A Slow ATP-induced Conformational Change Limits the Rate of DNA Binding but Not the Rate of β Clamp Binding by the Escherichia coli γ Complex Clamp Loader

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In Escherichia coli, the γ complex clamp loader loads the β-sliding clamp onto DNA. The β clamp tethers DNA polymerase III to DNA and enhances the efficiency of replication by increasing the processivity of DNA synthesis. In the presence of ATP, γ complex binds β and DNA to form a ternary complex. Binding to primed template DNA triggers γ complex to hydrolyze ATP and release the clamp onto DNA. Here, we investigated the kinetics of forming a ternary complex by measuring rates of γ complex binding β and DNA. A fluorescence intensity-based β binding assay was developed in which the fluorescence of pyrene covalently attached to β increases when bound by γ complex. Using this assay, an association rate constant of 2.3 × 10^7 M⁻¹ s⁻¹ for γ complex binding β was determined. The rate of β binding was the same in experiments in which γ complex was preincubated with ATP before adding β or added directly to β and ATP. In contrast, when γ complex is preincubated with ATP, DNA binding is faster than when γ complex is added to DNA and ATP at the same time. Slow DNA binding in the absence of ATP preincubation is the result of a rate-limiting ATP-induced conformational change. Our results strongly suggest that the ATP-induced conformational changes that promote β binding and DNA binding differ. The slow ATP-induced conformational change that precedes DNA binding may provide a kinetic preference for γ complex to bind β before DNA during the clamp loading reaction cycle.

Two accessory factors, a sliding clamp and a clamp loader, enhance the efficiency of DNA replication by increasing the processivity of DNA synthesis. The clamp loader assembles onto DNA. The clamp binds the DNA polymerase to increase the processivity of DNA synthesis from tens to thousands of nucleotides in a single binding event. These processivity factors are conserved from bacteria to humans (reviewed in Refs. 1 and 2). The Escherichia coli β clamp is composed of two identical protein monomers assembled into a ring-shaped dimer, which encircles duplex DNA (3, 4). The clamp slides freely along the DNA while tethering DNA polymerase III to the DNA template. The E. coli clamp loader is composed of seven subunits. At the replication fork, a complete clamp loader includes three copies of the dnaX gene product and one copy each of δ, δ', χ, and ψ (5–7). The dnaX gene produces two proteins, a full-length protein (τ) and a truncated protein (γ), which is created by a translational frameshift (8–10). Clamp loaders containing either three copies of the τ subunit (τ complex, τ,δδ'χψ) or three copies of the γ subunit (γ complex, γ,δδ'χψ) are fully active in clamp loading (6). The τ subunits have an additional C-terminal extension that coordinates the activities of the replisome (reviewed in Refs. 11 and 12). The clamp loader used in all experiments in this study is the γ complex.

In the presence of ATP, γ complex binds with high affinity to β and DNA to form a ternary complex. Binding to primed template DNA triggers the clamp loader to hydrolyze all three molecules of ATP (13–15). Upon ATP hydrolysis, β is released onto DNA and the clamp loader dissociates from the β-DNA complex. Thus the clamp loading reaction can be divided into two stages based on ATP requirements: 1) formation of a ternary complex requiring ATP binding, and 2) dissociation of the ternary complex to load β on DNA requiring ATP hydrolysis. This article focuses on steps in the first phase involving the formation of the ternary complex.

High affinity binding of the clamp loader to the β clamp and DNA requires ATP binding by the clamp loader (16–18). Presumably, ATP binding promotes conformational changes within the clamp loader that expose surfaces/residues that interact with the clamp and DNA. When equilibrated with ATP, the γ complex can bind either the clamp or DNA; however, a productive clamp loading reaction most likely requires the clamp loader to bind the clamp before binding DNA. When the γ complex alone binds p//DNA, the interaction with DNA triggers rapid ATP hydrolysis and dissociation of the clamp loader (19, 20). Therefore, clamp loading would be more efficient if there were some mechanism to favor clamp binding before DNA binding and productive clamp loading. One such mechanism would be a kinetic preference for the clamp loader to bind the clamp before DNA. This kinetic preference could be...
Rates of DNA Versus β Clamp Binding to the γ Complex Clamp Loader

established simply by the clamp loader binding the clamp at a faster rate than DNA. Alternatively, given that the clamp loader has three ATP binding sites, sequential filling of ATP sites and incremental conformational changes could allow the clamp loader to bind the clamp before DNA. In other words, a subset of sites bind ATP and induce conformational changes that promote β binding faster than the subset of sites that bind ATP and make conformational changes to induce DNA binding. In this article, the questions of whether there is a kinetic preference for γ complex to bind β prior to DNA and how the overall rates of ATP-induced conformational changes contribute to the rates of clamp and DNA binding are addressed.

EXPERIMENTAL PROCEDURES

Nucleotides and Oligonucleotides—Concentrations of ATP (Amersham Biosciences/GE Healthcare) diluted with 20 mM Tris-HCl (pH 7.5) and ATPγS (Roche Diagnostics) diluted with water (Roche Diagnostics) were determined by measuring the absorbance at 259 nm and using an extinction coefficient of 15,400 M⁻¹ cm⁻¹. Synthetic oligonucleotides were obtained from Integrated DNA Technologies (Coralville, IA) and purified by 12% denaturing polyacrylamide gel electrophoresis. The sequences of the 60-nucleotide template and the complementary 30-nucleotide primer are as follows: 5'--TTC AGG TCA GAA GGG TTC TAT CTC TGT GCA GAA TGT CCC TTT TAT TAC TGG TCG TGT-3' and 5'-ACA CGA CCA GTA ATA AAA GGG ACA TTC (C6dT) GG-3', where C6dT is a T with a C6 amino linker that was covalently labeled with 7-diethylaminocoumarin-3-carboxylic acid succinimidyl ester (Invitrogen) as described (16, 21). Primed templates were labeled. The QuikChange mutagenesis kit (Stratagene) was used for site-directed mutagenesis with the following primers: used for site-directed mutagenesis with the following primers: Q299C, 5'-CAT CTG GAA GCT GGC and Q299R, 5'-CAT CTG GAA GCT GGC TCA GAT CTC CTA AAG CAG CAG CC-3'. Site-directed mutagenesis was performed to change the glutamine 299 to cysteine in a β construct in which surface cysteines 260 and 333 were replaced with serine so that Cys-299 could be selectively labeled. The QuikChange mutagenesis kit (Stratagene) was used for site-directed mutagenesis with the following primers: Q299C, 5'-CAC CGC CAA CAA CCC GGA ATG CGA AGA AGC GGA AGA GAT C; C260S, 5'-CAT CTC GAA GCT GGC TCA GAT CTC CTC AAG CAG CAG CC-3'; and C333S, 5'-CTG AAC GCG CTC AAA CTG AGA ACG TCC GCA TGA TGC. Concentrations of γ complex were determined by measuring the absorbance at 280 nm in 6 M guanidine hydrochloride and using the calculated extinction coefficient (220,050 M⁻¹ cm⁻¹). The β clamp was covalently labeled on Cys-299 with N-(1-pyrene)maleimide (Molecular Probes) and purified as described previously (27, 28). The protein concentration of βPY was determined using the modified Lowry protein assay (Pierce). Based on the concentration of protein and the concentration of pyrene determined by its absorbance, the typical labeling efficiency was 60%.

Steady State Fluorescence Assays—Fluorescence emission spectra were measured using a QuantaMaster QM-1 fluorometer (Photon Technology International) or an Edinburgh Analytical Instruments FS900 fluorometer. 7-Diethylaminocoumarin-3-carboxylic acid (DCC)-labeled DNA was excited at 440 nm and emission scanned from 450 to 550 nm using a 3.6-nm bandpass on the Edinburgh SF900 fluorometer. PY-labeled β was excited at 345 nm and emission scanned from 355 to 455 nm using a 5-nm bandpass on the Photon Technology International fluorometer. Equilibrium binding of γ complex to βPY was done by adding reagents sequentially to the cuvette starting with 72 µl of assay buffer with ATP, followed by 4 µl of βPY, and finally 4 µl of γ complex for a total reaction volume of 80 µl. Stoichiometric binding of γ complex to βPY or DNA-DCC was done by adding reagents sequentially to the cuvette starting with 68 µl of assay buffer, followed by 4 µl of βPY or p/t-DNA-DCC, 4 µl of γ complex, and finally 4 µl of ATP (or ATPγS for DNA) for a total reaction volume of 80 µl. For both equilibrium and stoichiometric βPY binding assays, emission spectra were measured following each addition, and relative intensities at 375 nm were plotted as a function of γ complex concentration. The buffer (+ATP) background was subtracted from all spectra and made relative to the free PY or DCC signal during analysis. The dissociation constant (KD) of the βPY-γ complex was calculated by fitting the observed intensity data (Iobs) to Equation 1 in which γs is the concentration of γ complex, β is the concentration of βPY, Imin is the intensity of free βPY, and Imax is the intensity of γ complex-βPY. The intensity of γ complex-βPY (Imax) and KD were fit as adjustable parameters by nonlinear regression using KaleidaGraph.

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I_{\text{obs}} = \frac{(K_D + \gamma_s + \beta) - \sqrt{(K_D + \gamma_s + \beta)^2 - 4\gamma_s\beta}}{2\beta} (I_{\text{max}} - I_{\text{min}}) + I_{\text{min}}
\]

(Eq. 1)

Equation 1 was also used to fit the quench in DCC fluorescence as a function of γ complex in the DNA binding assay to calculate the relative intensity of a clamp loader-DNA complex and a KD value. In this case, the concentration of DNA was substituted for β, Imax represents the intensity of free DNA, and Imin represents the intensity of bound DNA.

Pre-Steady State Fluorescence Assays—Assays were done using an Applied Photophysics SX20MV stopped-flow apparatus at 20 °C. Single-mix reactions were performed by mixing equal volumes (50 µl) of reagents immediately before they entered the cuvette. Data were collected for a total of 5 s at intervals of 1 ms. Final concentrations for each experiment in assay buffer with 4% glycerol are indicated in the figure legends and/or under “Results.” Measurements of DCC fluorescence were made using a 455-nm cut-on filter to collect emission while exciting at 430 nm with a 3.72-nm bandpass. A 365-nm cut-on filter was used to collect PY emission when exciting at 345 nm using a 3.72-nm bandpass for βPY binding experiments.
Rates of DNA Versus β Clamp Binding to the γ Complex Clamp Loader

Time courses for \( \beta^\text{PY} \) binding were empirically fit to a single exponential rise, \( y = a(1 - e^{-k_{\text{obs}}t}) + c \), using KaleidaGraph to determine values for observed rate constants, \( k_{\text{obs}} \), as a function of γ complex concentration.

**Kinetic Modeling**—DNA binding data in Figs. 4C, 6, and 7 were globally fit to the model illustrated in Fig. 8 using DynaFit to calculate the rate constants given in the figure. All rate constants were treated as adjustable parameters; none were set at fixed values. The relative intensity for free DNA and for the initial clamp loader-DNA complex prior to the DNA-induced conformational change were set to 1, and the relative intensities for all other clamp loader-DNA complexes were set to the value of 0.58 derived from the titration in Fig. 4B. Concentrations of protein and DNA were fixed based on the concentrations determined experimentally from absorbances at 280 and 260 nm, respectively, as described above. In experiments in which γ complex was preincubated with ATP, the “incubate” and “dilute” functions in DynaFit were used to generate the equilibrium concentrations of the two conformational states of γ complex that exist in the presence of ATP (15). These functions were used to: 1) mix a solution of γ complex, twice as concentrated as in the final reaction, with ATP; 2) dilute the γ complex in half to the final concentration; and 3) incubate the solution for 1 s in silico to generate the equilibrium populations of conformational states that are added to DNA to initiate the binding reactions. In experiments in which the γ complex was not preincubated with ATP, reactions were initiated in silico by adding free γ complex to ATP and DNA.

**RESULTS**

**Equilibrium Binding of γ Complex to β**—The clamp loader when charged with ATP has a high affinity for the clamp. The equilibrium dissociation constant (\( K_D \)) of γ complex-β is in the low nanomolar range (18). In previous studies, an anisotropy assay was used to measure γ complex-β binding (21, 28), but a limitation of this assay is that it is not sensitive enough to work at low nanomolar protein concentrations, where \( K_D \) values are accurately determined. Therefore, a more sensitive intensity-based fluorescence assay was developed to measure clamp loader-clamp binding. In this assay, the Gln-299 in β was mutated to Cys, and Cys-299 was covalently labeled with PY. Based on available structural data and similarities between the *E. coli* and yeast clamps and clamp loaders (5, 29, 30), residue 299 in β is likely to be near a site where a γ subunit contacts the surface of the β clamp in a clamp loader-clamp complex (Fig. 1). Given the head-to-tail symmetry of the β dimer and the anticipated contacts between γ complex and β, the fluorescence of PY is likely to be affected by γ complex binding only at the position near the middle γ subunit (Fig. 1). When γ complex binds to \( \beta^\text{PY} \), PY fluorescence increases (Fig. 2A). At saturating concentrations of γ complex, the intensity of PY is just over two and a half times greater than for free \( \beta^\text{PY} \).

This increase in PY fluorescence was used to measure equilibrium binding of γ complex to \( \beta^\text{PY} \) and calculate the dissociation constant for the interaction. A \( K_D \) value of 5.0 ± 2.4 nM was calculated from three independent experiments at 10 nM \( \beta^\text{PY} \) (average values are shown in Fig. 2B). This \( K_D \) value is in agreement with the value of 3.2 nM previously reported (18).

Binding was measured at a higher concentration of \( \beta^\text{PY} \) (80 nM) to determine the dissociation constant for the weaker ATP-independent binding reaction (Fig. 2C, filled circles) and to determine the stoichiometry of binding in the presence of ATP (Fig. 2C, filled squares). The \( K_D \) value calculated for ATP-independent binding was 135 ± 32 nM, which is in agreement with the value of 151 nM reported previously (28). In the presence of ATP, γ complex binds β with the expected 1:1 stoichiometry as indicated by achieving saturation at a concentration of around 80 nM γ complex, and these data also give a calculated \( K_D \) value of 2.9 nM for ATP-dependent binding. Interestingly, the maximum increase in PY fluorescence in assays without ATP is less than 2-fold, whereas it is greater than 2.5-fold in the presence of ATP. This difference could reflect differences in conformational states of γ complex and/or β in the presence and absence of ATP. ATP is shown to cause a conformational change in γ complex that gives it a high affinity for the clamp (18). The clamp is likely to be open in clamp loader-clamp complexes in assays with ATP, whereas it is likely to be closed in clamp loader-clamp complexes in assays without ATP (31).

**Kinetics of γ Complex Binding β**—To determine the rate of γ complex binding β, the increase in PY intensity was measured in real time when a solution of γ complex and ATP was mixed with a solution of \( \beta^\text{PY} \) and ATP. Reactions contained 20 nM \( \beta^\text{PY} \) and γ complex concentrations of 10, 20, 40, and 80 nM (Fig. 3A). These time courses were empirically fit to exponential rises to calculate apparent rate constants, \( k_{\text{obs}} \). These \( k_{\text{obs}} \) values were plotted as a function of γ complex concentration and fit to line to calculate an apparent on-rate constant, \( k_{\text{on(app)}} \), of 2.3 × 10⁷ M⁻¹ s⁻¹ and an apparent dissociation rate constant, \( k_{\text{off(app)}} \), of 0.14 s⁻¹ from the slope and y intercept, respectively (32) (Fig.
Rates of DNA Versus β Clamp Binding to the γ Complex Clamp Loader

FIGURE 2. Equilibrium binding of γ complex to β. A, emission spectra of βγ were taken at excitation wavelength of 345 nm. The light gray trace is a scan of free βγ and the black trace is after the addition of γ complex and ATP. Final concentrations were: 80 nM βγ, 240 nM γ complex, and 0.5 mM ATP. B, equilibrium binding of γ complex to β was determined by measuring the intensity of PY as a function of γ complex concentration, where γ complex was titrated into βγ solutions in assay buffer containing ATP. Final concentrations were: 10 nM βγ and 0.5 mM ATP in assay buffer. C, stoichiometric binding of γ complex to β was measured in the absence (circles) and presence (squares) of ATP. The γ complex and ATP were added sequentially to a solution of βγ in assay buffer. The relative intensity of PY at 375 nm is plotted as a function of γ complex concentration. Final concentrations after the addition of ATP were 80 nM βγ and 0.5 mM ATP in assay buffer.

3B). The apparent $K_D$ value calculated from these kinetic constants was 6 nM, which is in agreement with the value determined from equilibrium measurements.

Binding of γ Complex to DNA—A fluorescence intensity-based assay was used in which the primer strand of p/t-DNA is labeled with DCC 3 nucleotides from the 3' end, p/t-DNA-DCC, was used to measure the kinetics of γ complex binding to DNA (21). When the clamp loader binds DNA at the primer/template junction, the environment of DCC is altered and the fluorescence decreases. The γ complex requires ATP binding for high affinity binding to DNA, but DNA binding triggers hydrolysis of ATP and dissociation of the clamp loader so that binding is transient (19, 33). Therefore, γ complex-DNA binding under equilibrium conditions was measured in assays with the non-hydrolyzable ATP analog, ATPγS. When γ complex binds p/t-DNA-DCC, the fluorescence is quenched and the emission maximum of DCC is blue-shifted by 5–6 nm (Fig. 4A). To determine the magnitude of the fluorescence quench and relative quantum yield of a clamp loader-DNA complex, p/t-DNA-DCC was titrated with γ complex in the presence of ATPγS (Fig. 4B). These titration data showed that the fluorescence of the protein-DNA complex is 58% of the fluorescence of free p/t-DNA-DCC.

Transient binding of γ complex to DNA in assays with ATP was measured in real time stopped-flow fluorescence experiments. A solution of γ complex and ATP was added to a solution of p/t-DNA-DCC and ATP, and the intensity of DCC was measured as a function of time (Fig. 4C). Binding rates were measured with increasing concentrations of γ complex and DNA at 125, 250, and 500 nM. In each case, a rapid decrease in DCC fluorescence due to γ complex binding p/t-DNA-DCC was followed by a rapid increase in fluorescence due to dissociation of γ complex from DNA following ATP hydrolysis. As the concentrations of γ complex and DNA were increased, the binding reaction became faster and the magnitude of the fluorescence quench increased. However, the magnitude of the decrease in fluorescence at the lowest point was not as large as measured in assays with ATPγS (Fig. 4B). Moreover, these reactions are not approaching a limiting value for the minimum fluorescence, but continue to decrease with increasing concentrations. Assuming that the magnitude of the quench in fluorescence is the same for clamp loader-DNA complexes in assays with ATP and ATPγS, this shows that the ATP hydrolysis-induced γ complex dissociation is fast relative to the DNA binding reaction so that only a fraction of p/t-DNA-DCC is bound when the fluorescence reaches a minimal value. Given a relative fluorescence of 58% for the clamp loader-DNA complex, about 14% of the p/t-DNA-DCC is bound when the intensity reaches a minimum value of about 0.94 in the reaction containing 500 nM γ complex and p/t-DNA-DCC.
Rates of DNA Versus β Clamp Binding to the γ Complex Clamp Loader

It is possible that the rates of these conformational changes govern the rates of clamp and DNA binding to the clamp loader. To test this possibility, the rate of γ complex binding βPY was measured in assays in which γ complex was preincubated with ATP (Fig. 5A, gray trace) and in which γ complex was added to ATP at the same time as βPY (Fig. 5A, black trace). After mixing, both reactions contained identical concentrations of βPY (200 nM), γ complex (200 nM), and ATP (0.5 mM). There was no difference between the binding rates when γ complex was preincubated or not with ATP before adding β. This suggests that ATP binding and ATP-induced conformational changes that give γ complex a high affinity for β are relatively rapid. When βPY binding was measured in the absence of ATP (Fig. 5B, gray trace), binding was relatively fast but a smaller fraction of β was bound than in a reaction with ATP (Fig. 5B, black trace). The smaller fraction bound is consistent with the weaker ATP-independent binding reaction (Fig. 2C). This result provides additional support that ATP binding and ATP-induced conformational changes that give γ complex a high affinity for β are relatively rapid.

DNA Binding with and without ATP Preincubation—To determine whether the rate of ATP binding and ATP-induced conformational changes could limit the rate at which γ complex binds DNA, DNA binding was measured in real time assays in which γ complex was preincubated with ATP or in which γ complex was added to ATP at the same time as p/t-DNA-DCC. When a solution of γ complex and ATP was added to a solution of p/t-DNA-DCC and ATP (ATP preincubation), a rapid quench in fluorescence occurred that reached a minimum value in about 60–70 ms and was followed by an increase in fluorescence that reached a steady state level within about 500 ms (Fig. 6, black trace, gray line). In contrast, when a solution of γ complex without ATP was added to a solution of p/t-DNA-DCC and ATP (no ATP preincubation), the decrease in DCC fluorescence is slower overall (Fig. 6, gray trace, black line). There is a short lag of at least 25 ms before fluorescence begins to decrease and the rate of decrease is slower, such that the maximal quench occurs between 150 and 200 ms. Final concentrations of γ complex (200 nM) and pt-/DNA-DCC (200 nM) were the same in both reactions. Because the overall binding rate is slower, but the subsequent rates of ATP hydrolysis and γ complex-DNA dissociation are likely to be the same, the amplitude of the change in fluorescence is smaller when the γ complex was not preincubated with ATP. This experiment was repeated with ATPγS instead of ATP to measure DNA binding in the absence of DNA-triggered ATP hydrolysis and subsequent clamp loader dissociation (supplemental Fig. S1).

Although the reaction in which γ complex was preincubated with ATPγS was slightly faster than the non-preincubated case, overall both reactions were much slower and took 6–8 s to reach completion. It is possible that DNA binding reactions are slower in assays with ATPγS because ATPγS does not have the efficacy of ATP in promoting conformational changes that allow the clamp loader to bind DNA. We have also observed a decrease in the affinity of γ complex for β in assays with ATPγS compared with ATP (21). For this reason, we chose to focus on assays with ATP even though the kinetics are complicated by...
hydrolysis. In any case, a slow ATP-dependent step is bypassed when γ complex is preincubated with ATP.

Effect of ATP Concentration on γ Complex-DNA Binding Kinetics—Either a slow ATP binding step or a slow ATP-induced conformational change could limit the rate at which γ complex binds DNA. To determine which step is slow, rates of γ complex-DNA binding were measured as a function of ATP concentration in reactions in which γ complex was not preincubated with ATP. A solution of γ complex and p/t-DNA-DCC was mixed with solutions of ATP, and DCC fluorescence was measured as a function of time. In Fig. 7, reactions containing 25 (A, red), 50 (B, green), 250 (C, orange), and 500 μM (D, blue) ATP are plotted in the same graph with a plot of a reaction containing 100 μM ATP (black trace). At the lowest concentration of ATP used, the reactions approached a maximal rate, becoming slightly faster with increasing concentrations of ATP, and reaching a maximum rate at 100 μM ATP. In all previous experiments, reactions contained 500 μM ATP, therefore, ATP binding was unlikely to be rate-limiting. The slower rates of DNA binding in assays in which γ complex was not preincubated with ATP must be due to a slow ATP-induced conformational change.

Kinetic Modeling of DNA Binding Reactions—DNA binding data were modeled to get an estimate of the rate of ATP-induced conformational change that precedes DNA binding to determine whether there could be a kinetic preference for binding β prior to DNA. A kinetic model from previous studies was adapted to fit these DNA binding data (15). The kinetic model was developed based on experiments measuring rates of ATP hydrolysis in reactions in which the γ complex was preincubated with ATP for varying periods of time before adding p/t-DNA. The earlier studies revealed that the rate of ATP hydrolysis was limited by an ATP-dependent conformational change and suggested that a slow ATP-induced conformational change activates the clamp loader for DNA binding. Experiments in Figs. 6 and 7 confirm this by establishing that a slow ATP-induced conformational change precedes DNA binding. The complete kinetic mechanism is quite complex and contains too many forward and reverse rate constants to define in a single set of experiments. Therefore, the kinetic model was simplified by including only forward rates for most of the steps and allowing
three molecules of ATP to bind and three molecules of ADP to dissociate as a unit in a single step. All of the rate constants shown in Fig. 8 were derived by globally fitting the data in Figs. 4C, 6, and 7 to the model shown using DynaFit (34). The forward and reverse rate constants for the ATP-induced conformational change obtained from DNA binding data were 3.3 and 1.7 s$^{-1}$ compared with values of 6.5 and 3.9 s$^{-1}$, respectively, obtained previously from fits of ATP hydrolysis data (15). After the conformational change, p/t-DNA binding occurred at a rate of 4.0 $\times$ 10$^7$ M$^{-1}$ s$^{-1}$. The rate of DNA binding is similar to the rate of clamp binding (2.3 $\times$ 10$^7$ M$^{-1}$ s$^{-1}$) and on the order of what would be expected for a diffusion-limited rate for two macromolecules interacting. However, in reactions in which the $\gamma$ complex is not preincubated with ATP, the rate of DNA binding is limited by the slow, 3.3 s$^{-1}$ conformational change, whereas the rate of $\beta$ binding is not.

The interaction with p/t-DNA triggers a change in the clamp loader that makes the ATP sites competent for hydrolysis, and in the model used here, three molecules of ATP are hydrolyzed sequentially before DNA is released. ADP release allows the clamp loader to go through the cycle again. The DNA binding data do not provide a direct measure of ATP hydrolysis, and in terms of these data, the ATP hydrolysis steps in the kinetic model provide a time lag between observed DNA binding and

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**FIGURE 5.** Rates of $\gamma$ complex binding $\beta$ with and without preincubation of $\gamma$ complex with ATP. A, the rate of $\gamma$ complex binding $\beta^{\gamma}$ was measured in assays with (gray trace) and without (black trace) preincubation of $\gamma$ complex and ATP. The relative intensity of PY is plotted as a function of time for a reaction in which a solution of $\gamma$ complex and ATP was added to a solution of $\beta^{\gamma}$ and ATP (gray trace) and a reaction in which a solution of $\gamma$ complex was added to a solution of $\beta^{\gamma}$ and ATP (black trace). The inset is the first 0.2 s of the reaction time courses. B, rates of $\gamma$ complex binding $\beta^{\gamma}$ were measured in the absence of ATP (gray trace) and in the presence of ATP but without preincubation of $\gamma$ complex with ATP (black trace). Final concentrations were: 200 nM $\beta^{\gamma}$ and $\gamma$ complex and 0.5 mM ATP in assay buffer with 4% glycerol.
release. The model used to fit DNA binding data here differs from that used to fit ATP hydrolysis data in that two molecules of ATP were hydrolyzed rapidly and the third slowly in the ATP hydrolysis model. The DNA binding data could be fit equally well by a model in which ATP hydrolysis occurred in two phases (data not shown), hydrolysis of two molecules of ATP rapidly before DNA release, and hydrolysis of one molecule of ATP slowly after DNA release as proposed previously based on ATP hydrolysis data (15). When DNA binding data were fit to this two-phase model for ATP hydrolysis, similar rates for ATP binding \( (k_{\text{on(\ATP)}} = 12.6 \text{ M}^{-1} \text{ s}^{-1}) \), the ATP-induced conformational change \( (k_{\text{conf}} = 3.3 \text{ s}^{-1}, k_{\text{rev(conf)}} = 1.7 \text{ s}^{-1}) \), DNA γ complex binds the clamp or DNA faster, and how preincubation of the clamp loader with ATP to form the ATP-activated state affects the rates of clamp loader-clamp and clamp loader-DNA binding.

To measure the rate of clamp binding, a sensitive fluorescence intensity-based assay was developed in which the β clamp was covalently labeled with PY. The intensity of PY increases when the clamp loader binds the clamp. An advantage of this assay is that clamp loader-clamp complex can be measured directly in solution and in real time. The assay is sensitive enough to measure the high affinity binding of γ complex to β under equilibrium conditions as well as measuring binding

**DISCUSSION**

Clamp loaders catalyze the assembly of sliding clamps onto DNA for use by DNA polymerases. To load clamps, the affinity of the clamp loader for the clamp and DNA must be modulated. Initially, the clamp loader must have a high affinity for the clamp and DNA to bring these macromolecules together, but then the affinity must decrease so that the clamp loader can release the clamp onto DNA. This affinity modulation is achieved by ATP binding and hydrolysis. ATP binding activates the clamp loader for binding the clamp and DNA, whereas ATP hydrolysis deactivates the clamp loader for releasing the clamp and DNA. Thus, the clamp loading reaction can be divided into two phases depending on the ATP requirements: 1) ATP binding-dependent formation of a ternary clamp loader-clamp-DNA complex, and 2) ATP hydrolysis-dependent decay of the ternary complex releasing the clamp on DNA. Much of our previous work focused on the second phase of the reaction by adding a preformed clamp loader-clamp complex to DNA to rapidly form the ternary complex via a single pathway. Here, we focused on the first phase of the reaction, formation of a ternary clamp loader-clamp complex. We asked whether

**FIGURE 7. Effect of ATP concentration on γ complex-DNA binding kinetics.** Rates of DNA binding were measured as a function of ATP concentration in a reaction in which γ complex was not preincubated with ATP. A solution of γ complex and p/t-DNA-DCC was mixed with a solution of ATP, and the DCC fluorescence was measured as a function of time. Final concentrations were: 200 nM p/t-DNA-DCC and γ complex; and A, 25 μM (red trace); B, 50 μM (green trace); C, 250 μM (orange trace); and D, 500 μM (blue trace) ATP relative to the same reaction at 100 μM ATP (black trace) in assay buffer with 4% glycerol.
induced conformational changes must precede clamp or DNA binding, the overall rate of these conformational changes will contribute to observed binding rates in experiments in which γ complex is not preincubated with ATP prior to adding the clamp or DNA. On the other hand, if the γ complex is preincubated with ATP before adding the clamp or DNA, these conformational changes can take place during the preincubation period and binding rates will not be influenced by the rate of the conformational changes. When the γ complex was preincubated with ATP, the rate of p/t-DNA binding was faster than for reactions in which γ complex was not allowed to bind ATP before p/t-DNA (see Fig. 6). DNA binding kinetics measured as a function of ATP concentration (see Fig. 7) demonstrated that the slower DNA binding kinetics were not due to a slow ATP binding reaction, but instead must be the result of slow ATP-induced conformational changes. In contrast, the observed rate for β binding was the same regardless of whether the γ complex was preincubated with ATP or not (see Fig. 5). The rapid β binding in experiments without ATP preincubation is unlikely to be due to an ATP-independent binding reaction because the ATP-independent binding interaction is weaker (see Fig. 2C) and could only account for a fraction of the binding events (see Fig. 5B). ATP binding and subsequent ATP-induced conformational changes are rapid relative to the rate of β binding. Given that only DNA binding rates are limited by a slow ATP-induced conformational change, this would suggest that a different set of conformational changes gives rise to β and DNA binding. It will be interesting to see which conformational changes promote clamp opening. The first “fast” conformational change could promote both clamp binding and clamp opening as illustrated in Fig. 9. Alternatively, the second slower conformational change could promote clamp opening and subsequent DNA binding.

There are a number of interesting observations in the literature that can be used to speculate about a mechanism by which two distinct sets of conformational changes could occur sequentially to promote β and then DNA binding. The γ complex has three ATP sites and sequential filling of these sites could promote sequential sets of conformational changes. A
subset of ATP sites could fill first to induce conformational changes rapidly that promote β binding. A slow conformational change could then allow the remaining sites to fill with ATP and promote DNA binding. Incubation of the γ complex with ATP prior to adding β or DNA would give all the sites time to fill with ATP and allow the conformational changes to occur so that the γ complex could bind either the clamp or DNA. Studies with arginine finger mutants of the E. coli and yeast clamp loaders, which bind but do not “sense” or respond to bound ATP, show that ATP binding at different sites has differential effects on clamp and DNA binding (28, 35). Similarly, studies of single Walker A mutants that reduce ATP binding to the yeast clamp loader showed that mutations of individual sites had differential effects on DNA binding (36). Both Arg finger (35) and Walker A (36) mutations that affect ATP sensing/binding to the replication factor C (RFC) 2 and RFC3 subunits of the yeast clamp loader showed that mutations of individual sites had differential effects on DNA binding. In a crystal structure of the minimal E. coli clamp loader (γ,δδ′), the middle γ subunit does not contain a bound ATPγS molecule, whereas the other two γ subunits are bound to ATPγS (37). Biochemical characterization showed that this minimal clamp loader (γ,δδ′) has greatly reduced DNA binding activity compared with γ complex (γ5δδ′ψχ), which is consistent with the idea that ATP binding to the middle γ subunit is important for DNA binding (21). That is not to say that a single subunit binds DNA, but instead that ATP binding at one (or a subset of sites) is important for inducing conformational changes within the clamp loader that as a whole increase DNA binding activity. If a slow conformational change were to open this site and cause it to fill last, then this could provide a mechanism for sequential binding of the clamp and DNA.

The Conformational Change That Gives γ Complex a High Affinity for DNA—The observation that the rate of p/t-DNA binding is limited by the rate of ATP-induced conformational changes is consistent with results from our previous work measuring rates of p/t-DNA-triggered ATP hydrolysis by the γ complex. As with the DNA binding kinetics, we found that rates of DNA-triggered ATP hydrolysis were faster when the γ complex was preincubated with ATP than when there was no ATP preincubation, and that the slower rate was not due to slow ATP binding (15). Based on rates of ATP hydrolysis as a function of preincubation time with ATP, a forward rate constant of 6.5 s⁻¹ and reverse rate constant of 3.9 s⁻¹ was calculated for the ATP-induced conformational changes. These values are in good agreement with the values of 3.3 and 1.7 s⁻¹ for the forward and reverse rate constants calculated from kinetic modeling of the DNA binding reaction (see Fig. 8) given the complexity of this reaction and that both data sets were fit independently. Both sets of experiments predict that after equilibration with ATP a little over 60% of the clamp loaders (63% from ATPase experiments and 66% from DNA binding experiments) exist in the conformational state that has high affinity for p/t-DNA. Interaction with the β clamp increases the affinity of the clamp loader for DNA, and it is interesting to speculate that the β clamp may do this by stabilizing the conformational state with a high affinity for DNA and shifting the equilibrium to favor this species (16). Studies with a minimal form of the clamp loader, γ,δδ′, missing the χ and ψ subunits support this idea. The minimal clamp loader is defective in ATP-dependent DNA binding activity. The β clamp can rescue the ATP-dependent DNA binding activity of the minimal clamp loader most likely by stabilizing or promoting the conformational state with high affinity for DNA (21). It is likely that the β clamp also increases the affinity of the clamp loader for DNA by directly interacting with the DNA duplex. The central cavity of the β ring is lined with positively charged amino acid residues (4) and these residues interact with the sugar-phosphate backbone of the duplex (3).

A Kinetic Preference for the Clamp Loader to Bind β before DNA—When equilibrated with ATP, the E. coli γ complex can bind either β or DNA. The same is true for the eukaryotic clamp loader, RFC (38). Although these clamp loaders can bind either the clamp or DNA, productive clamp loading is likely to require these clamp loaders to bind the clamp first. In contrast, the bacteriophage T4 clamp loader can productively load clamps by binding either DNA or the clamp first (39, 40). This may be due to differences in the solution structures of the clamps. The bacteriophage T4 gp45 clamp exists as an open ring in solution (41, 42), whereas the E. coli β clamp (43) and eukaryotic clamp are likely to exist as closed rings in solution (44). When bound to genomic DNA, the geometry of the clamp loader-DNA complex likely prevents the clamp loader from productively binding a closed clamp. In addition, binding of the bacterial and eukaryotic clamp loaders to p/t-DNA triggers rapid ATP hydrolysis and dissociation of the clamp loader from the DNA, such that the clamp loader-DNA complex is transient and not likely to be long-lived enough to efficiently bind clamps (19, 45). Nonproductive interactions between the clamp loaders and DNA would reduce the overall efficiency of clamp loading by engaging the clamp loader in futile cycles of DNA binding and ATP hydrolysis. A mechanism that favored clamp binding prior to DNA binding would increase the overall efficiency of clamp loading. It is possible that the rate-limiting ATP-induced conformational changes that precede DNA binding but not clamp binding provide a kinetic preference for the clamp loader to bind the clamp before DNA and increase the overall efficiency of clamp loading. The fastest rate at which the clamp loader can bind DNA is limited by rate of the ATP-induced conformational change (3.3 s⁻¹), but binding could be slower if DNA concentrations are limiting. The rate of the conformational change that promotes high affinity β binding is faster so that the rate of β binding may be a function of the concentration of β in the cell, 500 nm assuming 300 copies of β (46) in a 1 × 10⁻¹⁵ liter cell volume, and the bimolecular rate constant for binding, 2.3 × 10⁻¹⁵ m⁻¹ s⁻¹, to give an effective on-rate of 11 s⁻¹. This effective on-rate assumes that β on the lagging strand is rapidly recycled by excess δ subunit and/or clamp loader in the cell such that all the β is free to be loaded (47). Even at saturating concentrations of DNA, given these rates, the γ complex would bind β first at least 75% of the time. During active replication, the timing of primer synthesis on the lagging strand could also
Rates of DNA Versus β Clamp Binding to the γ Complex Clamp Loader

provide a mechanism for regulating the order of β and DNA binding. This could be accomplished by synthesizing primers at a rate that would allow the clamp loader to bind a clamp before a new primed template site is formed.

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DNA: Replication, Repair, Recombination, and Chromosome Dynamics: A Slow ATP-induced Conformational Change Limits the Rate of DNA Binding but Not the Rate of β Clamp Binding by the Escherichia coli γ Complex Clamp Loader

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