

# Multiple deprotonation paths of the nucleophile 3'-OH in the DNA synthesis reaction

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DNA synthesis by polymerases is essential for life. Deprotonation of the nucleophile 3'-OH is thought to be the obligatory first step in the DNA synthesis reaction. We have examined each entity surrounding the nucleophile 3'-OH in the reaction catalyzed by human DNA polymerase (Pol)  $\eta$  and delineated the deprotonation process by combining mutagenesis with steady-state kinetics, high-resolution structures of in crystallo reactions, and molecular dynamics simulations. The conserved S113 residue, which forms a hydrogen bond with the primer 3'-OH in the ground state, stabilizes the primer end in the active site. Mutation of S113 to alanine destabilizes primer binding and reduces the catalytic efficiency. Displacement of a water molecule that is hydrogen bonded to the 3'-OH using the 2'-OH of a ribonucleotide or 2'-F has little effect on catalysis. Moreover, combining the S113A mutation with 2'-F replacement, which removes two potential hydrogen acceptors of the 3'-OH, does not reduce the catalytic efficiency. We conclude that the proton can leave the O3' via alternative paths, supporting the hypothesis that binding of the third Mg<sup>2+</sup> initiates the reaction by breaking the  $\alpha$ - $\beta$  phosphodiester bond of an incoming deoxyribonucleoside triphosphate (dNTP).

polymerase | reactant alignment | electrostatic | three-Mg<sup>2+</sup>-ion | catalysis

NA polymerases catalyze the incorporation of deoxy-Dribonucleoside monophosphate (dNMP) into an existing primer after binding a deoxyribonucleoside triphosphate (dNTP) complementary to a templating base. The catalytic process is thought to begin with deprotonation of the primer 3'-OH by a general base, continue by nucleophilic attack of the  $\alpha$ -phosphate of the dNTP leading to a new phosphodiester bond, and finish with release of pyrophosphate after its protonation by a general acid (Fig. 1A) (1). This mechanism has long been established to require two  $Mg^{2+}$  ions (A and B) (2). However, visualizing the reaction intermediates using time-resolved X-ray crystallography reveals little movement of the protein but a third and transiently bound  $Mg^{2+}$  ion necessary for the DNA synthesis reaction (3, 4). Whether the reaction is driven by the third  $Mg^{2+}$  ion or the nucleophile 3'-OH, the identity of a general base responsible for deprotonating the primer 3'-OH remains unclear.

Crystal structures of various DNA polymerases bound to DNA and dNTP, known as ternary complexes, reveal a conserved catalytic center with two Mg<sup>2+</sup> ions coordinated by the conserved catalytic carboxylates, three phosphates of an incoming dNTP, and the primer 3'-OH (Fig. 1C) (5, 6). DNA polymerase has been likened to a right hand with the palm domain containing the catalytic residues, the finger domain closing on top of the nascent or replicating base pair, and the thumb domain binding the upstream DNA (product) duplex (7, 8). Many DNA polymerases undergo a large conformational change involving closing of the finger domain upon binding a correct dNTP (8), which was proposed to be a rate-limiting step (9). However, the rate of finger domain closing is much faster than the rate of chemical reaction (10-13). Moreover, the finger domain of the Y-family DNA polymerases is already closed even in the absence of an incoming nucleotide (14). Although intradomain motions have

been implicated in one example (15), the rate-limiting step is probably the chemical reaction (3, 4, 16, 17).

If nucleophilic attack is the first step, deprotonation of the primer 3'-OH by a general base would be essential to initiate the reaction (Fig. 1*A*). In the more than three decades since the first DNA polymerase structure was determined (7), no residues other than the conserved carboxylates that coordinate the two  $Mg^{2+}$  ions have been found to eliminate the catalytic activity when mutated. Computationally, the conserved carboxylates (1, 18) and the incoming nucleotide together with the water molecule bound to the A-site  $Mg^{2+}$  (WatA) have each been suggested to deprotonate the 3'-OH (12, 19–21). As the carboxylates and dNTPs are necessary for  $Mg^{2+}$  binding and the synthesis reaction, their role in deprotonation is nearly impossible to be experimentally tested.

A third  $Mg^{2+}$  ion (occupying the C site) has been observed to transiently bind to dNTP in the reactions catalyzed by different DNA polymerases (3, 22–24). Unlike the A- and B-site  $Mg^{2+}$ ions, which are coordinated by the catalytic carboxylates, the C-site  $Mg^{2+}$  does not contact the enzyme at all. Its low affinity ( $k_d$ ) matches the minimal  $Mg^{2+}$  concentration required for catalysis (3, 4). When two canonical  $Mg^{2+}$  ions are bound and reactants are aligned for the inline nucleophilic attack, the third  $Mg^{2+}$  ion does not bind readily (3, 4). Only when products are formed is the third  $Mg^{2+}$  ion observed to bind between the product DNA and pyrophosphate with four additional water ligands. Yet without the third  $Mg^{2+}$ , no product can form (4). Its

## Significance

To determine if there is a general base in the DNA synthesis reaction that deprotonates the nucleophile, we systematically removed potential hydrogen-bond acceptors of the nucleophile, the 3'-OH of the primer strand. We then characterized the activity of human DNA Pol  $\eta$  by kinetic, structural, and molecular dynamics simulation analyses. We found that no single or combined perturbations eliminate catalysis. Moreover, removal of two potential proton acceptors of the 3'-OH by mutating the conserved \$113 to Ala and addition of 2'-F to the primer end rescued the defects of the \$113A mutation alone. Our results support that there is no specific general base and the proton is prone to leave the O3' upon activation by the three Mg<sup>2+</sup> ions.

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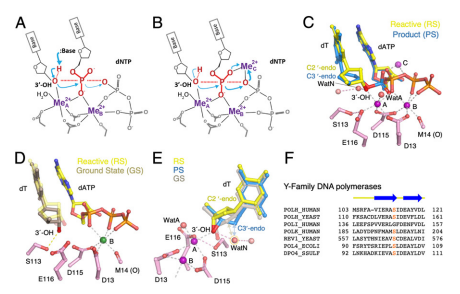
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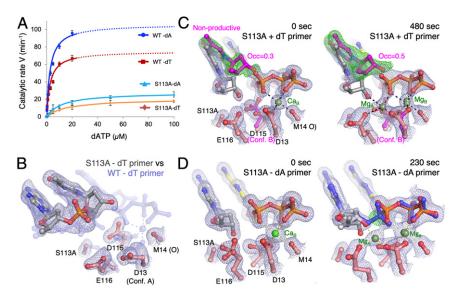
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**Fig. 1.** DNA synthesis reaction. (A) Deprotonation and activation of the primer-end 3'-OH for the nucleophilic attack is hypothesized to be the first step. (B) In the three– $Mg^{2+}$ -ion catalysis, the first step appears to break the phosphodiester bond between the α- and β-phosphates of the dNTP. (C) Structural superposition of the reaction-ready (RS) and product state of the WT enzyme in extending the dT primer. The two  $Mg^{2+}$  ions (A and B sites) are bound and the 3'-OH is hydrogen-bonded to the transient WatN, which in turn is linked to the bulk solvent via another water molecule. The third  $Mg^{2+}$  (C site) is observed only with the products. (D) In the ground state with one  $Ca^{2+}$  (green sphere), the 3'-OH is hydrogen-bonded with S113 and not aligned for the inline nucleophilic attack in the absence of the A-site  $Mg^{2+}$ . The aligned 3'-OH in the RS is shown as semitransparent sticks as a reference. (E) The transient WatN (circled in orange dashes) appears only in the RS and would clash with the C2' in the GS (wheat) and PS (blue) as indicated by double arrowheads. (F) Sequence alignment of active-site residues around S113 (highlighted in red) of human Pol η, *Saccharomyces cerevisiae* Pol η (yeast), human Pol ι, human Pol κ, S. cerevisiae Rev1, *Escherichia coli* Pol IV, and *Sulfolobus solfataricus* Dpo4. The conserved secondary structures are shown above the sequences.

binding is thermal energy (temperature)-dependent, and concurrent with the product formation. The third Mg<sup>2+</sup> ion may drive dNMP incorporation by breaking the  $\alpha$ - $\beta$  phosphodiester bond of dNTP and pushing the  $\alpha$ -phosphate toward the 3'-OH for the new bond formation (4) (Fig. 1*B*). With the C-site Mg<sup>2+</sup>, deprotonating the 3'-OH is likely favored and does not need a strong general base.

The in crystallo analysis of the DNA synthesis reaction catalyzed by human polymerase (Pol)  $\eta$  reveals three reaction states and two potential candidates to deprotonate the nucleophile.



**Fig. 2.** CS113 anchors the primer end. (A) Steady-state kinetic profiles of WT and S113A mutant Pol  $\eta$  extending primers terminating in dA or dT. Kinetic parameters for each profile are listed in Table 1. Error bars represent standard deviation of triplicate measurements. (*B*) Superimposed ternary-complex structures of S113A (solid multicolor) and WT (semitransparent blue) Pol  $\eta$  both with 0.1 mM dAMPNPP. In the S113A structure, carbon atoms of the DNA are shown in gray and of Pol  $\eta$  in salmon pink. 2Fo–Fc map of the S113A structure (1 $\sigma$ ; gray) is shown as meshes. (C) 2Fo–Fc (1 $\sigma$ ; gray) and Fo–Fc maps (3 $\sigma$ ; green; calculated by omitting the nonproductive primer conformation) are superimposed onto the 0- (*Left*) and 480-s (*Right*) structures of the S113A Pol  $\eta$ –dT-primer complex. The nonproductive conformations of the primer end and D115 are shown in magenta. (*D*) 2Fo–Fc maps (contoured at 1 $\sigma$ ) are superimposed on the 0-s S113A (*Left*) and 230-s S113A Pol  $\eta$ –dA-primer structure (*Right*). By 230 s, formation of the product (shown as blue sticks) and the new bond between the primer and dATP are evident in the green Fo–Fc map, which is calculated without the product and contoured at 3 $\sigma$ .

Initial dNTP binding with one divalent cation only (B site) leads to the ground state (GS) (Fig. 1D). Binding of the second  $Mg^{2+}$ converts the GS to the reactive state (RS), in which the 3'-OH is aligned for the inline nucleophilic attack, and appearance of the third  $Mg^{2+}$  (C site) is coupled with the product formation (product state; PS) (3, 4) (Fig. 1 C and E). In the GS, the 3'-OH is within 2.7 Å and hydrogen-bonded to the hydroxyl group of S113 (3). Although S113 is highly conserved among the Y-family polymerases (Fig. 1F) and may facilitate deprotonation of the 3'-OH, removal of the hydroxyl group by mutating S113 to Ala in human Pol n reduces the catalytic efficiency but does not eliminate catalysis (3). In the RS, the 3'-OH is 2.7 Å from a water molecule, which is termed WatN (N for nucleophile) and hypothesized to shuttle the proton off the 3'-OH to the bulk solvent (3). However, preliminary kinetic analysis showed that displacement of the water molecule by the 2'-OH of a ribonucleotide at the primer end leaves catalysis of DNA synthesis unaltered (3).

Here we investigate in detail how the 3'-OH is deprotonated in the reaction catalyzed by human Pol  $\eta$ . Combining S113A mutant Pol  $\eta$  (S113A) and modified nucleotides at the primer 3' end, we measured how the altered environment of the 3'-OH perturbed the steady-state kinetics and reaction process in crystallo. In addition, static structures and dynamics simulation are employed to explain observed kinetic parameters. In contrast to the hypothesis of a specific general base, we find that the proton of the 3'-OH can depart via multiple paths.

# Results

**S113A Exhibits Reduced**  $k_{cat}$  and Increased  $K_{M}$ . In the Pol  $\eta$  active site, S113 is hydrogen-bonded with the 3'-OH at the primer end in the GS and, when the 3'-OH shifts toward the incoming dNTP in the RS, S113 is hydrogen-bonded with the catalytic carboxylate E116 that chelates the A-site Mg<sup>2+</sup> ion (Fig. 1 *C* and *D*). The serine-to-alanine mutation was chosen to remove the functional group (OH), while keeping the protein structure minimally changed (3). An equivalent of S113 exists in the A- and B-family polymerases as His and Thr, respectively (*SI Appendix*, Fig. S1). As S113, the conserved Thr in the B-family polymerases can hydrogen-bond with the primer 3'-OH (25, 26), and the conserved His seems to insulate the active site and thereby stabilizes the primer 3' end (27–29). Although not essential for the catalysis, these equivalent residues improve catalytic efficiencies (30, 31).

The steady-state kinetic parameters of S113A were measured for nucleotide incorporation in comparison with wild-type (WT) Pol  $\eta$ . Different nucleotides at the primer 3' end have been shown to affect primer alignment and nucleotide incorporation (32). To include DNA sequence variations, both dA and dT at the primer 3' end were examined. Previously only a dAterminated primer (dA-primer) had been studied (3). On average, S113A lost ~20-fold catalytic efficiency ( $k_{cat}/K_M$ ) compared with WT (Fig. 2A and Table 1), with a 3- to 4-fold reduction in  $k_{\text{cat}}$  and 4- to 6-fold increase in  $K_{\text{M}}$ . The catalytic efficiency of the primer ending with dT was a little lower than with dA even with WT Pol  $\eta$  (1.4-fold), and the disparity became more pronounced with S113A (1.8-fold). Interestingly, WT Pol  $\eta$  had a similar  $K_{\rm M}$ for different primer ends but, with S113A,  $K_{\rm M}$  for dATP was higher with the dT- than the dA-primer. The kinetic data indicate that S113A may have a reduced ability to bind an incoming nucleotide when dT is at the primer end.

**S113A Fails to Dock the 3' dT in the Active Site.** To understand how S113 assists dNTP binding and facilitates catalysis, we first compared the ternary-complex structures of WT and S113A Pol  $\eta$  with a dT-terminated primer (dT-primer). To stop the polymerization reaction, the incoming nucleotide deoxyadenosine triphosphate (dATP) was replaced by a nonreactive analog, 2'-

deoxyadenosine-5'-[( $\alpha$ , $\beta$ )-imido]triphosphate (dAMPNPP). The structure of S113A with dT at the primer end was determined at 2.27-Å resolution (*SI Appendix, Methods* and Table S1). WT and S113A Pol  $\eta$  complexed with the same DNA substrate and 1 mM dAMPNPP were indistinguishable, and the rmsd of over 431 pairs of C $\alpha$  atoms was only 0.22 Å (*SI Appendix*, Fig. S2A). Nearly superimposable onto the RS of WT Pol  $\eta$  in catalysis, the static structures obtained with dAMPNPP have a slightly tighter packing around the 3'-OH (*SI Appendix*, Fig. S2B).

Because the  $K_{\rm M}$  for incoming nucleotides was higher with S113A, we determined crystal structures of WT and S113A Pol  $\eta$  with 0.1 mM instead of 1 mM dAMPNPP, as 0.1 mM is similar to physiological dNTP concentrations. The ternary complex of WT Pol  $\eta$  formed readily with 0.1 mM dAMPNPP, but S113A failed to make a ternary complex and remained as a binary Pol  $\eta$ –DNA complex (Fig. 2*B*). Without the hydrogen bond between S113 and the 3'-OH at the primer end, the last nucleotide of the primer shifted toward the dNTP-binding site. Although hydrogen-bonded with the templating base dA, the 3' terminal dT was no longer base-stacked with the rest of the primer strand (Fig. 2*B*).

We further compared in crystallo catalytic processes of S113A with WT Pol  $\eta$ . The in crystallo reactions were carried out as previously reported (3). Briefly, crystals were grown with a normal template/primer DNA and 1 mM dATP, but reaction was prevented by substituting Mg<sup>2+</sup> with Ca<sup>2+</sup>. After fully grown, the crystals were "washed" in a stabilization buffer containing no dATP or Ca<sup>2+</sup> first, and the reaction was initiated by transferring the crystals to a buffer containing 1 mM Mg<sup>2+</sup>. By freezing a series of crystals at various reaction time points in liquid nitrogen, the catalytic process was visualized by X-ray diffraction of these crystals (33).

In the GS (before soaking crystals in the Mg<sup>2+</sup> buffer), the active site of WT Pol  $\eta$  contains a fully occupied dATP and one Ca<sup>2+</sup> occupying the B site coordinated by all three phosphates of dATP, while the 3'-OH at the primer end is misaligned for nucleophilic attack (Fig. 1D) (3). In contrast, before soaking in the  $Mg^{2+}$  buffer, only 70% of the S113A active site contained an incoming dATP (Fig. 2C), and the 3'-OH appeared to assume an RS-like conformation instead of the GS in the absence of the A-site Mg<sup>2+</sup> (Fig. 2C). The remaining 30% of Pol  $\eta$ -DNA complexes were devoid of dATP because the primer ends shifted into the incoming nucleotide-binding site (Fig. 2C), as observed in the S113A-DNA binary complex formed with 0.1 mM AMPNPP (Fig. 2*B*). After exposure to  $Mg^{2+}$  for 60 s, WT Pol η starts to form products and peaks at 230 s (3). With S113A, despite 70% RS-like ternary complexes existing prior to  $Mg^{2+}$ exposure, no product was observed by 480 s, and instead 50% of the primer end became misaligned as dATP was partially diffused out (Fig. 2C). Removal of the hydroxyl group of S113 in Pol  $\eta$  resulted in defective docking of the primer 3' end in the GS and an inactive RS-like state upon Mg<sup>2+</sup> binding, which corresponded to the increased  $K_{\rm M}$  and reduced catalytic efficiency (Table 1).

**Defective Primer Docking Is Partially Rescued by the 3'-End dA.** The defective docking of the primer 3' end in the active site and absence of product formation of S113A are partially rescued by dA at the primer 3' end. The large adenine base favors base stacking (34), and dAMP is often incorporated by DNA polymerases without a template, a phenomenon known as the "A rule" (35). The ternary-complex structure of S113A, DNA with dA at the primer end, and dATP was identical to the GS structures of WT Pol  $\eta$  with the dT-primer (Fig. 2D). The non-productive primer conformation was eliminated by stable base stacking of dA. S113 may be more important for stabilizing a pyrimidine than purine due to the different intrinsic propensities of base stacking. By forming the hydrogen bond with the 3'-OH,

Table 1.	Kinetic data	of WT and	S113A Pol η
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Experi	ment	$k_{\rm cat}$ , min <sup>-1</sup>	<i>Κ</i> <sub>Μ</sub> , μΜ	$\begin{array}{c} \text{Catalytic efficiency,} \\ \mu M^{-1} \cdot \text{min}^{-1} \end{array}$	Relative efficiency
WT	dA	106 ± 5.0	2.7 ± 0.4	39.3	1
	rA	72.8 ± 2.3	$2.0 \pm 0.3$	36.4	0.93
	2′F-	95.1 ± 1.0	5.2 ± 0.2	18.3	0.47
	dA				
S113A	dA	28 ± 1.0	12.3 ± 1.7	2.3	1
	rA	79.5 ± 3.1	8.4 ± 1.2	9.5	4.16
	2′F-	109.2 ± 1.8	$21.4\pm0.9$	5.1	2.24
	dA				
WT	dT	75.0 ± 6.0	2.7 ± 0.7	27.8	1
	rU	38.1 ± 1.0	$4.3 \pm 0.4$	8.9	0.32
S113A	dT	$20.7 \pm 0.7$	16.3 ± 1.5	1.3	1
	rU	43.0 ± 1.6	7.7 ± 0.9	5.6	4.40

S113 allows WT Pol  $\eta$  similar  $K_M$  for correct incoming dNTPs regardless of whether a purine or pyrimidine is at the primer end (Table 1).

S113A was able to extend a dA-primer in crystallo (Fig. 2D), but more slowly than WT protein (Fig. 3). The Fo-Fc maps of the reaction intermediates calculated by omitting the 3'-OH and A-site  $Mg^{2+}$  revealed a persistent GS (50% of the ternary complexes up to 300 s) and an absence of the C2'-endo RS (Figs. 2D and 3A). Because previous in crystallo reactions were carried out with dT-primers (Fig. 1 C-E), we analyzed in crystallo extension of the dA-primers by WT Pol  $\eta$  (Fig. 3B). After soaking in the Mg<sup>2+</sup> reaction buffer, products began to form as early as 40 s and accumulated to over 65% by 230 s, which is similar to the catalytic extent of dT-primers (3, 4). During the reaction course the 3'-end deoxyribose was transformed from C2'-endo in the substrate state to C3'-endo in the product state (Fig. 1*E*). But the distinct RS with C2'-endo and the 3'-OH aligned and hydrogen-bonded to WatN (Fig. 1 C and E) was absent, and the substrate remained in the GS with the O3' hydrogen-bonded to S113 (Fig. 3B). As a result, the occupancy of the A-site  $Mg^{2+}$  was nearly constant (0.5) instead of reducing from 0.9 at 40 s to 0.5 at 300 s with the dT-primer (3, 4). We suspect that the dA-primer favors the GS and the RS is too transient to be observed.

While the GS persisted in the S113A–dA ternary complex after soaking in the  $Mg^{2+}$  reaction buffer, 30 (40 s) to 50% (300 s) of primer ends underwent conformational change from C2'-endo (GS) to C3'-endo, but only half of them were converted to product. In the C3'-endo conformation, the primer end was 3.1 to 3.2 Å from the  $\alpha$ -phosphorus of dATP and well-aligned for the

nucleophilic attack (Fig. 2D). However, the new phosphodiester bond was inexplicably slow to form (Figs. 2D and 3A), analogous to the inactive RS-like S113A–dT ternary complexes (Fig. 2C). The occupancy of the A-site  $Mg^{2+}$  was slightly higher with S113A than WT Pol  $\eta$  throughout the reaction, and the occupancy of the C-site  $Mg^{2+}$  was similar to the product amount as usual (4). In solution, measurement of  $Mg^{2+}$ -concentration dependence also showed that S113A and WT are similar (*SI Appendix*, Fig. S3). The in crystallo analyses of S113A indicate that the absence of this conserved hydroxyl group that anchors the 3'-OH not only changes the geometry but also the reactivity of the primer end, even when seemingly well-aligned.

Slow Catalysis of S113A Is Rescued by Ribonucleotide at the Primer 3' End. To differentiate conformation and reactivity, we examined Pol  $\eta$  catalysis on RNA-terminated primers. Deoxyribonucleotides predominantly adopt a C2'-endo pucker, while ribonucleotides exist in the C3'-endo conformation (36). We replaced dA with rA (adenylate) and dT with rU (uridylate) at the primer end and repeated kinetic measurements. With WT Pol  $\eta$  these RNAterminated primers were extended at a slightly reduced  $k_{cat}$ (Table 1). But with S113A, RNA-terminated primers restored the catalytic rate to the level of WT enzyme on both rU and rA primers (Fig. 44).

To visualize the effects of ribonucleotides on the active-site configuration, we determined the crystal structure of S113A bound to an rA-terminated primer (rA-primer) with Mg<sup>2+</sup> and dAMPNPP (SI Appendix, Table S1). The rA at the primer end adopted a C3'-endo pucker as expected, and the rest of the active site remained in the active RS conformation including the O3' location (Fig. 4B). The reaction intermediates of S113A extending the rA-primer were also examined. Because of the C3'-endo conformation with rA at the primer end, there was no GS and the O3' was aligned for nucleophilic attack prior to soaking in the  $Mg^{2+}$  buffer (Fig. 4 C and D). In addition, the phosphate of the rA adopted a singular conformation unlike those of dA and dT, which exhibited two alternate conformations (SI Appendix, Fig. S4). The in crystallo reaction took place at a rate matching the WT enzyme with the dA-primer. The occupancy of the incoming dATP was reduced at 0.85, and pyrophosphate product (PPi) was partially released, which was not observed with WT Pol η. Concurrently, both A- and C-site Mg<sup>2+</sup> occupancies were lower than with fully occupied dATP and PPi. Nevertheless, the catalytic activity of S113A on the rA-primers approached the level of WT Pol n on DNA (Fig. 4D and Table 1). The similar catalytic efficiency of WT Pol  $\eta$  with a dA- or rA-primer indicates that both C3'-endo and C2'-endo sugar

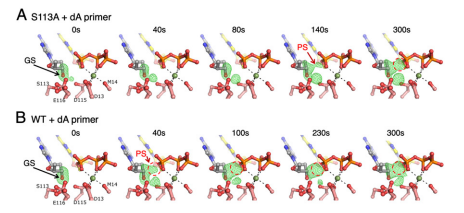
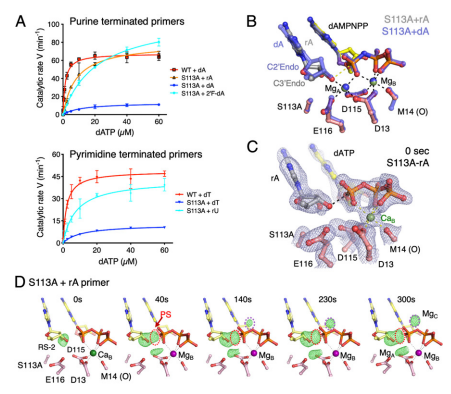


Fig. 3. GS-to-PS transition of a dA-primer. (A) Time course of the S113A mutant Pol  $\eta$ -catalyzed extension reaction of a dA-primer. (B) Time course of the WT Pol  $\eta$ -catalyzed extension of a dA-primer. Fo<sub>(S113A or WT)</sub>-Fc<sub>(WT, GS)</sub> maps (3.5 $\sigma$ ) calculated with C3' and O3' omitted are superimposed onto the GS structure of the WT Pol  $\eta$ -dA-dATP complex (t = 0 s). GS and PS positions are indicated with arrows. Product density is circled in red.



**Fig. 4.** RNA-terminated primer rescues the defects of S113A Pol  $\eta$ . (A) Steady-state kinetic profiles of WT and S113A mutant Pol  $\eta$  extending primers terminating in dA, rA, or 2'-fluoro-dA (*Top*) and dT or rU (*Bottom*). Kinetic parameters for each profile are listed in Table 1. Error bars represent standard deviation of triplicate measurements. (*B*) Superimposed structures of S113A Pol  $\eta$  bound to the dA- or rA-primer and dAMPNPP. The dA-primer structure is shown in semitransparent blue sticks and the rA-primer structure is shown in colored sticks (yellow dAMPNPP, salmon protein residues, and gray primer). The sugar pucker of the terminal primer base is labeled. (C) The 0-s structure of S113A Pol  $\eta$  bound to the rA-primer with the 2Fo–Fc map (1 $\sigma$ ) superimposed. The rA adopts the C3'-endo conformation and is reaction-ready (RS-2) prior to Mg<sup>2+</sup> binding in the A site. (*D*) The reaction time course of S113A extending the rA-primer extension by S113A. Fo<sub>(S113A-rA, 0-300 s)</sub>–Fc<sub>(WT, GS)</sub> maps (3.5 $\sigma$ ) with C3' and O3' omitted are superimposed onto the 0-s S113A Pol  $\eta$ –rA-primer complex at 0 s. The electron densities for the product and Mg<sup>2+</sup><sub>C</sub> are circled in red and purple, respectively.

puckers can participate in the nucleophilic attack efficiently. However, C3'-endo at the primer end renders the A-site  $Mg^{2+}$  unnecessary for the 3'-OH to be in the reaction-ready state. In the absence of hydroxyl groups of S113 and WatN surrounding the O3', the good catalytic efficiency of S113A with the rA-primer suggests that alternative proton-transfer pathways exist.

Molecular Dynamics Analysis of WT and S113A Pol  $\eta$ . To find out why the well-aligned 3'-OH of dA- and dT-primers is slow to react with S113A, we carried out classical molecular dynamics (MD) simulations of either the C2'- or C3'-endo RS form of WT or S113A Pol η on the scale of 50 to 100 ns (*SI Appendix, Methods*). As the C-site  $Mg^{2+}$  is necessary for catalysis (4), it is placed near the  $\alpha$ -phosphate (P $\alpha$ ) of dATP initially. In a large number of independent MD simulations of S113A, the C-site Mg<sup>2+</sup> appeared easily perturbed and moved between the  $\alpha$ - and  $\gamma$ -phosphates of dNTP complexed with S113A, in contrast to the stable binding to the  $\alpha$ -phosphate (P $\alpha$ ) with WT Pol  $\eta$  prior to the product formation (Fig. 5 A and B). Only one out of seven simulations of S113A with the dA-primer led to the same binding mode as the WT enzyme. Coupled with the changes of the C-site Mg<sup>2+</sup> binding mode, positions of the primer end and the incoming nucleotide were also perturbed, leading to a longer O3'to-P $\alpha$  distance (Fig. 5C). The observations are qualitatively consistent with the perturbed substrate binding by S113A. These results suggest that the C-site Mg<sup>2+</sup> is important for not only product formation but also the favorable nucleophilic attack configuration, thus playing an explicit role in catalysis.

With the C-site  $Mg^{2+}$  bound to the  $\alpha$ -phosphate, the activesite properties of S113A with the dA-primer were generally similar to the WT enzyme. However, subtle differences are observed in the distributions of O3'-Mg<sup>2+</sup><sub>A</sub> and O3'-WatA, which exhibit peak distributions at slightly longer distances in S113A than in the WT enzyme (Fig. 5D). Meanwhile, removal of the hydroxyl group of S113 exposes the active site and increases the frequency of water molecules (Ow) surrounding the 3'-OH (Fig. 5D). The S113A mutation seems to perturb interactions between the primer end and both A- and C-site Mg<sup>2+</sup>, leading to either less favorable nucleophilic attack geometries or weaker polarization of the 3'-OH by the A-site Mg<sup>2+</sup>, and thus a slightly lower catalytic activity.

To understand the observed rescuing effect of the 2'-OH at the primer end, we also carried out MD simulations for S113A with the rA-primer. In contrast to the unstable C-site Mg<sup>2+</sup> and increased distance between O3' and P $\alpha$  in the simulations of the S113A–dA complex, S113A-rA exhibited characteristics similar to WT Pol  $\eta$  (Fig. 5*C*). The results from classic MD analyses have established foundations for future in-depth analysis of catalysis of different enzyme and substrate complexes by quantum mechanics/molecular mechanics (QM/MM) studies.

**Deprotonation Path Is Flexible.** In addition to the altered sugarpucker conformation, the 2'-OH of the rA at the primer 3' end also displaced the transient WatN, which is hydrogen-bonded to the 3'-OH and potentially facilitates deprotonation of the nucleophile. The O2' is competent to accept a proton from the O3' and release it to solvent. Consistent with such a proton-transfer

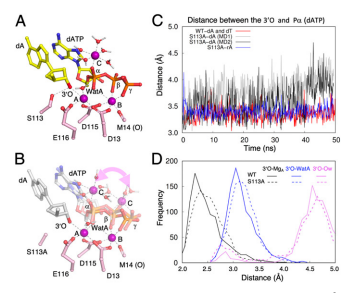


Fig. 5. MD simulation of the WT and S113A active sites. (A) The C-site Mg<sup>2+</sup> is stably bound to the  $\alpha\text{-phosphate}$  of dATP in the WT active site. Three  $M\tilde{g}^{2+}$ ions (A, B, and C) are shown as purple spheres. The primer-end dA, dATP, catalytic residues, and inner-shell water molecules of Mg<sup>2+</sup> ions are shown as sticks, and Mg<sup>2+</sup> coordinations are shown as dashed lines. (B) In multiple independent MD simulations of the S113A mutant, the C-site Mg<sup>2+</sup> binds either the  $\alpha$ - or  $\beta$ - $\gamma$  phosphates of dATP. The alternative binding mode is indicated by the curved double arrowhead. The incoming dATP also wobbles during simulations. Two representative intermediate states are shown. (C) During dynamics simulation, the distance between the primer O3' and P $\alpha$  of dATP is stable in the WT Pol  $\eta$  active site (red trace) but is varied and longer in the S113A active site. Two traces of the S113A simulation are shown in gray and black. In the S113A-rA simulation, the two reactants (O3' and P $\alpha$  of dATP) are as stable (blue trace) as with WT Pol  $\eta$ . (D) Comparison of distances between the O3' and  $\mathrm{Mg}^{2+}{}_{\mathrm{A}}$ , WatA, and surrounding water molecules in the WT (solid lines) and S113A mutant Pol  $\eta$  (dashed lines).

relay, a water molecule was observed to hydrogen-bond with the O2' in the ternary complex with an RNA-terminated primer and dAMPNPP (Fig. 64). The near-normal primer extension on the RNA-terminated primers by WT and S113A Pol  $\eta$  indicates that any role WatN may have in the nucleophile deprotonation can be replaced by the O2'.

To test if the 2'-OH of a ribonucleotide is required for deprotonation of the 3'-OH, we replaced rA with 2'-fluorodATP (2'F-dA) at the primer 3' end. Unlike a hydroxyl group, fluorine attached to C2' would displace the transient water but is unlikely to serve as a proton acceptor. To our surprise, the 2'F-dA-terminated primer was extended by WT enzyme as efficiently as normal DNA. The 2'F-dA even rescued the defects of S113A and improved its  $k_{cat}$  and  $k_{eff}$  to near-WT levels (Fig. 4A and Table 1). In the absence of the two most obvious suspects, S113 and the transient WatN (displaced by 2'-F), another path of 3'-OH deprotonation is needed to facilitate the DNA synthesis reaction.

To identify an alternative proton acceptor or relay path, the structure of WT Pol  $\eta$  bound to the 2'F-dA-terminated primer with Mg<sup>2+</sup> and dAMPNPP in the RS-like state was determined at 1.31-Å resolution (*SI Appendix*, Table S1). At this resolution, weakly bound groups and solvent were observable. Six potential proton acceptors of the O3' were found, all of which were within hydrogen-bond distance (3.1 Å). They are S113 (OH), D115 (Oδ), E116 (Oε),  $\alpha$ O-proRp, and O5' of dAMPNPP, and a water molecule that chelates the A-site Mg<sup>2+</sup> (WatA) (Figs. 1 *C*-*E* and 6*C*). As the  $\alpha$ O-proRp of dAMPNPP coordinates the A-site Mg<sup>2+</sup> and has an extremely low intrinsic pK<sub>a</sub> value, it is an

unlikely proton acceptor (37). The carboxylates D115 and E116 also directly coordinate the A-site  $Mg^{2+}$  and thus also have lowered pK<sub>a</sub> values. Although the residue equivalent to Pol  $\eta$ 's E116 in DNA Pol  $\beta$  has been suggested to be the proton acceptor during polymerase catalysis (18, 22), the loss of catalytic activity upon mutating the carboxylate can be due to defective A-site  $Mg^{2+}$  binding instead.

Aside from the three  $Mg^{2+}$  ligands and S113, only two candidates remain. Firstly, the O5' of an incoming nucleotide may take the proton and pass it to the bulk solvent. Lastly, WatA that coordinates the A-site  $Mg^{2+}$  ion is proximal to the O3' and the only nonmacromolecule chelators of the A- and B-site Mg<sup>2+</sup> ions. WatA has been previously proposed to directly accept the O3' proton in the Dpo4 and Pol ß catalysis using QM/MM simulations (19–21). WatA is usually more than 3.3 Å away from the 3'-OH in all in crystallo reaction time courses but is 3.05 Å away in the static 2'F-dA and rA structures (Fig. 6 C and D) and  $\sim$ 3.1 Å in MD simulations (Fig. 5D). In these two comparable structures, only S113 (OH) and the O5' of dAMPNPP became closer to the O3' with 2'F-dA than with rA, while the other four maintained the same distance (Fig. 6D). The last possible candidates are two water molecules hydrogen-bonded to the 2'-F (Fig. 6C). Although fluorine is not known to be a proton acceptor, it may be a stepping stone to pass the proton from the 3'-OH to the bulk solvent.

### Conclusion

Our combined structural, kinetic, and computational analyses suggest that human Pol  $\eta$  does not rely on a specific proton

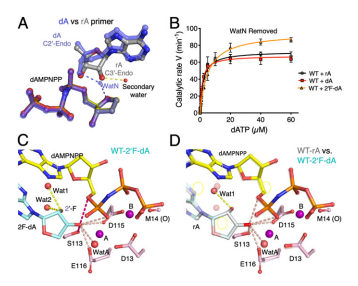


Fig. 6. Deprotonation route is flexible. (A) Structural superposition of the dAMPNPP-bound WT Pol  $\eta$  with dA- (semitransparent blue sticks) and rA-(grav and vellow sticks) primers. The transient water associated with the 3'-OH of dA is shown. The secondary water of the rA structure is shown as a red sphere. Hydrogen bonds between the terminal primer sugar hydroxyl groups and the water molecules are shown as dashes. (B) Steady-state kinetic profiles of WT Pol n extending primers terminating in dA, rA, or 2'F-dA. Kinetic parameters for each profile are listed in Table 1. Error bars represent standard deviation of triplicate measurements. (C) Possible 3'-OH deprotonation routes. The structure of WT Pol n bound to the 2'F-dA-terminated primer is shown. The six atoms within 3.1 Å of the 3'-OH of 2'F-dA are marked by pink dashed lines. Two (the hydroxyl of \$113 and O5' of dAMPNPP) are highlighted in hot pink because these distances are 0.1 to 0.15 Å shorter in the 2' F-dA complex than those in the rA complex. (D) The structure of the rAprimer is superimposed on 2'F-dA (semitransparent). The two are nearly identical. Two water molecules (Wat1 and Wat2) are within hydrogen-bond distance of the 2'-F. Only Wat1 is bound to the 2'-OH of rA.

pathway for catalysis. The conserved S113 plays an important role in the primer 3'-end docking and influencing the A- and C-site Mg<sup>2+</sup> and incoming dNTP binding. The hydroxyl group of S113, WatN transiently bound to the 3'-OH, WatA of the A-site  $Mg^{2+}$ , and O5' of the incoming dNTP each may be a proton acceptor and shuttle it out to the bulk solvent. In agreement with previous experimental and computational studies, neither the proton acceptor nor the departure path of the nucleophile deprotonation is fixed. When a convenient or preferred path is eliminated, multiple alternatives, including water molecules (38), a protein side chain, or incoming dNTP (12), may aid the proton to depart from the O3'. Such a flexible deprotonation mechanism is consistent with the three-metal-ion catalysis, by which the C-site Mg<sup>2+</sup> alters the electrostatic environment around the  $\alpha$ -phosphate and 3'-OH and effectively lowers the energy barrier to product formation (39) (Fig. 1B). Rather than requiring a specific general base to deprotonate the 3'-OH and initiate the nucleophilic attack, the  $Mg^{2+}$  likely drives the reaction forward and promotes dissociation of the proton.

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# **Materials and Methods**

Pol  $\eta$  1 to 431 amino acids was cloned, expressed, and purified as described previously (40). Oligos were obtained from Integrated DNA Technologies or the Food and Drug Administration Facility for Biotechnology Resources. The oligo sequences are listed in *SI Appendix*, Table S2. Kinetic measurements, X-ray crystallographic analyses, and MD simulations were carried out according to established protocols. Please see details in *SI Appendix*.

**Data Availability.** The structure coordinates and structure factors reported in this article have been deposited in the Protein Data Bank (ID codes 7M7L–7M7U, 7M7Y, 7M7Z, 7M8A–7M8D, 7M80–7M89; *SI Appendix*, Table S1).

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