Insights into the Determination of the Templating Nucleotide at the Initiation of φ29 DNA Replication*

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Background: The initiation of replication in φ29 mainly occurs opposite the 3’ penultimate nucleotide of the template.

Results: Mutations in the TP residue Phe-230 changed the specificity for the templating nucleotide.

Conclusion: The results suggest a role for Phe-230 in determining the templating nucleotide.

Significance: The results shed light on the positioning of the template nucleotide in organisms that initiate with protein-priming mechanism.

Bacteriophage φ29 from Bacillus subtilis initiates replication of its terminal protein (TP)-DNA by a protein-priming mechanism. To start replication, the DNA polymerase forms a heterodimer with a free TP that recognizes the replication origins, placed at both 5’ ends of the linear chromosome, and initiates replication using as primer the OH-group of Ser-232 of the terminal protein (TP)-DNA by a protein-priming mechanism and has become the paradigm in the study of TP-DNA replication. Phage φ29 initiates replication by a protein-priming mechanism and has become the paradigm in the study of TP-DNA replication. Phage φ29 has a linear dsDNA 19285 base pairs (bp) long with a 6-bp inverted terminal repeat (3’-TTTTCAT) that, together with the terminal 12-bp and with the phage-encoded TP covalently attached at each 5’ DNA end, assemble a minimal replication origin (9). The DNA polymerase and a free TP molecule (primer TP) form a heterodimer (10) that recognizes the replication origin. Then, the DNA polymerase catalyzes the incorporation of the initiating dAMP to the OH-group of the TP residue Ser-232 (8), a reaction directed by the 3’ penultimate dTMP of the template strand. Once the ini-

DNA polymerases synthesize DNA exclusively in the 5’-3’ direction and are unable to start de novo DNA synthesis, compelling the use of a priming 3’ OH-group generally provided by a short RNA or DNA molecule. These features of the DNA polymerases create the so-called end replication problem of linear chromosomes according to which, after successive rounds of replication, the short region of unreplicated ssDNA at the end of the chromosome would lead to a progressive shortening of the daughter DNA (1). Many mechanisms have evolved to maintain the DNA ends of linear chromosomes through subsequent replications rounds, most of them making use of the presence of repetitive sequences at the ends that allow the formation of long concatemers, as in phage T7 (2), to circularize, as in phage λ (3), or to form hairpin loops to fill the incomplete 5’ ends, as in poxvirus (4–6) or in the linear plasmid of Borrelia (7).

Several prokaryotic and eukaryotic viruses, as well as linear plasmids from bacteria, fungi, higher plants, and Streptomyces spp. have solved the end replication problem by using a protein, called terminal protein (TP),⁶ to prime DNA synthesis from the end of their linear genomes (TP-DNA) (8). Thus, a specific amino acid of the TP (serine, threonine, or tyrosine) provides the OH-group to prime initiation of DNA replication, the TP becoming covalently linked to the 5’ ends of the nascent DNA (parental TP).

Bacillus subtilis phage φ29 initiates replication by a protein-priming mechanism and has become the paradigm in the study of TP-DNA replication. Phage φ29 has a linear dsDNA 19285 base pairs (bp) long with a 6-bp inverted terminal repeat (3’-TTTTCAT) that, together with the terminal 12-bp and with the phage-encoded TP covalently attached at each 5’ DNA end, assemble a minimal replication origin (9). The DNA polymerase and a free TP molecule (primer TP) form a heterodimer (10) that recognizes the replication origin. Then, the DNA polymerase catalyzes the incorporation of the initiating dAMP to the OH-group of the TP residue Ser-232 (8), a reaction directed by the 3’ penultimate dTMP of the template strand. Once the ini-

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⁶ The abbreviations used are: TP, terminal protein; TPR, terminal protein region.
Template Nucleotide in Initiation of φ29 DNA Replication

The crystallographic structure of φ29 TP in complex with the DNA polymerase (19) showed that it consists of a disordered N-terminal domain (residues 1–73) responsible for DNA binding and TP nucleoid localization (20), an intermediate domain (residues 74–172) that confers specificity to the interaction with the DNA polymerase, connected through a hinge region to the C-terminal priming domain (residues 173–266) bearing the palm (containing the catalytic and DNA ligand residues), fingers (containing the dNTP ligands), and thumb (which confers stability to the primer) (15). Additionally, the polymerization domain has two insertions called terminal protein regions 1 and 2 (TPR1 and TPR2), specifically present in the protein-primed DNA polymerases subgroup (16, 17). The arch-like structure formed by the TPR2 insertion and the thumb subdomain confers a toroidal shape to the polymerization domain that encircles the nascent DNA, being responsible for the high processivity of the enzyme. In addition, the TPR2 subdomain together with the fingers subdomain and the exonuclease domain wrap the template strand, acting as a “molecular wedge” to unwind the parental DNA strands during replication (18). Despite the physical separation between the exonuclease and polymerization domains, these two opposite activities must act in concert to achieve a productive and accurate replication reaction.

The 3′-5′ exonuclease activity of Vent DNA polymerase dissociates from the TP (13) and the same DNA polymerase molecule catalyzes processive chain elongation via a strand displacement mechanism to fulfill TP-DNA replication (14). These two features of φ29 DNA polymerase, processivity and strand displacement capacity, enable the enzyme to replicate the viral genome without requiring the assistance of processivity and unwinding factors, unlike most replicases.

Previous studies performed with chimerical TPs allowed to describe (26). The incubation mixture contained, in 25 mM Tris-HCl, pH 7.5, 10 mM MgCl₂, 20 mM ammonium sulfate, 1 mM dithiothreitol (DTT), 4% (v/v) glycerol, 1.6 mM φ29 TP as template, 0.1 mg/ml of bovine serum albumin (BSA), 0.2 μM dATP (1 μCi of [α-32P]dATP), 13 nM either wild-type or mutant φ29 TP, and 13 nM φ29 DNA polymerase. Samples were incubated for 10 min at 30 °C. Reactions were stopped by adding 10 mM EDTA, 0.1% SDS, and the samples were filtered through Sephadex G-50 spin columns in the presence of 0.1% SDS and further purified by SDS-PAGE in 12% polyacrylamide gels. Quantification was done by densitometric analysis of the labeled band corresponding to the TP-dAMP complex detected by autoradiography.

For the reconstituted DNA the reaction was carried out essentially as described above for the TP-DNA but in the presence of 13 nM of either wild-type or mutant TP, 13 nM D12A/D66A exonuclease-deficient φ29 DNA polymerase, 0.4 nM

Experimental Procedures

Nucleotides and DNAs—Unlabeled nucleotides were supplied by Amersham Biosciences Pharmacia. [α-32P]dATP (3000 Ci/mmmol) and [α-32P]dTTP (3000 Ci/mmmol) were supplied by PerkinElmer Life Sciences. Oligonucleotides were obtained from Invitrogen.

Site-directed Mutagenesis of φ29 TP—TP mutants were obtained using the QuikChange site-directed mutagenesis kit provided by Stratagene, using as template the plasmid pT7-3 that contains the viral gene 3 coding for the TP (22). The presence of the desired mutations, as well as the absence of additional ones was determined by sequencing the entire gene.

Proteins—TP mutants were expressed in Escherichia coli BL21(DE3) cells harboring gene 3 cloned into plasmid pT7-3 and further purified, essentially as described for the wild-type TP (23). Both, the wild-type φ29 DNA polymerase and the exonuclease-deficient mutant D12A/D66A (24) were expressed in E. coli BL21(DE3) cells harboring gene 2 cloned into plasmid pJLPM (a derivative of pT7–4w2 containing the ribosomal binding site; the nucleotide T was changed into G in the Shine-Dalgarno sequence) and further purified essentially as described (25).

Preparation of Reconstituted TP-DNA—The 3′ ends of TP-DNA (6 nM) were partially degraded by the 3′-5′ exonuclease activity of Vent DNA polymerase in its reaction buffer during 15 min at 75 °C. Then, the degradation product was hybridized in an incubation mixture that contained 20 nM of the indicated 29-mer single-stranded oligonucleotide, 1.8 mM EDTA, 150 mM NaCl, and 0.02% Tween 20 during 5 min at 75 °C. The temperature was allowed to decrease to 55 °C during 99 min and remain for 5 min at 55 °C. Then, the temperature was lowered until 4 °C during 15 min.

Protein-primed Initiation Assay—The ability to carry out the initiation step of TP-DNA replication was analyzed as described (26). The incubation mixture contained, in 25 μl, 50 mM Tris-HCl, pH 7.5, 10 mM MgCl₂, 20 mM ammonium sulfate, 1 mM dithiothreitol (DTT), 4% (v/v) glycerol, 1.6 mM φ29 TP-DNA as template, 0.1 mg/ml of bovine serum albumin (BSA), 0.2 μM dATP (1 μCi of [α-32P]dATP), 13 nM either wild-type or mutant φ29 TP, and 13 nM φ29 DNA polymerase. Samples were incubated for 10 min at 30 °C. Reactions were stopped by adding 10 mM EDTA, 0.1% SDS, and the samples were filtered through Sephadex G-50 spin columns in the presence of 0.1% SDS and further analyzed by SDS-PAGE in 12% polyacrylamide gels. Quantification was done by densitometric analysis of the labeled band corresponding to the TP-dAMP complex detected by autoradiography.
reconstituted DNA, 1 mM MnCl₂, and 0.2 μM dATP (1 μCi of [α-32P]dATP). Samples were incubated for 30 min at 30 °C.

In the case of the TP-dAMP or TP-dTMP formation using as template 29-mer ssDNA instead of TP-DNA the reaction was carried out essentially as described above for the TP-DNA but in the presence of 2 μM of the indicated oligonucleotide, 130 nM of either wild-type or mutant TP, 60 nM of the D12A/D66A exonuclease-deficient φ29 DNA polymerase, 1 mM MnCl₂, and either 0.2 μM dATP (1 μCi of [α-32P]dATP) or 0.2 μM dTTP (1 μCi of [α-32P]dTTP). Samples were incubated for 1 h at 12 °C.

TP-DNA Replication Assay—The replication assay was performed as described (26). The incubation mixture contained, in 25 μL, 50 mM Tris-HCl, pH 7.5, 10 mM MgCl₂, 20 mM ammonium sulfate, 1 mM DTT, 4% (v/v) glycerol, 0.1 mg/ml of BSA, 20 μM each dNTP and [α-32P]dATP (1 μCi), 13 nM of either wild-type or mutant TP, 13 nM DNA polymerase, and 1.6 nM φ29 TP-DNA. After 10 min at 30 °C, the reaction was stopped by adding 10 mM EDTA, 0.1% SDS, and the samples were filtered through Sephadex G-50 spin columns in the presence of 0.1% SDS. The samples were analyzed by electrophoresis in SDS-12% polyacrylamide gels (360 × 280 × 0.5 mm) to obtain enough resolution to distinguish the TP bound to the first elongation product.

Results

Effect of the φ29 Parental TP in the Positioning of the First Templating Nucleotide—By using single-stranded synthetic oligonucleotides with the sequence of the replication origins (11) and TP-free dsDNA fragments containing the φ29 DNA terminal sequences (28) it was concluded that initiation of replication occurs mainly opposite the second nucleotide at the 3’ end of the template. However, the use of TP-free DNA ends precluded analysis of the influence of the parental TP in positioning the templating nucleotide during the initiation reaction. To address this question, full TP-DNA was partially digested with the 3’-5’ exonuclease activity of Vent DNA polymerase giving rise to TP-ssDNA regions (see scheme in Fig. 1). Replication origins were reconstituted after hybridization of 29-mer oligonucleotides containing either the wild-type sequence corresponding to the right replication origin (TTT) or single substitutions of the first (ATT), second (TAT), or third (TTA) thymine for adenine (see Fig. 1). These hybrid molecules were further used as substrate for the initiation reaction. Fig. 2A shows that, as expected, in the presence of the initiator [α-32P]dATP, the single-stranded 29-mer oligonucleotides TTT, ATT, and TTA supported formation of the initiation product TP-dAMP by the exonuclease-deficient φ29 DNA polymerase D12A/D66A (24) (the use of this mutant prevents the degradation of the template ssDNA). The small amount of TP-dAMP complex obtained with the template containing the TAT sequence is due to some initiation reaction directed by the third T (11). Essentially the same result was obtained with the reconstituted origins (Fig. 2B). The origins containing the

FIGURE 1. A, schematic representation of reconstituted TP-DNA. φ29 TP-DNA is represented with a TP linked at each 5’ end. At the first step, the 3’-5’ exonuclease activity of Vent DNA polymerase degrades the 3’ ends of the TP-DNA. The resulting ssDNA region is further hybridized to 29-mer single-stranded oligonucleotides containing the φ29 right replication origin sequence and variants in the 3’ terminal thymines. B, 29-mer oligonucleotides containing wild-type and variants of the φ29 right replication origin sequence. The thymines changed into adenine are indicated in black letters.

Template Nucleotide in Initiation of φ29 DNA Replication

A

| TP | AAAATGACCCCATTCACATGAT-3′ |
|---|---|
| 3′ | TTTCAATGGGACGCAT-GGT |

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GATAGTTGGCTATTGGAGAATGGATGATGATCTAGCTACATCAGACACGGACATGGATGAAT (3′)

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CTATCCAGCAGTTACCCCTTTACACATTACAACACGGACATGGATGAAT (TP)

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Hybridization

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TGTTGATATTTGCTTGGATCCCCCTATT (3′)

B

| Template Nucleotide | 29 Parental TP in the Positioning of the First Templating Nucleotide |
|---------------------|---------------------------------------------------------------|
| TTT | TTTTACATCAGCTGTCTTGGATGGT |
| ATT | ATTTCATCCAGCTGCTTGGATGGT |
| TAT | TATTCCATCCAGCTGTCTTGGATGGT |
| TTA | TTACCATCCAGCTGCTTGGATGGT |
The aromatic residue could be playing some role in the selection from the initiator residue. Thus, it is tempting to speculate that the priming domain (32) and the aromatic residue is four positions away between the last and penultimate helixes of the domain (19). Although the structure of this priming loop is disordered in the crystallographic structure of the φ29 DNA polymerase-TP complex, it should be positioned at the polymerization active site in this heterodimer to allow Ser-232 to prime the formation of the initiation product by the DNA polymerase. As shown in Fig. 3A the alignment of the corresponding priming loop of the TPs from different viruses that initiate the replication of their genomes with a TP shows the presence of aromatic residues close to the initiating (priming) residue. Interestingly, there seems to be a correlation between the distance of the aromatic residue to the priming amino acid and the initiation position. Accordingly, in phages φ29 and GA-1 that initiate opposite the second 3′ T (11, 29) the aromatic residue is located two positions before the priming residue. In Cp-1 and PRD1 an aromatic residue is three positions away from the initiator amino acid and the initiation reaction is directed by the third (30) and fourth (31) 3′ nucleotide, respectively. In adenovirus the replication occurs opposite the fourth nucleotide at the 3′ end of the template (32) and the aromatic residue is four positions away from the initiator residue. Thus, it is tempting to speculate that the aromatic residue could be playing some role in the selection of the templating nucleotide.

To study the role of the aromatic residue in positioning the template nucleotide we made site-directed mutagenesis in the priming loop of φ29 TP obtaining three mutants: E229F/F230V, E228F/F230V, and F230V (see Fig. 3A). In the single mutant F230V the aromatic residue Phe-230 of the priming loop is removed, whereas in the double mutants the loop is moved one (E229F/F230V) or two (E228F/F230V) positions N terminally. In the case of the mutant E229F/F230V the aromatic residue is in a position similar to that occupied in the TPs from PRD1 and Cp-1 and in the case of the mutant E228F/F230V the aromatic residue was moved N terminally to a position similar to that occupied in the TP from adenovirus.

To determine whether the introduced mutations affected the template nucleotide that directs the initiation reaction, the 29-mer single-stranded oligonucleotides TTT, ATT, TAT, and TTA were used as template. To detect the formation of TP-DTMP when the templating nucleotide is an adenine, [α-32P]dTMP was used as the initiator nucleotide. As it can be observed in Fig. 3B, the wild-type TP primes initiation mainly opposite the second 3′ nucleotide of the template (TAT) although some initiation reaction also takes place opposite the third 3′ nucleotide, as described (11). In contrast, with mutant TPs F230V, E229F/F230V, and E228F/F230V formation of TP-DTMP was mainly directed by the third nucleotide of the template strand (TTA). These results indicated that when the aromatic residue Phe-230 was removed or the distance to the priming residue was increased one (E229F/F230V) or two (E228F/F230V) positions the initiation occurred in a more internal position (third instead of second 3′ terminal nucleotide), suggesting that the location of aromatic residue Phe-230 within the priming loop could be a determinant to place the templating nucleotide used to direct the φ29 protein-primed initiation reaction.

Taking into account the presence of an additional aromatic residue in the φ29 TP priming loop (Phe-227) we analyzed whether it was playing any role in the initiation reaction. To address this question, φ29 TP residue Phe-227 was changed into valine (F227V mutant) and further evaluated in the initiation reaction described above. As it can be observed in Fig. 3C, when this mutant was used as primer, the initiation mainly occurred opposite the second nucleotide of the template and to some extent opposite the third nucleotide, like with the wild-type TP, indicating that the aromatic residue Phe-227 does not seem to contribute to the placement of the 3′ terminus of the template strand at the polymerization active site.

To study if the residues changed in the mutant TPs could be playing a role in further steps of φ29 TP-DNA replication, the formation of the TP-dAMP initiation complex and its further elongation was evaluated using as template φ29 TP-DNA. As shown in Fig. 4A and Table 1, when the DNA polymerase used the TP mutants as primer the formation of the TP-dAMP was impaired, especially when residue Phe-230 was removed or moved away from the priming residue. As mentioned in the Introduction, once formation of the TP-dAMP product was catalyzed, the same DNA polymerase elongates it via strand displacement to yield full-length φ29 TP-DNA. To ascertain to what extent the mutations introduced at the TP affect the replication process, assays were carried out using a minimal replication system based on φ29 TP-DNA, DNA polymerase, and TP (14). As observed in Fig. 4B and Table 1, when mutant TPs were used as primers, defects in the replication reaction were

**FIGURE 2.** Formation of the TP-dAMP initiation complex using as template ssDNA (A) and reconstituted TP-DNA (B). The assays were carried out essentially as described under “Experimental Procedures” in the presence of 1 mM MnCl₂. The position of the TP-dAMP is indicated. TP-DNA, initiation reaction using as template intact TP-DNA; C (control), initiation reaction using as template TP-DNA incubated at the same temperature and in the same reaction mixture as reconstituted DNA but in the absence of Vent polymerase and oligonucleotide; O (oligonucleotide), initiation reaction using as template the amount of oligonucleotide used in the hybridization but without TP-DNA; D (degraded), initiation reaction using as template TP-DNA with the 3′ ends degraded but not hybridized to the oligonucleotide.

TTT, ATT, and TTA sequences gave rise to the formation of the TP-dAMP initiation product, the TAT origin being the one that supported the lowest initiation reaction. The initiation levels observed with the reconstituted origins were not due either to potentially non-hybridized oligonucleotides (lane O) or to non-degraded TP-DNA (lane D). The results obtained indicate that the parental TP is not playing a significant role in determining the internal nucleotide used as template of the initiation reaction.

**Role of the Priming Loop Aromatic Residue Phe-230 in Positioning the Internal 3′ Nucleotide Used as Template**—Previous assays with chimerical TPs indicated that the priming domain is responsible for placing the templating nucleotide at the polymerization active site during initiation (21). The TP priming domain (residues 173–266) is comprised of a four-helix bundle structure, the priming loop (residues 227–233) being located close to the initiating (priming) residue. Interestingly, there seems to contribute to the placement of the 3′ terminus of the template (TAT) although some initiation reaction also takes place opposite the third 3′ nucleotide, as described (11). In contrast, with mutant TPs F230V, E229F/F230V, and E228F/F230V formation of TP-DTMP was mainly directed by the third nucleotide of the template strand (TTA). These results indicated that when the aromatic residue Phe-230 was removed or the distance to the priming residue was increased one (E229F/F230V) or two (E228F/F230V) positions the initiation occurred in a more internal position (third instead of second 3′ terminal nucleotide), suggesting that the location of aromatic residue Phe-230 within the priming loop could be a determinant to place the templating nucleotide used to direct the φ29 protein-primed initiation reaction.

Taking into account the presence of an additional aromatic residue in the φ29 TP priming loop (Phe-227) we analyzed whether it was playing any role in the initiation reaction. To address this question, φ29 TP residue Phe-227 was changed into valine (F227V mutant) and further evaluated in the initiation reaction described above. As it can be observed in Fig. 3C, when this mutant was used as primer, the initiation mainly occurred opposite the second nucleotide of the template and to some extent opposite the third nucleotide, like with the wild-type TP, indicating that the aromatic residue Phe-227 does not seem to contribute to the placement of the 3′ terminus of the template strand at the polymerization active site.
even more pronounced that during the initiation, mutant TPs E229F/F230V, E228F/F230V, F230V, and F227V being 11-, 100-, 27-, and 4-fold were less efficient in supporting DNA replication than the wild-type protein.

Furthermore, we investigated whether the heterodimer formed by the mutant TPs and DNA polymerase was able to recover the first and second 3\end{superscript}/H11032 nucleotides. To this end truncated elongation assays were performed in the presence of TP-DNA, dATP, dGTP, and ddTTP, to allow elongation to take place only up to the first A of the template. As mentioned above, initiation with the wild-type TP occurs using as template the penultimate 3\end{superscript}/H11032 T and then the TP-dAMP product translocates one position backwards to recover the information of the 3\end{superscript}/H11032 terminal T in a mechanism called sliding-back (11). Then, when the wild-type TP was used as primer, the elongation continued up to the incorporation of the ddTMP opposite the fifth nucleotide (Fig. 5), indicating that all the positions of the template, including the first nucleotide, were replicated. As it can be observed, 40% of the elongated initiation products stopped at the fourth position, most likely due to the inefficiency displayed by \(\phi29\) DNA polymerase to insert ddNMPs. When mutant F227V was used as primer, the DNA polymerase gave rise to a 5-nucleotide long product (60%), indicating a recovery of the first 3\end{superscript}/H11032 terminal nucleotide as it occurred with the wild-type TP.

In contrast, when aromatic residue Phe-230 was removed (F230V) or the distance was increased one (E229F/F230V) or two (E228F/F230V) positions the elongation mainly stopped opposite the fourth nucleotide (70, 55, and 70%, respectively). This suggests that after the initiation opposite the third nucleotide, most of the initiation products performed only a sliding-back step between the third and second positions. A possible explanation is that initiation at the third 3\end{superscript}/H11032 nucleotide may affect the performance of the additional sliding-back step between the second and the first position that would be required to recover the 3\end{superscript}/H11032 end information.

**Discussion**

The first step of \(\phi29\) DNA replication requires the formation of a heterodimer between the DNA polymerase and the primer TP that further recognizes the replication origins located at both ends of the linear genome. The DNA polymerase catalyzes the covalent linkage of the initial dAMP to the hydroxyl group of Ser-232 of the TP and its further elongation (8). Similarly, TP-DNA replication of bacteriophages PRD1 (31, 33), Cp-1 (30), and the \(\phi29\)-related phages GA-1 (22, 29) and Nf (22, 34) was also shown to occur by a protein-primed mechanism, involving its corresponding DNA polymerase and TP. Initiation of \(\phi29\) DNA replication occurs at the second 3\end{superscript}/H11032 nucleotide of the template. The first nucleotide is then recovered by a sliding-back mechanism to fulfill TP-DNA replication (11). After incorporation of the 10th nucleotide there is a dissociation of the DNA polymerase (13) that continues replication via a strand displacement mechanism to fulfill TP-DNA replication (14).

![Figure 3](image-url)
The complex between the replicative DNA polymerase and a free TP molecule interacts with the replication origins by specific recognition of the parental TP and DNA sequences. The presence of the parental TP, which is covalently linked to the 5’-ends of the non-template strand by a previous cycle of replication, is the main signal to be recognized by the polymerase-TP complex for the initiation of replication, because when terminal DNA fragments lacking parental TP were used as templates, the initiation reaction decreased 6–10-fold with respect to the activity obtained with TP-DNA (34, 35). Due to its important role in recruiting the φ29 DNA polymerase/TP heterodimer to the origins we analyzed whether the parental TP was condition-

TABLE 1
Activity of wild-type and mutant φ29 TPs

| φ29 TP          | TP-dAMP formation* | TP-DNA replication* |
|-----------------|--------------------|---------------------|
| Wild-type       | 100                | 100                 |
| E229F/F230V     | 26.5 ± 9.7         | 9.1 ± 3.5           |
| E228F/F230V     | 11.8 ± 6.5         | 1 ± 0.5             |
| F230V           | 15.1 ± 7.3         | 3.7 ± 1.9           |
| F227V           | 53.2 ± 18.4        | 28.2 ± 4.7          |

* TP-dAMP formation using φ29 TP-DNA as template.

Previous assays showed that the priming domain is the main structural determinant that dictates the internal 3’ nucleotide used as template during initiation (21). Here, we have analyzed the role of the aromatic residue Phe-230 of the TP priming loop, proximal to the priming Ser-232. The results showed that mutations in the aromatic residue Phe-230 changed the specificity for the templating nucleotide, as initiation occurred mainly opposite the third 3’ T instead of the second one. An appealing hypothesis would be that this aromatic residue acts as a structural barrier that limits the internalization of the 3’ end of the template strand, conditioning the placement at the polymerization active site of the correct templating nucleotide. Fig. 6 shows a schematic of the priming-loop of the φ29 and adenovirus TPs, with the corresponding aromatic residues Phe-230 and Tyr-576, respectively, and how they could be interacting with the 3’ terminal base. Such a contact would block further internalization, allowing the second and fourth position in φ29 and adenovirus, respectively, to be placed at the polymerization active site.

Residues Phe-230 and Phe-227 could be also involved in the proper stabilization/orientation of the priming residue Ser-232 at the polymerization active site of DNA polymerase, explaining the lower levels of initiation and replication exhibited by the mutants, specially F230V. During the transition step, whereas the priming domain must move in response to the translocation of the DNA after nucleotide incorporation, the intermediate domain would be in a fixed orientation on the polymerase by means of stable contacts with the TPR1 subdomain (36). Some displacements of the priming domain appear possible without dissociation due to the flexibility of the hinge region that connects the intermediate and priming domains. After incorpora-
tion of 6–7 nucleotides the proximity of the priming Ser to the hinge region would impede a further priming domain rotation, promoting complex dissociation (19). The ability of mutant TPs F230V, E229F/F230V, and E228F/F230V to accomplish the first sliding-back step better than the second one, could indicate that the \( \phi 29 \) DNA polymerase/TP heterodimer is structurally adapted to perform mainly one backward movement of the initiation product. During the sliding-back a set of interactions between the TP priming domain and the DNA polymerase TPR2 and thumb subdomains should be broken and new contacts should be established. This fact could compel the polymerase to move forward, explaining why, although mutant TPs can prime initiation opposite the third T, the recovery of the terminal 3’ T is hindered.

**Author Contributions**—A. P. conducted the experiments, analyzed the results, and wrote most of the paper. J. M. L. performed the pJLPM plasmid. E. L. performed the TP mutant E228F/F230V. L. V. purified the TP mutants. M. S. and M. V. conceived the idea for the project, discussed the results, and wrote the paper with A. P.

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