DNA polymerase (pol) ι is a Y-family polymerase involved in translesion synthesis, exhibiting higher catalytic activity with Mn$^{2+}$ than Mg$^{2+}$. The human germline R96G variant impairs both Mn$^{2+}$-dependent and Mg$^{2+}$-dependent activities of pol ι, whereas the Δ1–25 variant selectively enhances its Mg$^{2+}$-dependent activity. We analyzed pre-steady-state kinetic and structural effects of these two metal ions and genetic variations on pol ι using pol ι core (residues 1–445) proteins. The presence of Mn$^{2+}$ (0.15 mM) instead of Mg$^{2+}$ (2 mM) caused a 770-fold increase in efficiency ($k_{pol}/K_{dACTP}$) of pol ι for dCTP insertion opposite G, mainly due to a 450-fold decrease in $K_{dACTP}$. The R96G and Δ1–25 variants displayed a 53-fold decrease and a 3-fold increase, respectively, in $k_{pol}/K_{dACTP}$ for dCTP insertion opposite G with Mg$^{2+}$ when compared with wild type, substantially attenuated by substitution with Mn$^{2+}$. Crystal structures of pol ι ternary complexes, including the primer terminus 3′-OH and a non-hydrolyzable dCTP analogue opposite G with the active-site Mg$^{2+}$ or Mn$^{2+}$, revealed that Mn$^{2+}$ achieve more optimal octahedral coordination geometry than Mg$^{2+}$, with lower values in average coordination distance geometry in the catalytic metal A-site. Crystal structures of R96G revealed the loss of three H-bonds of residues Gly-96 and Tyr-93 with an incoming dNTP, due to the lack of an arginine, as well as a destabilized Tyr-93 side chain secondary to the loss of a cation–π interaction between both side chains. These results provide a mechanistic basis for alteration in pol ι catalytic function with coordinating metals and genetic variation.

Genomic DNA is continuously attacked by a variety of endogenous and exogenous agents in cells, and the persistent unrepaired lesions can lead to genomic mutations and related diseases such as cancer. DNA polymerases (pols), as well as DNA repair enzymes, are key enzymes for maintaining or altering genomic integrity against DNA lesions during various DNA transactions in organisms. The DNA replicative mechanisms linked to DNA damage and repair are believed to contribute to producing various mutational signatures in human cancer genomes (1). At least 17 different human DNA polymerases have been identified to date, which differ in their functions in DNA replication, repair, recombination, and damage tolerance (2,3).

Y-family DNA polymerases, including pols η, ι, κ, and REV1, are specialized in replicating through DNA lesions, so-called translesion DNA synthesis (TLS). These polymerases have low fidelity with undamaged DNA templates but have spacious and solvent-accessible active sites to allow the accommodation and replicative bypass of bulky and distorted DNA lesions (4). Individual Y-family polymerases play error-free or error-prone roles in TLS, depending on DNA lesion types in cells (5). At bulky carcinogen-derived N$^2$-G DNA lesions (e.g. benzo[a]pyrene-diol epoxide-G), both pol κ and REV1 catalyze error-free TLS, but both pols η and ι catalyze error-prone TLS (6–14). By contrast, at UV-induced cyclobutane thymine dimers (T-T), only pol η (but not the other Y-family pols) can catalyze error-free TLS (15,16). Therefore, the overall balance toward error-free TLS with all working polymerases at various DNA lesions might be crucial in preventing mutations from numerous genotoxic agents. Recently, we reported that catalytic (either hypoxic or hyperactive) alterations are found in a considerable number of human germline non-synonymous variants of Y-family pols κ, ι, and REV1 (17–19), which might potentially influence on the overall TLS capacity in the affected individuals. pol ι is exceptionally error-prone in DNA synthesis among polymerases, particularly opposite template bases G and T, due to its uniquely restricted active-site and related non-Watson-Crick base pairing (20–22). pol ι is able to catalyze nucleotide insertion opposite a variety of DNA lesions, including N$^2$- and O$^6$-alkyl and aralkyl G adducts, 8-oxo-7,8-dihydroG (8-oxoG),
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TABLE 1
Pre-steady-state kinetic parameters for dCTP incorporation opposite G by pol λ(1–445), pol λ(26–445), and R96G pol λ(1–445) in the presence of either Mg²⁺ or Mn²⁺.

| Metal       | pol λ (1–445) | kₚ₀ | Kₑ,dCTP | kₚ₀/Kₑ,dCTP | Relative efficiency* |
|-------------|---------------|-----|---------|-------------|----------------------|
| MgCl₂ (2 mM) | 0.43 ± 0.02   | 670 ± 100 | 6.4 × 10⁴ | 1           |
| MgCl₂ (2 mM) | 0.77 ± 0.024  | 390 ± 40  | 2.0 × 10⁵ | 3.1          |
| R96G pol λ (1–445) | 0.063 ± 0.007 | 5200 ± 1300 | 1.2 × 10⁷ | 0.019        |
| MnCl₂ (150 μM) | 0.74 ± 0.06   | 1.5 ± 0.4  | 4.9 × 10⁵ | 1           |
| MnCl₂ (150 μM) | 0.79 ± 0.03   | 1.8 ± 0.2  | 4.4 × 10⁵ | 0.90         |
| R96G pol λ (1–445) | 0.077 ± 0.002 | 1.4 ± 0.1  | 5.5 × 10⁴ | 0.11         |

* Relative efficiency, calculated as the ratio of the kₚ₀/Kₑ,dCTP of each pol λ for dCTP insertion opposite G to the kₚ₀/Kₑ,dCTP of pol λ (1–445) for dCTP insertion opposite G.

pyrimidine dimers, and abasic sites, but mediates largely mis-coding TLS (albeit occasionally accurate) with varied nucleotide selectivity depending on lesion type (13, 23, 24). Both C and T are inserted opposite template N²- and O⁶-alkyl and aralkyl G adducts by pol λ (13, 25). C is only slightly favored over G opposite template 8-oxoG by pol λ, A is favored opposite the 3’ T of (6-4) T-T photoproducts, and both G and T are favored opposite 5’ T of (6-4) T-T photoproducts and abasic sites (23, 24). A possible implication of pol λ in mutation and cancer has been suggested by substantial evidence from several knock-out mouse studies (26–28), as well as from multiple studies verifying frequent pol λ dysregulation in various types of human cancers (29–32). In this respect, the appropriate catalytic function of pol λ in cells might be required for preventing cancer. One of the distinctive catalytic properties of pol λ is the metal ion preference. Unlike other Y-family polymerases, pol λ is known to prefer Mn²⁺ over Mg²⁺ as the metal ion cofactor for catalysis (33, 34), but the structural mechanistic details remain speculative. In addition, substantial alterations were reported in metal-dependent DNA polymerase activity from two rare human germline pol λ variants, i.e. severe impairment of both Mg²⁺-dependent and Mn²⁺-dependent activities in the R96G variant and moderate enhancement of only the Mg²⁺-dependent activity in the Δ1–25 variant for matched and mismatched nucleotide incorporations opposite normal and lesion templates (17). Detailed structural and kinetic mechanisms of catalytic alterations of pol λ by these genetic variants still remain unclear. Any disease associations have not been reported yet, but the catalytically altered pol λ genetic variants might be of potential importance in that they would change the TLS capacity of pol λ and consequently modify mutation phenotypes to genotoxic agents in genetically predisposed individuals.

To elucidate both the kinetic and the structural basis for alterations in the catalytic function of pol λ by different two metal ions, Mg²⁺ and Mn²⁺, as well as by two human germline non-synonymous variants, R96G and Δ1–25, we performed pre-steady-state kinetic analysis for nucleotide insertion by pol λ and also determined x-ray crystal structures of ternary pol λ complexes in the presence of either Mg²⁺ or Mn²⁺ ions, using the recombinant human pol λ core (residues 1–445) proteins of wild type and two variants with a simple model of a correct dCTP incorporation opposite normal G. The combined pre-steady-state kinetic and structural results indicate that Mn²⁺ enables pol λ to adopt more ideal octahedral coordination in the active site and achieve much higher catalytic efficiency than Mg²⁺, whereas the R96G variant results in the loss of hydrogen bond interactions of residues Gly-96 and Tyr-93 with an incoming nucleotide as well as conferring a much greater reduction in its catalytic efficiency. Our detailed kinetic and structural results are discussed in the context of understanding the possible mechanistic and functional aspects of metal ions and genetic variations on pol λ.

Results

Pre-steady-state Kinetic Analysis of dCTP Incorporation Opposite G by pol λ(1–445), pol λ(26–445), and R96G pol λ(1–445) Enzymes in the Presence of Mg²⁺ or Mn²⁺—Pre-steady-state kinetic methods were used to quantify the alterations in catalytic efficiency and the apparent nucleotide binding affinity of pol λ by metal ions (Mg²⁺ or Mn²⁺) and known human genetic variants (Δ1–25 or R96G). Pre-steady-state kinetic parameters were determined for dCTP incorporation opposite template G into 18-mer/36-mer duplexes by pol λ (1–445), pol λ (26–445), and R96G pol λ (1–445) enzymes under single turn-over conditions (where pol λ was present in 10-fold excess over DNA substrate), in the presence of either 0.15 mM MnCl₂ or 2 mM MgCl₂, which is in the optimal range for pol λ activity (33, 34), using a rapid quench flow instrument. Analysis of the change of the observed rate (kₐ₀) as a function of increasing dCTP concentration yielded kₚ₀, the maximal rate of nucleotide incorporation, and Kₑ,dCTP, a measure of the binding affinity of dCTP to the pol-DNA binary complex to form a ternary complex poised for catalysis (Table I and supplemental Fig. S1). pol λ(1–445) displayed a kₚ₀ of 0.74 ± 0.06 s⁻¹ and a Kₑ,dCTP of 1.5 ± 0.4 μM in the presence of Mn²⁺. Thus, the catalytic efficiency (kₚ₀/Kₑ,dCTP) of pol λ(1–445) with Mn²⁺ was estimated to be 4.9 × 10⁵ M⁻¹ s⁻¹, which was 770-fold higher than that with Mg²⁺, mainly due to a 450-fold lower Kₑ,dCTP. Similar trends of kinetic results were also observed with pol λ (26–445) and the R96G variant, indicating that pol λ binds nucleotide much more tightly and catalyzes nucleotide insertion much more efficiently in the presence of Mn²⁺ than in the presence of Mg²⁺. pol λ (26–445) had a kₚ₀/Kₑ,dCTP value similar to that of pol λ (1–445) in the presence of Mn²⁺ but displayed a 3-fold increase in kₚ₀/Kₑ,dCTP in the presence of Mg²⁺ when compared with pol λ (1–445), indicating a slight enhancement only in the Mg²⁺-dependent catalytic efficiency of pol λ due to the deletion of N-terminal 25 residues. The R96G variant showed a 53-fold decrease in kₚ₀/Kₑ,dCTP for dCTP insertion opposite G in the presence of Mg²⁺ when compared with that with wild type, while showing a 9-fold decrease of that value in the presence of Mn²⁺. This mitigation effect of Mn²⁺ on the Mg²⁺-de-
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Crystal Structures of Human pol γ Pre-catalytic Ternary Complexes Incorporating Nonhydrolyzable dCTP Analogue (dCMPNPP) Opposite Template G in the Presence of Mg^{2+} or Mn^{2+}—To observe the structural alterations of pol γ by metal ions (Mg^{2+} or Mn^{2+}) and genetic variants (Δ1–25 or R96G), we determined six crystal structures of pre-insertion ternary complexes of pol γ (1–445), pol γ (26–445), and R96G pol γ (1–445) with DNA containing template G and dCMPNPP in the presence of either Mg^{2+} or Mn^{2+}. The strategy, employing a non-hydrolyzable dCTP analogue, dCMPNPP, as well as DNA substrate having an intact 3’-OH at the primer end, was utilized to capture pre-catalytic ternary pol γ complexes that preserve metal-coordinating ligands in the active site while preventing catalysis in the presence of active-site metal ions, as successfully applied with pol β (35). Crystals of ternary complexes of pol γ (26–445), pol γ (1–445), and R96G pol γ (1–445) diffracted to about 2.5–2.6, 3.2–3.6, and 2.8 Å resolution, respectively (Table 2). All ternary complex structures of pol γ contained two metal ions with relatively lower occupancy in the A-site metal ions in the active site, except for the R96G pol γ (1–445)-Mg^{2+} ternary complex that missed an Mg^{2+} ion at the metal A-site near the primer terminus 3’-OH. To our knowledge, our structures represent the first ternary pol γ structures containing the primer end 3’-OH entity, as well as defining the position of two active-site metal ions of either Mg^{2+} or Mn^{2+}, which provides geometric information in the pre-catalytic state. However, electron density was not observed for the N-terminal 25 residues in all pol γ (1–445) ternary complexes, suggesting the disordered nature of this negatively charged N-terminal region. Thus, all the refined structures of pol γ ternary complexes contained pol γ residues 51–439 as observed previously with the ternary pol γ (26–445) complex structure (36). The overall structures of six ternary pol γ complexes were almost identical, except for subtle variations near metal ions in the active site and at the amino acid substitution site (supplemental Fig. S2). Three pol γ structures with the Mn^{2+} ions were superimposed with root mean square deviations (RMSD) of 0.20–0.27 Å among the positions of Ca atoms (supplemental Fig. S2, A and B), whereas three pol γ structures with Mg^{2+} ions were superimposed with RMSD of 0.31–0.35 Å, indicating slightly more backbone variations in the presence of Mg^{2+} (supplemental Fig. S2, A–D). Superposition of relatively high resolution (2.5 and 2.6 Å, respectively) structures of the pol γ (26–445)-Mg^{2+} and pol γ (26–445)-Mn^{2+} ternary complexes showed almost identical overall structures between them with an RMSD of 0.20 Å over Ca atoms but displayed slight differences in positions of metal ions as well as side chains of nearby residues Asp-59 and Glu-152 in the active site (Fig. 1B), indicating possible differences between Mg^{2+} and Mn^{2+} coordination in the pol γ active site.

Coordination of Two Divalent Mg^{2+} or Mn^{2+} Ions in the Active Sites of Human pol γ (26–445) Ternary Complexes with Primer 3’-OH and dCMPNPP—To understand the differences in the metal coordination geometry between Mg^{2+} and Mn^{2+} ions in the pol γ active site, we compared the coordination dependence $k_{pol} / K_{d_{dCTP}}$ reduction in the R96G variant seemed to be mainly due to the full restoration of the apparent dCTP binding affinity to the level comparable with that of wild type by Mn^{2+}.

**Table 2**

| Complex | pol γ (26–445) GdCMPNPP (Mg^{2+}) | pol γ (26–445) GdCMPNPP (Mn^{2+}) | pol γ (1–445) GdCMPNPP (Mg^{2+}) | pol γ (1–445) GdCMPNPP (Mn^{2+}) | R96G pol γ (1–445) GdCMPNPP (Mg^{2+}) | R96G pol γ (1–445) GdCMPNPP (Mn^{2+}) |
|---------|---------------------------------|---------------------------------|---------------------------------|---------------------------------|---------------------------------|---------------------------------|
| Data collection | | | | | | |
| Wavelength (Å) | 0.97872 | 0.97872 | 0.97872 | 0.97872 | 0.97872 | 0.97872 |
| Space group | P6_122 | P6_122 | P6_122 | P6_122 | P6_122 | P6_122 |
| Unit cell (a, b, c) (Å) | 97.61, 97.61, 203.00 | 97.83, 97.83, 201.66 | 97.74, 97.74, 202.01 | 98.12, 98.12, 202.72 | 97.54, 97.54, 202.21 |

**Refinement**

| | $R_{work}$ | $R_{free}$ | Completeness [%] | No. of unique reflections | Wilson plot | $eta$ factor (Å²) | Root mean square deviations | Bond angles (°) | Ramachandran Favorable (%) | Allowed (%) | Outliers (%) | PDB code |
|---|---|---|---|---|---|---|---|---|---|---|---|---|
| | | | | | | | | | | | | |
| | 0.220 | 0.204 | 0.204 | 2928/329 | 68.9 | 21065 | 0.005 | 98.7 | 1.1 | 0.3 | 5KT2 |
| | 0.249 | 0.248 | 0.244 | 2928/329 | 68.9 | 21065 | 1.024 | 97.9 | 1.9 | 0.0 | 5KT3 |
| | | | | 2928/329 | 68.9 | 21065 | 0.003 | 97.1 | 2.9 | 0.0 | 5KT4 |
| | | | | | 2928/329 | | | | | | | 5KT5 |

$^a$ Values for highest resolution shell are given in parentheses.

$^b$ $R_{merge} / R_{linear} = \text{SUM}(|I(I−)|)/\text{SUM}(I)$, where $I$ is the integrated intensity of a given reflection.

$^c$ One non-catalytic metal ion is present in the structure.
structures of Mg$^{2+}$ and Mn$^{2+}$ ions in the pol $\iota$ active site utilizing relatively high resolution structures of the pol $\iota$(26–445)-Mg$^{2+}$ and pol $\iota$(26–445)-Mn$^{2+}$ ternary complexes. The $F_o - F_c$ omit maps for the incoming nucleotide and metal ions showed clear density for the incoming dCMPNPP and two Mg$^{2+}$ (or Mn$^{2+}$) ions bound in the pol $\iota$ active site (Fig. 2, A and B). Two Mg$^{2+}$ or Mn$^{2+}$ ions were present in both the catalytic metal site (A-site) and the nucleotide binding metal site (B-site) in the pol $\iota$ active site, as typically observed with polymerases (37), although having partial occupancies (63 and 67%, respectively) at the A-site. Both Mg$^{2+}$ and Mn$^{2+}$ at the B-site in the pol $\iota$ active site showed nearly perfect octahedral coordination geometry with six ligands, which had similar values of average coordination distance (2.14 and 2.06 Å, respectively) and distance RMSD (0.230 and 0.214 Å, respectively), albeit slightly lower with Mn$^{2+}$ (Table 3 and Fig. 2, C–E), as observed with the B-site Mg$^{2+}$ in the previously reported structures of normal GdCMPNPP ternary complexes of pols $\beta$ and $\eta$ (38, 39). The A-site Mn$^{2+}$ in the pol $\iota$ ternary complex also showed good octahedral coordination geometry involving five ligands, although missing one ligand (possibly a water molecule observed with pols $\beta$ and $\eta$ ternary complexes) (38, 39), yielding average coordination distance (2.29 Å) and distance RMSD (0.305 Å) values that were similar to those (2.22–2.34 and 0.187–0.293 Å, respectively) observed with the A-site Mg$^{2+}$ in the normal G-dCMPNPP ternary complexes of pols $\beta$ and $\eta$ (38,
39) (Table 3 and Fig. 2, D and E). In strong contrast, the Mg\(^{2+}\) at the A-site in the pol \(\lambda\) active site showed a considerable deviation from the ideal octahedral coordination geometry, yielding average coordination distance (2.45 Å) and distance RMSD (0.474 Å) values that were substantially higher than those with Mn\(^{2+}\) (Table 3 and Fig. 2, C and D). These more optimal fea-

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TABLE 3

Active site metal coordination distances and angles in normal G-dCMPNPP ternary complex structures of pols \(\iota\), \(\eta\), and \(\beta\) in the presence of divalent metal ions

| pol          | Metal ion | Inter-metal distance | \(n^a\) | A-site metal coordination | B-site metal coordination |
|--------------|-----------|---------------------|--------|--------------------------|--------------------------|
| \(\iota\) (26–445) | Mg\(^{2+}\) | 3.23                | 5      | 2.45 ± 0.12              | 0.474                    |
|              | Mn\(^{2+}\) | 3.43                | 5      | 2.29 ± 0.06              | 0.305                    |
| \(\eta\)    | Mg\(^{2+}\) | 3.02                | 5      | 2.37 ± 0.14              | 0.869                    |
|              | Mn\(^{2+}\) | 3.33                | 5      | 2.22 ± 0.18              | 0.409                    |
| \(\eta\)    | Mg\(^{2+}\) | 3.55                | 6      | 2.34 ± 0.06              | 0.256                    |
|              | Mn\(^{2+}\) | 3.72                | 6      | 2.22 ± 0.04              | 0.187                    |
| \(\beta\)   | Mg\(^{2+}\) | 3.44                | 6      | 2.23 ± 0.06              | 0.293                    |

\(a\) Coordination number.

\(b\) Average coordination distance ± standard error.

\(c\) Distance RMSD from the ideal octahedral geometry was estimated by UCSF Chimera (56).

\(d\) PDB ID 4DL2 (38).

\(e\) PDB ID 4DL3 (38).

\(f\) PDB ID 4TUQ (39).
tures with Mn$^{2+}$ than Mg$^{2+}$ for octahedral coordination geometry at the A-site were similarly observed with the refined pol $\iota$(1–445) ternary complex structures (Table 3), albeit having lower resolution (3.6 and 3.2 Å). The poor coordination with Mg$^{2+}$ at the A-site appeared to be related to subtle displacement of the side-chain carboxyl group of Asp-59 away from the A-site Mg$^{2+}$ as well as a slight shift of the A-site Mg$^{2+}$ to the B-site Mg$^{2+}$, yielding a 0.2 Å shortening of inter-metal distance when compared with that with Mn$^{2+}$ in the pol $\iota$ active site (Fig. 2, C, D, and F). Interestingly, the C$'$/endo conformation was equally observed at the sugar moieties at the 3$'$ primer end (Fig. 2), as well as the nucleotide 5$'$ to the primer end, and three nucleotide pairs at positions n-2 to n-4 of the primer/template duplex in both pol $\iota$ ternary complex structures, unlike the previously reported structures of the pol $\iota$ binary and ternary complexes lacking the primer end 3$'$-OH (Protein Data Bank (PDB) IDs 2FLP and 2ALZ) (36, 40), indicating a distinctive pattern of sugar pucker changes induced in the pre-catalytic pol $\iota$ ternary complex.

Active-site Structures of Human R96G pol $\iota$ (1–445) Ternary Complexes in the Presence of Mg$^{2+}$ or Mn$^{2+}$—To reveal the structural mechanism of severe catalytic impairments in the R96G pol $\iota$ variant, we compared the active-site structures of R96G pol $\iota$ (1–445) ternary complexes with those of pol $\iota$(26–445) ternary complexes of relatively high resolution. Interestingly, the R96G variant structures showed substantial alterations not only at the amino acid substitution site (Gly-96) but also at the nearby residue site (Tyr-93), when compared with...
The Mg$^{2+}$ ions are slightly improved in the aspect of distance RMSD (2.66 and 0.456 Å, respectively) of the A-site Mg$^{2+}$ ions (albeit involving only four coordinating atoms) in the previously reported structure (PDB ID 2ALZ) of pol $\iota$ ternary complex lacking the primer end 3'-OH (36). This Mg$^{2+}$-induced geometric alteration at the A-site in the pol $\iota$ active site contrasts with the near perfect octahedral coordination of the A-site Mg$^{2+}$ observed in ternary complex structures of other pols $\eta$ and $\beta$ with the correct incoming non-hydrolyzable nucleotide (Table 3 (38, 39)).

The Mn$^{2+}$ requirement for achieving the optimal octahedral coordination geometry at the A-site in ternary complex structures with the correct incoming nucleotide seems to be unique in pol $\iota$. Other DNA polymerases such as bacteriophage pol RB69 and human pol $\beta$ achieve good octahedral coordination geometry not only with Mg$^{2+}$ but also with Mn$^{2+}$ at the A-site, as observed in their ternary complex structures with the correct incoming non-hydrolyzable nucleotide (PDB IDs 3SJ J, 3SPY, 2FMS, and 3C2K) (35, 41, 42). It is notable that the superior coordinating ability of Mn$^{2+}$ when compared with Mg$^{2+}$ at the A-site is observed in the previously reported structures of ternary pol $\beta$ complexes (PDB IDs 4PGQ, 4PGX, 4PHA, and 4PHD) with an incoming incorrect non-hydrolyzable nucleotide (43). Our data and that of others suggest that Mn$^{2+}$ is more tolerant of atypical pol active sites (e.g., the inherently restricted pol $\iota$ active site and the distorted pol active site due to base pair mismatch) than Mg$^{2+}$ and thus able to form good octahedral coordination geometry not only at the B-site but also at the A-site in the pol active site. This effect might be attributed to a relaxed coordination requirement of Mn$^{2+}$ when compared with Mg$^{2+}$ (37). It is of interest to perform further studies to verify whether our results of pol $\iota$ with normal base pairs of template G and incoming dCTP are valid for other base pairs involving DNA lesions or mismatches such as G:T and T:G.

The Mg$^{2+}$-induced distortion of the A-site coordination geometry in the pol $\iota$ active-site structure seems to be closely related to a severe (220–770-fold) diminution in catalytic efficiency ($k_{cat}/K_{M,dCCTP}$) of pol $\iota$ in the presence of Mg$^{2+}$ when shorter than that with Mn$^{2+}$ ions (Table 3). Thus, in the pol $\iota$ active site, only Mn$^{2+}$ ions but not Mg$^{2+}$ ions seem to achieve an inter-metal distance similar to that with Mg$^{2+}$ ions usually observed in active sites of ternary complex structures of other human pols $\eta$ and $\beta$ with normal GdCMPNPP (38, 39) (Table 3).

Superposition of the active-site metal binding sites of pol $\iota$Mg$^{2+}$, pol $\iota$Mn$^{2+}$, and pol $\gamma$Mg$^{2+}$ (PDB ID 4DL3) ternary complexes appears to reflect some gradational changes in the aspect of positional shift of the A-site metal (toward the B-site metal) as well as in the angle of rotational displacement of the side-chain carboxyl group of Asp-59 (or Asp-13 for pol $\iota$), in the order of pol $\eta$Mg$^{2+} < \gamma$Mg$^{2+} < \iota$Mg$^{2+}$ (Fig. 2F). Consequently, Mg$^{2+}$ ions led to a considerable deviation from the ideal octahedral coordination geometry at the A-site, yielding the values of average coordination distance and distance RMSD (2.45 and 0.474 Å, respectively), which were quite higher than those with Mn$^{2+}$ (Table 3). These coordination parameters with Mg$^{2+}$ ions seem to be slightly improved in the aspect of average coordination distance but worsened in the aspect of distance RMSD by the presence of the primer terminus 3'-OH, when compared with those (2.66 and 0.456 Å, respectively) of the A-site Mg$^{2+}$ (albeit involving only four coordinating atoms) in the previously reported structure (PDB ID 2ALZ) of pol $\iota$ ternary complex lacking the primer end 3'-OH (36). This Mg$^{2+}$-induced geometric alteration at the A-site in the pol $\iota$ active site contrasts with the near perfect octahedral coordination of the A-site Mg$^{2+}$ observed in ternary complex structures of other pols $\eta$ and $\beta$ with the correct incoming non-hydrolyzable nucleotide (Table 3 (38, 39)).

The key differences between the pol $\iota$ (26–445) ternary complex structures (Fig. 3, B and D), which were obviously observed in those with the pol $\iota$ (26–445) complexes (Fig. 3, A and C). This collagen destabilization of the Tyr-93 side chain seems to be related to the loss of a cation-π interaction between Arg and Tyr side chains due to the loss of an Arg side chain in the R96G variant. Accordingly, two hydrogen bonds of the Arg-96 side chain with a β,γ-bridging oxygen and a γ-phosphate oxygen of dCMPNPP, as well as one hydrogen bond of the Tyr-93 side chain with a γ-phosphate oxygen, were lost in R96G variant structures when compared with those in pol $\iota$ (26–445) ternary complex structures (Fig. 3, E–H). Interestingly, only Mn$^{2+}$ but not Mg$^{2+}$ was observed at the A-site in the R96G pol $\iota$ active site (Fig. 3, B and D), possibly reflecting an inferior A-site binding ability of Mg$^{2+}$ when compared with Mn$^{2+}$, which might in part be attributed to a 6-fold greater decrease in catalytic efficiency of the R96G variant with Mg$^{2+}$ than that with Mn$^{2+}$ (Table 1).

Discussion

In this study, we provide structural and pre-steady-state kinetic evidence that Mn$^{2+}$ is more optimal for the two-metal ion binding site configuration and catalysis of pol $\iota$ than Mg$^{2+}$. Our pol $\iota$ ternary complex structures containing intact coordinating ligands such as the primer end 3’-OH reveal that Mn$^{2+}$ ions achieve more ideal octahedral coordination geometry than Mg$^{2+}$ ions in the pol $\iota$ active site, specifically in the catalytic metal A-site. Moreover, our pre-steady-state kinetic data demonstrate that Mn$^{2+}$ ions confer much higher (2–3 orders of magnitude) efficiency of pol $\iota$ catalysis than Mg$^{2+}$ ions, mainly through augmenting nucleotide binding affinity. We also confirmed that the R96G variant, displaying a severe reduction in catalytic efficiency, loses three hydrogen bond interactions of other pols $\eta$ and $\beta$ with the correct incoming non-hydrolyzable nucleotide (Table 3 (38, 39)).
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compared with Mn²⁺ (Table 1). This severe catalytic impairment with Mg²⁺ was mainly due to a severe reduction in the affinity of productive nucleotide binding of pol τ in the presence of Mg²⁺ when compared with Mn²⁺, as reflected by large (220–450-fold) increases in the apparent equilibrium dissociation constant ($K_d$) for incoming dCTP (Table 1). These results suggest that substitution of Mn²⁺ for Mg²⁺ might boost the catalytic efficiency of pol τ for correct nucleotide insertion, mainly through improving the binding affinity of nucleotide, by achieving optimal octahedral coordination at the A-site in the active site. Our finding is in good agreement with previous studies with other polymerases, e.g. RB69 and pol β (42, 44). It is also notable that the extent of increases (220–450-fold) in the superimposition of Mg²⁺ (220–450-fold) increases in the apparent equilibrium dissociation constant ($K_d$) for incoming dCTP (Table 1). These results suggest that substitution of Mn²⁺ for Mg²⁺ might boost the catalytic efficiency of pol τ for correct nucleotide insertion, mainly through improving the binding affinity of nucleotide, by achieving optimal octahedral coordination at the A-site in the active site. Our finding is in good agreement with previous studies with other polymerases, e.g. RB69 and pol β (42, 44), implying a more marked effect of Mn²⁺ on pol τ than other polymerases. Our combined structural and kinetic data suggest that the optimal octahedral coordination of two active-site metal ions is essential for proper catalytic function of pol τ and Mn²⁺ is superior in this respect when compared with Mg²⁺, particularly at the A-site in the pol τ active site. The superiority of Mn²⁺ for pol τ function is also supported by the biochemical property of pol (1–445) to more tightly bind DNA substrates in the presence of a low level of Mn²⁺ than Mg²⁺ (17). Although DNA polymerases most likely utilize physiologically abundant Mg²⁺ ions for catalysis in vivo, from our and previous studies (17, 33, 34), it may be relevant to postulate that pol τ would inherently employ physiologically low levels of Mn²⁺ for catalysis in kinetic and structural preference to Mg²⁺ in vivo. Similarly, pol λ has been also suggested to use Mn²⁺ as the preferred activating metal ion in vivo (45). Mn²⁺ has also been reported to increase the activity of non-canonical DNA polymerases such as Sulfolobus solfataricus Dpo4 and human PrimPol in vitro (46, 47). We also note a very recent study suggesting the requirement of a third metal ion for pol η catalysis, with slight preference for Mg²⁺ (48). It would be of interest to investigate whether this is the case in pol τ catalysis.

The loss of hydrogen-bonding interactions of two structurally altered residues, the substituted Gly-96 and the nearby destabilized Tyr-93, with the incoming nucleotide in the crystal structures of the R96G pol τ ternary complexes (Fig. 3) may provide a molecular explanation for severe diminution of the catalytic efficiency in the R96G variant (Table 1). The disturbed electron density of the Tyr-93 side chain in R96G pol τ crystal structures seems to be related to the destabilization of the Tyr-93 side chain due to the absent cation-π interaction with Gly-96. Both Arg-96 and Tyr-93 residues, which are conserved and important residues for nucleotide binding in all Y-family pols (20), seem to co-stabilize their side chain conformations by forming the cation-π interaction between their parallel side chains as observed in our pol τ ternary complexes (Fig. 3). In good accordance with our data, individual missense mutations of two homologous Arg-67 and Tyr-64 residues in yeast pol η severely diminish its catalytic activity (49). It can be postulated that the weakened interaction of the R96G variant with incoming nucleotide likely diminishes its nucleotide binding affinity and catalytic efficiency. Interestingly, the attenuating effect of the R96G variation on the apparent nucleotide binding affinity of pol τ was observed much more strongly in the presence of Mg²⁺ (Table 1). Mn²⁺ substitution fully restored the apparent nucleotide binding affinity of the R96G variant to a level similar to that of wild-type pol τ (Table 1), indicating a rescuing effect of Mn²⁺ to offset the destabilized nucleotide interaction in the R96G variant. Accordingly, the extent of reduction of catalytic efficiency in the R96G variant was considerably lessened in the presence of Mn²⁺ when compared with Mg²⁺ (Table 1).

The variant of pol (26–445) lacking the N-terminal 25 residues retained an Mn²⁺-dependent catalytic efficiency almost similar to that of the pol (1–445) but displayed a 3-fold increase in selectivity in Mg²⁺-dependent catalytic efficiency when compared with that of the pol (1–445) (Table 1). These results are in good agreement with our previous steady-state kinetic data (17). However, the structural effects of the N-terminal 25 residues are not clear due to their disordered nature in the x-ray crystal structure. Although it may not be obvious due to its low resolution (3.6 Å), interestingly, our refined crystal structure of pol (1–445) ternary complex in the presence of Mg²⁺ appeared to have slightly altered conformations of Arg-96 and Tyr-93 side chains when compared with the pol (26–445) ternary complex (supplemental Fig. S1C). This subtle structural alteration, observed only with Mg²⁺ ions in the pol (1–445) active site, may in part explain a slightly lower efficiency for Mg²⁺-dependent catalysis by pol (1–445) when compared with pol (26–445).

In summary, we have investigated the effects of different metal ions (Mg²⁺ and Mn²⁺) and genetic variations (R96G and Δ1–25) on both the structure and the catalytic function of pol τ. Here we report the first x-ray crystal structures acquired in the presence of either Mg²⁺ or Mn²⁺ of pol τ ternary complexes having intact coordinating metals and ligands such as the primer terminus 3′-OH. Comparisons of active-site conformations between pol τ-Mg²⁺ and pol τ-Mn²⁺ ternary complexes revealed that pol τ adopts near perfect octahedral coordination geometries for two metal ions in the active site only in the presence of Mn²⁺. This structural feature with Mn²⁺ is clearly consistent with the pre-steady-state kinetic observation that Mn²⁺ greatly bolsters the apparent nucleotide binding affinity and the catalytic efficiency of pol τ when compared with observations with Mg²⁺. Moreover, our combined structural and pre-steady-state kinetic analysis also revealed that the catalytic impairment in the R96G variant is related to the lack of hydrogen-bonding interactions of Gly-96 and Tyr-93 with incoming nucleotides in the active site. Our comparison between pol (1–445) and pol (26–445) also suggests a potential role of the disordered N-terminal 25 amino acids in selectively improving (albeit slightly) the Mn²⁺-dependent catalytic efficiency in the wild-type pol τ. Overall, our study provides insights into the delicate structural and kinetic features of different metal coordination and genetic variants that contribute to understanding of the molecular basis of the catalytic function of pol τ.

Experimental Procedures

Materials—T4 polynucleotide kinase and dNTPs were purchased from New England Biolabs (Ipswich, MA). [γ-³²P]ATP
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**Pre-steady-state Reactions**—The 18-mer primer (5′-AGC CAG CCG CAG ACG CAG-3′) was 5′ end-labeled using T4 polynucleotide kinase with [γ-32P]ATP and annealed with 36-mer template (3′-CGG AGC TCG GTC GGC GTG GTC GTC GCT CCT CGC GCT-5′). Rapid quench experiments were performed using a model RQF-3 KinTek Quench Flow instrument (KinTek Corp., Snow Shoe, PA). All DNA polymerase reactions were performed in 50 mM Tris-HCl (pH 7.5) buffer containing 5 mM dithiothreitol, 100 μg ml−1 BSA (w/v), 10% glycerol (v/v), and 2 mM MgCl2 (or 0.15 mM MnCl2). Reactions were initiated by rapid mixing of [32P]-primer/template/polymerase mixtures (18-mer/36-mer, 100 nM; pol λ, 1 μM, in 10-fold excess to DNA substrate to ensure single turnover conditions) with the metal-dCTP mixtures (2 mM MgCl2 or 0.15 mM MnCl2; dCTP, in varying concentrations), and then quenched with 0.15 M EDTA at time intervals from 0.15 to 30–120 s (or from 2 to 240–480 s for the R96G variant). MgCl2 was supplemented by as much as the increase of dCTP to counterbalance the Mg2+-chelating effect of dCTP for the reactions at high levels of dCTP. Reaction products were mixed with formamide-dye solution (20 mM EDTA, 95% formamide (v/v), 0.5% bromphenol blue (w/v), and 0.05% xylene cyanol (w/v)) and then separated using an 8 M urea-containing denaturing gel with 0.5% bromphenol blue (w/v), and then quantified by a Bio-Rad 16% polyacrylamide (w/v), and then quantified by a Bio-Rad

**Determination of kpol and Kd,dCTP**—The pre-steady-state kinetic parameters kpol and Kd,dCTP were estimated by analyzing the dCTP dependence on the observed pre-steady-state rates of dCTP insertion under single turnover conditions. A graph of the observed rate (kobs) versus dCTP concentration was fit to the hyperbolic equation $k_{obs} = [k_{pol}[dCTP]]/[([dCTP] + K_p)]$, where $k_{pol}$ is the maximal rate of nucleotide incorporation and $K_{d,dCTP}$ is the equilibrium dissociation constant for dCTP (50, 51).

Crystallization of pol λ-DNA: dCMPNPP Ternary Complexes in the presence of Mg2+ or Mn2+—Crystals were obtained under the conditions previously reported (34, 36) with slight modifications described below. The purified pol λ enzymes (0.22 mM) were incubated with annealed self-complementary 18-mer DNA (5′-TCT GGG GTG CTA GGA CCC-3′, 0.26 mM) and 20 mM dCMPNPP (or 4 mM dCMPNPP for MnCl2) in the presence of 10 mM MgCl2 (or 2 mM MnCl2) on ice. The ternary complexes were crystallized in 0.1 M MES (pH 6.5), 0.2–0.4 M (NH4)2SO4, and 10–22% polyethylene glycol 5000 (w/v) using the hanging-drop vapor diffusion method at 4°C. Crystals were typically observed in 1–3 days. Crystals started to form with reservoir solutions containing 13–15, 18, and 18–20% polyethylene glycol monomethyl ether 5000 (w/v), respectively, for pol λ (26–445), pol λ (1–445), and R96G pol λ (1–445) ternary complexes. Crystals were mounted in nylon loops and cryoprotected in a reservoir solution containing 25% glycerol (v/v), and then flash-frozen in liquid nitrogen.

**Structure Determination and Refinement**—X-ray diffraction data were collected on the 21-ID-F (Life Sciences Collaborative Access Team (LS-CAT)) beam line at the Advanced Photon Source (Argonne National Laboratory, Argonne, IL). Collected data were indexed, integrated, and scaled using HKL2000 (52). All crystal types belonged to space group P6322. Structures were determined by molecular replacement phasing using the program Phaser MR (53) and the pol λ structure (PDB ID 2ALZ) as a search model. Structure refinements and model building were performed using PHENIX (54) and COOT (55). Metal coordination geometry was analyzed with UCSF Chimera (56). Structural illustrations were prepared with PyMOL (Schrödinger, LLC).

**Author Contributions**—F. P. G. and J.-Y. C. conceived the study and designed the experiments. J.-Y. C. and M. Y. purified the enzymes. M. Y., J.-Y. C., and Q. Z. conducted the kinetic experiments. J.-Y. C. and A. P. crystallized the protein complexes. Y.-S. L., J.-Y. C., and A. P. solved the structures. J.-Y. C., F. P. G., M. E., and Y.-S. L. analyzed the data and wrote the paper.

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