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Characterization of the DNA Gyrase from the Thermoacidophilic Archaeon *Thermoplasma acidophilum*

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*Thermoplasma acidophilum* is sensitive to the antibiotic drug novobiocin, which inhibits DNA gyrase. We characterized DNA gyrases from *T. acidophilum* strains in vitro. The DNA gyrase from a novobiocin-resistant strain and an engineered mutant were less sensitive to novobiocin. The novobiocin-resistant gyrase genes might serve as *T. acidophilum* genetic markers.

The members of the domain *Archaea* are phylogenetically distinct from those of *Bacteria* and *Eukarya* (21). *Thermoplasma acidophilum* is a thermoacidophilic archaeon which was originally isolated from a self-heating coal refuse pile (5). This archaeon grows optimally at pH 1.8 and 56°C, has no cell wall, and is classified as a member of the *Euryarchaeota* (5). Strains of this archaeon have also been isolated from Japanese hot springs (22). Presently, there are no genetic tools that can be used to manipulate *T. acidophilum* genomes. A coumarin, novobiocin, competitively inhibits ATP binding to type IIA topoisomerase (3). Novobiocin inhibits *T. acidophilum* growth (5, 22), and a novobiocin-resistant strain (strain HO-62NIC) has been isolated (22). Therefore, it seems that a novobiocin-resistant DNA gyrase could be used as a genetic marker during the development of transformation methods.

Type II topoisomerases cleave both strands of a DNA duplex and pass a second duplex through the double-stranded break (3). The type II topoisomerases are classified into two types: type IIA, e.g., DNA gyrase and topoisomerase IV (TopoIV); and type IIB, e.g., topoisomerase VI (TopoVI). DNA gyrase introduces negative supercoils into closed circular duplex DNA in an ATP-dependent fashion. This supercoiling activity is essential for DNA replication, transcription, and recombination (3). Gyrase also relaxes supercoiled DNA in an ATP-independent manner (14). TopoIV decatenates interlinked daughter chromosomes after DNA replication and can relax positive and negative DNA supercoils (19). TopoVI also has relaxation and decatenation activities (2). In the domain of *Archaea*, TopoVI is often present and DNA gyrase is occasionally found (6). Only one kind of type II topoisomerase is found in *Thermoplastomates*, of which *T. acidophilum* is a member. Gadelle et al. suggested that the *T. acidophilum* type II topoisomerase is a DNA gyrase based on its phylogenetic position (6). However, to date, nothing is known about the biochemical characteristics of any archaeal DNA gyrase. Therefore, we have cloned, expressed, purified, and characterized a novobiocin-sensitive *T. acidophilum* strain and two resistant forms of DNA gyrase.

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**Bacterial strains and growth conditions.** *T. acidophilum* 122-1B2 was kindly provided by D. G. Searcy. *T. acidophilum* strains HO-01, HO-54, and HO-121 and the novobiocin-resistant strain HO-62NIC were isolated by Yasuda et al. (22). *T. acidophilum* culture medium was prepared as described previously (22).

**Sequencing the *T. acidophilum* HO-62NIC gyrase gene.** The archaeal gyrase B sequences were aligned automatically using the program Clustal X, version 1.81 (18), and then optimized manually. Degenerate primers were synthesized based on conserved nucleotide sequences identified using these alignments (Table 1). A partial gyrase B gene sequence was amplified by nested PCR using HO-62NIC genomic DNA. PCR was performed first with the Gyr-1F and Gyr-1R primers and then with the Gyr-2F and Gyr-2R primers. The PCR product was cloned and sequenced.

A restriction map, flanking the partial gyrase B gene, was constructed using Southern analysis. Based on the physical map, genomic HO-62NIC DNA was digested with either BamHI or SalI, and then the two types of linear fragments were each self-circularized. Inverse PCR was performed using the self-ligated products as templates and using the following primer pairs: L-gyr-1F and L-gyr-2R, L-gyr-2F and L-gyr-2R, L-gyr-A1 and L-gyr-A4, or L-gyr-A2 and L-gyr-A4. The PCR products were cloned and sequenced.

**Construction of gyrase A and B expression vectors.** The gyrase A (Ta1054) and gyrase B (Ta1055) genes of *T. acidophilum* 122-1B2 (referred to as gyrA*TA* and gyrB*TA* hereafter) and the gyrase A and B genes of *T. acidophilum* HO-62NIC (sequenced as described above and referred to as gyrA*62* and gyrB*62* hereafter) were amplified from genomic DNA by PCR. The primers used were gyrA-EX-F and gyrA-EX-RAR for gyrA*TA*; gyrA-EX-F and gyrA-EX-62R for gyrA*62*; gyrB-EX-F and gyrB-EX-R for gyrB*TA* and gyrB*62*. The products were cloned into pCR4-TOPO vectors (Invitrogen, Tokyo, Japan) and sequenced. The gyrase genes were then subcloned into pET-28a (+) at the NdeI and HindIII or EcoRI sites.

A PCR-based mutagenesis technique was used for site-directed mutagenesis of the gyrase B subunit (13). The primers used for the mutagenesis reactions were TABR136H and 62BR136H for gyrB*TA* and 62BR136H for gyrB*62*. T7P and T7T primers were used as the upstream and downstream primers, respectively.


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Construction of expression vectors

GyrB62 were heat-treated at 60°C for 20 min. Those of GyrB(R136H)TA and GyrB(R136H)62 were also treated for 20 min but at 55°C. The precipitates were removed by centrifugation at 82,800 × g for 20 min. The supernatants containing recombinant GyrA TA, GyrB TA, GyrA R136H, or GyrB R136H were heat-treated at 60°C for 20 min. Those of GyrB(R136H) TA and GyrB(R136H) R136H were also treated for 20 min but at 55°C. The precipitates were removed by centrifugation at 82,800 × g for 20 min. Each subunit was purified by column chromatography. Each supernatant was individually applied to Ni²⁺-NTA (Amersham Biosciences, Tokyo, Japan) equilibrated with buffer A. The columns were each washed with 3 column volumes of buffer A, and the glyrase subunits were eluted using a gradient of 10 to 300 mM imidazole in buffer B. The samples were each eluted with a linear elution gradient of 50 to 400 mM NaCl in buffer B. The sample fractions were combined and concentrated using a Vivaspin 1 20-ml concentrator (Vivascience, Hannover, Germany). The glyrase fractions were dialyzed against 20 mM KPi, pH 7.4, 50 mM NaCl, 10 mM 2-mercaptoethanol, and 50% glycerol and stored at −80°C.

DNA supercoiling, relaxation, and decatenation assays. DNA supercoiling activity was assayed by monitoring the concentration of relaxed pBR322 to its supercoiled form (7). Complete reaction mixtures (30 μl) contained 35 mM piperazine-1, 4-bis(2-ethanesulphonic acid), pH 6.5, 0.14 mM Na₂EDTA, 5 mM dithiothreitol, 3 mM spermidine, 0.01% (wt/vol) bovine serum albumin, 9.5 μg/ml E. coli tRNA, 1.4 mM ATP, 6 mM MgCl₂, 300 ng relaxed pBR322, and 2 units each of the glyrase A and B subunits. Reaction mixtures were incubated at 50°C. Reactions were terminated by addition of 30 μl phenol-chloroform. Twenty microfilters of each reaction mixture were mixed with 2 μl of loading buffer (1% sodium dodecyl sulfate, 50% glycerol, and 0.05% bromophenol blue) and then electrophoresed on 1% agarose gel that had been stained with 0.5 μg/ml ethidium bromide. The DNA was visualized by fluorescence and photographed with a digital camera system.

**TABLE 1. Primers**

<table>
<thead>
<tr>
<th>Primer name</th>
<th>Sequence*</th>
<th>Purpose</th>
</tr>
</thead>
<tbody>
<tr>
<td>GyrA-EX-F</td>
<td>GGAATTCCTAGATGGAGAAAGACGATTAAGTTG</td>
<td>Construction of expression vectors</td>
</tr>
<tr>
<td>GyrA-EX-R</td>
<td>ATAAAGCTTCACTTGCGAAGCGGCA</td>
<td></td>
</tr>
<tr>
<td>GyrB-EX-F</td>
<td>GGAATTCCTAGATGGAGAAAGACGATTAAGTTG</td>
<td></td>
</tr>
<tr>
<td>GyrB-EX-R</td>
<td>ATAAAGCTTCACTTGCGAAGCGGCA</td>
<td></td>
</tr>
<tr>
<td>Gyra-EX-F</td>
<td>GGAATTCCTAGATGGAGAAAGACGATTAAGTTG</td>
<td></td>
</tr>
<tr>
<td>Gyra-EX-R</td>
<td>ATAAAGCTTCACTTGCGAAGCGGCA</td>
<td></td>
</tr>
<tr>
<td>GyrA-EX-TAR</td>
<td>ATAAAGCTTCACTTGCGAAGCGGCA</td>
<td></td>
</tr>
<tr>
<td>GyrB-EX-TAR</td>
<td>ATAAAGCTTCACTTGCGAAGCGGCA</td>
<td></td>
</tr>
<tr>
<td>Gyra-EX-62R</td>
<td>ATAAAGCTTCACTTGCGAAGCGGCA</td>
<td></td>
</tr>
<tr>
<td>GyrB-EX-62R</td>
<td>ATAAAGCTTCACTTGCGAAGCGGCA</td>
<td></td>
</tr>
</tbody>
</table>

* Restriction recognition sites are underlined, and mutated codons are shown in boldface.
resed in a 0.8% agarose gel (135 by 135 by 10 mm) equilibrated with Tris-borate-EDTA buffer (16). One unit of enzyme activity is defined as the amount of gyrase that converts one-half of the initial 300 ng of relaxed pBR322 to the supercoiled form in 30 min at 50°C. DNA relaxation and decatenation activities were assayed under the same conditions as those for assay supercoiling, except that the substrates were supercoiled pBR322 or kinetoplast DNA, respectively, and the relaxation

<table>
<thead>
<tr>
<th>Residues in supercoiled pBR322...</th>
<th>175</th>
</tr>
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<tbody>
<tr>
<td>...relaxed pBR322 in the...</td>
<td>168</td>
</tr>
<tr>
<td>...supercoiled form in 30 min at 50°C. DNA relaxation and decatenation activities were assayed under the same conditions as those for assay supercoiling, except that the substrates were supercoiled pBR322 or kinetoplast DNA, respectively, and the relaxation...</td>
<td>168</td>
</tr>
</tbody>
</table>

![Fig. 2](image-url)
times and the amounts of enzyme differed. (See the legend of Fig. 3 for additional details.)

Effects of novobiocin on the growth of \textit{T. acidophilum} strains.\textit{Novobiocin is a potent growth inhibitor of \textit{T. acidophilum} (5). The novobiocin 50% inhibitory concentrations (IC$_{50}$) for the \textit{T. acidophilum} 122-1B2, HO-01, HO-54, and HO-121 strains are about 0.01 µg/ml (Fig. 1). However, the IC$_{50}$ for HO-62N1C is 1.0 µg/ml. Therefore, the effect of novobiocin on the growth of HO-62N1C is about 100-fold smaller than it is on the other strains.\textit{Cloning and sequencing of the \textit{T. acidophilum} HO-62N1C gyrase genes.} The positions of the gyrase subunit genes have been identified in the genome of \textit{T. acidophilum} 122-1B2 (15). Novobiocin inhibits ATP binding to type II topoisomerases (1, 8). To determine whether the resistance to novobiocin by HO-62N1C is caused by a decreased binding affinity between the antibiotic and the \textit{gyr} subunits, we purified recombinant 122-1B2 and HO-62N1C gyrases and assayed their supercoiling activities.

A comparison of the 122-1B2 and HO-62N1C sequences shows that the number of mutated amino acid residues is 8 for the \textit{A} subunit and 10 for the \textit{B} subunit (Fig. 2). A R136H mutation in the ATP-binding site of the \textit{E. coli} gyrase \textit{B} subunit causes novobiocin resistance, as does a R138H mutation in the "\textit{Haloferax alicantei}" \textit{B} subunit (9, 10). However, the corresponding amino acid sequences are the same for the ATP-binding regions of the \textit{B} subunits of 122-1B2 and HO-62N1C. The spontaneous mutation S127L, which is present in the \textit{Streptococcus pneumoniae} gyrase \textit{B} subunit, also causes novobiocin resistance (11). GyrB$_{TA}$ has a valine and GyrB$_{62}$ has a leucine at the corresponding position (residue 119) (Fig. 2). Accordingly, the mutation at position 119 may be responsible for the increased novobiocin resistance found for HO-62N1C. However, site-directed mutagenesis is needed to validate this proposal.

Construction of expression vectors for \textit{T. acidophilum} DNA gyrase and purification of recombinant His-tagged GyrA$_{TA}$, GyrB$_{TA}$, GyrA$_{62}$, and GyrB$_{62}$. For expression of the archaellae gyrase subunits, all genes were first PCR amplified and then cloned into plasmid pT7-1A(+), which is a T7 promoter-based expression vector. Hexahistidine tags were fused to the N termini of all expressed subunits. The recombinant gyrase subunits of \textit{T. acidophilum} 122-1B2 (GyrA$_{TA}$ and GyrB$_{TA}$) and GyrA$_{62}$ were expressed in \textit{E. coli} BL21(DE3) codon Plus-RIL cells, while GyrB$_{62}$ was expressed in \textit{E. coli} Rosetta cells. All expressed subunits were purified using heat treatment, Ni-chelating chromatography, and ion-exchange chromatography. From approximately 1 g of wet \textit{E. coli} cells, 0.9 to 1.6 mg of soluble protein was purified, with the exception of GyrB$_{62}$, for which 0.3 mg was purified (Table 2).

\begin{table}[h]
\centering
\begin{tabular}{|c|c|c|c|c|}
\hline
Strain & Subunit & Wet wt of cells (g) & Amt of purified protein (mg) & Sp act (U/mg$^a$) & IC$_{50}$ (µg/ml)$^b$ \\
\hline
TA & GyrA & 4.3 & 5 & $5 \times 10^4$ & 0.97 \\
 & GyrB & 7.3 & 8.1 & $3.5 \times 10^4$ & 0.57 \\
 & GyrBR136H & 12.2 & 20 & $1 \times 10^4$ & 1.6 \\
HO-62N1C & GyrA & 9.9 & 9 & $5 \times 10^4$ & \\
 & GyrB & 15.0 & 4.3 & $3.5 \times 10^4$ & 28 \\
 & GyrBR136H & 10.9 & 17 & $2 \times 10^4$ & 130 \\
\hline
\end{tabular}
\caption{Purification of DNA gyrase}
\end{table}

\textbf{FIG. 3.} Assays of \textit{T. acidophilum} gyrase activities. (A) Supercoiling activity. Supercoiling was measured in the absence of enzyme (-E); in the presence of all reaction components (all) as described in the text; and in the absence of one of the following components: GyrA$_{TA}$ (-GyrA); GyrB$_{TA}$ (-GyrB); ATP (-ATP); Mg$^{2+}$ (-Mg); or spermidine (-spe). The substrate was relaxed pBR322, and the reaction time was 5 minutes. The labels on the left mark the positions of relaxed (R) and supercoiled (S) pBR322 DNA. (B) Relaxation activity. Relaxation was measured in the absence of enzyme (-E); in the presence of all components, including 10 units each of GyrA$_{TA}$ and GyrB$_{TA}$, and ATP (all); and in the presence of all components except ATP (-ATP). The substrate was supercoiled pBR322, and the reaction time was 2 hours. The labels on the left mark the positions of relaxed (R) and supercoiled (S) pBR322 DNA. (C) Decatenation activity. Decatenation was measured in the absence of enzyme (-E); in the presence of all components including 4 units each of GyrA$_{TA}$ and GyrB$_{TA}$ (all); and in the absence of one of the following components: ATP (-ATP); GyrA$_{TA}$ (-GyrA); and GyrB$_{TA}$ (-GyrB). The substrate was intertwined kinetoplast DNA, and the reaction time was 30 min. The labels on the left mark the positions of relaxed (R) and supercoiled (S) monomeric kinetoplast DNA.
Gyrase supercoiling, relaxation, and decatenation activities.

The supercoiling activity of DNA gyrase requires a divalent cation, such as Mg\(^{2+}\), and is stimulated by spermidine (14). We found that the supercoiling activity of the *T. acidophilum* gyrase also required ATP and Mg\(^{2+}\) and was stimulated by spermidine when assayed with relaxed pBR322 DNA (Fig. 3A). Both A and B subunits are required for supercoiling. The optimal pH is 6.5, which is near the internal pH of *T. acidophilum* (17), and the optimal concentrations of Mg\(^{2+}\) and spermidine are 2 to 10 and 3 mM, respectively (data not shown).

Gyrases usually catalyze the relaxation of supercoiled DNA given the same conditions as used for supercoiling, except that ATP is not necessary; however, the relaxation reaction is about 20 to 40 times less efficient than supercoiling (14). We found that, within 2 hours, relaxation of negatively supercoiled DNA by 10 units of gyrase was detectable and that relaxation did not require ATP (Fig. 3B). ATP-dependent decatenation activity was also detected when assayed with kinetoplast DNA (Fig. 3C). Therefore, the activities of the archaeal gyrase are similar to those of bacterial gyrases (14).

These results and the absence of a TopoVI-like gene in the *T. acidophilum* genome support the possibility that the gyrases of thermoplasmalates perform several functions, including decatenation (6). Alternatively, the *Thermoplasmales* TopoIIIs, e.g., Ta0063 of *T. acidophilum*, may be responsible for decatenation of daughter chromosomes; the *E. coli* TopoIII has decatenation activity (12). However, to date, it has not been shown that any archaeal TopoIII has decatenation activity. Assigning an archaeal decatenation activity to one or more enzymes is an area open to investigation.

**Inhibition of gyrase DNA supercoiling activity by novobiocin.** To determine whether the affinity of GyrB62 towards novobiocin is less than that of GyrB TA, we tested the effect of novobiocin on the supercoiling abilities of the wild-type gyrase and a gyrase composed of GyrA TA and GyrB62 (Fig. 4). Because GyrA TA was present in both holoenzymes and because the reactions contained the same number of subunits (expressed as units of gyrase activity), any differences in novobiocin sensitivity could be attributed to GyrB62. The IC\(_{50}\), which is the drug concentration that inhibits supercoiling by 50%, is 0.57 μg/ml for the wild-type gyrase and 28 μg/ml for the gyrase containing GyrB62 (Fig. 4 and Table 2). Therefore, an increase in novobiocin resistance can be attributed to a mutation in GyrB62.

If a novobiocin-resistant gyrase is to be a reliable genetic marker, then spontaneous novobiocin-resistant mutations arising in the host cell gyB gene must not be allowed to produce false positives during transformation assays. Introduction of a second novobiocin-resistant mutation that further increases the level of antibiotic resistance would diminish the possibility that false positives would be selected. An R136H mutation in the B subunit of the *E. coli* gyrase increases the novobiocin IC\(_{50}\) 10-fold (4). We engineered this mutation into the genes for GyrB TA and GyrB62, and the resulting proteins are named, respectively, GyrB(R136H) TA and GyrB(R136H)62 (Fig. 2). The specific activities of both mutants are less than that of the wild-type archael gyrase, which is a phenomenon also found for the *E. coli* gyrase mutant (Table 2) (4). When the R136H mutation is present in GyrB(R136H) TA, the resistance to novobiocin is 2.8-fold greater than that found for the wild-type gyrase, and when the mutation is present in GyrB(R136H)62, the resistance is 4.6-fold greater than that found for the gyrase containing GyrB62.

GyrB(R136H)62 has the greatest resistance to novobiocin. The GyrB(R136H)62 gene may prove to be a useful selectable marker during *T. acidophilum* genetic studies. We have identified a plasmid in *T. acidophilum* (22). This plasmid, with the gene for GyrB(R136H)62 inserted, should allow us to develop a transformation method for *T. acidophilum.*

**Nucleotide sequence accession number.** The nucleotide sequence of the gyrase gene from HO-62N1C has been deposited.
into the DNA Data Bank of Japan and is identified by the accession number AB206999.

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