A polar filter in DNA polymerases prevents ribonucleotide incorporation

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ABSTRACT

The presence of ribonucleotides in DNA can lead to genomic instability and cellular lethality. To prevent adventitious rNTP incorporation, the majority of the DNA polymerases (dPols) possess a steric filter. The dPol named MsDpo4 (Mycobacterium smegmatis) naturally lacks this steric filter and hence is capable of rNTP addition. The introduction of the steric filter in MsDpo4 did not result in complete abrogation of the ability of this enzyme to incorporate ribonucleotides. In comparison, DNA polymerase IV (PolIV) from Escherichia coli exhibited stringent selection for deoxyribonucleotides. A comparison of MsDpo4 and PolIV led to the discovery of an additional polar filter responsible for sugar selectivity. Thr43 represents the filter in PolIV and this residue forms interactions with the incoming nucleotide to draw it closer to the enzyme surface. As a result, the 2'-OH in rNTP will clash with the enzyme surface, and therefore ribonucleotides cannot be accommodated in the active site in a conformation compatible with productive catalysis. The substitution of the equivalent residue in MsDpo4–Cys47, with Thr led to a drastic reduction in the ability of the mycobacterial enzyme to incorporate rNTPs. Overall, our studies evince that the polar filter serves to prevent ribonucleotide incorporation by dPols.

INTRODUCTION

DNA polymerases (dPols) synthesize DNA through template-dependent addition of dNTPs to the growing primer strand. All dPols share a similar core structure with thumb, fingers, and palm domains. Three acidic residues constitute the catalytic center of these enzymes, and these residues are present in the palm domain. The catalytic residues co-ordinate a divalent metal ion, usually Mg²⁺, that is critical for DNA synthesis activity (1–3).

In the cellular environment, dPols encounter rNTPs much more frequently than dNTPs. The cellular concentrations of rNTPs are up to 10–100-fold higher than those of dNTPs depending on the type of cell, and stage of cell cycle (4–9). Like dNTPs, rNTPs can form Watson-Crick base pairs with DNA nucleotides, as they are sister molecules of dNTPs with an additional 2'-OH on the sugar moiety (4,10,11). Hence it is possible, that the dPols may inadvertently add rNTPs to the growing primer strand. The presence of ribonucleotides in DNA is detrimental to the integrity of the genome as they are less stable due to the reactive nature of the hydroxyl group on the C2' atom (7–10,12,13). The presence of ribonucleotides in DNA can lead to the spontaneous appearance of double-strand breaks (5,8,13,14). The misincorporation of rNTPs into DNA is known to slow down the rate of DNA replication by Polβ, Polε and PolIII (5,6,12,15). Ribonucleotides can alter the conformation of DNA from B to A form, and thus adversely affect target sequence recognition by proteins such as transcription factors (10,16,17). Overall, the presence of ribonucleotides in the genomic DNA is detrimental to cellular physiology and, represents a significant challenge for the basic genomic processes such as replication, repair, and transcription (13,17). It is, therefore, imperative to reduce the frequency of incorporation of rNTPs into the genome during DNA replication.

Most dPols employ a steric filter to minimize rNTP incorporation (4,18–20). The steric filter is represented by an amino acid residue with a bulky side chain which will clash with the 2'-OH group on the ribose ring of the incoming nucleotide (Supplementary Figure S1) (9,19,21). As a result, rNTPs cannot bind stably in the active site of the dPol. The steric gate is represented by Glu in A-family polymerases and Tyr or Phe in members of the B, X, Y and RT families (11,18–20,22–36).

MsDpo4 is a Y-family dPol from Mycobacterium smegmatis (37,38). MsDpo4 does not possess the steric filter and can incorporate ribonucleotides (28,39). In comparison, DNA polymerase IV (PolIV) from Escherichia coli, exhibits stringent sugar selectivity. A rigorous comparison between MsDpo4 and PolIV shows the presence of a strateg-
Figure 1. Comparison of rNTP incorporation by MsDpo4-WT, MsDpo4-L14Y, and PolIV. (A) Sequence alignment of the N-terminal region of MsDpo4 and homologous proteins from bacteria, archaea, and humans shows that MsDpo4 lacks the steric filter as it has a Leu residue instead of an aromatic residue at the appropriate position (highlighted in red). Primer extension assays show that (B) MsDpo4-WT is capable of rNTP and dNTP incorporation. (C) MsDpo4-L14Y can incorporate rNTP and dNTP and (D) PolIV-WT from E. coli is unable to incorporate rNTP and therefore exhibits stringent sugar selectivity.

The PolIII gene (E. coli) was amplified from E. coli genomic DNA and cloned into a modified pET-28b vector (pDJN1) (41). The exonuclease deficient mutant of PolIII (D335A) (42) was generated using QuickChange Lightning site-directed mutagenesis kit (Agilent). The protein was expressed in the C41(DE3) strain of E. coli. The freshly transformed cells were grown to an OD$_{600}$ = 0.8, at 37°C. The expression of PolIII was induced with 0.4 mM IPTG followed by incubation at 18°C for 16 h. The cells were then harvested by centrifugation, lysed by sonication and the lysate was clarified by centrifugation. The purification was done using pre-packed Ni-NTA column (GE-Healthcare). The binding buffer (Buffer A) composition was 25 mM Tris pH 8.0, 250 mM NaCl, and 5% glycerol. The elution buffer (Buffer B) composition was 25 mM Tris pH 8.0, 250 mM NaCl, 5% glycerol and 1 M Imidazole. The bound protein was eluted using a step gradient of elution Buffer B. The eluted protein was then dialyzed overnight in Buffer A and then, concentrated and stored in $-80^\circ$C.
Figure 2. Structural mechanism of rCTP and dCTP incorporation by MsDpo4-WT. (A) The structures of MsDpo4-WTDNA(dG):dCTP (colored yellow) and MsDpo4-WTDNA(dG):rCTP (colored violet) superimpose with an RMSD of 0.36 Å. The enzyme structure is shown in cartoon representation, DNA is shown as ribbons, and the incoming nucleotide and the Leu14 residue (in green) are shown in stick representation. A close-up of the region surrounding the incoming nucleotide, is displayed for (B) MsDpo4DNA(dG):rCTP and (C) MsDpo4DNA(dG):dCTP. The comparison shows that the dCTP and rCTP bind the MsDpo4 active site in the same location and conformation with marginal differences in the enzyme structure.

Figure 3. Mechanism of rCTP incorporation by MsDpo4-L14Y. (A) A close-up of the region surrounding the incoming nucleotide is displayed for MsDpo4-L14YDNA(dG):rCTP structure. (B) The structures of MsDpo4-L14YDNA(dG):dCTP (colored orange) and MsDpo4-L14YDNA(dG):rCTP (colored white), superimpose with an RMSD of 0.37 Å. The comparison shows that the rCTP is repositioned and undergoes a change in the sugar pucker to prevent a steric clash between the 2′-OH and the engineered steric filter. (C) The surface of the fingers domain (in yellow) from the MsDpo4-L14YDNA(dG):rCTP structure is shown along with space filling model of the bound rCTP (colored according to element). The figure shows that there is no steric clashes between the rCTP molecule and the enzyme surface in this structure.

Primer extension assay and steady-state kinetic analysis

To prepare the DNA substrate, a 6FAM labeled 15mer primer (5′-6FAM-CGTACTCGTAGGCAT-3′) was annealed to 50mer template (5′-TCCTACCGTGCCTACCTGAACAGCTGGTCACACA XATGCCTACGATCG-3′). The nucleosides deoxyadenosine, deoxycytidine, deoxythymidine and deoxyguanosine occupied the underlined position for the templates A, C, T and G respectively. The reaction mixture was composed of enzyme (MsDpo4/PollV/PollII), dsDNA, dNTPs or rNTPs (Jena Bioscience), 25 mM Tris pH 8.0, 0.2 mM DTT, 0.1 mM (NH₄)₂SO₄, 2.5 mM MnCl₂ and 0.05 mg/ml BSA in a final volume of 20 µl. The reaction was carried out at 37°C for 60 min. The reaction was terminated by adding 10 µl of stop buffer, composed of 80% formamide, 10% bromophenol blue and 10% 0.5 M EDTA (40,43). The reaction products were denatured by heating at 95°C for 10 min and quick cooling in ice for 10 min. The products were then resolved in a 20% polyacrylamide gel with 8 M urea, in 1× TBE buffer and visualized using Typhoon scanner FLA-7000 (GE Healthcare). The band intensities were quantified using ImageQuant software. Relative percentage incorporation (RPI) in each group was calculated with respect to the highest in the group. RPI was calculated using the following equation

\[
\text{RPI} = \left( \frac{I}{I'} \right) \times 100,
\]

where I is the percentage incorporation by the mutant and I’ is the percentage incorporation by the most active version of the protein. For steady-state kinetic analysis, the reaction was carried out for varying time points (30 s to 30 min) at 37°C. The time point at which 20% of the primer had been extended was used for further analysis. Reactions were carried out with varying concentrations of incoming rCTP. After quantification of the resolved products, apparent \(K_m\) and \(V_{max}\) values were calculated using a Lineweaver–Burk plot. The measurements were carried out in triplicate, and the standard deviation values were calculated.
The X-ray diffraction data for MsDpo4 DNA(dG):dCTP, X-ray generator with a PhotonII detector (Bruker Inc.).

Crystallization, structure determination & crystallographic refinement

An 18mer self-complementary oligonucleotide (5′-TCTGGGTGCTCTAGGACC 3′) was purified by ion-exchange chromatography using a monoQ column. The purified oligonucleotide was desalted, lyophilized, solubilized in water and annealed. The ternary complexes were reconstituted by mixing MsDpo4 (0.3 mM) with annealed dsDNA in a molar ratio of 1:1.2 followed by addition of 5 mM dCpNHpp or CpCpp (Jena Bioscience) and MgCl2 or MnCl2, respectively. Crystallization trials were carried out by the hanging drop method using commercial screens. After screening and optimization, the best crystals were obtained in 0.2 M Bis-Tris Propane (pH 6.0), 15–25% (w/v) PEG 3350 or PEG 2KMME and 0.1–0.7 M NaCl.

The PolIV-F13A+T43C protein in complex with DNA (template dG) and incoming rCpCpp or dCpNHpp was crystallized using the same conditions as that for wt-protein described previously (40,43). The initial characterization of the crystals was carried out using a Metaljet X-ray generator with a PhotonII detector (Bruker Inc.). The X-ray diffraction data for MsDpo4DNA(dG):dCTP, MsDpo4-L14YDNA(dG):dCTP and MsDpo4-L14YDNA(dG):rCTP were collected at the BM14 beamline of the European Synchrotron Radiation Facility (ESRF) in Grenoble. Data corresponding to MsDpo4-C47T_DNA(dG):dCTP, PolIV-F13A+T43C_DNA(dG):dCTP and PolIV-F13A+T43C_DNA(dG):rCTP were collected at ID30B and ID29 beamline at ESRF, Grenoble, France. The data for MsDpo4-C47T+L14Y_DNA(dG):dCTP was collected at Eletra Sincrotrone in Trieste.

All images were integrated using iMOSFLM of CCP4 (44). The structure of the SeMet–MsDpo4_DNA(dG):dCTP complex was determined by single-wavelength anomalous dispersion method using the AUTOSOL program in PHENIX (45). The initial model was built using the AUTOBUILD program in PHENIX, and subsequent manual rebuilding was done using COOT (46). The other structures of ternary complexes of wt- or mutant versions of MsDpo4 were determined by Molecular Replacement using PHASER in CCP4 (47,48). Similarly, the structure of PolIV-T43C+F13A_DNA(dG):rCTP was determined by PHASER using the structure of PolIV_DNA(dG):dCTP (41RC) as a search model. All the structures were refined using PHENIX until convergence after which TLS refinement was carried out using Refmac in CCP4 (49).

Growth assays

The DNA sequence corresponding to the β-clamp binding peptide from E. coli PolIV (342QMERQLVLG351) (50) was added to the 3′ end of the MsDpo4 gene, and this gene construct (msdpo4β) was cloned into a modified version of the pET-28b vector (41). The C47T mutation of the msdpo4β gene was generated using QuickChange Lightning site-directed mutagenesis kit (Agilent). The presence of the mutation was confirmed by gene sequencing. The growth assays were conducted using the B834 strain of E. coli. The spot assay was carried out with a serial dilution of culture harvested at the log phase (A600 = 0.6). LB-Ampicillin plates with 0.1 mM IPTG were used to induce the expression of the different constructs and assess their effect on cell survival. LB-Ampicillin plates with 1% glucose were used as the control. The plasmids bearing wt- and mutant msdpo4β gene constructs, and the empty vector were transformed into fresh competent cells of B834, and the transformed cells were grown overnight. Single colonies were picked and used to inoculate a starter culture which was harvested at the log phase (A600 = 0.6). The following serial dilutions were prepared- 10−1, 10−2 and 10−3. For each dilution, 5 μl was spotted on LB ampicillin plates with 0.1 mM IPTG or 1% glucose. The plates were incubated in 37°C for 12 h after which the observations were recorded.

The standard plate assay described by McDonald et al. was performed to quantitate the effect of the C47T mutation on bacterial survival (51). The transformed cells were grown to an OD600 = 0.6, and each culture was induced with 0.1 mM IPTG. Subsequently, the cultures were grown at 37°C for 5 h, following which the cells were plated and incubated overnight. The colonies obtained were counted, and the experiment was repeated three times. The fold change in the number of colony forming units (CFUs) was then calculated with respect to the wild type. The assays were repeated.
Figure 5. The polar residue reduces rCTP incorporation by MsDpo4. The results of primer extension assays conducted using different concentrations of wt- MsDpo4 and variants (MsDpo4-L14Y, MsDpo4-C47T & MsDpo4-C47T+L14Y) at different concentrations of rCTP or dCTP are shown. The concentrations of enzyme used were 5 nM (Lanes 1, 2 and 3), 10 nM (Lanes 4, 5 and 6) and 15 nM (lanes 7, 8 and 9). The concentrations of incoming nucleotide (rCTP or dCTP) used were 1/2 M (lanes 1, 4 and 7), 2/2 M (lanes 2, 5, and 8) and 4/2 M (lanes 3, 6 and 9). The assays show that mutation of Cys47 to Thr enhances the ability of MsDpo4 to exclude ribonucleotides. The MsDpo4-C47T and MsDpo4-C47T+L14Y proteins show higher selectivity for dCTP than MsDpo4-WT or MsDpo4-L14Y. These experiments shows that engineered residue (Thr47) enhances sugar selectivity.

RESULTS

Presence of the steric gate does not abrogate the ability of the MsDpo4 to incorporate ribonucleotides

The steric filter is naturally absent in MsDpo4, and it has a Leu at the 14th position, instead of Tyr or Phe (Figure 1A). Primer extension assays show that MsDpo4 can incorporate both dNTPs and rNTPs (Figure 1B). The aromatic steric gate was engineered in MsDpo4, by mutating the Leu14 residue to Tyr. Surprisingly, primer extension assays showed that the MsDpo4-L14Y enzyme still exhibited substantial ability to incorporate rNTP (Figure 1C). In comparison, PolIV-WT did not show significant incorporation of rNTPs and thus exhibited stringent sugar selectivity (Figure 1D). Steady-state kinetic analysis carried out to compare the ability of MsDpo4-WT and MsDpo4-L14Y to incorporate ribonucleotides showed that addition of the steric gate led to only a six-fold reduction in the catalytic efficiency of rCTP incorporation (Table 1).

To understand the structural basis of ribonucleotide incorporation by MsDpo4-WT and the L14Y mutant, we first determined the structure of the wt- enzyme in complex with DNA bearing dG at the templating position and rCTP or dCTP as the incoming nucleotide to a resolution of 2.5 and 2.3 Å, respectively (Supplementary Table S2A and B). The
The presence of the polar residue results in drastic reduction in incorporation of all rNTPs. The level of rNTP incorporation by MsDpo4-WT and variants (MsDpo4-C47T, MsDpo4-L14Y & MsDpo4-L14Y+C47T) are displayed. (A) The level of incorporation of rATP opposite template dT by MsDpo4-WT and variants is displayed. (B) The figure exhibits the level of incorporation of UTP opposite template dA by MsDpo4-WT and variants. (C) The level of incorporation of rGTP opposite template dC by MsDpo4-WT and variants is displayed. (D) The ability of MsDpo4 and variants to incorporate rCTP opposite dG is displayed. For all graphs, the error bars indicate standard deviation (n = 3). For MsDpo4, the presence of the polar filter (C47T) results in a drastic reduction in rNTP incorporation and the presence of both filters in MsDpo4 gives rise to stringent sugar selectivity.

Table 1. Steady-state kinetics of rCTP incorporation by MsDpo4-WT and MsDpo4-L14Y

| Protein            | $k_{cat}$ (s⁻¹) | $K_m$ (μM) | $k_{cat}/K_m$ (μM⁻¹ s⁻¹) |
|--------------------|-----------------|------------|--------------------------|
| MsDpo4-WT          | 0.66 ± 0.07     | 55.3 ± 4.0 | 0.01 ± 0.001              |
| MsDpo4-L14Y        | 0.1 ± 0.01      | 59.54 ± 13.4 | 0.0016 ± 0.0003           |

*The numbers after ± denote standard deviation (n = 3).*

The structure of MsDpo4-WT showed that both dCTP and rCTP form Watson-Crick base pairs with the template dG, and are present in the location and conformation compatible with productive catalysis (Figure 2A). The two structures superimposed onto each other with an RMSD of 0.36 Å and showed no difference in the position and conformation of incoming nucleotide. These structures, therefore, show that MsDpo4 can bind rCTP and dCTP without any distortion in enzyme and DNA structure (Figure 2B and C). The structure of MsDpo4-L14Y was also determined with DNA bearing dG at the templating position and dCTP or rCTP as the incoming nucleotide to a resolution of 2.16 and 2.06 Å, respectively (Supplementary Table S2C and D). The structure showed that the rCTP was accommodated in the active site of MsDpo4-L14Y without any steric clash between the aromatic residue and 2’-OH (Figure 3A).

The structures of MsDpo4-L14Y in complex with DNA (template dG) and incoming rCTP or dCTP superimposed onto each other with an RMSD of 0.37 Å (Figure 3B). This comparison showed that, unlike in the case of the wt- enzyme, the incoming rCTP was repositioned. The rCTP is situated slightly upwards such that the C2’ atom of the sugar ring of rCTP is present 1.3 Å above that of the dCTP. Also, the sugar of the rCTP attains a C3’ endo conformation, and due to this, the 3’-OH group is present 2.4 Å above the corresponding location in the case of dCTP (Figure 3B).
Due to these structural differences, the 2'-OH of the incoming rCTP is present in the same location as the C2' atom of dCTP and therefore will not clash with the τ-electron cloud over the aromatic side chain of the engineered Tyr14 residue (Figure 3C). The location of the triphosphate moiety is similar in the case of rCTP and dCTP and the changes in the position and conformation of the sugar do not adversely affect the ability of the rCTP to form a productive Watson-Crick base pair with the template dG. Overall, due to the repositioning of the rCTP and change in the sugar pucker, rCTP can bind to the MsDpo4-L14Y enzyme in a conformation compatible with catalysis and without any steric clashes with the enzyme surface (Figure 3C). Thus, the comparison shows that the presence of a steric filter is not adequate to achieve stringent sugar selectivity.

The presence of a polar filter prevents ribonucleotide incorporation
Since PolIV exhibits stringent ribonucleotide exclusion, the structures of ternary complexes of MsDpo4-WT and PolIV-WT (PDB ID 4IRC) (40) were compared to identify additional determinants of sugar selectivity. The structures of MsDpo4-WT and PolIV-WT, in complex with DNA (tem-
Figure 8. Presence of the polar filter enhances bacterial survival. Spot Assay was conducted to assess the effect of MsDpo4-WT and MsDpo4-C47T on bacterial survival. The pDJN1 (vector alone), MsDpo4-WT and MsDpo4-C47T are shown in rows from top to bottom respectively. (A) Serial dilutions of cells expressing MsDpo4-WT and variants were spotted on LB ampicillin plates with 0.1mM IPTG and it was seen that bacteria expressing MsDpo4-C47T exhibit better survival than those expressing MsDpo4-WT. (B) Serial dilutions of cells expressing MsDpo4-WT and MsDpo4-C47T were spotted on LB ampicillin plates with 1% glucose and the displayed picture shows that equal amount of cells were spotted for all the samples tested. (C) Quantitative growth assay was done to assess the contribution of the polar filter towards bacterial survival. The effect of expression of wt- and mutant versions of MsDpo4 on the survival of the E. coli were measured and compared. The cells expressing MsDpo4-C47T exhibited 10-fold higher number of Colony Forming Units than cells expressing MsDpo4-WT. The error bars indicate standard deviation ($n = 3$).

To test this hypothesis, two additional mutant versions of the MsDpo4 enzyme were prepared- one wherein the Cys47 residue was mutated to Thr, and another with the double mutation C47T+L14Y. It was observed that the MsDpo4-C47T enzyme exhibited substantially higher sugar selectivity than the wt- enzyme. The double mutant of MsDpo4 with both the steric filter and the polar residue showed almost no ribonucleotide incorporation activity at different concentrations of enzyme and nucleotide (Figure 5). The ability of the mutant versions of MsDpo4 to prevent ribonucleotide incorporation was true for all four rNTPs (Figure 6). There was no difference in the ability of the wt- and mutant enzymes to incorporate dNTPs (Supplementary Figure S2).

To ascertain if the engineered polar residue interacts with the incoming dNTP, the structure of MsDpo4-C47T in complex with DNA (with dG in the templating position) and dCTP was determined to a resolution of 1.8 Å (Supple-
Figure 9. The absence of polar and aromatic steric filter in PolIV results in a drastic reduction in sugar selectivity. (A) Primer extension assays showed that the mutant version of PolIV without the steric and polar filter exhibits a 50-fold enhancement in rCTP incorporation. The error bars indicate standard deviation (n = 3). (B) Primer extension assays showed that the absence of the steric and polar filters did not affect the ability of PolIV to incorporate dCTP. The error bars indicate standard deviation (n = 3). (C) The PolIV-F13A+T43C DNA-dCTP (peach) structure superimposed with PolIV-F13A+T43C DNA-dCTP (violet) structure with an RMSD of 0.26 Å. The position and conformation of the dCTP and rCTP in the two structures are identical.

Figure 10. The contribution of polar and steric filters towards sugar selectivity in B-family DNA polymerases. Y424 and N497 represent the steric and polar filters in PolII (E. coli), a representative member of the B-family of dPolI. (A) Primer extension assays were carried out to compare the ability of PolII-WT, PolII-Y424L and PolII-Y424L+N497A to incorporate rCTPs. These assays show that the polar filter contributes substantially towards ribonucleotide exclusion in this family. (B) Further, primer extension assays showed that the mutations Y424L and Y424L+N497A did not have a significant effect on the ability of PolII to incorporate dCTPs. For both the graphs, the error bars indicate standard deviation (n = 3).

Further, the structure of MsDpo4-L14Y+C47T in complex with DNA (template dG) and incoming dCTP was determined to a resolution of 2.5 Å (Supplementary Table S2E) and compared with that of the similar complex of PolIV (PDB ID: 4IRC (40)). This comparison showed that the orientation and conformation of incoming nucleotide in the two structures were nearly identical (Figure 7D–F). The presence of the steric and polar filters in MsDpo4, therefore, led to a similar interactions as that for PolIV-WT and complete loss of ribonucleotide incorporation activity.

Presence of polar filter reduces cellular lethality due to ectopic expression of MsDpo4

It has been observed previously that, incorporation of rNTPs into genomic DNA, leads to cellular lethality in E.
coli and higher organisms (7,10,13,33,51,52). To assess the importance of the polar filter in ribonucleotide incorporation, growth assays were performed using E. coli as a surrogate host. To enable the recruitment of the MsDpo4 protein to the replication fork in E. coli, the β-clamp binding motif of PolIV (50) (342QMERQLVLGL351) was added to C-terminus of MsDpo4. Wt- and mutant versions (C47T) of this construct were prepared.

It was seen that when the expression of these constructs was induced in E. coli, fewer colonies were obtained in case of wt- protein as compared to the construct bearing the C47T mutation or empty vector (Figure 8A-B). In the quantitative plate assay, cells expressing the C47T mutant exhibited 10-fold higher survival with respect to wild type (Figure 8C). The only difference between the wt- and mutant constructs is the presence of the polar filter in the latter. Hence, the low level of survival observed in the case of wt- enzyme may be due to the higher incorporation of rNTP into the genome.

**PolIV lacking polar and steric filter can incorporate ribonucleotides**

The variant of MsDpo4 with the polar filter showed substantially enhanced sugar selectivity. To observe the effect of removal of the polar filter in PolIV on ribonucleotide incorporation, the polar residue Thr43 was replaced with Cys (corresponding to Cys47 in MsDpo4) and Phe13 (steric filter) was replaced by Ala. The ability of the mutant proteins (PolIV-F13A and PolIV-F13A+T43C) to incorporate rCTP and dCTP was compared to that of PolIV-WT using primer extension assays. The level of rCTP incorporation by PolIV-F13A+T43C was 50-fold higher than the wt- enzyme; while that of the F13A mutant was ~35-fold higher (Figure 9A). However, the mutations did not show a significant difference in the ability of PolIV to incorporate dCTP (Figure 9B). The structure of the complex of PolIV-F13A+T43C with DNA (template G) and incoming rCTP or dCTP was determined to a resolution of 2.77 and 2.44 Å, respectively (Supplementary Table S2G and H). The comparison of the two structures shows that the dCTP and rCTP superimpose well on each other (Figure 9C). These structures show that the loss of polar residue helps accommodation of rCTP in a catalytically competent conformation without any steric clash between the 3′-OH and the enzyme surface. Thus, the mutant version of PolIV without the polar and steric filters exhibits a considerably heightened ability to incorporate ribonucleotides as seen in the case of MsDpo4-WT. These experiments also show that the polar and aromatic filters act in tandem to achieve stringent dNTP selectivity in PolIV-WT.

The polar filter is conserved in B and Y family of DNA Polymerases

The structures of ternary complexes of different dPols (enzyme:DNA:dNTP) available in PDB were analyzed to ascertain if the presence of the polar filter is a general feature of dPols. It was observed that the polar and aromatic residues were conserved in the majority of the dPols belonging to Y- and B- families.

For the Y- family, it was seen that the polar filter is represented by a Ser or Thr residue, and a Tyr or Phe residue is present as the steric filter (Supplementary Figure S3A) (53–56) with both filters present in the fingers domain. In the B-family, Asn (fingers domain) represents the polar filter, and Tyr (palm domain) is present as the steric filter (Supplementary Figure S3B) (29,42,57–60).

To validate the role of the polar filter in ribonucleotide exclusion by B-family dPols, DNA polymerase II (PolII) from E. coli was utilized as a model enzyme. In this enzyme, Tyr424 is present as the steric filter, and Asn497 is predicted to represent the polar filter (Supplementary Figure S3B). Primer extension assays showed that, compared to PolII-WT, the mutant proteins PolII-Y424L and PolII-Y424L+N497A exhibit considerably enhanced ability to incorporate ribonucleotides (Figure 10A). The levels of rCTP incorporation by the single and double mutants were 3 and 4.5 times higher than the wild type. In comparison, the ability of the mutant proteins to incorporate dNTPs was unaffected (Figure 10B). The results show that polar residue Asn497 contributes significantly towards ribonucleotide exclusion by PolII. In summary, the studies on MsDpo4, PolII, and PolII described here point to the presence of a polar filter in dPols that acts in tandem with the steric filter to exclude rNTPs during DNA synthesis.

**DISCUSSION**

Our results evince that, to achieve stringent sugar selectivity, dPols possess two filters. The first one, known as the steric filter, is composed of a bulky aliphatic or aromatic side chain that is present underneath the ribose sugar and will clash with the 2′-OH of the rNTP molecule to reduce the chances of it occupying the active site in a conformation compatible with productive catalysis. The present study shows the existence of a second polar filter, which interacts with the 3′-OH and triphosphate moiety of the incoming nucleotide and orients it closer to the surface of the fingers domain. As a result, the presence of the polar filter ensures that the 2′-OH of an rNTP molecule will clash with the surface of the fingers domain and consequently there is a limited possibility of the rNTP binding in a catalytically competent conformation. The steric filter is present below the sugar, and the polar filter is present close to the 3′-OH and the triphosphate moiety of the incoming nucleotide. Hence, the filters fall on nearly perpendicular planes to each other, and synergize to ensure high sugar selectivity.

The presence of the steric and polar filters that adopt chemically distinct strategies to prevent ribonucleotide incorporation appears to be a conserved feature of the Y- and B-family of dPols. Polθ is a Y-family dPol and exhibits a heightened ability to incorporate ribonucleotides (61). It is believed that this is because it has an extraordinarily spacious active site (30). However, the ability of the enzyme to incorporate ribonucleotides may also be due to the fact that it does not possess a polar filter. An alignment of hPolθ with PolIV (RMSD = 2.36 Å) shows that an Ala residue (Ala49) is present at the position equivalent to the Thr43 residue (Supplementary Figure S4). The alignment of Polθ incorporating rCTP and dCTP (RMSD = 0.19 Å) shows that the rCTP is repositioned away from the aromatic fil-
ter (Supplementary Figure S4D and E) (30). The presence of a polar filter that interacts with the rNTP will prevent such a repositioning. It has been reported before Polη has a heightened ability to incorporate ribonucleotides, and these observations are in line with the present study (61).

The strategy utilized by Y- and B- family dPols to exclude ribonucleotides is distinct from that observed in the case of A, X and C- family members. In A-family, it is suggested that dual steric filters exist in the form of Glu residue below the sugar of the incoming nucleotide and an aromatic residue lining the fingers domain (19,62–64). The two steric filters are positioned such that the 2′-OH of an rNTP molecule will clash with these residues, thus preventing its incorporation (65,66). Based on the structures of DNA polymerase III (PolIII) from *E. coli* and *Geobacillus kaustophilus*, C-family dPols may utilize a histidine residue to perform the function of the steric filter. It was seen that the side chains of His760 and His 1275 in PolIII from *E. coli* and *G. kaustophilus*, respectively (67,68) stack against the ribose sugar of the incoming nucleotide. A number of X-family members exhibit lower selectivity for dNTPs and even though, some members show the presence of an aromatic residue at the position equivalent to the steric filter, the structures show that it does not stack with the ribose sugar (18,69). Instead, it appears that the polypeptide backbone in the region which is present close to the C2′ atom of the sugar acts as a steric filter to provide some level of sugar selectivity (18). For the PrimPol family, it is suggested that the main chain of residues 289–291 in human PrimPol may act as a steric filter to prevent ribonucleotide incorporation (32). Also, Y100 present in the fingers domain, which does not stack with the sugar of the incoming nucleotide is known to play an important role in nucleotide selectivity (70,71). The members of the RT family of dPols do possess the steric filter to reduce ribonucleotide incorporation (72–74). The D-family of dPols represents the least studied family of dPols and strategy utilized by these enzymes to prevent ribonucleotide incorporation is unclear (75).

B-family dPols are highly accurate and responsible for DNA synthesis during replication and repair, especially in eukaryotes (76–80). In humans, members of the B-family include Polα, Polβ, Polε, and Polκ. A thorough examination of different databases such as NIH-NCI GDC data portal, NCB Clinv and BIH-DB SNP Short Genetic Variation, revealed a correlation between mutations at the polar filter site and disease in human dPols δ and ξ (Supplementary Table S1). Thermostable members of B-family dPols have been used as enzymes for PCR-based applications (81,82). The results presented in the present study may be useful in the development of engineered enzymes that can directly amplify RNA without the need for cDNA formation. Such engineered dPols may be a valuable tool for a number of applications in biotechnology (83).

### DATA AVAILABILITY

The structure factors and the refined co-ordinates have been deposited in the PDB with the following codes: 6JUL MsDpo4-WT DNA(dG):dCTP, 6JUM MsDpo4-C47T DNA(dG):dCTP, 6JUN MsDpo4-C47T+L14Y DNA(dG):dCTP, 6JUP PolIV-F13A+T43C DNA(dG):dCTP, 6JUQ PolIV-F13A+T43C DNA(dG):dCTP, 6JUR MsDpo4-L14Y DNA(dG):rCTP and 6JUS MsDpo4-WT DNA(dG):rCTP.

### SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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