

Dynamics of the E. coli β -Clamp Dimer Interface and Its Influence on DNA Loading

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ABSTRACT The ring-shaped sliding clamp proteins have crucial roles in the regulation of DNA replication, recombination, and repair in all organisms. We previously showed that the Escherichia coli β -clamp is dynamic in solution, transiently visiting conformational states in which Domain 1 at the dimer interface is more flexible and prone to unfolding. This work aims to understand how the stability of the dimer interface influences clamp-opening dynamics and clamp loading by designing and characterizing stabilizing and destabilizing mutations in the clamp. The variants with stabilizing mutations conferred similar or increased thermostability and had similar quaternary structure as compared to the wild type. These variants stimulated the ATPase function of the clamp loader, complemented cell growth of a temperature-sensitive strain, and were successfully loaded onto a DNA substrate. The L82D and L82E I272A variants with purported destabilizing mutations had decreased thermostability, did not complement the growth of a temperature-sensitive strain, and had weakened dimerization as determined by native trapped ion mobility spectrometry-mass spectrometry. The β L82E variant had a reduced melting temperature but dimerized and complemented growth of a temperature-sensitive strain. All three clamps with destabilizing mutations had perturbed loading on DNA. Molecular dynamics simulations indicate altered hydrogen-bonding patterns at the dimer interface, and cross-correlation analysis showed the largest perturbations in the destabilized variants, consistent with the observed change in the conformations and functions of these clamps.

SIGNIFICANCE The relationship between protein dynamics and function is still poorly understood. The bacterial β sliding clamp displays dynamic behavior at the dimer interface, raising the possibility that the clamp is transiently open. To test the role of dynamics in clamp function, we designed stabilizing and destabilizing mutations at and near the dimer interface. Clamp variants with destabilized dimer interfaces generally had altered thermal stability and dimerization and did not support bacterial growth. Molecular dynamics simulations support the model of interface perturbation because the variants with destabilizing mutations have dramatically altered hydrogen-bonding patterns at the dimer interface.

INTRODUCTION

The ring-shaped sliding processivity clamp protein of the DNA replication machinery has crucial roles in the regulation of DNA replication, recombination, and repair in all living organisms and some viruses. DNA polymerases replicate the genomes of entire organisms, and the sliding processivity clamp provides the speed and processivity required by DNA polymerases for efficient DNA replication $(1-3)$.

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The sliding clamp binds the polymerase and tethers it to DNA, allowing the polymerase to replicate DNA without dissociation from the template. Structural studies reveal that sliding clamps have a pseudo-sixfold symmetry and individual subunits are arranged in a head-to-tail conformation $(1,4–8)$ $(1,4–8)$ $(1,4–8)$. Clamps from bacterial species (4) (4) are homodimeric, with three domains in each monomer, whereas eukaryotic and archaeal species $(5,9-11)$ are trimers with two domains in each monomer. Each domain consists of two α -helices and two four-stranded antiparallel β -sheets; adjacent α -helices and a long curved β -sheet provide the interdomain interactions ([Fig. 1\)](#page-1-0). The Escherichia coli β -clamp increases the processivity and speed of replication

FIGURE 1 Design of β -clamp mutations to stabilize or destabilize the dimer interface. Monomer A is depicted in gray and Monomer B in purple on the three-dimensional representations. The locations of mutations in Domain 1 (T45 and T47 variants) are depicted with blue beads, while the mutations at the dimer interface (L82D, L82E, L82E I 272A, S107R, and S109R variants) are depicted with green beads on the protein.

by \sim 5000-fold and 100-fold, respectively [\(12–15](#page-13-0)). Clamp proteins slide on DNA and serve as platforms for multiple proteins to bind. The β -clamp interacts with all five E. coli DNA polymerases ([16–26\)](#page-13-1) and coordinates the polymerase switch during translesion synthesis [\(27–29](#page-13-2)). In addition, β interacts with replication initiation factors DnaA [\(30](#page-13-3)) and Had ([31\)](#page-13-4); mismatch repair proteins MutS and MutL; and DNA ligase A, which is responsible for Okazaki fragment maturation ([16\)](#page-13-1). Client interactions occur on the front face of the clamp at the $3'$ recessed end of DNA at primertemplate junctions.

Clamps are loaded on DNA by clamp-loader complexes, which are members of the $AAA+$ family ATPases ([32–35\)](#page-13-5). The loader functions as a switch $(36,37)$ $(36,37)$ $(36,37)$ and first binds ATP to bind to the clamp; ATP hydrolysis is not needed for clamp opening but rather for the clamp-loader ejection from the clamp and DNA ([38,39](#page-13-7)). The E. coli clamp loader, the γ -complex, contains one copy of each δ and δ' and three copies of γ or τ subunits. The γ/τ subunits harbor the ATPase function, and once ATP-binding sites are occupied, a spiral arrangement is triggered that complements the right-handed helix of DNA. The rearrangement of the clamp loader exposes a region of the δ subunit previously occluded by δ' and allows binding of δ to the β -clamp and the primer-template DNA ([40\)](#page-13-8). Available structural data of the γ -clamp-loader complex in its inactive state show an asymmetric conformation and the steric clashes precluding it from binding the clamp [\(40](#page-13-8)). The structure of the γ -complex bound to DNA demonstrates the active conformation of a symmetrically aligned clamp loader with all three ATPbinding sites occupied, elucidating the mechanism of ATP hydrolysis induced by the γ -complex binding DNA ([41\)](#page-13-9).

The stability of clamps as closed structures varies in different organisms, and trimers tend to be less stable than dimers [\(42–45](#page-13-10)). The ability of a clamp loader to open a clamp is determined by the stability of the clamp ring structure. The T4 bacteriophage clamp gp45 tends to be open in solution and unload readily from DNA as a trimer and monomerize [\(42–46](#page-13-10)), eliminating the need for the clamp loader gp44/62 to open the clamp. Heterotrimeric Sulfolobus solfataricus PCNA samples ring-open states in solution in the absence of its loader [\(47](#page-13-11)). The human clamp PCNA is less stable in solution than its nearly identical structural homolog from Saccharomyces cerevisiae [\(48](#page-13-12)). Evidence exists both for and against opening of yeast PCNA by its replication factor C (RFC) loader complex. It has been shown experimentally that PCNA opening is enhanced by \sim 20,000-fold when bound to the RFC complex compared to unbound PCNA ([49–52\)](#page-13-13). However, computational modeling of a mutated RFC that is unable to form a spiral arrangement appears to stabilize the open conformation of PCNA rather than destabilize its closed conformation ([53\)](#page-13-14). Human PCNA and the β -clamp have been shown to dissociate slowly from DNA, with $t_{1/2}$ of 24 and 60 min, respectively [\(42](#page-13-10)). The β -clamp forms a dimer with a K_D of $65-\sim600$ pM at a concentration three orders of magni-tude lower than yeast PCNA [\(52,54\)](#page-13-15), making the β -clamp the most stable clamp in solution and suggesting the need for clamp opening by the clamp loader. Other work indicates that Domain 1 at the dimer interface can undergo local unfolding in solution, suggesting that at least transient spontaneous opening is possible ([55,56](#page-14-0)).

In dimeric β , the helix harboring residues I272 and L273 of Domain 3 is unwound and distorted to facilitate a hydrophobic interaction with Domain 1 of the other β monomer to create a well-packed dimer interface [\(Fig. 1\)](#page-1-0). The crystal structure of a monomeric β -clamp (β I272A L273A) bound to the δ subunit of the γ -complex reveals that this helix is undistorted, and the formation of a well-packed dimer interface is prevented [\(57](#page-14-1)). In addition, a five-residue loop (274– 278) that is part of a turn in the dimeric β structure and therefore occluded from interaction with δ in the closed clamp provides an interaction surface for δ and monomeric β ([57\)](#page-14-1). Although δ does not directly interact with the dimer interface of closed β , the above rearrangements are attributed to the interaction of δ with monomeric β . The δ subunit does not open β at the dimer interface but rather stabilizes one monomer, preventing ring closure. The strained conformation of the clamp monomers held together by interactions at the dimer interface can provide the energy for δ to induce β to open or trap β in an open conformation. The monomeric β shows greater flexibility in solution compared to the dimeric structure in HXMS experiments ([55\)](#page-14-0), and this dynamic behavior is confirmed by steered molecular dynamics (MD) simulations ([58\)](#page-14-2). The simulated monomeric β is dynamic and opens more in solution relative to the dimer. The tilt observed in Domain 3 relative to the other domains of the monomeric clamp is the same as in the crystal structure of the monomeric β - δ complex. Fluorescence-based experiments show that clamp binding is fast, whereas clamp opening is a slower process that does not require ATP hydrolysis ([39\)](#page-13-16). The δ subunit alone or the γ -complex depleted of ATP has been shown to unload dimeric β in vitro ([59\)](#page-14-3). Binding to δ stabilizes the open conformation, shifting the equilibrium between closed and open β toward an open conformation [\(57,58\)](#page-14-1).

We previously found that the β -clamp undergoes opening transitions in solution and occupies conformational states in which Domain 1 of β is particularly dynamic, undergoing local cooperative unfolding ([55,56](#page-14-0)). These findings, along with other work demonstrating that β is flexible and under tension in the dimeric state, suggest that β may transiently sample multiple conformations, some of which could be more favorable for clamp-loader binding. The goal of this study is to better understand how the stability of the β -clamp dimer interface influences its dynamic properties and the opening and closing of its ring structure underlying clamp loading. We made a broad set of mutations aimed at stabilizing the closed conformation or destabilizing the dimer interface. Biochemical, biophysical, and computational analyses reveal correlations between dimer stability and clamp opening and loading functions of the β -clamp variants.

MATERIALS AND METHODS

Protein expression and purification

The *dnaN* gene encoding the β -clamp was previously cloned into pET11T and expressed and purified generally as described ([55\)](#page-14-0). Detailed methods can be found in the [Supporting Materials and Methods.](#page-12-3) Mutations in this construct were introduced using the QuikChange II Site-Directed Mutagenesis Kit (Agilent Technologies, Santa Clara, CA). Mutations were confirmed by DNA sequencing (Eton, Charlestown, MA). The γ -clamploader complex was expressed and purified as previously described [\(41](#page-13-9)).

For isothermal titration calorimetry (ITC) experiments, the dnaN gene and a fragment of the holA gene encoding the N-terminal domain of the E. coli γ -clamp-loader complex δ subunit (residues 1–140; referred to as mini- δ) were subcloned into the pET28b+ vector (Novagen, Darmstadt, Germany) using the NdeI and XhoI sites, as described elsewhere [\(60](#page-14-4)). Point mutations in the β -clamp gene, encoding T45R, T47R, S107R, S109R, L82D, L82E, L82E I272A, and I272A L273A, were introduced following the modified inverse PCR procedure [\(61](#page-14-5)). The nucleotide sequence of the resulting constructs was confirmed by sequencing (Genewiz, Cambridge, MA). Recombinant mini- δ , wild-type (WT) β , and variant β -clamps T45R, T47R, S107R, S109R, L82E, L82E I272A, and I272A L273A were expressed and purified as described previously ([60\)](#page-14-4) and in the [Sup](#page-12-3)[porting Materials and Methods.](#page-12-3) The β -clamp L82D variant could not be purified in sufficient amounts for ITC experiments.

To test for a potential interaction between β and SSB, E. coli BL21 (DE3) strain containing pEAW134 (a gift from Dr. Michael Cox from the University of Wisconsin, Madison) expressing single-stranded DNA binding protein (SSB) [\(62](#page-14-6)) was transformed with pET28b expressing His-tagged β -clamp. Transformants were selected on Luria broth plates supplemented with ampicillin and kanamycin. A 50-mL starter culture in a shake flask (200 rpm) was grown overnight at 37° C. A 1-L culture was seeded and grown at 37° C until OD₆₀₀ of 1.0 was reached. Protein expression was induced with isopropyl β -D-1-thiogalactopyranoside to a final concentration of 1 mM at 30 $^{\circ}$ C for 4 h. Cells were harvested by centrifugation for 10 min at 6750 \times g at 4°C and frozen at -80°C or lysed immediately. Protein overexpression was confirmed by sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE). Affinity chromatography was used to investigate possible β and SSB interactions [\(63,64](#page-14-7)). Cell pellets containing overexpressed β and SSB were resuspended on ice in 50 mM HEPES (pH 7.5), 500 mM NaCl, 30 mM imidazole, 1 mM dithiothreitol, 15% glycerol (buffer A) supplemented with lysozyme and $+/-$ DNase I. Cells were lysed by repeated freeze/thaw cycles $(-80^{\circ}$ C followed by 37°C). Cell debris was pelleted by centrifugation at $14,000 \times g$ at 4° C for 1 h. The lysate was loaded onto a 2×5 mL HisTrap HP (GE Healthcare, Marlborough, MA) column and eluted with buffer A and increasing imidazole concentration to 0.5 M. Fractions were analyzed by SDS-PAGE.

Differential scanning fluorimetry

The melting temperature of the WT and variant β -clamps was determined using a thermofluor-based assay ([56,65](#page-14-8)). Optical 96-well clear PCR plates (Thermo Fisher Scientific, Waltham, MA) were used to analyze the melting temperature (T_m) of β proteins. Samples containing 5 μ M β protein and a final concentration of $10 \times$ Sypro Orange dye (Invitrogen, Carlsbad, CA) were prepared in a $20-\mu L$ reaction volume containing 20 mM HEPES (pH 7.5). The plates were sealed with an optical adhesive film (Applied Biosystems, Foster City, CA). The temperature in a CFX 96 Real-Time System (Bio-Rad, Hercules, CA) was raised from 4 to 100° C in 0.2°C increments every 10 s, and the fluorescence was detected. Samples were analyzed three times in duplicate, and average T_m and standard deviations are reported.

Native and denaturing PAGE

Purified WT β and variants (1 μ g each) were incubated in 1 \times loading buffer (62.5 mM Tris-HCl (pH 6.8), 0.01% Bromophenol Blue, 2.5% β -mercaptoethanol, 10% glycerol) before loading onto 12% PAGEr precast gels (Lonza, Basel, Switzerland) for native PAGE. Proteins were incubated in loading buffer supplemented with 2% SDS at 95° C for 10 min before loading on 16% SDS-PAGE. Gels were stained with Coomassie gel stain.

Native nESI-TIMS-MS

Ion mobility mass spectrometry (MS) experiments were performed on a custom built nanoelectrospray ionization (nESI)-trapped ion mobility spectrometry (TIMS) unit coupled to an Impact Q-TOF mass spectrometer (Bruker, Billerica, MA) [\(66,67](#page-14-9)). The TIMS unit is run by custom software in LabView (National Instruments, Austin, TX) synchronized with the MS platform controls [\(67](#page-14-10)). Aliquots (10 μ L) of WT and clamp variants, dissolved in 10 mM NH₄Ac (native conditions) to 5 μ M, were loaded in a pulled-tip capillary biased at 1500 V to the MS inlet. Briefly, the ion mobility separation in a TIMS device is based on holding the ions stationary using an electric field (E) against a moving buffer gas (68) . TIMS separation depends on the gas flow velocity (v_e) , elution voltage $(V_{elution})$, ramp time (t_{ramp}) , and base voltage (V_{out}) [\(66](#page-14-9)). The mobility, K, is defined by

$$
K = \frac{v_g}{E} \tilde{=} \frac{A}{(V_{elution} - V_{out})}.
$$
 (1)

The mobility calibration was performed by varying the ramp time to determine $V_{elution}$ as described ([69\)](#page-14-12). The mobility calibration constant A was determined using known reduced mobilities of Tuning Mix components from Agilent analyzed under the same experimental conditions. TIMS experiments were carried out using nitrogen (N_2) as a buffer gas at ambient temperature (T), with v_g set by the pressure difference between the funnel entrance (P1 = 2.6 mbar) and exit (P2 = 1.1 mbar). A radio frequency voltage of 200 V_{pp} at 880 kHz was applied to all electrodes. Ions were softly transferred and injected into the TIMS analyzer section injection to avoid collisional induced activation (i.e., $V_{\text{deflection}} = 80 \text{ V}$, $V_{\text{capillary}} = 70$ V, $V_{\text{funnel in}} = 0$ V, and $V_{\text{out}} = 60$ V; TIMS cell schematic shown in [Fig. S1\)](#page-12-3). The measured mobilities were converted into collision cross sections (CCSs, \AA^2) using the Mason-Schamp equation:

$$
\Omega = \frac{(18\pi)^{1/2}}{16} \frac{q}{(k_B T)^{1/2}} \left(\frac{1}{m} + \frac{1}{M}\right)^{1/2} \frac{1}{N} \times \frac{1}{K},\qquad(2)
$$

where q is the ion charge, k_B is the Boltzmann constant, N is the gas number density, m is the ion mass, and M is the gas molecule mass [\(68](#page-14-11)).

ITC

The interaction between the WT β or variant clamps with mini- δ was analyzed by ITC performed at 25°C (20°C for L82E, L82E I272A, and I272A L273A β variants) on a Nano-ITC (TA Instruments, New Castle, DE). The ITC experiment consisted of $20 \times 2.5 \mu L$ injections of mini- δ (0.6–0.7 mM) to the β -clamp solution (0.1 mM) repeated with 300 s interval between injections. The ITC data were corrected for heat measured in corresponding blank titrations of mini- δ into buffer. The data were analyzed using the NanoAnalyze software (TA Instruments) to obtain the dissociation constant K_D , association enthalpy ΔH , and binding stoichiometry parameter n.

ATPase assay and clamp loading assay

The ATPase assay was performed as per the BIOMOL Green manual (Enzo Life Sciences, Farmingdale, NY). The clamp loading assay was carried out by separating biotinylated-DNA-bound proteins from unbound proteins using streptavidin magnetic beads (NEB, Ipswich, MA). Both assays are described in detail in the [Supporting Materials and Methods.](#page-12-3)

Complementation assay

Chemically competent E. coli MS120 strain ([70](#page-14-13)) bearing temperature-sensitive mutations in the chromosomal dnaN gene was transformed with pET11T empty vector, plasmid encoding WT, or variant β -clamp using a standard transformation procedure (71) (71) . After outgrowth at 30°C, cells were plated on Luria broth agar plates in triplicate, supplemented with ampicillin, and grown at 30 or 37° C for 16–18 h. Experiments were repeated three times. Average colony-forming units and standard deviations are reported. Despite the likely low level of expression with pET vectors, β expressed from pET11T is found to complement a dnaN^{Ts} strain and was detected by immunoblot. Indeed, we previously observed sufficient expression from a pET vector of replication protein DnaE to complement a strain harboring a temperature-sensitive allele of $dn a E$ [\(72](#page-14-15)).

MD simulations

Seven different systems were prepared using the x-ray crystallographic structure of the E. coli DNA polymerase III β subunit (Protein Data Bank [PDB]: 1MMI) ([8\)](#page-12-4) containing two monomers (A and B) and subjected to MD simulations. The residues that are located on Monomer B are depicted with a prime sign. Each monomer within the proteins contains three different domains that are named alphanumerically with capital ''D'' and the domain numbers, D1, D2, and D3. The domains on Monomer B are indicated with a prime sign as DI' , $D2'$, and $D3'$.

The topology and coordinate files were built after hydrogenation of the samples using the tLEaP module in AMBER16 ([73\)](#page-14-16). All of the amino acids were protonated in aqueous solution (TIP3P water), and the ff14SB forcefield parameters were used to model the amino acids (74) . The total charge of the WT protein was -22 , the total charge of the variants containing negatively charged amino acid (Asp and Glu) mutations was -24 , and the total charge of the variants containing Arg mutations was -20 . To neutralize the total negative charge of the samples, potassium counterions were added. The waters in the crystallographic structure were deleted, and the system was solvated using TIP3P water molecules. The distance between the solute and the edge of the box was chosen as 12 Å , resulting in an average box dimension of $118.7 \times 84.0 \times 128.2$ Å.

All simulations were performed using the CUDA version of the AMBER 16 pmemd program ([73,75,76](#page-14-16)). Berendsen temperature coupling [\(77\)](#page-14-18) was employed, and a time step of 1.0 fs was used in all simulations. Periodic boundary conditions were imposed on the samples, and the particle mesh Ewald summation technique (78) (78) was used with 8- \AA cutoff distance. The SHAKE algorithm ([78\)](#page-14-19) was used for the bonds involving hydrogen atoms. The systems were equilibrated with an 11-stage process after a 1000-step minimization for each system. First, a 10-ps MD simulation was performed at 10 K using a harmonic restraint of 250 kcal/mol/ A^2 on all residues. Second, the harmonic restraint was reduced to 100 kcal/mol/ $A²$ in a simulation of 10,000 steps at 10 K. In the third and fourth stages, the harmonic restraints were reduced by half in simulations, each of which were 20 ps at the same temperature. Fifth, the temperature was gradually increased to 300 K in a 10-ps simulation using a harmonic restraint of 500 kcal/mol/ A^2 . Sixth, a 40-ps simulation was performed at 300 K using the same harmonic restraint for all systems. Seventh, the restraints were reduced to 250 kcal/mol/ \AA^2 in a 20,000-step simulation, which was followed by eighth, another 20,000 steps of simulation with a harmonic restraint of 100 kcal/mol/ $A²$ for each system. The ninth and 10th stages were performed with the same harmonic constraints as in the third and the fourth stages, respectively, but at 300 K with 20-ps-long simulations for both stages. Finally, 11th, a 20-ps simulation was performed without harmonic restraints at 300 K. Production simulations were performed in the NVT ensemble at 300 K. Seven different MD trajectories were simulated, each of which was 100 ns.

Trajectory analysis and visualizations

Root mean-square deviation (RMSD), root mean-square fluctuations (RMSFs) of the residues, dynamic cross-correlation analysis, and hydrogen-bond analysis of the trajectories were performed using the cpptraj module [\(79](#page-14-20)). All three-dimensional representations were obtained using VMD [\(80](#page-14-21)), and the two-dimensional representations were prepared using MarvinSketch [\(http://www.chemaxon.com\)](http://www.chemaxon.com) [\(81](#page-14-22)).

RESULTS

Motivations for this study and design of the β clamp variants

The ability of clamp proteins to open and close is essential to their function as protein hubs for recruitment of interacting protein partners that regulate traffic on DNA during replication and repair [\(2](#page-12-5)). The dimer interface of the E. coli β -clamp is stabilized by four antiparallel β -strands and two α -helices, yet there is evidence that the β -clamp undergoes opening events and populates conformational states in which Domain 1 at the dimer interface in particular is dynamic ([55\)](#page-14-0). Several hydrophobic residues line the dimer interface, and previous attempts to destabilize the interface to a considerable extent have proven unsuccessful, except for the double mutation I272A L273A, which is a monomer at low micromolar concentrations ([55,82\)](#page-14-0). The β -clamp point-mutation variants F104W, F106A, F106W, and L273A are dimers in solution ([82–84](#page-14-23)). Newer work suggests that electrostatic interactions play a key role in stabilizing the interface $(52,54)$ $(52,54)$ $(52,54)$. Our goal in this work was to design and characterize mutations at the dimer interface to test the hypothesis that the clamp transiently populates open conformational states captured by the high-affinity binding by the clamp loader in the process of clamp loading.

Our approach included constructing a broad range of mutations altering the stability of the dimer interface ([Fig. 1](#page-1-0)). Starting with the crystal structure of the clamp ([4,8](#page-12-1)) as a guide, we chose a set of amino acid substitutions with the goal of either stabilizing or destabilizing its dimeric structure. We avoided making mutations at L273 because this residue lies on the side of the β -clamp involved in binding DNA polymerases and the clamp loader and could interfere with these interactions. Furthermore, we designed mutations away from residues E93 and

L98 because they are important for interactions with DinB and UmuC ([26,85–87\)](#page-13-17).

To weaken the dimer interface, we introduced charge at residue L82, specifically L82D and L82E, to remove existing hydrophobic interactions and increase the repulsion at the dimer interface. Additionally, we combined the L82E mutation with an I272A mutation that replaces a larger hydrophobic residue with a smaller one. We hypothesized that mutating S107 and S109 to charged arginine residues would cause the formation of a salt bridge across the dimer interface and thereby stabilize the β -sheet at the dimer interface. Furthermore, we wanted to stabilize Domain 1 of the clamp to reduce the conformational flexibility we previously observed in this region. In the hydrogen-deuterium exchange mass spectrometry (HXMS) experiments, we observed increased deuterium uptake, consistent with β transiently populating open states in which Domain 1 is more flexible and prone to unfolding. Indeed, the peptide including residues 1–34 exhibited EX1 kinetics, indicating local unfolding in that region of the clamp [\(55,56\)](#page-14-0). Thus, we made mutations near this region, introducing charged Arg at positions T45 and T47 to increase ionic interactions with E52 and stabilize β -strand 4 and α -helix 1. Although these mutations likely stabilize Domain 1 in the context of both the closed state and the open states with disrupted dimer interface, we expect they also increase stability of the β dimer by reducing the entropic penalty associated with the loss of Domain 1 flexibility upon formation of the closing ring.

Thermostability gives insight into the effect of dimer interface mutation

To assess whether we had altered the thermostability of the resultant variant clamps, we used a thermofluor assay, in which WT and variant β -clamp proteins incubated with a fluorescent dye were subjected to increasing temperatures. Initially, no fluorescent signal is observed because the fluorescence of the dye is quenched by water, but as the protein denatures, more of its hydrophobic regions are exposed and more dye binds until maximal fluorescent signal is reached ([Fig. 2](#page-5-0) A). From this, we can calculate the melting temperature of the protein, which is defined as the temperature at which 50% of the protein is unfolded. A stabilizing mutation is expected to increase the melting temperature of the protein, and a destabilizing mutation is expected to decrease the melting temperature. Indeed, all mutations that were designed to be stabilizing, namely T45R, T47R, S107R, and S109R, have melting temperatures similar to or a few degrees higher than WT β (69.0°C) ([Fig. 2](#page-5-0) B). Compared to WT β , β L82D and β L82E I272A have significantly lower melting temperatures of 49.5 and 45.3 $^{\circ}$ C, respectively, whereas the L82E mutation more modestly decreases the melting temperature to 62.3° C. Overall, following the above strategy, we have successfully designed a set of

FIGURE 2 (A) Differential scanning fluorimetry of WT and variant β -clamps. Melting temperatures were observed by monitoring Sypro Orange dye fluorescence. (B) Bar graph showing average melting temperatures and standard deviations for WT β (light blue), β T45R (red), β T47R (green), β S107R (purple), β S109R (cyan), β L82D (orange), β L82E (dark blue), and β L82E I272A (maroon) from triplicate experiments.

both stabilizing and destabilizing β -clamp mutations that we subsequently use to correlate the β -dimer stability with opening dynamics, clamp-loader binding, and loading efficiency.

Native gel electrophoresis and native nESI-TIMS-MS shows altered dimerization state of variants

A representative denaturing gel [\(Fig. 3](#page-5-0) A) shows the purity of WT and variant β -clamps and monomeric conformations on SDS-PAGE. The dimerization state of WT and variant β -clamps was evaluated by native gel electrophoresis ([Fig. 3](#page-5-0) B). WT β and the T45R, T47R, S107R, and S109R variants have a pronounced dimer band, whereas mutations at position L82 only or when in combination with I272A result in a less prominent dimer band. Indeed, the L82D variant shows a very weak dimer band and instead shows mainly monomeric and multimeric conformations, which are presumably monomers forming multimeric structures.

FIGURE 3 Native gel electrophoresis to assess dimerization state of variants. (A) Denaturing PAGE showing purified WT and variant clamp proteins as monomers in the presence of SDS. (B) Native PAGE shows monomeric, dimeric, and multimeric conformations of WT and variant clamp proteins at 1μ g each.

Mass spectra of the WT β -clamp and the L82E variant displayed a major dimer distribution, ranging from $[M + 16H]^{16+}$ to $[M + 20H]^{20+}$ [\(Fig. 4](#page-6-0), A and B; [Table](#page-12-3) [S1](#page-12-3)), and a minor tetramer distribution, ranging from $[M + 25H]^{25+}$ to $[M + 28H]^{28+}$ and $[M + 24H]^{24+}$ to $[M + 28H]^{28+}$, respectively [\(Fig. 4;](#page-6-0) [Table S1](#page-12-3)), when using native nESI. In the case of the L82E I272A variant, the mass spectrum displayed an equivalent monomer and dimer distribution, ranging from $[M + 10H]^{10+}$ to $[M + 12H]^{12+}$ and $[M + 16H]^{16+}$ to $[M + 19H]^{19+}$, respectively ([Fig. 4](#page-6-0) C ; [Table S1](#page-12-3)). However, only a monomer distribution, ranging from $[M + 10H]^{10+}$ to $[M + 12H]^{12+}$, was obtained in the MS spectrum of the L82D variant [\(Fig. 4](#page-6-0) D ; [Table S1\)](#page-12-3). These MS data were thus found consistent with our other observations indicating that the L82E I272A and L82D variants destabilize the clamp structure, whereas the glutamic acid mutation at residue L82 (L82E) does not disrupt the clamp as much. Typical TIMS spectra for the multiply protonated species of the *n*-mers of the clamp variants are shown [\(Fig. 4\)](#page-6-0), and the measured CCSs are listed in [Table S1](#page-12-3). Inspection of the TIMS spectra for dimers and tetramers of the WT clamp exhibited higher CCS values as compared to the dimers and tetramers of the L82E variant for the same charge states. This means that the additional negative charge provided by the L82E mutation leads to a more compact conformation than the WT clamp, probably by making additional hydrogen bonds around the mutation site (see below, [MD Simulations\)](#page-3-0). In addition, the presence of some dimers for the L82E I272A variant at similar CCS values, as compared to the L82E variant dimers, suggests that the hydrogen-bond network is only partially perturbed by substituting a larger hydrophobic residue with a smaller one. Furthermore, the observation of only monomers for the L82D variant at similar CCS values, as compared to some monomers observed for the L82E I272A variant,

FIGURE 4 MS and TIMS spectra for the multiply protonated species of the β -clamp (A) WT, (B) L82E, (C) L82E I272A, and (D) L82D variants. (E) The distribution of monomers/dimers/tetramers [\(Table S1](#page-12-3)). To see this figure in color, go online.

suggested that the two monomer variant structures adopt similar conformations despite the different mutations. The native TIMS-MS experiments enabled us to conclude that the mutation most responsible for the disruption of the dimers is provided by the negatively charged L82D, in which the hydrogen-bond network is strongly perturbed around the mutation site as compared to the L82E and L82E I272A variants in the gas phase. Our observations from native gel electrophoresis and TIMS-MS indicate that WT β , the stabilizing T45R, T47R, S107R, and S109R variants, and the moderately destabilizing β L82E variant remain dimeric at micromolar concentrations, whereas β L82E I272A and β L82D are at least partially monomeric.

Designed β variants bind mini- δ with affinity similar to that of WT β

To probe the effect of altered stability of the β -clamp variants on their affinity for the clamp-loader complex, we characterized interaction of mini- δ with WT β ; with the stabilizing variants β T45R, β T47R, β S107R, and β S109R; and with the destabilizing variants β I272A L273A, β L82E, and β L82E I272A using ITC. Note that we were unable to express and purify the least stable variant, β L82D, in the amount necessary for ITC binding studies with mini- δ . The raw ITC data and the fits for the β -clamp variants are shown in [Fig. S2](#page-12-3). The binding parameters for mini- δ with all considered β -clamp variants, including the equilibrium dissociation constant K_D , association enthalpy ΔH , binding stoichiometry parameter *n*, and association entropy ΔS , are listed in [Table 1.](#page-6-1)

Previous studies demonstrated that the equilibrium dissociation constant K_D for δ binding to WT β (determined by surface plasmon resonance) is one to two orders of magnitude weaker than the K_D for δ binding to β I272A L273A [\(82](#page-14-23));

TABLE 1 Binding Parameters for Mini- δ and β -Clamp Variants Obtained by ITC

β -clamp variant	$K_{\rm D}$ (μ M)	\boldsymbol{n}	ΔH (kJ/mol) (J/mol/K) T (°C)	ΔS	
WТ	5.6 ± 0.69	$1.01 \pm$ 2.0×10^{-4}	$-21.6 + 0.4$	27.7	25
I272A	$0.53 + 0.08$	$0.71 + 0.01$	$-34.6 + 0.4$	2.0	20
L273A					
L82E	$6.6 + 1.8$	$0.83 + 0.03$	$-17.8 + 1.1$	38.6	20
L82E	$5.5 + 1.0$	$0.41 + 0.01$	$-34.0 + 2.0$	-15.3	20
I272A					
S107R	4.4 ± 0.9	0.96 ± 0.02	-20.4 ± 0.8	34.2	25
S109R	$5.0 + 1.1$		$1.06 + 0.03 -19.4 + 0.9$	36.5	25
T45R	$6.7 + 2.0$		0.95 ± 0.03 -22.5 ± 1.5	23.4	25
T47R	$6.0 + 1.7$	$0.95 + 0.03$	-22.5 ± 1.4	24.3	25

Mini- δ (0.6–0.7 mM) was titrated into 0.1 mM β -clamp solution. The parameters include the dissociation constant K_D , association enthalpy ΔH , binding stoichiometry parameter *n*, and association entropy ΔS obtained at 25°C (20°C for L82E, L82E I272A, and I272A/L273A β variants).

therefore, this variant was included here as a control. We were able to confirm this result by ITC [\(Fig. S2](#page-12-3); [Table 1](#page-6-1)): titration of mini- δ into a solution of WT β resulted in a K_D of \sim 6 μ M, whereas K_D for mini- δ binding to β I272A L273A was 0.5 μ M, over one order of magnitude stronger. Furthermore, the association enthalpy for β I272A L273A was \sim 1.6-fold higher than for WT β (ΔH of -35 kJ/mol vs. -22 kJ/mol for WT β). It is remarkable that mini- δ binding to WT β also resulted in positive (favorable) change in entropy ($\Delta S \sim$ 28 J/mol/K), presumably due to dehydration of the binding interface accompanied by an increase in solvent entropy. On the other hand, the association entropy for mini- δ binding to β I272A L273A was much smaller than for WT β $(\Delta S \sim 2.0 \text{ J/mol/K})$. This is consistent with higher flexibility of the I272A L273A β mutant, presumably due to increased sampling of a more dynamic open conformation, which is reduced upon mini- δ binding.

On the other hand, ITC analysis for mini- δ binding to stabilized β variants T45R, T47R, S107R, and S109R resulted in K_D values of 4–7 μ M and ΔH about -20 kJ/mol, similar to those obtained for mini- δ binding to WT β . However, the association entropy for two of the stabilized variants (S107R, S109R; $\Delta S \sim 34-37$ J/mol/K) was slightly higher than for WT β ($\Delta S \sim 28$ J/mol/K), potentially pointing to a smaller entropic penalty associated with a decrease in β -clamp flexibility upon mini- δ binding.

Interestingly, the K_D (6.6 μ M) and association enthalpy and entropy ($\Delta H \sim -18$ kJ/mol and $\Delta S \sim 39$ J/mol/K, respectively) obtained for the destabilized L82E variant are similar to those for β S107R and β S109R, likely suggesting that at the concentrations used in our ITC measurements (\sim 100 μ M), β L82E predominantly resides in the closed state. However, β L82E I272A, although displaying only a marginally stronger K_D (5.5 μ M), shows similar trends as β I272A L273A, i.e., higher association enthalpy than for WT β (ΔH of -34 kJ/mol vs. -22 kJ/mol) and a much smaller association entropy $(\Delta S \text{ of } -15.3 \text{ J/mol/K} \text{ vs.}$ 28 J/mol/K). These ITC data suggest that stabilizing (T45R, T47R, S107R, S109R) and moderately destabilizing (L82E) β variants reside in the closed state at our experimental conditions (\sim 100 μ M β) and exhibit K_D and ΔH for mini- δ binding similar to those observed for WT β . On the other hand, more destabilizing β L82E I272A and β I272A L273A variants increasingly populate open-like conformation(s) and exhibit larger (favorable) association enthalpy ΔH and smaller (less favorable) association entropy ΔS , which is consistent with the formation of more stabilizing intermolecular contacts and greater flexibility loss for open β -clamp upon mini- δ binding.

Lower ATPase activity of the γ -complex is observed in the presence of β L82D and DNA

We next assessed whether the variant β -clamps stimulate the ATPase activity of the γ -complex in the presence of primed DNA, using a colorimetric inorganic phosphate quantitation assay. We incubated each of the reaction components alone or in combination and measured the absorbance to assess color formation; the increase in inorganic phosphate in solution is brought about by the ATPase activity of the γ -complex ([Fig. 5](#page-8-0) A). No significant color change is observed when ATP, β , γ -complex, or DNA alone are incubated in assay buffer or when ATP and γ -complex or ATP, β , and γ -complex are incubated in the absence of DNA. ATPase activity is detected when primer-template DNA is added to ATP, β , and the γ -complex. The β L82D variant does not enhance the ATPase activity of the γ -complex, β S109R shows a modest decrease in conferring enhancement of the ATPase activity, and other variants result in similar levels of ATPase activity compared to the level induced by the presence of WT β -clamp ([Fig. 5](#page-8-0) B). ATP hydrolysis occurs upon release of β on DNA by the γ -complex, so reduced ATPase activity could be due to defects in clamp binding to the loader, clamp loading, or in release of the β -clamp on DNA.

Stabilized variants are efficiently loaded onto primer-template DNA

To evaluate clamp loading of the β variants, we adapted an assay (88) (88) utilizing a 5['] biotinylated primer-template substrate with a $5'$ single-stranded overhang for loading of the clamp and streptavidin-coated magnetic beads that allow for separation of bound and unbound protein. To prevent β from sliding off the linear DNA, SSB is added to the loading reactions, which traps β on single-stranded DNA (ssDNA; [Fig. S3](#page-12-3)). After the loading reactions are washed with buffer, bead resuspensions are applied to the magnet, the pellet (P) fractions indicate loaded β , and the supernatant (S) fractions indicate unloaded β , shown at 0 and 60 min.

Representative anti- β immunoblots show the loading efficiencies of the WT and variant β -clamps ([Figs. 5](#page-8-0) C and [S3\)](#page-12-3). Overall, the amount of loaded β and the amount still present after 60 min is greater in the variant clamps with intended stabilizing mutations T45R, T47R, S107R, and S109R, as indicated by the greater amount of β in the pellet at 0 and 60 min ([Fig. 5](#page-8-0) C). Less of the L82D variant is loaded on DNA than WT β , whereas the β L82E and L82E I272A variants appear to remain loaded on DNA, as indicated by the absence of β in the supernatant at 0 and 60 min and the presence of β in the pellet at 60 min. These observations could indicate altered interactions with the clamp loader and, in the case of β L82D, are consistent with a less stable β dimer. The ITC experiments [\(Fig. S2;](#page-12-3) [Table 1](#page-6-1)), which indicate similar equilibrium dissociation constants for these clamps as for WT β binding to mini- δ , were carried out with only a truncation of the δ subunit of the clamp loader, whereas the ATPase and loading assays are carried out

FIGURE 5 ATPase activity of the γ -complex measured using a BIOMOL Green colorimetric reagent. (A) A schematic depicting loading of the β -clamp onto a primer-template DNA with biotinylated 5'-ends. The template is longer than the primer, resulting in a $3'$ recessed end onto which the γ -complex loads β in the presence of ATP. ATP hydrolysis ejects the clamp loader. The increase of inorganic phosphate following the loading reaction is measured by addition of the BIOMOL Green reagent. (B) A bar graph showing the amount of phosphate released (nmol) during the loading reaction after the intrinsic γ -complex ATPase activity is subtracted (γ -complex hydrolyzes ATP in the presence of DNA, β enhances the ATPase activity of the γ -complex). Average nmol of phosphate released and standard deviations from at least three replicates are reported. (C) Immunoblot detection of β loaded onto a primer-template substrate using a bead-based assay for WT β and the variants indicated. Input control (L) indicates the amount of β in each loading reaction. After three wash steps (W1, W2, and W3), pellet (P) fractions indicate loaded β and supernatant (S) fractions indicate unloaded β , respectively, for both 0- and 60-min time points (also see [Fig. S3\)](#page-12-3)

with the full five-subunit clamp loader. We did not observe in our simulations that the destabilizing β variants have more extensive contacts with DNA than WT β ; however, altered or additional hydrogen-bonding interactions at the dimer interface could explain the persistence of the destabilizing β variants on DNA ([Fig S3](#page-12-3); see below, [MD](#page-3-0) [Simulations](#page-3-0)).

β L82D and β L82E I272A variants do not support cell growth in complementation assay

To determine whether the variant β -clamps support cell growth, we utilized an E. coli MS120 strain [\(70](#page-14-13)) bearing mutations conferring temperature sensitivity in the chromosomal *dnaN* gene. The MS120 strain grows uninhibited at 30° C but is unable to grow at temperatures of 37° C or higher unless complemented with a functional β gene. We transformed these cells with either empty vector or plasmids expressing WT or variant β -clamps, grew them at 30 and 37°C, and determined the number of colony-forming units at each temperature ([Fig. 6](#page-8-1)). All transformants grew at 30° C, but only those that produced functional clamps were able to support growth at 37 $^{\circ}$ C. The β L82D and β L82E I272A variants did not support cell growth in the complementation assay. Interestingly, the β L82E variant was able to complement growth, but the resultant colonies were much smaller, suggesting these clamps were compromised, resulting in a growth defect. Thus, all stabilizing clamp variants confer growth on the temperature-sensitive strain, whereas the destabilizing variants β L82D, β L82E I272A, and β L82E are unable or less able to rescue growth at the nonpermissive temperature. We analyzed protein levels at the nonpermissive temperature by immunoblotting, which shows that the variants were expressed at detectable levels, except the L82E I272A variant, which does not support growth at 37° C and therefore was not analyzed ([Fig. S5](#page-12-3)).

MD simulations reveal altered conformations and dynamics of β -clamp variants

MD simulations with WT β -clamp and the seven variants were performed to evaluate the effect of the mutations on the dynamics of the protein. The stability of the trajectories has been investigated with RMSD analysis of the backbone atoms (C, C_{α} , and N) of β on Monomers A and B separately. In all cases, the average RMSD for both the full-length protein and backbone atoms on Monomers A and B was at or below 3 Å [\(Figs. S6](#page-12-3) and [S7](#page-12-3)).

FIGURE 6 Ratio of colony-forming units at 37° C vs. 30° C for WT and variant β -clamps in complementation assay. EV indicates the empty vector. Data represent the average and error bars represent the standard deviation of at least three replicates. To see this figure in color, go online.

Changes in RMSF relative to WT β were determined to assess the effect of the point mutations on the fluctuations for the different variants. Nine residues with a change in relative RMSF value greater than ± 1 Å were observed for the different variants (Fig. $S11$). P20', L21', and G22' on Monomer B (indicated by the prime symbol) become more mobile in the L82E variant [\(Fig. S11](#page-12-3), green) relative to $WT \beta$. D120', which is located on the interdomain connector loop between D1 and D2, becomes more mobile in the S107R variant [\(Fig. S11](#page-12-3), cyan), whereas the mobility of $D120'$ is observed to be either very similar to that of the WT protein or calculated to be less than that of WT β for all other variants. The interdomain connector loop has been previously reported to have regions that are rapidly deuterated in HXMS experiments with different clamp proteins ([56\)](#page-14-8).

The effect of different point mutations on the correlations between the residues located at the dimer interface was studied by calculating the deviation of the cross correlations with respect to WT β ([Fig. S12\)](#page-12-3). Helix-3 (H3) and helix-21 (H21) are observed to be mostly positively correlated in the L82E, T45R, and S107R variants ([Fig. S12](#page-12-3) A, region colors: blue-cyan), whereas negative correlations are found for the remaining systems relative to WT β .

For the Domain 3-Domain $1'$ interface, helix-10 (H10) and helix-14 (H14) are observed to be correlated in the T45R and S107R variants [\(Fig. S12](#page-12-3) B, region colors: blue-cyan). The L82E and L82E I272A variants increase the number of residues in β sheets that are correlated with each other on the domain interface between Domain 1 and Domain 3'. The region composed of residues from L82 to R96, in particular, is found to be highly correlated with the region composed of residues Y282' to N295' in these variants. It is noteworthy that the magnitude of the correlations between these β -sheets is the highest in the L82E I272A variant relative to the other systems (see [Supporting Materials and Methods](#page-12-3) for detailed results).

The two-dimensional representation of the hydrogen bonds that are observed within the dimer interface based on a hydrogen-bond analysis of the simulations is given in [Fig. S15,](#page-12-3) in which each hydrogen-bond interaction is depicted by a different color. Several changes in the interdomain contacts are observed, depending on the individual mutation. The L82D and L82E variants introduce negatively charged residues into the protein, which alter the hydrogen-bond network around the mutation site. The total fraction of frames that had hydrogen bonds around the mutation site is depicted in [Fig. 7](#page-10-0). It is observed that the hydrogen bond between the backbone oxygen atom of G81 and the side chain of Q265 of the neighboring monomer is conserved in both interfaces in a higher percentage of the sampled frames of β L82D, whereas the Domain 1-Domain $3'$ interface exhibits the interaction for a higher percentage than on the Domain 3 -Domain $1'$ interface of the β L82E variant ([Fig. 7](#page-10-0) A) relative to WT β . This interaction is conserved with a higher percentage in at least one dimer interface of β L82E I272A variant relative to WT protein [\(Fig. 7](#page-10-0) A). There is almost no alteration in these interactions with other mutations. In the WT protein, the backbone oxygen atom of $G81$ and $G81'$ forms a hydrogen bond with the side chains of $R269'$ and $R269$, respectively ([Fig. 7](#page-10-0) B). However, the L82D variant results in a smaller total fraction of frames with this hydrogen bond present. On the other hand, the L82E variant increases the time this interaction is observed, especially on the Domain 3-Domain 1' interface.

A similar pattern is seen in the β L82E I272A variant but with longer conservation time on the $D1-D3'$ interface compared with the Domain 3 -Domain $1'$ interface. These results indicate that the mutation of L82 to a negatively charged residue, especially Glu, increases the total fraction of time that the backbone oxygen atoms of $G81$ and $G81'$ can receive a hydrogen bond from a neighboring residue. The mutations in the T45R and S107R variants also increase the number of frames in which $G81$ and $G81'$ have hydrogen bonds with R269' and R269, respectively, relative to WT β ([Figs. 7](#page-10-0) B and [S15](#page-12-3)).

The most dramatic alteration observed is for the L82D, L82E, and L82E I272A variants, for which a new contact is formed only in these systems and not any other. In these systems, the carboxyl moiety on the side chains (D or E) form hydrogen bonds with the positively charged side chain of R269 and R269' ([Fig. 7](#page-10-0), C and D). This interaction is maintained for at least 60% of the simulation time in the β L82D variant. In the β L82E variant, the two dimer interfaces appear to be asymmetric with respect to hydrogen bonding between residues 82 and 269. The introduction of the additional mutation, I272A, reduces the distinctive feature between the two dimer interfaces of β L82E I272A, in which the interactions are present at least 60% of the simulation time at both dimer interfaces. These results are consistent with the fact that this variant remains loaded on DNA. Similar differential effects at other locations in β are observed for other sites, including R96 and R96', S109 and S109', and T45R and T45'R (see [Support](#page-12-3)[ing Materials and Methods](#page-12-3)). As with the cross-correlation analysis, the largest perturbations to hydrogen bonding are found in the destabilizing β L82D, L82E, and L82E I272A variants.

DISCUSSION

Structural and biochemical observations have provided many of the mechanistic details regarding the clamp loading process and reveal variations across species. The crystal structure of E. coli γ -complex in its inactive conformation demonstrates its inability to bind to the β -clamp because of steric clashes [\(40](#page-13-8)). The crystal structure of the γ -complex with all its ATPase sites occupied bound to DNA reveals its

FIGURE 7 The total fraction of the frames that a hydrogen bond exists between (A) the backbone oxygen atom of G81 and side chain Q265' (solid $brown$) and the backbone oxygen atom of $G81'$ and side chain $Q265$ (*pattern brown*), (*B*) the backbone oxygen atom of G81 and side chain $R269'$ (solid blue) and the backbone oxygen atom of G81' and side chain R269 (pattern blue), and (C) the side chain of $D82/E82$ and side chain R269^o (solid cyan) and the side chain of $D/EB2'$ and side chain R269 (pattern cyan). The backbone oxygen atoms are depicted with subscript "O," while the side chain is depicted with subscript "SC." (D) The three-dimensional representation of the depicted interactions obtained from the trajectory at 85 ns of variant L82E I272A.

active conformation and the mechanism of ATP hydrolysis ([41\)](#page-13-9). However, the conformation of the clamp loader when it binds the clamp is unknown because the crystal structure of the clamp-clamp loader complex has not been solved, possibly because such an intermediate may be short-lived, as the clamp may sample open and closed states in solution.

Clamp-loader-facilitated opening of the clamp can be an active process in which the clamp loader binds a closed clamp and opens it or the clamp loader can bind to and stabilize a transiently open clamp ([42–45\)](#page-13-10). The T4 bacteriophage clamp gp45 is frequently open in solution and therefore does not need a dedicated clamp loader for opening; instead, the gp44/62 stabilizes the open conformation [\(42–46](#page-13-10)). The crystal structure of the T4 gp45 clamp bound to gp44/62 clamp loader and DNA ([89\)](#page-14-25) reveals an open clamp-clamp loader assembly, whereas in the structure of the yeast PCNA-RFC complex, the clamp is closed ([90\)](#page-14-26). The yeast closed-clamp structure may be an artifact because an arginine finger variant of the clamp loader was used that renders the loader unable to form a helical conformation and bind the clamp.

The β -clamp has a $t_{1/2}$ of 60 min on DNA ([42\)](#page-13-10). In E. coli, the δ subunit moderates the affinity of the complex for the clamp [\(91](#page-14-27)), and the δ subunit alone has been shown to unload the clamp ([59\)](#page-14-3). The δ subunit has been demonstrated to have a higher affinity for the β monomer rather than the dimer, suggesting that δ prefers the open conformation of β [\(57](#page-14-1)). The structure of mini- δ bound to a β variant that is not a stable dimer shows a relaxed curvature that results in a \sim 15-A opening at one dimer interface, a distortion wide enough to accommodate ssDNA ([57\)](#page-14-1). The δ subunit does not destabilize β at the dimer interface but rather binds to the hydrophobic pocket where most β client interactions occur [\(57](#page-14-1)). The helix containing residues I272 and L273 in β Domain 3 is under strain and allows for an intermonomer hydrophobic interaction with Domain 1, forming a tight dimer interface. In the crystal structure of the monomeric β -clamp bound to the δ subunit, these residues are mutated to alanine, and these smaller hydrophobic residues prevent dimer formation in the low micromolar range ([57\)](#page-14-1). Furthermore, the five-residue loop immediately following those residues provides an interaction surface for monomeric β and δ , whereas the loop is occluded in dimeric β . The monomer shows more dynamic behavior than WT β in the HXMS experiments [\(55](#page-14-0)). This greater flexibility is supported by steered MD simulations as the monomer opens up more relative to the dimer ([58\)](#page-14-2), and this region is perturbed most in the β L83E variant [\(Fig. S11\)](#page-12-3). A 7° interdomain angle is observed in the simulated monomeric clamp ([58\)](#page-14-2), which is similar to the angle observed in the crystal structure of the β monomer complexed to the δ subunit ([57\)](#page-14-1).

We previously observed that residues 1–34 in Domain 1 undergo a local cooperative unfolding as they are deuterated faster and display EX1 kinetics in HXMS experiments ([55,56](#page-14-0)). The solvent accessibility of these residues signifies the greater flexibility in this region and suggests that β transiently opens in solution. The crystallographic data, along with HXMS experiments, suggest that the γ -complex traps β in an open conformation as the strain held at the dimer interface is alleviated and the δ subunit serves to prevent ring closure. The δ subunit does not open β at the dimer interface but rather stabilizes one monomer, preventing ring closure. Here, we further investigate the dynamic behavior at the dimer interface and its effect on the opening and closing of the ring structure. Using the crystal structure of WT β as a guide, we designed mutations with the intention of stabilizing or destabilizing the dimer interface.

The mutations we hypothesized to be stabilizing, T45R, T47R, S107R, and S109R, had similar or increased thermostability compared to WT β and similar functions as WT β except for having greater residence on DNA after loading ([Fig. 8](#page-11-0)). The atomic fluctuations that the mutations produced in the MD simulations were examined by performing RMSF analysis. The highest increase in the mobility of residues G209, G210, and D211 was observed in the T45R variant. The same residues on the opposite monomer were less mobile in all other variants except for the T47R variant. The hydrogen-bond analysis of the MD simulations determined that residues R45 and R47 within T45R and T47R variants formed new interactions with E52 relative to WT β ([Fig. S17](#page-12-3)). Additionally, in the S107R variant, R107 formed interactions with E93 [\(Fig. S20](#page-12-3) E). Lastly, R96 was observed to interact with Q299 and S107 in the S109R variant ([Fig. S20\)](#page-12-3).

The mutations designed to be destabilizing disrupted the dimer interface to varying degrees ([Fig. 8](#page-11-0)). The L82D variant produced the greatest destabilization effect by far, and in comparison, L82E had milder effects. When the L82E mutation was combined with I272A, the effect closely resembled the L82D variant. The β L82D and L82E I272A variants had significantly reduced thermostability, whereas β L82E moderately reduced the melting temperature of the protein by $\sim 7^{\circ}$ C ([Fig. 8\)](#page-11-0). The dimerization state of all

FIGURE 8 Summary of results presented, showing the different effects of each of the mutations made.

three variants was altered, with β L82D detected as a monomer at concentrations in the μ M range, β L82E I272A as monomer and dimer, and β L82E as dimer and multimer by native PAGE and native TIMS-MS. The β L82D and β L82E I272A variants did not complement cell growth in the temperature sensitivity assay. The β L82E variant supported cell growth but resulted in smaller colonies that grew to a normal size with longer incubation time. The intrinsic ATPase activity of the clamp loader was not stimulated by the addition of β L82D, whereas β L82E and β L82E I272A stimulated the ATPase activity to nearly the same extent as the WT β -clamp. The destabilized β L82E and β L82E I272A variants did not appear to unload from the linear, end-blocked DNA substrate. One possibility that would explain this observation is that these variants might remain bound to the γ -complex, as the δ subunit has been shown to have greater affinity for an open clamp. The L82E and L82E I272A variants had similar affinity for mini- δ as WT β , but those experiments were carried out only with mini- δ , whereas the clamp-loading experiments were performed with the complete clamp-loader complex and DNA. The β L82D variant did not stimulate the ATPase activity of the γ -clamp loader and was loaded onto DNA less efficiently than WT β . The RMSF of L248, which is in close proximity to the β client proteinbinding pocket ([19,57,92\)](#page-13-18), was observed to increase in one of the monomers in the L82E I272A variant. $P20'$, $L21'$, and $G22'$ on the second monomer were observed to be more mobile in the L82E variant. These residues are located on the complementary region consisting of residues 1–34, which was previously reported to undergo a local cooperative unfolding and to display EX1 kinetics in the HXMS experiments [\(55,56\)](#page-14-0). Difference cross-correlation analysis for these destabilizing variants suggests that at least one of the two dimer interfaces (both interfaces in L82D) exhibits higher anticorrelated movement in the interface region compared to WT β , as well as higher changes in correlation overall. The hydrogen-bond analysis revealed that changing the leucine at the 82 position to either an aspartic acid or glutamic acid resulted in new interactions between D/E82 and R269 and between G81 and Q265. These new hydrogen-bond interactions also reduce a contact between the G81 backbone and R269 side chain in WT β . Taken together, the MD simulations suggest that the changes in specific hydrogen-bond contacts and other noncovalent interactions induced by the point mutations result in significant changes in conformation and dynamics that are consistent with the observed experimental results for both the stabilizing and destabilizing mutations.

Our rational approach resulted in variants with stabilizing and tunable destabilizing mutations that likely render the clamp closed or open, respectively. We also report here MD simulations that rationalize our observations of the effects of these mutations on the biological activities of the resulting clamp variants, as well as the complete hydrogen-bond-interaction map. We are now working to create stable monomers in solution by designing the next generation of mutations.

SUPPORTING MATERIAL

Supporting Material can be found online at [https://doi.org/10.1016/j.bpj.](https://doi.org/10.1016/j.bpj.2019.06.035) [2019.06.035](https://doi.org/10.1016/j.bpj.2019.06.035).

AUTHOR CONTRIBUTIONS

B.N.K., H.G., A.A.R., F.F.-L., D.M.K., G.A.C., and P.J.B. designed research. B.N.K., H.G., A.A.R., S.L., K.J.D.F., A.C., and M.L.L. performed research and analyzed data. B.N.K., H.G., A.A.R., S.L., K.J.D.F., M.L.L., F.F.-L., D.M.K., G.A.C., and P.J.B. wrote the manuscript.

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