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1 **INTRINSIC PROPERTIES OF THE TWO REPLICATIVE DNA POLYMERASES OF**
2 **PYROCOCCUS ABYSSI IN REPLICATING ABASIC SITES: POSSIBLE ROLE IN DNA**
3 **DAMAGE TOLERANCE ?**

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16

17 Running Tittle : DNA synthesis by *Pabpols* of DNA containing abasic sites

18

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25 **SUMMARY**

26 Spontaneous and induced abasic sites in hyperthermophiles DNA have long been
27 suspected to occur at high frequency. Here, *P.abysyi* was used as an attractive model to analyse
28 the impact of such lesions onto the maintenance of genome integrity. We demonstrated that
29 endogenous AP sites persist at a slightly higher level in *P.abysyi* genome compared to *E.coli*.
30 Then, the two replicative DNA polymerases, *PabpolB* and *PabpolD*, were characterized in
31 presence of DNA containing abasic sites. Both *Pabpols* had abortive DNA synthesis upon
32 encountering AP sites. Under running start conditions, *PabpolB* could incorporate in front of the
33 damage and even replicate to the full-length oligonucleotides containing a specific AP site , but
34 only when present at a molar excess. Conversely, bypassing activity of *PabpolD* was strictly
35 inhibited. The tight regulation of nucleotide incorporation opposite the AP site was assigned to
36 the efficiency of the proofreading function, because exonuclease-deficient enzymes exhibited
37 effective TLS. Steady-state kinetics reinforced that *Pabpols* are high-fidelity DNA polymerases
38 onto undamaged DNA. Moreover, *Pabpols* preferentially inserted dAMP opposite an AP site,
39 albeit inefficiently. While the template sequence of the oligonucleotides did not influence the
40 nucleotide insertion, the DNA topology could impact on the progression of *Pabpols*. Our results
41 are interpreted in terms of DNA damage tolerance.

42 INTRODUCTION

43 The genome of a living cell continuously undergoes a plethora of both exogenous or
44 endogenous genotoxic attacks. Among the myriad of DNA lesions, the abasic
45 [apurinic/apyrimidinic (AP)] sites are one of the most common lesions arising at high steady-
46 state levels, yielding up to 2,000-10,000 lesions per human cell per day by spontaneous
47 hydrolysis of the N-glycosylic bond (Lindahl and Nyberg, 1972; Lindahl, 1993). These lesions
48 can be generated by direct elimination of bases via free radical attacks, as a consequence of cells
49 exposure to chemical and physical agents (Breen and Murphy, 1995; Cadet *et al.*, 1999; Loeb *et*
50 *al.*, 1986). Furthermore, AP sites appear transiently as intermediates of Base Excision Repair
51 (BER) by DNA N-glycosylases (Loeb *et al.*, 1986; Scharer and Jiricny, 2001). Despite the fact
52 that it could be considered as an attractive model, identification and determination of the
53 mutagenic properties of AP sites in hyperthermophilic archaea (HA) remains poorly understood.
54 Presumably, life at high temperature inflicts additional stress to genomic DNA in each cell and
55 very high rates of potentially mutagenic DNA lesions (deamination, depurination, oxidation by
56 hydrolytic mechanisms and subsequent strand breakage) should be expected. However, and
57 interestingly, it was demonstrated that the hyperthermophilic crenarchaeon *Sulfolobus*
58 *Acidocaldarius* exhibits a modest rate of spontaneous mutations nearly close to that of
59 *Escherichia coli* (*E. coli*), raising the question of how HA do to preserve their genome intact in
60 such deleterious environmental conditions (Grogan *et al.*, 2001; Jacobs and Grogan, 1997).
61 To cope with the huge spectrum of impediments that result in genome destabilizing lesions,
62 multiple DNA repair mechanisms have evolved in all organisms to ensure genomic stability
63 (Friedberg *et al.*, 2006; Grogan, 2004; Hoeijmakers, 2001). However, situations can arise in
64 which DNA damage escapes to DNA repair and persists into the genome. Cells have developed

65 DNA damage tolerance mechanisms to tolerate hurdles in DNA either by post-replicative gap
66 filling, copy-choice DNA synthesis or translesion DNA synthesis (TLS) (Friedberg, 2005;
67 Friedberg *et al.*, 2006). Both bacteria and eukaryotes can tolerate arrested DNA replication by
68 template switching, therefore avoiding accumulation of mutations (Courcelle *et al.*, 2003;
69 McGlynn and Lloyd, 2002). Whereas template switching systems remain unknown in archaea,
70 TLS appears to be conserved within the three kingdoms of life (Boudsocq *et al.*, 2001; Friedberg
71 *et al.*, 2000; Hubscher *et al.*, 2002; Nohmi, 2006; Pages and Fuchs, 2002; Shimizu *et al.*, 2003;
72 Yang and Woodgate, 2007). Kinetically, this damage tolerance mechanism can be divided in two
73 steps: (i) nucleotide insertion opposite the DNA lesion; (ii) extension beyond the lesion.
74 Depending on the nature of the lesion, the bypass may involve a single or the concerted action of
75 DNA polymerases (Friedberg, 2005; Friedberg *et al.*, 2005). High-fidelity replicative DNA
76 polymerases in crenarchaea, bacteria and eukaryotes are intrinsically severely blocked upon
77 incorporation opposite a lesion such as an abasic site, thus recognizing the illegitimate formed
78 base pair and entering into futile cycles of insertion/excision (Gruz *et al.*, 2003; Pages *et al.*,
79 2005; Tanguy Le Gac *et al.*, 2004; Zhao *et al.*, 2004). This phenomenon called 'idling' is
80 relevant to replicative DNA polymerases harbouring the proofreading 3'-5' exonuclease and
81 reflects the partitioning of a mispaired DNA template between the exonuclease/polymerase
82 active sites (Villani *et al.*, 1978). The exonuclease activity acts as a kinetic barrier to TLS by
83 preventing the stable incorporation of bases opposite the DNA lesion and, therefore, confers the
84 exquisite accuracy of replicative DNA polymerases to preserve the integrity of the genome
85 (Khare and Eckert, 2002). In the absence of coding information due to the base loss, most of
86 replicative DNA polymerases obey to the A-rule, preferentially incorporating a dAMP opposite
87 the abasic site (Haracska *et al.*, 2001; Lawrence *et al.*, 1990; Shibutani *et al.*, 1997).

88 Conceivably, the DNA sequence context, the structure of the DNA primer lesion and the
89 replicative DNA polymerase examined can account for the preferential dAMP insertion opposite
90 an abasic site. Currently, the A-rule for replicative DNA polymerases remains under intensive
91 debates (Hogg *et al.*, 2004; Kroeger *et al.*, 2006; Taylor, 2002).

92 Here, we used *Pyrococcus abyssi* (*P. abyssi*), as a model for studying the genomic maintenance
93 at high temperature. This euryarchaeote grows at an optimum of 95°C and is faced to
94 environmental fluctuations imposed by hydrothermal vents (Erauso *et al.*, 1993; Jolivet *et al.*,
95 2003). Interestingly, *P. abyssi* is able to duplicate bidirectionally its 1.7 million base-pairs from a
96 single origin as fast as 45 minutes (Myllykallio *et al.*, 2000) and DNA replication is thought to
97 be achieved by the two high-fidelity DNA polymerases (*PabpolD* and *PabpolB*) and their
98 accessory factors (Henneke *et al.*, 2005; Rouillon *et al.*, 2007). Consistent with the existing
99 translesional systems and the lack of specialised DNA polymerases in *P. abyssi*, we speculate
100 that one or both *Pabpols* could be involved in damage tolerance. In this study, we determine the
101 steady-state level of AP sites in *E. coli* and *P. abyssi* at different growth stages. Secondly, we
102 examine the bypass properties of the exonuclease proficient and deficient replicative *Pabpols*
103 across an abasic site by varying the DNA topology and sequence context. Finally, steady-state
104 kinetic was employed to give substantial insights into the role of the proofreading activity of
105 *Pabpols* for nucleotide incorporation on damaged in comparison with intact DNA templates.
106 Potential mutagenicity of abasic sites and more generally genomic maintenance in *P. abyssi* are
107 discussed.

108 **RESULTS**

109 **Rate of endogenous AP sites in *P. abyssi* and *E. coli***

110 Before dissecting the *in vitro* behaviour of the *Pabpols* in the presence of abasic sites, we
111 investigated whether such DNA lesions were present into the genome of *P. abyssi*. The
112 mesophilic bacteria *E. coli* was used as a control. The steady-state level of abasic sites was
113 evaluated during the exponential and stationary phases of growth (Figure 1). In the exponential
114 phase, 2 and 25 abasic sites per 100,000 bp were calculated for *E. coli* and *P. abyssi*, respectively
115 (Figure 1). This value moderately increased to reach the number of 4 and 42 abasic sites per
116 100,000 bp at the stationary phase, for *E. coli* and *P. abyssi*, respectively. Taken together, these
117 data provided evidence for the first time that the genome of the hyperthermophile *P. abyssi* has
118 to deal with the presence of abasic sites. Further, the level of AP sites in *P. abyssi* genome is
119 approximately 10-fold higher than in *E. coli*.

120

121 **Replication of AP sites containing M13mp18 DNA template by *Pabpols***

122 We first checked the capacity of *Pabpols* to duplicate a circular AP-containing
123 heteropolymeric M13mp18 DNA template. Preparation of this damaged AP-M13mp18 templates
124 is depicted in Figure 2A. Under the conditions employed, 11 apurinic (the predominant lesion)
125 and apyrimidinic sites are introduced per molecules (Schaaper and Loeb, 1981). *Pabpols* were
126 tested in a primer extension assay in the presence of either abasic or undamaged M13mp18
127 templates. DNA elongation of the 5'-end fluorescein labelled oligonucleotide 6 (Table 1) was
128 visualized by product analysis on alkaline agarose gel. In the presence of undamaged DNA
129 template, both *Pabpols* (wild-type or exonuclease-deficient, respectively, *exo+* and *exo-*) could
130 extend the primer but with distinct efficiencies. While *PabpolB* *exo+/exo-* carried out DNA

131 synthesis to the full-length of the unmodified M13mp18 (7,249-nt), *PabpolD* *exo*+/*exo*- did only
132 extend the primer to 3,600-nt likely due to its sensitivity to secondary structures as already
133 observed (Henneke *et al.*, 2005) (Figure 2B, compare lanes 2 and 4 to 7 and 9). However, when
134 *PabpolD* *exo*- elongated the undamaged template, a faint band corresponding to the full-length
135 product was observed (Figure 2B, lane 9), consistent with the lower sensitivity of the *PabpolD*
136 *exo*- to secondary structures. DNA synthesis reactions of the wild-type *Pabpols* with AP
137 templates gave patterns similar to those obtained with undamaged templates, but with a lower
138 amount of elongated products (Figure 2B, lanes 3 and 8). Therefore, the presence of abasic sites
139 has an inhibitory effect on *Pabpols* activities. The results obtained with *Pabpols* were compared
140 to those of T4 DNA polymerase, used as a control. While the relative distribution of the products
141 of DNA replication was different with damaged versus undamaged M13mp18 DNA template,
142 the lower efficiency of the T4 DNA polymerase (family B) in the presence of DNA lesions was
143 confirmed (Figure 2B, lanes 12 and 13), as already described (Blanca *et al.*, 2007; Tanguy Le
144 Gac *et al.*, 2004). It should be noted that, when reactions were carried out in the presence of
145 *Pabpols* *exo*- with damaged DNA, a higher amount of replicated DNA products appeared
146 (Figure 2B, compare lanes 3 to 5 for *PabpolB* and lanes 8 to 10 for *PabpolD*) indicating that the
147 proofreading activity of *Pabpols* acts as a kinetic barrier to translesion synthesis onto damaged
148 M13mp18. To further address the inhibitory effect of AP sites onto the DNA polymerising
149 activity of *Pabpols*, quantitative analyses were performed by acid precipitable assay as described
150 under Experimental procedures. Figure 2 C shows the results of replicating these DNA templates
151 by *Pabpols*. Both *Pabpols* could discriminate between damaged and intact M13mp18 DNA
152 template. Indeed, reduced synthetic rates were observed for *Pabpols* independently of the
153 proficient or deficient exonuclease activity, with damaged DNA templates. The presence of AP

154 sites caused a more pronounced inhibition of replication by *PabpolD*, showing a 183-fold and
155 55-fold reduction of synthetic rates, respectively, for *PabpolD* *exo*⁺ and *PabpolD* *exo*⁻. In the
156 case of *PabpolB*, a 6-fold and 5-fold reduction of synthetic rates, respectively, for *PabpolB* *exo*⁺
157 and *PabpolB* *exo*⁻, were caused by AP sites. T4 DNA polymerase (family B), used as a control,
158 exhibited a reduced replicating activity comparable to that of *PabpolB*. Taken together, these
159 results argued that *Pabpols* discriminate between undamaged or damaged DNA templates, with
160 distinct reduced polymerising activities, suggesting that the presence of AP sites is rate-limiting.

161

162 **Replication of AP-containing mini-circular and linear oligonucleotides DNA templates by**
163 **the wild-type or exonuclease-deficient *PabpolB***

164 Since we provided evidence that DNA synthesis was severely impaired on damaged
165 M13mp18 for both *Pabpols*, we further investigated the insights of such reduced activities.
166 Translesional synthesis of a unique AP site from wild-type or exonuclease-deficient *Pabpols*
167 were examined under running start conditions using either linear or mini-circular
168 oligonucleotides DNA templates. The presence of the AP site was controlled by using T4 DNA
169 polymerase as already described (Blanca *et al.*, 2007). While *PabpolB* *exo*⁺ could bypass the
170 abasic site with moderate efficiency (63%) at a ratio enzyme / DNA of 4:1, under these
171 conditions *PabpolB* *exo*⁻ displayed almost full translesion synthesis (93%) (Figure 3A, compare
172 lanes 7 and lanes 14, respectively for *PabpolB* *exo*⁺ and *PabpolB* *exo*⁻). Moreover, *PabpolB*
173 *exo*⁺ had an increased capacity to stall at the AP site, as showed by the more marked presence of
174 pausing sites at the lesion (Figure 3A, lanes 3-7). The efficiency of the bypass was dependent on
175 the amount of the DNA polymerase used and the presence or absence of the 3'- to 5'
176 exonuclease but, in all cases, a plateau could be reached near the equimolar enzyme / DNA

177 concentrations (Figure 3A, lanes 4-5 and lanes 11-12, respectively for *PabpolB* *exo*⁺ and
178 *PabpolB* *exo*⁻). The tolerance of the abasic site was reproducible and confirmed by using a linear
179 73-mer with a different DNA sequence context (data not shown) as already described (Tanguy
180 Le Gac *et al.*, 2004). When replicating the circular DNA template, the translesion ability of
181 *PabpolB* *exo*⁺/*exo*⁻ was reduced at all enzyme concentrations tested. Interestingly, even at the
182 highest concentrations of *PabpolB* over the minicircle template, translesion synthesis across the
183 AP site for *PabpolB* *exo*⁺ only reached 32% at its peak (Figure 3B, lane 7) whereas it was much
184 more effective, reaching up to 86% for *PabpolB* *exo*⁻ (Figure 3B, lane 13). In addition, under
185 these saturating conditions, there was a stimulation of strand displacement activity of the two
186 *PabpolB* versions, as indicated by the proportion of products longer than the 87-mer (Figure 3B,
187 lane 6-7 and 12-13). Taken together these results indicate that *PabpolB* has the capacity to
188 replicate past an AP site at high pol/DNA ratio. The proofreading activity of *PabpolB* influences
189 its translesion capacity and progression of both exonuclease-deficient and proficient *PabpolB* are
190 reduced in the presence of the mini-circular DNA template.

191

192 **Replication of AP-containing mini-circular and linear oligonucleotides DNA template by** 193 **the wild-type or exonuclease-deficient *PabpolD***

194 The ability of *PabpolD* *exo*⁺/*exo*⁻ to bypass an abasic site onto the linear and circular
195 oligonucleotides DNA templates was analysed and the results are presented in Figure 4. As it can
196 be seen, *PabpolD* *exo*⁺ could incorporate in front of the AP site but, contrary to *PabpolB*, could
197 not extend past the lesion both on linear and circular DNA templates at all the enzyme's
198 concentrations tested (lanes 3-7 in Figure 4A, and in Figure 4B). In addition, accumulation of a
199 shorter product at position +32 could be detected, indicating the 'idling' activity of the *PabpolD*

200 *PabpolD* *exo-* was also blocked at the AP site in the presence of the linear template but
201 longer products past the AP site at position + 34, + 35, and + 36 could be detected, although they
202 never reached the full-length of the 87-mer, even at saturating enzyme concentrations (Figure
203 4A, lanes 9-13). Interestingly, 35% of bypass could be measured at a ratio *PabpolD* *exo-* / DNA
204 of 4:1 on the linear DNA template. Similarly, when the experiments were repeated in the
205 presence of the minicircle DNA template, a strong block at the AP site (+ 33) could be observed
206 for *PabpolD* *exo-* present at lower concentration to the DNA template (Figure 4B, lanes 9-10).
207 Moreover, DNA synthesis continued past the AP site at position + 34, + 35, and + 36, but was
208 also able to reach up to the full-length 87-mer (Figure 4B, lanes 11-13), when the concentration
209 of the enzyme was higher than that of the minicircle DNA template. Therefore, in the case of
210 *PabpolD*, its exonuclease activity prevents translesion synthesis of an abasic site independently
211 of the structure of DNA template, while the *exo-* mutant shows some bypass capacity that
212 seems enhanced in the case of a mini-circular DNA.

213

214 **Steady-state kinetic analysis of nucleotide incorporation of wild-type and exonuclease-** 215 **deficient *Pabpols* opposite undamaged bases**

216 *Pabpols* have been designated as replicative DNA polymerases (Henneke *et al.*, 2005).
217 This designation supposes that *Pabpols* must endow with high selectivity for each incoming
218 nucleotide depending on the nature of the base-containing template. Nucleotide incorporation
219 kinetics were measured in standing start reactions as described in Experimental procedures.
220 Incorporation efficiency (k_{cat}/K_m) were measured for the wild-type and exonuclease-deficient
221 *Pabpols* and the frequency of nucleotide misinsertion was calculated as the ratio of the efficiency
222 (k_{cat}/K_m) of incorrect nucleotide incorporation to the efficiency (k_{cat}/K_m) of correct nucleotide

223 incorporation (Table 2). Both wild-type *Pabpols* exclusively incorporated the correct dGMP
224 opposite template C and no misincorporation could be detected. In these conditions, a 5-fold
225 reduced incorporation efficiency for *PabpolD* compared to *PabpolB* was observed as indicated
226 by the k_{cat}/K_m values. The results obtained with the exonuclease-deficient *Pabpols* at template C
227 showed that the correct dGMP was preferentially incorporated but their efficiencies were
228 dramatically reduced compared to the wild-type enzymes, as judged from the dropped k_{cat}/K_m
229 values, 55- and 103-fold, respectively, for *PabpolB* and *PabpolD*. In addition, misinsertion
230 events by the exonuclease-deficient *Pabpols* at template C were comparable, with preference for
231 insertion of dTMP > dAMP > dCMP (Table 2). Taken together, these results demonstrate that,
232 while wild-type *Pabpols* monitor the instructional base of the template and discriminate between
233 correct and incorrect nucleotides insertion, exonuclease-deficient *Pabpols* are much less
234 efficient.

235

236 **Steady-state kinetics analysis of nucleotide incorporation of wild-type and exonuclease-** 237 **deficient *Pabpols* opposite an AP site**

238 The efficiency for deoxynucleotide insertion opposite an abasic site for the wild-type
239 *PabpolB* followed the order dAMP > dTMP and to a lesser extent dGMP. Interestingly, *PabpolB*
240 *exo +* incorporated a dGMP 8355-fold less efficiently opposite an AP site than opposite the
241 template C (Table 2). Similarly, as judged by the k_{cat}/K_m values, *PabpolD* *exo +* incorporated
242 preferentially a dAMP over a dTMP opposite the abasic site with a 20-fold higher magnitude
243 efficiency. It is noteworthy that the apparent K_m values from the incorporation of nucleotides
244 opposite the abasic site were always higher than from the insertion of a correct dGMP at
245 template C. Thus, proficient proofreading *Pabpols* are sensitive to abasic sites and are not

246 efficient at inserting a nucleotide opposite such a non-instructive lesion. As observed from the
247 k_{cat}/K_m values, *Pabpol*s *exo* - inserted a dAMP more efficiently than other deoxynucleotides
248 opposite the AP site. While incorporation efficiencies of dAMP, dGMP and dTMP were higher
249 when the proofreading function of *Pabpol*B was deficient, no striking difference was observed
250 between the wild-type and exonuclease-deficient *Pabpol*D. Therefore, *Pabpol*B and *Pabpol*D
251 exonuclease-deficient resemble each other in their ability to insert nucleotides opposite an abasic
252 site with higher efficiencies for dAMP incorporation. Taken together, the data show that, while
253 wild-type *Pabpol*B does not significantly discriminate among nucleotides for incorporation
254 opposite an AP site, the exonuclease-deficient *Pabpol*B and both wild-type and exonuclease-
255 deficient *Pabpol*D were much sensitive to a non-coding lesion that seems to govern the dAMP
256 incorporation rather than other dNTPs.

257

258 **Effect of varying the downstream template base on nucleotide incorporation opposite an**
259 **AP site by the *Pabpol*s**

260 The primer-templates used in these experiments were designed in order to examine the
261 effect of changing the template base (N) at the 5' side of the AP site (X) on the initiation of the
262 extension of the primer. Nucleotide incorporation was measured in standing start reactions under
263 standard *Pabpol*s assay conditions. The 32-mer primer (oligonucleotide 6) was annealed to the
264 four templates differing by the nature of the base N = A, T, G and C at the 5' side of the AP site
265 (Oligonucleotides 2, 1, 3, 4, respectively) (Figure 5). Extension of the primer in the four
266 duplexes was conducted in the presence of each of the natural dNTPs. On the one hand, the wild-
267 type *Pabpol*s exclusively inserted a dAMP opposite the abasic lesion, independently of the
268 sequence context (Figures 5A and 5B). While the percentages of incorporation of a dAMP

269 ranged from 46 to 58% for *PabpolD* *exo+*, they only reached 18% for *PabpolB* *exo+*. The
270 *Pabpols* *exo-* also preferentially incorporated a dAMP opposite the AP site with a higher order of
271 magnitude compared to the *Pabpols* *exo+*. For example, the percentages of incorporation of a
272 dAMP ranged from 66 to 72% for *PabpolD* *exo-* and 46 to 58% for *PabpolD* *exo+* (Figure 5D
273 and Figure 5B, respectively). However, unlike *Pabpols* *exo+*, the *Pabpols* *exo-* also inserted
274 other dNMPs opposite the non instructional lesion, following the order:
275 dAMP>dGMP>dTMP>dCMP for *PabpolB* *exo-* and dAMP>dTMP~dCMP>dGMP for *PabpolD*
276 *exo-*, independently of the sequence context (Figures 5C and 5D, respectively). Thus, these
277 results demonstrate that nucleotide incorporation opposite an AP site by the *Pabpols* *exo+/exo-*
278 is not directed by the nature of the base located at the 5' side of the AP site.

279 **DISCUSSION**

280 In hyperthermophiles, cellular and environmental stresses encountered by cells are likely
281 to be exacerbated by adaptation to extreme conditions. Cell survival is ensured by a complex
282 network of DNA events that contributes to the maintenance of the functional integrity of nucleic
283 acids at high temperature. Here, we have focused the study on *P. abyssi*, an anaerobe
284 hyperthermophilic euryarchaeota (HA) that thrives at elevated temperature in the immediate
285 surroundings of deep-sea hydrothermal vents. This is the first report that establishes a
286 relationship between the specific genomic level of abasic sites and the resulting impact on the
287 intrinsic properties of replicative DNA polymerases in archaea. These features show that *P.*
288 *abyssi* can be used as an informative model to analyse the biological relevance of DNA damage
289 accumulation in the hyperthermophilic chromosome and the underlying genomic maintenance
290 mechanisms.

291 The investigation presented here shows that the steady-state level of AP sites into the 1.7
292 million base-pairs of *P. abyssi* ranged from 25 to 42 AP sites per 100,000 bp during the
293 exponential and stationary phases, respectively. Similarly, a 2-fold increase of AP sites is
294 observed in *E. coli*. These findings corroborate the higher efficiency of DNA protection and
295 repair mechanisms in proliferating cells, compatible with a low level of AP sites. However, it is
296 important to precise that our expectation to detect a drastic increased number of abasic sites in
297 the hyperthermophilic chromosome was not fulfilled. Indeed, the level was only 10-fold higher
298 than in the mesophilic bacteria *E. coli*. Comparatively, the frequency of endogenous AP sites in
299 mammalian cells reached 10 to 12 AP sites per 100,000 bp (Zhao *et al.*, 2006). Indeed, in the
300 literature, the number of AP sites in the genome of thermophilic and hyperthermophilic
301 microorganisms is always suspected to increase drastically compared to the mesophilic

302 counterparts (Grogan, 1998; Grogan, 2000). This assumption takes into account of the intrinsic
303 properties of the primary structure of DNA at elevated temperatures, corresponding to a 3,000-
304 fold increase in DNA decay at 100°C (Lindahl, 1993). Conceivably, *P. abyssi* has evolved to
305 adjust genetically the level of endogenous AP sites in its genome that could be detrimental for
306 genome integrity of mesophiles. Accumulation of AP sites and, more generally, others DNA
307 lesions can be envisaged, suggesting that hyperthermophiles are adapted to survive pre-existing
308 mutations. Clearly, these results establish that the number of AP sites seems to evolve respect to
309 the growth stage without affecting cell growth and viability. However, further studies are
310 required to determine the threshold that hyperthermophiles can support to ensure cell survival.

311 Evolution has produced multiple DNA polymerases able to replicate undamaged or
312 damaged DNA. Sixteen DNA polymerases have been described in human, nine in
313 *Saccharomyces cerevisiae*, five in *E. coli* (Bebenek and Kunkel, 2004; Hubscher *et al.*, 2002;
314 Rothwell and Waksman, 2005) and up to five in archaea (Barry and Bell, 2006; Yang, 2005).
315 Interestingly, the genome of the euryarchaeota *P. abyssi* encodes only two DNA polymerases,
316 families B and D, required to faithfully duplicate the genetic information (Henneke *et al.*, 2005).
317 In this study, steady-state kinetic analyses of nucleotide insertion show for the first time that
318 *PabpolD* is endowed with high-fidelity onto undamaged DNA as its replicative counterpart,
319 *PabpolB* (Table 2). Incorporation efficiency of *PabpolD* was reduced to five fold compared to
320 *PabpolB*, demonstrating that *Pabpols* possesses distinct kinetic properties.

321 Our results give evidence that the presence of AP sites strongly inhibited the DNA polymerising
322 activity of both *Pabpols* and that the absence of their proofreading function correlate with
323 enhanced bypass of AP sites. The degree of inhibition of DNA synthesis was dependent upon the
324 *Pabpol* examined. *PabpolD* could insert a nucleotide opposite the AP site and, in all conditions

325 tested, was not able to extend beyond the 3' primer lesion. *PabpolB* inserted opposite the AP site
326 and extended the DNA template to the full-length, only when present at a molar excess over the
327 template. This result is comparable to previous studies showing that molar excess of enzyme
328 versus DNA template could enhance translesion synthesis by DNA polymerases past an abasic
329 site (Blanca *et al.*, 2007; McCulloch and Kunkel, 2006; Tanguy Le Gac *et al.*, 2004). While
330 translesion synthesis of both exonuclease-proficient and deficient *PabpolB*s and the exonuclease-
331 deficient *PabpolD* were differently affected by the topology of the DNA template, the template
332 sequence context did not significantly influence the bypass properties of *Pabpols*. Further, both
333 *Pabpols* inserted dAMP opposite the AP site independently of the nature of the 5' template base
334 to the AP site. Despite this nucleotide selectivity, steady-state kinetics showed that dAMP
335 incorporation was not efficient. These observations show for the first time that archaeal
336 replicative (families B and D) DNA polymerases follow the 'A-rule' (Strauss, 1991; Taylor,
337 2002) like eukaryal and bacterial counterparts (Haracska *et al.*, 2001; Shibutani *et al.*, 1997). The
338 physical basis of the 'A-rule' is still an intensive debate (Kool, 2002; Zahn *et al.*, 2007) and
339 structural studies reported the molecular level of replication blockage that produced catalytically
340 inactive DNA polymerases (Freisinger *et al.*, 2004; Hogg *et al.*, 2004). Whether the molecular
341 and physical bases are conserved through archaeal replicative DNA polymerases would have to
342 be unravelled. Accordingly, the ability of abasic sites to inhibit *Pabpols* could reflect steric
343 constraints imposed by the "tightness" of the active site. Furthermore, the capacity to partition
344 the mismatches away from the polymerase domain into the exonuclease active site might exhibit
345 structural rearrangements that are differentially influenced by the dynamic features of the DNA
346 polymerase. To this point, the major distinction between the two *Pabpols* is the subunits
347 composition. While *PabpolB* is a monomeric enzyme with associated exonuclease and

348 polymerase activities, *PabpolD* is an heterodimeric enzyme with the large and the small subunits
349 carrying, respectively, the polymerase and the exonuclease activities (Gueguen *et al.*, 2001).
350 Therefore, it is reasonable to suggest that the architecture of the two DNA polymerases may
351 account for the subtle differences observed within the polymerase and exonuclease efficiencies.
352 However, a complete detailed functional analysis must await the crystal structure of the
353 individual *Pabpols*. The balance between polymerization and excision was recently described in
354 B-family DNA polymerase in archaea (Kuroita *et al.*, 2005) but never in D-family. The distinct
355 kinetic partitioning of insertion and edition of mispairs observed within *Pabpols* corroborates
356 with eukaryal and bacterial homologues properties (Jin *et al.*, 2003; Jin *et al.*, 2005; Pages *et al.*,
357 2005) and confers that replicative DNA polymerases are high-fidelity enzymes (Bloom *et al.*,
358 1997; Chen *et al.*, 2000; Shimizu *et al.*, 2002).

359 The down-regulation of the proofreading function of *PabpolB* could favour TLS in order to
360 overcome the block imposed by AP sites. Bypass of AP sites could generate either single-base
361 substitutions or frameshift mutations (Baynton and Fuchs, 2000). *PabpolB* appears to proceed
362 through single-base substitution upon completion of DNA template containing an AP site. The
363 molar excess of the enzyme over the DNA template accounted for TLS under our *in vitro*
364 conditions. On the other hand, a replicative *Pabpol* idling at a DNA lesion could be a crucial
365 factor to trigger cellular responses to DNA damage in *P. abyssi*. The mechanism by which
366 proofreading activities of archaeal DNA polymerases could be regulated *in vivo* (dNTPs balance,
367 role of accessory proteins and enzymes switching) and their contribution in some cases to
368 counteract genomic DNA lesions has to be elucidated. Recently, it was described that *P. abyssi*
369 has evolved with efficient DNA strategies to cope with ionizing radiations and elevated
370 temperatures (Jolivet *et al.*, 2003). Biochemical evidence for relevant DNA repair mechanisms

371 has not been demonstrated in *P. abyssi* yet. More striking is the lack of identification of any
372 mismatch repair genes and the complete set of damage excision genes (Cohen *et al.*, 2003).
373 Interestingly, homologous recombination genes (RecA/Rad51) have been identified into the
374 genome sequence of *P. abyssi* together with the fact that exponentially growing thermophilic
375 archaea contain several copies of the chromosome (Bernander and Poplawski, 1997; Breuert *et*
376 *al.*, 2006). This might be particularly consistent for the repair of strand breaks. Furthermore, it is
377 not excluded that the recently characterised primase from *P. abyssi* could also play a role in
378 damage avoidance since it possesses sequence and structural similarities with the family X DNA
379 polymerases (Le Breton *et al.*, 2007). Ultimately, the process of such DNA lesions would have to
380 be explored by the *P. abyssi* replisome in the context of genomic mutagenicity.

381 **EXPERIMENTAL PROCEDURES**

382 **Strains and cell culture techniques**

383 *P. abyssi* GE5 (Brittany Culture Collection, <http://www.ifremer.fr/souchotheque>) were
384 grown in 50 ml YPS medium under anaerobic conditions at 95°C (Erauso *et al.*, 1993). The *E.*
385 *coli* CIP 54.8 strain (CRBIP) was cultivated in 1 L of Luria-Bertani (LB) broth (1% tryptone,
386 0.5% yeast extract, 1% NaCl) at 37°C and pH 7.3, in a shaking incubator (170 rpm). Growth was
387 monitored by density measurements with a cell Thoma counting chamber (0.02 mm depth).
388 Samples for DNA extraction were collected in the exponential and stationary growth phases as
389 indicated in Figure 1. The samples for DNA extraction were centrifuged at 6,000 g for 15 min at
390 4°C and the pellets were stored at –20°C.

391

392 **Genomic DNA Isolation and Detection of AP sites**

393 Genomic DNA from *P. abyssi* was isolated using the extraction method as described
394 (Charbonnier *et al.*, 1995) and optimised in order to avoid the formation of additional AP sites.
395 Briefly, cell pellets were suspended in 800 µl TE-Na-1 X lysis buffer (100 mM Tris-HCl, 50 mM
396 EDTA, 100 mM NaCl, pH 8.0). This was followed by successive additions of 50 µl proteinase K
397 (20 mg/ml), 100 µl Sarcosyl (10%), 100 µl SDS (10%). The applied lysis treatment were
398 performed at 37°C for 1.5 hours and isolation of the total DNA was accomplished by adding an
399 equal volume of buffered (pH 8.0) PCI (Phenol/Chloroform/Isoamyl Alcohol: 25/24/1). The
400 samples were gently mixed and the aqueous phases were collected by centrifugation at 10,000 g
401 for 10 min at 4°C. 10 µl RNase (10 mg/ml) were added and incubation was performed at 37°C
402 for one hour. DNAs were purified with an equal volume of PCI and centrifuged. The upper phase
403 was extracted with an equal volume of pure chloroform and centrifuged. DNA precipitation was

404 obtained by mixing the final aqueous phase with 0.7 volume of 100% isopropanol followed by
405 incubation for one hour at room temperature. After a 30 min centrifugation at 15,000 g at 4°C,
406 the DNA pellets were washed once with 0.5 ml of 70% ice cold ethanol. Finally, the DNA pellets
407 were air-dried during 1 hour before solubilization in TE-1X buffer (10 mM Tris-HCl, 2 mM
408 EDTA, pH 7.5).

409 Genomic DNA extractions from *E. coli* was performed following the CTAB (Hexadecyl
410 trimethyl-ammonium bromide) method for Gram-negative bacteria as described (Park, 2007).

411 The level of AP sites in genomic DNA was measured using the DNA damage
412 quantification –AP site Counting kit from Dojindo Molecular Technologies (Gaithersburg, MD).
413 DNA pellets were dissolved in TE buffer supplied by the kit and DNA concentrations were
414 exactly adjusted to 100 ng/μl. Briefly, DNA samples were incubated with the Aldehyde Reactive
415 Probe (ARP) reagent (N'-aminooxymethylcarbonylhydrazino-D-biotin) that specifically reacted
416 with the aldehydic ring-opened AP sites (Kow and Dare, 2000; Kubo *et al.*, 1992). The AP sites
417 tagged with biotin interacted with horseradish peroxidase-streptavidin and AP sites were
418 colorimetrically detected. For each condition, the average of three measurements per sample was
419 used in the statistical analyses.

420

421 **Chemicals and Enzymes**

422 Unlabelled dNTPs were from MP Biomedicals. T4 polynucleotide kinase, DNA ligase
423 and T4 DNA polymerase were from New England Biolabs. *PabpolD* was cloned, expressed and
424 purified as described (Henneke *et al.*, 2005). *PabpolB* (Isis DNA polymerase) and *PabpolB*
425 exonuclease-deficient (Pyra DNA polymerase) were purchased from MP Biomedicals. 1 unit of
426 *Pabpols* corresponds to the incorporation of 1 nmol of total dTMP into acid-precipitable

427 material per min at 65°C in a standard assay containing 0.5 µg (nucleotides) of
428 poly(dA)/oligo(dT)_{10:1}. All other reagents were of analytical grade and purchased from Sigma-
429 Aldrich and Fluka.

430

431 **Construction, expression and purification of the recombinant wild-type and exonuclease-** 432 **deficient His tag *PabpolD***

433 The pET26b expression vector containing the *PabpolD* large subunit (DP2) (Gueguen *et*
434 *al.*, 2001) was digested with *NdeI* and *Sall* and the resulting fragment was inserted into the
435 pET28a expression vector (Novagen) in order to introduce a histidine tag (His tag) at the N-
436 terminus. To render the *PabpolD* exonuclease-deficient, site-directed mutagenesis was carried
437 out by introducing the H451A point mutation onto the *PabpolD* small subunit (DP1) (Gueguen *et*
438 *al.*, 2001). The two site-specific complementary primers, reverse H451A 5'-
439 TGGCCTAGCGGCATCGGCATTTCTGGCCCTAT-3' and forward H451A 5'-
440 ATAGGGCCAGGAAATGCCGATGCCGCTAGGCCA-3' were used to PCR amplify the
441 pARHS expression vector containing DP1 according to the protocol of the Quick change
442 Mutagenesis kit (Stratagene, La Jolla, CA). DNA sequencing was used to confirm that no
443 spurious mutations had been introduced during PCR. The constructed expression vectors
444 pET28a/DP2 and either the wild-type or the exonuclease-deficient pARHS/DP1 were co-
445 introduced into host *E. coli* HMS174 (DE3). The transformed cells were grown in 1.5 liters Luria
446 Bertani (LB) medium containing ampicillin (100 µg/ml) and kanamycin (30 µg/ml) at 37 °C.
447 When A_{600} reached 0.7, 1 mM isopropylthio- β -D-galactoside was added to induce expression of
448 active DNA polymerases. After being cultured 4 hours at 37 °C with gentle shaking (160 rpm),
449 the cells were harvested by centrifugation, resuspended in 25 ml buffer A (20 mM sodium

450 phosphate, pH 6.6, 1 mM DTT, 20 mM Imidazole) containing the protease inhibitor, disrupted
451 on ice by French press and then heat-treated at 80°C for 15 min. Denatured host proteins were
452 removed by centrifugation. The clarified supernatant was applied further onto Ni²⁺-HisTrap
453 column (5 ml of bed volume) pre-equilibrated with buffer A. Proteins were eluted with buffer B
454 (20 mM sodium phosphate, pH 6.6, 1 mM DTT, 500 mM Imidazole and active fractions were
455 pooled and dialysed against buffer C (20 mM sodium phosphate, pH 6.6, 1 mM DTT). The
456 dialysate was loaded onto a heparin column (5 ml of bed volume) pre-equilibrated with buffer D
457 (20 mM sodium phosphate, pH 6.6, 1 mM DTT, 0.15 M NaCl). The column was developed with
458 a linear gradient from buffer D to buffer E (20 mM sodium phosphate, pH 6.6, 1 mM DTT, 1 M
459 NaCl). Eluted protein showed over 98% purity. Pure His-*Pabp*oD (wild-type and exonuclease-
460 deficient) were dialyzed against storage buffer (25 mM Tris-HCl, pH 8.0, 1 mM DTT, and 50%
461 glycerol) and stored at -20 °C until use. We checked by acid precipitable assay, as described
462 (Henneke *et al.*, 2005), that the addition of the His tag at the N terminus of DP2 had no effect on
463 the DNA polymerization activities. Moreover, the 3'-5' exonuclease deficiency for the mutant
464 H451A was confirmed (data not shown). Sequence alignment of exonuclease domain in
465 representative euryarchaeal DNA polymerases highlighting residues critical for proofreading
466 function is shown in Figure S1.

467

468 **Nucleic Acid Substrates**

469 Single-stranded (ss) M13mp18 was purchased (Amersham Biosciences, GE Healthcare).
470 In order to create AP sites into natural DNA templates, the ssM13mp18 viral DNA was
471 incubated at a final concentration of 0.18 pmol/μl in 30 mM potassium chloride, 10 mM sodium
472 citrate, pH 3.0 at 70°C for 45 min (Schaaper *et al.*, 1983). These conditions introduced one AP

473 sites per molecule in 4 min, measured by survival (Schaaper and Loeb, 1981). After treatment,
474 the damaged M13mp18 was purified with the QIAquick® PCR Purification Kit from Qiagen
475 (Germany).

476 The sequences of the DNA primers / templates used in the present study are depicted in
477 Table 1. All oligonucleotides, including those containing a tetrahydrofuran moiety mimicking an
478 abasic site, were chemically synthesized and gel-purified (Eurogentec, Belgium). Primers were
479 labelled at their 5'-end by fluorescein with the 5' End Tag kit labelling system from Vector
480 Laboratories (California). Free fluorescein was removed through the Microspin G-25 column
481 (Amersham Biosciences, GE Healthcare) and the labelled primers were hybridised to the
482 respective templates at equimolar concentrations.

483 The minicircle template was prepared as described (Tanguy Le Gac *et al.*, 2004). Briefly,
484 the linear 5'-phosphorylated oligonucleotide 1 (intact or containing the tetrahydrofuran moiety)
485 was intramolecularly ligated under dilute conditions using a scaffold 40-mer oligonucleotide (5'-
486 ATATTCCTACCCTCCCGATCTATCCACCATACTACCCTCC- 3'). Minicircles were gel-
487 purified and their concentration was determined spectrophotometrically, followed by annealing
488 with their complementary 5'-fluorescein labelled primer at equimolar concentration.

489

490 **Primer extension onto intact or damaged primed-oligonucleotides**

491 “Standing start” and “Running start” assays were catalysed into a final volume (15 µl)
492 containing the following components: (i) for *PabpolD*: 8.3 nM of labelled primers / templates, 20
493 nM of *PabpolD* *exo*⁺/*exo*⁻ unless otherwise specified, 10 mM Tris-HCl (pH 9.0), 50 mM KCl,
494 10 mM MgCl₂ and 200 µM dNTPs; (ii) for *PabpolB*: 8.3 nM of labelled primers / templates, 13
495 nM of *PabpolB* *exo*⁺/*exo*⁻ unless otherwise mentioned, 50 mM Tris-HCl (pH 8.8), 50 mM KCl,

496 1 mM DTT, 2 mM MgCl₂ and 200 μM dNTPs. Reactions were performed at 55°C for 30
497 minutes and quenched by the addition of 15 μl of stop buffer (98% formamide, 10 mM EDTA).
498 Samples were heated at 95°C for 5 minutes. The reactions products were resolved on 15%
499 polyacrylamide, 7 M urea gels and visualized with a Mode Imager Typhoon 9400 (Amersham
500 Biosciences, GE Healthcare). Quantification of the results was performed using ImageQuant 5.2
501 software. The extent of the bypass reaction was calculated as the ratio of the intensity of the
502 bands downstream of the AP site to the intensity of the bands opposite the lesion.

503 Effect of sequence context on AP site bypass was analysed under standing start
504 conditions. The fluorescein-labelled primer (oligonucleotide 6) was annealed right before the
505 template AP site that is indicated by X. Different template bases 5' to the AP site are depicted by
506 N (N=A, oligonucleotide 2; N=T, oligonucleotide 1; N=G, oligonucleotide 3; N=C,
507 oligonucleotide 4) (Table 1). Bypass assays were performed as described above excepted that 16
508 nM of DNA templates were used when the template base 5' to the AP site was: N=T and N=C
509 for *PabpolB* exo-, N=T and N=G for *PabpolD* exo+, N=C for *PabpolB* exo+. Quantification of
510 nucleotides insertion opposite the AP site are calculated for wild-type and exonuclease-deficient
511 *Pabpols* in triplicate but only the more resolving gel was quantified.

512

513 **Steady-state Kinetic Analyses**

514 A 5'-fluorescein labelled primer, annealed to either a correct or damaged template, was
515 extended in the presence of increasing concentrations of a single dNTP. *Pabpols* concentrations
516 and reaction times were set so that maximal product formation was $\leq 20\%$ of the substrate
517 concentration. The linear primer-template (oligonucleotides 6 and 1) was extended with dNTP at
518 55°C in the presence of 6.6-33.3 nM enzyme for 1 or 5 min, depending on the proper utilization

519 efficiency and substrate utilisation. All reactions (15 μ l) were carried out at various dNTP
520 concentrations (in triplicate) and quenched with 2 volumes of a solution of 20 mM EDTA in 95%
521 formamide (v/v). Products were resolved using a 15% polyacrylamide (w/v) electrophoresis gel
522 containing 7 M urea and visualized using a Mode Imager Typhoon 9400. Bands were quantified
523 with ImageQuant 5.2 software (Amersham Biosciences, GE Healthcare). The observed rates of
524 deoxynucleotide incorporation as a function of dNTP concentration were firstly determined from
525 Lineweaver-Burk plots. The data were fit by nonlinear regression using the Marquardt-
526 Levenberg algorithm (EnzFitter 2.0, BioSoft) to the Michaelis-Menten equation describing a
527 hyperbola, $v = (V_{max} \times [dNTP]/K_m + [dNTP])$ as already described (Le Breton *et al.*, 2007).
528 Apparent K_m and V_{max} kinetic parameters were obtained from the fit and were used to calculate
529 the efficiency of deoxynucleotide incorporation (k_{cat}/K_m). The kinetics values are the average of
530 at least triplicate determinations and are shown with standard deviations (SD). ND, means that
531 no detectable incorporation was observed. Gel patterns and quantitation of single nucleotide
532 incorporation reactions are shown in Figure S2.

533

534 **Primer extension onto intact or damaged M13mp18 DNA template**

535 *Product analysis.* The oligonucleotide 6 was annealed to either the damaged AP-
536 M13mp18 or undamaged M13mp18 at a molar ratio 3:1. Standard *Pabpols* reactions (10 μ l) were
537 conducted into their respective buffer containing 200 μ M each of dNTPs, 7 nM of DNA template
538 and 2 pmol of *Pabpols*. Reactions were carried out at 60°C for 30 minutes. T4 DNA polymerase
539 reactions were performed at 37°C for 30 minutes into the 1 X T4 pol buffer (according to the
540 manufacturer's protocol) with 7 nM of DNA template, 100 μ M each of dNTPs and 2 pmol of T4
541 DNA polymerase. Reaction mixtures were stopped by the addition of 10 μ l of 30 mM EDTA and

542 the samples were heated to 100°C for 10 min. Reactions mixtures were subjected to a 0.8% (w/v)
543 denaturing alkaline agarose gel electrophoresis, and replication products were visualized with a
544 Mode Imager Typhoon 9400 (Amersham Biosciences, GE Healthcare). DNA ladders (Raoul
545 markers, MP Biomedicals) were run into the same gel and revealed separately.

546 *Acid precipitable assay.* The reaction buffers composition were identical to those
547 described in product analysis for *Pabpols* and T4 DNA pol. The final volume of 10 µl contained
548 200 µM of unlabeled dNTPs, 20 µM [³H]dTTP, 7 nM of DNA template (AP-M13mp18 or
549 undamaged M13mp18) and 2 pmol of enzyme to be tested. Reactions were carried out at 60°C
550 and 37°C, respectively, for *Pabpols* and T4 DNA pol for 30 minutes. DNA was precipitated with
551 10% trichloroacetic acid (TCA). Insoluble radioactive material was determined by scintillation
552 counting as described (Henneke *et al.*, 2005; Rouillon *et al.*, 2007).

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- 733
- 734

735 **FIGURE LEGENDS**

736 **Table 1. Damaged or intact oligonucleotides used in this study.** X represents the position of
737 the correct base, template C, or a tetrahydrofuran moiety designed to functionally mimic an
738 abasic site.

739
740 **Table 2. Incorporation kinetics by wild-type and exonuclease-deficient *Pabpols*.** Single
741 nucleotide insertion assays were performed as described in Experimental procedures. The
742 observed rates of deoxynucleotide incorporation as a function of dNTP concentration were firstly
743 determined from Lineweaver-Burk plots. The data were fit by nonlinear regression using the
744 Marquardt-Levenberg algorithm (EnzFitter 2.0, BioSoft) to the Michaelis-Menten equation
745 describing a hyperbola, $v = (V_{max} \times [dNTP]/K_m + [dNTP])$ as already described (Le Breton *et*
746 *al.*, 2007). Apparent K_m and V_{max} kinetic parameters were obtained from the fit and were used
747 to calculate the efficiency of deoxynucleotide incorporation (k_{cat}/K_m). The kinetics values are the
748 average of at least triplicate determinations and are shown with SD. The f (misinsertion
749 frequency) is the ratio k_{cat}/K_m for the incorrect nucleotide to k_{cat}/K_m for the correct nucleotide. ND
750 means no detectable incorporation observed.

751
752 **Figure 1. Rate of endogenous AP sites into *P. abyssi* and *E.coli* genomes at different growth**
753 **stages.** Steady-state level of AP sites per 100,000 bp was calculated during the exponential and
754 stationary phases of growth. The number of AP sites per 100,000 bp represents the mean of
755 triplicate experiments and error bars show the standard deviations of each measurement.

756

757 **Figure 2. Replication of AP sites containing M13mp18 DNA template by *Pabpols*.** **A,**
758 Chemical treatment to induced AP sites into M13mp18 DNA. **B,** Primer extension assays were
759 performed with 5'-fluorescein end labelled primer (oligonucleotide 6) hybridised to either the
760 damaged or undamaged M13mp18 DNA template, *Pabpols* and T4 DNA polymerase used as a
761 control experiment. The elongated products were separated on a 0.8 % (w/v) denaturing alkaline
762 agarose gel. Lanes 1, 2, 4, 6, 7, 9, 11, 12 are the undamaged extended products; lanes 3, 5, 8, 10,
763 13 are the damaged extended products. **C,** dNTPs incorporation into the damaged and
764 undamaged M13mp18 DNA primed-templates were tested by acid precipitation and incubation
765 was performed according to the dependent polymerase reactions with [³H]dTTP as the substrate
766 (as outlined in Experimental procedures).

767

768 **Figure 3. Replication of AP-containing mini-circular and linear oligonucleotides DNA**
769 **templates by the wild-type or exonuclease-deficient *PabpolB*.** Primer extension assays were
770 performed at the indicated *PabpolB* concentrations with 8.3 nM of primer-template
771 (oligonucleotides 1 and 5), 200 μ M dNTPs at 55°C for 30 min as described in the Experimental
772 procedures. Quantifications of the extended products from the AP site are mentioned below the
773 gels. The extent of the bypass reaction was calculated as the ratio of the intensity of the bands
774 downstream of the AP site to the intensity of the bands opposite the lesion. **A**, Replication onto
775 the AP site-containing linear template. The position of the abasic site is indicated by X. Lanes 2
776 and 9 correspond to the positive control with 8.3 nM of intact template (X=C). **B**, Replication of
777 the AP site-containing circular template. Lanes 2 and 8 correspond to the positive control with
778 8.3 nM of intact template (X=C). 32 mer indicates the position of the base preceding the AP site,
779 while 33 mer is the position of the AP site.

780

781 **Figure 4. Replication of AP-containing mini-circular and linear oligonucleotides DNA**
782 **templates by the wild-type or exonuclease-deficient *PabpolD*.** Primer extension assays were
783 performed at the indicated *PabpolD* concentrations with 8.3 nM of primer-template
784 (oligonucleotides 1 and 5), 200 μ M dNTPs at 55°C for 30 min as described in the Experimental
785 procedures. Quantification of the extended products from the AP site is mentioned below the
786 gels. The extent of the bypass reaction was calculated as the ratio of the intensity of the bands
787 downstream of the AP site to the intensity of the bands opposite the lesion. **A**, Replication onto
788 the AP site-containing linear template. The position of the abasic site is indicated by X. Lanes 2
789 and 8 correspond to the positive control with 8.3 nM of template (X=C). **B**, Replication of the
790 AP site-containing circular template. Lanes 2 and 8 correspond to the positive control with 8.3

791 nM of template (X=C). 32 mer indicates the position of the base preceding the AP site, while 33
792 mer is the position of the AP site.

793

794 **Figure 5. Effect of varying the downstream template base on nucleotide incorporation**

795 **opposite an AP site by *Pabpols*.** Standing start reactions were performed with four DNA

796 templates that varied by the nature of the 5' template base. The fluorescein-labelled primer

797 (oligonucleotide 6) was annealed right before the template AP site that is indicated by X.

798 Different template bases 5' to the AP site are depicted by N (N=A, oligonucleotide 2; N=T,

799 oligonucleotide 1; N=G, oligonucleotide 3; N=C, oligonucleotide 4) (Table 1). Single nucleotide

800 incorporations were carried out as described in Experimental procedures with the different

801 primed-templates, 13 nM of *PabpolB* *exo*⁺/*exo*⁻, 20 nM of *PabpolD* *exo*⁺/*exo*⁻, 200 μM of each

802 dNTP at 55°C for 30 min. A. Reaction with 13 nM *PabpolB* *exo*⁺. B. Reaction with 20 nM

803 *PabpolD* *exo*⁺. C. Reaction with 13 nM *PabpolB* *exo*⁻. D. Reaction with 20 nM *PabpolD* *exo*⁺.

804

Oligonucleotide 15' -CAGGAAACAGCTATGACCATGATTACGAATTCGAGCTCGGTACCCGGGGATCC**T**XTAGAGTCGACCTGCAGGCATGCAAGCTTGGCA-3'**Oligonucleotide 2**5' -CAGGAAACAGCTATGACCATGATTACGAATTCGAGCTCGGTACCCGGGGATCC**A**XTAGAGTCGACCTGCAGGCATGCAAGCTTGGCA-3'**Oligonucleotide 3**5' -CAGGAAACAGCTATGACCATGATTACGAATTCGAGCTCGGTACCCGGGGATCC**G**XTAGAGTCGACCTGCAGGCATGCAAGCTTGGCA-3'**Oligonucleotide 4**5' -CAGGAAACAGCTATGACCATGATTACGAATTCGAGCTCGGTACCCGGGGATCC**C**XTAGAGTCGACCTGCAGGCATGCAAGCTTGGCA-3'**Oligonucleotide 5**

5' -TGCCAAGCTTGCATGCC-3'

Oligonucleotide 6

5' -TGCCAAGCTTGCATGCCTGCAGGTCGACTCTA-3'

805

806 **Table 1 : Damaged or intact oligonucleotides used in this study**

807

	DNA polymerase	dNTP	K_m (μM)	k_{cat} (min^{-1})	k_{cat}/K_m ($\mu\text{M}^{-1} \text{min}^{-1}$)	$f(\text{misinsertion frequency})$
Insertion opposite C	<i>PabpolB</i> exo+	dATP	ND	ND	ND	
		dTTP	ND	ND	ND	
		dGTP	0.25±0.01	425.27 ±0.12	1671.5	
		dCTP	ND	ND	ND	
	<i>PabpolD</i> exo+	dATP	ND	ND	ND	
		dTTP	ND	ND	ND	
		dGTP	0.19±0.02	66.70±0.69	346.36	
		dCTP	ND	ND	ND	
	<i>PabpolB</i> exo-	dATP	18.70±1.23	61.09±0.43	3.27	0.11
		dTTP	5.46±0.05	35.05±0.86	6.42	0.21
		dGTP	1.98±0.35	60.29±0.15	30.41	1.00
		dCTP	32.01±2.11	23.07±0.37	0.72	0.02
	<i>PabpolD</i> exo-	dATP	74.47±9.16	39.71±1.28	0.53	0.16
		dTTP	17.00±3.50	32.50±1.30	1.91	0.57
		dGTP	13.93±1.08	46.81±8.60	3.36	1.00
		dCTP	374.69±129.32	10.41±8.20	0.03	0.01
Insertion opposite AP site	<i>PabpolB</i> exo+	dATP	100±9	87±4	0.79	
		dTTP	121±7	91±4	0.75	
		dGTP	610±40	120±8	0.20	
		dCTP	ND	ND	ND	
	<i>PabpolD</i> exo+	dATP	36±3	114±4	3.17	
		dTTP	650±20	102±2	0.16	
		dGTP	ND	ND	ND	
		dCTP	ND	ND	ND	
	<i>PabpolB</i> exo-	dATP	4.19±0.31	89.42±1.23	21.35	
		dTTP	15.02±0.18	62.92±0.11	4.19	
		dGTP	8.88±2.35	73.68±4.71	8.30	
		dCTP	ND	ND	ND	
	<i>PabpolD</i> exo-	dATP	12.00±2.41	29.03±1.32	2.42	
		dTTP	76.19±0.85	25.55±0.07	0.34	
		dGTP	70.41±7.61	12.43±0.36	0.18	
		dCTP	82.08±6.12	30.72±0.54	0.37	

808

809

810 **Table 2 : Incorporation kinetics by wild-type and exonuclease-deficient *Pabpols***

811

812

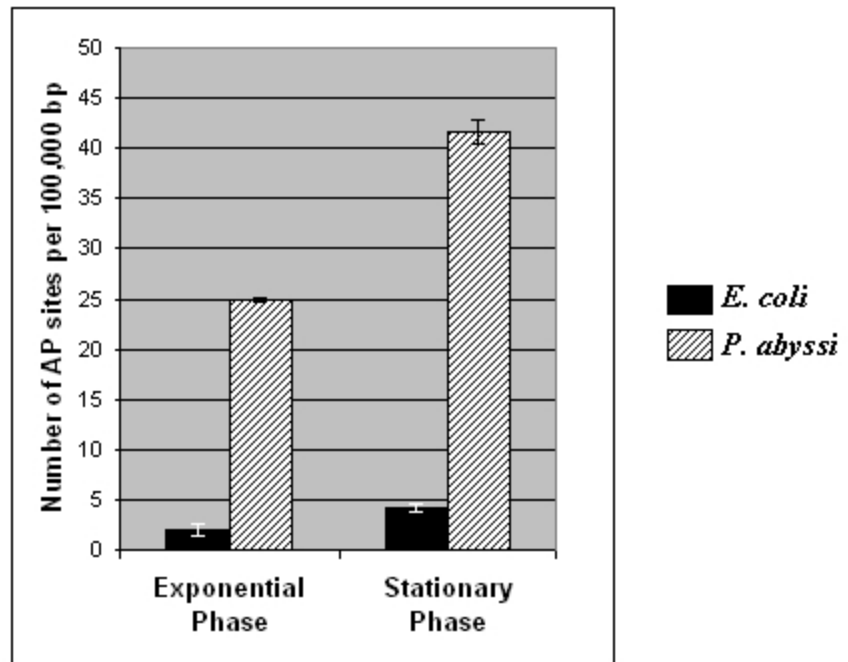
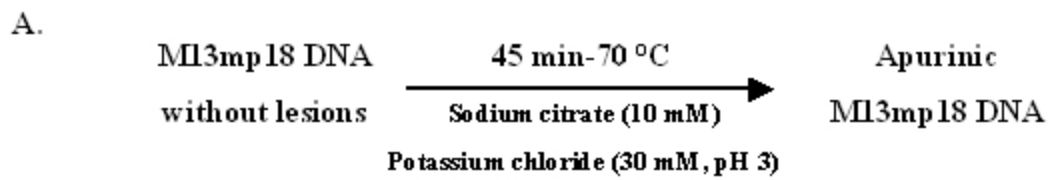


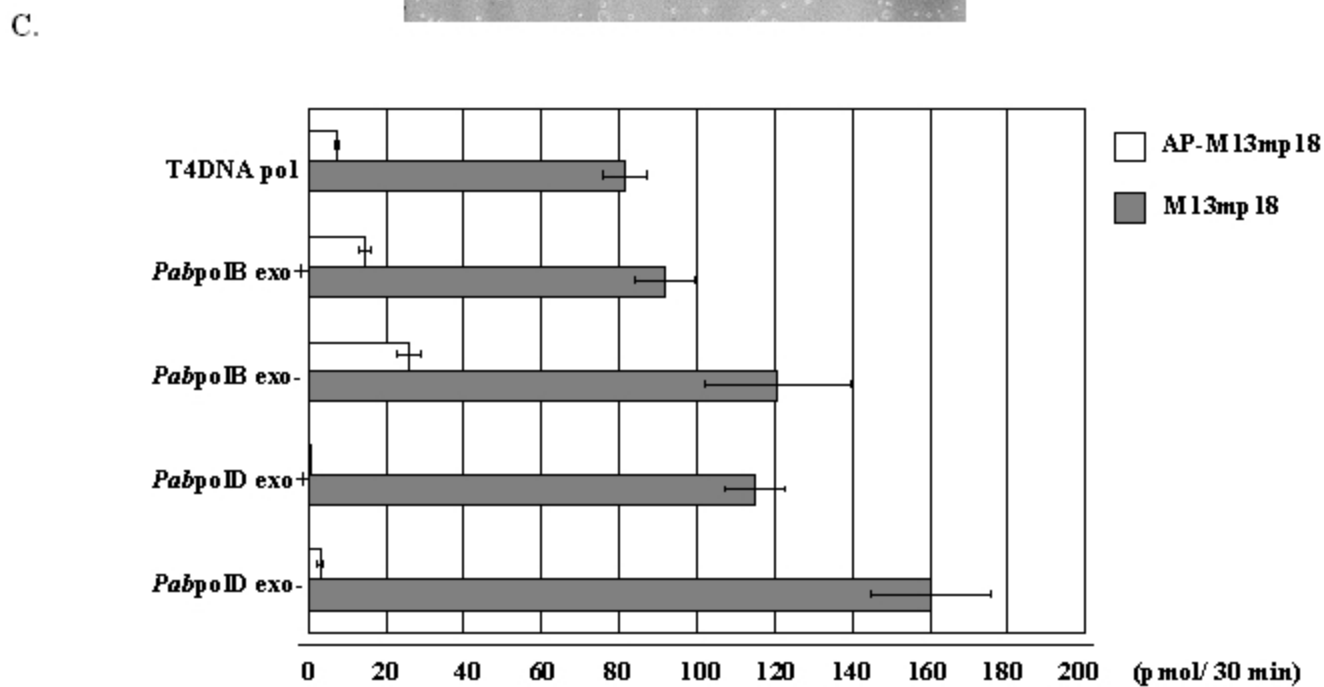
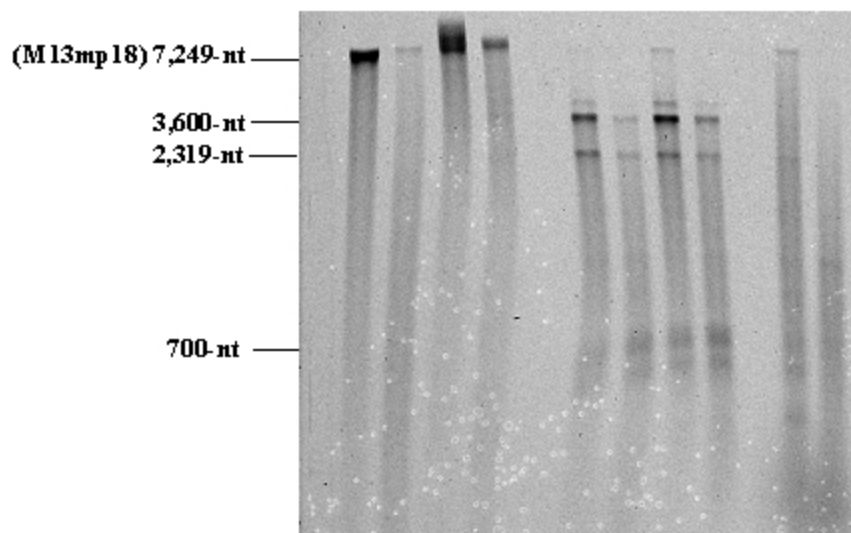
Figure1

Figure 2



B.

	1	2	3	4	5	6	7	8	9	10	11	12	13
M13mp18	+	+	-	+	-	+	+	-	+	-	+	+	-
AP-M13mp18	-	-	+	-	+	-	-	+	-	+	-	-	+
T4DNA pol	-	-	-	-	-	-	-	-	-	-	-	+	+
<i>PabpoD</i> exo+	-	-	-	-	-	-	+	+	-	-	-	-	-
<i>PabpoD</i> exo-	-	-	-	-	-	-	-	-	+	+	-	-	-
<i>PabpoB</i> exo+	-	+	+	-	-	-	-	-	-	-	-	-	-
<i>PabpoB</i> exo-	-	-	-	+	+	-	-	-	-	-	-	-	-



5'-CAGGAAACAGCTATGACCATGATTACGAATTCGAGCTCGGTACCCGGGGATCCTXTAGAGTCGACCTGCAGGCATCAAAGCTTGGCA-3'
 3'-CCGTACGTTCGAACCGT-5'

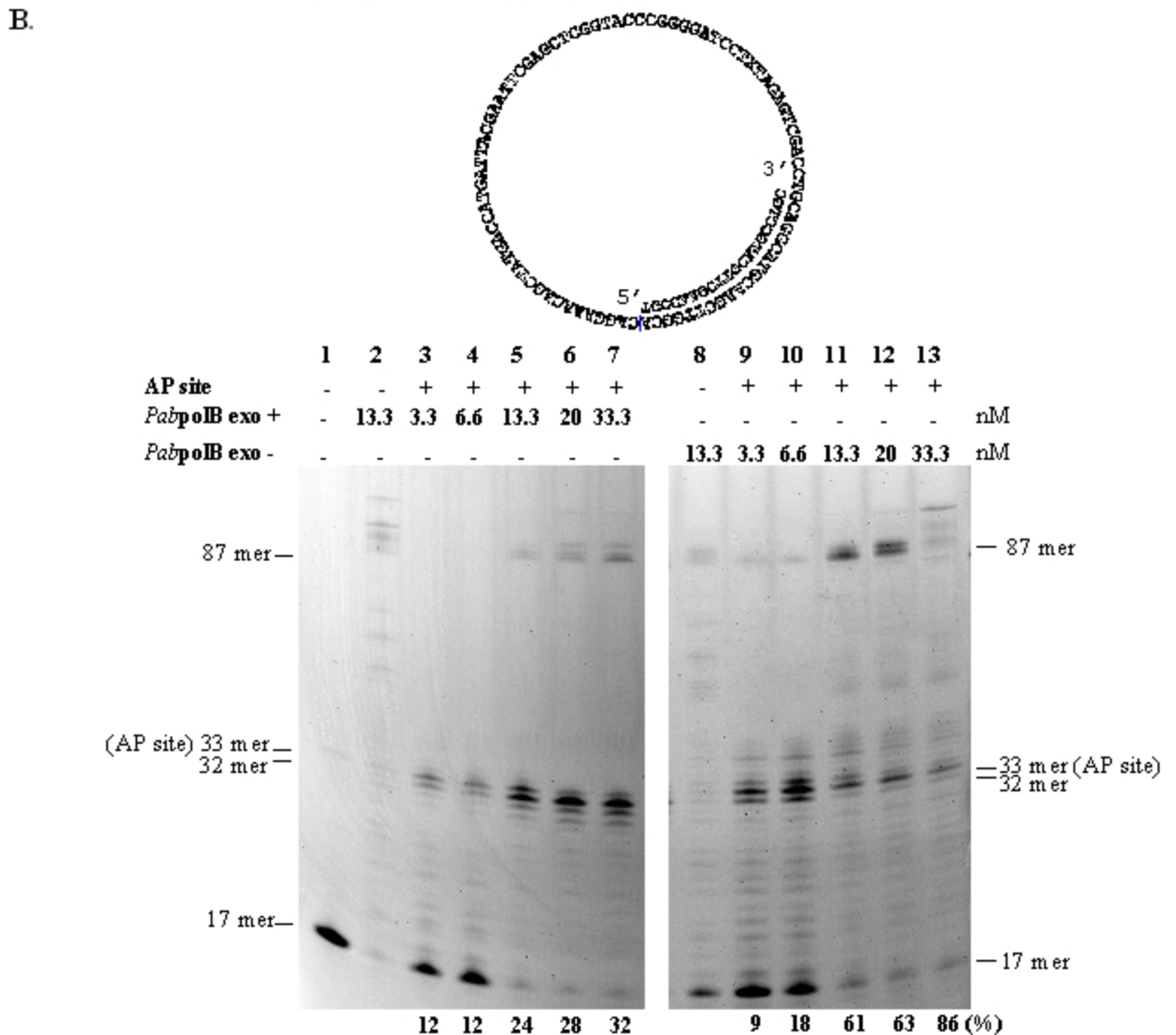
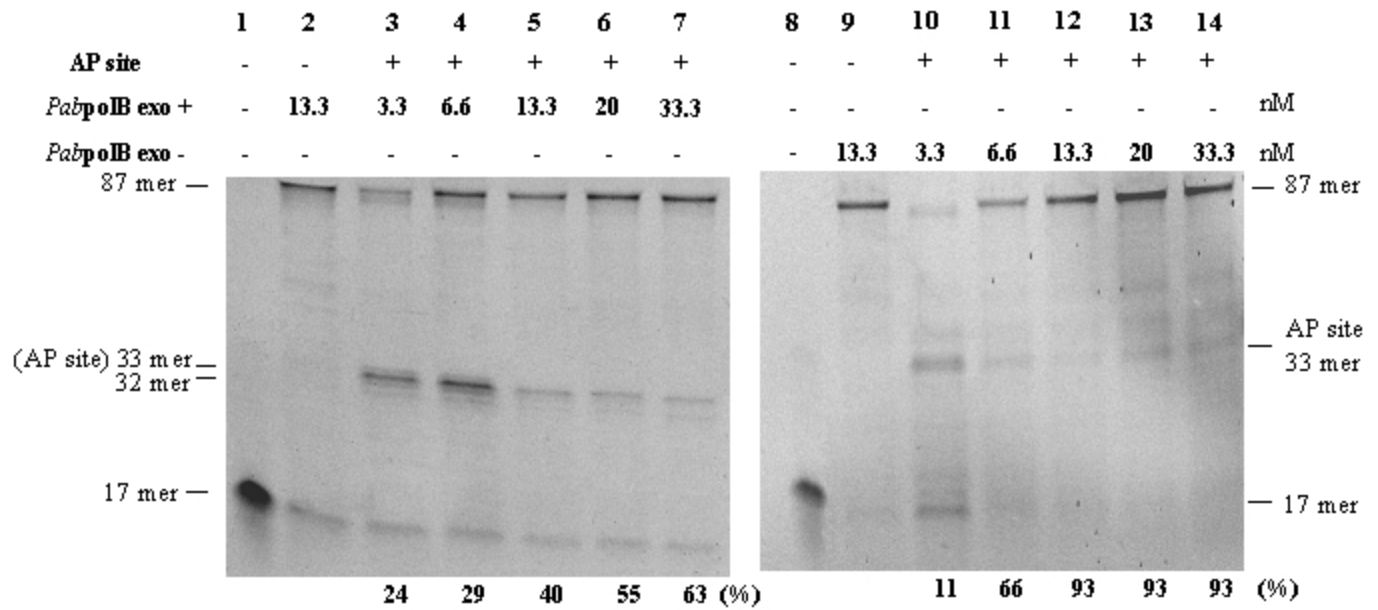
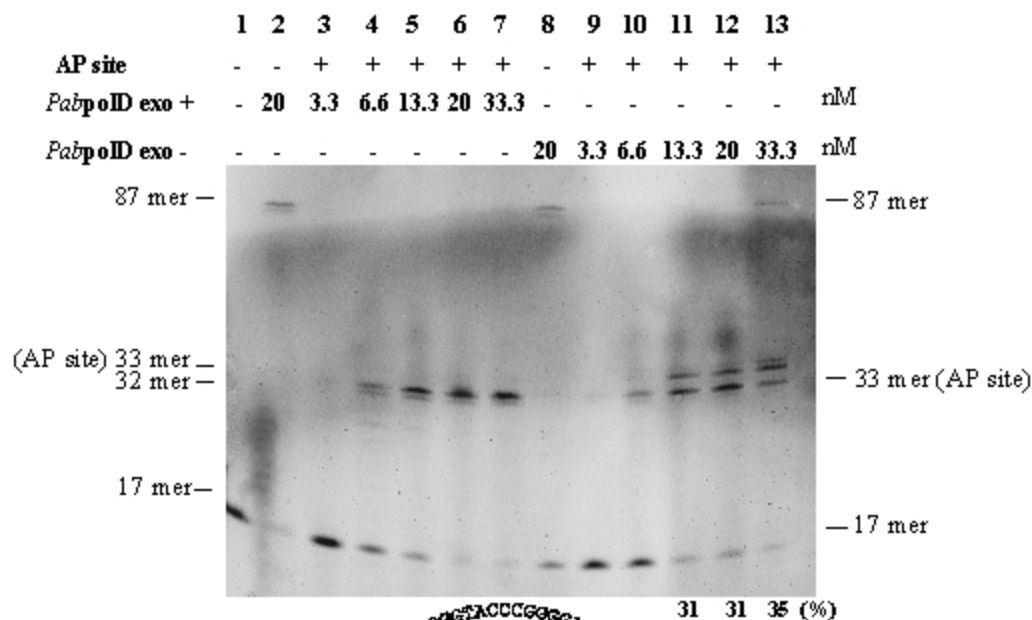


Figure 3

A.

5'-CAGGAAACAGCTATGACCATGATTACGAATTCGAGCTCGGTACCCGGGGATCCTXTAGAGTCGACCTGCAGGCATGCAAGCTTGCCA-3'
 3'-CCGTACGTTCGAACCGT-5'



B.

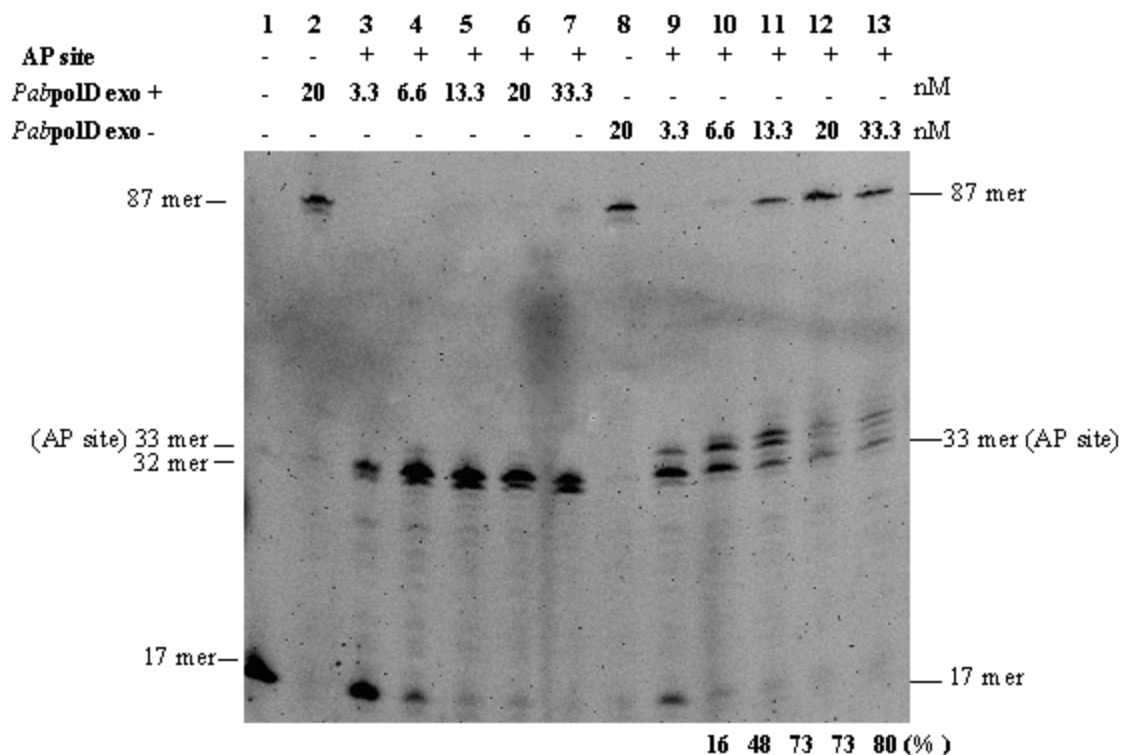
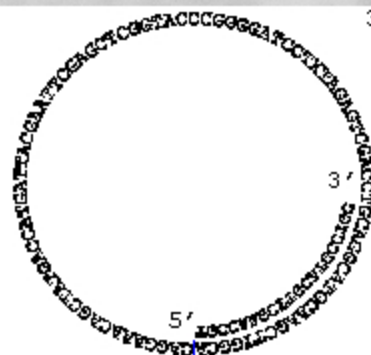
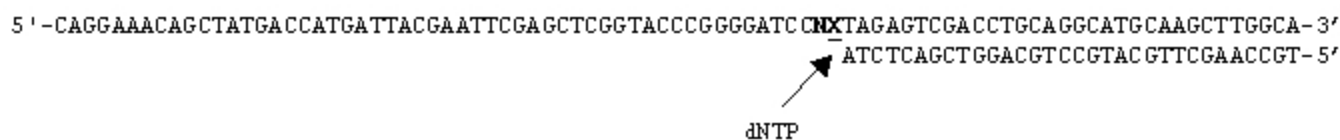
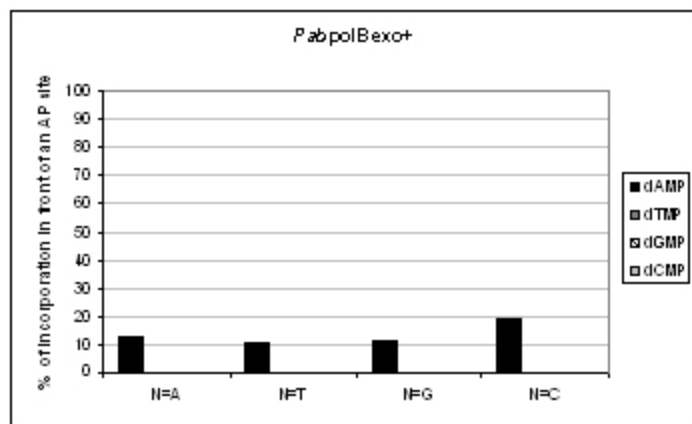


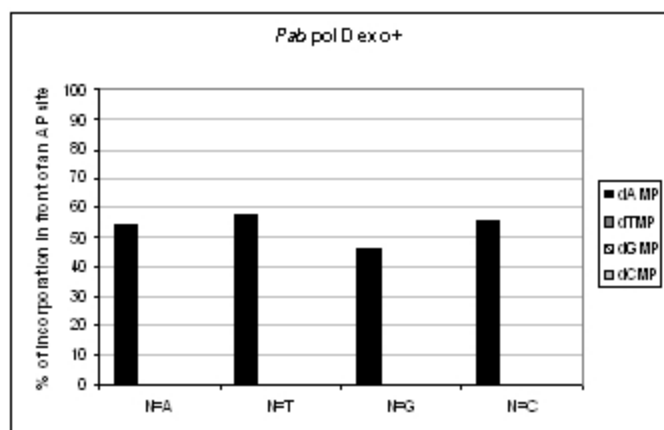
Figure 4 : Replication of AP-containing mini-circular and linear oligonucleotides DNA templates by the wild-type or exonuclease-deficient *Pabp*oD



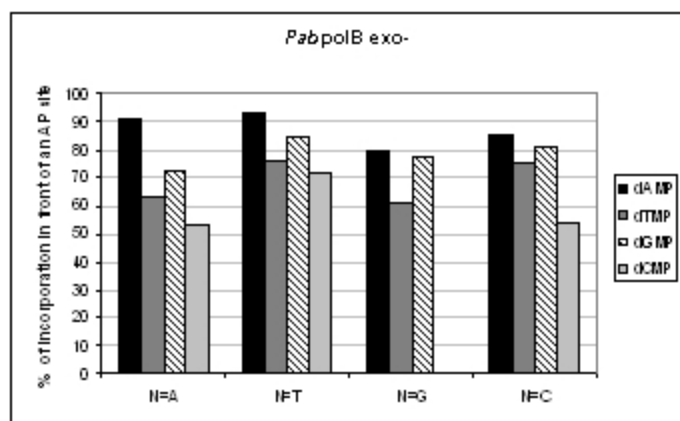
A.



B.



C.



D.

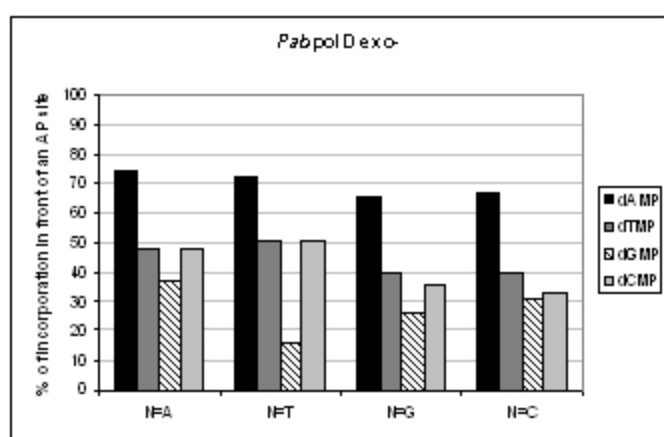


Figure 5

A

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131                                     191
Pab GNEEITFLAVDITFLYHEGE.FFGKGGPTIMISYADREGAKVITWKSIDL.P.YVEVVSSEEMIKRRLVIVT
Pfu GEEETKILAFDITFLYHEGE.FFGKGGPTIMISYADENEAKVITWKNIDL.P.YVEVVSSEEMIKRRLVIT
Pho GNEKLTFLAVDITFLYHEGE.FFGKGGPTIMISYADREGAKVITWKKIDL.P.YVEVVSSEEMIKRRLVIT
Tko GDEELKMLAFDITFLYHEGE.FFAGEGPTLMISYADREGARVITWKNVDLP.YVDVVSSEEMIKRRLVIVV
Mth GAHGLDILSDITFVRNPHGMPDPEKDDIIVMIGVAGNMGYE.SVISTAGDHLDFVEVVEDRELLERFALIVT
Mja EIPKLSVAFDITFVRNPDTEPPEKDDIILMASFWDENGGKVITYKEFNHPN.IEVVKNSEKELIKKIITL
Afu EFPPLKMLVDFDITFVRLSSGMPPEKDDIIVISVKTNDDE.....ILTGDSEKILISDFVRLI
Tac FNPPLKMLVDFDITFVRLSSGMPPEKDDIIVISVKTNDDE.....ILTGDSEKILISDFVRLI

201                                     261
Pab REKDDPDIIVYNGDNDFPFYLLKRAEKLGIKLLPLGGRDNSEPKMQRMGDSLAVEIKKGRHFDLFPVIRRTPI
Pfu REKDDPDIIVYNGDNDFPFYLLKRAEKLGIKLLPLGGRDNSEPKMQRMGDSLAVEIKKGRHFDLFPVIRRTPI
Pho REKDDPDIIVYNGDNDFPFYLLKRAEKLGIKLLPLGGRDNSEPKMQRMGDSLAVEIKKGRHFDLFPVIRRTPI
Tko REKDDPDIIVYNGDNDFPFYLLKRAEKLGIKLLPLGGRDNSEPKMQRMGDSLAVEIKKGRHFDLFPVIRRTPI
Mth IDKKPDLIVYNGDNDFPFYITRAAILGAEGLDLGNDGSKIRTMRRGFANATAIKKGVHFDLFPVIRRTPI
Mja KEY..DIVIYNGDNDFPFYLRARAKIYGIIDINLGDGSEELKIKRGGMEYRSYIPGRVHIDLYPISRRRL
Afu KSYDDPDIIVYNGDNDFPFYLRARAKIYGIIDINLGDGSEENVVFR...GGRPKITGRRLNVDLITAMRIS
Tac RAEDDPDIIVYNGDNDFPFYLRARAKIYGIIDINLGDGSE...IPRRFMNQFVRVHGRLLISDTMWSVKRRLI

271                                     321
Pab NLPYTTDEAYEAI.FGKPKKVVYADEITAKAWETGKGLERVAKVSMEDAKVTYELGKKEF
Pfu NLPYTTDEAYEAI.FGKPKKVVYADEITAKAWESGLENLVARVAKVSMEDAKVTYELGKKEF
Pho NLPYTTDEAYEAI.FGKPKKVVYADEITAKAWETGKGLERVAKVSMEDAKVTYELGKKEF
Tko NLPYTTDEAYEAV.FGQPKKVVYAEITTTAWETGENLVARVAKVSMEDAKVTYELGKKEF
Mth NLDRYTDEAYEQL.FGEEKIDLPGLDRLEWYWRDRDLRDLFRVGLDVAVTRIRAEKI
Mja KLTXYTDEAVVYLL.FGIEKLIKPHTKIVDYWANNDK..TLIEVSLQDAKYVYKIKKYI
Afu DIKIKKDEAVAEPLGTLKELADIEAKDIYRYVSRGK..EKVNLARQDAINPLIKKEL
Tac HPKHESEAVYANMLL.DGEGKDNIDRLHIDDEK...KRRREYVAVGILKADDLRIPEKEL

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B

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352                                     422
Pab VYAILLSDIHVGSKEPCEKAFIKFLEWLNCEVNESKEEELIVSRVRYLLIAGDVVVDSGCHVPGQYSDLVVTP
Pfu VYAILLSDIHVGSKEPCEKAFIKFLEWLNCEVNESKEEELIVSRVRYLLIAGDVVVDSGCHVPGQYSDLVVTP
Pho VYAILLSDIHVGSKEPCEKAFIKFLEWLNCEVNESKEEELIVSRVRYLLIAGDVVVDSGCHVPGQYSDLVVTP
Tko VYAILLSDIHVGSKEPCEKAFIKFLEWLNCEVNSRTEELIVSRVRYLLIAGDVVVDSGCHVPGQYSDLVVTP
Mth FSVAFISDVIHGSOTFLEDAFPMKPVKWLINGDFGSEEQRSLAADVRYLVVAGDVIHDSGCHVPGQYSDLVVTP
Mja LYMAFISDVIHGSKEPCEKAFIKFLEWLNCEVNESKEEELIVSRVRYLLIAGDVVVDSGCHVPGQYSDLVVTP
Afu FYIVPLSDIHVGSKEPCEKAFIKFLEWLNCEVNESKEEELIVSRVRYLLIAGDVVVDSGCHVPGQYSDLVVTP
Tac VYVAFISDVIHGSKEPCEKAFIKFLEWLNCEVNESKEEELIVSRVRYLLIAGDVVVDSGCHVPGQYSDLVVTP

432                                     492
Pab DIFDQYBALANLLANVPEHITMFIQPGNHDAARPAIPOPPEFYKVEYAKPIYKLRNAIITSNPAVIRLHGRD
Pfu DIFDQYBALANLLSNVPHKHTMFIQPGNHDAARPAIPOPPEFYKVEYAKPIYKLRNAIITSNPAVIRLHGRD
Pho DIFDQYBALANLLSNVPHKHTMFIQPGNHDAARPAIPOPPEFYKVEYAKPIYKLRNAIITSNPAVIRLHGRD
Tko DIFDQYBALANLLKQVPHHTMFIQPGNHDAARPAIPOPPEFYKVEYAKPIYKLRNAIITSNPAVIRLHGRD
Mth DIHQYREARLPLGDIRSDIKIIVMIPGNHDSRRIARPOPPEFYKVEYAKPIYKLRNAIITSNPAVIRLHGRD
Mja DIIHQYREARLPLGDIRSDIKIIVMIPGNHDAARPAIPOPPEFYKVEYAKPIYKLRNAIITSNPAVIRLHGRD
Afu DIHQYREARLPLGDIRSDIKIIVMIPGNHDAARPAIPOPPEFYKVEYAKPIYKLRNAIITSNPAVIRLHGRD
Tac NPLERQYANLAEVLYVDVPEVDVYVFMIPGNHDAARPAIPOPPEFYKVEYAKPIYKLRNAIITSNPAVIRLHGRD

512                                     572
Pab FLVHAGRGITDDVVSFVPGLTTHHKPGLPMVELLKMRLHLDAPFCGKVPFIAPDFEDDLVIEVPPDLVDMGHVH
Pfu FLVHAGRGITDDVVSFVPGLTTHHKPGLPMVELLKMRLHLDAPFCGKVPFIAPDFEDDLVIEVPPDLVDMGHVH
Pho FLVHAGRGITDDVVSFVPGLTTHHKPGLPMVELLKMRLHLDAPFCGKVPFIAPDFEDDLVIEVPPDLVDMGHVH
Tko FLVHAGRGITDDVVDVFPNRSHHHRAEAAMVELLKLRLHLDAPFCGKVPFIAPDFEDDLVIEVPPDLVDMGHVH
Mth TLIVHGRSFDMMAMSVNGLSHERSDLIWEELKRRHLDAPFVGRTPPLASELEDLVIEVPPDLVDMGHVH
Mja TLIVHGRSFDLVLGQIRAAASYENPVTMKELIKRRLHLDAPFVGRTPPLASELEDLVIEVPPDLVDMGHVH
Afu TLIVHGRSFDLVLGQIRAAASYENPVTMKELIKRRLHLDAPFVGRTPPLASELEDLVIEVPPDLVDMGHVH
Tac VLVVHGRSFDLVLGQIRAAASYENPVTMKELIKRRLHLDAPFVGRTPPLASELEDLVIEVPPDLVDMGHVH

581
Pab VYDAVVVRG
Pfu VYDAVVVRG
Pho VYDTAVVRG
Mth VMQYKTEG
Tko INAYKKEG
Mja INGVGLVRG
Afu TYGTGFVRG
Tac SHYIGNVKG

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Figure S1: Mapping of conserved residues involved in 3'-5' exonuclease activity among representative euryarchaeal DNA polymerases.

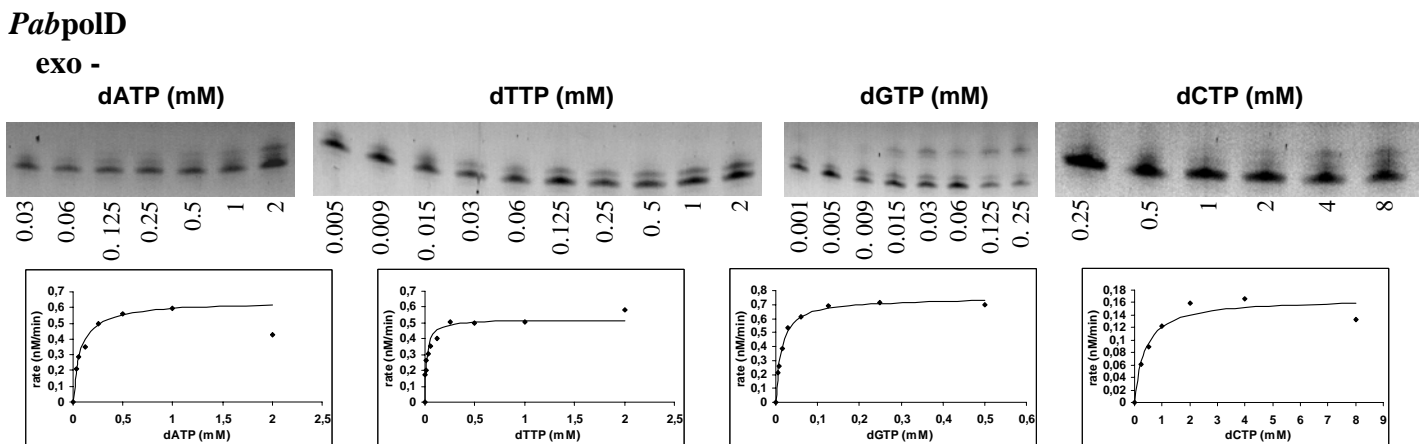
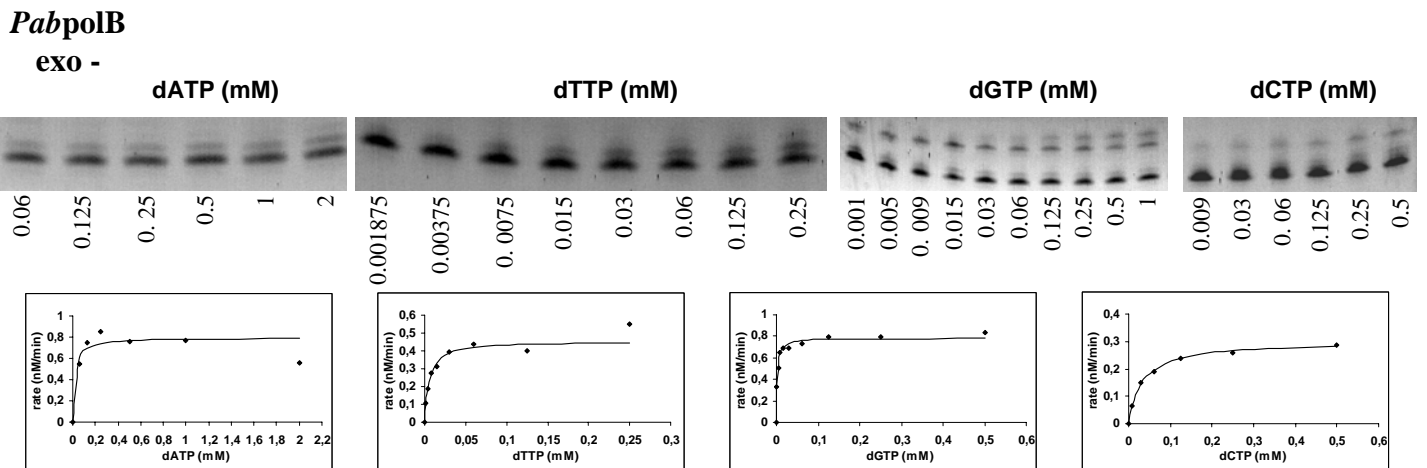
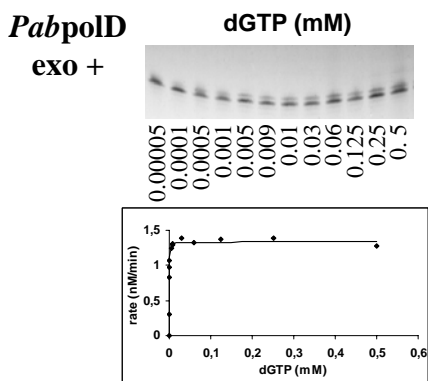
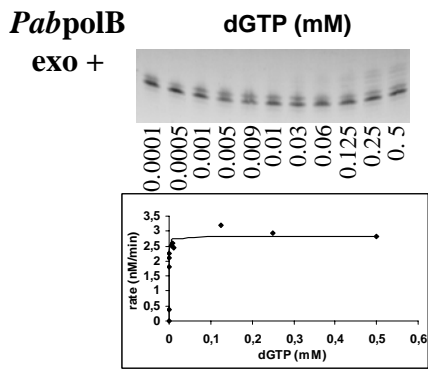
A, Sequence alignment of the exonuclease domain in representative family B DNA polymerases. The sequences are from eight euryarchaeota species, *P. abyssi* (Pab, accession number, gi:14521919), *P. furiosus* (Pfu, accession number, gi:18976584), *P. horikoshii* (Pho, accession number, gi:14591688), *T. kodakarensis* KOD1 (Tko, accession number, gi:57639936), *M. thermoautotrophicum* (Mth, accession number, gi:15679219), *M. jannashii* (Mja, accession number, gi:15669075), *A. fulgidus* (Afu, accession number, gi:11498108), *T. acidophilum* (Tac, accession number, gi:16081956). The star indicates the identified residue in *Pab*polB responsible for proofreading activity. **B**, Partial sequence alignment of the small subunits (DP1s) of the family D DNA polymerases. The sequences are from eight euryarchaeota species, *P. abyssi* (Pab, accession number, gi:14520339), *P. furiosus* (Pfu, accession number, gi:18976390), *P. horikoshii* (Pho, accession number, gi:14590067), *T. kodakarensis* KOD1 (Tko, accession number, gi:57641837), *M. thermoautotrophicum* (Mth, accession number, gi:15679404), *M. jannashii* (Mja, accession number, gi:15668883), *A. fulgidus* (Afu, accession number, gi:11499379), *T. acidophilum* (Tac, accession number, gi:16081371). The star indicates the identified residue in *Pab*DP1 responsible for proofreading activity. Amino acid sequence alignments have been constructed by ClustalW2. Numbering refers to *P. abyssi* amino acid sequences.

A.

Figure S2

5' - CAGGAAACAGCTATGACCATGATTACGAATTCGAGCTCGGTACCCGGGATCCCTCTAGAGTCGACCTGCAGGCATGCAAGCTTGGCA - 3'
 ATCTCAGCTGGACGTCCGTACGTTTGAACCGT - 5'

↑
 dNTP incorporation opposite undamaged nucleotide C



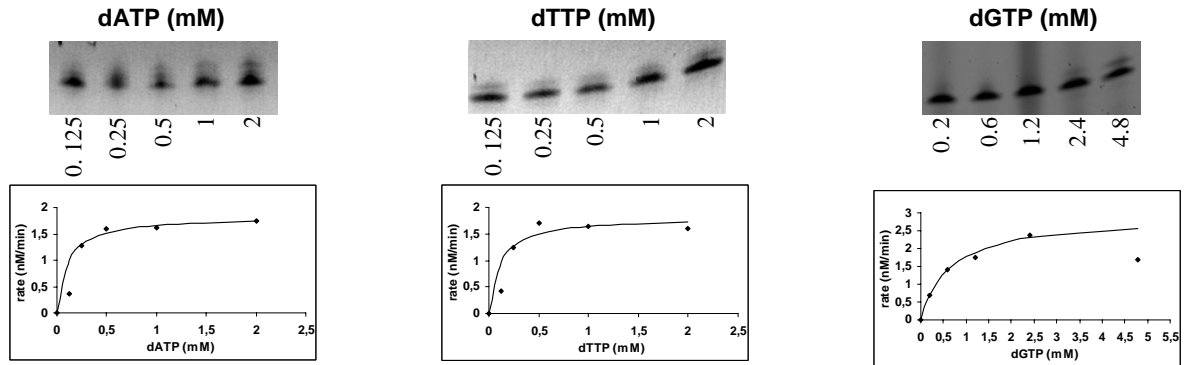
B.



dNTP incorporation opposite an AP site

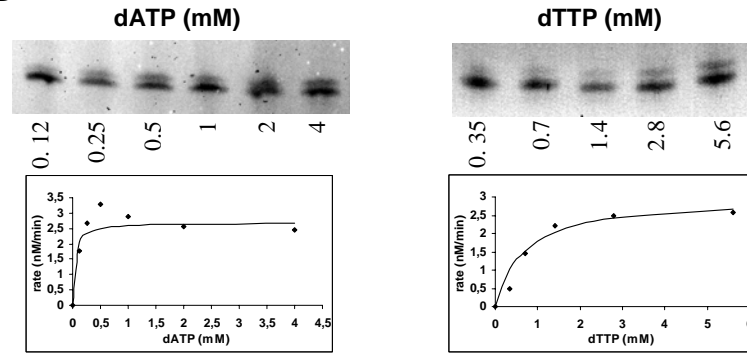
PabpolB

exo +



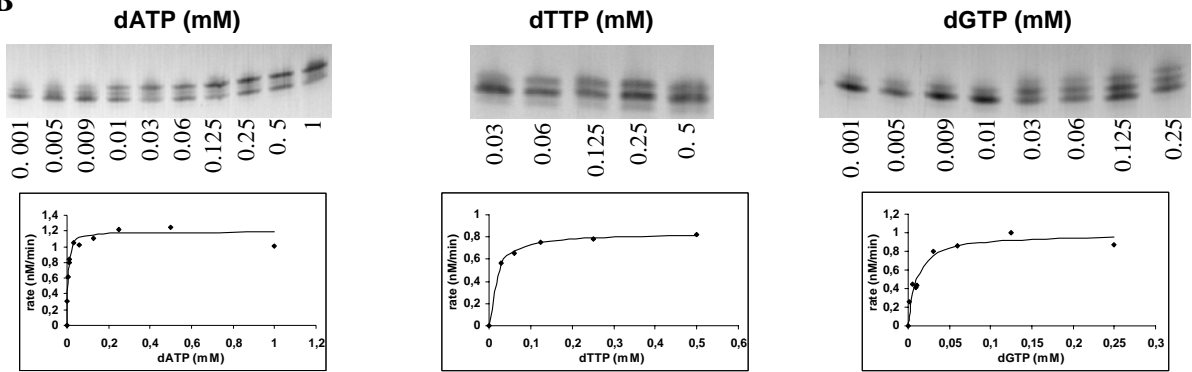
PabpolD

exo +



PabpolB

exo -



PabpolD

exo -

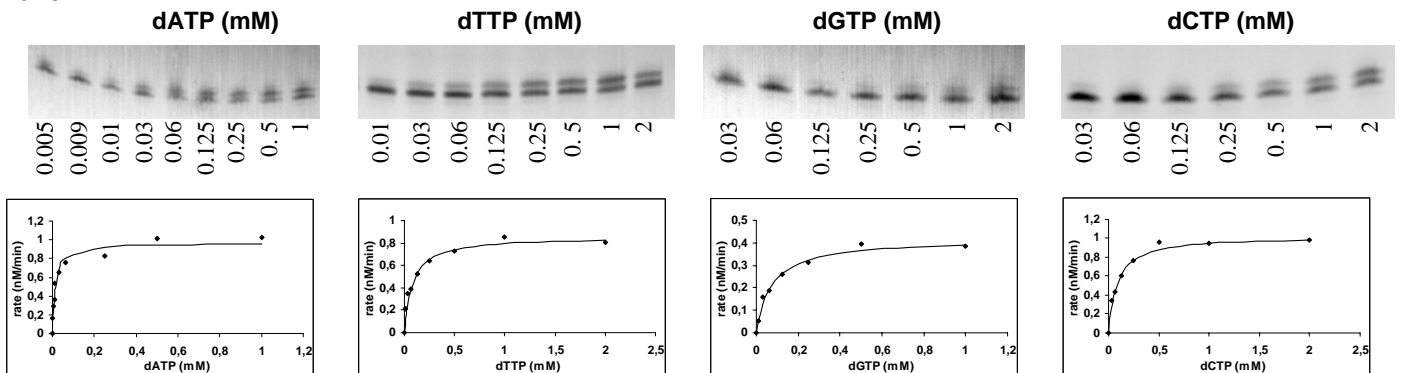


Figure S2: Steady-state kinetics analyses of deoxynucleotide incorporation by *Pabpols*.

A, Incorporation of dNTP opposite undamaged C. The primer-template duplex (8.3 nM) was incubated individually with *PabpolB* *exo+* (6.6 nM), *PabpolD* *exo+* (20 nM), *PabpolB* *exo-* (13 nM) or *PabpolD* *exo-* (16 nM), in the presence of a single dNTP at the indicated concentrations. **B**, Incorporation of dNTP opposite an AP site. The primer-template duplex (8.3 nM) was incubated individually with *PabpolB* *exo+* (24 nM), *PabpolD* *exo+* (20 nM), *PabpolB* *exo-* (13.3 nM) or *PabpolD* *exo-* (33.3 nM), in the presence of a single dNTP at the indicated concentrations. Reactions were carried out at 55°C under standard *Pabpols* assay conditions and the products were resolved by denaturing PAGE. The rate of incorporation was graphed as a function of dNTP concentration and the data were fit to the Michaelis-Menten equation. The k_{cat} et K_m parameters obtained from the fit are listed in Table 2.