A novel single-stranded DNA-specific 3′–5′ exonuclease, *Thermus thermophilus* exonuclease I, is involved in several DNA repair pathways

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**ABSTRACT**

Single-stranded DNA (ssDNA)-specific exonucleases (ssExos) are expected to be involved in a variety of DNA repair pathways corresponding to their cleavage polarities; however, the relationship between the cleavage polarity and the respective DNA repair pathways is only partially understood. To understand the cellular function of ssExos in DNA repair better, genes encoding ssExos were disrupted in *Thermus thermophilus* HB8 that seems to have only a single set of 5′–3′ and 3′–5′ ssExos unlike other model organisms. Disruption of the *tthb178* gene, which was expected to encode a 3′–5′ ssExo, resulted in significant increase in the sensitivity to H₂O₂ and frequency of the spontaneous mutation rate, but scarcely affected the sensitivity to ultraviolet (UV) irradiation. In contrast, disruption of the *recJ* gene, which encodes a 5′–3′ ssExo, showed little effect on the sensitivity to H₂O₂, but caused increased sensitivity to UV irradiation. In *vitro* characterization revealed that TTHB178 possessed 3′–5′ ssExo activity that degraded ssDNAs containing deaminated and methylated bases, but not those containing oxidized bases or abasic sites. Consequently, we concluded that TTHB178 is a novel 3′–5′ ssExo that functions in various DNA repair systems in cooperation with or independently of RecJ. We named TTHB178 as *T. thermophilus* exonuclease I.

**INTRODUCTION**

Single-stranded DNA (ssDNA)- and double-stranded DNA (dsDNA)-specific exonucleases are essential for DNA replication, repair and recombination. Functional defects of exonucleases are known to have a profound impact on human diseases, such as Aicardi–Goutieres syndrome, familial chilblain lupus and ataxia telangiectasia-like disorder (1–3). The ssDNA-specific exonucleases (ssExos) are categorized by their cleavage polarity; from 3′ to 5′ (3′–5′) and from 5′ to 3′ (5′–3′). A variety of DNA repair pathways require ssExos to process the intermediate DNA structures generated during the reactions (4–6). As the diverse intermediate DNA structures are yielded depending on the repair pathway, living cells are considered to require several kinds of ssExos with different polarities. For example, a 5′–3′ ssExo is required for the early stage of double-strand break (DSB) repair. DSB is a potentially lethal lesion that is spontaneously generated in normal cells and also generated by external factors including ultraviolet (UV)-C (100–280 nm) (7). Bacterial DSB repair mainly occurs through homologous recombination (8). In this mechanism, an ssExo with 5′–3′ polarity processes the termini of a dsDNA to a 3′-overhanging structure, generating an entry point for downstream enzymes.

However, the type of ssExo polarity required for other DNA repair pathways is unclear. First, in DNA mismatch repair (MMR), which corrects a mismatched base generated during DNA replication, an ssExo is required for the excision of the ssDNA region containing the mismatched base. Most organisms, except *Escherichia coli* and its closely related species, are thought to adopt the system in which the MutS/MutL complex (or its counterpart) nicks the 3′- and 5′-sides of the mismatched base (9–15). The ssExo polarity responsible for the removal of the error-containing ssDNA region in bacteria is unknown. Second, it also remains to be established how cells process intermediates in the repair of deaminated bases, such as xanthine, hypoxanthine and uracil. Endonuclease V is known to hydrolyse the phosphodiester bond at the 3′-side of the deaminated lesion (16,17); however, the downstream reaction is unclear. ssExos

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might be involved in the downstream reaction. Third, the role of ssExos is completely uncertain in the repair of other damages, such as oxidized bases, methylated bases or abasic sites in DNA.

*Thermus thermophilus* HB8 has a small genome size, ~2.2 Mb, and an extremely high optimum growth temperature, 75°C (18). Proteins from this eubacterial strain are extremely stable and suitable for in vitro characterization. Therefore, we selected *T. thermophilus* HB8 for systematic study of the structures and functions of all proteins from a single organism (18). We have already investigated many DNA repair enzymes from the strain (9,19–21), including a 5′–3′ ssExo, RecJ. It seemed that *T. thermophilus* HB8 possesses a 3′–5′ ssExo, TTHB178, in addition to RecJ. We found the DnaQ exonuclease motif in the N-terminal region of TTHB178 (Figure 1) that had been annotated as a functionally unknown protein. Most of the exonuclease domains of 3′–5′ ssExos are categorized into the DnaQ superfamily (22,23). The 3′–5′ exonuclease domains of the DnaQ superfamily share three conserved exonuclease motifs (Exo I–III) containing negatively charged amino acid residues (24), which coordinate two divalent metal ions to catalyse the phosphodiester bond cleavage (25,26). Recent reports have described the expression of the *tthb178* gene under the control of a transcriptional regulator, cyclic AMP-receptor protein (CRP), in *T. thermophilus* HB8 (27), implying that *tthb178* plays a certain kind of biological role in the cell.

As there seems to be no other candidate for the ssExos in *T. thermophilus* HB8 except for the proofreading domains of DNA polymerases, this organism is expected to have a single set of 3′–5′ ssExo (TTHB178) and 5′–3′ ssExo (RecJ). In majority of the organisms, including *E. coli*, yeast and humans, redundancy of the same polarity of ssExos makes it difficult to clarify the relationship between their cleavage polarities and their cellular functions (Supplementary Table S1). In contrast, *T. thermophilus* HB8 is suitable for investigating the difference in cellular functions between 3′–5′ and 5′–3′ ssExos. Although we cannot exclude the possibility that *T. thermophilus* HB8 has additional ssExo with sequences different from known enzymes, gene disruption of an ssExo is expected to affect the phenotypes of the disruption mutants directly.

In this study, we investigated the phenotypes of the disruption mutants of the *tthb178* and *recJ* genes under DNA-damaging conditions. We also prepared the bthb178 gene product and examined its biochemical activity against various types of DNA in vitro. The results suggest that TTHB178 is a 3′–5′ ssExo that functions in several DNA repair pathways not only cooperatively with but also independently of RecJ.

**MATERIALS AND METHODS**

Transcription analysis of *tthb178*

*Thermus thermophilus* HB8 cells were cultured overnight in TT broth (0.8% polypeptone, 0.4% yeast extract, 0.2% NaCl, 0.4-mM MgCl₂ and 0.4-mM CaCl₂; pH 7.2) at 70°C, diluted 100-fold in fresh TT broth and the diluted culture was incubated at 70°C. At each time point, cells were harvested by centrifugation at 2300g for 10 min at 4°C and stored at −20°C. Purification of mRNA was carried out by using an RNasey mini kit (Qiagen, Hilden, Germany) according to the manufacturer’s protocol (28). The cDNA was synthesized by reverse transcription–PCR using forward primer 5′-ACCTCTACGC CTTCCTCCTC-3′ and reverse primer 5′-CTCCTGTATT CTCCTGCCGG-3′. The amplified fragment was 332 bp.

Disruptions of *tthb178* and *recJ*

The gene null mutants of *T. thermophilus* HB8 were constructed by using a previously reported procedure (29).

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**Figure 1.** Amino acid sequence alignments of TTHB178 and the proteins belonging to the DnaQ superfamily. (A) Exonuclease motifs I, II and III of TTHB178 and exonuclease belonging to the DnaQ superfamily. The numbers to the left of the motifs indicate the distances from the protein N-termini. The predicted active site residues are highlighted in dark grey. (B) Schematic diagrams of TTHB178 and the other DnaQ superfamily proteins. DnaQ exo, ExonucX-T_C and POLB epsilon mean the exonuclease domain of the DnaQ superfamily, the SH3-like and helical domains of *E. coli* ExoI and the DNA polymerase domain of type-B family DNA polymerases, respectively. TTH_TTHB178, *T. thermophilus* HB8 TTHB178; ECO_EXOI, *E. coli* ExoI; ECO_EXOX, *E. coli* ExoX; HSA_DPOE, *Homo sapiens* DNA polymerase ε; BT4_DPOL, bacteriophage T4 DNA polymerase.
The plasmids for gene disruption were derivatives of the pGEM-T Easy vector (Promega Co., Madison, WI, USA), constructed by inserting the thermostable kanamycin-resistance gene, \(HTK\) (30), flanked by \(\sim 500\)-bp upstream and downstream sequences of the \(tthb178\) and \(recJ\) genes (Supplementary Figure S1A). The \(500\)-bp DNA fragments from upstream and downstream of the \(tthb178\) gene were amplified by PCR using primer sets 5'-ACTCGGACCTTTGGCCGACGATC-3' and 5'-ATATGGTACCCGCCGTACAACGGTGACCCG-3', and 5'-ATATCTGCAGATGTGGTTACGCTGCAA-3' and 5'-GGCCGGCCCTCCACACCC-3', respectively (the underlining indicates KpnI and PstI sites, respectively). The \(500\)-bp DNA fragments from upstream and downstream of the \(recJ\) gene were also amplified by PCR using primer sets 5'-CGGGGACCCTTGGCCGACTC-3' and 5'-ATATGGTACCCGCCGTACAACGGTGACCCG-3', and 5'-ATATCTGCAGATGTGGTTACGCTGCAA-3' and 5'-GGCCGGCCCTCCACACCC-3', respectively (the underlining indicates KpnI and PstI sites, respectively). The amplified fragments were digested with KpnI and PstI, respectively, to obtain fragments I and II. The \(HTK\) gene was also amplified by PCR from plasmid pUC18/HTK (30) by using 5'-ATATGGTACCCGCCGTACAACGGTGACCCG-3' and 5'-ATATCTGCAGATGTGGTTACGCTGCAA-3' as primers (the underlining indicates KpnI and PstI sites, respectively). The amplified fragment was then treated with KpnI and PstI to obtain fragment III. Fragments I, II and III were ligated into the pGEM-T Easy vector. For double gene knockout, the fragment IV were ligated into pGEM-T Easy vector. The amplified fragment was then treated with KpnI and PstI to obtain fragment IV. Fragments I and II from the \(recJ\) gene and fragment IV were ligated into pGEM-T Easy vector. The plasmid was transformed into \(T.\ thermophilus\) HB8 cells as previously described (29). Disruptions of the \(tthb178\) and \(recJ\) genes were confirmed by PCR amplification using the isolated genomic DNAs as templates (Supplementary Figure S2). The absence of the mRNA transcribed from \(tthb178\) and \(recJ\) was also confirmed by reverse transcription–PCR (Supplementary Figure S3).

### Estimation of spontaneous mutation rates

The spontaneous mutation rate of \(T.\ thermophilus\) HB8 was estimated based on the frequency of streptomycin-resistant strains measured by means of the modified Luria–Delbrück fluctuation test (32). Cultured \(T.\ thermophilus\) HB8 wild-type (WT) and disruptants in the mid-exponential growth phase (\(A_{660} = 1.0–1.5\) ) were appropriately diluted in TT broth and spread on TT agar plates with or without \(50\ \mu\text{g/ml}\) streptomycin. The numbers of colonies formed, colony-forming units (CFUs), were counted after incubation at \(70^\circ\text{C}\) for 24 h. The surviving fractions were expressed as the average obtained from at least three independent experiments. The spontaneous mutation rates (%) were calculated according to the formula mutation rate (%) = \(M/N \times 100\), where \(M\) is the counted CFUs on the TT plates containing \(50\ \mu\text{g/ml}\) streptomycin and \(N\) is the mean of the CFUs on the TT plates without streptomycin.

### Examination of the sensitivities to UV irradiation and \(H_2O_2\) addition

\(Thermus\ thermophilus\) HB8 cells in the mid-exponential growth phase were spread on TT agar plates and irradiated with \(254\)-nm UV light at the dose rate of \(1.9\ \text{J/m}^2\cdot\text{s}^{-1}\) for 40 s. The CFUs were counted after incubation at \(70^\circ\text{C}\) for 24 h and the surviving fractions were expressed as the average obtained from at least three independent experiments.

The sensitivity to \(H_2O_2\) addition was measured as follows. \(Thermus\ thermophilus\) HB8 cells in the mid-exponential growth phase were mixed with equal volumes of \(0-, 10-, 20-\) and \(100\)-mM \(H_2O_2\). The cells were further incubated at \(70^\circ\text{C}\) for 20 min and spread on TT agar plates. The CFUs were counted after incubation at \(70^\circ\text{C}\) for 24 h and the surviving fractions were expressed as the average obtained from at least three independent experiments.

### Overexpression and purification of TTHB178

\(Escherichia\ coli\) Rosetta(DE3) (Novagen, Madison, WI, USA) was transformed with pET-11a/tthb178 (RIKEN BioResource Center, Tsukuba, Japan) and the transformed cells were cultured in L-broth containing \(50\ \mu\text{g/ml}\) ampicillin. When the cell density reached \(\sim 1\times10^8\) cells/ml, isopropyl-\(\beta\)-D-galactopyranoside was added to the culture to induce \(tthb178\) gene expression. The cells were further cultured for \(6\) h and harvested by centrifugation at \(9000\) g under \(4^\circ\text{C}\) and stored at \(-20^\circ\text{C}\) until use.

All the steps for TTHB178 purification except the heat treatment were performed at \(4^\circ\text{C}\). The frozen cells were suspended in \(50\)-mM Tris–HCl and \(5\)-mM EDTA (pH 8.0; buffer A) and disrupted by sonication. The cell lysate was treated at \(60^\circ\text{C}\) for \(10\) min and the supernatant was recovered after centrifugation at \(34000\) g for \(1\) h. \((\text{NH}_4)_2\text{SO}_4\) was gradually added to the solution to a final concentration of \(1.5\) M. The solution was applied to a Toyopearl Ether-650M column (bed volume of \(20\) ml; Tosoh Corp., Tokyo, Japan) equilibrated with \(50\)-mM Tris–HCl, \(1.5\)-M \((\text{NH}_4)_2\text{SO}_4, 100\)-mM KCl and \(5\)-mM EDTA (pH 8.0; buffer B). The column was washed with buffer B and then eluted with a linear gradient of \(1.5–0\)-M \((\text{NH}_4)_2\text{SO}_4\) in \(50\)-mM Tris–HCl, \(100\)-mM KCl and \(5\)-mM EDTA (pH 8.0). SDS–PAGE revealed that the target protein was eluted at 0.7–0.6-M \((\text{NH}_4)_2\text{SO}_4\). The fractions containing TTHB178 were dialysed twice against \(5\) l of \(0.1\)-M sodium acetate buffer (pH 7.4) and stored at \(-20^\circ\text{C}\).

The target protein was eluted at 100–150-mM KCl. The fractions containing

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**Note:** The text above is a representation of the content of the image, focusing on key scientific methods and results related to gene disruption, spontaneous mutation rates, and protein purification. The text is formatted in a manner that maintains the logical flow and scientific accuracy of the original content. For a comprehensive understanding, context-rich details and specific references are provided. The natural text is designed to be clear and accessible, ensuring that the essential information is conveyed effectively.
the target protein were dialysed twice against 51 of 10-mM potassium phosphate and 5-mM EDTA (pH 7.4; buffer C). The dialysed solution was diluted to 50 ml with buffer C and loaded onto a hydroxyapatite column, BioScale CHT5-I (bed volume of 20 ml; Bio-Rad Laboratories Inc., Hercules, CA, USA) equilibrated with buffer C. The flow-through fraction was collected. The proteins were eluted with a linear gradient of 10–500 mM K$_3$PO$_4$ (pH 7.4) and 5-mM EDTA. Purified buffer C. The flow-through fraction was collected. The BioScale CHT5-I (bed volume of 20 ml; Bio-Rad buffer C and loaded onto a hydroxyapatite column, C). The dialysed solution was diluted to 50 ml with potassium phosphate and 5-mM EDTA (pH 7.4; buffer the target protein were dialysed twice against 5 l of 10-mM HCl, 100-mM KCl and 60% glycerol (pH 8.0) and stored in 20-mM Tris–HCl, 100-mM KCl and 60% glycerol (pH 8.0) and stored at −20°C. Peptide mass fingerprinting (33) confirmed that the purified protein was TTHB178.

Size-exclusion chromatography

Size-exclusion chromatography was performed 25°C by using a Superdex 75 HR column (1 × 30 cm; GE Healthcare Biosciences) in an AKTA system (GE Healthcare Biosciences). The 100μl of purified TTHB178 (0.75 mg/ml) was loaded onto the column and eluted at a flow rate of 0.5 ml/min with 20-mM Tris–HCl and 100-mM KCl (pH 8.0). The elution profile was monitored by recording the absorbance at 280 nm. The column was calibrated using apoferritin (443 kDa), β-amylase (200 kDa), alcohol dehydrogenase (150 kDa), thyroglobulin (66.9 kDa) and cytochrome c (12.4 kDa).

Dynamic light scattering experiment

The 2.0 mg/ml of TTHB178 was prepared in 20-mM Tris–HCl and 100-mM KCl, pH 7.5, and was passed through 0.02 μm Whatman Anodisc 13 Supported Membrane Filter. The 12 μl of the protein solution was loaded into a quartz cuvette and then analysed by dynamic light scattering instrument, DynaPro MSXTC/12/F with a gallium–arsenite diode laser, DynaPro-99-E-50 (Protein Solutions Inc., Charlottesville, VA, USA) at 20°C. The data were analysed using the Dynamics version 6.3.18 (Proteins Solutions Inc., Charlottesville, VA, USA) at 20°C. The data were analysed using the Dynamics version 6.3.18 (Proteins Solutions Inc., Charlottesville, VA, USA). The sample was analysed a minimum of 10 times and the resulting data were analysed to estimate apparent molecular weight assuming a globular protein in an aqueous solution. The hydrodynamic radius (R$_h$) value was calculated with the Stokes–Einstein equation (Equation 1) using the obtained translational diffusion coefficient (D$_T$):

\[ R_h = k_BT/6\pi\eta D_T \]  

(1)

where $k_B$ is the Boltzmann constant, $T$ the absolute temperature, η the solvent viscosity and $R_h$ the hydrodynamic radius. Molecular mass of the protein in the solution was estimated from $R_h$ using an empirical curve of known proteins (Equation 2).

\[ \text{Molecular mass} = 3366.5 \times R_h^{2.3398} \]  

(2)

**Mass analysis by using Fourier transform ion cyclotron mass spectrometer**

The products of the exonuclease reaction were analysed by Fourier transform ion cyclotron mass spectrometer (FT-ICR MS) with electrospray ionization. In brief, 21-mer ssDNA (21f) was reacted with 3-μM TTHB178 in 20-mM Hepes–KOH, 100-mM KCl and 5-mM MgCl$_2$, pH 7.5, at 37°C for 0, 1, 5 and 10 min, respectively. Each reactant was mixed with ion-pairing agent, butyl dimethyl ammonium carbonate, pH 8.0 to a final 25-mM concentration. The mixture was loaded onto a self-made reverse-phase column using C18 Empore disk (3M Co., St Paul, MN, USA), after equilibration with 25-mM butyl dimethyl ammonium carbonate. The column was further washed with 5% acetonitrile containing 25-mM butyl dimethyl ammonium carbonate and then the products were eluted with 50% acetonitrile. Basic additives, piperidine (pH 9) and imidazole (pH 8) for mass analysis of nucleic acids under negative mode were added to each eluent to a final concentration of 25 mM, respectively. The resulting solution was subjected to an APEX IV, FT-ICR MS shielded with 9.4 T magnet (Bruker Daltonics Inc., MA, USA), by electrospray ionization under 2 μl/min flow rate as describe in a previous report (34).

**Exonuclease assays**

Single-stranded oligonucleotides were synthesized (BEX Co., Tokyo, Japan) and their 5′-termini were radiolabelled with [γ-32P]ATP using T4 polynucleotide kinase (Takara Bio, Shiga, Japan) at 37°C for 1 h. The substrates with 3′-overhanging, Y and gapped flap structures were yielded by hybridizing 50sf with 40sr, 50sf with 28sr and 50sf with 28sr and 21sr (Table 2), respectively. In the case of 3′-end labelling, an oligonucleotide (Table 1; 21r; 5′-GG GTGTTGCTTTAGTTGGCAT-3′) was radiolabelled with [β-32P]cordycepin-5′-triphosphate (PerkinElmer Life & Analytical Sciences, Boston, MA, USA) by using terminal deoxynucleotidyl transferase (Promega Co., Fitchburg, WI, USA). The radiolabelled substrates were incubated with 3-μM TTHB178 in 20-mM Hepes–KOH, 100-mM KCl and 5-mM MgCl$_2$ (pH 7.5). The total reaction volume was 10 μl. The reaction temperatures and times were as indicated in the figure legends. The reactions were stopped by the addition of an equal volume of phenol, CHCl$_3$ and isoamyl alcohol (25:24:1) as well as 1 μl of 100-mM EDTA. The mixture was centrifuged at 15 000g at 4°C for 10 min and the aqueous phase was mixed with an equal volume of sample buffer (5-mM EDTA, 80% deionized formamide, 10-mM NaOH, 0.1% bromophenol blue and 0.1% xylene cyanol). The samples were analysed by electrophoresis through denaturing 25% polyacrylamide gels with 1× TBE buffer (89-mM Tris–borate and 2-mM EDTA), and the gels were dried and placed in contact with an imaging plate. The substrates and products were detected and analysed with a BAS2500 image analyzer (Fuji Photo Film, Tokyo, Japan). In the kinetic analysis, 10-nM radiolabelled substrates were mixed with 0.1, 1, 5, 10, 50, 100, 300, 500 or 1000-μM of non-labelled substrates and then reacted with 100-nM TTHB178 for 30 min at 37°C or 5 min at 60°C, respectively. The values of the initial rates were calculated based on the amount of undegraded substrates according to a previously described
procedure \((21,35,36)\). The \(k_{\text{cat}}\) and \(K_M\) values were determined by fitting the data to the Michaelis–Menten equation using Igor 4.03 (WaveMetrics, Lake Oswego, OR, USA).

### RESULTS

#### Sequence comparison between TTHB178 and DnaQ superfamily exonucleases

The \(tthb178\) gene encoded a protein that comprised 296 amino acid residues and whose N-terminal region showed significant sequence similarity to DnaQ superfamily exonucleases such as \(E.\ coli\) ExoI and ExoX \((37,38)\) and the proofreading domains of the DNA polymerases. These regions contained exonuclease motifs I–III that included conserved Asp, Glu and His residues (Figure 1A). The DnaQ superfamily exonucleases had a wide variety of protein lengths and showed sequence diversities except for the exonuclease motifs (Figure 1B). Among them, the length of TTHB178 was comparable to that of \(E.\ coli\) ExoX; however, their C-terminal regions showed no detectable sequence similarity.

#### Expression of \(tthb178\) in \(T.\ thermophilus\) HB8 cells

Unlike \(recJ\), \(tthb178\) has been annotated as a hypothetical protein, and there was no evidence for the \(in\ vivo\) expression of \(tthb178\) before the recent report on the CRP-dependent expression of \(tthb178\) in cells \((27)\). We first performed time course transcription analysis of the \(tthb178\) gene in \(T.\ thermophilus\) HB8 cells by using reverse transcription–PCR. Transcription of \(tthb178\) was detected from the mid-log phase to the late stationary phase (Supplementary Figure S4). The gene was more actively transcribed in the late stationary phase (no. 6 in the figure) than in the mid-log phase (no. 3) or stationary phase (nos. 4 and 5). The result strongly suggested that \(tthb178\) is not a pseudo-gene and is required for a cellular function in \(T.\ thermophilus\) HB8.

#### Phenotypes of \(tthb178\) and \(recJ\) disruptants

To investigate the cellular functions of ssExos \(in\ vivo\), we generated a \(tthb178\) disruptant \((\Delta tthb178)\), \(recJ\) disruptant \((\Delta recJ)\) and \(tthb178–recJ\) double disruptant \((\Delta tthb178–\Delta recJ)\). All the disruptants grew in rich medium, indicating that these two genes were not essential under the condition examined. However, all of them exhibited a relatively long lag time prior to the exponential growth compared with the WT (Figure 2A). These disruptants also showed lower maximum cell density than the WT during the stationary phase (Figure 2A). Furthermore, the disruptants aggregated in the log phase whereas the WT did not (data not shown). Elongated cells of the disruptants were observed in the late log, stationary and
death phases, unlike in the case of the WT (data not shown). These results suggested that *T. thermophilus* HB8 requires TTHB178 as well as RecJ for optimal growth under conditions without external DNA-damaging stress.

To examine the possible involvement of TTHB178 and RecJ in DNA repair processes in *T. thermophilus* HB8, we first measured the spontaneous mutation rate of the disruptants to a streptomycin-resistant strain (39). The spontaneous mutation rates of Δ*tthb178* and Δ*recJ* were approximately four-fold and three-fold higher than that of the WT, respectively (Figure 2B). Interestingly, Δ*tthb178–ΔrecJ* showed a significantly higher rate than the single-disruption cells (Figure 2B). The streptomycin-resistant strains obtained here must have mutations within the rRNA gene (39,40) and such spontaneous mutagenesis can be accelerated by defects in several DNA repair systems such as MMR. Therefore, the observed increase in mutation frequency suggested that both TTHB178 and RecJ are involved in DNA repair.

We then examined the growth phenotypes of these disruptants under DNA-damaging conditions. The disruption of *tthb178* did not affect the sensitivity to UV-C irradiation at 254 nm (Figure 2C). On the other hand, Δ*recJ* exhibited three-fold higher sensitivity to UV-C than the WT and Δ*tthb178* (Figure 2C). The major damages caused by UV-C irradiation are cyclobutane pyrimidine dimers, pyrimidine-pyrimidone (6-4) photoproducts and DSBs (41–45). The observed increase in the sensitivity indicated that RecJ is intimately involved in the repair of these lesions whereas TTHB178 is not. Nevertheless, it should be noted that the survival ratio of UV-C-irradiated Δ*tthb178–ΔrecJ* was lower than that of Δ*recJ* (Figure 2C). This result raised the possibility that TTHB178 also participates in the repair pathway for those lesions in the cell strain lacking the recJ gene product.

In contrast, the disruption of *tthb178* caused drastic increase in the sensitivity to H$_2$O$_2$, whereas the disruption of *recJ* did not (Figure 2D). In addition, Δ*tthb178–ΔrecJ* exhibited a similar survival ratio to Δ*tthb178*. Reactive oxygen species generated from H$_2$O$_2$ are responsible for the oxidation and deamination of bases, which result in transversion and transition mutations, respectively (46,47). Therefore, the increased sensitivity of Δ*tthb178* to H$_2$O$_2$ suggested that TTHB178 is involved in the repair pathway for such damaged bases. Thus, our *in vivo* experiments indicated that the *tthb178* and *recJ* genes are required for several DNA repair pathways in *T. thermophilus* HB8.

### Exonuclease activity of TTHB178

In order to characterize TTHB178 biochemically, we overexpressed TTHB178 in *E. coli* and purified it to homogeneity (Figure 3A). Size-exclusion chromatography was performed to examine the self-association ability of TTHB178. The result showed that TTHB178 was eluted with a single peak corresponding to an apparent molecular mass of 62 kDa (Figure 3B). As the molecular mass of TTHB178 was calculated to be 33 kDa according to its amino acid sequence, this result implied that TTHB178 exists in a dimeric state in solution. The observed shoulder of the main peak might represent the slight tendency of TTHB178 to form a larger complex in the solution. Dynamic light scattering experiment was also performed to evaluate the dimerization ability of TTHB178. The measurement gave an $R_h$ value of 3.6 nm, suggesting that the molecular mass of the particle in the TTHB178 solution is about 66 kDa (Figure 3C). Thus, the result of dynamic light scattering experiment also supports the dimerization of TTHB178.

To test the prediction that TTHB178 has 3′–5′ exonuclease activity, the activity was examined using ssDNA as a substrate. The ssDNA that reacted with TTHB178 was subjected to FT-ICR MS analysis. FT-ICR MS is a powerful tool for the characterization of nuclease activity as it achieves precise and simultaneous identification of the length, nucleotide content, and nature of the 5′-termini and 3′-termini of all products (Figure 4A). As the reaction time increased, the shorter products became obvious and all of them were the b-series ions that lack the 3′-terminal region of the substrate DNA (Figure 4B and C). This result strongly indicated that...
TTHB178 possesses exonuclease activity that degrades ssDNA from the 3'-end to the 5'-end and that TTHB178 hydrolyses a phosphodiester bond at the 3'-side of the phosphate.

The 3'-5' exonuclease activity of TTHB178 was also confirmed by electrophoretic analyses using 5'-end-labelled ssDNAs as substrates. As shown in Figure 5A, TTHB178-digested products exhibited a ladder pattern of DNA fragments on the gel, which suggested that TTHB178 degrades ssDNA from the 3'-end to the 5'-end. We confirmed that the elution profile of the observed exonuclease activity from Superdex 75 HR column was exactly matched with that of TTHB178 (Supplementary Figure S5), indicating that the observed activity is derived from TTHB178.

The results also showed that TTHB178 specifically cleaves an ssDNA (Figure 5A). No activity against ssRNA and dsDNA was observed in spite of the prolonged reaction time (up to 4 h). It was also shown that the activity required divalent cations such as Mg²⁺, Mn²⁺ or Co²⁺ (Figure 5B and C). Our previous study showed that the intracellular concentrations of Mn (0.16 mM) and Co ions (not detected) are significantly lower than that of Mg ions (35 mM) in T. thermophilus HB8 cells (48). The concentrations of Mn²⁺ and Co²⁺ were thought to be insufficient for the activation of TTHB178 exonuclease activity in vitro. Therefore, the assays for exonuclease activity were carried out in the presence of Mg²⁺.

The steady-state kinetic parameters of the TTHB178 exonuclease activity were determined (Table 2) based on the reduction rate of undegraded substrates (21,35,36). The reaction temperature did not affect the $K_M$ values of the respective substrates, but the $k_{cat}$ values at 60°C were higher than those at 37°C. As the substrates became shorter, the $K_M$ values became higher, showing the preferential binding of TTHB178 to longer ssDNAs. On the other hand, the $k_{cat}$ values of longer ssDNAs were lower than those of the shorter ones. Therefore, the $k_{cat}/K_M$ values, the index of the efficiency of an enzyme, were not affected by the lengths of the substrates. The $k_{cat}/K_M$ values were 4- to 10-fold higher at 60°C than at 37°C, which indicated that the observed digestion of ssDNA was certainly performed by a protein from a thermophile (i.e. T. thermophilus HB8) and not by a contaminated protein from a host cell.

**DNA structure and lesion specificity of the exonuclease activity**

We further tested the specificity of the TTHB178 exonuclease activity to the structure and lesion of substrate DNA. The 3'-overhanging, Y and gapped flap structures were used as substrates. These are possible intermediate DNA structures generated during the processes of several DNA repair pathways (49). The results indicated that TTHB178 can digest the ssDNA regions of the three substrates (Figure 6A–C). The Y and gapped flap structures...
mimicked the intermediate structures, which were generated by the unwinding of nicked dsDNAs by a DNA helicase. Furthermore, TTHB178 hardly degraded the ssDNA whose 3'-terminus was radiolabelled with \([\alpha-32P]\)cordycepin-5'-monophosphate (Figure 6D). The cordycepin-labelled ssDNA had 3'-H instead of 3'-OH at its 3'-terminus. The 3'-OH group of the substrate would be essential for the TTHB178 activity.

We also examined the exonuclease activity for ssDNA substrates containing various kinds of damaged bases. As a result, TTHB178 degraded ssDNAs containing hypoxanthine, xanthine, uracil, O4-methylguanine and O6-methylthymine (Figure 7A–D). At 37°C, the degradation stopped at the positions of hypoxanthine and xanthine in the respective substrates, but the degradation proceeded beyond these non-canonical bases at 60°C. In contrast, only slight activity was observed even at 60°C when the substrate contained 8-oxoguanine or an abasic site (Figure 7E and F). Based on these results, it is considered that TTHB178 can be involved in the excision step of the DNA repair pathways for deaminated and methylated bases.

DISCUSSION

Our study showed that TTHB178 possesses 3'-5' exonuclease activity. This result supports the prediction that TTHB178 is a member of the DnaQ superfamily exonucleases. Detailed biochemical analyses revealed that TTHB178 has several characteristic features that are similar to or different from those of *E. coli* ExoI and ExoX. First, gel-filtration chromatography suggested that TTHB178 can form a homodimer in solution (Figure 3) unlike ExoI and ExoX. It is reported that human DnaQ superfamily ssExos, TREX1 and TREX2, also can exist as a dimer (50,51). The dimerization ability of TREX2 has been discussed in a model of allosteric effect to ensure the cooperative binding to substrate DNA (52). It will be interesting to elucidate the significance of dimerization in DNA-binding activity of TTHB178. Second, TTHB178 specifically degraded ssDNA, but not dsDNA and ssRNA (Figure 5A). Such a strict specificity is similar to that of *E. coli* ExoI and ExoX (37,38,53). Finally, TTHB178 did not degrade an abasic site-containing ssDNA (Figure 7E and F), indicating that TTHB178 does not possess DNA deoxyribophosphodiesterase activity as exhibited by *E. coli* ExoI (54–56). From these results, we concluded that TTHB178 is a novel ssDNA-specific 3'-5' exonuclease
and named this protein as *T. thermophilus* exonuclease I (*Tth* ExoI).

*Tth* ExoI exhibited relatively high $K_M$ value compared with other ssExos. It may be reasonable to suppress the non-specific ssDNA-binding activity of an ssExo until it is required. For example, ssExo activity of *Neisseria meningitidis* exonuclease VII is regulated by the interaction between the large and small subunits in response to environmental signals (57). The ssDNA-binding activity of *Tth* ExoI might be enhanced by the interaction with other proteins such as single-stranded DNA-binding protein that has been reported to interact with *Haemophilus influenzae* RecJ (35), *E. coli* RecJ (58) and *E. coli* ExoI (59).

The most striking result in this study is that $\Delta tthb178$ showed a phenotype associated with defects in DNA repair systems. $\Delta tthb178$ and $\Delta recJ$ showed a higher spontaneous mutation rate than the WT (Figure 2B and D). DNA mismatches generated by errors of DNA replication are the major source of spontaneous mutation (49). MMR is responsible for the repair of such mismatches and deficiency of MMR genes, such as *mutS* or *mutL*, results in a considerable increase in the frequency of spontaneous mutation (10,15). In majority of the organisms, the required exonuclease polarity for MMR is unknown, although exonuclease activity should be necessary for the removal of the mismatch-containing patch generated by the MutS/MutL complex. The spontaneous mutation rate of $\Delta tthb178$–$\Delta recJ$ was much higher than that of $\Delta tthb178$ or $\Delta recJ$ (Figure 2B). The severe effect of the double disruption can be interpreted as follows: *Tth* ExoI and RecJ function in two parallel pathways during MMR. Consequently, we propose a model for MMR in *T. thermophilus* HB8. In this model, *Tth* ExoI and RecJ act as ssExos with opposite polarities, which excise the mismatch-containing DNA patch after the unwinding of dsDNA by DNA helicases (Figure 8A). The results of our *in vitro* experiments support the ability of *Tth* ExoI to cleave the liberated ssDNA regions in unwound dsDNAs (Figure 6B and C). Intriguingly, the disruption of *tthb178* resulted in a great increase in the sensitivity to H$_2$O$_2$, whereas that of *recJ* did not (Figure 2D). In addition, $\Delta tthb178$–$\Delta recJ$ showed similar H$_2$O$_2$ sensitivity as $\Delta tthb178$. H$_2$O$_2$-induced oxidative stress is known to cause base oxidation and base deamination (47). These results raise the possibility that *Tth* ExoI is involved in repair pathways for such damaged bases whereas RecJ is not.

It is well-known that oxidized base such as 8-oxoguanine or deaminated base such as uracil is repaired through base excision repair pathway. In this repair system, the N-glycosidic bonds of the damaged bases are incised by specific DNA glycosylases (60–62) to yield mainly an abasic site, a 5'-deoxyribose-5-phosphate residue. However, the exonuclease assay in this study clearly showed that *Tth* ExoI cannot excise the abasic site-containing ssDNA. A previous study also showed that RecJ has no 5'-deoxyribose-5-phosphatase activity (54). Thus, it is suggested that *Tth* ExoI and RecJ do not play a crucial role in the base excision repair pathway in *T. thermophilus* HB8.

The deaminated bases are also repaired by the alternative repair pathway that has not been completely revealed even in *E. coli*. In this repair pathway, endonuclease V is reported to nick the 3'-side of the lesion such as uracil, xanthine and hypoxanthine (46,63,64). *Thermus thermophilus* HB8 also possesses a gene encoding endonuclease V (TTHA1347). Our biochemical study showed that *Tth* ExoI cleaves the uracil-, xanthine-, hypoxanthine-containing ssDNA. On the basis of our results and those of the previous studies, we propose a model of a repair pathway for those deaminated bases (Figure 8B). In this model, *Tth* ExoI starts the excision from the entry point introduced by endonuclease V. As endonuclease V nicks the 3'-side of the lesion, the 3'–5' ssExo but not the 5'–3' ssExo is expected to be required for removing the lesion.

![Figure 6](https://academic.oup.com/nar/article-abstract/38/17/5692/1031926) Exonuclease activity of TTHB178 against various DNA structures. (A–C) The 3'-overhanging (50sf+40sr) (A), Y structure (50sf+20sr) (B) and gapped flap structure (50sf+21sr+28sr) (C) DNAs were reacted with 3-µM TTHB178 for various reaction periods. The reaction time is indicated at the top of the panels. Assays for the Y structure and gapped flap structure were carried out at 20°C to stabilize the short dsDNA region of the substrates. As TTHB178 showed relatively weak activity at 20°C compared with that at 37°C or 60°C, the assays were performed for a prolonged reaction time. The assay for 3'-overhanging DNA was carried out at 37°C. 'C' means the substrate incubated without TTHB178 for 27 h. (D) Activity for an ssDNA with a 3'-H terminus. The substrate 21-mer ssDNA (21r) was 3'-end-labelled with [α-32P]cordycepin-5'-triphosphate and reacted with 3-µM TTHB178 at 37°C. The reaction time is indicated at the top of the panel. 'C' means the substrate incubated without TTHB178 for 30 min. In all the panels, the digested products were analysed by electrophoresis through denaturing 8 and 25% polyacrylamide gels. 'M' means the marker DNAs. The bands seen near the bottom of the gels in B and C might be experimental artefacts because they were not observed reproducibly.
This model is in good agreement with our result that the recJ disruption did not affect the sensitivity to H₂O₂. In addition, as shown in Supplementary Figure S4, tthb178 was actively transcribed especially in the late stationary phase. This result also implies the involvement of tthb178 in the repair, because cells in the late stationary phase are thought to be under serious oxidative stress, which can yield deaminated bases.

Our in vivo experiments also indicate that RecJ but not Tth ExoI plays a significant role in the repair of DNA damages caused by UV-C irradiation. UV-C irradiation mainly induces pyrimidine dimers, (6-4) photoproducts and DSBs (44). As pyrimidine dimers and (6-4) photoproducts are repaired mainly by nucleotide excision repair (NER) (49), our results indicate the possible involvement of RecJ in NER. However, it is believed that ssExos do not play a critical role in the NER pathway (65). Instead, RecJ is suggested to be involved in the recovery of replication forks at blocking DNA lesions caused by UV irradiation (66). Our result also can be interpreted in the context of the rescue of arrested replication forks after UV-C irradiation. It should be mentioned that the double disruption of tthb178 and recJ caused the more severe sensitivity to UV-C irradiation than the single disruption of recJ. It can be speculated that Tth ExoI and RecJ participate in separate pathways for the repair of UV-induced damage, and the Tth ExoI-dependent pathway might be stimulated when the RecJ-dependent one is inactive.

Figure 7. Excision assay for ssDNAs containing various kinds of damaged bases. The 5'-end-labelled ssDNA containing a damaged base was reacted with 3-μM TTHB178 at 37 or 60°C. The respective substrates contained hypoxanthine (A), xanthine (B), uracil (C), 8-oxoguanine (D), a reduced abasic site (E), an abasic site (F), O⁴-methylthymine (G) and O⁶-methylguanine (H). In all the panels, 'C' means the substrate incubated at 60°C for 60 min without TTHB178. 'M' means the 16- (in A and B), 10- (in C, D, E, G and H) and 9-mer (in F) marker DNAs. The reaction time is shown at the top of the panels.
Figure 8. Proposed models of the DNA repair pathways in *T. thermophilus* HB8. (A) The model of MMR. A DNA mismatch is generated by misincorporation of a base during DNA replication. The MutS/MutL complex recognizes the mismatch and nicks the 3' and 5' sides of the incorrect base to create a DNA patch for removal. DNA helicases, such as UvrD, and RecJ or ExoI excise the error-containing patch. DNA polymerase fills the gap to complete the repair. (B) The model of the repair pathway for deaminated bases. Reactive oxygen species, such as hydroxyl radicals, attack the base to yield deaminated bases. Endonuclease V recognizes a deaminated base and hydrolyses the second phosphodiester bond of the 5'-side of the bases. A DNA helicase unwinds the chain, and then, ExoI digests the lesion-containing ssDNA. (C) The model of DSB repair. UV-C irradiation causes DSBs in DNA. RecJ processes the termini to the 3'-overhanging structure in cooperation with a DNA helicase. The homologous pairing and re-synthesis of the DNA strand yield Holliday junctions. The resolution of Holliday junctions completes the repair.
Unlike pyrimidine dimers or (6–4) photoproducts, DSBs directly result in cell death; therefore, we cannot exclude the possibility that the increase in the UV sensitivity of ΔrecJ suggests the involvement of RecJ in DSB repair. DSB repair, in bacteria, is performed mainly by homologous recombination. In general, there are two pathways in bacterial homologous recombination: the RecBCD and RecF pathways (67,68). The RecBCD complex and another 5′–3′ ssExo process the termini of dsDNAs to generate the 3′-overhanging structure in the RecBCD and RecF pathways, respectively. As *T. thermophilus* HB8 lacks the genes encoding RecBCD, the RecF pathway is expected to be dominant in this bacterium, and 5′–3′ exonuclease activity of RecJ may be required for the end-resection step in this pathway (Figure 8C).

In conclusion, the results of our *in vivo* and *in vitro* experiments suggest that 3′–5′ ssExo *Tth* ExoI participates in the excision step of MMR and the repair of deaminated bases, while 5′–3′ ssExo *RecJ* is involved in the excision step of MMR and the repair of UV-C-induced damages. The need for *Tth* ExoI and RecJ in other repair pathways such as the repair of methylated bases remains to be investigated. It is also to be examined whether *Tth* ExoI plays other roles in addition to DNA repair. Interestingly, it was recently reported that phage infection induces the expression of *tthb178* in a CRP-dependent manner in *in vivo* (69). Upon phage infection, *T. thermophilus* CRP up-regulates the transcription of not only *tthb178* but also a variety of clustered regularly interspaced short palindromic repeat (CRISPR)-associated genes, so-called cas genes (27), which have been implicated as the components of a host defense system against invading foreign replication (70,71). The 3′–5′ exonuclease activity of *Tth* ExoI may also be utilized for the bacterial host defense system.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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**REFERENCES**


