New DNA polymerase from the hyperthermophilic marine archaeon Thermococcus thioreducens

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Abstract The family B DNA polymerase gene of Thermococcus thioreducens, an archaeon recently isolated from the Rainbow hydrothermal vent field, was cloned and its protein product expressed, purified and characterized. The gene was found to encode a 1,311 amino acid chain including an intein sequence of 537 residues. Phylogenetic analysis revealed a predominantly vertical type of inheritance of the intein in the Thermococcales order. Primary sequence analysis of the mature protein (TthiPolB) showed significant sequence conservation among DNA polymerases in this family. The structural fold of TthiPolB was predicted against the known crystallographic structure of a family B DNA polymerase from Thermococcus gorgonarius, allowing regional domain assignments within the TthiPolB sequence. The recombinant TthiPolB was found to have moderate thermal stability and fidelity, and a high extension rate, consistent with an extremely low $K_m$ corresponding to the dNTP substrate. TthiPolB performed remarkably well in a wide range of PCR conditions, being faster, more stable and more accurate than many commonly used enzymes.

Keywords DNA polymerase · Archaea · Hyperthermophilic · PCR · Thermococcus

Introduction

DNA polymerases play an essential role in the replication and repair of genetic material of all living organisms. They are classified into families according to their sequence similarities. The first four families (Ito and Braithwaite 1991) were defined by homology to Escherichia coli DNA polymerases (family A for Pol I homologs, family B for Pol II homologs, family C for Pol III homologs) and to the eukaryotic DNA polymerase $b$ (family X). Family D was later proposed to account for a new heterodimeric DNA polymerase found in the Euryarchaeota subdomain of archaea (Cann and Ishino 1999). The most recently described DNA polymerase family is the Y family (Ohmori et al. 2001), which consists of phylogenetically related DNA polymerases that are characterized by low fidelity DNA synthesis and their ability to bypass DNA lesions. While bacterial replicative DNA polymerases belong to the C family and eukaryal ones belong to the B family (except for mitochondrial DNA polymerases which are family A), the situation in archaea (which have no family A or C DNA polymerases) is more complex. According to available genome sequences, crenarchaeotes have 2 or 3 family B and no family D DNA polymerases. This implies that chromosomal replication is performed by family B DNA polymerases in the Crenarchaeota kingdom, while euryarchaeotes have a single family B polymerase and a single
family D DNA polymerase, both being likely involved in replication (Henneke et al. 2005).

Family B DNA polymerases from hyperthermophilic archaea have attracted considerable interest as PCR enzymes because of their remarkable thermal stability, their proofreading 3′-5′ exonuclease activity and their monomeric nature making them relatively easy to produce. Several have been characterized so far, especially from species belonging to the Thermococcales order such as *Pyrococcus furiosus* (Lundberg et al. 1991), *Pyrococcus abyssi* (Dietrich et al. 2002; Gueguen et al. 2001) *Thermococcus litoralis* (Kong et al. 1993), *Thermococcus kodakarensis* (Takagi et al. 1997), *Thermococcus kodakensis* (Katan et al. 1997), *Thermococcus gorgonarius* (Bonch-Osmolovskaya et al. 1996) and *Thermococcus fumicolans* (Camberon-Bonavita et al. 2000). Family B DNA polymerases from other euryarchaeota include *Methanobacterium thermoautotrophicum* thermoautotrophicus (Kelman et al. 1999) and crenarchaeotes such as *Pyrobaculum islandicum* (Kahler and Antranikian 2000). In contrast with their high sequence similarities, these DNA polymerases show important variations in their properties such as fidelity, thermal stability, kinetic parameters and processivity, even within the same genus.

A novel hyperthermophilic, obligately sulfur-reducing archaeon, *Thermococcus thioreducens* (*T. thioreducens*) was isolated from samples collected at the Rainbow hydrothermal vent site (Mid-Atlantic ridge) and described by our group (Pikuta et al. 2007). Its genomic material was used for the search of a family B DNA polymerase gene. An open reading frame encoding the targeted enzyme was identified, cloned and sequenced. The recombinant DNA polymerase *T. thioreducens* PolB (*TthiPolB*) was expressed, purified and characterized. The structural fold of TthiPolB was also predicted against known crystallographic structures of family B DNA polymerases to highlight relationships between functional properties and structural features.

**Materials and methods**

**Strains and culture conditions**

*Thermococcus thioreducens* (ATCC: BAA-394, DSM: 14981) was grown as described (Pikuta et al. 2007). Recombinant plasmids were propagated in *E. coli* strain TOP10 (Invitrogen, USA) for those based on pCR4-TOPO and strain TOP10F’ (Invitrogen, USA) for those based on pCRT7/CT-TOPO. *E. coli* strain Rosetta(DE3) (Novagen, USA) was the host for protein expression. All *E. coli* cultures were grown in Luria–Bertani medium containing 100 mg/l ampicillin (to select for the expression vector) at 37°C with vigorous shaking; chloramphenicol was added to a final concentration of 35 mg/l in expression cultures (to retain the Rosetta phenotype).

**Gene cloning**

Genomic DNA of *T. thioreducens* was isolated from 400 mg of fresh biomass through a standard phenol/chloroform extraction followed by ethanol precipitation (Sambrook et al. 1989). Two DNA polymerase gene fragments were amplified by PCR from the genomic DNA using primers pairs (5′-ATGATCTCGAYACTGACTAC/5′-CTCGCCCTCGTGTGTAAGGC and 5′-AGAAAGTAGCCGGTGTAG-AGACG/5′-GATGTAGCTTTACTCAGTCKCC) designed from conserved regions in an alignment of six available family B DNA polymerase sequences from Thermococcales species. A 105 bp linker DNA was amplified from the pCR4-TOPO vector (Invitrogen, USA), using T3 (5′-AACCCTCACCTAAAGGG) and T7 (5′-TAATACGACT CACTATAGGG) primers. Genomic DNA from *T. thioreducens* was digested with Eco RV and ligated to the linker with T4 ligase. Both enzymes were from Promega (USA) and used according to the manufacturer’s instructions. The product was used as a template in nested PCR with DNA polymerase-specific primers (designed from the previously amplified fragments) and linker-specific primers. Amplified fragments were excised from an agarose gel and inserted into a pCR4-TOPO vector for sequencing. The entire coding region of the DNA polymerase gene was then amplified with the high-fidelity DNA polymerase KOD (Novagen, USA) using primers (POLB-1F: 5′-ATGATCTCTCGATGCCGAC TACATCACCAG, POLB-1R: 5′-TCACCTCATTGTTTCCA CCTTCAGCCGCAGCC) based on the exact sequence of the gene ends, and cloned into a pCRT7/CT-TOPO (Invitrogen, USA) expression vector. The insert from two clones as well as the PCR product were completely sequenced. An inteinless version of the gene was constructed following the overlap extension method (Horton et al. 1989), amplifying the overlapping fragments from genomic DNA using primers POLB-1F with 5′-GTAGTAACTGTTTGCCAGA TCTTGATGGCGCGTG and POLB-1R with 5′-TC TGGCAACAGTTACTACGGCTACTACGGCTACGCC respectively. The inteinless gene was cloned into a pCRT7/CT-TOPO expression vector and its sequence verified.

**Sequence determination and analysis**

Sequence determinations were performed using an ABI 310 Genetic Analyzer and the DYNEnamic ET Terminator Cycle Sequencing Kit (Amersham-Pharmacia, USA). Raw sequences were manually edited with Chromas (http://www.technelysium.com.au/) and assembled in GCG.
(Accelrys, USA). Codon usage analyses, translations and calculation of molecular weight were performed using the Sequence Manipulation Suite (Stothard 2000). Sequence alignments were performed using ClustalW (Thompson et al. 1994). Phylogenetic analyses were conducted using MEGA version 2.1 (Kumar et al. 2001). Phylogenetic trees were constructed with the neighbor-joining method (Saitou and Nei 1987) with 1,000 replications of bootstrap resampling.

Nucleotide sequence accession numbers

The sequence of the family B DNA polymerase from *T. thioreducens* described in this study has been deposited in GenBank under accession number [GenBank:EF058197].

The sequence of the PCNA gene described in the exonuclease assay has been deposited under accession number [GenBank:EF058196]. Accession numbers of DNA polymerase sequences from other *Thermoccocus* species mentioned in this study are as follows: *T. aggregans*, [GenBank:Y13030]; *T. fumicola*, [GenBank:Z69882]; *T. hydrothermalis*, [GenBank:AJ245819]; *T. litoralis*, [GenBank:M74198]; *T. peptonophilus*, [GenBank:E13953]; *T. sp. 9°N-7*, [GenBank:U47108]; *T. sp. GE8*, [GenBank:AJ250333]; *T. kodakarensis*, [GenBank:D29671].

Sequence and domain assignments

The functional domains of TthiPolB were identified within its primary sequence by superimposing the homologous residues onto the known crystallographic structure of thermostable family B DNA polymerase (TgoPolB) from *Thermococcus gorgonarius* [PDB:1tgo]. TthiPolB residues were positioned by chemical based conformational sampling coupled to the usage of energy restraints derived from the force field available in the crystallographic and NMR System (CNS) program suite (Brunger et al. 1998). The resulting detailed model was refined with conjugate gradient minimization with no experimental energy terms used. All backbone atoms of the molecules were fixed while the remaining atoms were minimized for 500 steps. The resulting structure was visualized with PYMOL (http://www.pymol.org).

Expression and purification

A culture of *E. coli* Rosetta(DE3) harboring the recombinant expression vector containing the inteinless version of the TthiPolB gene was grown in a 10 l fermenter at 37°C. When the optical density at 600 nm reached 0.5, expression was induced by the addition of isopropyl thio-β-D-galactoside (IPTG) at a final concentration of 0.5 mM. After 16 h, cells were harvested by centrifugation. All subsequent steps were performed at 4°C. Cell were resuspended in buffer P (10 mM sodium phosphate, pH 7.0, 100 mM NaCl, 0.1 mM EDTA, 0.5 mM PMSF and 0.5 mM dithiothreitol) and disrupted by sonication (6 cycles of 45 pulses) using a Branson Sonifier 250 (VWR Scientific, USA). Cell debris were removed by centrifugation (12,000g, 20 min). The supernatant was heated for 25 min at 80°C and the precipitate was removed by further centrifugation. The supernatant was loaded into a heparin column (HiTrap Heparin HP, Amersham, USA) that had been pre-equilibrated with buffer P and eluted with a 0.1–1.5 M NaCl linear gradient in buffer P, using an Äkta Explorer FPLC system (Amersham-Pharma, USA). Fractions containing DNA polymerase (showing a band of the expected size on a SDS-PAGE gel) were pooled, dialyzed against buffer B (25 mM Bis-Tris-Propane pH 9.5, 0.1 mM EDTA, 50 mM KCl, 0.5 mM PMSF and 0.5 mM dithiothreitol) and applied to an anion-exchange column (HiTrap Q HP, Amersham, USA) that had been pre-equilibrated with the same buffer. Bound proteins were eluted with a 0.0–0.75 M NaCl linear gradient in buffer B. Fractions containing DNA polymerase were pooled, concentrated using a Centricon Plus PL-30 centrifugal filter device (Millipore, USA), applied to a Sephacryl S-200 gel filtration column (Pharmacia, USA) pre-equilibrated with buffer T (50 mM Tris pH 7.5, 0.1 mM EDTA, 50 mM KCl, 0.5 mM PMSF and 0.5 mM dithiothreitol) and eluted with the same buffer. Fractions containing DNA polymerase were pooled and mixed with glycerol and Triton X-100 at a final concentration of 50 and 0.5%, respectively. Aliquots were stored at −20°C in 1.5 ml tubes. Protein concentrations were determined according to Bradford (Bradford 1976), using BSA as a standard.

DNA polymerase activity assay

A non-radioactive, fluorescence-based DNA polymerase activity assay was used which we adapted from Tveit and Kristensen (2001), and which was originally described by Seville et al. (1996). It was based on the specific binding affinity of the dye PicoGreen (Molecular Probes, Invitrogen) with dsDNA. Annealed primer-template was prepared as follows: a solution containing 200 μg/ml M13mp18 ssDNA (Bayou Biolabs, USA) and 170 nM UPlong primer (5’-TTCCCAGTCAGGACGTTCTTAAACAGACGGGCTCAC GTG) was heated at 70°C for 5 min, cooled to room temperature over 20 min and stored at −20°C until needed. A typical DNA polymerase assay was performed as follows: 1.2 μl of the annealed primer-template was mixed with 2 μl of dNTP solution (at 2.0 mM each) and 1.8 μl of 10× reaction buffer in a final volume of 18 μl in a 500 μl PCR tube. Unless specified otherwise, the 10× reaction buffer consisted of 100 mM Tris-HCl pH 8.25, 600 mM KCl, 20 mM MgCl₂ and 1% Triton X-100. After a 1-min
equilibration at the assay temperature (74°C unless specified otherwise), the reaction was started by the addition of 2 μl of DNA polymerase solution (diluted in the same reaction buffer). Incubation took place in a thermal cycler (Mastercycler gradient, Eppendorf, USA). The reaction was stopped after a precise amount of time by the addition of 2 μl of 250 mM EDTA, after which the reaction tube was immediately put on ice until all reactions were completed. Reactions were diluted with 150 μl TE (10 mM Tris-HCl pH 7.2, 1 mM EDTA) and transferred into 3.5 ml methacrylate cuvettes (Perfector Scientific, USA), where 1.5 μl of PicoGreen and 850 μl TE were added. Cuvettes were kept in the dark until the fluorescence was measured in a FluoroMax-3 fluorometer (Jobin Yvon, USA). Excitation and emission wavelengths were 485 and 525 nm, respectively. Because of the small volume, a 12-mm platform was placed below the sample in the cuvette compartment in order to get reproducible readings. Fluorescence values were converted into dsDNA mass amounts using a standard curve obtained with known amounts of added dsDNA. Incorporation of dNTPs (in pmol per unit of time) was calculated as the increase in dsDNA amount in pg divided by 649 (average molecular weight of a deoxyribonucleoside monophosphate pair). One unit of DNA polymerase activity was defined as the amount required to incorporate 10 nmol of dNTP into dsDNA in 30 min (extrapolated from 2-min measurements) at 74°C in the reaction buffer described above.

In the case when DNA polymerase activity was measured as a function of temperature, MOPS buffer was used instead of Tris in order to minimize pH variations with temperature. When relative DNA polymerase activity was measured as a function of buffer type and concentration, the pH of buffers were adjusted at room temperature so that they would all be 7.5 at 72°C, as calculated from the temperature coefficients (Tris: −0.028, Heps: −0.014, MOPS: −0.011, phosphate: −0.003 pH unit per °C). In addition, Triton X-100 was not included in the buffers used in these experiments.

Structural thermostability

Thermal stability of the DNA polymerase structure was investigated through circular dichroism measurements as a function of time during incubation at 95°C. DNA polymerase solution was dialyzed overnight against sodium phosphate buffer pH 8.9 and diluted to a final concentration of 50 μg/ml in the same buffer. A volume of 2.4 ml was used in a 10 mm cylindrical quartz cell (Starna Cells, USA). Circular dichroism at 200–250 nm (10 reads per points) was measured at regular time intervals during a total of 35 h, on an OLIS RSM 1000 CD spectrophotometer (Olis, USA).

Functional thermostability

Functional thermostability was evaluated by measuring residual DNA polymerase activity after various pre-incubation times at 95°C. A solution of TthiPolB at 17 μg/ml in the assay reaction buffer described above was incubated at 95°C for over 3 h. Samples were taken at various times and cooled to room temperature, before being diluted ten times in reaction buffer. DNA polymerase activity was measured as described with four replicates per time point using 2 μl of the diluted DNA polymerase per reaction. The assay temperature was 72°C. Taq polymerase (New England Biolabs, USA) was used as a control in parallel under the same conditions except for the reaction buffer, which was the one provided by the manufacturer.

Fidelity assay

The error rate generated by TthiPolB was measured using a modified version of the yeast-based PCR fidelity assay described by Flaman et al. (1995), in which the reporter gene HIS3 was replaced by the ADE2 gene. Wild-type TP53 cDNA was PCR amplified using TthiPolB. The PCR product was co-transformed with a linearized expression vector into a TP53 transcriptional activity reporter strain, and cloned in vivo by homologous recombination. After selection of colonies expressing the TP53 protein, white colonies expressed the wild type protein whereas red colonies expressed mutant proteins (Flaman et al. 1994). Therefore, the percentage of red colonies was used to calculate the error rate per nucleotide per cycle (ER) using the following formula: ER = % red colonies/(d × del × 100), where d is the amount of effective doublings and del is the effective target size (542 bp). The amount of effective doublings was calculated using the equation 2d = (amount of PCR product)/(amount of starting target). PCR amplifications were performed in a 50 μl volume using 200 μM dNTP, 1.5 mM MgCl2, 10 mM Tris-HCl pH 9.0 (at 25°C), 50 mM KCl, 0.1% Triton-X100, 50 pmol of P3 and P4 primers (Flaman et al. 1994) and 13 pg of input DNA. The cycling profile consisted of 30 cycles of 94°C 20 s, 62°C 30 s and 72°C 2 min, preceded by 3 min at 95°C and followed by 5 min at 72°C. The fidelity assay was performed in parallel with Taq polymerase (ABgene, France) as a control.

Exonuclease assay

The substrate for the exonuclease assay was a 750 bp long double-stranded DNA fragment encoding the T. thioreducens PCNA, amplified by PCR using TthiPolB, separately with PTO-modified primers and unmodified primers. The primers (forward: 5’-ATGCCCCGGAGATGTTTTTGA TG, reverse: 5’-TCAGTCCCTCGACGGCGGCGGAG
1.8 ng of substrate (purified PCR product) was mixed with primers. Exonuclease assay was performed as follows: GeneClean Turbo (Qbiogene, USA) in order to remove free modified primers. The PCR products were purified using GeneClean Turbo (Qbiogene, USA). The PTO Extremophiles Km 10.0 nM in the final reaction). For the determination of the apparent Km parameter and the dNTP parameter experiments. The apparent Km parameters were determined from a Lineweaver–Burk plot (Lineweaver and Burk 1934).

DNA polymerase activity assays were performed as described in four replicates for each data point and three incubation times per condition. For the determination of the Km corresponding to the DNA substrate, the reaction mix contained a variable amount of primed template (0.5–10.0 nM in the final reaction). For the determination of the Km corresponding to the dNTP substrate, the reaction mix contained dNTP concentrations ranging from 1 to 200 µM each. The DNA polymerase amount was 1.7 ng per reaction and 1.13 ng per reaction, respectively for the DNA parameter and the dNTP parameter experiments. The apparent Km parameters were determined from a Lineweaver–Burk plot (Lineweaver and Burk 1934).

Extension rate

Extension rate was determined by applying the DNA polymerase activity assay described above to monitor the extent of conversion of the M13 template into dsDNA under a large excess of DNA polymerase. A template concentration of 1 nM instead of 6 nM was used with polymerase to template molar ratios of 2–50, incubation times from 0 to 5 min and four replicates per data point.

Polymerase chain reaction

Unless specified otherwise, PCR using TthiPolB was performed as follows. Reaction mixture typically contained 10% of the 10× reaction buffer described above, 200 µM each dNTP, 600 nM each primer, a variable amount of template and between 140 and 270 ng/ml DNA polymerase (corresponding to 1 to 2 units per 50 µl reaction). The cycling profile was: initial denaturation, 2 min at 95°C; 30 cycles of 15 s denaturation at 95°C, 20 s annealing at 50–65°C and 30–60 s extension at 72°C; final extension, 3 min at 72°C.

Results and discussion

Cloning and sequence analysis of a family B DNA polymerase gene

A comparison of family B DNA polymerase sequences from the Thermococcales order revealed a remarkable high similarity at several conserved regions, including at the nucleotide level, allowing the design of PCR primers with limited degeneracy. Two non-overlapping fragments, respectively, 450 and 330 base pairs (bp) long, were amplified with such primers, and their sequence confirmed that they belonged to a family B DNA polymerase gene. An approach similar to the vectorette PCR technique (Riley et al. 1990), yet simpler, was devised to determine the ends of the gene sequence. A linker was designed from a region of the pCR4-TOPO vector that could easily be amplified using readily available T3 and T7 primers and which was verified not to correspond to any archaeal genomic sequence. Digested genomic DNA from T. thioreducens was then ligated with this linker and used as a template for PCR using DNA polymerase-specific primers designed from the previously amplified fragments and linker-specific primers. The resulting amplified fragments were determined to contain sequences corresponding to the DNA polymerase gene termini. The fragments encoded 131 and 182 amino acid residues of the N- and the C-termini, respectively, as well as additional upstream and downstream sequences. The entire gene, amplified using primers designed from the newly determined gene terminal sequences, was found to be 3,936 bp long and to encode a 1,311 amino acid (aa) polypeptide chain (Fig. 1a).

Alignments with homologous DNA polymerase gene sequences revealed the presence of an intein gene within the coding sequence, encoding a 537 aa protein and splitting the DNA polymerase sequence into a 491 aa N-terminal and a 283 aa C-terminal fragments (Fig. 1b). Inteins are “selfish” genetic elements that are post-translationally excised from a protein precursor through a self-catalytic protein splicing process (Perler 2002; Pietrokovski 2001). First discovered in yeast (Hirata et al. 1990; Kane et al. 1990), over 300 have been identified so far in all domains of life (http://www.neb.com/neb/inteins.html). In addition to their splicing domains, most have a homing endonuclease domain that confers them potential mobility. Archaeal family B DNA polymerase genes have three known specific insertion sites for intein sequences designated pol-a, pol-b and pol-c. The...
majority of archaeal DNA polymerase genes sequenced so far include one to three intein elements (Fig. 2a). According to InBase nomenclature, the intein within the T. thioreducens family B DNA polymerase belongs to the “DNA polymerase motif B” group, typified by the prototype allele Tli Pol-1, and should be designated Tthi Pol. Among Thermococcus DNA polymerases that have been studied so far, Tthi PolB is the only one with a single intein at the pol-b insertion site. Tthi Pol’s closest homologue, with 77.1% identity, is Thy Pol-1 from T. hydrothermalis, which has a second intein at the pol-c insertion site. A phylogenetic tree illustrating the position of Tthi Pol among other known inteins of the DNA polymerase motif B group is shown in Fig. 2b. A comparison with the phylogenetic tree of family B DNA polymerases belonging to the Thermococcales order (Fig. 2c) suggests a predominantly vertical type of inheritance of the motif B inteins. Indeed, with the exception of Pyrococcus sp. GB-D, the tree topology is remarkably conserved between the DNA polymerases and the motif B inteins, with 92.4, 93.4 and 92.5% identity, respectively. Alignments of complete protein sequences (not available for T. hydrothermalis) show 92.9% identity with T. 9°C/N-7 and 92.3% with T. GE8. The mature form of Tthi PolB is 774 amino acid residues long and has a calculated molecular weight of 89.98 kDa. Tthi PolB will be discussed from here on as the mature T. thioreducens family B DNA polymerase without its intein sequence. The amino acid sequences corresponding to the functional domains are shown against the known crystallographic structure of family B DNA polymerase from Thermococcus gorgonarius [PDB:1tgo] (Hopfner et al. 1999), designated as TgoPolB from here on, that shares 91.0% sequence identity with Tthi PolB (Fig. 1c).
While work to pursue the three-dimensional structure of TthiPolB by X-ray crystallography is in progress, the primary sequence of TthiPolB was aligned and modeled against the crystallographic structure of TgoPolB to predict the protein’s tertiary structure. Among the family B DNA polymerase structures determined to date, TthiPolB has the highest sequence identity to that of TgoPolB. We have therefore assumed that the three-dimensional structure of TthiPolB would share very similar topological and structural features with TgoPolB (Hopfner et al. 1999). The projected structural domains within the mature TthiPolB enzyme include the N-terminal domain (M1-E130, F327-A368), the exonuclease (3'-5') domain (G131-F326), the palm (P369-I449, A501-F588), fingers (P450-Y500) and thumb (F589-R775) (Fig. 1c). These are the general structural characteristics of family B DNA polymerases and examples of these are readily observed in the structures of gp43 from bacteriophage RB69 (Wang et al. 1997), 9°N-7 pol from Thermococcus sp. 9°N-7 (Rodriguez et al. 2000) and KOD polymerase from T. kodakarensis (Hashimoto et al. 2001). Moreover, the exonuclease and palm domains share structural similarities and have comparable active sites to those of family A DNA polymerases and they both show metal dependence for exonuclease and replication mechanisms (Hopfner et al. 1999; Wang et al. 1997).

Expression and purification

Failed attempts to express TthiPolB in common E. coli hosts suggested sensitivity to codon bias, as a comparison of codon usage in the TthiPolB gene and in the E. coli genome (Nakamura et al. 2000) revealed significant
differences. Consequently, a Rosetta strain, carrying a plasmid providing the host with an additional supply of rare tRNA genes (argU, ileX, glyT, leuW and proL) was used for expression. Recombinant TthiPolB could then be expressed at high levels.

TthiPolB was purified to >95% purity, as estimated by SDS-PAGE analysis, after a heat treatment followed by three chromatographic steps: heparin, anion-exchange and gel filtration. Approximately 20 mg of purified TthiPolB can be routinely obtained per liter of culture with a specific activity of 146 U/µg. The specific activity is remarkably high compared to other reports on archaeal family B DNA polymerase purification [66 U/µg with *Pyrococcus abyssi* (Dietrich et al. 2002), 25 U/µg with *Thermococcus sp. 9°N-7* (Southworth et al. 1996)] and consistent with a higher extension rate (see below), although some of the differences can probably be attributed to differences in purity and to the way units are defined and measured.

Biochemical properties

The enzymatic activity of TthiPolB was measured under various conditions of pH, temperature and medium composition (Fig. 3), using a DNA polymerase assay adapted from Tveit and Kristensen (2001). More than 80% of maximal activity was obtained between pH 7.8 and 9.0, measured at 25°C, which corresponds to pH 6.5–7.7 at 72°C when Tris is used as a buffer (Fig. 3a). In contrast with *T. thioreducens’* optimal growth temperature of 82–85°C and upper growth limit of 94°C (Pikuta et al. 2007), maximal DNA polymerase activity was observed between 72 and 76°C, above which activity decreased abruptly (Fig. 3b). This is probably the result of thermal denaturation of the DNA substrate rather than the enzyme itself. At temperatures above the apparent optimum for activity, most primer molecules are free rather than annealed to the M13 template, which prevents determination of the actual optimum temperature.

The enzyme was active under a wide range of salt concentrations, and showed a slight preference for potassium chloride over sodium chloride (Fig. 3c). Over 80% of maximal activity was obtained with KCl concentrations of 50–80 mM, with a maximum at 60 mM. Magnesium ions are known cofactors for DNA polymerases. As expected, no activity was detected in the absence of magnesium. Activity was maximal between 1.5 and 3.0 mM magnesium chloride (Fig. 3d).

The effects of buffer type and concentration on DNA polymerase activity were examined (Fig. 3e). Tris was

![Fig. 3](https://via.placeholder.com/150)

Fig. 3 Effects of pH, temperature, salts and buffers on TthiPolB activity. Relative DNA polymerase activity was measured as a function of pH (a), temperature (b), potassium and sodium chloride concentration (c), magnesium chloride concentration (d) and buffer type and concentration (e). Activity assays were carried out at 72°C as described except for the varied reaction condition and the absence of Triton X-100. MOPS buffer instead of Tris was used in experiment b because of its lower temperature coefficient, in order to minimize pH variation with temperature. Buffer pH were adjusted at room temperature so they would all reach the value of 7.5 at 72°C, as calculated from their temperature coefficients. Error bars represent standard deviation calculated from four replicates.
found to allow the highest activity. The enzyme was 20% less active in HEPES and MOPS and 50% less active in phosphate buffer at concentrations of 10 mM. In all cases, activity decreased with increased buffer concentration. The decrease was stronger in the case of phosphate, possibly because of a reduced availability of magnesium ions through complexation with phosphate. The same experiment was done with decreased concentrations of potassium chloride while the buffer concentration was increased, in order to minimize ionic strength fluctuations, and gave very similar results (data not shown), suggesting that even Tris has an inhibitory effect on DNA polymerase activity and may not be the ideal buffer for this DNA polymerase, although it was the best among the ones we tested.

The DNA polymerase activity was also measured with other additives commonly used in DNA polymerase buffers. The presence of bovine serum albumin (tested at 0–200 µg/ml) or ammonium sulfate (tested at 0–20 mM) in the Tris-containing reaction buffer did not have any significant effect on activity, while addition of Triton X-100 at a concentration of 0.1% had a notable effect, increasing DNA polymerase activity by as much as 30% (data not shown).

Thermal stability

Thermal stability of TthiPolB was investigated from both a structural and a functional point of view. Circular dichroism (CD) spectroscopy was used to detect structural changes during incubation at 95°C, at the wavelengths typically used to monitor β-sheets (212 nm, Fig. 4a) and α-helices (222 nm, Fig. 4b). The data obtained suggest an extreme structural stability at this temperature, with no detectable change during the first 20 h and an apparent half-life of 22 h. Interestingly, evaluation of functional thermostability as the measure of residual DNA polymerase activity after incubation at 95°C showed a quite different result (Fig. 4c), with a half-life at that temperature of 125–130 min, similar to published values for other thermostable DNA polymerases (Table 1). The result obtained with Taq as a control further validated the method used, giving a half-life of 45 min, consistent with the manufacturer’s claim of 45 min at 94°C (New England Biolabs). The absence of correlation between structural and functional thermal stability observations suggests that the activity loss of TthiPolB with time may result from local irreversible changes affecting the active site rather than protein denaturation on a global scale.

Fidelity

The fidelity of TthiPolB was compared to that of Taq polymerase using a yeast-based PCR fidelity assay (Flaman et al. 1994). The error rate exhibited by TthiPolB was $1.4 \times 10^{-5}$ errors per nucleotide and per replication event, while it was $2.4 \times 10^{-5}$ for Taq polymerase. In other words, TthiPolB has a 1.7 times higher fidelity than Taq polymerase. This value is similar to those obtained with other archaeal DNA polymerases Tfu from T. fumicolans (Cambon-Bonavita et al. 2000) and Deep Vent from Pyrococcus sp. GDB (Flaman et al. 1994) using the same assay under similar conditions, giving error rates of $1.6 \times 10^{-5}$ and $1.2 \times 10^{-5}$, respectively. However, the fidelity of TthiPolB does not match that of Pfu (from P. furiosus), which is reported to exhibit an error rate that is almost an order of magnitude lower than Taq (Cline et al. 1996; Flaman et al. 1994).

![Fig. 4](image-url) Thermal stability of TthiPolB. Structural thermostability was monitored by circular dichroism, measured as a function of incubation time at 95°C. The unfolding of β-sheets (a) and α-helices (b) was recorded at 212 and 222 nm, respectively. There were no apparent structural changes until about 20 h. Thermal stability of TthiPolB’s DNA polymerase activity was also measured (c). TthiPolB and Taq DNA polymerase residual activities were measured and compared at 72°C as a function of pre-incubation time at 95°C. Error bars represent standard deviation calculated from four replicates. The half-life values of TthiPolB and Taq were 125–130 min and 45 min, respectively.
Table 1  TthiPolB parameters compared with those of other thermostable DNA polymerases

<table>
<thead>
<tr>
<th>Source species</th>
<th>Pol family</th>
<th>Half-life at 95°C (min)</th>
<th>Relative fidelity</th>
<th>$K_{m}^{DNA}$ (nM)</th>
<th>$K_{m}^{dNTP}$ (µM each)</th>
<th>Extension rate (nt/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thermococcus thioreducens</td>
<td>B</td>
<td>127±1</td>
<td>1.7a</td>
<td>0.66 ± 0.02a</td>
<td>10.5 ± 1.5a</td>
<td>≥96±2</td>
</tr>
<tr>
<td>Thermococcus sp. 9°N-7</td>
<td>B</td>
<td>–</td>
<td>–</td>
<td>0.05 ± 0.03b</td>
<td>75 ± 36b</td>
<td>–</td>
</tr>
<tr>
<td>Thermococcus fumicolans</td>
<td>B</td>
<td>200±2</td>
<td>1.5±1</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Thermococcus kodakarenensis</td>
<td>B</td>
<td>720±4</td>
<td>3.7±1</td>
<td>–</td>
<td>–</td>
<td>106–138±d</td>
</tr>
<tr>
<td>Thermococcus litoralis</td>
<td>B</td>
<td>300±2, 480±2</td>
<td>2.9±1</td>
<td>0.07–0.12e</td>
<td>41–57e</td>
<td>17±1</td>
</tr>
<tr>
<td>Pyrococcus furiosus</td>
<td>B</td>
<td>1,140±1</td>
<td>6.2±1</td>
<td>0.7f</td>
<td>16 ± 2f</td>
<td>9.3 ± 1.7f</td>
</tr>
<tr>
<td>Thermus aquaticus</td>
<td>A</td>
<td>45±1</td>
<td>1</td>
<td>4.0 ± 0.7f</td>
<td>24 ± 2f</td>
<td>47 ± 1f</td>
</tr>
</tbody>
</table>

Relative fidelity is defined as the error rate of Taq divided by the error rate of the DNA polymerase tested, when both were determined in the same experiment under the same conditions. For consistency, only fidelity values obtained through experiments in which all reaction mixtures contained 200 µM each dNTP (standard PCR conditions) are shown. References: a(this study); b(Southworth et al. 1996); c(Cambon-Bonavita et al. 2000); d(Takagi et al. 1997); e(Kong et al. 1993); f(Hogrefe et al. 2001); g(Cline et al. 1996)

Exonuclease activity

Exonuclease assays commonly use radiolabeled templates and measure either the release of radioactive nucleotides from double-stranded DNA (dsDNA) or the decrease in size of end-labeled oligonucleotides through autoradiography (Kong et al. 1993; Blöndal et al. 2001; Hogrefe et al. 2001). In this study, a non-radioactive approach was chosen and an assay was designed based on the one used for DNA polymerase activity determination (Tveit and Kristensen 2001) where variation in dsDNA concentration is measured using a fluorescent dye. An exonuclease-resistant template was used to discriminate between 3’-5’ and 5’-3’ directionality. Phosphorothioate (PTO) modified oligonucleotides in which a non-bridging oxygen is replaced by a sulfur in the phosphate backbone have been shown to confer resistance to exonuclease activity (Kunkel et al. 1981; Putney et al. 1981). For this reason, PTO-modified oligonucleotides have been used in gene therapy (Agrawal et al. 1988) and even in PCR to improve specificity by preventing primer degradation by proof-reading DNA polymerases (de Noronha and Mullins 1992; Skerra 1992). In our assay, a DNA sequence was amplified using 5’-PTO modified primers, generating a double-stranded fragment that was resistant to 5’-3’ exonuclease degradation while being susceptible to 3’-5’ degradation. The same sequence was amplified by unmodified primers, generating an unprotected double-stranded molecule. We anticipated that in the absence of dNTP, a decrease in dsDNA concentration over time would indicate the presence of exonuclease activity, and would demonstrate 3’-5’ directionality when the 5’-PTO protected substrate is used. In the presence of dNTP, any 3’-5’ exonuclease activity would be masked by the presence of the polymerase activity, therefore a decrease in dsDNA concentration with the unprotected substrate could only be the result of a 5’-3’ exonuclease activity, while the PTO-modified substrate would serve as a negative control. The limitation of our method is that quantitation of 3’-5’ exonuclease activity cannot be performed in the presence of dNTP, preventing the determination of an accurate polymerase to exonuclease activity ratio under physiological conditions.

The results of our exonuclease assay are shown in Fig. 5. In the absence of dNTP, exonuclease activity is demonstrated by the decrease in dsDNA concentration comparatively to the control. The same level of decrease when the template is 5’-protected indicates that the directionality of the exonuclease activity is 3’-5’ (Fig. 5a). We calculated the degradation rate to be approximately two nucleotides per second per enzyme molecule under the assay conditions.

In the presence of dNTP, the assay showed a surprising increase in dsDNA concentration with the unprotected and a somewhat smaller increase with the PTO-protected templates, while the negative control containing no DNA polymerase did not show any significant change (Fig. 5b). Since no family B DNA polymerase was ever reported to exhibit a 5’-3’ exonuclease activity so far, we expected to see no variation in dsDNA level over time. The hypothesis that the intriguing increase in dsDNA concentration could be explained by the extension of partially extended chains in the PCR product could be dismissed because nucleotide incorporation was an order of magnitude slower than when a regular primed template was present. In addition, during the course of the experiment, the amount of dsDNA increased by 80%, which would indicate that as much as half of the PCR product would have had to be single-stranded at the start of the experiment, a highly unlikely scenario considering the small size of the amplified sequence and the long extension times used during its amplification. An extensive investigation of this phenomenon was outside the scope of this report and will be the
object of future studies. However, it is strongly suspected that the observed result is due to the extension of self-primed 3'-0 ends. Because of the high assay temperature, the DNA present is probably in dynamic equilibrium between different secondary structures and the 3'-0 ends can form transient hairpins which can become templates for regular DNA polymerizing activity. Extension of such priming sites requires simultaneous strand displacement, which would explain the lower extension rate.

While the exonuclease assay described here allowed the demonstration of a 3'-5' exonuclease activity in TthiPolB, no conclusion could be drawn on a possible 5'-3' activity because of the unexpected DNA synthesis that was observed (a decrease in dsDNA amount with the unprotected template and no variation with the PTO-protected template, both in presence of dNTP, would have indicated the presence of a 5'-3' exonuclease activity). However, such an activity in TthiPolB is very unlikely because of its homology to family B DNA polymerases with known structures (Hashimoto et al. 2001; Hopfner et al. 1999; Rodriguez et al. 2000). In contrast with family A DNA polymerases which contain both 5'-3' and 3'-5' exonuclease domains, known family B DNA polymerases only have a single exonuclease domain of 3'-5' directionality.

Kinetic parameters

Apparent Michaelis constants were determined for each of the two substrates, annealed primer-template DNA and dNTP, in the presence of an excess of the other substrate (Fig. 6a, b). The K_m for the DNA parameter, 0.66 ± 0.02 nM, was an order of magnitude higher than that of other archaeal family B DNA polymerases while being significantly lower than that of Taq polymerase (Table 1). However, the K_m for the dNTP parameter, 10.5 ± 1.5 μM each, was lower than any published constant for a thermophilic DNA polymerase.

As opposed to the determination of K_m values, which does not require knowledge of enzyme concentration nor even precise activity measurements (relative activity can suffice), it is crucial to have a high precision in both the knowledge of enzyme molar concentration and the ability to measure absolute activity values in order to determine the k_cat accurately from Michaelis–Menten kinetic experiments. We devised an experiment that would allow a more reliable estimation of the k_cat by being completely independent from a precise knowledge of enzyme or DNA concentration. When a template of known size (the primed M13 template has a 7,215 nucleotide long single-stranded region) is incubated with excess amounts of polymerase and variation of dsDNA concentration is measured over time, the concentration of dsDNA is expected to reach a plateau corresponding to the conversion of all available single stranded DNA (ssDNA) into dsDNA. The height of the plateau should be independent of polymerase concentration while the slope of the initial part of the curve should increase with increasing enzyme concentration until reaching a maximum corresponding to the polymerase’s maximal extension rate (in other words, the k_cat). Under these conditions, the extent of conversion from ssDNA to dsDNA can reasonably be expressed in terms of template length rather than template concentration. The experimental results (Fig. 6c) indicate that at least 80% of the template 7,215 nucleotides were converted into dsDNA.

![Fig. 5 Exonuclease activity assay. Changes in dsDNA amounts were measured as a function of incubation time at 72°C in the absence (a) and presence (b) of dNTP. Experiments using TthiPolB are indicated with filled squares. Negative controls with no DNA polymerase present are shown as open squares. Solid lines correspond to unmodified template while dashed lines correspond to 5'-PTO modified template. Error bars represent standard deviation calculated from three replicates. The decrease in dsDNA content (expressed in amount of base pairs destroyed) in the absence of dNTP in the reaction medium (a) indicates exonuclease activity. The same level of activity with 5'-PTO modified (exonuclease-resistant) template suggests a 3'-5' directionality of the exonuclease activity. In the presence of dNTP (b), an increase in dsDNA content (expressed in base pairs created) is observed, indicating an unexpected DNA synthesis, partially inhibited when the template is 5'-PTO modified.](image-url)
after 1 min, which corresponds to an extension rate of 96 nucleotides per second. TthiPolB is therefore among the fastest thermophilic DNA polymerases described so far (Table 1).

**PCR performance**

TthiPolB was tested successfully in various PCR applications using templates such as genomic DNA, plasmid DNA, PCR products, *E. coli* colonies and yeast colonies. It performed remarkably well in cloning applications using degenerate primers including the RSO-PCR technique (Sarkar et al. 1993; Weber et al. 1998), even when commercial DNA polymerases such as Taq or KOD failed to generate PCR products. Examples of PCR amplification are shown in Fig. 7.

**Conclusion**

The family B DNA polymerase gene of *T. thioreducens* encodes a 774 aa DNA polymerase and a 537 aa intein. The DNA polymerase is most active in 10 mM Tris pH 7.8–9.0 (measured at 25°C), 60 mM KCl, 1.5–3.0 mM MgCl₂, 0.1% Triton X-100 and at temperatures between 72 and 76°C (under laboratory conditions). It has a proofreading 3′–5′ exonuclease activity. Its fidelity is 1.7 times higher than that of Taq polymerase and within the range of other DNA polymerases from the *Thermococcus* genus while being significantly lower than the high-fidelity polymerase *Pfu*. Its thermal stability is somewhat lower than that of other archaeal DNA polymerases but higher than that of Taq polymerase. The apparent Michaelis constant for the dNTP parameter is the lowest among published values for thermostable DNA polymerases and the extension rate is among the highest. TthiPolB appears to be more comparable to Taq than to most commercially available archaeal DNA polymerases in terms of robustness, versatility and reliability, while exhibiting higher extension rate, fidelity and thermal stability. It could therefore advantageously replace Taq in most PCR applications. Some of the properties of TthiPolB are summarized in Table 1 and compared with published results for other thermostable DNA polymerases.

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