Central role of the RNA polymerase trigger loop in intrinsic RNA hydrolysis

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The active center of RNA polymerase can hydrolyze phosphodiester bonds in nascent RNA, a reaction thought to be important for proofreading of transcription. The reaction proceeds via a general two Mg$^{2+}$ mechanism and is assisted by the 3′ end nucleotide of the transcript. Here, by using *Thermus aquaticus* RNA polymerase, we show that the reaction also requires the flexible domain of the active center; the trigger loop (TL). We show that the invariant histidine (β His1242) of the TL is essential for hydrolysis/proofreading and participates in the reaction in two distinct ways: by positioning the 3′ end nucleotide of the transcript that assists catalysis and/or by directly participating in the reaction as a general base. We also show that participation of the β′ His1242 of the TL in phosphodiester bond hydrolysis does not depend on the extent of elongation complex backtracking. We obtained similar results with *Escherichia coli* RNA polymerase, indicating that the function of the TL in phosphodiester bond hydrolysis is conserved among bacteria.

Catalysis by the active center of RNA polymerase (RNAP) is thought to proceed through a general two Mg$^{2+}$ ion mechanism (1–4). Structural and biochemical studies, however, revealed that during phosphodiester bond synthesis a flexible domain of the active center, the trigger loop (TL), is also required for catalysis (5–10). The TL was proposed to act by orienting the triphosphate moiety of the incoming NTP for efficient attack by the 3′ hydroxyl of RNA (5–7, 10). In crystal structures of RNAP elongation complexes, the TL was observed in two distinct states, folded and unfolded. Only in the folded conformation are amino acids of the TL close enough to the catalytic center to be able to participate in the reaction, so the folded form of the TL is assumed to be the catalytically active state (6–9).

The active center of RNAP is also able to perform hydrolysis of the phosphodiester bond of the transcript (3, 11, 12). This reaction almost exclusively proceeds at the penultimate (second) phosphodiester bond at the 3′ end of RNA transcript (12, 13). Such preference is because of the fact that the rate of hydrolysis of the second phosphodiester bond is hugely increased by assistance from the 3′ end NMP of the RNA (transcript-assisted hydrolysis) (12). Transcript-assisted hydrolysis proceeds in a 1-bp backtracked conformation of the elongation complex, in which the 3′ end NMP of the transcript disengages from the complementary template base to assist with the second phosphodiester bond hydrolysis (12, 14). The 3′ end NMP was proposed to provide its chemical groups to the active center of RNAP to increase the affinity to the second Mg$^{2+}$ ion (Mg$^{2+}$II) and/or to position the attacking water molecule (12). Reaction is especially efficient when RNAP incorporates an erroneous nucleotide, given that the misincorporated elongation complex is stabilized in the 1-bp backtracked state. Hydrolysis of the second phosphodiester bond, therefore, is thought to be important for proofreading of transcription (12).

The data on participation of the TL in the cleavage reaction are controversial. The TL was proposed to be required for RNA hydrolysis on the basis of the fact that RNA cleavage by *Thermus aquaticus* RNAP was inhibited by the antibiotic Streptolydigin (Stl) (6). According to crystallographic data, Stl binds to the TL and freezes it in the unfolded, inactive, conformation (6, 15). This proposal, however, was inconsistent with the results of a recent crystallographic study of several backtracked elongation complexes formed by yeast RNAP II (backtracked by 1, 2, 3, 4, 7, and 13 bp) (14). The TL residues in these complexes were too far from the catalytic Mg$^{2+}$ and the reacting groups, suggesting that the TL may not participate in RNA cleavage. A recent biochemical study on *Escherichia coli* RNAP also suggested that the TL is not required for intrinsic hydrolysis (10). The authors used RNAP that lacked the TL (ΔTL RNAP) and showed that WT and ΔTL RNAPs cleaved RNA in 2-bp backtracked elongation complexes at similar rates. Notably, these rates were 1–2 orders of magnitude slower than those observed in earlier studies with *T. aquaticus* RNAP (12). These results suggested that the mechanisms of RNA hydrolysis used by *E. coli* and *T. aquaticus* RNAPs may differ. It was also suggested that the features of the structure of elongation complexes (extent of complementarity of the DNA strands and/or length of the RNA/DNA hybrid) may influence the mechanism of hydrolysis (10). Taken together, the role of the TL in RNA hydrolysis, and consequently in proofreading of transcription, remains unclear.

**Results**

**H1242 of the TL Is Essential for Phosphodiester Bond Hydrolysis.** We investigated the mechanism whereby the TL participates in phosphodiester bond hydrolysis and the impact of this on proofreading of transcription. We used *T. aquaticus* RNAP, for which several crystal structures have been solved (16–18). Its active center is identical in amino acid sequence to that of *Thermus thermophilus* RNAP, for which several crystal structures are also available (6, 7, 15, 19–22). We constructed mutant RNAPs either lacking the whole TL (ΔTL RNAP) (5, 6) or bearing single alanine substitutions of the TL amino acids, R1239, H1242, and Q1235 (5). In the crystal structure of the *T. thermophilus* RNAP elongation complex with NTP bound in the active center, these amino acids are in the close proximity to the reacting groups (7). During phosphodiester bond formation, R1239 and H1242 were proposed to stabilize the transition state of the reaction (5, 7, 10), whereas Q1235 was suggested to properly orient R1239 and H1242 (5). These amino acids may potentially occupy similar positions close to the reactants during phosphodiester bond hydrolysis and thus participate in the reaction (Fig. L4). We used elongation complexes in which the 3′ end NMP of the RNA was noncomplementary to the corresponding template base, but with fully complementary template and nontemplate strands (Fig. S1). Such complexes are stabilized in the 1-bp backtracked conformation, thus eliminating possible translational effects on the rate of second phosphodiester bond hydrolysis. Given that the properties of the reactions depend on the identity

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of the 3' end NMP of the transcript, we used elongation complexes bearing either AMP or UMP at the 3' end [MEC(A) and MEC(U), respectively; Fig. S1]. The 3' OH and the phosphate group of UMP, and the 3' OH, the phosphate, and the N7 atom of AMP have been proposed to participate in chelation of the second catalytic Mg$^{2+}$ ion, as inferred from the effects of substitutions of these groups on the Mg$^{2+}$ dependence of the second phosphodiester bond hydrolysis in these complexes (12). The pH profiles of hydrolysis in these complexes were shown to be different, suggesting that the base moieties of the AMP and UMP may influence the reaction (12).

Deletion of the TL drastically decreased the rate of second phosphodiester bond hydrolysis in both MEC(A) and MEC(U) (Fig. 1B and C). This result is consistent with data on inhibition of the cleavage reaction by the antibiotic Stl, which inhibits TL folding (6), and indicates that the TL is indeed required for phosphodiester bond hydrolysis. Next, we tested single amino acid substitutions in these reactions. Substitutions R1239A and Q1235A had no or minor effects on the reactions (Fig. 1B and C). Interestingly, alanine substitution of H1242 decreased the hydrolysis rates in both MEC(U) and MEC(A) by ~130- and ~200-fold, respectively, effects close to those of the whole TL deletion (~140- and ~430-fold, respectively; see also Fig. 1B and C). These results thus indicate that the function of the TL in hydrolysis is determined by H1242. Moreover, if during phosphodiester bond hydrolysis and phosphodiester bond synthesis the TL folds in a similar way, H1242 may be positioned in close proximity to the scissile phosphodiester bond (Fig. L1).

Two Roles of H1242 in Phosphodiester Bond Hydrolysis. To investigate the role that H1242 plays during hydrolysis, we examined the Mg$^{2+}$ and pH dependences of phosphodiester bond hydrolysis catalyzed by WT and H1242A RNAPs in MEC(U) and MEC(A) (Fig. 2A–C). As seen from Fig. 2A, the H1242A substitution did not change the $K_M$[Mg$^{2+}$] during hydrolysis in MEC(U). The result therefore suggests that H1242 does not participate in Mg$^{2+}$ chelation and must assist the reaction via another mechanism. In contrast, the pH profile of hydrolysis in MEC(U) was strongly altered by the H1242A mutation (Fig. 2C), suggesting that H1242 directly participates in the reaction. The pH profile of hydrolysis in MEC(U) by WT RNAP reveals that the apparent $pK_a$ of the reaction is around 7.8 (Fig. 2C), which is in the $pK_a$ range for a histidine residue. This result suggests that H1242 participates in hydrolysis as a general base, presumably by deprotonating the attacking water molecule. Note, however, that the slope of the log-log curve is less than 1, indicating that more than one $pK_a$ is being titrated even in the absence of H1242, and some other groups, possibly UMP, still contribute to general acid/base catalysis of phosphodiester bond hydrolysis.

Unexpectedly, the $K_M$[Mg$^{2+}$] for hydrolysis in MEC(A) by H1242A RNAP was ~35 times higher than that by WT RNAP (Fig. 2B). This effect of the H1242A substitution on the $K_M$[Mg$^{2+}$] was of similar magnitude to that of a carbon substitution of N7 of the 3' end AMP (12) N7 being proposed to participate in the chelation of Mg$^{2+}$ (12, 14). The above result therefore suggests that, in the absence of H1242, the 3' end AMP of the RNA cannot be oriented properly in the active center to assist RNAP in chelating the Mg$^{2+}$ ion via its N7 and, possibly, via its 3' OH groups.

The pH profile of hydrolysis by H1242 RNAP in MEC(A) was strongly altered as compared to that of WT RNAP (Fig. 2C). However, unlike MEC(U), the slope of the pH profile of hydrolysis by H1242A RNAP in MEC(A) was close to 1, indicating that only one $pK_a$ is being titrated ($pK_a$ of Mg$^{2+}$–OEF complex) and that the reaction is no longer assisted by either H1242 or the 3' end AMP. This result is consistent with the above proposal that H1242 is responsible for proper orientation of the AMP moiety in the active center. We cannot distinguish if H1242 acts as a general base during hydrolysis in MEC(A), as it does in MEC(U), or if the altered pH profile is because of the loss of assistance from the 3' AMP. However, disappearance of the inflection point at pH ~ 7.5 in the pH profile of H1242A RNAP argues in favor of the former possibility. From the above data, we conclude that during phosphodiester bond hydrolysis H1242 of the TL directly participates in the reaction and that it does so in two ways: (i) as a general base and (ii) by positioning the 3' NMP moiety of the RNA for transcript-assisted catalysis. It is also possible that the 3' end AMP and the TL may mutually affect their conformations in the active center and, as a result, their modes of participation in the reaction.

Previously, we have proposed that in the correct elongation complexes (with a correctly paired 3' end of the RNA) cleavage
still proceeds almost exclusively at the second phosphodiester bond, because of assistance from the 3′ end NMP of the RNA, which makes slow unassisted cleavage of the ultimate phosphodiester bond at the 3′ end of the RNA undetectable (12). Following from the above, H1242 of the TL is essential to orient the 3′ end AMP of the RNA for AMP-assisted hydrolysis of the second phosphodiester bond. Therefore, in an elongation complex with the 3′ end AMP of the RNA correctly paired, the H1242A and ΔTL RNAPs should lose their preference for second phosphodiester bond cleavage. In agreement with this prediction, the H1242A mutation reduced, and deletion of the whole TL eliminated the preference for second phosphodiester bond hydrolysis (Fig. 2D). The differences in the magnitude of the effects of the H1242A substitution versus deletion of the TL were also consistent with their effects on the rate of AMP-assisted hydrolysis (Fig. 1C).

Role of the TL in Proofreading of Transcription. Proofreading via RNA cleavage was proposed to be an important property of transcription (12), because misincorporation leads to the production of erroneous RNA messages and results in stalling of elongation (5, 12, 13, 23). Given that the TL is required for efficient transcript-assisted cleavage, the intrinsic way of correcting errors, we expected that mutation of the catalytic H1242 would strongly diminish proofreading activity. To test that, we supplied elongation complexes with NTPs complementary for incorporation at positions +2, +3, and +4 but not +1, thus promoting a misincorporation event followed by extension of the erroneous transcript (see the scheme in Fig. 3A). Transcripts in the initial elongation complexes were labeled at the 3′ end, allowing us to simultaneously observe misincorporation, erroneous transcript extension, and transcript-assisted proofreading via dinucleotide cleavage. As seen from Fig. 3A and B, H1242A RNAP readily extended about half of the transcripts beyond the misincorporated nucleotide, whereas WT RNAP removed almost all (~90%) of the wrong NMP from the RNA without transcribing through the erroneous position. Note also that the rate of incorporation by H1242A RNAP was ~20 times slower than that of WT RNAP, which gives H1242A RNAP more time for proofreading. Therefore, the actual contribution of the TL to proofreading is expected to be an order of magnitude stronger than observed in the experiment. From this we propose that the TL plays a critical role in transcript-assisted proofreading.

The TL Is Required for RNA Hydrolysis in Complexes Backtracked by >1 bp. In the course of carrying out the above experiments, we uncovered complexes that had undergone efficient cleavage at the third phosphodiester bond. Formation of such complexes required misincorporation of NMP that was complementary to position +2. In this case, misincorporation was followed by efficient extension by one more NMP, to position +2 (Fig. 4A and Fig. S3A). Almost no proofreading of the misincorporation event was detected, but instead efficient cleavage occurred at the third phosphodiester bond (Fig. 4A and Fig. S3A). This result indicates that, after extension to +2 position, the elongation complex backtracks by 2 bp. To investigate the role of H1242 in hydrolysis of the third phosphodiester bond, we made use of such an elongation complex, MEC(UU) 2n (*n* for naturally backtracked), in which RNAP is backtracked by 2 bp and hydrolyzes the third phosphodiester bond with release of the 3′ terminal tri- mer pGpUpU (cartoon in Fig. 4A). We used MEC(UU) 2n obtained through misincorporation of UMP [rather than via assembly with mismatched RNA like MEC(UU)] as a control. As expected, hydrolysis by H1242A RNAP in MEC(UU) 2n was strongly diminished as compared to WT RNAP (Fig. 4A; compare lanes 1–7 and 8–14). The third phosphodiester bond in MEC(UU) 2n was also cleaved much more slowly by H1242A RNAP than by WT RNAP (Fig. 4A; compare lanes 15–21 and 22–28). Therefore, our results suggest that participation of H1242 of the TL in phosphodiester bond hydrolysis is independent of the extent of elongation complex backtracking. Whether or not the hydrolysis in complexes backtracked by more than 1 bp is assisted by the transcript requires further investigation. However, the comparable rates of hydrolysis of the second and the third phosphodiester bonds (compare lanes 1–7 and 15–21 in Fig. 4A and lanes 1–7 and 8–14 in Fig. S3A) argues in favor of participation of the transcript in the reaction.

The Role of the TL in RNA Cleavage Is Conserved Among Bacteria. To test if the role of H1242 of the TL in RNA hydrolysis is conserved among bacterial RNAPs, we used *E. coli* RNAP, which is only distantly related to *T. aquaticus*. We constructed mutant *E. coli* RNAP (EcH936A RNAP) bearing an alanine substitution of H936, the homologue of H1242 in *T. aquaticus*. Using EcWT and EcH936A RNAPs in an assembled
Fig. 3. Impact of TL catalyzed hydrolysis on the proofreading of transcription. (A) The scheme above the gels shows the experimental setup. RNA in the elongation complex (ECZ in Fig. S1) is labeled at the 3′ end by incorporation of [α-32P]UTP (asterisk in the scheme). The 3′ end labeling of RNA enables the monitoring of misincorporation, readthrough, and proofreading simultaneously. Misincorporation of 1 mM ATP, readthrough in the presence of 100 μM CTP, and proofreading (cleavage of pUpA) by WT and H1242A RNAPs are shown. The cleavage products, migrating slower than a dinucleotide, originate from hydrolytic cleavage in readthrough complexes (with RNA longer than 15 nucleotides) after backtracking (see Fig. 4 and Fig. S3) (30). Products are designated in colors the same as those used in the scheme of the reaction shown above the gels. A black vertical line separates lanes originating from the same gel that were brought together. (B) Graphic representation of the data from A. The color code for the plots is the same as in A.

Discussion
Hydrolysis of the second phosphodiester bond of the transcript is assisted by the 3′ end NMP of the transcript. This function seems to be similar to the one proposed for the TL in phosphodiester bond synthesis on the basis of biochemical and crystallographic studies: The TL was suggested to orient the triphosphate moiety of the NTP for the reaction to proceed (5, 7, 10). Therefore, TL-assisted hydrolysis cooperates with the riboyme-like transcript-assisted hydrolysis to increase the efficiency of the reaction. This situation resembles modern ribozymes, whose activities are frequently assisted by protein components.

Earlier we proposed that the dependence of the pH profiles of transcript-assisted hydrolysis on the identity of the RNA 3′end NMP may be because of participation of the RNA 3′NMP in the reaction as a general acid/base (12). The discovery of the new player in the reaction, the TL, suggests that the 3′ end NMP may also change the microenvironment for TL-assisted hydrolysis or influence the orientation of the catalytic histidine of the TL. However, the different pH profiles of hydrolysis in the MEC(A) and MEC(U) complexes, even in the absence of the TL (Fig. S4), argue that the 3′ end NMP may still directly participate in the reaction.

Our discovery of the important role of the TL in phosphodiester bond hydrolysis contradicts the conclusions of the recent study on E. coli RNAP (10). The authors showed that deletion of the whole TL did not influence the rate of the reaction. However, they noted that, because of extensive noncomplementarity of the template and nontemplate strands and a short RNA-DNA hybrid, the elongation complexes they used might be “locked” in a configuration that precludes the TL from functioning (10). This idea is supported by the fact that the rates of hydrolysis by both E. coli and T. aquaticus WT RNAPs observed in our study (Figs. 1B and 4B) were about an order of magnitude faster than those observed by Zhang et al. (10), even though we used a lower pH (pH 8.0 versus 9.0) and lower Mg2+ concentration (10 versus 20 mM). In addition to using complexes assembled with mispaired 3′ end of RNA, we also used elongation complexes that had undergone misincorporation prior to cleavage [MEC(U)n, MEC(G)n, MEC(UU)n, and MEC(GG)n], a more physiological
way to bring and stabilize complexes in 1- and 2-bp backtracked states (Fig. 4 A and C).

Notably, in the crystal structures of yeast RNAP II in the backtracked state the invariant histidine residue of the TL, H1085 (homologue of H1242 of T. aquaticus), is far from the scissile phosphodiester bond, making it unlikely that it participates in the cleavage reaction (14). However, the rate of phosphodiester bond hydrolysis by yeast RNAP II is about 2 orders of magnitude slower than that of T. aquaticus RNAP (12, 14). This rate difference is close to the effect of TL deletion in T. aquaticus RNAP on hydrolysis (see above). Taken together, these data suggest that RNAP II may not use its TL for hydrolysis. Nevertheless, it remains unclear whether the eukaryotic TL has lost its function in RNA hydrolysis or the bacterial TL has acquired it after these multisubunit RNAPs diverged early in evolution.

Recently, it was suggested that misincorporation at template position +1 occurs faster if the noncognate NTP is complementary to the template base in position +2 (25). The authors proposed that in this situation the +2 template base may shift into the active center (template misalignment) thus leading to recognition of the noncognate NTP as a cognate one. We did not observe any dependency of the rate of misincorporation of a noncognate NTP in position +1 on the identity of the +2 position of the template for either T. aquaticus or E. coli RNAPs (Fig. S3B). We, however, observed the following propensity. When there is no complementarity of the +1 misincorporated NMP and +2 position of the template, the misincorporated complex is stabilized in a 1-bp backtracked state and thus can only slowly be extended. In contrast, when incoming NTP is noncomplementary to position +1 but complementary to position +2 of the template, after misincorporation at +1, the mismatched RNA is quickly extended to position +2. These observations suggest that the noncognate NMP at the 3′ end of the RNA in such a misincorporated complex is recognized by RNAP as a correct one, thus allowing its efficient extension. Though disagreeing with Kashkina et al. (25) on the mechanism of the phenomenon, our results agree with their proposal that misincorporation of the active center (template misalignment) thus leading to recognition of the noncognate NTP as a cognate one.

Material and Methods

Mutant RNAP Construction and Purification. T. aquaticus RNAP lacking the TL (β′ residues 1238–1254 substituted by glycine residue) was constructed as described (5, 6). T. aquaticus mutant RNAPs bearing single alanine substitutions of the β′ Q1235, R1239, and H1242 (Q1235A, R1239A, and H1242A RNAPs, respectively) were obtained by site-directed mutagenesis in a coexpression system for recombinant RNAP (5, 26). WT and mutant core RNA polymerases were purified as described (27) and their purification was performed as described (27). To obtain E. coli EcH936A RNAP, the mutation was introduced in the β′ subunit coded under inducible promoter in plasmid pRL663. WT and mutant RNAPs production was performed as described (27) and (28).

Transcription Assays. Elongation complexes were assembled and immobilized on Ni-nitrilotriacetate beads (Qiagen) as described (5, 12), in transcription buffer containing 40 mM KCl and 20 mM Tris pH 8.0, unless otherwise specified. Sequences of the oligos used for the elongation complexes assembly are shown in Fig. S1. RNA was either kinased at the 5′ end (12) or labeled at the 3′ end after elongation complex assembly by incorporation of [α-32P]GTP or [α-32P]UTP (PerkinElmer), with subsequent removal of unincorporated nucleotide through beads washing with transcription buffer. Cleavage reactions were initiated by addition of 10 mM MgCl2 (final concentration), unless otherwise specified. Misincorporation was initiated by addition of a mixture 10 mM MgCl2, 1 mM noncognate NTP, and, where specified,
100 μM NTPs complementary to the following positions (see Fig. 3a) (final concentrations). Reactions were incubated at 40 °C for times indicated in the figures and were stopped by addition of loading buffer containing formamide. Products were resolved by denaturing (8 M Urea) 23% PAGE, revealed by PhosphorImaging (GE Healthcare), and analyzed by using ImageQuant software (GE Healthcare). To determine the rate of RNA hydrolysis, a proportion of the cleaved RNA was plotted against time and fitted to a single exponential equation by using nonlinear regression (5, 12). To determine the rate of misincorporation, the proportion of complexes that had undergone misincorporation (and subsequent cleavage) was plotted against time and fitted as above. To determine the $K_{\text{M}}^{\text{Mg}^{2+}}$ for cleavage in MEC(A) and MEC(U), the reaction rates obtained in various MgCl$_2$ concentrations were fitted to the Michaelis–Menten equation (5, 12).

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