A thermostable endonuclease III homolog from the archaeon Pyrobaculum aerophilum

Hanjing Yang1, Isabella T. Phan1, Sorel Fitz-Gibbon12, Mahmud K. K. Shivji3, Richard D. Wood2, Wendy M. Clendenin1, Elizabeth C. Hyman1 and Jeffrey H. Miller1,*

1Department of Microbiology and Molecular Genetics and the Molecular Biology Institute, 2IGPP Center for Astrobiology, University of California, Los Angeles, CA 90095, USA and 3Imperial Cancer Research Fund, Clare Hall Laboratories, South Mimms, Hertfordshire EN6 3LD, UK

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ABSTRACT

Pyrimidine adducts in cellular DNA arise from modification of the pyrimidine 5,6-double bond by oxidation, reduction or hydration. The biological outcome includes increased mutation rate and potential lethality. A major DNA N-glycosylase responsible for the excision of modified pyrimidine bases is the base excision repair (BER) glycosylase endonuclease III, for which functional homologs have been identified and characterized in Escherichia coli, yeast and humans. So far, little is known about how hyperthermophilic Archaea cope with such pyrimidine damage. Here we report characterization of an endonuclease III homolog, PaNth, from the hyperthermophilic archaeon Pyrobaculum aerophilum, whose optimal growth temperature is 100°C. The predicted product of 223 amino acids shares significant sequence homology with several [4Fe-4S]-containing DNA N-glycosylases including E.coli endonuclease III (EcNth). The histidine-tagged recombinant protein was expressed in E.coli and purified. Under optimal conditions of 80–160 mM NaCl and 70°C, PaNth displays DNA glycosylase/β-lyase activity with the modified pyrimidine base 5,6-dihydrothymine (DHT). This activity is enhanced when DHT is paired with G. Our data, showing the structural and functional similarity between PaNth and EcNth, suggests that BER of modified pyrimidines may be a conserved repair mechanism in Archaea. Conserved amino acid residues are identified for five subfamilies of endonuclease III/UV endonuclease homologs clustered by phylogenetic analysis.

INTRODUCTION

Pyrimidine adducts in cellular DNA arise from modification of the pyrimidine 5,6-double bond by oxidation, reduction or hydration. So far, more than 20 stable pyrimidine adducts have been discovered (1,2). The biological consequences of some of these modified pyrimidine bases, including 5-hydroxycytosine, 5-hydroxypyrimidine, uracil glycol, thymine glycol and 5,6-dihydrothymine (DHT), have been studied in detail regarding miscoding potential and effect on DNA replication (1,3,4). With only a few exceptions, they all cause either miscoding or polymerase blocking. Both lead to increase in mutation rate or possible cell death.

Repair of pyrimidine adducts is mainly via the base excision repair (BER) process which is initiated by DNA N-glycosylases (1,5,6). In Escherichia coli, endonuclease III (EcNth) is responsible for the excision of most modified pyrimidine bases (7–14). It is bifunctional, having both DNA N-glycosylase activity that removes modified pyrimidine bases, and an intrinsic apurinic/apyrimidinic (AP) lyase activity that cleaves the phosphodiester backbone of DNA 3’ to AP sites. This generates 5’-deoxyribophosphate and a 3’-terminal unsaturated sugar derivative. The abnormal 3’-abasic residue is then removed by an AP endonuclease, another BER enzyme, to generate a 3’-OH nucleotide residue. Subsequently the gap is repaired by other BER components restoring the original genetic information. Functional endonuclease III homologs have been identified in organisms as diverse as yeast and humans (15–18), which suggests that BER of modified pyrimidines has been crucial for maintaining DNA stability during evolution.

Structurally, EcNth belongs to the HhH DNA N-glycosylase superfamily, which contains a helix–hairpin–helix (HhH) motif for DNA binding and a proline/glycine-rich loop (GPD motif). This GPD motif contains the catalytic aspartic acid residue corresponding to D138 in EcNth (19). Other members in this HhH superfamily are T/G and U/G mismatch-specific glycosylases (20–22), a few subgroups of methylpurine DNA glycosylase (23–25), UV endonuclease (26,27), adenine-DNA glycosylase (28–30) and 7,8-dihydro-8-oxoguanine DNA glycosylase (31–35). They all catalyze N-glycosyl bond cleavage of different unwanted bases from DNA. Many of the proteins that belong to the HhH DNA N-glycosylase superfamily also contain an additional conserved structural element—an iron–sulfur cluster loop (FCL motif). This motif contains four cysteine residues for binding of a [4Fe-4S] cluster (36). The crystal structures of members in the HhH superfamily reveal the nucleotide flipping mechanism for the base removal. The
unwanted base is flipped out of the DNA helix into the glycosylase active site pocket and cleaved (37–42).

In thermophiles, whose optimal growth temperature is above 65°C, the rate of spontaneous deamination of cytosine and 5-methylcytosine is thermally enhanced, giving increased yields of U/G and T/G mismatches (20,21,43–46). Little is known about other types of pyrimidine lesions in thermophiles or their corresponding repair pathways. However, putative DNA N-glycosylase sequence homologs have been identified in thermophiles and hyperthermophiles whose genomes have been completely sequenced (47). This allows us to study, in vitro, the enzymatic activities of these putative homologs as a first step toward understanding the DNA repair pathways for pyrimidine lesions in these organisms. So far, the only published study for a thermophilic endonuclease III homolog is an NMR structure description of a homolog from the hyperthermophilic archaeon *Archaeoglobus fulgidus* (48). Here, we report cloning and characterization of a putative DNA N-glycosylase homolog from the hyperthermophilic archaeon, *Pyrobaculum aerophilum*, whose optimal growth temperature is 100°C (49). Using defined DNA substrates we demonstrate that this protein is an endonuclease III homolog (PaNth), which efficiently cleaves a modified pyrimidine, DHT, opposite G. Our data, showing the structural and functional similarity between PaNth and EcNth, suggests that BER of modified pyrimidines may be a conserved repair mechanism in Archaea. Conserved amino acid residues are identified for five subfamilies of endonuclease III/UV endonuclease homologs clustered by phylogenetic analysis.

**MATERIALS AND METHODS**

**Identification of the candidate protein**

The candidate protein coding region, pag5_880, was identified in the recently completed whole genome sequence of *P. aerophilum* (50; S.T.Fitz-Gibbon, H.Ladner, U.-J.Kim, K.O.Stetter, M.I.Simon and J.H.Miller, unpublished data) by analyzing sequence similarity to the public database using FASTA (51).

**Expression and purification of PaNth**

The pag5_880 DNA was amplified by polymerase chain reaction (PCR). The PCR product was cloned into pCR2.1-TOPO vector using TOPO TA Cloning kit (Invitrogen, Carlsbad, CA) and subsequently subcloned into the bacterial expression vector pQE30 (Qiagen, Chatsworth, CA) between the BamHI and HindIII sites. Transformants of *E. coli* CSH100 were grown at 37°C in 1.5 l of Luria–Bertani medium with ampicillin (200 µg/ml). The culture was induced with 0.1 mM isopropyl-β-D-thiogalactopyranoside (IPTG) when it reached optical density 1.5 at 600 nm, and allowed to continue growing overnight. The procedure for purification by Ni²⁺-NTA affinity chromatography was the same as previously described (21). The peak fractions that contained the olive color were combined and dialyzed overnight in Buffer B (50 mM Tris–HCl, pH 7.5, 1 mM EDTA, 1 mM DTT, 30 mM NaCl and 50% glycerol). A clear protein sample was obtained after centrifugation of the dialyzed sample and stored at −80°C. The protein concentration, determined by Bio-Rad Protein Assay (Bio-Rad, Hercules, CA), was 19 mg/ml. The method for determination of molecular mass by electron mass spectrometry was the same as previously described (21).

To prepare heat-treated PaNth, 50 µl of 10-fold-diluted purified recombinant PaNth (1.9 mg/ml) was incubated at 70 or 90°C for 10 min followed by centrifugation at 4°C for 15 min. The clear supernatant was collected and the protein concentration determined by Bio-Rad Protein Assay (Bio-Rad).

**Oligonucleotide substrates**

Sequences of two 96mer oligonucleotides, which were complementary to each other except at position 60, were previously described (21). The oligonucleotide that contains a single modified base, DHT at position 60, was synthesized and PAGE purified (DNAgency, Malvern, PA). The 96mer double-stranded DNA (dsDNA) substrates, containing a base pair X/Y at position 60 [X = A, C, G, T, U or DHT; Y = A, C, G, T or GO (7,8-dihydro-8-oxoguanine or 8-oxoG)], were prepared in an annealing buffer containing 70 mM Tris–HCl, pH 7.6, and 10 mM MgCl₂, as previously described (21). Unless otherwise stated, the X-containing strand was 32P-labeled at the 5′-end.

**Glycosylase activity assay**

The glycosylase activity assay was essentially carried out as previously described (21). Briefly, the standard reaction mixture contained 20 mM Tris–HCl, pH 7.5, 1 mM DTT, 1 mM EDTA, 3% glycerol, 1 µl of 32P-labeled 96mer dsDNA substrate (final concentration of 1 nM) and 2 µl of the 90°C heat-treated recombinant PaNth in a total volume of 20 µl. Unless otherwise stated, the PaNth reaction mixture had 120 mM NaCl (a few cases contained 80 mM NaCl). The 2 µl of 90°C heat-treated recombinant PaNth was diluted to the desired concentrations using Buffer B (50 mM Tris–HCl, pH 7.5, 1 mM EDTA, 1 mM DTT, 30 mM NaCl and 50% glycerol). In the reactions with no PaNth, 2 µl Buffer B was used. The reaction was carried out for 15 min at 70°C (unless otherwise stated) and terminated by cooling to 4°C. Then 8 µl of loading buffer was added. The reaction mixture was heated to 94°C for 1 min and analyzed on a 15% polyacrylamide denaturing gel. Gels were dried and then visualized and quantified by PhosphorImager 445 SI (Sunnyville, CA) using the program Molecular Dynamics ImageQuaNT Version 4.2a. The percentage of cleaved product was determined by dividing the cleaved product intensity by the total intensity. Total intensity was the sum of intact DNA intensity and cleaved product intensity. All intensities were stated after subtracting background (a blank area of identical size).

EcNth (2 mg/ml) was a gift from Dr Richard M. Cunningham (Department of Biological Sciences, State University of New York at Albany, Albany, NY). For reactions containing EcNth, no NaCl was added to the glycosylase reaction buffer, unless otherwise stated. EcNth (2 µl) diluted in Buffer B to the desired concentration was used and the reactions were carried out at 37°C for 15 min. In the reactions with no EcNth, 2 µl Buffer B was used. The subsequent steps were similar to those described in the PaNth glycosylase activity assay (see above).

Reactions to determine the specific activity of the enzyme with DHT/G substrate were carried out using 1 nM DHT/G, 0.25 nM PaNth (at 70°C) or 5 nM EcNth (at 37°C) for 4 min. The reaction products were analyzed as described above.
AP lyase activity assay
A dsDNA substrate containing an AP site opposite G (AP/G) at position 60, was prepared from uracil-DNA glycosylase-treated U/G substrate as described previously (21). Increasing amounts of PaNth were incubated with 1 nM AP/G substrate at 70°C for 15 min followed by analysis on a 15% polyacrylamide denaturing gel as described in the glycosylase activity assay.

Escherichia coli endonuclease IV (endoIV; 0.157 mg/ml) was a gift from Dr Bruce Demple (Department of Cancer Cell Biology, Harvard School of Public Health, Boston, MA). The following experiments were carried out in the reaction buffer containing 20 mM Tris–HCl, pH 7.6, and 50 µg/ml bovine serum albumin (BSA). EcNth (10 nM) was incubated with 1 nM DHT/G substrate for 15 min at 37°C, followed by the addition of 5 nM endoIV and continued incubation for 15 min at 37°C. The reactions were analyzed on a 15% polyacrylamide denaturing gel as described in the glycosylase activity assay.

Oligonucleotide substrate containing 5,6-dihydrouracil (DHU) and activity assay
The 55mer dsDNA substrate, containing DHU paired with G at position 19, was prepared as described by Klungland et al. (52). Standard reaction mixtures (10 µl) contained 100 fmol of DNA substrate, 45 mM HEPES–KOH, pH 7.8, 70 mM KCl, 5 mM MgCl2, 1 mM DTT, 0.4 mM EDTA, 2 mM ATP, 1 mM NAD, 20 µM each of dATP, dGTP, dCTP and dTTP, 3.6 µg BSA, 2% glycerol and either EcNth or PaNth. Prior to adding the glycosylases to the reactions, the proteins were heat-treated as indicated by incubating at 80°C for 20 min followed by centrifugation to remove any precipitate. The reactions were incubated at 37°C for 30 min and terminated by the addition of SDS and EDTA to final concentrations of 0.6% and 25 mM, respectively. The proteins were digested with proteinase K (500 µg/ml) at 37°C for 1 h. The DNA was purified and resolved by electrophoresis on a 20% polyacrylamide denaturing gel. The reaction products were visualized by autoradiography.

UV endonuclease activity assay
The UV-damaged DNA substrates were produced by irradiating plasmid pBluescript KS+ DNA with 450 J/m² UV light. Both the UV-treated plasmid DNA and the non-damaged pHM14 plasmid DNA were treated with EcNth protein to remove pyrimidine hydrates. This was followed by repeated purification of closed circular forms of plasmid DNA as described (53). The UV-damaged DNA was mixed with an equal amount of non-damaged closed circular pHM14 plasmid DNA. The repair reactions (25 µl) containing either PaNth or T4 endoV (as a positive control) and 500 ng DNA mixture (±UV) were incubated in buffer containing 40 mM HEPES–KOH, pH 7.8, 0.5 mM DTT, 4 mM EDTA, 70 mM KCl and 0.36 mg/ml of BSA at either 37°C for 30 min or 70°C for 15 min. The DNA was purified by first treating with proteinase K, phenol–chloroform and precipitating the DNA in absolute ethanol at the presence of 10 µg yeast RNA. After centrifugation, the DNA pellet was washed with 70% ethanol and dried under vacuum. The DNA was resuspended in TE and incubated at 37°C for 15 min. RNase A (80 µg/ml) was added to the DNA to digest the carrier yeast tRNA prior to loading the DNA on a 1% agarose gel containing 0.25 µg/ml ethidium bromide. After electrophoresis, the gel was viewed under UV light.

Preparation of crude extract from *Pyrobaculum aerophilum*
*Pyrobaculum aerophilum* cells (400 ml) were grown in BSY medium (49) at 90°C for 3 days to optical density 0.2 at 600 nm. Cells were harvested by centrifugation. The cell pellet was washed once with Buffer P (10 mM NaPO₄, pH 7.0, 50 mM NaCl), centrifuged and then resuspended in 4 ml Buffer P and sonicated. The crude extract was obtained following centrifugation. The protein concentration of the crude extract was determined by Bio-Rad Protein Assay (Bio-Rad).

Adduct formation in the presence of NaBH₄
The DNA-trapping reaction was carried out using 120 mM NaBH₄ (instead of 120 mM NaCl). After incubation at 70°C for 10 min the reaction mixture was denatured at 90°C for 3 min in standard SDS–PAGE loading buffer and was then separated by electrophoresis on a 10% SDS–polyacrylamide gel. The gel was dried and analyzed using a phosphoimager. In the case of EcNth, 25 mM NaBH₄ was used and the reaction was carried out at 37°C for 10 min. The subsequent steps were similar to those described above.

Phylogenetic analysis
Distance analysis was performed using neighbor joining in the PAUP program (54). A multiple sequence alignment of the ‘HhH–GPD’ domain was retrieved from the Pfam Web site (http://pfam.wustl.edu; 55–57). Sequences containing a lysine residue in the HhH motif corresponding to Lys120 in EcNth were selected from the Pfam alignment. Sixteen additional sequences were manually added (Aa4, Ap1, Buch1, Cj1, Dr1, Dr2, Dr3, MjOgg, Nn1, Pab1, Pae1, PaNth, Tn1, Tm2, Vc1 and Xf1; see Fig. 9 legend).

**RESULTS**

Homology between PaNth and other DNA N-glycosylase homologs
Open reading frame (ORF), pag5_880, from the hyperthermophilic archaeon *P. aerophilum*, was identified as encoding a putative DNA N-glycosylase by FASTA (51). The amino acid alignments show that it contains all the structural attributes of a [4Fe-4S]-containing HhH DNA N-glycosylase (Fig. 1): a HhH motif, a strictly conserved aspartic acid residue at position 148 (GPD motif) and four cysteine residues for binding of a [4Fe-4S] cluster (FCL motif). Overall, amino acid sequence similarity was seen when comparing this putative DNA N-glycosylase to other characterized [4Fe-4S]-containing HhH DNA N-glycosylases using Bestfit (58): EcNth (37.4% identity), *Micrococcus luteus* UV endonuclease (MIUVendo, 37.6% identity), *Thermotoga maritima* methylenetetrahydrofolate dehydrogenase II (TmMgPorII, 36.5% identity), *P. aerophilum* T/G and U/G mismatch glycosylase (PaMig, 33.3% identity) and *E. coli* adenine-DNA glycosylase (EcMutY, 30.4% identity over 100 amino acid residues). The biochemical data that follows demonstrates that the purified recombinant protein can catalyze the base removal of a modified pyrimidine, a substrate for endonuclease III, but lacks detectable activity for substrates of UV endonuclease, T/G and U/G mismatch glycosylase, and adenine-DNA glycosylase. Therefore, we designate it as *P. aerophilum* endonuclease III homolog (PaNth).
The amino acid alignment shown in Figure 1 reveals that PaNth contains a lysine residue at position 131 (arrowhead indicates position). It appears that this is a strictly conserved residue in the endonuclease III family and is also present in the UV endonuclease from *M.luteus* (26). In contrast, at the corresponding position, methylpurine glycosylase homologs have a strictly conserved glutamic acid residue (24), and the T/G and U/G mismatch glycosylase/adenine-DNA glycosylase homologs have a conserved tyrosine or a serine residue (21,29). In addition, PaNth lacks the adenine-DNA glycosylase-specific C-terminal domain (59).

**Expression and purification of the recombinant PaNth**

The histidine-tagged recombinant PaNth was expressed in *E.coli* and was purified to near homogeneity by affinity chromatography on a Ni²⁺-NTA column (Fig. 2, lane 3). Although the apparent molecular mass, estimated from the SDS gel, was ~30 kDa (Fig. 2), the mass spectrometry study showed the recombinant PaNth to be 26 624 Da (data not shown). This is in close agreement with the size of 26 594 Da predicted from the histidine-tagged pag5_880 ORF sequence. The majority of the purified recombinant PaNth remained soluble after heat treatment at either 70 or 90°C (Fig. 2, lanes 4 and 5) suggesting that it is heat stable. Therefore, 90°C heat treatment was used as a second step purification to further reduce the amount of contaminants from *E.coli* that precipitated at this temperature. The purified PaNth displays a distinct olive color due to the presence of [4Fe-4S] clusters with a corresponding absorption peak at 410 nm (data not shown).

**PaNth glycosylase activity**

Defined oligonucleotide substrates were used for the characterization of PaNth glycosylase activity. Endonuclease III substrates containing a single modified pyrimidine (DHT) paired with each of the four normal DNA bases, were first tested (Fig. 3). Efficient removal of DHT opposite G (Fig. 3A, lane 8), along with weak activity for removal of DHT opposite A, C and T, was observed (Fig. 3A, lanes 7, 9 and 10). The concentration dependence curve of PaNth with DHT/G substrate is shown in Figure 3B and its specific activity for DHT/G is 11 fmol/µg/min at 70°C. *EcNth* was used as a positive control and shows a similar opposite base preference on DHT substrates (Fig. 3C and D). Its specific activity for DHT/G is 0.6 fmol/µg/min at 37°C and is apparently much less efficient than PaNth.

Optimal conditions were studied for PaNth activity with DHT/G. The temperature dependence of the PaNth glycosylase activity is shown in Figure 3B and its specific activity for DHT/G is 11 fmol/µg/min at 70°C. *EcNth* was used as a positive control and shows a similar opposite base preference on DHT substrates (Fig. 3C and D). Its specific activity for DHT/G is 0.6 fmol/µg/min at 37°C and is apparently much less efficient than PaNth.

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activity with DHT/G substrate is shown in Figure 4A. While weak PaNth activity was detected at temperatures below 37 °C (Fig. 4A, lanes 9–11), maximum cleavage was observed at temperatures between 60 and 70 °C (Fig. 4A, lanes 13 and 14). At temperatures above 70 °C, the activity of PaNth decreases (Fig. 4A, lanes 15 and 16).

Figure 4B shows the salt dependence of PaNth with DHT/G substrate. The optimal NaCl concentration for PaNth is between 80 and 160 mM, while little activity was observed in the absence of NaCl (Fig. 4B). In contrast, EcNth prefers low NaCl conditions and even performs best in the absence of NaCl (Fig. 4C).

PaNth AP lyase activity

AP lyase activity is present in DNA N-glycosylases that have a conserved lysine residue in the HhH motif (Fig. 1, arrowhead indicates position). This conserved lysine residue is also present in PaNth suggesting that it may have AP lyase activity. Indeed, in the presence of suitable substrate, PaNth was able to generate a cleaved DNA product rather than just an AP product. To confirm the AP lyase activity, we looked at PaNth’s ability to cleave AP containing DNA derived from UDG-treated U/G substrate. We found that PaNth can efficiently cleave an existing AP site (Fig. 5A). So far, three types of glycosylase/lyase actions have been identified: β-elimination, 3′-OH hydrolysis and β,δ-elimination. The products of these three actions can be distinguished on a polyacrylamide gel due to different mobility: the β,δ-elimination product is faster followed by the 3′-OH product and finally the β-elimination product (60). To define PaNth’s glycosylase/lyase action, the size of the PaNth product was compared with the β-elimination product generated by EcNth. The 3′-OH product, generated by the combination of EcNth and endoIV (61), was also included as a size marker. The PaNth product is the same size as the EcNth β-elimination product (Fig. 5B, lanes 2 and...
and both migrated more slowly than the EcNth/endoIV 3′-OH product (Fig. 5B, lane 4). Therefore, we conclude that PaNth’s glycosylase/lyase action is likely via β-elimination.

Other substrates for PaNth

Another known endonuclease III substrate is dsDNA containing DHU. We used a 55mer containing DHU paired with G at position 19 to test for PaNth cleavage (Fig. 6). The reactions were carried out at 37°C using both non-heat-treated and heat-treated PaNth samples to evaluate whether the observed PaNth activity was heat resistant. In the non-heat-treated PaNth samples as little as 2 ng PaNth was able to nick the DNA at the lesion site to generate a 19mer product (Fig. 6, lane 4). In addition, nuclease activity was observed when using a 100-fold excess of non-heat-treated PaNth, which caused DNA degradation (Fig. 6, lane 2). In the heat-treated PaNth samples the glycosylase/lyase activity was retained (Fig. 6, lanes 5–7), but the nuclease activity was not (Fig. 6, lane 5) indicating that the latter activity was from an E.coli contaminant. EcNth was used as a positive control and also generated the 19mer product (Fig. 6, lane 8), and its activity was heat labile (Fig. 6, lane 9).

Since PaNth shares similar amino acid identity with both EcNth (37.4%) and M1UVendo (37.6%), UV-damaged DNA containing cyclobutane pyrimidine dimers was also tested as a substrate. PaNth did not show any nicking activity towards the DNA containing cyclobutane pyrimidine dimers (data not shown).

T/G and U/G mismatch glycosylase substrates were tested. Only minor cleavage of T/G substrate by PaNth was detected (Fig. 7A, lane 4). The adenine-DNA glycosylase substrate, A/GO, was also tested for possible cleavage by PaNth (Fig. 7B). While only minor activity was observed with the T/GO substrate (Fig. 7B, lane 8), no activity was detected with G/GO, C/GO, A/GO or U/GO substrates (Fig. 7B, lanes 2, 4, 6 and 10). These results are consistent with the structural analysis described earlier, where PaNth lacks some amino acid residues that are conserved either in the T/G and U/G mismatch glycosylases or in the adenine-DNA glycosylases.

We also looked at 8-oxoG DNA glycosylase substrate C/GO (where GO is labeled) and other unmodified base pair mismatches (A/A, A/C, A/G, C/A, C/C, C/T, G/A, G/G, G/T, T/C and T/T). No activity was observed with any of these substrates (data not shown).

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The presence (+) or absence (–) of 2.24 µ (also known as DNA-trapping) by bifunctional N-glycosylase, we utilized the method of DNA adduct formation. For DHT substrates was also observed with EcNth at its optimal growth temperature is 100°C. We designated the protein as an endonuclease III homolog from *P. aerophilum* (PaNth) because it displays glycosylase/lyase activity on a modified pyrimidine, DHT. At optimal NaCl concentration, between 80 and 160 mM, PaNth can remove DHT. Efficient removal is achieved when the lesion is specifically paired with G. A similar opposite base preference for DHT substrates was also observed with EcNth at its optimal low NaCl concentrations, although the efficiency was less than PaNth. While this manuscript was in preparation, Asagoshi et al. published a paper showing significant activity of EcNth and a murine endonuclease III homolog (mNth1) when DHT was paired with G (65). These findings are consistent with our observations for EcNth. The fact that DHT/G preference is observed for endonuclease III homologs from organisms in each of the three kingdoms (Bacteria, Archaea and Eucarya) suggests that DHT/G preference is a common endonuclease III characteristic. Apparently DHT/G mispairs are not found under physiological conditions and therefore what we may be seeing is DHT’s resemblance to another pyrimidine lesion that pairs with G. It may be this unknown paired lesion that is biologically relevant. It is most likely generated by direct modification of an original cytosine base or one of its derivatives (such as 5-methylcytosine), or by a replication error which inserts this unknown lesion opposite G. The observed high activity of PaNth could reflect a more efficient repair of such lesions in *P. aerophilum*, and may have evolved due to the additional load of DNA damage caused by an extreme environment, such as high temperature.

In *E. coli*, two other DNA glycosylases, endonuclease VIII and MutM (formamidopyrimidine DNA glycosylase or Fpg), have substrate spectra that overlap with that of EcNth (66,67). We searched the *P. aerophilum* genome for putative sequence homologs of *E. coli* endonuclease VIII and Fpg, but found none. *Pyrobaculum aerophilum* has a U/G and T/G specific three are obligate parasitic bacteria. Notably, characterized DNA N-glycosylases that have different substrate preference (i.e. endonuclease III homologs and 8-oxoG DNA glycosylase homologs) cluster separately on the tree. When we look only at the endonuclease III/UV endonuclease homologs, we find that they cluster into five distinct groups. One of these contains PaNth and two other putative DNA N-glycosylase homologs from *Aeropyrum pernix* (Ap, 43.5% identity) and from *Salinobacter solfatarius* (Ss, 41.8% identity).

**DISCUSSION**

In this study, we report characterization of a putative DNA N-glycosylase homolog from a hyperthermophilic archaeon *P. aerophilum*, whose optimal growth temperature is 100°C. When we perform the phylogenetic analysis of region A, we obtained the same branching pattern seen in Figure 10A. However the tree for the base binding region, region B, fails to discriminate the five groups (data not shown), apparently due to both less divergence in this region among these groups and the shorter length of region B in comparison with region A.

**Phylogenetic study of endonuclease III homologs**

DNA N-glycosylase homologs, which contain a conserved lysine residue in the HhH motif corresponding to Lys120 in EcNth (Fig. 1, arrowhead indicates position), were used to construct a phylogenetic tree (Fig. 10A). So far, among 34 completed archael and bacterial genomes, it appears that only three lack endonuclease III homologs: *Mycoplasma genitalium*, *Mycoplasma pneumoniae* and *Ureaplasma urealyticum*. All
DNA glycosylase (PaMig; 21) which has extensive shared amino acid sequence identity (33.3%) to PaNth. It seems PaMig is quite specific for U or T opposite G but has no detectable activity on DHT/G with T/G and T/GO substrates. The definitive solution for finding a backup system for PaNth in *P. aerophilum* would be to construct a PaNth knockout strain and look for any residual activity with DHT/G substrate.

Our phylogenetic study, using completely sequenced whole genomes, shows that all except three obligate parasitic bacteria contain an endonuclease III/UV endonuclease homolog, underlining the importance of BER in maintaining DNA stability in both mesophilic and thermophilic organisms. Endonuclease III/UV endonuclease homologs are clustered into five major groups, each with unique conserved amino acid residues. One example of this is seen in region A, which is marked by amino acid residues occurring >80% of the time within a single group and outside the group only once or not at all. The arrow marks amino acid residues as follows: red, AVFPMILW (small and hydrophobic); blue, DE (acidic); magenta, RK (basic); green, STYHCNGQ (hydroxyl and amine).
regions A, our work helps for future annotation of putative DNA N-glycosylase sequence homologs.

At this point, the nature of these multiple endonuclease III/UV endonuclease homolog clusters, identified by phylogenetic analysis, is not clear but many possibilities exist. One possibility is that these clusters may be subfamilies of endonuclease III/UV endonuclease, each having a different function in terms of substrate preference. A possible example of this clustering by enzymatic function is the branch containing the single characterized UV endonuclease. Despite the fact that UV endonuclease and endonuclease III share close sequence homology, they catalyze different glycosylase reactions. In our phylogenetic analysis six putative endonuclease III homologs cluster with M.luteus UV endonuclease. They also contain some amino acid residues in region A that are unique to this cluster. The next logical step is to see whether these homologs have similar catalytic function to UV endonuclease. Another possibility for the multiple endonuclease III/UV endonuclease homolog clusters is that the pattern of clustering reflects amino acid sequence differences that occur either due to interaction with other protein (52) or simply due to independent evolution along the organism lineage. Further characterization of BER pathways and all their related cofactors will help clarify any biochemical basis for the clustering of endonuclease III/UV endonuclease homologs.

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