Single molecule measurement of the “speed limit” of DNA polymerase

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Although DNA replication is often imagined as a regular and continuous process, the DNA polymerase enzyme is a complicated machine and can pause upon encountering physical and chemical barriers. We used single molecule measurements to make a detailed characterization of this behavior as a function of the template’s secondary structure and the sequence context. Strand displacement replication through a DNA hairpin by single DNA polymerase molecules was measured in real time with near single base resolution and physiological concentrations of nucleotides. These data enabled the measurement of the intrinsic “speed limit” of DNA polymerase by separating the burst synthesis rate from pausing events. The strand displacement burst synthesis rate for Escherichia coli DNA Polymerase I (KF) was found to be an order of magnitude faster than the reported bulk strand displacement rate, a discrepancy that can be accounted for by to sequence specific pausing. The ability to follow trajectories of single molecules revealed that the burst synthesis rate is also highly stochastic and varies up to 50-fold from molecule to molecule. Surprisingly, our results allow a unified explanation of strand displacement and single strand primer extension synthesis rates.

FRET | pausing | DNA replication | DNA hairpin

Fast and accurate DNA replication is required to ensure the faithful transfer of genetic information to daughter cells. Replication forks can be slowed or paused by encountering DNA-binding ligands (1), genomic tRNA gene sites (2), stalled ternary complexes of RNA polymerase (3), and DNA-bound protein (4–5). DNA polymerase can also pause due to specific DNA sequences, including palindromic DNA capable of forming hairpin secondary structure (6–8), slow zones (9), and trinucleotide repeats of (CGG)$_n$(C)$n$(CG) or (CTG)$_n$(C)$n$(AG) (10–11). While these factors are thought to prevent fork movement along the template by steric hindrance and may be regulated by the replisome itself, other factors, including temperature, contaminants, nucleotide analogs (12), template tension (13–14), and nonclassic pause sites such as Pyr-G-C (15), may also interfere with steps in the DNA polymerase reaction pathway.

To measure the intrinsic “speed limit” of DNA polymerase itself, there must be sufficient spatial and temporal resolution to separate pausing events from burst synthesis. Although the kinetics of the Klenow fragment of Escherichia coli DNA Polymerase I (Pol I(KF)) have been studied using a variety of approaches (13, 15–20), there are conflicting reports of the synthesis rate and it has not yet been possible to dissect the interplay between pausing and burst synthesis. Rapid quenching and stopped-flow bulk techniques have determined that during primer extension synthesis, the first nucleotide is incorporated by Pol I(KF) with a rate ranging from 40–87 nt sec$^{-1}$ (16–17, 19) while subsequent nucleotides are incorporated at a slower rate of 13–15 nt sec$^{-1}$ (17, 20). This bulk processive rate is twice as fast as that found by single molecule primer extension synthesis rate measurements (7 nt sec$^{-1}$) (13). Primer extension synthesis is generally thought to be faster than strand-displacement synthesis, and this idea is supported by a bulk rapid quenching study which found that the strand displacement synthesis rate of Pol I(KF) was even slower at 1.2 nt sec$^{-1}$ (20).

We developed a FRET-based approach (21) to study single polymerase molecules undergoing strand displacement replication through a DNA hairpin to further investigate this conundrum between the various reported rates. This assay allowed us to measure polymerase activity in real time at saturating nucleotide concentrations with near single base resolution, thereby enabling separation of burst synthesis rates from pausing effects. Our results show that the only literature strand displacement synthesis rate for Pol I(KF) (20) is not the true burst synthesis rate because it likely includes sequence specific pausing effects due to the template used in the experiments. These pausing events are not unique to Pol I(KF) and were also observed with φ29 DNA polymerase. The existence of the pausing effect is known from bulk studies (15), but without the characterization of its heterogeneity and kinetics enabled by the single molecule experiments described here, its central role in determining the speed limit of DNA polymerase was not previously appreciated. The burst synthesis strand displacement rates obtained from our single molecule measurements were also highly heterogeneous from molecule to molecule, with rates varying from 1 nt sec$^{-1}$ to over 50 nt sec$^{-1}$. This variation in rates was previously hidden due to the ensemble averaging of bulk measurements and could only be accessed with a single molecule approach. Surprisingly, our characterization of the pausing heterogeneity in strand displacement also allows us to explain the discrepancies in rates found in primer extension experiments and shows that strand displacement rates are not slower than primer extension rates.

Results and Discussion

A 259-nt single-stranded DNA molecule was designed and synthesized containing an internal double-stranded 33-bp hairpin flanked by 94-nt single-stranded tails, one of which had a 3′-biotin group (Fig. 1A). The 3′ base of the hairpin contained an internal Cy3 FRET donor and the 5′ base contained an internal nonfluorescent FRET acceptor (Black Hole Quencher-2, BHQ-2) so that when the hairpin is fully folded, quenching of Cy3 by BHQ-2 prevents any fluorescence emission (Fig. 1B). The length of the hairpin was chosen to make full use of the dynamic range for this FRET pair: when the last base pair of the hairpin is broken due to strand displacement replication, the distance between fluorophores is just over twice the Förster radius ($R_0 = 5.02$ nm, 33 bp of dsDNA = 11.2 nm). An 80-nt primer was annealed to the 3′ end of the template and the DNA constructs were attached to the surface of a glass coverslip via a 3′ biotin-streptavidin linker. The coverslip was placed on an inverted Nikon TE-2000S TIRF microscope and images were collected with a 60 × 1.45 NA objective at 5 Hz with a Photometrics Cascade II EMCCD. Each field of view contained, 1To whom correspondence should be addressed. E-mail: quake@stanford.edu.

Author contributions: J.J.S. and S.R.Q. designed research; J.J.S. performed research; J.J.S. and S.R.Q. analyzed data; and J.J.S. and S.R.Q. wrote the paper.

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on average, over 1,000 individually resolvable single molecules (for full experimental details, see Materials and Methods).

To correlate the FRET signal with template position, a series of primer extension experiments were performed by adding DNA polymerase and only 3 of the 4 deoxynucleotide triphosphates (dNTPs). After the first extension and imaging step, the flow cell was thoroughly washed to remove any unincorporated nucleotides, and the process was repeated multiple times with a different set of 3 dNTPs. This allowed us to extend the primer to known positions on the template and observe the corresponding reduction in FRET signal as the donor and acceptor were forced apart. Before extension begins, the distance $R$ between the 2 dyes is small and the FRET efficiency is $E_{\text{FRET}} = 1/[1+(R/R_0)^2] \approx 1$. As the primer extends and the hairpin unwinds, $R$ increases and $E_{\text{FRET}}$ decreases, leading to recovery of Cy3 signal. The observed Cy3 intensity is thus given by $I_{\text{Cy3}} = I_{\text{MAX}} (1-E_{\text{FRET}}) = I_{\text{MAX}} [1-(1/[1+(R/R_0)^2])]$, where $I_{\text{MAX}}$ is the maximum Cy3 intensity achievable when the dyes are as far apart as possible (assuming 0.34 nm per base pair, the maximum dsDNA separation is $2 \times 33$bp + 5bp loop = 71 bp $\approx 24$ nm). A weighted least-squares fit of the data to this equation gave $I_{\text{MAX}} = 987$ a.u. and $R_0 = 13.5$ bp = 5.2 nm, which is consistent with the vendor’s reported Förster radius for Cy3 and BHQ-2. There is some width to the distributions of Cy3 intensities at each position due to signal to noise variation and Cy3-BHQ-2 orientation effects (supporting information (SI) Fig. S1 and Table S1).

However, by performing this alignment procedure on an ensemble of records at each position, a calibration curve relating the observed fluorescent intensity and the polymerase position along the template was created (Fig. 1C). Using this calibration curve and its 95% confidence limits, any arbitrary fluorescent intensity could be converted into a corresponding polymerase location along the template (Fig. 1D).

The real-time kinetics of DNA replication were examined in the presence of all four dNTPs with two different DNA polymerases: the Klenow fragment of DNA Polymerase I (exo-) from *E. coli* (Pol I(KF)) and the replicative polymerase from the *Bacillus subtilis* bacteriophage φ29. The trajectories for both polymerases exhibited heterogeneous behaviors that could be classified into 4 categories: fast replication without pausing (Fig. 2A), fast replication with a single pause (Fig. 2B), fast replication with multiple pauses (Fig. 2C), and slow replication. Slow replication occurred in less than 6% of the traces, and while its origin is unclear, it could stem from the template sticking to the surface or unfavorable polymerase-surface interactions. Approximately 66% of the DNA polymerases extended the entire template without a pause while about 28% of the polymerases paused at least once. The pauses occurred at highly stereotyped positions and we took advantage of the single molecule approach to measure both the distribution between pausing and nonpausing polymerase molecules and the relationship between the pauses and the burst synthesis speeds of the polymerase, data that would be impossible to observe with conventional bulk techniques.

The positional accuracy of individual trajectories depended on both template location and signal-to-noise with a maximum attainable resolution of approximately 2 bp. For example, a well-behaved trajectory that paused with a mean intensity of $562 \pm 47$ a.u. (SD) was called at $+15$ or $+16$ bp; a noisier trajectory that paused with a mean intensity of $447 \pm 168$ a.u. (SD) was called at $+14 \pm 2$ bp. By aligning an ensemble of trajectories for Pol I(KF) and φ29, polymerase pause positions were localized with near single base accuracy over the template from $-8$ bp to $+18$ bp (Fig. 3A and B). Outside of this window the overlapping confidence limits of the calibration curve prevented pause localization accuracy better than $\pm 3$ bp. For Pol...
I(KF), over 85% of the pauses were located between +13 and +17 bp, a region that is GC-rich and includes a Pyr-G-C motif at +16 bp. Similarly, over 82% of the φ29 pauses were located between +14 and +18 bp. Pyr-G-C motifs at +8 bp and +26 bp showed weak pausing for Pol I(KF) and no pausing for φ29, but the identification accuracy of pause events is lower at those locations.

A significant reduction in the frequency of pausing for both enzymes was observed with the addition of 1 M betaine to the DNAP/dNTP mix (Fig. 3A and B). Betaine is a zwitterionic osmoprotectant that is thought to alter DNA stability so that GC-rich regions melt at AT-rich temperatures (22), a finding that has led to its inclusion in PCR formulations for improved amplification of difficult templates (23–24). Betaine has also been used in bulk studies to suppress replication pausing at Pyr-G-C sequences (15). Increasing the temperature of the Pol I(KF) reaction from 23 °C to 37 °C also reduced the frequency of pausing significantly, which is in agreement with previous bulk studies by Mytelka et al. (15). These authors suggested that pauses at Pyr-G-C sequences might be caused by difficulties in the polymerase fingers-closing conformational change, as at the time this transition was thought to be rate-limiting and the most sensitive to changes in temperature. However, a recent report (25) shows that the slow prechemistry step is likely not the fingers-closing transition, raising the possibility that these pauses are associated with an earlier DNA template rearrangement step that might be sequence dependent.

To verify that the observed pausing was sequence dependent and not due to the hairpin secondary structure, a control DNA molecule was constructed with the same overall structure and length as the original (Fig. 1B Top) but with a different stem sequence (Fig. 1B Bottom). The control sequence removed all three occurrences of the Py-G-C motif while maintaining as much similarity to the original sequence as possible. Importantly, only 2 bases were changed in the region where most pausing was found to occur: 5′-TAGGCCCA-3′ was changed to 5′-TAGGATCCA-3′. Both polymerases showed over a 5-fold reduction in pause frequency with the control sequence compared to the original sequence, with less than 5% of the trajectories showing residual pausing (Fig. 3C and D). The source of the residual control sequence pausing could be the high GC content (66%) from +15 to +20 where the majority of the pauses occurred. In addition, Mytelka et al. (15) found that Pol I(KF) paused at two nonconsensus sequences that were within 1 nt of the Pyr-G-C consensus (TCC and TGT). The control sequence did contain a TCC motif at +18 which may have contributed to some weak pausing. The control also had a TGG motif at +14, which is within 1 nt of the consensus and was not previously identified to cause pausing, but may have also contributed. Taken together, the control experiments suggest that the secondary structure of the hairpin was not playing a role in the observed pausing. It also suggests that the central 5′-CG-3′ motif, either by itself or in the context of the surrounding sequence, was responsible for causing the majority of the pauses.

The distribution of pause lifetimes is consistent with single-step Poisson statistics. At 23 °C Pol I(KF) exhibited a mean pause lifetime of 13.2 sec for the sample template (Fig. 4A). In the reduced population of trajectories that showed pausing even with the addition of betaine or heat, the mean pause lifetime decreased 30% (Fig. 4B and Fig. S2). A similar reduction in pause lifetime was observed for Pol I(KF) acting on the control template (Fig. 4C). The mean pause lifetimes for φ29 with the sample template were less than for Pol I(KF) (Fig. 4E), and a reduction in lifetimes was not observed with betaine addition (Fig. 4F) or on the control sequence (Fig. 4G). Short time period pauses were likely undersampled due to limitations of the sampling bandwidth and signal to noise, which resulted in some of the pause lifetime distributions to have a distinct rise and decay, and so they were excluded from the fit. The pause intensities had a mean signal to noise ratio of 2.6 while the
full extension intensities had a mean signal to noise ratio of 5.6 (Fig. S3). Pause frequencies and lifetimes are impossible to obtain through ensemble studies and this knowledge may help constrain the biophysical models of the polymerase kinetic pathway.

The pauses seen here are clearly distinct from those observed in previous single molecule DNA polymerase experiments. T7 and Pol I(KF) polymerases have been shown to exhibit long pauses of heterogeneous length under template tension (13–14). These pauses were thought to be due to fluctuating hairpins, exogenous DNA hybridization, or template damage. The entire T7 replisome was also shown to halt leading-strand synthesis due to primase activity on the lagging strand (26). Similar experiments with the E. coli replisome identified a relationship between pause times and core polymerase concentration, suggesting the pauses can be caused by enzyme dissociation events (27).

A recent study on real-time DNA sequencing with single ε29 polymerase molecules identified pause sites that corresponded to regions with predicted template secondary structure (28). However, these experiments were not performed at saturating dNTP concentrations, which complicates the interpretation. We show here that ε29 and Pol I(KF) are susceptible to sequence-dependent pausing at saturating dNTP concentrations and that these pauses are not due to template secondary structure.

DNA synthesis is therefore a combination of high activity rapid synthesis that reflects the intrinsic “speed limit” of DNA polymerase interspersed with low activity pauses; our single molecule measurements enable us to separate these contributions and measure their kinetics. DNA synthesis burst rates were obtained by calculating the slopes of the trajectories between pauses (Fig. 5A Inset; see Materials and Methods). The time-averaged strand displacement rate for Pol I(KF) at 23 °C without betaine was 14 ± 3 nt sec⁻¹ (SEM) (Fig. 5A). The mean rate increased with either the addition of betaine [17 ± 2 nt sec⁻¹ (SEM), Fig. 5B] or with increasing temperature; at 37 °C, Pol I(KF) had a mean replication rate of 17 ± 3 nt sec⁻¹ (SEM) (Fig. 5C). For ε29 at 23 °C, a wide distribution of synthesis burst rates was found ranging from 1 nt sec⁻¹ to over 50 nt sec⁻¹ with a mean rate of 28 ± 5 nt sec⁻¹ (SEM) (Fig. 5D). Our window of sensitivity and data acquisition rate placed an upper limit of approximately 50 nt sec⁻¹ on our rate measurements, with faster rates being discarded. This is likely why our ε29 rate is slightly less than the reported ε29 bulk rates of 38 nt sec⁻¹ (29).

A recent bulk study of Pol I(KF) reported a strand displacement synthesis rate over an 18-bp template of 1.2 nt sec⁻¹ (20), which is an order of magnitude slower than our time-averaged burst synthesis rate. However, the template used in the bulk study contained a few Pyr-G-C pause motifs. Based on the results of our single molecule experiments, we would predict Pol I(KF) to have 1.3 sec of synthesis time and 10.3 sec of pause time over this template, giving an estimated ensemble rate of 1.6 nt sec⁻¹, which is in good agreement with their observations.

Although strand displacement synthesis and single stranded primer extension synthesis are expected to have substantial differences in their energetics, we compared our results to experiments that measured single stranded primer extension synthesis and found surprising agreement. Our observed rate of 14 nt sec⁻¹ is consistent with bulk primer extension experiments (17, 20). The rates measured here are 2- to 3-fold faster than a Pol I(KF) single molecule measurement (13), which was a low tension force spectroscopy study on a long template that did not have sufficient time resolution to separate pauses from bursts. Assuming that Pyr-G-C pause sites occur once every 32 bp in a random template and that the pause motif is 25% efficient, our data predict that the expected net synthesis rate over long templates is approximately 6 nt sec⁻¹, which is in agreement with the previous single molecule measurement (7 nt sec⁻¹).

For Pol I(KF) molecules that paused, the mean replication rate over the template sequence until the pause site was 11 ± 1 nt sec⁻¹ (SEM); after the pause site, the mean rate increased to 18 ± 2 nt sec⁻¹ (SEM). Similar to Pol I(KF), ε29 molecules that paused underwent faster replication after arrest [41 ± 3 nt sec⁻¹ (SEM)] than before [24 ± 4 nt sec⁻¹ (SEM)]. Accelerating synthesis rates were also observed in trajectories that did not pause (Fig. S4). The small difference in rates before and after pausing may be due to the transition near the end of the hairpin from strand displacement replication to primer extension replication, or it could be an artifact related to the presence of the fluorophore in the template. Given the predictive accuracy of...
our results described above, we believe this change is more likely the replication transition than a dye-enzyme effect.

There was a surprising amount of heterogeneity in the strand displacement synthesis rates for both Pol I(KF) and e29 from molecule to molecule over identical sequences (Fig. 5), with values ranging from 1 to 50 nt sec−1 to 0.1 nt sec−1. The stochasticity of the synthesis rates is hidden in time-averaged or bulk rates and can only be revealed with a single molecule approach. The source of this heterogeneity could be of great interest for future polymerase studies and has not received significant attention in the literature to date.

One potential drawback of a quencher-based approach is that blinking fluorophores might incorrectly be interpreted as real extension events. If a Cy3-Cy5 FRET pair were used, this would certainly be a concern as Cy5 is known to exhibit frequent blinking (30–31). In this FRET system, a false positive extension could be caused by either BHQ-2’s quenching ability switching off before extension begins or a dark Cy3 fluorophore switching on after extension has finished. BHQ-2 blinking was not significantly observed while imaging folded hairpins in the absence of polymerase or dNTPs and has not been reported in the literature. Cy3 fluorophores have a dark-state lifetime of approximately 2 sec and the on-blink and off-blink rates have been reported to be 0.36 and 0.26 times per molecule over 120 sec, respectively (32). Given the frequent blinking and short off times of Cy3, the probability of it being off during any real extension is very low. It is therefore not surprising that only approximately 1% of the trajectories exhibited identifiable blinking after full extension (signal going from on to off and then back to on). Importantly, for the kinetics experiments we observed intermediate intensity pauses in approximately 50% of the trajectories, a phenomenon that cannot be produced by fluorophore blinking. The calibration experiments (Fig. 1C) with polymerases stalled on the template did not show any significant deviation from the expected FRET behavior for a single acceptor system, so protein-induced quenching was not a significant source of error.

The real-time single molecule observation of Pol I (KF) revealed that the polymerase’s intrinsic strand displacement “speed limit” is an order of magnitude faster than reported by previous bulk processive synthesis measurements. Replication was found to be interrupted by heterogeneous sequence-dependent pauses that were independent of template secondary structure, and knowledge of these pause frequencies and lifetimes would otherwise be impossible to obtain from ensemble studies ranging from 1 to 50 nt/sec. The probability of Cy3 being off during any real extension is very low. It is therefore not surprising that only approximately 1% of the trajectories exhibited identifiable blinking after full extension (signal going from on to off and then back to on). Importantly, for the kinetics experiments we observed intermediate intensity pauses in approximately 50% of the trajectories, a phenomenon that cannot be produced by fluorophore blinking. The calibration experiments (Fig. 1C) with polymerases stalled on the template did not show any significant deviation from the expected FRET behavior for a single acceptor system, so protein-induced quenching was not a significant source of error.

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and distinguished from gradual nonstepped growth using a custom MATLAB step-fitting algorithm (34). This algorithm generates “S-values” which are a measure for the quality of a step-fit for a given number of steps over an entire trajectory, and the maximum value of S should correspond to the best fit. For noisy trajectories, step trains containing a slight excess of steps were chosen to ensure that all true steps were included. Some trajectories like 1C were scored as having multiple distinct pauses using this algorithm. Dwell times were calculated as the time between steps. To improve the accuracy of the rate calculations, trajectories were oversampled 20-fold and fit to a cubic spline in MATLAB. Synthesis rates were then calculated by measuring the trajectory slope between the mean intensities of 2 consecutive pauses or a pause and full extension. Due to the approximately 10 bp window of sensitivity and the 5 Hz acquisition rates, synthesis rates in excess of 50 nt/sec were discarded.

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Correction

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The authors note that, due to a printer error, the legend for Fig. 4 appeared incorrectly. The second sentence, “The red curves are normalized single exponential fits given by \( f = \tau^{-1} \exp(-t/\tau) \), where \( \tau \) is the mean pause lifetime,” should instead appear as “The red curves are normalized single exponential fits given by \( f = \tau^{-1} \exp(-t/\tau) \), where \( \tau \) is the mean pause lifetime.” The figure and its corrected legend appear below.

![Fig. 4](image-url)

Fig. 4. Pause lifetimes for Pol I(KF) and φ29 were measured under different conditions for the sample and control sequences. The red curves are normalized single exponential fits given by \( f = \tau^{-1} \exp(-t/\tau) \), where \( \tau \) is the mean pause lifetime. Bins excluded from the fit due to undersampling are shown in white. (A) Pol I(KF) at 23°C with the sample template; (B) Pol I(KF) at 23°C with 1 M betaine with the sample template; (C) Pol I(KF) at 23°C with the control template; (D) φ29 at 23°C with the sample template; (E) φ29 at 23°C with 1 M betaine with the sample template, and (F) φ29 at 23°C with the control template.