Hyperthermophilic Topoisomerase I from *Thermotoga maritima*

A VERY EFFICIENT ENZYME THAT FUNCTIONS INDEPENDENTLY OF ZINC BINDING*

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Topoisomerases, by controlling DNA supercoiling state, are key enzymes for adaptation to high temperatures in thermophilic organisms. We focus here on the topoisomerase I from the hyperthermophilic bacterium *Thermotoga maritima* (optimal growth temperature, 80 °C). To determine the properties of the enzyme compared with those of its mesophilic homologs, we overexpressed *T. maritima* topoisomerase I in *Escherichia coli* and purified it to near homogeneity. We show that *T. maritima* topoisomerase I exhibits a very high DNA relaxing activity. Mapping of the cleavage sites on a variety of single-stranded oligonucleotides indicates a strong preference for a cytosine at position −4 of the cleavage, a property shared by *E. coli* topoisomerase I and archaeal reverse gyrase. As expected, the mutation of the putative active site Tyr 288 to Phe led to a totally inactive protein. To investigate the role of the unique zinc motif (Cys-X-Cys-X16-Cys-X-Cys) present in *T. maritima* topoisomerase I, experiments have been performed with the protein mutated on the tetracysteine motif. Strikingly, the results show that zinc binding is not required for DNA relaxation activity, contrary to the *E. coli* enzyme. Furthermore, neither thermostability nor cleavage specificity is altered in this mutant. This finding opens the question of the role of the zinc-binding motif in *T. maritima* topoisomerase I and suggests that this hyperthermophilic topoisomerase possesses a different mechanism from its mesophilic homolog.

Hyperthermophilic organisms have evolved a variety of strategies to adapt their physiology to extreme environments. In particular, molecular machineries in charge of genetic information have to function at temperatures near the boiling point of water.

Topoisomerases, a class of enzymes specialized in manipulating DNA, are involved in the maintenance and control of DNA supercoiling. In mesophilic bacteria (*i.e. Escherichia coli*), the DNA supercoiling control is achieved mainly through the balance between negative supercoiling impelled by gyrase and DNA relaxation by topoisomerase I (1, 2). In hyperthermophilic organisms, this control appears particularly important for life at high temperatures. Indeed, at such temperatures, the opening of the double helix is favored, and the control of supercoiling is crucial to maintaining the stability of the DNA duplex while ensuring initiation of essential processes such as replication, transcription, and recombination.

Little information about the DNA supercoiling control in hyperthermophilic bacteria is available. To get a better view of this process, we used *Thermotoga maritima* as a model organism. In this bacterium, which grows at 80 °C, we have already described three topoisomerasers: a gyrase (3) and a topoisomerase I (4), as found in all other bacteria, but also a reverse gyrase (5) as found in hyperthermophilic archaea (6). This third topoisomerase is able to introduce positive supercoils into DNA. Surprisingly, in *T. maritima*, episomal DNA (3, 7) is negatively supercoiled as in *E. coli*, contrary to the hyperthermophilic archaea in which episomal DNA is positively supercoiled (8). Thus, in *T. maritima*, gyrase and topoisomerase I probably control the global level of DNA supercoiling as in mesophilic bacteria.

In the present work, we have focused on the study of the *T. maritima* topoisomerase I overexpressed in *E. coli* and used mutational analysis to explore some of its biochemical properties. The protein is composed of 633 amino acids. Sequence analysis (4, 9) has shown previously that this protein is clearly related to the topoisomerase IA subfamily, the representative of which is topoisomerase I from *E. coli*. Similarities are located particularly within the first 540 amino acids of the protein. In this region, all the conserved sequence motifs defined for the other members of the family are found in the *T. maritima* enzyme, and Tyr238 was proposed to be the active site tyrosine (4).

Here, we show that *T. maritima* topoisomerase I relaxes DNA at 75 °C with an unusually high specific activity compared with the enzyme from *E. coli* and is also able to decatenate nicked plasmid DNA networks. Analysis of the DNA cleavage reaction, which constitutes the first step of the topoisomerization cycle, shows a strong preference for cytosine at position −4 of the cleavage point as is the case for *E. coli* topoisomerase I (10) and archaeal reverse gyrase (11, 12). We have also confirmed that Tyr238 is the active site residue involved in this reaction. Interestingly, although relaxation is highly sensitive to temperature, the cleavage step by *T. maritima* topoisomerase I is very efficient at 24 °C.

Examining the C-terminal part of the protein including the last 93 amino acids, no similarity with the other topoisomerases of the family was found (4), except a putative zinc-binding motif with the structure Cys-X-Cys-X16-Cys-X-Cys (Fig. 1). This type of tetracysteine motif is present in the C-terminal part of most bacterial topoisomerases I sequences with a variable number of copies of up to four. Interestingly, all of these motifs share a significant similarity (Fig. 1). In *E. coli* topo-
isomerase I, it has been shown that zinc binding to the three tetracysteine motifs present in the protein is required for DNA relaxation activity (13, 14). Here we show that, contrary to the case of E. coli, the unique zinc-binding motif of T. maritima topoisomerase I is not required for the topoisomerization reaction. Indeed, the enzyme mutated on the tetracysteine motif loses the ability to bind zinc but retains wild type activities.

The function of the zinc motif in T. maritima topoisomerase I remains unknown, but our results support the idea that the mechanism of the hyperthermophilic topoisomerase I may be somewhat different from that of its E. coli mesophilic homolog.

EXPERIMENTAL PROCEDURES

Cloning and Protein Expression—A DNA fragment encoding T. maritima topoisomerase I was prepared by polymerase chain reaction using genomic DNA as template, Fnu DNA polymerase (Expand High Fidelity kit, Roche Molecular Biochemicals) and the following primers: 5’-AGAGGGTGGAGGAGTCCGGACAGAAAGCTTCGGCTGGAAAGGAAATGAGG-3’ as forward primer containing a BamHI restriction site and 5’-TCTACCCGAGTGTGACTACAGAGCCTTTTATCCGTT-3’ as reverse primer containing a SalI restriction site. Polymerase chain reaction products were inserted into the pET29b expression vector (Novagen) at the BamHI/SalI sites, and constructs were checked by DNA sequencing.

The resulting expression plasmid was transformed into E. coli BL21(DE3). A colony of transformed cells was picked and grown at 37 °C in LB medium with 30 µg/ml kanamycin until A600 reached 0.5–0.6. Induction was carried out with 1 mM isopropyl-1-thio-β-D-galactopyranoside for 3 h at 37 °C (A600 1.0–1.2), and the cells were harvested and stored at –80 °C until required. About 15 g of cells were obtained from 4 liters of culture. The same procedure was followed to express the mutant enzymes.

Protein Purification—Frozen cells were resuspended in 10 volumes of buffer A (50 mM Tris-HCl, pH 7.5, 0.1 mM NaCl), lysed by sonication, and centrifuged at 40,000 × g for 20 min at 4 °C. To remove DNA, polymer F was added to the supernatant to a final concentration of 0.5%. After stirring for 1 h, the solution was centrifuged for 1 h at 90,000 × g. Ammonium sulfate (70% saturation) was added to the supernatant, and the solution was stirred overnight at 4 °C before centrifugation (24,000 × g for 30 min). The pellet was dissolved in 7 ml of buffer A and dialyzed extensively against the buffer containing 50 mM Tris-HCl, pH 8, 0.5 mM DTT, and 100 µM BSA. After 30 min of incubation at 75 or 50 °C (or as otherwise indicated), the reaction was stopped by adding 2 µl of stop solution containing 50 mM EDTA, 2.5% SDS, 25% glycerol, and 0.2% bromphenol blue. Samples were electrophoresed through 1.2% agarose gel in TEP buffer (90 mM Tris-phosphate, pH 8.0, 2 mM EDTA) for 2.5 h at 4 V/cm. Gels were stained with 1 µg/ml ethidium bromide and photographed under UV light.

Decatenation of Catenated Kinetoplast Minicircular DNA (kDNA)—The reaction mixture was identical to that used in the DNA relaxation assay except that negatively supercoiled pTZ 18 was replaced by 200 ng of kinetoplast DNA (TopoGEN). The activity was measured using serial dilutions of enzymes. Incubations were performed for 30 min at 75 °C as described in the paragraph above. The incubation products were analyzed by treatment with 0.5 units of XhoI (New England Biolabs) for 30 min at 37 °C. For the control of kDNA decatenation with human topoisomerase II (TopoGEN), 2 units of enzyme and 1 mM ATP were added to the reaction mixture, and the incubation was performed at 37 °C for 30 min. After the addition of the stop solution, samples were electrophoresed and stained as described above. When required, 1 µg/ml ethidium bromide was added in gel and electrophoresis buffer.

Oligonucleotides Radiolabeling—Oligonucleotides were synthesized and purified by MWG Biotech. 10 pmol were 5’-end-labeled by 5 units of T4 polynucleotide kinase (New England Biolabs) in the presence of 20 µCi of [γ-32P]ATP in 10 µl of the buffer recommended by the manufacturer. After 30 min of incubation at 37 °C, the reaction was stopped by heating the samples at 75 °C for 15 min. The oligonucleotide size standards were labeled (Amersham Pharmacia Biotech) and labeled following the manufacturer’s protocol.

Oligonucleotide Cleavage Assay—0.1 pmol of 5’-end-labeled oligonucleotide was incubated with 1 pmol of T. maritima topoisomerase I in a buffer containing 50 mM Tris-HCl, pH 8, 0.5 mM DTT, 30 µg/ml BSA, and 60 mM NaCl for 15 min at 75 °C (unless otherwise indicated) in a final volume of 10 µl. The reaction was stopped by adding 10% SDS and 0.5 µl of loading buffer (97.5% formamide, 10 mM EDTA, 0.3% bromphenol blue, 0.3% xylene cyanol). The samples were denatured by heating at 95 °C for 3 min and loaded onto a 15% polyacrylamide gel (19:1) containing 50% (w/v) urea in TBE buffer (90 mM Tris, 90 mM boric acid, 1 mM EDTA). The gel was electrophoresed at 1300 V for 2 h and autoradiographed. Quantification was performed with the ImageQuant v1.2 software.

RESULTS

Overexpression and Purification of T. maritima Topoisomerase I and Its Mutants—After amplification of the T. maritima topoisomerase I gene by polymerase chain reaction from genomic DNA and subcloning into the pET29b expression vector, the recombinant protein was expressed in E. coli. Expression and purification steps are summarized in Fig. 2. Induction with isopropyl-1-thio-β-D-galactopyranoside at 37 °C led to the production of an additional protein migrating to the expected molecular mass of the recombinant T. maritima topoisomerase.

Zinc Content Measurements—The zinc content of wild type and mutant proteins was determined both by a colorimetric method and flame atomic absorption spectroscopy. The colorimetric assay was conducted as follows. A solution of 2.5 mM p-hydroxy(mercuri)benzenesulfonate (PMPS) was added by increasing of 1 µl to a 500 µl sample of 10 µM of enzyme in 10 mM Tris-HCl, pH 7.5, 200 mM NaCl, 5% glycerol containing 0.1 mM 4-(2-pyridylazo)resorcinol (PAR). Zinc release from the enzyme was monitored by absorbance at 500 nm of the PAR-Zn(II) complex for which molar absorptivity is 6.6 × 10^4 M^-1 cm^-1 (16). The zinc content determination using flame atomic absorption was performed by using a Varian AA775 spectrometer. The protein samples (49 µl) were mixed against a buffer containing 10 mM Tris-HCl, pH 7.5, 0.1 mM EDTA, and 150 mM KCl with a protein concentration of 0.5 µM.

DNA Relaxation Assay—The reaction mixture (10 µl) contained 50 mM Tris, pH 8.0, 0.5 mM DTT, 30 µg/ml BSA, 10 mM MgCl2, 120 mM NaCl, and 200 ng of negatively supercoiled plasmid DNA (pTZ 18). The reaction was initiated by adding purified topoisomerase I, which was diluted in a buffer containing 50 mM Tris-HCl, pH 8.0, 1 mM DTT, and 100 µM BSA. After 30 min of incubation at 75 or 50 °C (or as otherwise indicated), the reaction was stopped by adding 2 µl of stop solution containing 50 mM EDTA, 2.5% SDS, 25% glycerol, and 0.2% bromphenol blue. Samples were electrophoresed through 1.2% agarose gel in TEP buffer (90 mM Tris-phosphate, pH 8.0, 2 mM EDTA) for 2.5 h at 4 V/cm. Gels were stained with 1 µg/ml ethidium bromide and photographed under UV light.

The abbreviations used are: BSA, bovine serum albumin; PMPS, p-hydroxy(mercuri)benzenesulfonate; PAR, 4-(2-pyridylazo)resorcinol; DTT, dithiothreitol; kDNA, kinetoplast minicircular DNA.

The data shown are representative of at least three independent experiments.

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I (76 kDa) on SDS-polyacrylamide gel (lane 2). A 15-min heating step at 75°C led to the elimination of most of the E. coli proteins, which precipitate at this temperature, whereas the thermophilic topoisomerase remained soluble (lane 4). This fraction was further purified on a heparin-Sepharose column, and topoisomerase I eluted from the column at around 0.6 M NaCl. The eluate contained nearly pure topoisomerase (lane 5).

To eliminate traces of a remaining nuclease contaminant, we finally performed an exclusion gel chromatography (Superdex S200). The resulting purified protein is shown in lane 6. We obtained about 2–3 mg of T. maritima topoisomerase I from 1 liter of culture.

To determine the functional relevance of the predicted zinc-binding domain described in Fig. 1, the two first cysteines of the tetracysteine motif were replaced by two alanines (C559A/C561A mutant) using site-directed mutagenesis, thus preventing a potential coordination of a zinc atom by the cysteine residues. We also performed a mutagenesis on Tyr288 of the active site (Y288F mutant) to produce an inactive form of the enzyme used as a control. The two mutant proteins were purified to near homogeneity (Fig. 2, lanes 7 and 8) by the same procedure used to purify the wild type enzyme.

**Fig. 1.** Sequence alignment of tetracysteine motifs in 19 bacterial topoisomerases I. The alignment was performed with Clustal X software. When a topoisomerase sequence contains more than one tetracysteine motif, each of them is numbered (from 1 to 4). The positions of the motifs in the sequences are indicated. Cysteines are boxed in red, and amino acids that are similar are boxed in blue. Organism names and their NCBI gene identification numbers are as follows: Tm, Thermotoga maritima, 1174737; Fi, Fervidobacterium islandicum, 7531209; Hi, Hemophilus influenzae, 1174735; Vc, Vibrio cholerae, 11387201; Ec, Escherichia coli, 730966; Ba, Buchnera aphidicola, 11135225; Pa, Pseudomonas aeruginosa, 14423974; Bs, Bacillus subtilis, 730965; Bh, Bacillus halodurans, 11135320; Tp, Treponema pallidum, 6094493; Hp, Helicobacter pylori J99, 7531219; Mm, Mycoplasma genitalium, 7190053; Mg, Mycoplasma galliseptium, 14423975; Cj, Campylobacter jejuni, 14423978; Nm, Neisseria meningitidis, 7225334; Mp, Mycoplasma pneumoniae, 2501239; Uu, Ureaplasma urealyticum, 10956714.

**Fig. 2.** Purification of recombinant T. maritima topoisomerase I wild type and mutant enzymes. Aliquots of the different steps of the purification of the wild type enzyme were analyzed by 9% SDS-polyacrylamide gel electrophoresis (lanes 1–6). Lane 1, uninduced cells; lane 2, induced cells; lane 3, crude extract; lane 4, extract after heating; lane 5, heparin-Sepharose eluate; lane 6, purified Superdex S200 fraction; lane 7, purified fraction of C559A/C561A mutant enzyme; lane 8, purified fraction of Y288F mutant enzyme. Lane M, molecular mass markers.
wavelength and a maximal absorbance of 0.658 for the wild type enzyme, the calculated stoichiometry was around one zinc atom (0.98) per molecule of topoisomerase I. Similar results were obtained by using the atomic absorption spectroscopy (results not shown).

By contrast, the zinc content analysis of the C559A/C561A mutant revealed no detectable zinc bound to the enzyme. From this analysis, we conclude that T. maritima topoisomerase I is indeed able to coordinate one zinc atom and that alteration of the zinc-binding domain totally abolishes the ability of the protein to bind zinc.

Conformational Changes in the Double Cysteine Mutant Compared with Wild Type Enzyme—To investigate the potential conformational changes that could affect the mutant proteins, fluorescence emission spectra upon excitation at 295 nm of wild type and mutant enzymes were compared (Fig. 4). At this wavelength, the fluorescence essentially monitors the local environment of tryptophan residues. Four tryptophans are present in T. maritima topoisomerase I at positions 94, 156, 365, and 410 (4).

The wild type enzyme fluorescence spectrum is characterized by a maximum emission wavelength at 335 nm (Fig. 4, curve a). Whereas no significant change was observed in the emission fluorescence spectrum of the active site Y288F mutant (curve b), the spectral properties of the C559A/C561A mutant were significantly affected. A red shift from 335 to 344 nm in the fluorescence spectrum of the active site Y288F mutant (curve d) shows that as little as 0.1 fmol of topoisomerase I is sufficient to completely transform 100 fmol of the plasmid DNA after 30 min of incubation at 75 °C. This very high activity level is at least 100-fold greater than that of E. coli topoisomerase I (18–20). The optimal temperature of incubation for the assay was around 75 °C (Fig. 5B), although a significant activity was still observed at 37 °C when 1 pmol of topoisomerase I was added to the reaction mixture (results not shown).

An examination of the effect of mutations on the relaxation activity showed that the Y288F mutant is totally inactive (Fig. 5A) even with an amount of protein 10,000-fold higher than that used to observe activity of the wild type enzyme. This result indicates that Tyr288, defined as the active site tyrosine from sequence analysis, is probably the amino acid involved in the DNA cleavage step of the topoisomerase reaction.

An unexpected result was obtained with the C559A/C561A mutant, which exhibited an activity similar to that of the wild type enzyme (Fig. 5A) with the same optimal temperature for activity (Fig. 5B). This result contrasts with E. coli topoisomerase I for which the apoenzyme without zinc is totally inactive in DNA relaxation (13, 14). Moreover, the thermostability of the C559A/C561A mutant is comparable with that of the wild type enzyme (Fig. 6). In these assays, enzymes were incubated at 80 °C for 30 min prior to measurement of the remaining DNA relaxation activity. To increase the accuracy of measurements, the test temperature was lowered to 50 °C allowing an increase in the amount of enzyme (i.e. reduction of the dilution factors). The results show that both wild type and C559A/C561A mutant enzymes remained fully active after 30 min at 80 °C, suggesting that zinc binding is not involved in the thermostability of T. maritima topoisomerase I.

T. maritima Topoisomerase I Decatenates Nicked Kinetoplast DNA—In regard to the relaxation results, we were interested in analyzing other enzymatic properties of the T. maritima topoisomerase I and comparing them with those of the mutant enzymes. Besides relaxation, another test for type I topoisomerases is their ability to decatenate nicked DNA. In E. coli, topoisomerase I is able to decatenate double-stranded DNA interwoven dimers provided that one of the circular DNA molecules contains a nick (21). In addition, another type I topoisomerase present in this organism, topoisomerase III, was identified as a potent decatenase (22) efficient in the resolution of plasmid replication intermediates in vitro (23). In the case of T. maritima, an analysis of the whole genome sequence (24) did not reveal the presence of a topoisomerase III equivalent. We therefore investigated the ability of T. maritima topoisomerase I and its mutant forms to decatenate kinetoplast DNA (Fig. 7), a substrate usually dedicated to topoisomerase II assays and never tested with a topoisomerase I. This particular DNA forms a network of interlinked minicircles in which a signifi-
A fraction can be nicked. When kDNA was incubated with \textit{T. maritima} topoisomerase I at 75 °C, a product appeared that migrated as nicked circles in an agarose gel containing ethidium bromide (Fig. 7A, lane 1). A fraction of kDNA was not transformed (material remaining in the well). As a control and to confirm the nature of the decatenation product observed, we performed a decatenation assay with human topoisomerase II, which is known to decatenate interlinked minicircles (25). The electrophoretic pattern (Fig. 7A, lane 2) showed that the totality of the kDNA was transformed and yielded two products, one of them migrating as nicked circles and the other exhibiting a higher electrophoretic mobility presumably corresponding to closed circular monomers. This assay confirmed the presence of both intact and nicked minicircles in the kDNA. The products of incubation of kDNA with \textit{T. maritima} topoisomerase I were further treated by \textit{XhoI} endonuclease, which cuts once by minicircle; linear monomers were obtained that migrated as the digestion product of kDNA by \textit{XhoI} (Fig. 7A, lanes 3 and 4). From these results, we conclude that \textit{T. maritima} topoisomerase I is able to decatenate interlinked DNA if a nick is present on the molecules to be decatenated.

Again, the decatenase activity of wild type and C559A/C561A mutant enzymes showed no significant difference (Fig. 7B). Nicked DNA circles were still produced by incubation with 0.2 pmol of either protein. As expected, the Y288F mutant was totally inactive (Fig. 7B).

\textbf{Cleavage by \textit{T. maritima} Topoisomerase I Shows a High Preference for Cytosine at Position -4 of the Cleavage Site}—In their catalytic mechanism, type I topoisomerases transiently cleave one DNA strand to permit strand passage through the break and then reseal the cleaved strand. Although all type IA topoisomerases cleave single-stranded DNA by linking to the 5’-phosphoryl end of cleaved DNA, the specificity of the cleavage sequence is somewhat variable. The derived consensus sequence is CTT for \textit{E. coli} topoisomerase III (26), CANNN for human topoisomerase III (27), ANN for yeast topoisomerase III (28), and CNNN for \textit{E. coli} topoisomerase I (10) and archaeal reverse gyrase (11, 12). We therefore investigated the cleavage of a single-stranded oligonucleotide at 75 °C by \textit{T. maritima} topoisomerase I. A 22-base oligonucleotide (22N, sequence shown in Fig. 8B), already described as a substrate for
E. coli topoisomerase I and topoisomerase III (29, 30), was labeled at its 5′ terminus and used as a substrate for T. maritima topoisomerase I. The results (Fig. 8) showed that the enzyme strongly cleaves two sites leading to the production of two labeled fragments of 11 and 17 nucleotides (Fig. 8A, lane 2). Cleavage positions are indicated by arrows on the substrate sequence (Fig. 8B). A minor cleavage site was also observed at position 14. In all cases, a cytosine was present in position −4 of the cleavage point, and the efficiency of cleavage appeared highly dependent on its presence. Indeed, when cytosine 8 or 14 was changed to thymine (oligonucleotides 22N C8T and 22N C14T; sequences and cleavage positions shown in Fig. 8B), the corresponding cleavage sites (positions 11 and 17, respectively) totally disappeared (Fig. 8A, lanes 4 and 6). Furthermore, when both cytosines 8 and 14 were mutated (oligonucleotide 22N C8T/C14T; sequence and cleavage positions shown in Fig. 8B) only several minor sites were observed (Fig. 8A, lane 8). Finally, we tested an oligonucleotide (21N RG; sequence shown in Fig. 8B) known to contain a strong cleavage site for Sulfolobus shibatae reverse gyrase (12). An 11-nucleotide labeled fragment was predominantly produced (Fig. 8A, lane 10). The corresponding cleavage position (Fig. 8B) was the same as the one found for reverse gyrase (12) and exhibited again a cytosine in position −4. Taken together, these results indicate that T. maritima topoisomerase I cleaves DNA with a strong preference for sites located 4 bases after a cytosine, as observed for E. coli topoisomerase I and archaeal reverse gyrase.

Guanine at Position −1 Decreases Cleavage Efficiency—Although cleavage by T. maritima topoisomerase I showed a high preference for cytosine in position −4 of the cleavage site, it appeared that all cytosines were not equivalent, since only two sites are cleaved almost exclusively in the 22N, whereas this oligonucleotide contains 5 cytosines. This prompted us to investigate the importance of the bases located at the −1 and +1 positions of the cleavage for their specificity. We therefore defined 11 oligonucleotides derived from 22N C8T (exhibiting only one cleavage site), in which the bases adjacent to the cleavage site were changed. The results are summarized in Table I. It appeared that the base in position +1 has very little influence on the cleavage efficiency. On the other hand, the presence of a guanine in position −1 decreased by 2–3-fold the level of cleavage. Similarly, in the case of the E. coli enzyme, guanine residues were not found at position −1 in 98% of the cleavage sites studied (10). This result could explain the absence of observable cleavage at position 20 of 22N, but other structural or sequence reasons may be advanced to explain the strong preference for only two sites, while five potential sites are possible. Nevertheless, we can propose the sequence 5′-CCNH↓N (H corresponding to A, C, or T) as a preferential sequence for cleavage by T. maritima topoisomerase I.

Cleavage Step of the Hyperthermophilic T. maritima Topoisomerase I Is Not Dependent on Temperature—We have shown that the DNA relaxation activity, which reflects the whole topoisomerization reaction, is highly dependent on temperature with an optimum around 70–75 °C (Fig. 5B). By contrast, when we tested the influence of temperature on the cleavage, which is the first step of the reaction, we observed that the cleavage efficiency was not significantly different in a range of temperature going from 24 °C to 75 °C (Fig. 9). This result suggests that the sensitive temperature step in the topoisomerization is not the trans-esterification reaction but more likely the strand passage event.

Cleavage Specificity Remains Unchanged in the Double Cysteine Mutant—To know whether the loss of the ability to bind zinc changes the specificity of the cleavage, and to confirm that Tyr288 of T. maritima topoisomerase I is the residue involved in the cleavage step, the cleavage pattern of the 22N oligonucleotide by the wild type T. maritima topoisomerase I was compared with mutant enzymes. The results (Fig. 10) showed that the Y288F mutant is totally inefficient in cleavage, attesting to the fact that Tyr288 is the catalytic amino acid of the active site. On the other hand, cleavage positions obtained with the C559A/C561A mutant were similar to those observed with the wild type enzyme. Preference for cytosine at position −4 was retained, and cleavage intensities were comparable. Alteration of the zinc-binding domain therefore does not affect the efficiency and specificity of the cleavage reaction.

![Fig. 7. Decatenation activity of wild type T. maritima topoisomerase I and mutant enzymes.](http://www.jbc.org/)

**A** kDNA (lane 1) or kDNA digested with XhoI (lane 2). Samples were electrophoresed through agarose gel containing 1 μg/ml ethidium bromide. Lane C corresponds to the kDNA control without protein. **B** A comparison of kDNA decatenation activities of the wild type enzyme, C559A/C561A mutant, and Y288F mutant enzymes. Activities were measured by serial dilutions of enzymes. Dilution factors of 1, 2, 4, and 10 correspond to 2, 1, 0.5, and 0.2 pmol, respectively.
DISCUSSION

This work describes the purification and biochemical properties of the hyperthermophilic topoisomerase I from *T. maritima* and two of its mutants. One was altered in the active site of the enzyme on the putative tyrosine involved in the cleavage step of DNA. The other was mutated on the two first cysteines of the tetracysteine motif (Cys-X-Cys-X_X-Cys-X-Cys) (Fig. 1) located in the C-terminal part of the protein. The thermophilic character of *T. maritima* enzyme and its mutants allowed us to obtain all of the proteins in large amounts and at a high degree of purity by introducing a heating step in the purification procedure.
Based on sequence similarities, it was proposed previously that topoisomerase I from *T. maritima* belongs to the *E. coli* topoisomerase I family (4, 9). The results presented here clearly confirm this hypothesis, even though significant differences in their properties are pointed out.

A striking result of our study was the exceptionally high DNA relaxation activity of *T. maritima* topoisomerase I, with a specific activity at least 100-fold greater than that of the enzyme from *E. coli*. One molecule of *T. maritima* topoisomerase I can catalyze about 500–1000 strand-passage events/min as compared with about 5 for *E. coli* (20). This suggests that a large number of strand passages might occur during a given topoisomerization cycle, as in the case of topoisomerases IB (about 1200 rotations/min for the vaccinia virus topoisomerase I) (31). For this family of topoisomerases, a “controlled rotation” mechanism has been proposed, in which the end of the cleaved strand that is non-covalently attached to the enzyme, rotates around the uncleaved strand in a controlled way (32).

Today, insufficient arguments are available to propose such a mechanism for *T. maritima* topoisomerase I. The reasons for such a high specific activity remain unknown, but it may be required in vivo to locally increase, with great efficiency, the DNA linking number and thus prevent the double-helix opening at high temperature.

Besides its DNA relaxation activity, *T. maritima* topoisomerase I exhibits an efficient decatenation activity of nicked microrings present in kinetoplastid DNA. In *E. coli*, topoisomerase III, rather than topoisomerase I, is efficient to decatenate multiply inter-twisted nicked dimers (22, 33). The lack of a topoisomerase III equivalent in *T. maritima* (24) suggests that in this organism, topoisomerase I could also be involved in the resolution of replicating daughter DNA molecules at the end of replication.

An examination of oligonucleotide cleavage by the wild type enzyme from *T. maritima* showed the constant presence of a cytosine in position –4 of the cleavage point, as is the case for *E. coli* topoisomerase I (10) and archaeobacterial reverse gyrase (11, 12). Although cleavage efficiency was not affected by the identity of the base in position +1, a guanine in position –1 clearly reduced cleavage intensity. Interestingly for a thermophilic enzyme, cleavage did not appear to be dependent on temperature, suggesting that this step of the topoisomerization cycle is not rate-limiting. We also show that Tyr288 is the amino acid involved in the trans-esterification reaction, because its mutation to phenylalanine totally abolishes cleavage.

The major differences between *T. maritima* and *E. coli* topoisomerases I are located in the C-terminal region of the enzymes (4) (9). Indeed, the *T. maritima* topoisomerase I C terminus is much shorter than the one from *E. coli* enzyme, and it contains a unique tetracysteine motif instead of three tetracysteines in the *E. coli* topoisomerase I. The present results show that this putative zinc-binding motif is functional because *T. maritima* topoisomerase I does coordinate one Zn(II). Moreover, the double cysteine C559A/C561A mutant was unable to bind zinc, and its fluorescence properties were modified, suggesting a change of conformation in the protein lacking zinc.

Surprisingly, analysis of the enzymatic properties of the C559A/C561A mutant showed that the relaxation as well as decatenation activities are similar to those of the wild type enzyme. Furthermore, the thermal stability and DNA cleavage properties are not affected in this mutant despite the loss of zinc. These results contrast with those found for *E. coli* topoisomerase I, in which zinc binding to the three tetracysteine motifs is required for DNA relaxation activity (13, 14) and disruption of the second zinc motif deeply affects thermal stability and cleavage specificity (18). Indeed, it was proposed that the zinc binding domain would contribute to the stabilization of the association of the 3‘-end of the cleaved DNA with topoisomerase I and would assist the passage of uncleaved strand across the break (14, 34). This model is difficult to imagine with *T. maritima* topoisomerase I, because we have showed that the unique zinc motif is not involved in the topoisomerase activity at least in vitro. Taken together, our results suggest that the two enzymes may have distinct ways of functioning.

The absence of involvement of the zinc-binding domain of *T. maritima* topoisomerase I in the topoisomerization reaction stresses the question of the putative role of this domain in other processes. The tetracysteine motif present in *T. maritima* topoisomerase I is close to those of 19 bacterial topoisomerases I examined (Fig. 1). Strikingly, all of these conserved motifs are located within the highly divergent C-terminal part of these enzymes. The similarity of the tetracysteine motif includes the high frequency of a Gly following Cys^2, the number of residues between Cys^2 and Cys^3 (15–17 residues), and a hydrophobic region just upstream of Cys^3 (Fig. 1). It was recently proposed that these tetracysteine motifs could be members of the zinc ribbon domain superfamily (35), also found in transcription factors TFII B, TFII S, TFII E large subunit (36–38), RNA polymerase II subunit 9 (RPB9) (39), adenovirus E1A transactivator (40), and bacteriophage T7 DNA primase (41). All of these proteins are part of multipartite-nucleic acid complexes, but the proposed function of the zinc-binding domain appears very variable. Some are involved in interactions with single-stranded DNA, RNA, or other proteins (40–43), whereas in other cases, the role of this motif remains unclear (44).

Thus, the question of the role of the zinc-binding motif in *T. maritima* topoisomerase I is largely open. One hypothesis could be that it interacts with other proteins of the replication or transcription machinery, given the involvement of topoisomerases in these processes. We are presently interested in searching for the potential protein partners of *T. maritima* topoisomerase I.

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Hyperthermophilic Topoisomerase I from *Thermotoga maritima*: A VERY EFFICIENT ENZYME THAT FUNCTIONS INDEPENDENTLY OF ZINC BINDING

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