Catalytically inactive T7 DNA polymerase imposes a lethal replication roadblock

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Bacteriophage T7 encodes its own DNA polymerase, the product of gene 5 (gp5). In isolation, gp5 is a DNA polymerase of low processivity. However, gp5 becomes highly processive upon formation of a complex with Escherichia coli thioredoxin, the product of the trxA gene. Expression of a gp5 variant in which aspartate residues in the metal-binding site of the polymerase domain were replaced by alanine is highly toxic to E. coli cells. This toxicity depends on the presence of a functional E. coli trxA allele and T7 RNA polymerase-driven expression but is independent of the exonuclease activity of gp5. In vitro, the purified gp5 variant is devoid of any detectable polymerase activity and inhibited DNA synthesis by the replisomes of E. coli and T7 in the presence of thioredoxin by forming a stable complex with DNA that prevents replication. On the other hand, the highly homologous Klenow fragment of DNA polymerase I containing an engineered gp5 thioredoxin-binding domain did not exhibit toxicity. We conclude that gp5 alleles encoding inactive polymerases, in combination with thioredoxin, could be useful as a shutoff mechanism in the design of a bacterial cell-growth system.

The minimal replication machinery of bacteriophage T7 provides a superb model system to dissect the molecular mechanisms of DNA replication. Only four proteins are required to reconstitute coordinated leading- and lagging-strand DNA synthesis in vitro (1–3). The primase-helicase, gp4, unwinds dsDNA via its C-terminal helicase domain, providing the templates for DNA synthesis; the N-terminal primase domain of gp4 synthesizes tetraribonucleotides to prime lagging-strand synthesis. T7 DNA polymerase holoenzyme, a complex of the phage-encoded DNA polymerase, gp5, and its host-encoded processivity factor, thioredoxin (Trx), is responsible for the synthesis of leading and lagging strands. Finally, the T7 ssDNA-binding protein, gp2.5, interacts with ssDNA, T7 DNA polymerase, and primase-helicase gp4 to coordinate the enzymatic events between the leading and lagging strands (2–4).

All members of the DNA polymerase superfamily employ two invariant carboxylic acid-containing side chains to coordinate catalytic metal ions in the palm domain of the polymerase active site (5, 6). The A family of polymerases includes T7 gp5, the eukaryotic mitochondrial polymerase γ, and Escherichia coli DNA Pol I. In E. coli Pol I, the carboxylates are present as aspartate 705 and aspartate 882. Mutation of either of these residues to alanine results in undetectable polymerase activity (7). Mechanistic experiments show that these residues function in steps prior to the chemical step (8). Aspartate 882 functions in the finger-closing transition, whereas aspartate 705 functions to facilitate entry of a metal associated with the dNTP delivered into the active site. Although no similar mutants for gp5 have been identified, it seems likely that metal-coordinating aspartate residues in gp5 will play an analogous role as asp 705 and 882 in DNA Pol I because of the conservation of the nucleotidyl transfer step among all DNA polymerases (9, 10).

While investigating the transfer of primers from T7 DNA primase to T7 DNA polymerase (11), we constructed a gp5 variant that lacks polymerase activity. The alteration consisted of replacing the two aspartate residues, corresponding to those described above in the metal-binding site for E. coli DNA Pol I, with alanine. (8, 12). Expression of this inactive polymerase (gp5-D475A/D654A) was deleterious to E. coli, as evidenced by a lack of growth of E. coli containing gene 5 expression plasmids. This toxic effect depended on the co-expression of thioredoxin by E. coli. In the present study we show that both in vivo and in vitro, the inactive T7 gp5 and thioredoxin form a complex that inhibits DNA synthesis mediated by the E. coli and T7 replisomes.

Results

Polymerase-inactive gp5 is toxic to E. coli in a thioredoxin-dependent manner

The X-ray crystallographic structure of T7 DNA polymerase in complex with a primer/template shows that aspartate residues at position 475 and 654 in gp5 coordinate the two Mg2+ ions observable in the polymerase domain (Fig. 1) (12). We constructed gp5 variants with aspartate-to-alanine substitution at positions 475 and 654 and examined their ability to complement a T7 phage with a deletion in gene 5 (T7Δ5). None of the gp5 alleles we constructed containing alanine substitutions of the invariant Mg2+-binding active site aspartates were able to complement T7Δ5 (Table 1), suggesting that the mutations disrupted the enzymatic activity of gp5.

To our surprise we were unable to transform the E. coli expression strain BL21 (DE3) with plasmids containing mutant gene 5 alleles under the control of the T7 promoter. Interestingly, transformation was successful with E. coli A307 (DE3) (data not shown), a protein overexpression strain that lacks the...
inactive lacUV5 promoter and is equally capable of expressing plasmid-encoded genes under the control of a T7 promoter as the parental BL21 (DE3) (data not shown). Plasmids harboring polymerase-inactive gp5 alleles can be transformed into *E. coli* BL21(DE3) ΔtrxA as efficiently as the plasmid encoding WT gp5 (Table 1). This result confirms the necessity of thioredoxin in *E. coli* for inhibition of *E. coli* growth by expression of polymerase-inactive gp5 (gp5-D475A/D654A) (Table 1). Transformation efficiency in *E. coli* BL21, a strain that lacks the gene for T7 RNA polymerase in its chromosome, was also examined. This strain would be unable to express the plasmid-encoded gene 5, and comparison of transformation efficiency shows that the lack of expression of gp5-D475A/D654A or gp5-D475A does not produce a toxic effect, suggesting that the toxicity requires expression of mutant gp5. However, we observed a lower transformation efficiency of BL21 with a plasmid encoding the single gp5-D654A mutation (Table 1).

The expression levels resulting from leaky basal expression of T7 RNA polymerase in (DE3) host cells are sufficient to drive the production of gp5-D475A/D654A to toxic levels, because isopropyl β-D-thiogalactoside was not present to induce high levels of expression. Toxicity is observed only when host cells containing both the trxA gene and a λDE3 lysogen are transformed with plasmids encoding gp5-D475A/D654A (Table 1). The genetic requirement for trxA for toxicity and the strong interaction between the two proteins (13) suggest that gp5-D475A/D654A binds *E. coli* trx in vivo to form a protein with a deleterious effect on *E. coli* growth. Additionally, a single substitution of alanine for aspartate at positions 475 or 654 in gp5 is sufficient for inhibition of *E. coli* growth (Table 1). The 3′–5′ exonuclease activity of gp5 is not required for the toxic effect, because gene 5 alleles expressing gp5 with or without exonuclease activity exhibit toxicity (data not shown).

Further, no toxicity is observed in control experiments when *E. coli* is transformed by a polymerase-inactive version of *E. coli* Klenow fragment (KF-D705A/D882A-TBD) bearing an insertion of the gp5 thioredoxin-binding domain (TBD), which we previously showed confers increased processivity (Table 1) (14). The observation that KF-D705A/D882A-TBD did not exhibit toxicity could be because although this chimeric protein has enhanced processivity in the presence of thioredoxin compared with WT Klenow fragment, it binds thioredoxin with a 10-fold lower affinity, its polymerase activity is only stimulated ~8-fold by thioredoxin (compared with >2,500-fold for gp5), and it shows a 3-fold lower specific activity than gp5 in the presence of thioredoxin (14).

In addition we found that gp5-D475A/D654A is similarly toxic for T7 phage growth. WT T7 phage is unable to grow in *E. coli* cells containing an expression plasmid bearing alleles encoding polymerase-inactive variants of gp5, whereas the presence of an expression plasmid with WT gp5 shows no inhibition of growth (Table 1).

gp5-D475A/D654A is devoid of polymerase activity and inhibits DNA synthesis mediated by T7 and *E. coli* replisomes

We expressed and purified gp5-D475A/D654A as well as an exonuclease-deficient variant (gp5-D5A/D65A) as well as an exonuclease-deficient variant (gp5-D5A/D65A/D475A/D654A)
In E. coli A307 (DE3), an expression strain in which the endo-
geneous trxA gene is deleted. Substitution of Asp475 and Asp654 to 
Ala selectively inactivated the polymerase activity of gp5. Both 
purified gp5-D475A/D654A versions have no detectable poly-
merase activity (<0.2%) as compared with WT T7 DNA poly-
merase using primed M13 (Fig. 2A). The selectivity of the muta-
tions in targeting polymerase activity is underscored by the 3’→5’
exonuclease activity displayed by purified exonuclease-compe-
tent gp5-D473A/D654A (Fig. 2B).

A plausible explanation for the inhibition of E. coli growth 
by the presence of an inactive T7 DNA polymerase is that the 
inhibition results from an interference of the E. coli replisome 
by the inactive polymerase. To determine whether this is the 
case, the effect of T7 gp5-D475A/D654A on an in vitro E. coli 
replication system was examined. This reconstituted system 
results in coordinated synthesis of leading and lagging 
strands, with rates comparable with those determined for E. coli 
DNA replication in vivo (15). The addition of a complex 
of gp5-D475A/D654A and Trx into DNA replication reac-
tions catalyzed by the E. coli replisome led to a greater than 
95% reduction in DNA synthesis (Fig. 3A). We conclude that 
gp5-D475A/D654A inhibits E. coli growth by interfering 
directly with DNA synthesis mediated by the bacterial 
replicosome.

The inhibition of DNA synthesis by gp5-D475A/D654A 
depends on the presence of thioredoxin. The addition of gp5-
D475A/D654A in the absence of thioredoxin to DNA synthesis 
reactions mediated by the E. coli replisome, up to a concen-
tration of 320 nM, does not result in any significant decrease 
in DNA synthesis (Fig. 3). Likewise, the presence of thiore-
doxin alone fails to inhibit E. coli Pol III–dependent DNA 
synthesis (Fig. 3B). Although WT gp5/Trx was able to 
attenuate E. coli replication at high concentrations, it did so 
with markedly lower potency than gp5-D75A/D654A/Trx. 
At concentrations greater than 80 nM, gp5-D475A/D654A/
Trx essentially eliminated DNA synthesis by E. coli Pol III, 
whereas significant levels of DNA synthesis by Pol III per-
sisted even at the highest concentration of WT gp5/Trx 
tested (Fig. 3A). The decrease in DNA synthesis by WT gp5/
Trx may arise from an indirect effect on DNA synthesis (see 
“Discussion”).

Given that polymerase-inactive gp5 alleles are also toxic for 
T7 phage growth (Table 1), we examined the effect of gp5-
D475A/D654A on the T7 DNA replication system. The pre-

cence of gp5-D475A/D654A partially inhibits the activity of WT 
T7 DNA polymerase and the T7 replisome. We first tested the 
effect of gp5-D475A/D654A on primer extension by T7 DNA 
polymerase. There is significant inhibition of the extension of a 
primer annealed to M13 ssDNA by WT T7 DNA polymerase 
only at gp5-D475A/D654A concentrations equivalent or higher 
than the concentration of WT polymerase present in the reac-
tion (Fig. 4A). In the absence of gp5-D475A/D654A, partially 
extended intermediates that arise from polymerase pausing 
at regions of secondary structure accumulate at earlier time 
points but diminish as synthesis progresses through the struc-
tured region (Fig. 4A, second and fourth lanes, black arrow). At 
concentrations of gp5-D475A/D654A approximately equimo-
lar or in excess to WT gp5, these intermediates persist even 
at the longest time points, with a concomitant decrease in fully 
extended products (Fig. 4A, eighth through sixteenth lanes, 
white arrow).

gp5-D475A/D654A inhibits leading strand DNA synthesis, 
catalyzed by T7 DNA polymerase and the T7 helicase gp4 
(Fig. 4B). During leading strand DNA synthesis, gp4 unwinds 
upstream dsDNA, generating the single-stranded template 
for nucleotide polymerization by T7 DNA polymerase. To-
gether, gp4 and T7 DNA polymerase display a processivity of 
>17 kb (3, 4). The presence of gp5-D475A/D654A leads to a 
dose-dependent decrease in deoxyribonucleotide incorpora-
tion. At a concentration of 30 nM, gp5-D475A/D654A inhib-
its DNA synthesis by ~50%. However, at the highest concen-
tration of gp5-D475A/D654A tested (320 nM), synthesis of 
the leading strand still occurs at a value of ~20% of control 
reactions (Fig. 4C).

The effect of gp5-D475A/D654A was also examined under 
conditions where leading and lagging strands are synthesized in 
a coordinated manner (Fig. 4C). The presence of gp5-D475A/ 
D654A leads to a similar degree of inhibition of DNA synthesis 
of leading and lagging strands (Fig. 4C). Similar to the results 
observed for leading strand synthesis, significant DNA syn-
thesis persists even at high concentrations of gp5-D475A/D654A. 
In conclusion, gp5-D475A/D654A also inhibits T7 DNA rep-
lication in vitro, but to a lesser extent than the inhibition 
observed for the E. coli replisome in vitro.

gp5-D475A/D654A/trxA displays a more rapid association to 
primer/template DNA.

T7 DNA polymerase has a high affinity for a primer template, 
with a $K_d$ of 20 nM (9, 16). One possibility for the inhibitory 
effect of gp5-D475A/D654A/Trx is that it retains its ability to 
bind to a primer template but is locked in position, because it is 
unable to covalently link the incoming nucleotide to the 
primer. We compared the ability of WT gp5 and gp5-D475A/
D654A to bind to a primer template and to ssDNA using an
electrophoretic mobility shift assay (EMSA), in both the presence and the absence of Trx (Fig. 5A). gp5-D475A/D654A exhibits the same binding characteristics as WT gp5. In the case of both gp5 variants, no significant binding is observed to either ssDNA or primer/template in the absence of Trx (Fig. 5A, lanes 2–5 and 15–18). Like WT gp5, gp5-D475A/D654A binds the primer/template stably only in the presence of thioredoxin (Fig. 5A, lanes 19–26). This strong binding is specific for a primer/template configuration because only weak binding to ssDNA is observed (Fig. 5A, lanes 2–5 and 15–18). The presence of thioredoxin slightly increases the affinity of gp5-D475A/D654A for ssDNA (Fig. 5A, lanes 4, 5, and 10–13). However, the polymerase–DNA complex formed is unstable under these electrophoretic conditions. Nevertheless, a decrease in the signal for free, labeled ssDNA and a smear representing labile protein–DNA complexes during electrophoresis is observed (Fig. 5A, lanes 2–13).

We used surface plasmon resonance to obtain a more quantitative description of the DNA-binding characteristics of gp5-D475A/D654. We used a series of binding experiments in which a concentration series of either WT gp5/Trx or gp5-D475A/D654A/Trx was followed over an immobilized primer template in the absence of deoxyribonucleotides and determined the kinetic parameters for their interaction (Fig. 5B and C). The results show that gp5-D475A/D654A/Trx binds a primer/template DNA with a 4-fold greater affinity than WT gp5/Trx. The increased affinity of gp5-D475A/D654A/Trx for primer/template DNA is solely due to enhanced association kinetics. As suggested in the experiment presented in Fig. 5B, the dissociation rate constant ($k_{\text{off}}$) is identical between both gp5 variants. However, the association rate constant ($k_{\text{on}}$) of gp5-D475A/D654A/Trx for primer/template DNA is roughly four times greater than for WT gp5/Trx. These results suggest that although the active site in gp5-D475/D654A/Trx cannot incorporate a nucleotide because it lacks the catalytic aspartates for polymerization, it associates with primer/template DNA faster, resulting in a higher affinity compared with WT gp5/Trx (17, 18).

Figure 3. gp5-D475A/D654A inhibits DNA synthesis by the reconstituted E. coli replisome in a thioredoxin-dependent manner. A, DNA synthesis by the E. coli replisome is inhibited by the presence of gp5-D475A/D654A/Trx. Denaturing agarose gel showing replication by the E. coli replisome in the presence of various concentrations of gp5 (+/− Trx) and gp5-D475A/D654A (+/− Trx). The E. coli replisome was assembled onto a rolling circle DNA substrate, and DNA synthesis was initiated as described under “Experimental procedures.” 10 s after addition of the initiation mixture containing [α-32P]dATP, the reaction was supplemented with various concentrations of WT gp5 or gp5-D475A/D654A in the absence and presence of Trx. Quenched samples were loaded on a denaturing agarose gel, and the radiolabel incorporated into DNA was detected by phosphorimaging. Bottom panel, the intensity of DNA replication products was plotted as a function of the concentration of gp5 variant added. Intensities were normalized to the average intensity of the replication product in no addition controls. B, inhibition of the E. coli replisome is not mediated by thioredoxin. Denaturing agarose gel showing replication by E. coli replisome in the presence of various concentrations of thioredoxin alone (Trx) and gp5-D475A/D654A (+/− Trx). Bottom panel, the intensity of DNA replication products was plotted as a function of the concentration of Trx or gp5-D475A/D654A/Trx added. Intensities were normalized to the average intensity of the replication product in no addition controls. RL, Resolution Limit.
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**Figure 4. Inhibition of WT T7 DNA polymerase by gp5-D475A/D654A.** A gp5-D475A/D654A inhibits primer extension by T7 DNA polymerase. A $^{32}$P-5' -end-labeled primer (50 nM) was annealed to M13 ssDNA (10 nM) and incubated with 100 nM WT T7 DNA polymerase, in the presence of 0.3 mM dNTPs, and increasing concentrations (0.03, 0.13, 0.5, or 2 μM) of gp5-D475A/D654A. Samples were removed at 10, 30, and 90 s after addition of MgCl$_2$ and separated on a denaturing polyacrylamide gel. Examples of extension intermediates accumulating (black arrow) or decreasing (white arrow) as the concentration of gp5-D475A/D654A increases. B, inhibition of strand-displacement DNA synthesis by gp5-D475A/D654A. Strand-displacement reactions rely on the helicase activity of gp4 to displace dsDNA encountered during DNA synthesis by T7 DNA polymerase. The reaction mixtures were supplemented with gp5-D475A/D654A (0, 1, 9, 3, 7, 15, 30, or 120 nM). The samples were incubated for 5 min at 30°C after initiating the reaction by addition of MgCl$_2$ and quenched with 50 mM EDTA, and the incorporation of dGMP was measured by liquid scintillation counting. C, gp5-D475A/D654A inhibits coordinated leading and lagging-strand DNA synthesis. gp5-D475A/D654A (0, 5, 20, 80, or 320 nM) was incubated in a standard minicircle reaction (see Experimental Procedures) for 5 min prior to addition of MgCl$_2$. The reactions were incubated at 30°C for 5 min and quenched with 50 mM EDTA, and incorporation of dGMP (for leading strand synthesis) or dCMP (for lagging-strand synthesis) was measured by liquid scintillation counting.

**Discussion**

DNA polymerases employ two carboxylic acid-containing residues to coordinate a catalytic metal cation, usually Mg$^{2+}$, in the active site (5, 10, 17). In the case of DNA polymerases from family A, two conserved aspartate residues are responsible for the coordination of metal ligands and are indispensable for catalysis (7, 12). From the crystal structure of T7 DNA polymerase (12), we identified Asp$^{475}$ and Asp$^{654}$ as the metal-coordinating residues in gp5 equivalent to those previously identified in the Klenow fragment of *E. coli* DNA polymerase I (7, 8). We targeted these postulated catalytic residues by replacing each or both aspartates with alanine.

We were surprised when we were unable to obtain transformants of singly or doubly substituted aspartate-to-alanine gene 5 alleles using common *E. coli* expression strains. We find that expression of polymerase-inactive gp5 is toxic if the host cells contain a functional allele for thioredoxin; no toxicity is observed in cells lacking thioredoxin. In addition, we find that the toxic effect is not observed in *E. coli* strains lacking the (DE3) lysogen, which encodes T7 RNA polymerase. This finding suggests that the levels of gp5-D475A/D654A protein that accumulate as a result of “leaky expression” of T7 RNA polymerase from the lacUV5 promoter in the (DE3) lysogen are sufficient to inhibit *E. coli* growth. In other words, our genetic analysis indicates that the toxic effect is manifest with very low expression of the altered polymerase. Combined, these results show that a complex of polymerase-inactive gene 5 DNA polymerase with thioredoxin is toxic for *E. coli* growth. We do not observe a toxic effect with WT gp5 or other polymerase active site gp5 variants. Toxicity depends on the presence of thioredoxin and is specific for gp5; no toxicity is observed in *trxA* strains. Furthermore, a *pol* Klenow fragment bearing homologous mutations corresponding to the magnesium-coordinating residues in the polymerase active site of gp5 and containing the TBD of gp5 did not exhibit toxicity. This latter result is not too surprising, considering that the chimeric protein shows a marginal level of stimulation in the presence of thioredoxin compared with gp5, has a 10-fold lower affinity for Trx than does T7 gp5, and is 3-fold less active than gp5/Trx, and its complex with Trx has a lower processivity compared with gp5/Trx (14).

Thioredoxin is essential for the T7 DNA replication machinery. It binds tightly to T7 gp5 and confers processivity to the polymerization reaction (16, 19, 20). T7 gp5 alone has a low processivity of ~5–20 nucleotides per binding cycle, and the acquisition of thioredoxin increases the processivity to ~800 nucleotides per binding event (19). This increase in processivity arises from an increased affinity of gp5/Trx for a primer/template: The $K_d$ of gp5/Trx for a primer/template determined by incorporation assays is 20 nM, whereas it is at least 20-fold higher in the case of gp5 alone (9, 16). Thioredoxin also provides gp5 with contacts for the T7 gene 4 helicase-primase and to the gene 2.5 ssDNA-binding protein (13, 21). Binding of thioredoxin to gp5 leads to the formation of two basically charged unstructured regions in gp5 that interact with the negatively charged unstructured C-terminal domains of both the ssDNA-
SSB supports T7 growth and this inhibition depends on the replisome. E. coli DNA replication. The E. coli A and loaded on a native 0.5 M Tris, 45 mM Borate, 0.5 mM EDTA, pH 8.0) gel. After drying, the gel was exposed to a phosphorimaging screen. B, sensograms obtained using a biotinylated primer/template coupled to streptavidin chip under flow of solutions containing WT gp5 (red) or gp5-D475A/D654A (dark blue) in the absence of nucleotides. C, kinetic and binding constants for the interaction of WT gp5 or gp5-D475A/D654A to an immobilized primer/template ligand determined by surface plasmon resonance.

Figure 5. gp5-D475A/D654A/Trx shows increased DNA-binding stability. A, EMSA gp5 or gp5-D475A/D654A in the presence and absence of thioredoxin. Left panel, lanes 1–12, 5'-32P-end-labeled ssDNA (19 nt in length). Right panel, lanes 14–26, ds 5'-22P-end-labeled primer/template (19-nt primer and 26-nt template). The samples were incubated as described under “Experimental procedures” and loaded on a native 0.5 × TBE (45 mM Tris, 45 mM Borate, 0.5 mM EDTA, pH 8.0) gel. After drying, the gel was exposed to a phosphorimaging screen. B, sensograms obtained using a biotinylated primer/template coupled to streptavidin chip under flow of solutions containing WT gp5 (red) or gp5-D475A/D654A (dark blue) in the absence of nucleotides. C, kinetic and binding constants for the interaction of WT gp5 or gp5-D475A/D654A to an immobilized primer/template ligand determined by surface plasmon resonance.

The molecular mechanism for the toxicity of gp5-D475A/D654A/Trx on E. coli DNA replication is difficult to dissect. The simplest mechanism is that the high affinity of gp5/Trx for DNA polymerase bound to DNA could be locked into place as a result of its inability to add the next incoming nucleotide. Such a scenario is similar to the stable polymerase complex created by the incorporation of a dideoxy-nucleotide, where the absence of condensation of the next nucleotide locks the polymerase in a polymerizing mode (12). An alternative, nonmutually exclusive, possibility is that the toxic effect exerted by polymerase-inactive gp5/Trx could be due to the generation or propagation of toxic DNA damage. DNA replication machineries encounter multiple barriers during genomic replication, resulting in stalled or abandoned replication forks that, if left unchecked, result in highly deleterious dsDNA breaks (25). The reactivation of replication forks is a major pathway in the maintenance of genomic integrity (26). Polymerase-inactive gp5/Trx could hinder the assembly of replication fork restart pathways, resulting in catastrophic genomic damage and triggering cell death, whereas any inhibition by WT gp5/Trx would be likely less potent (27, 28).

We envision that pol− gene 5 alleles could have applications in synthetic biology or industrial settings as a cell host shutoff mechanism. Bacterial cellular toxicity triggered through inhibition of DNA synthesis can form the basis for the removal of binding protein (gp2.5) and helicase-primase (gp4) of phage T7 (22). Of these two functions, it seems most likely that the toxic effect arises from the increases affinity for DNA, because it is unlikely that proteins in the E. coli replisome have specific contacts with thioredoxin. However, we cannot exclude the possibility that gp5/Trx might interact with E. coli SSB, because this protein has an acidic C-terminal tail with similar length, charge distribution, and a C-terminal phenylalanine, as do the acidic N-terminal tails of T7 gp2.5 and gp4. In fact, a chimeric gp2.5 containing the C terminus of E. coli SSB supports T7 growth and interacts with T7 DNA polymerase. However, a chimeric SSB protein containing the C-terminal tail of gp2.5 does not support T7 growth (23).
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selected cells in a population in a specific manner by designing a regulatory circuit that depends on mutant gene 5 or trxA co-expression in a temporally controlled manner.

Experimental Procedures

Construction of plasmids and protein purification

Mutations were simultaneously introduced into codons for aspartate 475 and 654 of T7 gene 5 in plasmids encoding WT gene 5 (pGP5-3) and exonuclease-deficient gene 5 (D5A, D65A) (pGP5-3 exo−) using Gibson assembly (29). Site-directed mutagenesis was used to introduce single mutations into T7 gene 5. Similarly, we used Gibson assembly to construct a double substitution (D705A/D882A) variant of Klenow fragment containing an insertion of the T7 gp5 TBD using pKLN-TBD as template (14). BL21 (DE3) ΔtrxA was constructed by replacing trxA with a gene encoding kanamycin resistance using the λ RED system (30). T7 gp5 variants were expressed in E. coli A307 (DE3), which lacks endogenous thioredoxin and the overexpressed gene 5 proteins were purified as previously described (19). WT E. coli thioredoxin was prepared as previously described (31). E. coli clamp loader complex (∆add’Δχγ), β2 clamp, DnaG, DnaB, and ssDNA-binding protein (SSB) were purified as previously described (32, 33). αεθ, αεQθ, and αεLθ were provided by Nicholas Dixon (University of Wollongong).

Genetic assays

E. coli BL21 (DE3), BL21, and E. coli BL21 (DE3) ΔtrxA were transformed with plasmids containing the genes for WT or variant T7 gene 5 or the Klenow fragment of E. coli DNA polymerase I containing the T7 gp5 TBD according to standard methods. Transformation mixtures were plated onto LB agar plates containing 100 µg/ml ampicillin and grown overnight at 37°C. The number of colonies after incubation were counted. For phage growth and complementation assays, E. coli DH5α transformed with gene 5–expressing plasmid were grown in LB to exponential phase. Cell culture (0.3 ml) was mixed with 3 ml of soft agar (LB broth + 0.7% agar) and plated onto LB-agar/ampicillin plates. Aliquots of serially diluted WT or gene 5-deletion (Δ5) T7 phage were spotted onto the plates to determine phage titer.

DNA polymerase assays

Primed M13 ssDNA (7 nM) was incubated with increasing concentrations (3–2,000 nM) of purified WT gp5 or gp5-D475A/D654A in the presence of a 10-fold molar excess of thioredoxin in a buffer consisting of 40 mM Tris-HCl pH 7.5, 50 mM potassium glutamate, 10 mM MgCl₂, 5 mM DTT, and 0.3 mM dTTP, dATP, dCTP, and [α-32P]dTTP at 25°C for 2 min. The reactions were quenched with EDTA and spotted onto DE81 filters, and the amount of nucleotides incorporated into DNA was determined as previously described (1). 3’–5’ exonuclease activity of gp5 variants was measured using an internally [32P]-labeled 3-kb dsDNA generated by PCR in the presence of [α-32P]dGTP from a plasmid template. Reaction mixtures contained 40 mM Tris-HCl, pH 7.5, 10 mM MgCl₂, 5 mM DTT, 50 mM potassium glutamate, 3 µM uniformly labeled dsDNA (expressed as nucleotides) and varying concentrations of gp5 variants. Thioredoxin was included at a 10-fold molar excess over gp5 concentrations in the assay. Samples were taken after incubation at 37°C and quenched with 18% (w/v) TCA and 2 mg/ml BSA. The acid-soluble radioactivity in the supernatant was measured by liquid scintillation counting.

The primer template for rolling-circle DNA synthesis mediated by the E. coli replication proteins was prepared and purified as described (32). A 73-mer tailed oligonucleotide (T36GAATTTCCGACCGCTCCACAGTACGT—GAATCATG) was annealed to a modified M13mp7(L2) circular closed ssDNA and extended with T7 DNA polymerase (New England Biolabs) at 37°C for 20 min. The filled M13mp7(L2) was purified by phenol-chloroform extraction. The E. coli replisome was assembled by mixing 1 nM template, 20 nM τ, δ’, χ’, 20 nM β’, clamp dimer, 50 nM DnaB hexamer, 60 nM Pol III core complex (αεθγ), 50 µM ATPγS, and 60 µM dCTP and dGTP in 50 mM Hepes-KOH, pH 7.9, 12 mM Mg (OAc)₂, 0.1 mg/ml BSA, 10 mM DTT on ice and incubating the mixture in a water bath at 37°C for 6 min. Replication was initiated by adding prewarmed 10X initiation mixture to the assembly reactions. The 10X initiation mixture contained 1 mM ATP, 250 µM rNTPs, 60 µM dTTP, and [α-32P]dATP, 500 nM SSB tetramer, and 600 nM DnaG. All concentrations are the final concentrations in the reactions. 10 s after initiation, the indicated concentrations of either WT T7 gp5 or its variants were added to the reaction mixtures. Replication reactions took place in a 37°C water bath. Replication reactions were quenched by adding EDTA to a final concentration of 25 mM. Quenched reactions were applied to a 0.6% alkaline denaturing agarose gel, and replication products were resolved by running the gel in an alkaline electrophoresis buffer (50 mM NaOH, 2 mM EDTA) at 21 V for 15 h. Replication products were visualized by autoradiography (32).

To assess the effect of gp5-D475A/D654A on the activity of WT gp5, we incubated various concentrations of gp5. D475A/D654A(exo−)/Trx along with 100 nM gp5(exo−)/Trx in the presence of M13 DNA annealed to a 5’-32P-labeled primer in 40 mM Tris-HCl, pH 7.5, 50 mM potassium glutamate, 5 mM DTT, and 0.3 mM dNTPs. DNA synthesis was initiated by adding MgCl₂ to a final concentration of 10 mM. At 10, 30, or 90 s, the reactions were stopped by the addition of an equal volume of formamide loading dye (93% formamide, 50 mM EDTA, 0.01% xylene cyanol, and 0.01% bromphenol blue). The samples were heated to 95°C for 5 min and loaded onto a denaturing 5% polyacrylamide gel. The gel was dried and exposed to a phosphorimaging screen.

Leading-strand DNA synthesis was measured in a reaction containing 10 nM of a DNA minicircle bearing a replication fork (1), 50 mM Tris-HCl pH 7.5, 50 mM potassium glutamate, 5 mM DTT, 0.6 mM dATP, dCTP, dGTP, and [α-32P]dGTP, 60 nM gp4 (monomer), 80 nM WT gp5/Trx, and varying concentrations of gp5-D475A/D654A/Trx. The reaction was initiated by adding MgCl₂ to a final concentration of 10 mM, incubated at 25°C for 5 min before quenching with EDTA, and processed as described (34).

Coordinated leading- and lagging-strand DNA synthesis reactions were assembled in the presence of varying concentrations
of gp5-D475A/D654A/Trx. The reaction mixtures contained 40 mM Tris-HCl, pH 7.5, 50 mM potassium glutamate, 10 mM MgCl₂, 0.6 mM dNTPs (with either [α-32P]dGTP to measure synthesis of the leading strand or [α-32P]dCTP to measure lagging-strand synthesis), 60 mM gp4 (monomer), 80 nM WT T7 DNA polymerase, 3 μM gp2.5, and 10 nM minicircle DNA. The sequence bias of minicircle DNA allows for monitoring of synthesis of the leading and lagging strands by measuring the incorporation of dGMP or dCMP, respectively (1).

**DNA-binding assays**

An EMSA was used to measure binding of DNA polymerase to both ssDNA and a primer/template DNA duplex. A 19-nt ssDNA (5’-TGCGACGATACATCA-3’) was 5’-end-labeled with T4 polynucleotide kinase and [γ-32P]ATP and served as the primer strand. A dsDNA containing a 5’-overhang representing a primer/template duplex was formed by annealing the labeled primer strand with a 26-nt ssDNA (5’-CAGT-GACCAGGCTGTTATCGTCGGCA-3’). Binding reactions contained 3 nM 5’-32P-primer or primer/template in 40 mM Tris-HCl, pH 7.5, 50 mM potassium glutamate, 5 mM DTT, 10 mM MgCl₂, and various concentrations of gp5 exo⁻ or gp5-D475A/D654A exo⁻ in the presence or absence of Trx. The samples were incubated at room temperature for 10 min and loaded onto a native 10% polyacrylamide gel with 0.5X TBE (45 mM Tris, 45 mM Borate, 0.5 mM EDTA, pH 8.0) as the electrophoretic buffer. The gel was run in a cold room, dried, and visualized by phosphorimaging.

In surface plasmon resonance assays, the 19-nt primer used for EMSA was annealed to the 26-nt template strand containing a biotin attached to the 3’-end. dsDNA was coupled to a streptavidin chip at a concentration of 0.25 μM in 10 mM Heps, pH 7.4, 150 mM NaCl, 0.005% Tween 20 (vol./vol.) at a flow rate of 10 μl/min. Free streptavidin on the flow cells was then blocked by washing with biotin. Binding studies were carried out by flowing 200 nM gp5 exo⁻/Trx or gp5-D475A/D654A exo⁻/Trx over 600 RU immobilized primer/template in 20 mM Heps, pH 7.5, 5 mM MgCl₂, 2.5 mM DTT, 200 mM potassium glutamate, 1% glycerol (w/v) at a flow rate of 10 μl/min. For determination of kinetic parameters, 50 RU of primer/template was immobilized and various concentrations (12.5, 25, 50, 75, and 100 nM) of gp5 exo⁻/Trx or gp5-D475A/D654A exo⁻/Trx were flowed over the chip. Kinetic data were obtained by fitting binding sensorgrams using BIA-Evaluation software. All experiments were repeated at least four times, with the exception of those whose data are shown in Figs. 3B and 5C, which were done in duplicate.

**Data availability**

All of the data presented are contained in the manuscript and are available upon request from Alfredo J. Hernandez, Harvard Medical School, alfredo_hernandez@hms.harvard.edu.

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**Abbreviations**—The abbreviations used are: Trx, thioredoxin; Pol, polymerase; EMSA, electrophoretic mobility shift assay.

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**References**


