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Structure-specific nuclease activities in *Pyrococcus abyssi* RNase HII

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Running Title: Substrate structure requirements of PabRNase HII

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ABSTRACT

Faithful DNA replication involves the removal of RNA residues from genomic DNA prior to ligation of nascent DNA fragments in all living organisms. Because the physiological roles of archaeal type 2 RNase H are not fully understood, substrate structure requirements for detection of RNase H activity need further clarification. Biochemical characterisation of a single RNase H detected within the genome of *Pyrococcus abyssi* showed that this type 2 RNase H is a Mg- and alkaline- dependent enzyme. *Pab*RNase HII showed ribonuclease activity and acted as a specific endonuclease on RNA-DNA/DNAs. This specific cleavage, one nucleotide upstream of the RNA-DNA junction, occurred on a substrate in which RNA initiators had to be fully annealed to the complementary DNA template. On the other hand, a 5′-RNA flap Okazaki fragment intermediates impaired *Pab*RNase HII endonuclease activity. Furthermore, introduction of mismatches in the RNA portion near the RNA-DNA junction decreased both specificity and efficiency of cleavage by *Pab*RNase HII. Additionally, *Pab*RNase HII could cleave a single ribonucleotide embedded in a double-stranded DNA. Our data revealed *Pab*RNase HII as a dual-function enzyme likely required for completion of DNA replication and DNA repair.
INTRODUCTION

DNA replication in all living organisms takes place concurrently on two separate strands. The lagging strand consists of multiple discontinuous segments called Okazaki fragments, whereas the leading strand comprises one large continuous segment. Production of each individual lagging strand by DNA polymerase is primed by a short stretch of RNA. Later on, these RNA primers are eliminated and the resulting gap is filled with deoxyribonucleotides prior to ligation. Priming and DNA elongation at the replication fork involve a set of specialized polymerising enzymes which differ from replicative DNA polymerases, and one another, to correct erroneously inserted nucleotides. In archael cells, the priming complex lacks proofreading 3’-5’ exonuclease activity (13, 14) present in the replicative DNA polymerases B and D (1, 8). Consequently, mismatches in the vicinity of the RNA-DNA junction could arise in replicating cells, as already observed in eukaryotes (28, 31). Similarly, single ribonucleotides incorporated during DNA replication (20, 26) or by external agents (32) would represent another source of erroneous nucleotides. Persistence of residual RNA during DNA replication would be detrimental for the cells, suggesting that a combination of specific and efficient nucleolytic processes is absolutely required to preserve DNA integrity.

Ribonucleases H (RNase H) are enzymes which degrade the RNA portion of RNA/DNA or RNA-DNA/DNA duplexes (29). RNases H are classified into two major families, type 1 and type 2, based on amino acid sequence (21). The type 1 family includes bacterial RNase HI, mammalian RNase HII, the RNase H domain of reverse transcriptase and archael RNase HI and the type 2 contains bacterial RNase HII and RNase HIII, mammalian RNase HI and archael RNase HII (21). While type 2 RNase H enzymes are universally conserved in the three domains of life, their physiological role remains elusive. Much less is known about the type 2 family compared to the type 1 RNase H enzymes. The multiplicity of RNases H within
a single cell complicates the situation, although presumable roles in DNA replication, DNA
repair and transcription have been assigned as recently reviewed (2, 30). In archaea, structural
and biochemical characterization of type 2 RNases H (3, 4, 10, 12, 16, 19) suggested they can
initiate RNA removal from DNA duplexes, based on their ability to specifically cleave 5’ to
the junctional ribonucleotide. However, despite this information, the physiological role of
type 2 RNases H still remains elusive. They could be involved in the completion of either
leading or lagging strands or both. Additional biochemical experiments with catalytic
intermediates should provide valuable knowledge on the participation of these archaeal
cellular enzymes at the replication fork and in DNA repair.

To investigate this question, we have designed a set of RNA/DNA duplex, cognate RNA-
DNA/DNA duplex (15), a single ribonucleotide embedded in a DNA duplex (DNA-1RNA-
DNA/DNA) as substrates, and employed type 2 RNase H from the hyperthermophilic deep-
sea euryarchaeon Pyrococcus abyssi (Pab), PabRNase HII. Since a single rnh gene exists in
the genome of P. abyssi (6), RNase HII is likely the key enzyme involved in RNA elimination
in this organism. Thus, PabRNase HII can be considered as the representative of type 2
RNase H.

Here, we analysed the cleavage specificity of PabRNase HII for substrates with Okazaki
fragment-like structure. We also tested PabRNase HII activity on Okazaki fragment-like
substrates in the presence of mismatched base pair in order to assess the molecular
mechanism of recognition of the RNA-DNA junction and the subsequent cleavage specificity.
In addition, we examined whether PabRNase HII can incise the DNA backbone on the 5’-side
of a single ribonucleotide embedded in a DNA duplex. Our data provide substantial evidences
that the single RNase H in P. abyssi has a dual role in maintenance of genome integrity. The
results from this study are further discussed to define potent roles of type 2 RNase H from P.
abyssi in the resolution of RNA fragments at the replication fork and in the repair of single
embedded ribonucleotides.
MATERIALS AND METHODS

Nucleic acid substrates.
Gel-purified oligonucleotides for preparing the substrates for RNase HII assays were purchased from Eurogentec (Belgium) and their sequences are listed in Table 1. Fluorescent labelling at the 5’- end was performed with the 5’ End Tag kit labelling system from Vector Laboratories (California). Free fluorescent dyes were removed on MicroSpinTM G-25 columns. For some experiments, 5’-end- or 3’-end-fluorescent labelled oligonucleotides were chemically synthesized and HPLC-purified by Eurogentec (Belgium). To generate the substrates for the RNase HII assays, the appropriate oligonucleotides were mixed in 1:1 molar ratio in 20 mM Tris–HCl (pH 7.4), 150 mM NaCl, heated to 75°C and slowly cooled to room temperature.

Cloning, production and purification of PabRNase HII.
The gene encoding PabRNase HII (PAB0352) was cloned into the pQE-80L expression vector (Qiagen). PabRNase HII was overexpressed in E. coli strain BL21-CodonPlus-RIL strain (Stratagene) as a histidine-tagged protein and purified to near homogeneity via Ni-NTA beads (Qiagen) and S200 gel filtration using fast protein liquid chromatography (GE Healthcare) as previously described (16). Protein integrity was analyzed by MALDI-TOF analyses (Innova Proteomics, France). PabRNase HII purity was controlled by SDS–PAGE gradient gel (4-20 %) electrophoresis (Thermo Scientific).

Amino acid sequence alignments and secondary structure.
Amino acid sequence alignments have been constructed by ClustalW2 (available at [www.ebi.ac.uk/clustalW2/](http://www.ebi.ac.uk/clustalW2/)). Secondary structure elements calculated with the program
ESPript 2.2 (available at http://espript.ibcp.fr/ESPript/ESPript/) refer to the structure of TkoKOD1RNase HII (PDB: 1IO2).

**Assays for RNase HII activity.**

Assays to monitor cleavage by *Pab*RNase HII were performed in RNase HII buffer (10 µl) containing: 50 mM Tris-HCl (pH 8), 5 mM dithiothreitol, 5 mM MgCl₂ and 50 nM of DNA substrates. Enzymes were diluted from concentrated stocks in 20 mM Tris-HCl (pH 7.5), 20 % glycerol prior to usage. Enzyme concentrations for a typical reaction ranged from 4 to 400 nM, unless otherwise specified. After addition of *Pab*RNase HII, reactions were incubated at 60°C for 30 minutes and stopped on ice with 15 µl of stop buffer (98 % formamide, 10 mM EDTA). Samples were heated at 95°C for 5 minutes. A base hydrolysis ladder was prepared by incubation of the labelled RNA-DNA strand (10 µM) with snake venom phosphodiesterase I (0.018 units) for 10 minutes at 37°C. Product analysis was carried out by electrophoresis on 15 % denaturing polyacrylamide gels. After visualisation with a Mode Imager Typhoon 9400 (GE Healthcare), quantification of the results was performed using ImageQuant 5.2 software.

In all cases, the percentage of substrate hydrolysis was determined by the products / (products + substrate) ratio, allowing a correction for loading errors and a comparison of cleavage efficiency irrespective of the different products generated.

To analyse divalent cations or pH dependence, RNase HII assays were carried out with 50 nM of *Pab*RNase HII and 50 nM of the S1 substrate at 60°C for 30 min. Data are the average of triplicate measurements and are shown with standard deviations (SD).

To determine the kinetic parameters, steady-state kinetic reactions were carried out in the same conditions as described above by using substrate 1 at concentrations ranging from 0.03 to 3 µM. Initial velocity experiments were monitored as a function of time with 60 nM of *Pab*RNase HII at 60°C such that the rate of converted substrate did not exceed 20 % of the
Velocity measurements were reported as the amount of hydrolysed substrate (µM) over time (min). The observed rates of converted substrates with \( \text{PabRNase HII} \) were firstly determined from Lineweaver-Burk plots. The data were fit by nonlinear regression using the Marquardt-Levenberg algorithm (EnzFitter 2.0, BioSoft) to the Michaelis-Menten equation. Kinetic parameters, \( K_m \) and \( V_{\text{max}} \), were obtained from the fit and were used to calculate the catalytic efficiency \( (k_{\text{cat}}/K_m) \) of \( \text{PabRNase HII} \). The kinetics values are the average of at least triplicate determinations and are shown with standard deviations (SD). Any adjustments to the above are noted in the Figure Legends.
RESULTS

Archaeal RNase HII homologues

PabRNase HII showed amino acid sequence similarities of 75.2 % with TkoRNase HII, and 65 % with AfuRNase HII. Analysis of protein primary structures and related secondary structures outlined subtle differences between the three proteins (Fig. 1A). On the one hand, PabRNase HII and AfuRNase HII are isoelectric at basic pH (isoelectric point values of 9 and 7.6, respectively), whereas TkoRNase HII is isoelectric at acidic pH (isoelectric point of 5.5). On the other hand, PabRNase HII, TkoRNase HII and AfuRNase HII exhibit conserved secondary structure elements, with the exception that the α9-helix is incomplete in AfuRNase. This secondary structure element is important for TkoRNase HII to bind the substrate (19).

Structural alignments resulted in the identification of conserved active site residues (Asp7, Glu8, Asp105 and Asp135) in PabRNase HII, suggesting a similar catalytic mechanism between the three enzymes.

Enzymatic properties of P. abyssi RNase HII

His-tagged PabRNase HII was overproduced in E. coli and purified to give a single band on SDS-PAGE (Fig. 1B, lane 2). Reactions were carried out at 60 °C, the optimum temperature for PabRNase HII activity (16). PabRNase HII was assayed under different pH and ionic conditions, varying both the nature and the concentration of divalent cations, according to the general procedure described in Materials and methods using the RNA-DNA/DNA substrate (S1). The optimum pH for its activity was observed between pH 8.0 and 8.5 (Fig. 1C). However, at pH 7.5 and 9, the enzyme retained about 60 % of the activity measured at pH 8.
PabRNase HII exhibited enzymatic activity in the presence of MgCl₂, MnCl₂ and CoCl₂ (Fig. 1D). While PabRNase HII activity was entirely dependent on the presence of a divalent cation, the enzyme was not active in the presence of NiCl₂, FeCl₂, ZnCl₂, CaCl₂ or CuSO₄. The most preferred metal ion for PabRNase HII was MgCl₂ but the MnCl₂ and CoCl₂ could substitute for the MgCl₂ with reduced cleavage activity. As shown in Fig. 1D, the metal concentrations which gave the highest enzymatic activity were 5 mM for MgCl₂ and 2 mM for MnCl₂ and CoCl₂. Substrate hydrolysis in the presence of 5 mM for MgCl₂ was 1.5- and 3.2- fold higher than those determined at 2 mM for MnCl₂ and CoCl₂, respectively.

The kinetic parameters of PabRNase HII were determined in the presence of RNA-DNA/DNA substrate (S1) and 5 mM MgCl₂. The results are summarized in Table 2 and compared to that of type 2 RNase H archaeal homologue from Archaeoglobus fulgidus (AfuRNase HII) as already described (4). Interestingly, AfuRNase HII showed stronger substrate binding affinity than PabRNase HII, as attested by a 10-fold lower $K_m$ value, while the catalytic rate constants of the two enzymes were similar (Table 2). As a consequence, PabRNase HII displayed a lower catalytic efficiency on RNA-DNA/DNA substrate as compared to AfuRNase HII (Table 2).

Ribonuclease activity of P. abyssi RNase HII

Firstly, we examined whether PabRNase HII could cleave the RNA strand of a RNA/DNA duplex (S11). As shown in Fig. 2, different 5’-terminal RNA products accumulated depending on the enzyme concentration, indicating that PabRNase HII exhibited endoribonuclease activity. Because control assays without enzyme showed background degradation of the RNA primer (Fig. 2A, lane 2), they were subtracted from cleavage products signal. Less than 19 % intact RNA was present upon incubation with 100 nM PabRNase HII for 30 min at 60°C (Fig.
2A, lane 6). Multiple cleavage sites were detected (Fig. 2A, lanes 2-8) and comparative analysis of products with those from the snake venom phosphodiesterase digest of 5’-end labelled 32 nucleotide (nt) RNA ladder (Fig. 2A, lane 1) pointed out main cleavage events (6-, 8-, 9-, 12-, 13- and 17-nt). Moreover, further processing of short RNA fragments could be observed by increasing enzyme concentrations. It is important to note that PabRNase HII used in this study did not exhibit nuclease activity on single-stranded RNA (Fig. 2B). All together, these results provided evidence that PabRNase HII acts as an endoribonuclease on RNA/DNA duplexes.

**Structure-specific cleavage activities in P. abyssi RNase HII**

The absence of cleavage specificity of RNA/DNA duplexes prompted us to look for digestion of other relevant physiological substrates. We hypothesize that PabRNase HII participates in the mechanism of RNA primer removal, an activity which can occur once at the leading strand or much more frequently at the lagging strand. Therefore, we explored the cleavage specificity of PabRNase HII for substrates with Okazaki fragment-like intermediates.

*PabRNase HII specifically cleaves RNA-initiated DNA segments fully annealed to a DNA template*

We initially began to examine whether PabRNase HII could hydrolyse a cognate double-stranded Okazaki fragment (15). A single strand composed of 12 nucleotides of RNA (RNA12nt) followed by 18 nucleotides of DNA (DNA18nt), fluorescently labelled at the 5’-end, was annealed to a complementary 30 nucleotides DNA template to form the S1 substrate as shown in Fig. 3A. When this substrate was incubated in the presence of increasing PabRNase HII amounts, a major product appeared (Fig. 3B, lanes 2-5). This oligomer was
shown to correspond to 11 nucleotides RNA by migration with respect to a snake venom phosphodiesterase-generated digest of RNA12ntDNA18nt (Fig. 3B, lane 1). In addition to this main cleavage site, minor cleavage sites characteristic of non-specific nuclease activity were also found throughout the length of the RNA (Fig. 3B, lanes 2-5). Over time, PabRNase HII activity released the same oligomer which was basically free of any additional shorter fragments, suggesting that this product was not transiently formed and prevailed during the reaction (data not shown). Interestingly, PabRNase HII did not hydrolyse single-stranded RNA12ntDNA18nt (Fig. 3B, lanes 6-9) indicating that cleavage is dependent on the heteroduplex structure. It is of note that the pale band at ∼9-nt present at relatively constant levels did not correspond to a specific cleavage product (Fig. 3B, lanes 2-9). Absence of specific cleavage was also observed with substrates lacking the complementary DNA template to either the RNA12nt or the DNA18nt sequence (data not shown). While a fully annealed RNA12ntDNA18nt/DNA is definitely required to detect cleavage specificity, PabRNase HII did not hydrolyse the complementary DNA template (Fig. 3B, lanes 10-13). Shorter bands were not due to cleavage activity since they were detectable in all lanes with equal intensities even in the absence of enzyme. These data demonstrated that PabRNase HII can act endonucleolytically on initiator RNA and displays a specific cleavage activity dependent on the heteroduplex structure.

PabRNase HII specifically cleaves the fully annealed RNA strand of Okazaki fragment-gapped intermediates but not a 5’-RNA flap

Both at the leading and lagging strands, sequential enzymatic steps are thought to be part of the RNA primer elimination mechanism in P.abyssi. As a consequence, diverse structural Okazaki fragment-like substrates would arise. Therefore, we examined whether structural intermediates (S4, S5 and S6) which can be captured during the process (outlined in Fig. 4A)
could direct the cleavage activity of *Pab*RNase HII. On the 40-gapped S4 intermediate composed of an upstream DNA primer and a downstream RNA-DNA fragment fully annealed to the complementary DNA template, *Pab*RNase HII specifically cleaved the RNA segment, releasing one ribonucleotide attached to the DNA segment (Fig. 4B, lanes 1-4). During the elongation step, the size of the gap would decrease to reach the next RNA initiator. By reconstitution of model transient substrates, we demonstrated that both the 20-nt gapped S5 intermediate (Fig. 4B, lanes 5-8) and a nicked intermediate (data not shown) were specifically cleaved. Collectively, cleavage efficiencies of the gapped and nicked Okazaki fragment intermediates were comparable to those of double-stranded RNA-DNA fragments (Fig. 3B, lanes 2-5). However, on a 5’-RNA flap which can result from strand displacement activity by *Pab*polD of the next Okazaki fragment (11), *Pab*RNase HII did not significantly release oligomers (Fig. 4B, lanes 9-12). It is of note that a faint intensifying band at 8-nt (Fig. 4B, lanes 9-12) did not correspond to a specific cleavage product. These results clearly indicated that *Pab*RNase HII is not involved in the cleavage of single-stranded RNA initiator despite the presence of surrounding DNA duplexes. These data are consistent with our observations from Fig. 3B that *Pab*RNase HII exclusively cuts double-stranded RNA-DNA/DNA substrates. Importantly, we demonstrated that *Pab*RNase HII cleaves the RNA initiator fully annealed to the complementary DNA template independently of the size of the gap. In addition, we provided evidence that a 5’-RNA flap is not an appropriate substrate for *Pab*RNase HII, suggesting the requirement of additional enzymes to fully ensure the removal of Okazaki fragment intermediates at the lagging strand.

*Pab*RNase HII specifically cuts the RNA-DNA/DNA when the RNA is completely annealed to the DNA template
The above results indicated that PabRNase HII specifically cleaves the RNA in an RNA-DNA/DNA duplex one ribonucleotide upstream of the RNA-DNA junction. Based on this observation, we attempted to gain further information about the structure-specific recognition of the RNA-DNA junction. We predicted that mismatches located either downstream or upstream of the site of cleavage would alter the structure of the junction and prevent PabRNase HII from recognizing and cutting the substrate. Such substrates, which can be created during priming and DNA synthesis in eukaryotes (28, 31), could also be relevant in P. abyssi cells. In particular, the priming heterodimeric polymerase in P. abyssi, Pabp46/41 complex, does not possess 3’-5’ exonucleolytic activity and can consequently misincorporate nucleotides, creating mismatched base pairs at or near the RNA-DNA junction.

The complementary DNA template was designed to produce specific mismatches with the RNA12ntDNA18nt strand (Fig. 5A). When the mismatch was the deoxynucleotide downstream of the site of cleavage, PabRNase HII efficiently cleaved the S7 substrate and cut at one site into the RNA segment, leaving a monoribonucleotide attached to the DNA18nt strand (Fig. 5B, lanes 2-5). Cleavage efficiencies were still comparable to those of model Okazaki fragments described above. This result seems to point out that a deoxynucleotide mismatched Okazaki fragment does not affect recognition and specific cleavage by PabRNase HII. We next considered that ribonucleotide mismatches positioned downstream (Fig. 5A, S8 substrate) or upstream (Fig. 5A, S9 substrate) of the cutting site would be crucial for directing the cleavage specificity of PabRNase HII. Interestingly, the presence of the ribonucleotide just downstream of the cutting site induced random endonucleolytic cleavage with predominant products (Fig. 5B, lanes 6-9 and Fig. 5C, S8 substrate) and the percent of hydrolysed products was equivalent to that of Okazaki fragment-like substrates. When the ribonucleotide mismatch was positioned upstream of the site of cleavage, random endonucleolytic activity was enhanced but cleavage efficiencies were lowered (Fig. 5B, lanes
10-13). Multiple cleavage sites due to the loss of specificity appeared (Fig. 5C, S9 substrate). Taken together these data showed for the first time that an archaean RNase HII requires complete hybridization of the RNA segment to the DNA template in order to confer specific cleavage of RNA-DNA/DNA duplexes.

**PabRNase HII specifically cuts a single embedded ribonucleotide in a DNA duplex**

We anticipated that *Pab*RNase HII nuclease could act on a single ribonucleotid embedded in DNA. A single ribonucleotide in a DNA duplex could arise via misincorporation of ribonucleotide during DNA synthesis or by ligation of the monoribonucleotide attached to the DNA after cleavage of Okazaki fragments by type 2 RNase H (26). To determine whether an embedded ribonucleotide in DNA (S10 substrate) is a hydrolysable substrate, endonuclease activity of *Pab*RNase HII was carried out. Fig. 6B, lanes 8-11 demonstrated that *Pab*RNase HII was able to recognise and to cleave endonucleolytically on the 5'-side of an embedded monoribonucleotide. Additional fragments, shorter than the released 11-nt, were faintly detectable. Basically, cleavage efficiencies of a single embedded ribonucleotide were similar to those of model Okazaki fragment S1 substrate (Fig. 6B, compare lanes 2-5 and lanes 8-11).

Overall, we showed that *Pab*RNase HII is active on single embedded ribonucleotides in a DNA duplex and releases a major product consisting of a single ribonucleotide on the 5’-end of the downstream DNA segment.
Two types of RNase H, type 1 and type 2, have been identified in a multiplicity of archaeal genomes. While most archaeal microorganisms have only one type of RNase H, a few archaea such as *Sulfolobus tokodaii* and *Haloferax volcanii* possess both types of RNase H. Although the physiological significance of multiple rnh genes in single archaeal genomes is not well understood, RNases H are thought to be involved in important cellular processes (3, 10, 16, 23, 24). Interestingly, archaeal type 2 RNase H appears more universal because the encoding gene is distributed in almost all archaeal genomes. Sequence comparison within archaeal type 2 RNases H revealed a high degree of sequence similarity with conserved active site residues, suggesting that these enzymes may have common biochemical properties (3, 9). In this report, we demonstrated that *Pab*RNase HII, type 2 RNase H from *P. abyssi*, is as an alkaline enzyme. This property seems to be a hallmark of type 2 thermostable RNases H (3, 9, 22). In addition, *Pab*RNase HII appeared to prefer the Mg$^{2+}$ ion for RNase activity rather than Mn$^{2+}$ or Co$^{2+}$. Distinct metal dependencies have been described for *Archaeoglobus fulgidus* and *Thermococcus kodakaraensis KOD1* RNase HII with Mn$^{2+}$- or Co$^{2+}$-preference, respectively (3, 9). Metal ion usage by archaeal RNases HII may be a consequence of the environmental conditions they thrive. It may also dictate the substrate requirement for hydrolysis and confer a specialised function to the enzyme in the maintenance of genome integrity. Determination of kinetic parameters highlighted that the homologous archaeal enzymes, *Afu*RNase HII and *Pab*RNase HII, showed distinct catalytic efficiencies for RNA-DNA/DNA substrates. These results mainly reflected differences in substrate binding affinity. In general, biochemical discrepancies observed between the three enzymes are possibly related to variations in secondary structure elements and physicochemical parameters (e.g., isoelectric point). Despite these subtle differences, archaeal RNase HII seem to possess conserved structural features
required to specifically recognize a comparable region of the substrates, and to produce similar products. Like other type 2 archaeal RNase H, PabRNase HII behaved as an efficient endoribonuclease on RNA/DNA duplexes, stalling at particular sites (3, 9). Moreover, most of the biochemical features of PabRNase HII overlapped those of the eukaryotic equivalent, type 2 RNase H, described as a key enzyme in Okazaki fragment processing (17).

With diverse constructs representing replication-fork intermediates, PabRNase HII made structure-specific endonucleolytic cleavage in the RNA initiator, leaving a single ribonucleotide at the 5′-end of the RNA-DNA junction. Cleavage 5′ to the junctional ribonucleotide required the presence of double-stranded substrates with the RNA segment fully annealed to the complementary strand. Gapped double-stranded substrates containing RNA-DNA junctions did not alter cleavage specificity. However, a single-stranded 5′-RNA flap was resistant to cleavage activity, indicating that PabRNase HII does not carry out this reaction at the replication fork. On the other hand, other results have demonstrated that the structure-specific nuclease, Flap endonuclease I (Fen I), can cleave substrates with RNA flap structures, bypassing the need for Rnase HII in Okazaki fragment processing (18, 27).

Furthermore, we demonstrated that mismatches in the RNA portion, produced by erroneous priming and polymerising activities during initiation of DNA replication in eukaryotes (28, 31), resulted in loss of specificity by PabRNase HII. These results demonstrate, for the first time, that the RNA residues in the vicinity of the RNA-DNA junction are key structural determinants for cleavage specificity of type 2 archaeal RNase H. Notably, archaeal type 2 RNase H seems to differ from eukaryotic type 2 (17) in that it recognizes the RNA strand rather than the RNA-DNA junction. Possibly, the RNA portion of the RNA-DNA junction annealed to DNA template adopts an intermediate helical structure, which might target RNase HII recognition and induce specific cleavage. This hypothesis is sustained by the observation that RNA/DNA and DNA/DNA duplexes form A-type and B-type helices, respectively (5, 7).
We recently proposed a model of DNA replication in *P. abyssi* that involves the family B DNA polymerase, *Pab*polB, at the leading strand and the family D DNA polymerase, *Pab*polD, at the lagging strand (11). This model is reinforced by complementary studies demonstrating that *Pab*polB is likely the leading strand DNA polymerase (25). Typically, *Pab*polD has the capacity to displace the downstream fragment including the RNA initiator, while *Pab*polB is not active on this substrate. In this situation, RNA-initiated DNA segments fully annealed to a DNA template would arise only at the leading strand. Because *Pab*RNase HII cannot cleave 5’-RNA flap templates, *Pab*RNase HII would recognize the annealed RNA primer at the leading strand and promotes its endonucleolytic cleavage. The resulting 5’ phosphorylated junction ribonucleotide attached to the DNA would be subsequently displaced by *Pab*polB and cleaved by *Pab*Fen I, prior to ligation by *Pab*DNA ligase I. Thus, the functional importance of RNase HII in the completion of leading strand DNA replication in *P. abyssi* awaits the *in vitro* reconstitution of this multi-step enzymatic process (manuscript in preparation). Despite common biochemical properties with the eukaryotic type 2 RNase H, single archael RNases HII could be cellular enzymes involved in the removal of RNA residues at the leading strand rather than at the lagging strand. Such biological assumptions would indicate that these microorganisms have evolved differently by targeting analogous enzymes to unrelated biological functions.

Moreover, we demonstrated that *Pab*RNase HII is able to cleave at the 5’-end of single embedded ribonucleotides with similar efficiency as at cognate Okazaki fragments (15). Since such structural substrates can appear *in vivo* during Okazaki fragment processing from intrinsic RNA ligation activity or erroneous nucleotide incorporation (26) and during exposure to external damaging agents (32), we suggest that *Pab*RNase HII can participate in the removal of inappropriate ribonucleotides from the hyperthermophilic chromosome. These biochemical characteristics would imply that *Pab*RNase HII promotes the initial step of the
repair process as already observed in eukaryotes (26). However, reconstitution of the complete enzymatic process awaits further assessment.
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FIGURE LEGENDS

Fig. 1 Enzymatic properties of PabRNase HII. (A) Alignment of the amino acid sequences of archaeal RNase HII homologues. Sequences are from the three euryarchaeota species, *P. abyssi* (Pab, accession number, gi: 14520734), *T. kodakaraensis KOD1* (Tko, accession number, gi: 57640740), *A. fulgidus* (Afu, accession number, gi: 11498229). Conserved amino acid residues are shaded black. Similar amino acid residues are framed black. Proposed active sites residues are indicated by asterisks. Secondary structure is shown above the sequences, denoting β-sheets (arrows) and α-helices (ribbons). (B) SDS-PAGE gradient gel (4-20 %) of purified, recombinant His6-tagged PabRNase HII (0.5 µg; lane 2) and molecular mass markers (lane 1) stained with Coomassie Blue (C) pH dependence. The enzymatic activities were determined at 60°C for 30 min in reaction buffer containing 50 mM Tris-HCl, 5 mM dithiothreitol, 5 mM MgCl2, 50 nM of PabRNase HII and 50 nM of RNA-DNA/DNA substrate (S1) with pH values ranging from 5 to 10. Data are the average of triplicate measurements. (D) Divalent cation dependence. The enzymatic activities were determined at 60°C for 30 min in reaction buffer containing 50 mM Tris-HCl (pH 8), 5 mM dithiothreitol, 50 nM of PabRNase HII and 50 nM of RNA-DNA/DNA substrate (S1) at the indicated concentrations of MgCl2 (◆), MnCl2 (Δ) and CoCl2 (■). Data are the average of triplicate measurements.

Fig. 2 Ribonuclease activity by PabRNase HII. (A) Indicated amounts of PabRNase HII were incubated with the S11 substrate (lanes 2-8) and a base-hydrolysed ladder (lane 1) was prepared as described (see Materials and methods section). 5’-end fluorescently labelled products were visualised with a Mode Imager Typhoon 9400 (GE Healthcare) and quantification was performed using Image Quant 5.2 software. (B) PabRNase HII was
incubated with the 32-base single-stranded RNA oligonucleotide at the indicated amounts (lanes 1-4). An 8-nt RNA oligonucleotide was used as a ladder (lane 5). 5’-end fluorescently labelled products were visualised with a Mode Imager Typhoon 9400 (GE Healthcare).

Fig. 3 *Pab*RNase HII specifically cleaves RNA-initiated DNA segments fully annealed to a DNA template. (A) Substrate structure representations of S1, S2, and S3. The thick line and the closed circle represent the RNA portion and the fluorescent label, respectively. (B) Indicated amounts of *Pab*RNase HII were incubated with S1 substrate (lanes 2-5), S2 substrate (lanes 6-9) and S3 substrate (lanes 10-13). A base-hydrolysed ladder (lane 1) was prepared as explained in the Materials and methods section. Fluorescent-labelled products were visualised with a Mode Imager Typhoon 9400 (GE Healthcare) and quantification was performed using Image Quant 5.2 software.

Fig. 4 *Pab*RNase HII specifically cleaves the fully annealed RNA strand of Okazaki fragment-gapped intermediates but not a 5’-RNA flap. (A) Substrate structure representations of S4, S5, and S6. The thick line and the closed circle represent the RNA portion and the fluorescent label, respectively. (B) Indicated amounts of *Pab*RNase HII were incubated with S4 substrate (lanes 1-4), S5 substrate (lanes 5-8) and S6 substrate (lanes 9-12). An 18-nt nucleotide was used as an appropriate ladder (lane 13). Fluorescent-labelled products were visualised with a Mode Imager Typhoon 9400 (GE Healthcare) and quantification was performed using Image Quant 5.2 software.

Fig. 5 *Pab*RNase HII specifically cuts the RNA-DNA/DNA when the RNA is completely annealed to the DNA template. (A) Substrate structure representations of S7, S8, and S9. The thick line and the closed circle represent the RNA portion and the fluorescent label,
respectively. (B) Indicated amounts of PabRNase HII were incubated with S7 substrate (lanes 2-5), S8 substrate (lanes 6-9) and S9 substrate (lanes 10-13). An 11-nt nucleotide was used as an appropriate ladder (lane 1). Fluorescent-labelled products were visualised with a Mode Imager Typhoon 9400 (GE Healthcare) and quantification was performed using Image Quant 5.2 software. (C) Graphical representation of sites and extents of cleavage in mismatches RNA-DNA/DNA substrates. Cleavage sites are denoted by different bars. Deoxyribonucleotides and ribonucleotides are shown by uppercase and lowercase letters, respectively.

Fig. 6 PabRNase HII specifically cuts single embedded ribonucleotide in a DNA duplex.

(A) Substrate structure representations of S1 and S10. The thick line and the closed circle represent the RNA portion and the fluorescent label, respectively. (B) Indicated amounts of PabRNase HII were incubated with S1 substrate (lanes 2-5) and S10 substrate (lanes 8-11). Both substrates and the corresponding hydrolysed products were manually labelled. Lanes 1 and 6 are appropriate 11-nt and 12-nt ladders for hydrolysed S1 substrates. Lanes 7 and 12 are suitable 11-nt and 12-nt ladders for hydrolysed S10 substrates. Fluorescent-labelled products were visualised with a Mode Imager Typhoon 9400 (GE Healthcare) and quantification was performed using Image Quant 5.2 software.

TABLE 1 Oligonucleotide sequences used to create structural duplex substrates

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Primes</th>
<th>Template</th>
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<tr>
<td>S1</td>
<td>2, 2</td>
<td>8</td>
</tr>
<tr>
<td>S2</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td>S3</td>
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</tr>
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<td>S4</td>
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<td>S7</td>
<td>2</td>
<td>12</td>
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<tr>
<td>S8</td>
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<td>11</td>
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<tr>
<td>S9</td>
<td>2</td>
<td>10</td>
</tr>
<tr>
<td>S10</td>
<td>2</td>
<td>10</td>
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</table>

S1 substrate comprises primers 2 and template 8; S2 substrate is primer 2; S3 substrate comprises primers 2 and template 8; S4 substrate consists of primers 2, 3 and template 7; S5 substrate contains primers 2, 4 and template 7; S6 substrate includes primers 2, 5 and template 9; S7 substrate consists of primer 2 and template 12; S8 substrate is composed of primer 2 and template 11; S9 substrate consists of primer 2 and template 10; S10 substrate comprises
TABLE 2  Kinetic parameters of archaeal RNase HII. Hydrolyses of RNA-DNA/DNA substrates (S1) were carried out at 60°C in PabRNase HII reaction buffer as described in the Materials and methods section. The data were fit by nonlinear regression using the Marquardt-Levenberg algorithm (EnzFitter 2.0, BioSoft) to the Michaelis-Menten equation. Kinetic parameters, $K_m$ and $V_{max}$, obtained from the fit were used to calculate the catalytic efficiency ($k_{cat}/K_m$) of PabRNase HII. The kinetics values are the average of at least triplicate determinations and are shown with standard deviations (SD). Kinetic parameters of AfuRNase HII were extracted from previous studies (4).
Fig. 1
Fig. 3
Fig. 4
**Fig. 5**

**A**

[Diagram of RNA secondary structures labeled S7, S8, and S9]

**B**

[Image of gel electrophoresis showing substrate hydrolysis percentages for S7, S8, and S9 across different concentrations (nM)]

<table>
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<tr>
<th>Lanes</th>
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<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
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<th>8</th>
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<th>10</th>
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<tr>
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<td>0</td>
<td>12.8</td>
<td>45.2</td>
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**C**

[Diagram of RNA sequences for S7, S8, and S9]

5'-auucguaucauGGTCAGCTGTTTCCGT- 3'  
5'-auucguaucauGGTCAGCTGTTTCCGT- 3'  
5'-auucguaucauGGTCAGCTGTTTCCGT- 3'
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<td>32-nt</td>
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<td>2</td>
<td>30-nt</td>
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<td>17-nt</td>
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<td>5</td>
<td>12-nt</td>
<td>TGGGTGGGGTGG</td>
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<td>6</td>
<td>30-nt</td>
<td>ATTCGTAATCAuGGTCATAGCTGTTCCTG</td>
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<tr>
<td>Templates</td>
<td>Length</td>
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<td>30-nt</td>
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<td>10</td>
<td>30-nt</td>
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<td>11</td>
<td>30-nt</td>
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<tr>
<td>12</td>
<td>30-nt</td>
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<td>Enzymes</td>
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<td>$k_{cat}$ (min$^{-1}$)</td>
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<tr>
<td><em>Pab</em>RNase HII</td>
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<td><em>Afu</em>RNase HII</td>
<td>0.06±0.15</td>
<td>8.0±0.23</td>
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Table 2