Dpo4 (Sso) and the five newly identified polymerases; *S. tengchongensis* (Ste), *S. shibatae* (Ssh), *A. infernus* (Ain), *S. azoricus* (Saz) and *S. ohwakuensis* (Soh). Primer extension of primer M13HTP (5'-CCT TAG AAT CCT TGA AAA CAT AGC GA-3') annealed to M13mp18 from base pairs 4101 to 4126 was performed at 60°C for 5 min utilizing 10 nM of the M13HTP/M13mp18 primer-template and increasing concentrations of the Dpo4-like enzymes; from left to right 0.2, 2.0 and 20 nM. Replication products were separated on 12%/8 M urea polyacrylamide gels and visualized by PhosphorImager analysis. Primer location and number of nucleotides added are indicated to the left of the gel. (C) M13mp18 primer extension reactions using the 'little finger' domain chimeric polymerases; *A. infernus*/S. solfataricus (Ain/Sso) and *A. infernus*/S. tengchongensis (Ain/Ste).
novel enzymes. This analysis revealed that the orthologs most closely related to *S. solfataricus* Dpo4 are, in order of relatedness, *S. shibatae* Dpo4-like, *S. tengchongensis* Dpo4-like and *A. infernus* Dpo4-like, and as we show below, these polymerases possess enzymatic properties somewhat similar to Dpo4. The *S. azoricus* and *S. ohwakuenensis* polIV-like genes are more distantly related to Dpo4 and have properties somewhat similar to Dbh. Thus, a simple phylogenetic analysis appears to be a good indicator of the overall enzymatic properties of the novel archaeal Y-family polymerases.

**Overexpression, purification and processivity of novel Dpo4-like polymerases**

In order to characterize the biochemical properties of the new polIV-like enzymes, they were overexpressed in an *E. coli* strain, RW382, lacking *umuDC* (polV) (22) and purified utilizing a similar, but somewhat simplified scheme, to that was previously reported (Figure 2A) (6). Such an approach allowed us to readily purify anywhere between 1 and 5 mg of each polymerase from just 1–2 l of cell culture. Once the polymerases were purified to >95% homogeneity, we initially characterized them for their relative degree of processivity. The primer extension assays were performed at 60°C, with various concentrations of each enzyme and utilized a primer annealed to a circular 7.2 kb M13mp18 template (Figure 2B). Under these conditions, the Saz and Ssh polIV-like proteins exhibited weak polymerase activity. Indeed, even at the highest enzyme concentration, the Saz and Ssh enzymes added only few nucleotides to the primer. As a consequence, we did not characterize the Saz and Ssh enzymes further. In contrast, the Ssh Dpo4-like and the Ste Dpo4-like enzymes exhibited a similar level of polymerase activity and processivity to Sso Dpo4, demonstrating that they are robust thermostable polymerases. Intriguingly, while the Ain Dpo4-like enzyme showed a somewhat similar level of primer utilization compared with Dpo4, Ssh Dpo4-like or Ste Dpo4-like, the size of the replication products was substantially shorter, suggesting that the Ain Dpo4-like polymerase is less processive than the other three enzymes. As noted above, however, the LF domain of the polymerase has been shown to be the major determinant of polymerase processivity (6). In an attempt to increase the processivity of the Ain Dpo4-like enzyme, we interchanged the LF domain of the Ain Dpo4-like enzyme with the LF domain of either Sso Dpo4 or Ste Dpo4-like polymerase to create the Ain/Sso Dpo4-like and the Ain/Ste Dpo4-like chimeras. As anticipated, the processivity of each chimera was significantly enhanced relative to the native Ain Dpo4-like enzyme (Figure 2C).

**Ability of Dpo4-like enzymes to facilitate PCR**

Although the thermostable Dpo4-like enzymes are typically less processive than replicative polymerases, they are nevertheless capable of replicating hundreds of bases in the M13mp18 primer extension assay and we have previously shown that Dpo4 could effectively amplify an ~1300 bp pre-denatured linear target DNA molecule at high template concentrations (~400 ng) in a PCR (3). The ability of Dpo4 and our new Dpo4-like enzymes to substitute for *Taq* in PCRs was reexamined under more optimal reaction conditions. As shown in Figure 3, Dpo4-like enzymes can efficiently amplify an ~1100 bp target DNA in a ‘closed-tube reaction’ when the denaturation temperature is lowered from the typical ≥90 to 85°C. Using this lower denaturation temperature, we could also eliminate the pre-denaturation step, amplify non-linear DNA, reduce the amount of DNA to about 10 ng and reduce the primer concentration from 30 μM of each primer used previously, to 1 μM. Under these conditions, it appears that the Ain/Sso Dpo4-like and Ain/Ste Dpo4-like chimeras are the most robust PCR enzymes belonging to the Y-family examined to date. Further optimization of PCR amplification by Dpo4-like polymerases is certainly possible, e.g. by altering the reaction buffer components such as salts, protein stabilizers or reagents that lower melting temperature.

**Translesion synthesis activity of the novel Dpo4-like enzymes**

The major biological function of Y-family polymerases is thought to be the synthesis of damaged DNA refractory to replication by the cell’s conventional replicases. We therefore compared the lesion bypass capabilities of our new enzymes with that of Dpo4 in a primer extension assay using DNA templates that contain lesions such as a *cis*-syn CPD, an abasic site (Ab) (Figure 4) or a 5-hydroxy-5-methylhydrantoin (5-OH-5-MeHyd) (hyd) (Figure 5). These lesions arise endogenously as a result of exposure to ultraviolet (UV) radiation in the case of CPDs, or due to the action of base-excision repair (BER) mechanism(s) or spontaneous hydrolysis of the *N*-glycosyl bond in the case of abasic sites (23). In addition, exogenous DNA, such as forensic or ancient DNA (aDNA) samples, can suffer the same types of insults due to exposure to environmental factors including UV-light, heat, humidity and other genotoxic substances. Indeed, hydantoin derivatives
of pyrimidine bases, which are the most prevalent lesions encountered in aDNA (24), can be produced via several oxidative processes as a result of prolonged environmental exposure (25). Using approximately equal activities of Taq and the Dpo4-like polymerases, we found that all three lesions block Taq polymerase (Figures 4 and 5). However, in dramatic contrast, all of the Dpo4-like enzymes not only preferentially incorporated the correct nucleotides opposite each lesion, but they also promoted full lesion bypass, albeit with varying efficiencies (Figures 4 and 5). Of particular note, is the fact that the Ain/Sso and Ain/Ste Dpo4-like chimeras appear to be the most efficient of the novel enzymes at bypassing all three DNA lesions.

Dpo4-like enzymes and their ability to PCR amplify damaged DNA templates

The ability to detect DNA polymorphisms using molecular genetic techniques has revolutionized the forensic analysis of biological evidence. DNA typing now plays a critical
role within the criminal justice system, but one of the limiting factors with the technology is that DNA isolated from biological stains recovered from the crime scene is sometimes so damaged as to be intractable to analysis. Currently no methods exist to rectify such damaged genomic DNA. The development of a method to restore and recover DNA profiles from damaged DNA would thus represent a significant advance with widespread applications in forensic casework and mass disaster investigations.

We reasoned, therefore, that a potential solution to this problem would be to perform the PCR of damaged DNA in the presence of a thermostable Y-family polymerase. To specifically test this hypothesis, human genomic DNA was irradiated with UVC light at a flux rate of 0.197 J/cm²/min for 6 min. Exposure of naked DNA in this manner causes a multitude of lesions including significant quantities of CPDs and (6-4) photoproducts (26). Genetic profile recovery from the UVC damaged substrate was assessed by amplification of Alu short interspersed elements comprising 10% of the genome that contain a limited number of bipyrimidine sites known to form photoproducts (20). The ubiquity of Alu sequences throughout the genome thus renders the repair substrate a de facto whole genome scan of UVC-induced DNA damage and its repair.

*Taq* Gold alone gave an Alu signal that was barely detectable (Figure 6A, top panel). Remarkably, a blend of *Taq* with 100 nM Sso Dpo4 in the PCR permitted the recovery of an Alu signal from the damaged genomic DNA substrate (Figure 6A, bottom panel). In four replicate experiments, there was a 36- to 45-fold enrichment of Alu sequences recovered from the damaged template by the *Taq*/Dpo4 blend. Alternative PCR conditions were employed with AmpliTaq to test the other Dpo4-like thermostable Y-family enzymes described above and these included a lower denaturation temperature (85°C), an increased extension time to permit efficient polymerase switching at damaged sites, and amplifying a smaller *Alu* fragment. Using these conditions in replicate experiments, Ste and Ain Dpo4-like enzymes produced a 14- to 18-fold and a 10- to 15-fold enrichment of *Alu* sequences from the damaged template, respectively (Figure 6B). As controls, the *Taq* Gold or AmpliTaq concentrations were doubled in the absence of the thermostable Y-family enzyme and this resulted in an average 1.5-fold increase in *Alu* signal. No further improvement in signal intensity was obtained with additional increases...
in AmpliTaq or Taq Gold concentration (data not shown). Thus, the Alu enrichment observed in these polymerase blend experiments is consistent with bona fide lesion bypass and not an increase in PCR efficiency mediated by the presence of more DNA polymerase. An amplification negative control was run with each experiment in which all PCR reactants, including the Dpo4 enzyme, were present with the exception of the human DNA template. All such amplification negative controls indicated the absence of human DNA contamination in any of the PCR reactants, including the Dpo4-like enzyme preparations. Sequencing of the repaired substrates was not conducted, since it would not provide confirmation that repair of UV-damaged DNA had taken place. Some of the original starting molecules will have been bypassed, some will remain undamaged and some will be damaged but be permissive of lesion bypass. If this population was sequenced, it is likely that the vast majority would possess that wild-type sequence, including even some that were repaired and possess the wild-type sequence. Thus, the chances of obtaining a few sequences that differ from the wild-type are remote and, if sequenced subsequent to cloning, could be confused with the normal PCR mutational processes.

Experiments are currently in progress to assess the success of this same strategy for the recovery of locus specific microsatellite marker profiles from damaged blood, semen and other biological stains, with Dpo4 and the Ain/Sso and Ain/Ste chimeras. Possible improvements in the efficiency of the repair of genomic DNA templates possessing multiple types of DNA lesions include the development of an in vitro system that combines translesion synthesis and base-excision repair.

CONCLUSIONS

Here, we report a quick and simple method for the identification of polIV-like polymerases from Archaea. Due to their thermostable nature, large quantities of the recombinant enzymes can be readily purified from E.coli. Of the five novel enzymes reported here, three were Dpo4-like in their processivity and lesion bypass properties, while two were Dhh-like and exhibited poor processivity and lesion bypass. Second generation recombinant enzymes, with improved enzymatic properties, were simply generated by interchanging the LF domain of the respective polymerase. Last, but most significantly, we have shown that by using a thermostable Y-family polymerase in a blend with a conventional PCR enzyme like Taq, we can PCR amplify damaged DNA samples. This has immediate applications in forensic science and the analysis of ancient DNA samples, which are often degraded and are not easily amplified by Taq alone.

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Conflict of interest statement. J.P.M and R.W are co-inventors on the International Patent Application No. PCT/US2005/017941 by the National Institutes of Health, which was filed on May 20, 2005.

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