The ribosome uses cooperative conformational changes to maximize and regulate the efficiency of translation

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One of the most challenging unanswered questions regarding the structural biology of biomolecular machines such as the two-subunit ribosome is whether and how these machines coordinate seemingly independent and random conformational fluctuations to maximize and regulate their functional efficiencies. To address this question, we have used ribosome mutagenesis or a ribosome-targeting antibiotic to predictably perturb the dynamics of intersubunit rotation, a structural rearrangement of the ribosome that is essential for the translocation and ejection of ribosomal-bound tRNAs during translation. Concomitantly, we have used single-molecule fluorescence resonance energy transfer (smFRET) to characterize the effects of these perturbations on the dynamics of ribosomal L1 stalk movements and ribosome-bound tRNA reconfigurations, conformational changes that are likewise essential for the translocation and ejection of tRNAs during translation. Together with the results of complementary biochemical studies, our smFRET studies demonstrate that the ribosome uses cooperative conformational changes to maximize and regulate the efficiency with which it translocates and ejects tRNAs during translation. We propose that the ribosome employs cooperative conformational changes to efficiently populate global conformational states that are productive for translation, that translation factors exploit this cooperativity as part of their mechanisms of action, and that antibiotics exploit it to maximize the potency with which they inhibit translation. It is likely that similar cooperative conformational changes underlie the function and regulation of other biomolecular machines.

During the catalytic cycle of many enzymes, multiple, spatially distant enzyme structural elements must often undergo functionally important conformational changes (1). Consistent with the view that such structural rearrangements must be rapidly organized and executed to maintain catalytic efficiency, many recent studies strongly suggest that small, monomeric protein enzymes have evolved complex networks of cooperative conformational changes that coordinate the inherently stochastic conformational fluctuations of multiple structural elements in a manner that is optimal for catalysis (2). Within the context of energy landscape theory (3), such enzymes can be thought of as having evolved dynamic energy landscapes that bias enzyme conformational sampling in such a manner to maximize the efficiency of catalysis (2). Related proposals suggest that binding of allosteric effectors to enzymes remodels such energy landscapes, altering enzyme conformational sampling as part of the mechanisms through which these effectors regulate enzymatic activity (4).

Unfortunately, the conformational dynamics of large, macromolecular complexes remain much more challenging to characterize than those of small, monomeric proteins (5), making it very difficult to elucidate the role that cooperative conformational changes play in the function and regulation of biomolecular machines such as the ribosome (6). Following each round of aminoacyl–tRNA incorporation and peptide bond formation by the translating ribosome, the resulting ribosomal pretranslocation (PRE) complex must rapidly translocate the newly deacylated tRNA from the ribosomal peptidyl–tRNA binding (P) site to the ribosomal deacylated (or exit) tRNA binding (E) site and the newly formed peptidyl–tRNA from the ribosomal aminoacyl–tRNA binding (A) site to the P site. Concurrent with translocation of the tRNAs, the PRE complex advances along the mRNA by precisely one codon and ultimately ejects the E-site tRNA, producing a ribosomal post-translocation (POST) complex that is ready for the next round of the elongation cycle.

During the first step of translocation, the A- and P-site tRNAs are reconfigured from their classical P/P (denoting the small ribosomal subunit P site/large ribosomal subunit P site) and A/A configurations into their hybrid P/E and A/P configurations. Numerous studies suggest that formation of the hybrid tRNA configuration is accompanied by large-scale conformational changes of the ribosome itself (6–9). Indeed, relative to x-ray crystallographic and cryogenic electron microscopy (cryo-EM) structures of ribosomal complexes carrying classically configured tRNAs, structures of ribosomal complexes carrying hybrid-configured tRNAs exhibit a ratchet-like relative rotation of the ribosomal subunits as well as a closure of the L1 stalk element of the large subunit [defined as 23S ribosomal RNA (rRNA) helices 76, 77, and 78 (H76–78) and ribosomal (r)-protein L1] that allows the L1 stalk to establish a physical interaction with the P/E-configured tRNA (10, 11). Notably, single-molecule fluorescence resonance energy transfer (smFRET) studies have shown that in the absence of elongation factor G (EF-G), the guanosine triphosphatase translation factor that promotes further steps along the translocation pathway, PRE complexes and...
PRE complex analogs lacking an A-site tRNA (PRE-\(^{\text{A}}\) complexes) undergo thermally activated, stochastic, and reversible fluctuations between classical and hybrid tRNA configurations (classical=hybrid) (12), nonrotated and rotated subunit orientations (NR=S=R) (13), and open and closed L1 stalk conformations (L1,=,L1) (14, 15). Unfortunately, however, whether and how these stochastic tRNA, intersubunit, and L1 stalk dynamics are coordinated within PRE/PRE-\(^{\text{A}}\) complexes to facilitate translocation and/or allosteric regulation of translocation remains unknown, severely limiting our understanding of the fundamental physical processes that drive and control the rapid and precise translocation of the tRNAs through the ribosome during protein synthesis.

To determine whether tRNA, intersubunit, and L1 stalk movements are coordinated within PRE/PRE-\(^{\text{A}}\) complexes and to characterize the structural basis for such cooperative conformational changes, here we have used smFRET to characterize the L1 stalk and tRNA dynamics of Escherichia coli PRE-\(^{\text{A}}\) complexes assembled using ribosomes that have been strategically mutagenized or treated with a ribosome-binding inhibitor to predictably perturb intersubunit rotation. Together with the results of complementary translocation studies, our findings directly demonstrate that the ribosome uses cooperative conformational changes to maximize and regulate the efficiency of translocation and tRNA ejection during translation. Using structural and phylogenetic analyses of ribosomes to rationalize our results leads us to propose a structure-based model for the observed cooperativity. Collectively, the work presented here strongly suggests that coordination of spatially remote conformational changes is a fundamental aspect of the mechanism and regulation of translocation, other steps of the translation elongation cycle, other stages of protein synthesis, and other biomolecular machines.

Results and Discussion

Intersubunit Rotation Can Be Predictably Perturbed by Disrupting Electrostatic Interactions at the Subunit Interface. Intersubunit rotation requires extensive remolding of a collection of highly conserved, noncovalent, intersubunit RNA\(\text{A}RNA\text{}\), RNA\(\text{B}RNA\text{}\)-protein, and r-protein\(\text{A}r\text{-protein}\) interactions known as intersubunit bridges (11). Intersubunit rotation and the attendant remodeling of intersubunit bridges have been shown to play critical roles in enabling and regulating tRNA translocation (16, 17). Indeed, deletion of the functionally critical bridge B1h, which underlies the largest rearrangement during intersubunit rotation (18), has been shown to increase the propensity of PRE complexes to undergo EF-G-independent translocation (19) and disruption of single interactions across several intersubunit bridges, including bridge B1b, has been shown to alter the rate of PRE complex translocation and/or the propensity of PRE complexes to slip by one nucleotide and shift their reading frame on the mRNA during translocation (17, 19, 20). Similarly, the tetracyclinomycin family of antibiotic inhibitors of protein synthesis (e.g., viomycin, capreomycin, etc.) acts by binding to bridge B2a and strongly inhibiting translocation, presumably by blocking the remodeling of bridge B2a that accompanies intersubunit rotation (8).

Given its particularly critical and well-studied role in enabling and regulating translocation, we began by developing a series of mutant ribosomes in which bridge B1b was systematically mutagenized to predictably perturb intersubunit rotation. In E. coli, bridge B1b is composed of interactions between r-protein S13 on the small, or 30S, subunit and r-protein L5 on the large, or 50S, subunit. By comparing the structures of ribosomal complexes in the NR- and R-subunit orientations (11), we identified two highly conserved (Table S1), charged amino acids in S13, arginine 3 [(R3)S13] and aspartic acid 82 [(D82)S13], whose electrostatic interactions with oppositely charged residues in L5 are dramatically remodeled upon intersubunit rotation (Fig. 1).

When ribosomal complexes are in their NR-subunit orientation, (R3)S13 is involved in an electrostatic interaction with the highly conserved (Table S1) aspartic acid 143 of L5 [(D143)L5], whereas (D82)S13 is positioned too far away to interact with any residues within the 50S subunit. Upon intersubunit rotation, however, this electrostatic interaction is disrupted such that when ribosomal complexes are in their R-subunit orientations, (R3)S13 is now too far away to interact with any residues within the 50S subunit and (D82)S13 is involved in an electrostatic interaction with the highly conserved (Table S1) arginine 111 of L5 [(R111)L5]. Because (R3)S13–(D143)L5 and (D82)S13–(R111)L5 interactions should primarily alter the stabilities of the NR- or R-subunit orientations and the corresponding rates of intersubunit rotation, respectively, in PRE/PRE-\(^{\text{A}}\) complexes. Guided by these analyses, we set out to perturb intersubunit rotation by constructing and purifying wild-type 30S subunits (WT), 30S subunits lacking S13 [(WTS)S13], and 30S subunits reconstituted with recombinantly expressed and purified wild-type S13 [(WT)S13] or each of five S13 mutants [(R3A)S13, (R3D)S13, (D82A)S13, (D82K)S13, and (D3D/D82K)S13 (SI Materials and Methods)].

All PRE-\(^{\text{A}}\) Complexes Exhibited Dynamic L1,\(\text{=}\)L1, and L1\text{\textasteriskright}tRNA Equilibria. L1 stalk and tRNA dynamics of PRE/PRE-\(^{\text{A}}\) complexes can be monitored using two previously reported smFRET signals. The first signal, denoted smFRET\(_{1.1-1.9}\), derives from Cy3 FRET donor-labeled r-protein L9 [L9(Cy3)] and Cy5 FRET acceptor-labeled r-protein L1 [L1(Cy5)] and reports on movements of the L1 stalk between its L1, and L1, conformations (14). The second signal, denoted smFRET\(_{1.1-2.8}\), derives from Cy3-labeled P-site tRNA [tRNA(Cy3)] and L1(Cy5) and reports on rearrangements of the PRE/PRE-\(^{\text{A}}\) complex between a conformation in which the L1 stalk is in its L1, conformation and is too far away to physically interact with the P/P-configured tRNA (L1,tRNA) and a conformation in which the L1 stalk is in its L1, conformation and is able to physically interact with the P/E-configured tRNA (L1,tRNA) (21). Using each of the 30S subunits described in the previous section, we therefore assembled eight PRE-\(^{\text{A}}\) complexes harboring L1(Cy5)- and L9(Cy3)-labeled 50S subunits for smFRET\(_{1.1-1.9}\) experiments and eight PRE-\(^{\text{A}}\) complexes harboring L1(Cy5)-
labeled 50S subunits and tRNA\textsuperscript{Phe} (Cy3) for smFRET\textsubscript{L1-tRNA} experiments (SI Materials and Methods and Fig. 2).

smFRET imaging of PRE\textsuperscript{A} complexes was accomplished using total internal reflection fluorescence microscopy as previously described (14, 21) (SI Materials and Methods). Consistent with previous studies, the smFRET\textsubscript{L1-tRNA} signals from all PRE\textsuperscript{A} complexes exhibited fluctuations between a FRET state with a FRET efficiency (E\textsubscript{FRET}) = 0.54 ± 0.02, assigned to the L1\textsubscript{o}–L1\textsubscript{o} conformation of the L1 stalk, and a FRET state with an E\textsubscript{FRET} = 0.35 ± 0.01, assigned to the L1\textsubscript{c} of the L1 stalk, thus reporting on the L1\textsubscript{o}–L1\textsubscript{o} equilibrium (Fig. 2 and Fig. S1A) (14). Likewise consistent with previous studies, the smFRET\textsubscript{L1-tRNA} signals from all PRE\textsuperscript{A} complexes exhibited fluctuations between a FRET state with an E\textsubscript{FRET} = 0.15 ± 0.02, assigned to the L1\textsubscript{tRNA} conformation of the PRE\textsuperscript{A} complex, and a FRET state with an E\textsubscript{FRET} = 0.74 ± 0.03, assigned to the the L1\textsubscript{tRNA} conformation of the PRE\textsuperscript{A} complex, thus reporting on the L1\textsubscript{tRNA}→L1\textsubscript{tRNA} equilibrium (Fig. 2 and Fig. S1B) (21).

L1-Stalk and P-Site tRNA\textsuperscript{Phe} Dynamics Are Coupled to Intersubunit Rotation. Relative to PRE\textsubscript{WT} (where the subscript denotes the identity of the 30S subunit of the PRE\textsuperscript{A} complex), PRE\textsubscript{S13} exhibits dramatic shifts of the L1\textsubscript{o}–L1\textsubscript{o} and L1\textsubscript{tRNA}→L1\textsubscript{tRNA} equilibria toward L1\textsubscript{c} and L1\textsubscript{tRNA}, respectively, increasing the equilibrium constants (K\textsubscript{L1} and K\textsubscript{L1-tRNA}, respectively) by sixfold each (Tables S2 and S3). These sixfold increases in K\textsubscript{L1} and K\textsubscript{L1-tRNA} correspond to free-energy difference changes between L1\textsubscript{o} and L1\textsubscript{c} (ΔΔG\textsubscript{L1}) and L1\textsubscript{tRNA} and L1\textsubscript{tRNA} (ΔΔG\textsubscript{L1-tRNA}) in PRE\textsuperscript{A}\textsubscript{S13} vs. PRE\textsuperscript{A} of ~1.0 kcal/mol\textsuperscript{-1} (where the negative values indicate shifts of the equilibria toward L1\textsubscript{c} and L1\textsubscript{tRNA}, respectively) (Fig. 2 A and B and Tables S2 and S3). Survival probability analyses (SI Materials and Methods) reveal that the observed equilibrium shifts are predominantly driven by fivefold increases in the rates of L1\textsubscript{o}→L1\textsubscript{c} and L1\textsubscript{tRNA}→L1\textsubscript{tRNA} (k\textsubscript{L1-o-L1-c} and L1\textsubscript{tRNA}→L1\textsubscript{tRNA} (k\textsubscript{L1-tRNA-L1-tRNA}) that are further augmented by 30% decreases in the rates of L1\textsubscript{o}→L1\textsubscript{o} (k\textsubscript{L1-o-L1-o}) and L1\textsubscript{tRNA}→L1\textsubscript{tRNA} (k\textsubscript{L1-tRNA-L1-tRNA}) (Tables S2 and S3). A similar comparison demonstrates that in vitro reconstitution of recombinant wild-type S13 into (−)S13 30S subunits yields a PRE\textsuperscript{A} complex [i.e., PRE\textsuperscript{A}\textsuperscript{WT}S13] with thermodynamic and kinetic properties that are virtually indistinguishable from those of PRE\textsuperscript{A} complexes formed using completely wild-type 30S subunits (i.e., PRE\textsuperscript{A}\textsuperscript{WT}S13) (21).

Because S13 deletion disrupts both S13–L5 and S13–P-site tRNA interactions (22), we next sought to investigate how single S13 substitution mutations designed to exclusively disrupt S13–L5 interactions modulate the thermodynamic and kinetic properties of PRE\textsuperscript{A} complexes. From the structural analyses described above (Fig. 1), we expected that PRE\textsuperscript{A}\textsuperscript{(D82K)S13} and PRE\textsuperscript{A}\textsuperscript{(R3D)S13}, which contain mutations designed to disrupt the

Fig. 2. (A–H) Steady-state smFRET measurements and changes in the free-energy differences of PRE\textsuperscript{A} complexes formed using (A) WT, (B) (−)S13, (C) WT S13, (D) (D82A)S13, (E) (D82K)S13, (F) (R3A)S13, (G) (R3D)S13, and (H) (R3D/D82K)S13 30S subunits. Cartoon representations of PRE\textsuperscript{A} complexes labeled 50S subunits and tRNA\textsuperscript{Phe} (Cy3) for smFRET\textsubscript{L1-tRNA} experiments (SI Materials and Methods and Fig. 2).
favorable (D82)S13–(R111)L5 interaction observed in the R-subunit orientation while having no effect on the NR-subunit orientation, would primarily destabilize the R-subunit orientation relative to PRE\(^{WT}\)S13 (note that the thermodynamic and kinetic properties of all PRE\(^{−}\) complexes harboring S13 mutants are reported relative to PRE\(^{WT}\)S13). As shown in Fig. 2 D and E and Tables S2 and S3, PRE\(^{−}\)D82K\(_{S13}\) and PRE\(^{−}\)D82K\(_{L1c}\) both exhibited twofold decreases of \(K_L\) and \(K_{1,1}\), corresponding to \(\Delta G_{L1}\) and \(\Delta A_{1,1}\) values of \(\sim 0.4\) kcal\(\cdot\)mol\(^{-1}\) that favor the L1, and L1-tRNA conformations. In both cases, the equilibrium shifts were almost exclusively driven by twofold increases in \(k_{L1c-L1o}\) and \(k_{1,1-R}\) with no detectable changes in \(k_{L1o-L1c}\) and \(k_{1,1-R}\) (Tables S2 and S3). It is remarkable that disruption of a single, noncovalent, electrostatic interaction across the subunit interface of the ribosome can alter the thermodynamic and kinetic properties of the entire PRE\(^{−}\) complex; it is clear from this observation that the collection of noncovalent interactions that comprise the intersubunit bridges has evolved to ensure the metastability of the PRE complex.

The effects of disrupting the (D82)S13–(R111)L5 interactions, which are discussed in greater detail in SI Results and Discussion, demonstrate that disruption of an S13–L5 interaction that is predicted to destabilize the R-subunit orientation and consequently result in an increased rate of R→NR transitions (\(k_{NR-R}\)), also destabilizes the L1\(_o\) conformation of the L1 stalk and the P/E conformation of the P-site tRNA\(^{Phc}\), resulting in increases in \(k_{L1o-L1c}\) and \(k_{1,1-R}\). Correspondingly, disruption of an S13–L5 interaction that is predicted to have no effect on the NR-subunit orientation or the rate of NR→R transitions (\(k_{NR-R}\)) has virtually no effect on the stability of the L1\(_o\), conformation of the L1 stalk and the P/E conformation of the P-site tRNA \((k_{P-E})\). The decreased stability of the L1\(_o\), state is consistent with a model in which L1 stalk and P-site tRNA\(^{Phc}\) dynamics are coupled to intersubunit rotation in PRE/PRE\(^{−}\) complexes (21). It is important to note, however, that the identity of the P-site tRNA can influence the conformational equilibria of PRE/PRE\(^{−}\) complexes (21, 23) and that the PRE\(^{−}\) complexes investigated here all contained exclusively tRNA\(^{Phc}\) at the P site. Thus, future experiments using tRNAs other than tRNA\(^{Phc}\) will be needed to determine whether and how the coupling of P-site tRNA dynamics to intersubunit rotation observed here depends on the identity of the P-site tRNA.

Based on the structural analysis described above and in Fig. 1, we likewise expected that PRE\(^{−}\)R3A\(_{S13}\) and PRE\(^{−}\)R3D\(_{S13}\), which were designed to disrupt the favorable (R3)S13–(D143)L5 interaction observed in the NR-subunit orientation while having no effect on the R-subunit orientation, would primarily result in a destabilization of the NR-subunit orientation. As shown in Fig. 2 F and G and Tables S2 and S3, PRE\(^{−}\)R3A\(_{S13}\) and PRE\(^{−}\)R3D\(_{S13}\) both exhibited four- to sixfold increases in \(K_L\) and \(K_{1,1}\), corresponding to \(\Delta G_{L1}\) and \(\Delta A_{1,1}\) values of \(\sim 1.0\) kcal\(\cdot\)mol\(^{-1}\) that favor the L1, and L1-tRNA conformations. In both cases, the equilibrium shifts were predominantly driven by two- to fivefold increases in \(k_{L1o-L1c}\) and \(k_{1,1-R}\) that were augmented by 30–40% decreases in \(k_{L1c-L1o}\) and \(k_{1,1-R}\) (Tables S2 and S3).

The effects of disrupting the (R3)S13–(D143)L5 interactions, which are discussed in greater detail in SI Results and Discussion, demonstrate that disruption of an S13–L5 interaction that is predicted to destabilize the NR-subunit orientation and consequently result in an increase in \(k_{NR-R}\), also destabilizes the L1\(_o\) conformation of the L1 stalk and the P/E configuration of the P-site tRNA\(^{Phc}\), resulting in increases in \(k_{L1o-L1c}\) and \(k_{1,1-R}\). Once again, these observations are consistent with a model in which L1 stalk and P-site tRNA\(^{Phc}\) dynamics are coupled to intersubunit rotation in PRE/PRE\(^{−}\) complexes (21). In contrast to the data obtained using PRE\(^{−}\)D82K\(_{S13}\) and PRE\(^{−}\)D82K\(_{L1c}\), however, the data obtained using PRE\(^{−}\)D82K\(_{S13}\) and PRE\(^{−}\)D82K\(_{L1c}\) exhibited small, but detectable and reproducible, decreases in \(k_{L1o-L1c}\) and \(k_{1,1-R}\) (Tables S3 and S4). These small effects likely result from an (R3)S13 mutation-mediated increase in the stability of the R-subunit orientation or a change in the stability of one or more of the transition-state subunit orientations that are sampled during R→NR transitions (SI Results and Discussion and Fig. S3).

Interestingly, PRE\(^{−}\)D82K\(_{S13}\), which harbors an (R3D/ D82K)S13 double mutation, exhibits thermodynamic and kinetic properties that are essentially the sum of those observed for PRE\(^{−}\)D82K\(_{S13}\) and PRE\(^{−}\)D82K\(_{L1c}\) strongly suggesting that, within wild-type ribosomes, (R3)S13–(D143)L5 and (D82)S13–(R111)L5 interactions largely act independently to regulate the intersubunit, L1 stalk, and tRNA dynamics of PRE complexes (SI Results and Discussion; Fig. 2 E, G, and H; and Tables S2 and S3).

The Architecture of the Ribosome Itself Couples L1-Stalk Dynamics to Intersubunit Rotation. The results presented in Fig. 2 demonstrate that destabilization of the NR-subunit orientation also destabilizes the L1\(_o\) conformation of the L1 stalk and increases \(k_{L1o-L1c}\), whereas destabilization of the R-subunit orientation also destabilizes the L1\(_c\) conformation of the L1 stalk and increases \(k_{L1c-L1o}\). This cooperativity between intersubunit rotation and L1-stalk dynamics is remarkable given that the (R3) S13 mutations are \(\sim 20\) Å away from the nearest residue within the L1 stalk (D102 in L1) in the R-subunit orientation and \(\sim 90\) Å away from the nearest residue within the L1 stalk (G2112 in H76) in the NR-subunit orientation (24, 25). Because structures of P-site tRNA-bound ribosomal complexes reveal that P-configured tRNAs directly contact residues in S13 and L5 whereas P-configured tRNAs directly contact residues in S13 and in the L1 stalk (10, 11), it is possible that P/Ps/P/E transitions serve to physically couple intersubunit rotation with L1-stalk movements. To test whether a P-site tRNA is strictly required for the cooperativity that we observe between intersubunit rotation and L1 stalk movements, we prepared vacant 70S ribosomes using wild-type 30S subunits and L1-L9-labeled 50S subunits, and lacking any bound tRNAs (70S\(_{WT}\), where the subscript identifies the 30S subunit in the vacant 70S ribosome) (SI Materials and Methods).

Compared with PRE\(^{−}\)D82K\(_{S13}\), 70S\(_{WT}\) exhibited an eightfold decrease in \(K_{L1}\) dramatically shifting the L1,\(_{o} \leftrightarrow L1c\) equilibrium toward L1\(_{o}\) (Fig. 2A, Fig. S4, and Tables S3 and S4). Taken together with previous smFRET studies demonstrating that, relative to a PRE\(^{−}\) complex similar to PRE\(^{−}\)D82K\(_{S13}\), the NR\(_{WT}\) equilibrium in vacant 70S ribosomes is shifted by a similar factor of 8 toward the NR-subunit orientation (13), this result already strongly suggests that the architecture of the ribosome itself serves to physically couple L1-stalk movements to intersubunit rotation. To test this possibility further, we prepared a series of seven vacant 70S ribosomes, using the same 30S subunits that were used to prepare the PRE\(^{−}\) complexes described above. Notably, all of the vacant 70S ribosomes that we prepared exhibited decreases in \(K_{L1}\), \(\Delta A_{L1}\), \(K_{L1o-L1c}\), and \(K_{L1c-L1o}\) compared with 70S\(_{WT}\) that mirrored those observed when the corresponding PRE\(^{−}\) complexes were compared with PRE\(^{−}\)S13 (Tables S3 and S4).

To confirm that the changes in L1-stalk dynamics that we observe upon disruption of S13–L5 interactions across bridge B1b are indeed due to changes in intersubunit rotation, we conducted a series of smFRET\(_{1b-1b}\) experiments using PRE\(^{−}\)D82K\(_{S13}\) and 70S\(_{WT}\) in which we altered the dynamics of intersubunit rotation by perturbing a different intersubunit bridge, using a ribosome-targeting antibiotic. The tuberactinomycin antibiotic viomycin is a potent translocation inhibitor that binds within bridge B2a and stabilizes the ribosome in its R-subunit orientation (26), thereby perturbing intersubunit rotation in a manner.
Changes in the Relative Stabilities of Two Global Conformations of the PRE Complex Regulate the Rate and Extent of EF-G–Independent Translocation. As described above, the smFRET data obtained using PRE\(^{\text{WT}}\)\(_{13}\) and PRE\(^{\text{D82K}}\)\(_{13}\) reveal that S13 suppresses EF-G–independent translocation in wild-type PRE complexes (19) by shifting the conformational equilibria of the PRE complex toward L1\(_c\), L1\(_c\)tRNA, and, presumably, NR such that the PRE complex is in a conformation that is incompatible with translocation. In addition, we have characterized single substitution mutations within S13 (e.g., D82K\(_{13}\)) that likewise shift the conformational equilibrium of the PRE\(^{\text{D82K}}\) complex toward L1\(_c\), and L1\(_c\)tRNA by stabilizing an NR/L1\(_c\)/P(P)/ global conformation of the PRE\(^{\text{D82K}}\) complex in which the ribosome, L1 stalk, and P-site tRNA\(^{\text{Phe}}\) occupy the NR-subunit orientation, the L1\(_c\) configuration, and the P/P configuration, respectively.

Conversely, we have characterized single substitution mutations within S13 (e.g., R3D\(_{13}\)) that shift the conformational equilibria of the PRE\(^{\text{R3D}}\) complex toward L1\(_c\), and L1\(_c\)tRNA by stabilizing an R/L1\(_c\)/P(E) global conformation of the PRE\(^{\text{R3D}}\) complex in which the ribosome, L1 stalk, and P-site tRNA\(^{\text{Phe}}\) occupy the R-subunit orientation, the L1\(_c\) configuration, and the P/E configuration, respectively.

To test whether the stabilities of these two global conformations of the PRE\(^{\text{A}}\) complex regulate the propensity of PRE complexes to undergo EF-G–independent translocation, we prepared PRE complexes analogous to PRE\(^{\text{R3D}}\)\(_{13}\) and PRE\(^{\text{D82K}}\)\(_{13}\) and assessed the propensity of each of these PRE complexes to undergo EF-G–independent translocation. To do this, we used two independent biochemical assays: a puromycin reactivity assay and a tripeptide synthesis assay (SI Materials and Methods and Fig. S6). The results of these experiments demonstrate that, relative to PRE\(^{\text{WT}}\)\(_{13}\), PRE\(^{\text{R3D}}\)\(_{13}\) complexes exhibit a shift toward R/L1\(_c\)/P(E) [i.e., PRE\(^{\text{R3D}}\)\(_{13}\)–PRE\(^{\text{R3D/DS2K/S13}}\) and PRE\(^{\text{R3D/DS2K/S13}}\)–PRE\(^{\text{R3D/DS2K/S13}}\)], whereas PRE\(^{\text{D82K}}\)\(_{13}\) displays a corresponding increase in the rate and extent of EF-G–independent translocation, whereas the PRE complex exhibiting a shift toward NR/L1\(_c\)/P(P) [i.e., PRE\(^{\text{D82K/DS2K/S13}}\)] displays a corresponding decrease in the rate and extent of EF-G–independent translocation. Collectively, these results demonstrate that S13 indeed suppresses EF-G–independent translocation, at least in part, by stabilizing the NR/L1\(_c\)/P(P) global conformation of PRE complexes via bridge B1b. Similar results were obtained when we varied the experimental conditions (i.e., the Mg\(^{2+}\) concentration) to alter the relative stabilities of the NR/L1\(_c\)/P(P) and R/L1\(_c\)/P(E) global conformations of the PRE complex within a representative PRE complex, PRE\(^{\text{R3D/S13}}\), and assessed the propensity of this PRE complex to undergo EF-G–independent translocation (Fig. S6). Thus, any factor that can modulate the relative stabilities of the NR/L1\(_c\)/P(P) and R/L1\(_c\)/P(E)
Cooperative Conformational Changes Enable the Ribosome to Maximize and Regulate the Efficiency of Translation. Numerous experimental and computational studies strongly suggest that the architectures and corresponding conformational free-energy landscapes of small monomeric protein enzymes have evolved such that complex networks of cooperative conformational changes facilitate catalysis and allosteric regulation (1, 2). Our findings here demonstrate that the architecture and corresponding conformational free-energy landscape of the ribosome have evolved such that cooperative intersubunit rotation and L1-stalk dynamics ensure that vacant 70S ribosomes can efficiently sample NR/L1, and R/L1 global conformations (Fig. 3A). The presence of a P-site tRNA in within a PRE complex remodels this landscape, further increasing the cooperativity between intersubunit rotation and L1-stalk dynamics and enabling the PRE-complex to efficiently sample NR/L1/(P/P) and R/L1/(P/E) global conformations (Fig. 3B). Within the PRE complex, such cooperativity maximizes the efficiency with which the PRE complex samples the R/L1/(P/E) global conformation that is productive for translocation, thereby ensuring that the translating ribosome can undergo rapid translocation during the elongation cycle. Likewise, because POST complexes predominantly sample the NR-subunit orientation (13), such cooperativity maximizes the efficiency with which the POST complex samples the NR/L1/(P/P) global conformation in which the L1 stalk occupies the L1i conformation that enables ejection of the decacylated tRNA that was just translocated into the E site from the ribosome (27), thereby ensuring that the translating ribosome can rapidly progress into the next round of the elongation cycle.

In addition to maximizing the efficiency of mechanical processes such as the translocation and ejection of tRNAs, cooperative conformational changes can also be exploited for the allosteric regulation of protein synthesis by translation factors and antibiotic inhibitors. Previously, (14) and others (15) have shown that in addition to stabilizing the R-subunit orientation (13), binding of EF-G to PRE-complexes stabilizes the L1i conformation of the L1 stalk. This occurs despite the fact that EF-G does not directly contact either the L1 stalk or the intervening P-site tRNA within the PRE complex (14). The present study now allows us to rationalize this observation, strongly suggesting that EF-G uses the cooperative conformational changes that we have characterized here to allosterically regulate the dynamics of the L1 stalk by directly modulating the dynamics of intersubunit rotation. Similarly, the results of the viomycin experiments shown in Fig. 3B demonstrate that antibiotic inhibitors of translocation can exploit the cooperative conformational changes that we observe here as part of the mechanism through which they so strongly inhibit translocation. More generally, cooperative conformational changes such as the ones that we have characterized here are likely to play important roles in regulating the efficiency of many of the other steps of protein synthesis as well as the efficiency of the numerous biological processes that are carried out by other biomolecular machines.

Methods

smFRET Experiments. All smFRET experiments were performed in Tris-polyimine buffer (50 mM Tris-OAc (pH 7.5), 100 mM KCl, 5 mM NH4OAc, 0.5 mM Ca(OAc)2, 0.1 mM EDTA, 10 mM β-mercaptoethanol, 5 mM putrescine-HCl, and 1 mM spermidine, free base) containing 15 mM Mg(OAc)2 and supplemented with an oxygen-scavenging system (protocatechuate/proocatechuate-3,4-dioxygenase) (28) and a triplet-state quencher mixture (1 mM 1,3,5-tricyclooctatetraene (Sigma-Aldrich) and 1 mM 3-nitrobenzyl alcohol (Fluka)) (29). Further details regarding data acquisition, processing, and analysis are in SI Materials and Methods.

EF-G–Indepedent Translocation Experiments. Puromycin reactivity and tripeptide synthesis assays were performed in translocation buffer (50 mM Tris-HCl (pH 7.5), 70 mM NH4Cl, 30 mM KCl, 1 mM DTT) containing 7 mM MgCl2. Further details regarding data acquisition, processing, and analysis are in SI Materials and Methods.

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