Structure and Function of an RNA-Reading Thermostable DNA Polymerase**

Nina Blatter, Konrad Bergen, Oliver Nolte, Wolfram Welte, Kay Diederichs, Jutta Mayer, Markus Wieland, and Andreas Marx*

The high substrate specificity of DNA-dependent DNA polymerases is essential for the stability of the genome as well as many biotechnological applications.1 The discrimination between ribo- and deoxyribonucleotides and between RNA and DNA, particularly in cells, is fundamental since the concentration of ribo moieties by far exceeds that of deoxyribo analogues. Although the selection mechanisms for the incorporation of nucleotides have been investigated intensively for DNA and RNA polymerases,2 much less is known on how DNA-dependent DNA polymerases discriminate between the different nucleic acid templates (DNA versus RNA). Some viral DNA polymerases (e.g. reverse transcriptases) can utilize both DNA and RNA as a template for nucleic acid synthesis. Crystal-structure analysis of these enzymes, complexed either to an RNA or DNA template, have contributed significantly to our understanding of this process.3 However, similar structural data of DNA-dependent DNA polymerases that have a poor propensity to process RNA templates are lacking, presumably because the crystallization is hampered by the formation of unstable complexes that leads to structural heterogeneity.

Structure analysis of KlenTaq DNA polymerase, a shortened form of Thermus aquaticus DNA polymerase, has added significant contributions to the understanding of how DNA polymerases recognize the cognate substrate,3 process abasic sites,3 and non-native nucleotides.4-6 Since our attempts to obtain suitable crystals of KlenTaq complexed to RNA failed, we set out to engineer the enzyme in such a way that it is capable of processing an RNA template more efficiently, which might result in improved crystallization properties. Indeed, we were able to obtain a significantly improved KlenTaq variant from which we obtained structural insights into a DNA-dependent DNA polymerase while processing RNA as a template for the first time. Furthermore, the generated KlenTaq variant turned out to be a thermostable DNA polymerase with significant reverse transcriptase activity, thus resulting in it being a valuable tool for crucial applications in clinical diagnostics and molecular biology, such as transcriptome analysis as well as the detection of pathogens and disease-specific markers.

For the generation of an improved enzyme variant the mutation sites of two KlenTaq (KTq) variants M1 (L322M, L459M, S515R, I638F, S739G, E773G)6 and M747K7 were recombined by DNA shuffling. Both variants, M1 and M747K, were reported to possess either some reverse transcriptase activity or an expanded substrate spectrum. DNA shuffling was employed since the M1 variant, previously evolved in error-prone PCR, comprises six mutations distributed over the enzyme scaffold, but the individual contributions of the mutations are unknown. We generated a library of 1570 clones to obtain high coverage of all mutation combinations (as described in the Supporting Information). Proteins were expressed in 96-well plates and, after lysis and heat denaturation of the host proteins, used in a real-time PCR activity screen with DNA as the template. As approximately 80% of the expressed enzymes were found to be PCR active, we subsequently screened for reverse transcriptase activity by using MS2 RNA as the template, as described earlier.8 In this screening, the two variants RT-KTq1 and RT-KTq2 were identified and selected for further characterization as they showed significantly higher reverse transcriptase activity compared to the parental enzymes as well as a minimal number of mutations. RT-KTq1 possesses three mutations (S515R, I638F, and M747K), whereas RT-KTq2 displays the three mutations of RT-KTq1 and an additional one (L459M, Figure 1a). For comparison, we also combined all seven mutations in one protein scaffold (KTqM1/M747K) by site-directed mutagenesis. KTq wild-type, parental enzymes M1 and M747K, KTqM1/M747K, and RT-KTq1 and 2 were expressed, purified, adjusted to equal protein concentrations (see Figure S1 in the Supporting Information), and further characterized.

First, reverse transcriptase activity of the wild-type and all variants were examined in primer extension experiments with a 5’-[32P]-radiolabeled DNA primer annealed to an RNA template. Reactions were incubated at 72°C for 1 min and subsequently analyzed by denaturing PAGE (Figure 1b). When RNA was used as the template, we observed no primer elongation for the wild-type enzyme. KTq M747K incorporates up to three nucleotides, whereas KTq M1 was able to synthesize longer extension products with up to seven

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[6] M. Sc. N. Blatter, Dipl.-Biol. K. Bergen, Prof. Dr. W. Welte, Prof. Dr. K. Diederichs, Dr. J. Mayer, Dr. M. Wieland, Prof. Dr. A. Marx
Departments of Chemistry and Biology, Konstanz Research School Chemical Biology, University of Konstanz
Universitätstrasse 10, 78457 Konstanz (Germany)
E-mail: andreas.marx@uni-konstanz.de
Priv.-Doz. Dr. O. Nolte
Labor Dr. Brunner Konstanz
Mainaustrasse 48 a/b, 78464 Konstanz (Germany)

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Specific activities of KTq variants on DNA or RNA template.

Table 1:

<table>
<thead>
<tr>
<th>Variant</th>
<th>Specific activity [min⁻¹] DNA template</th>
<th>Specific activity [min⁻¹] RNA template</th>
</tr>
</thead>
<tbody>
<tr>
<td>KTq wt</td>
<td>145 ± 14</td>
<td>n.a. [b]</td>
</tr>
<tr>
<td>KTq M1</td>
<td>252 ± 21</td>
<td>1.67 ± 0.05</td>
</tr>
<tr>
<td>KTq M747K</td>
<td>279 ± 18</td>
<td>0.31 ± 0.01</td>
</tr>
<tr>
<td>KTq M1/M747K</td>
<td>342 ± 11</td>
<td>32.2 ± 1.8</td>
</tr>
<tr>
<td>M747K</td>
<td>181 ± 12</td>
<td>15.7 ± 0.6</td>
</tr>
<tr>
<td>RT-KTq 1</td>
<td>349 ± 13</td>
<td>34.9 ± 1.3</td>
</tr>
<tr>
<td>RT-KTq 2</td>
<td>349 ± 13</td>
<td>34.9 ± 1.3</td>
</tr>
</tbody>
</table>

[a] Quantified conversion of dNTPs per minute was plotted against different polymerase amounts; the resulting slope yielded the specific activity of the respective enzyme. [b] n.a.: not accessible, since no significant activity on an RNA template was observed.

As a result of the higher overall activity of RT-KTq2 compared to RT-KTq1 (Table 1) and a higher stability compared to KTq M1/M747K (see Figure S2 in the Supporting Information), we focused on RT-KTq2 for further in-depth analysis. We next aimed to obtain crystals of RT-KTq2 for structural insights. Therefore, we crystallized RT-KTq2 with an 11 mer DNA primer and a 16 mer RNA template and, for comparison, with the respective primer/template DNA duplex. Details on sample preparation, crystallization, and data statistics are listed in the Supporting Information. We successfully solved structures of RT-KTq2 bound to a DNA primer/template complex and an incoming 2',3'-dideoxyxycytidine-5'-triphosphate (ddCTP) at a resolution of 1.55 Å (termed RT-KTq2DNA, PDB ID 4BWJ).

The overall structure of RT-KTq2DNA is almost identical to closed ternary structures already described for KTq wild-type with a DNA duplex ([4] e.g. PDB ID 3RTV) and shows a low root-mean square deviation (rmsd) of 0.33 Å for Cα (see Figure S3a,b in the Supporting Information). We were also able to obtain crystals of the enzyme variant in complex with a DNA/RNA hybrid at a resolution of 1.75 Å (termed RT-KTq2RNA, PDB ID 4BWJ). This structure shows a half-open state, presumably caused by crystal contacts. Interestingly, a comparison of RT-KTq2RNA with RT-KTq2DNA revealed major changes in the thumb domain as a result of the altered geometry of the hybrid duplex (Figure 2a). The sugar moieties in the RNA template thereby adopt a puckering typical of an A-form conformation, whereas the DNA primer reveals different puckers that deviate from the typical B form (see Table S1 in the Supporting Information). [11] The terminal three base pairs of a DNA/DNA duplex, which are adjacent to the O helix of the DNA polymerase, already adopt an A form. [4a] Therefore, structural changes in the protein scaffold were only required in regions neighboring the nucleic acids downstream of these three terminal base pairs to adapt to the changed geometry of a hybrid DNA/RNA duplex (Figure 2b). The overlay of the DNA/DNA duplex and the hybrid duplex in RT-KTq2DNA demonstrates that part of the thumb domain would have clashed with the hybrid DNA/RNA duplex, which required further structural adaption of the protein scaffold. Consequently, we observe a reduced number of contacts compared to RT-KTq2DNA between the enzyme and the hybrid duplex, mainly in the 3' region of the template, and several changes in the interaction pattern (see Figure S4 in the Supporting Information). Remarkably, the major part of the reordered protein scaffold originates in the thumb domain, downstream of L459M in αH (Figure 2c; see also Figure S3c in the Supporting Information). This shift is probably caused by the mutation of L459 to the sterically less demanding and more flexible side chain of methionine, which facilitates the bending of αH and the subsequent movement of αH1 and αH2 to come in to proximity with the hybrid duplex, thereby maintaining the nucleic acid binding characteristics of this motif (Figure 2c). [4d] This motif is stabilized by the S515R mutation located in αH. The arginine residue displays two conformations in the RT-KTq2DNA structure, thus allowing interactions with the hydroxy group of S513 and the peptide backbone of K305 (Figure 2d). The role of I638F is less apparent, although it probably accounts for a tighter
folding of the finger domain through hydrophobic and π-π stacking interactions with neighboring amino acids. In contrast to the RT-KTq2 DNA structure, R515 and F638 are resolved in simulated annealing omit maps, indicating a stabilization of the corresponding region and further corroborating our observations (see Figure S3d,e in the Supporting Information). Finally, the mutation M747K enhances the positively charged surface potential in proximity to the negatively charged backbone of the DNA. This might promote binding of the nucleic acid backbone and thus, foster the ability to accept aberrant substrates. Interestingly, this is analogous to a finding where a positive charge introduced into the thumb subdomain of a thermostable DNA polymerase allows more efficient processing of a noncognate RNA primer/DNA template duplex by incorporation of ribonucleotide substrates.

Next, the potential of RT-KTq2 to be applied in molecular biology or clinical diagnostics was investigated by performing reverse-transcription polymerase chain reaction (RT-PCR) experiments. RNA is the template for transcriptome analysis, for pathogen detection, as well as for disease-specific marker recognition in RT-PCR. This method is based on the detection and quantification of RNA by the enzyme-mediated reverse transcription of RNA to its complementary DNA and a subsequent amplification by PCR, thereby allowing the abundance of specific RNA molecules to be monitored in real time. Since the DNA polymerases employed in standard PCR are lack sufficient reverse transcriptase activity, two enzymes are required for RT-PCR: a reverse transcriptase and a PCR-competent DNA polymerase. Although these two enzyme mixtures are state of the art, several drawbacks arise from the heat-instability of commonly used retroviral mesophilic reverse transcriptases such as unspecific priming, low yield on complex targets, for example, from secondary structure formation of the mRNA template, and premature reaction termination. In addition, the reverse transcription step usually adds up to 30 min to the PCR protocol, a time addition which during outbreak situations—when hundreds of swabs need to be analyzed—provides a critical bottleneck. Thus, thermostable DNA polymerases capable of reverse transcription and PCR would suprise the use of two enzymes and reduce the time and work. First, the purified enzymes were used in real-time RT-PCR experiments using MS2 bacteriophage RNA as the template. Under the conditions studied, RT-KTq2 leads to product formation approximately 10 cycles earlier than the parental enzyme KTq M1, whereas the wild-type enzyme has no detectable activity (Figure 3a). Formation of the expected product was confirmed by analysis on an agarose gel. Additionally, we demonstrated that RT-KTq2 is able to generate significantly longer products than the parental enzyme KTq M1 (see Figure S5 in the Supporting Information). To investigate the application of RT-KTq2 for the detection of various mRNA transcripts from extracted human total RNA isolated from Jurkat cells, products ranging from 71 to 111 base pairs (bp) were amplified by RT-PCR (Figure 3b). Amplification through DNA contamination could be excluded by choosing intron-spanning targets. Ultrafast two-step cycling was employed in this experiment, with only 30 s initial denaturation at 98°C and 30 cycles of two-step cycling with denaturation for 2 s and 15 s for combined annealing/extension at the temperature indicated.
Also, an additional reverse transcription step was omitted, thus saving further time. Thus, RNA detection was achieved in approximately 20 min. Expected product formation was observed for all targets (Figure 3b). Additionally, we investigated the sensitivity of RT-KTq2 for RNA detection by performing real-time RT-PCR experiments. A 90 nucleotide (nt) RNA stretch of human \(\beta\)-actin mRNA was targeted and amplified from total RNA isolated from Jurkat cells (Figure 3c). Again, two-step cycling was used for the PCR, and product formation was visualized by the binding of SYBRGreen I. We demonstrated that RNA detection was linear over a broad range of concentrations, from 100 ng to as low as 1 pg of RNA, thus competing with commercially available enzymes (see Figure S6 in the Supporting Information).

Next, we set out to detect the influenza virus A in TaqMan-based real-time RT-PCR by using RT-KTq2. Influenza viruses have been the cause of several pandemics\[15\]; therefore, rapid detection methods are required to identify new outbreaks early for subsequent medical intervention. As 5'-nuclease activity is needed for the hydrolysis of dual-labeled TaqMan probes, we used an enzyme blend of Taq DNA polymerase wild-type and RT-KTq2 in our assay. RNA extracts were from respiratory swab samples of patients known to be influenza A positive. A 141 nt RNA stretch coding for the influenza A matrix protein was targeted in real-time RT-PCR by using two-step cycling. Furthermore, a TaqMan probe conjugated with minor groove binders\[16\] was used to increase the reaction specificity in addition to optimized primers recommended for the detection of influenza A. An increase in the fluorescence signal was observed for seven different specimens; correct product formation was visualized on an agarose gel (Figure 3d).

In summary, we have reported the first structure of a DNA-dependent DNA polymerase, whose wild-type ancestor has no significant reverse transcriptase activity, incorporating a nucleotide against an RNA target. The results highlight the altered structure of the DNA/RNA duplex compared to the DNA/DNA duplex. Additionally, crucial amino acid sites were identified that prevent the wild-type enzyme from efficiently using RNA as the template. Whereas the mutation M747K might increase binding affinity to any negatively charged nucleic acid, S515R stabilizes the nucleic acid binding motif in the thumb domain. L459M results in an increased flexibility of the thumb domain, which contributes to preventing a clash with a DNA/RNA hybrid duplex. Furthermore, we have reported some very promising properties of the variant that allow applications such as fast detection and quantification of RNA by RT-PCR in approximately 20 min. The enzyme facilitates multiple applications such as rapid transcription analysis as well as the detection of RNA and DNA pathogens. Since RT-KTq2 exhibits reverse transcriptase and PCR activity in a single enzyme, it has great potential for minimizing labor and time-consuming optimizations without compromising reliable analysis.

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