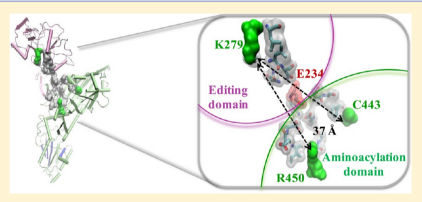


# Multiple Pathways Promote Dynamical Coupling between Catalytic Domains in Escherichia coli Prolyl-tRNA Synthetase

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Supporting Information



ABSTRACT: Aminoacyl-tRNA synthetases are multidomain enzymes that catalyze covalent attachment of amino acids to their cognate tRNA. Cross-talk between functional domains is a prerequisite for this process. In this study, we investigate the molecular mechanism of site-to-site communication in Escherichia coli prolyl-tRNA synthetase (Ec ProRS). Earlier studies have demonstrated that evolutionarily conserved and/or co-evolved residues that are engaged in correlated motion are critical for the propagation of functional conformational changes from one site to another in modular proteins. Here, molecular simulation and bioinformatics-based analysis were performed to identify dynamically coupled and evolutionarily constrained residues that form contiguous pathways of residue-residue interactions between the aminoacylation and editing domains of Ec ProRS. The results of this study suggest that multiple pathways exist between these two domains to maintain the dynamic coupling essential for enzyme function. Moreover, residues in these interaction networks are generally highly conserved. Site-directed changes of onpathway residues have a significant impact on enzyme function and dynamics, suggesting that any perturbation along these pathways disrupts the native residue-residue interactions that are required for effective communication between the two functional domains. Free energy analysis revealed that communication between residues within a pathway and cross-talk between pathways are important for coordinating functions of different domains of Ec ProRS for efficient catalysis.

lass II prolyl-tRNA synthetases (ProRSs) catalyze covalent attachment of proline to tRNA Pro in a two-step

$$Pro + ATP + ProRS \leq Pro-AMP \cdot ProRS + PP_i$$
 (i)

$$Pro-AMP \cdot ProRS + tRNA^{Pro}$$

$$\rightarrow$$
 Pro-tRNA<sup>Pro</sup> + AMP + ProRS (ii)

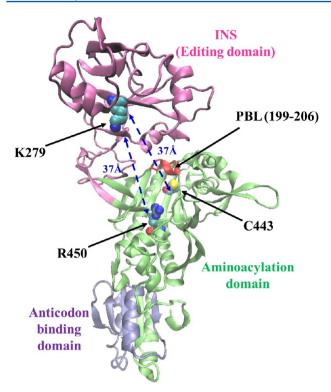
ProRSs are modular proteins and are divided into two evolutionarily distinct groups based on sequence alignment and structural architecture. The "prokaryotic-like" ProRSs contain an insertion domain (INS) between motifs 2 and 3 of the catalytic domain, whereas "eukaryotic-like" ProRSs have Cand/or N-terminal extension domains.<sup>3</sup> ProRSs from all three kingdoms of life have been shown to misactivate noncognate alanine and cysteine, resulting in mischarged tRNAPro.4,5 Many

aminoacyl-tRNA synthetases (AARSs) have evolved proofreading capabilities to correct mistakes in amino acid activation (pre-transfer editing) and charging (post-transfer editing).<sup>6</sup> It has been shown that a domain (INS) of approximately 180 residues inserted within the catalytic core of prokaryotic-like ProRSs (Figure 1) is a post-transfer editing active site that hydrolyzes specifically mischarged Ala-tRNA Pro 5,7,8 In contrast, Cys-tRNA<sup>Pro</sup> is hydrolyzed by an INS homologue known as YbaK, which is encoded as a single-domain protein by many bacteria. 9,10 Unlike prokaryotic-like ProRSs, most eukaryoticlike ProRSs do not possess the INS domain but in some cases encode free-standing editing-domain homologues. Some lower-

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**Figure 1.** Ribbon representation of the three-dimensional structural model of the monomeric form of Ec ProRS. The homology model was derived from the X-ray crystal structure of Ef ProRS. <sup>15</sup> The  $C_{\alpha}$ – $C_{\alpha}$  distances between the starting residues (C443 and R450) and the end residue (K279) are shown.

eukaryotic ProRSs encode an N-terminal domain that displays a low degree of homology with bacterial INS. This domain exhibits post-transfer editing activity against Ala-tRNA<sup>Pro</sup> in *Plasmodium falciparum*<sup>11</sup> but is a defunct editing domain in *Saccharomyces cerevisiae*.<sup>12</sup>

Bacterial ProRSs are modular enzymes, and efficient catalysis and editing requires effective communication between distant domains. Escherichia coli (Ec) ProRS, a representative member of the prokaryotic-like group, contains three distinct domains (Figure 1). The aminoacylation domain (motifs 1–3, consisting of residues 64-81, 128-164, and 435-465, respectively) catalyzes the activation of proline and the aminoacylation of tRNA Pro, as well as pre-transfer editing; 13 the anticodon binding domain (residues 506-570) is critical for recognition of cognate tRNA, and the INS (residues 224-407, located between motifs 2 and 3 of the catalytic domain) is the posttransfer editing active site.<sup>7,8</sup> The aminoacylation domain and the INS of Ec ProRS have been observed to depend on each other in terms of their individual catalytic activities. For example, mutation of a highly conserved aspartate (D350) in INS to alanine resulted in reduced aminoacylation activity. Previous studies have also indicated that covalent connectivity between domains is a prerequisite for efficient aminoacylation and editing functions by ProRSs and other AARS systems.<sup>14</sup> Deletion of the INS of Ec ProRS ( $\Delta$ INS) has a severe impact on catalysis; the amino acid activation efficiency of the deletion variant was reduced 1200-fold.<sup>13</sup> The addition of a separately cloned and purified INS in trans failed to stimulate the amino acid activation efficiency of the  $\Delta$ INS construct (S. Hati and K. Musier-Forsyth, unpublished data). The requirement of covalent connectivity between domains for efficient function

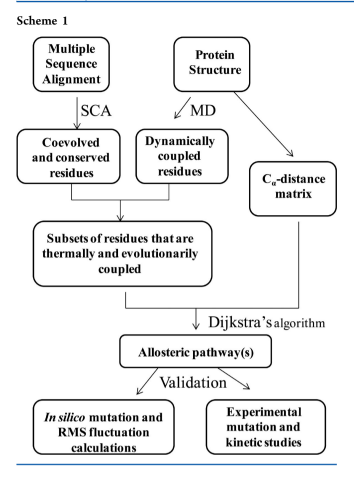
of Ec ProRS suggests the existence of interdomain communication in this enzyme. Moreover, structural studies revealed that the catalytically important proline-binding loop (PBL, residues 199–206) of bacterial ProRSs undergoes a conformational change from the "open" to "closed" state upon substrate binding. <sup>15</sup> A recent study showed that any perturbation in the surrounding structural elements has a significant impact on PBL dynamics. <sup>16</sup> The pathway by which substrate-induced conformational change propagates from the activation center to the distant protein segments that modulate PBL dynamics and conformational change is unknown.

For multidomain proteins like Ec ProRS, domain—domain communication is achieved by coupled-domain dynamics. <sup>17–20</sup> Recent MD simulation results revealed that the INS of Ec ProRS is engaged in coupled motion with structural elements of the catalytic domain. <sup>16</sup> The collective dynamics of the PBL is altered by the deletion of the INS or point mutation at the INS—aminoacylation domain junction. <sup>16</sup> To understand the molecular mechanism by which different structural elements of Ec ProRS coordinate their function, it is important to identify networks defined by key residue—residue interactions that promote coupled-domain dynamics in this enzyme.

In this study, a bioinformatics-based computational method, statistical thermal coupling analysis (STCA), was employed to trace pathways of site-to-site communication in Ec ProRS. Previous application of this method revealed that dynamically coupled and evolutionarily constrained residues are important for maintaining coupled-domain dynamics in Thermus thermophilus leucyl-tRNA synthetase (Tt LeuRS).<sup>20</sup> Here STCA was used to identify the residue-residue interaction networks between the INS and aminoacylation domains of Ec ProRS. In addition, mutational and kinetic studies, as well as thermal fluctuation analyses, were performed to validate the predicted networks of residue-residue interactions. In summary, this study demonstrates that a modular protein like Ec ProRS employs multiple pathways of residue-residue interactions to communicate between distant functional sites. Moreover, networks of these pathways involve residues that are evolutionarily constrained and engaged in correlated motion.

## MATERIALS AND METHODS

General Strategy. On the basis of reported experimental and structural data, the starting and ending points of interaction networks between the aminoacylation domain and the INS were selected; these active sites are >30 Å apart (Figure.1). The single cysteine residue (C443), which is important for amino acid activation, was chosen as one of the starting points within the amino acid activation site. The other starting residue is R450, found to be important for substrate binding <sup>15</sup> (Figure 1). The end point was chosen to be K279 in the INS, which has been shown to be required for the editing function<sup>8</sup> and is 37 Å from C443 and R450 (Figure 1). STCA was conducted in three discrete steps (Scheme 1). First, statistical coupling analysis (SCA)<sup>21,22</sup> was performed to identify conserved and co-evolved residues in the ProRS family. In a parallel study, the collective motions of various domains of the protein were studied by performing a long time scale molecular dynamics (MD) simulation of Ec ProRS. In the second step, the evolutionary dependence of the coupled-domain dynamics was explored. In this step, results of the SCA and MD simulation were computationally integrated (Scheme 1). This analysis identified a subset of residues that are not only dynamically coupled but also evolutionarily constrained. Next, distinct networks of



interacting residues between the two distant sites were identified from this subset of residues using Dijkstra's algorithm.<sup>23</sup> Finally, the role of these selected residues in site-to-site communication was probed experimentally by conducting site-directed mutagenesis and kinetic studies. In addition, *in silico* mutations were introduced, and their impact on protein dynamics was examined by comparing root-mean-square (rms) fluctuations of the WT and mutated variants.<sup>16</sup>

Proteins were visualized using VMD.<sup>24</sup> SCA was conducted using a MATLAB script obtained from the Ranganathan lab (http://systems.swmed.edu/rr\_lab/sca.html). The MD plots and SCA plots were created and all data processing conducted using MATLAB R2006b (The MathWorks Inc., Natick, MA).

#### COMPUTATIONAL METHODS

Molecular Dynamics Simulations. MD simulations were conducted using the three-dimensional homology model of monomeric Ec ProRS (residues 1–567) (provided by S. Cusack). The model was generated using the crystal structure of *Enterococcus faecalis* ProRS, <sup>15</sup> which is 48% identical to the Ec enzyme. Mutant proteins were built using the Mutator plugin of VMD.<sup>24</sup> Simulations were performed in water (TIP3P model<sup>25</sup>) with substrate-free enzymes using the all-atom CHARMM22 force field<sup>26</sup> within the NAMD package.<sup>27</sup> All simulations were conducted with a 500 ps equilibration step followed by a 25 ns production MD run. The details of the MD simulation protocol were as described previously.<sup>16</sup> To evaluate the statistical significance of the MD simulation analysis, three replicates were generated for each protein system, as described in the protein simulation studies by Roy and Laughton.<sup>28</sup>

The correlated motions between residue pairs of distant structural elements were studied by principal component analysis (PCA, also known as essential dynamics analysis) of collective motions, <sup>29</sup> as described previously. <sup>16,30</sup> PCA is a procedure by which the low-frequency (high-amplitude) collective motions of a biomolecule, which are often more relevant for its functions, are extracted from a MD simulation trajectory. <sup>31</sup> This method has been described in detail in our previous study with this enzyme. <sup>16</sup> Briefly, a covariance matrix of the  $C_{\alpha}$  atoms was generated using the simulated MD data set. The diagonalization of this covariance matrix produces eigenvectors and eigenvalues representing the direction and magnitude of the collective motion of the whole protein or protein segments of interest.

In this study, the last 20 ns of the MD simulation data was used to generate principal components of atomic (backbone  $C_{\alpha}$  atoms) fluctuations using Carma.<sup>32</sup> The first three principal components were used to perform PCA-based cluster analysis, which produced a new trajectory of conformations representing the predominant conformational fluctuations. Finally, these conformations were used to generate dynamic cross-correlation matrix  $\mathbf{C}$ , in which the ijth element,  $C_{ij}$ , represents the cross-correlation coefficient between fluctuations of residues at sites i and j during the simulation:

$$C_{ij} = \frac{\langle (x_i - \langle x_i \rangle)(x_j - \langle x_j \rangle) \rangle}{\sigma_{x_i} \sigma_{x_j}}$$
(1)

The atomic  $(C_{\alpha})$  displacements of residues i and j are represented by  $x_i$  and  $x_j$  respectively; the angular brackets represent ensemble averages, and  $\sigma_{x_i}$  and  $\sigma_{x_j}$  represent the standard deviations of these displacements.

The rms fluctuations of  $C_\alpha$  atoms averaged over three replicate simulations were also obtained for the WT and mutants. In these calculations, the last 20 ns of MD simulation data was used, each comprising an ensemble of 200000 conformations. PCA was also conducted for the WT and mutant proteins to compare the effect of mutations on coupled dynamics. For each of these protein systems, an ensemble of 600000 conformations, obtained by combining three replicate trajectories, was used to perform PCA. The first three clusters representing the predominant conformational fluctuations were considered in this study. In addition, the collective dynamics of the catalytically important PBL of the WT and various mutants were also analyzed following the method described previously.  $^{16,30}$ 

**Statistical Coupling Analysis.** SCA is based on the assumption that the coupling of two sites in a protein, whether for structural or functional reasons, should cause those two sites to co-evolve. SCA was conducted using the standard protocol, details of which have been published and are available at http://www.hhmi.swmed.edu/Labs/rr/sca.html. In this work, a multiple-sequence alignment of 492 protein sequences of the ProRS family was generated using BLAST. Only ProRS sequences that are significantly identical in sequence (>67%) with Ec ProRS were included in this study. Conservation constant  $\Delta G_i^{\rm stat}$  and coupling constant  $\Delta \Delta G_{ij}^{\rm stat}$  were obtained using standard procedures described previously.

Statistical Thermal Coupling Analysis: Integration of Evolutionary and Dynamic Information. In our earlier study of Tt LeuRS, we observed that coupled-domain motions are facilitated by networks of thermally and evolutionarily

constrained residues.<sup>20</sup> To identify residue clusters that are important for maintaining coupled-domain dynamics, we extracted a subset of residues that are simultaneously coupled through evolution and dynamics (Scheme 1). The motional coupling information from MD was integrated with the evolutionary conservation and co-evolution data set obtained from the SCA study.

The conserved and co-evolved residues were treated separately. The conserved and dynamically coupled residues were chosen by selecting only those residues that exhibit significant conservation ( $\Delta G_i^{\rm stat} \geq 0.5$ ;  $\Delta G_i^{\rm stat}$  values range from 0 to 1.0), as well as motional coupling  $[C_{ij} \geq 0.8; C_{ij}$  values range from -1.0 (anticorrelated motion) to 1.0 (correlated motion)] with each other. On the basis of recent studies of the significance of correlated motion in long-range communication, only positive correlations were considered. The value of  $C_{ij}$  was set to  $\geq 0.8$  to obtain residues that are engaged in strong correlated motions.

To select the co-evolved and dynamically coupled residues, the dynamic cross-correlation matrix (C matrix) was truncated by including only those columns that are present in the normalized SCA-derived G matrix. Next, the co-evolutionary dynamic coupling, CDC, matrix was created by multiplying each *ij*th element of the G matrix with the corresponding element of the truncated C matrix:

$$CDC_{ij} = \Delta \Delta G_{ij}^{\text{stat}} \times C_{ij}$$
 (2)

The CDC matrix therefore contains the covariance information of residue pairs that are co-evolved as well as dynamically coupled. In this study, the co-evolved and dynamically correlated residues are extracted by choosing only those residues for which CDC<sub>ij</sub>  $\geq$  0.4 ( $\Delta\Delta G_{ij}^{\text{stat}} \geq$  0.5;  $C_{ij} \geq$  0.8).

**Identification of Interaction Networks Using Dijkstra's Algorithm.** From the short-listed residues (Scheme 1), residue—residue interaction networks between C443 or R450 (aminoacylation domain) and K279 (INS) of Ec ProRS were identified using Dijkstra's algorithm.<sup>23</sup> In this method, each of the  $C_{\alpha}$  atoms of the protein backbone represents a node. The connectivity between two adjacent nodes was described by a binary connection matrix **P** of inter-residue  $(C_{\alpha}-C_{\alpha})$  contacts. The  $C_{\alpha}-C_{\alpha}$  distance matrix, **D**, was computed from the Cartesian coordinates of all  $C_{\alpha}$  atoms of the protein. Based on a  $C_{\alpha}-C_{\alpha}$  cutoff distance  $D_{ij}^{o}$ ,  $P_{ij}$  is equal to 1 if  $D_{ij} < D_{ij}^{o}$  and zero otherwise. The interaction networks (paths) between two functional sites were identified and listed in terms of a "cost", which is equal to the sum of all  $C_{\alpha}-C_{\alpha}$  distances between adjacent residues in a given path.

## EXPERIMENTAL METHODS

**Materials.** All amino acids (Sigma) were of the highest quality (>99% pure) and used without any further purification.  $[\gamma^{-32}P]ATP$  and  $[^{32}P]PP_i$  were from Perkin-Elmer. Primers for site-directed mutagenesis and polymerase chain reaction were from Integrated DNA Technologies.

**Enzyme Preparation.** Overexpression and purification of histidine-tagged WT and mutant Ec ProRS were performed as described previously. Plasmids encoding D198A, E234A, H302A, N305A, G412A, F415A, H302A/G412A, N305A/G412A, and E218A/N305A Ec ProRS were generated by site-directed mutagenesis of pCS-M1S<sup>38</sup> using primers listed in Table S1 of the Supporting Information. Results of mutagenesis were confirmed by DNA sequencing (Biotechnology Center,

University of Wisconsin—Madison, Madison, WI). Protein expression was induced in Ec SG13009 (pREP4) competent cells with 0.1 mM isopropyl  $\beta$ -D-thiogalactoside for 4 h at 37 °C. Histidine-tagged proteins were purified using a Talon cobalt affinity resin, and the desired protein was eluted with 100 mM imidazole. Protein concentrations were determined initially by the Bio-Rad assay (Bio-Rad Laboratories) followed by active site titration.<sup>39</sup>

RNA Preparation. Ec tRNA<sup>Pro</sup> was transcribed using T7 RNA polymerase from BstNI-linearized plasmid as described previously<sup>40</sup> and purified by denaturing 12% polyacrylamide gel electrophoresis.

**Enzyme Assays.** The ATP $-PP_i$  exchange assay was performed at 37 °C according to the published method. The concentrations of proline ranged from 0.05 to 2 mM. The enzyme concentrations used were 10–40 nM for proline activation. Kinetic parameters were determined from Lineweaver–Burk plots and represent the average of at least three determinations.

ATP hydrolysis reactions for monitoring pre-transfer editing were conducted as described previously  $^{13}$  using 500 mM alanine and 4  $\mu$ M ProRS.

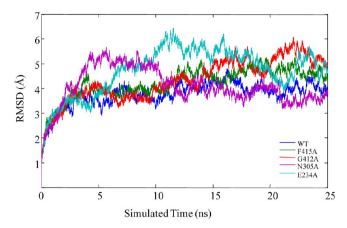
Aminoacylation assays were performed at room temperature in a reaction mixture containing 50 mM HEPES (pH 7.5), 20 mM KCl, 25 mM MgCl<sub>2</sub>, 0.1 mg/mL bovine serum albumin, 20 mM 2-mercaptoethanol, and 4 mM ATP. Cognate aminoacylation reaction mixtures also contained 23  $\mu$ M [³H]proline, 0.5  $\mu$ M tRNA<sup>Pro</sup>, and 100 nM Ec ProRS. When charged tRNA<sup>Pro</sup> was prepared to be used in deacylation assays, the same reaction mixture was used, and [¹⁴C]alanine was used to acylate G1:C72/U70 tRNA<sup>Pro</sup> (8  $\mu$ M) by Ec alanyl-tRNA synthetase (8  $\mu$ M). After incubation for 1.5 h, 10% acetic acid was used to quench the reaction, and the mischarged tRNA was purified by repeated phenol/CHCl<sub>3</sub> extraction (5:1 solution, pH 4.5), followed by ethanol precipitation.

Deacylation assays were conducted at room temperature in reaction mixtures containing 50 mM HEPES (pH 7.5), 5 mM MgCl<sub>2</sub>, and 1  $\mu$ M G1:C72/U70 [ $^{14}$ C]Ala-tRNA $^{Pro}$  and initiated with 0.5  $\mu$ M Ec ProRS. A buffer-only background curve was also performed and subtracted from each curve.

## RESULTS

**MD Simulation.** For the substrate-free WT ProRS, the root-mean-square-deviation (rmsd) of each frame of the 25 ns MD simulation trajectory was computed relative to the initial coordinates (Figure 2). The rmsd values of these frames fluctuated with a mean value of 1.5–2.0 Å during the last 20 ns of the simulations, and similar fluctuations in rmsd values were obtained for all three replicate simulations. These variations are consistent with previous studies of substrate-unbound AARSs. 42–44

**Dynamic Cross-Correlation Analysis.** The dynamic cross-correlation matrix (C) for the WT enzyme (Figure S1 of the Supporting Information) was generated using the first three principal components. Analysis of the cross-correlation of fluctuations of residues revealed both inter- and intradomain dynamic correlation. The aminoacylation domain and the INS are mainly engaged in anticorrelated motions; i.e., their displacements are in opposite directions ( $C_{ij} < 0$ ). However, various structural elements within the aminoacylation domain and the INS are engaged mainly in correlated motion ( $C_{ij} > 0$ ). The C matrix (Figure S1 of the Supporting Information) was



**Figure 2.** rmsd of the  $C_{\alpha}$  atoms from their initial coordinate as a function of time for WT Ec ProRS and four variants. Calculations of rmsds were performed using 25 ns MD simulation data.

used to extract residues that are engaged in strong correlated motion ( $C_{ij} \ge 0.8$ ).

Conserved and Co-Evolved Residues. Earlier, we showed that evolutionarily constrained residues are important for maintaining coupled-domain dynamics in Tt LeuRS. To identify the conserved residues responsible for coupled-domain dynamics between aminoacylation and editing active sites of Ec ProRS, the  $\Delta G_i^{\rm stat}$  value of each residue, a quantitative measure of the conservation of a residue at the *i*th position of the sequence, was calculated using SCA. The  $\Delta G_i^{\rm stat}$  values were obtained as a normalized one-dimensional vector (Figure S2 of the Supporting Information). The number of conserved residues in Ec ProRS obtained at different  $\Delta G_i^{\rm stat}$  cutoffs is reported in Table 1. In addition, a 567 × 152 coupling matrix

Table 1. Numbers of Conserved and Co-Evolved Residues within Ec ProRS That Are Engaged in Correlated Motion<sup>a</sup>

$\Delta G_i^{ ext{stat}}$	≥0.50	≥0.55	≥0.60	≥0.65	≥0.70
no. of conserved residues	207	182	151	104	61
$CDC_{ij}$	≥0.40	≥0.45	≥0.50	≥0.55	≥0.60
no of so avalved residues	06	60	10	20	10

"These residue numbers were determined by varying cutoff values for  $\Delta G_i^{\text{stat}}$  (evolutionary conservation constant) and  $\text{CDC}_{ij}$  ( $\text{CDC}_{ij} = \Delta \Delta G_{ij}^{\text{stat}} \times C_{ij}$ ; co-evolutionary dynamic coupling constant) while maintaining a correlation coefficient of residue—residue fluctuations ( $C_{ij}$ ) of  $\geq 0.8$ .

containing the evolutionary coupling indices  $(\Delta \Delta G_{ij}^{\text{stat}})$  of the 567 total residues for 152 perturbation sites<sup>21,22</sup> was obtained from SCA (Figure 3a). It is evident that only a small fraction of Ec ProRS residues have high  $\Delta \Delta G_{ij}^{\text{stat}}$  values. The multiplication of this 567 × 152 SCA matrix with the truncated 567 × 152 MD matrix yields the CDC matrix (eq 2, Figure 3). The numbers of evolutionarily and dynamically coupled residues determined for various CDC $_{ij}$  cutoffs are listed in Table 1. These pools of conserved and co-evolved residues were further used to identify long-range interaction networks described below.

**Interaction Networks across Domains.** To map the interaction networks between the aminoacylation domain (C443/R450) and the INS (K279), Dijkstra's algorithm<sup>23</sup> was applied to the pool of dynamically and evolutionarily (conserved and/or co-evolved) coupled residues (Scheme 1), as described in the methods sections. Because the average van

der Waals radius of an amino acid is ~3 Å and the acceptable distance for the noncovalent interactions is 2.0–3.0 Å, a strong noncovalent interaction will be prevalent when two  $C_{\alpha}$  atoms are separated by a distance of 8.0–9.0 Å. Therefore, in this study, distance cutoff  $D_{ij}^{o}$  for the potential interactions between the neighboring  $C_{\alpha}$  atoms was varied between 8.0 and 9.0 Å.

Several residue—residue interaction networks were identified between the aminoacylation and INS domains. These contiguous interaction networks were obtained by using various cutoff values for  $\Delta G_i^{\rm stat}$ ,  $C_{ij}$ , and  $D_{ij}^{\rm o}$  as listed in Table 2 while maintaining a CDC $_{ij}$  value of  $\geq$ 0.4 (CDC $_{ij} = \Delta\Delta G_{ij}^{\rm stat} \times C_{ij}$ ;  $\Delta\Delta G_{ij}^{\rm stat} \geq$  0.5;  $C_{ij} \geq$  0.8). The four probable contiguous pathways (paths I–IV), identified on the basis of the distance between the starting and ending residues (cost), as well as the degree of dynamic correlation between networking residues (average correlation coefficient value), are listed in Table 2 and shown in Figure 4. These predicted paths (interaction networks) are dominated by polar residues. Moreover, close scrutiny of residues in the predicted pathways revealed that conserved residues are dominant over co-evolved residues.

To evaluate the proposed communication pathways, we conducted mutational studies on some of the conserved residues (Table 2, underlined) that are present in more than one of the predicted pathways. The impact of altering pathway residues on amino acid activation, tRNA aminoacylation, and editing reactions was investigated.

Amino Acid Activation. First, the WT enzyme and six single mutants [F415A, G412A, and D198A (aminoacylation domain); N305A, H302A, and E234A (INS)] were examined for their role in amino acid activation. Using the ATP-PP, exchange reaction, we found that mutation of some of these residues to alanine had a considerable impact on proline activation (Table 3). Alanine substitution of F415 of the catalytic domain, which undergoes a conformational change upon prolyl-adenylate binding, 15 resulted in an ~400-fold decrease in  $k_{\rm cat}/K_{\rm M}$ . Mutation of residues in the INS domain also had a significant impact on amino acid activation. For example, significantly lower levels of amino acid activation efficiency were observed for N305A (~50-fold reduced) and E234A (~10-fold reduced) variants. To examine if residues in the predicted pathways are indeed coupled, amino acid activation efficiencies of double mutants were also evaluated. The H302A/G412A variant, obtained by alanine substitution of residues that belong to the two separate pathways (H302, path I, and G412, path III), exhibited a >4-fold decreased proline activation efficiency (Table 3); the H302A and G412A single mutants exhibited a 2.1- and 1.6-fold reductions in amino acid activation efficiency, respectively. The free energy analysis  $[\Delta \Delta G = -RT \ln(\text{fold decrease in } k_{\text{cat}}/K_{\text{M}})$ , where R is the gas constant, 1.986 cal  $K^{-1}$  mol<sup>-1</sup>, and T is 310 K]<sup>42</sup> showed that the free energy change of the double mutant (0.88 kcal/mol) is somewhat greater than the sum of the free energy change of the H302A (0.46 kcal/mol) and G412A (0.29 kcal/mol) mutants (Table 3). This difference in free energy change implies weak coupling between H302 and G412.<sup>45</sup> Proline activation was not detected for the other two double mutants tested, N305A/ G412A (path III) and E218A/N305A (path IV). The lack of activity of the path III variant suggests strong coupling between N305A and G412A because the effect of the double mutant  $(\Delta \Delta G > 3.7 \text{ based on the level of detection of this assay})$  is greater than expected on the basis of the single mutant effects (2.4 and 0.29 kcal/mol for N305A and G412A, respectively). In a recent study, we showed that the proline activation efficiency

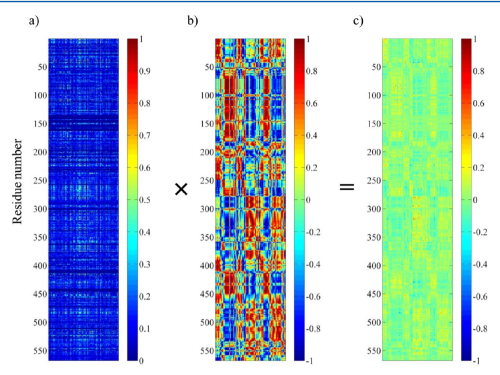


Figure 3. Generation of the co-evolutionary dynamic coupling (CDC) matrix from the SCA coupling matrix and truncated dynamic cross-correlation matrix. (a) Unclustered SCA coupling matrix from SCA of the ProRS family. The *X*-axis represents the 152 perturbation sites, while the *Y*-axis corresponds to residues 1–567 of Ec ProRS. The color gradient, as indicated in the color bar, is as follows: blue squares for the lowest and red squares for the highest statistical coupling energies,  $\Delta\Delta G_{ij}^{\text{stat}}$ . (b) Truncated MD cross-correlation matrix generated by taking only those columns of perturbation sites (residues) that are present in the SCA matrix. The cross-correlation values range from 1 (correlated) to -1 (anticorrelated). (c) The SCA-MD plot obtained as the CDC matrix by multiplying individual elements of the SCA matrix with the corresponding elements of the truncated MD matrix. Values range from 1.0 (co-evolved and thermally correlated) to -1.0 (co-evolved and thermally anticorrelated).

Table 2. Probable Pathways of Communication between the Aminoacylation and INS Domains in Ec ProRS As Identified Using  ${\rm STCA}^a$ 

pathway	parameters	residue networks	cost (Å)
C443 → K279			
I (8 residues)	$C_{ij} \ge 0.8$ $\Delta G_i^{\text{stat}} = 0.5$ $\text{CDC}_{ij} = 0.4$	$C443 \rightarrow \underline{F415} \rightarrow T199 \rightarrow \underline{D198} \rightarrow \underline{E234} \rightarrow N389 \rightarrow \underline{H302} \rightarrow K279$	43.9 (0.62, 0.51, 60)
	$D_{ii}^{o} = 8.0$		
II (9 residues)	$C_{ij} \ge 0.8$ $\Delta G_i^{\text{stat}} = 0.6$ $CDC_{ij} = 0.4$ $D_{ij}^{\text{o}} = 8.0$	$C443 \rightarrow \underline{F415} \rightarrow T199 \rightarrow \underline{D198} \rightarrow \underline{E234} \rightarrow \underline{N305} \rightarrow L304 \rightarrow L281 \rightarrow K279$	46.6 (0.66, 0.59, 0.62)
R450 → K279	211 010		
III (10 residues)	$\Delta G_i^{\text{stat}} = 0.6$ $CDC_{ij} = 0.4$	$\mathbf{R450} \rightarrow \mathbf{V411} \rightarrow \underline{\mathbf{G412}} \rightarrow \mathbf{E209} \rightarrow \underline{\mathbf{D198}} \rightarrow \underline{\mathbf{E234}} \rightarrow \underline{\mathbf{N305}} \rightarrow \mathbf{L304} \rightarrow \mathbf{L281} \rightarrow \mathbf{K279}$	51.5 (0.47, 0.60, 0.63)
( )	$D_{ij}^{o} = 8.0$		/
IV (10 residues)	$\Delta G_i^{\text{stat}} = 0.5$	$\mathbf{R450} \rightarrow \mathbf{I409} \rightarrow \underline{\mathbf{E218}} \rightarrow \mathbf{D219} \rightarrow \mathbf{N232} \rightarrow \underline{\mathbf{E234}} \rightarrow \underline{\mathbf{N305}} \rightarrow \mathbf{L304} \rightarrow \mathbf{L281} \rightarrow \mathbf{K279}$	53.3 (0.74, 0.65, 0.64)
	$\Delta G_i^{\text{stat}} = 0.5$ $CDC_{ij} = 0.4$ $D_{ij}^{\text{o}} = 8.0$		

<sup>&</sup>quot;Abbreviations:  $C_{ij}$  correlation coefficient of residue—residue fluctuations;  $\Delta G_i^{\rm stat}$ , evolutionary conservation constant;  $\Delta \Delta G_{ij}^{\rm stat}$  evolutionary coupling constant;  $CDC_{ij} = \Delta \Delta G_{ij}^{\rm stat} \times C_{ij}$ , co-evolutionary dynamic coupling constant;  $D_{ij}^{\rm o}$  distance cutoff. Numbers in parentheses in column 4 represent average  $C_{ij}$  values between adjacent residues in a given path obtained from dynamic cross-correlation matrix C for the three replica simulations. Residues shown in bold represent the terminal residues. All residues except those shown in italics are conserved residues; residues shown in italics represent the co-evolved residues. The underlined residues were chosen for mutational studies.

was reduced by ~50-fold in the E218A variant. <sup>16</sup> Thus, the lack of activity of the path IV double mutants is not surprising. No conclusions can be drawn regarding energetic coupling between

E218A and N305A because the combined  $\Delta\Delta G$  (5 kcal/mol) for the two single mutants is beyond the detection limit of this assay and no activity was detected for the double mutant.

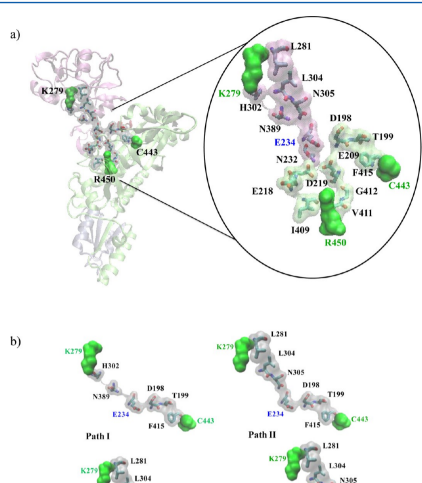


Figure 4. (a) Representation of residue—residue interaction networks between the substrate binding domain (R450/C443) and the INS (K279) identified in this study. Terminal residues in these pathways are shown in green space-filling surface representation. (b) Paths I—IV are shown in space-filling surface representation.

Path IV

Table 3. Kinetic Parameters for Amino Acid Activation by WT and Mutant Variants of Ec ProRSa

Path III

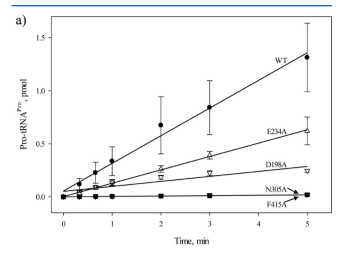
Ec ProRS	$k_{\rm cat}~({ m s}^{-1})$	$K_{\mathrm{M}}$ (mM)	$k_{\rm cat}/K_{\rm M}~({\rm mM}^{-1}~{\rm s}^{-1})$	$k_{\rm cat}/K_{ m M}$ (relative)	fold decrease	$\Delta\Delta G$ (kcal/mol)
WT	$12.6 \pm 4.9$	$0.18 \pm 0.03$	71	1.0	_	_
D198A	$6.98 \pm 0.37$	$0.33 \pm 0.01$	21	0.30	3.4	0.75
$E218A^b$	$4.4 \pm 2.3$	$3.40 \pm 0.68$	1.3	0.02	55	2.5
E234A	$6.7 \pm 1.9$	$1.03 \pm 0.25$	6.5	0.09	11	1.5
H302A	$7.3 \pm 1.9$	$0.22 \pm 0.04$	33	0.46	2.1	0.46
N305A	$0.61 \pm 0.18$	$0.45 \pm 0.18$	1.4	$2.0 \times 10^{-2}$	51	2.4
G412A	$12.8 \pm 0.57$	$0.300 \pm 0.003$	43.0	0.61	1.6	0.29
F415A	$0.131 \pm 0.010$	$0.76 \pm 0.29$	0.17	$2.4 \times 10^{-3}$	400	3.7
H302A/G412A	$10.7 \pm 0.80$	$0.62 \pm 0.26$	17	0.24	4. 2	0.88
N305A/G412A	ND	ND	ND	_	_	_
E218A/N305A	ND	ND	ND	_	_	_

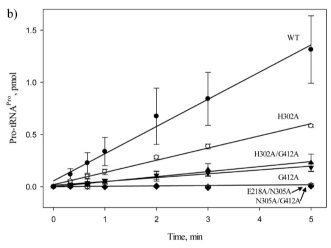
<sup>&</sup>quot;Results are the average of three trials with the standard deviation indicated.  $\Delta\Delta G$  was calculated according to the equation  $\Delta\Delta G = -RT \ln(\text{fold decrease in } k_{\text{cat}}/K_{\text{M}})$ , where R is the gas constant, 1.986 cal K<sup>-1</sup> mol<sup>-1</sup>, and T is 310 K. ND indicates not detectable under the experimental conditions used. <sup>b</sup>Data from ref 16.

**Aminoacylation.** The aminoacylation activity of single and double mutants was examined by steady-state kinetic assays

under conditions where synthesis of Pro-tRNA<sup>Pro</sup> was linear with time and the initial rate of reaction was proportional to

 $k_{\rm cat}/K_{\rm M}$ . The aminoacylation activity of F415A and N305A variants was nearly abolished with rates 70-fold slower than the WT rate (Figure 5a). The overall aminoacylation activity was





**Figure 5.** Initial rates of aminoacylation of tRNA<sup>Pro</sup> with proline by WT and single- and double-point variants of Ec ProRS. For the sake of clarity, the results are presented in two panels (a and b). The assays were performed at room temperature with 0.5  $\mu$ M tRNA<sup>Pro</sup> and 100 nM Ec ProRS. Linear fits of the data are shown.

also slightly reduced for the H302A and E234A variants (2fold) and the D198A (5.5-fold) variant (Figure 5a,b). Interestingly, mutation of G412 to alanine resulted in significantly weakened aminoacylation capability for both the single (G412A, 7-fold) and double (H302A/G412A, 5.5-fold) mutants (Figure 5b), although an only ~2-4-fold reduction in proline activation efficiency was observed for these variants, which were weakly coupled (Table 3). When  $\Delta \Delta G$  is calculated for the aminoacylation reaction (Table 4), strong coupling is observed between H302 and G412 because the  $\Delta\Delta G$  for double mutant H302A/G412A (1.0 kcal/mol) is significantly smaller (subadditive) than the sum of the two single mutants H302A and G412A ( $\Delta\Delta G = 0.4 \text{ kcal/mol} + 1.2 \text{ kcal/mol} = 1.6$ kcal/mol). Aminoacylation activity was not detected for double mutants N305A/G412A and E218A/N305A (Figure 5). On the basis of the detection limit of the aminoacylation assays, no conclusions can be drawn regarding the coupling of these residues. Taken together, these observations suggest that

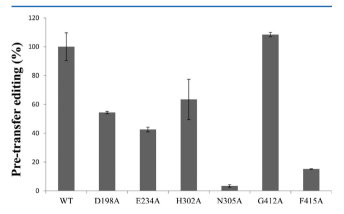
Table 4. Kinetic Parameters for Aminoacylation by WT and Mutant Variants of Ec ProRS<sup>a</sup>

Ec ProRS	$k_{\rm cat}/K_{ m M}$ (relative)	fold decrease	$\Delta\Delta G$ (kcal/mol)
WT	1	1	_
D198A	0.18	5.5	1.0
$E218A^b$	0.71	1.4	0.20
E234A	0.48	2.1	0.43
H302A	0.45	2.2	0.46
N305A	0.014	70	2.5
G412A	0.14	7.1	1.2
F415A	0.014	70	2.5
H302A/G412A	0.18	5.5	1.0
N305A/G412A	ND	_	_
E218A/N305A	ND	_	_

<sup>a</sup>Aminoacylation assays were performed under conditions where the initial rates were proportional to RNA concentrations. This indicated that  $V_0/[S]$  was an accurate reflection of  $k_{\rm cat}/K_{\rm M}$ . The relative  $k_{\rm cat}/K_{\rm M}$  values were normalized and set to 1 for WT.  $\Delta\Delta G$  was calculated according to the equation  $\Delta\Delta G = -RT \ln({\rm fold\ decrease\ in\ }k_{\rm cat}/K_{\rm M})$ , where R is the gas constant, 1.986 cal K<sup>-1</sup> mol<sup>-1</sup>, and T is 298 K. ND indicates not detectable under the experimental conditions used. <sup>b</sup>Data from ref 16.

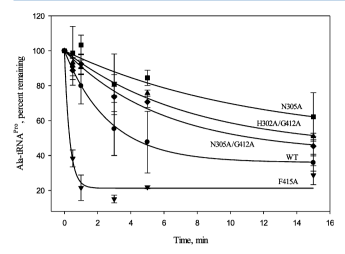
mutation of putative pathway residues has distinct impacts on the two steps of the tRNA aminoacylation reaction.

**Pre-Transfer Editing.** The pre-transfer editing reaction was studied by monitoring enhanced hydrolysis of ATP in the presence of alanine <sup>13</sup> relative to that observed in the absence of the amino acid. Reduced pre-transfer editing activity was observed for all mutants except G412A, which demonstrated levels of pre-transfer editing similar to that of WT ProRS (Figure 6). This result may be due to the variant's ability to activate alanine slightly more efficiently than the WT enzyme (data not shown).



**Figure 6.** Relative alanine pre-transfer editing activity of WT and mutant variants of Ec ProRS. The assay was performed at 37  $^{\circ}$ C using 4  $\mu$ M ProRS and 500 mM alanine. Results are reported as percent activity relative to WT, which was set to 100%.

**Post-Transfer Editing.** The impact of mutations on AlatRNA<sup>Pro</sup> hydrolysis was also studied (Figure 7). A significant reduction in post-transfer activity was observed for the N305A mutant (~4-fold reduced relative to that of WT). This residue resides at the interface between the INS and activation domains, far from the editing active site. Most of the other mutant ProRS variants tested (D198A, E234A, H302A, G412A, and E218A/N305A) displayed editing activity that was similar to that of WT ProRS (data not shown). The H302A/G412A

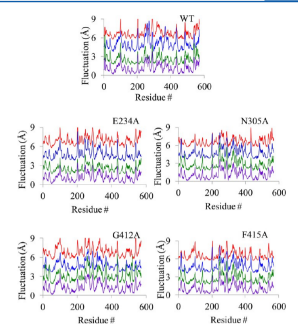


**Figure 7.** Deacylation of Ala-tRNA<sup>Pro</sup> by WT and mutant variants of Ec ProRS. The assays were performed at room temperature with 1  $\mu$ M G1:C72/U70 [ $^{14}$ C]Ala-tRNA<sup>Pro</sup> and 0.5  $\mu$ M Ec ProRS.

and N305A/G412A double mutants exhibited small ( $\sim$ 3- and  $\sim$ 2-fold, respectively) decreases in activity. Interestingly, the F415A mutant, which was severely defective in proline activation and aminoacylation, hydrolyzed mischarged AlatRNA<sup>Pro</sup> at a rate nearly 9-fold greater than that of the WT enzyme (Figure 7).

Protein Flexibility and PBL Dynamics. To examine if the alteration in enzyme activity observed for the ProRS variants correlates with a change in protein dynamics caused by mutation of on-pathway residues, we analyzed the backbone flexibility and collective dynamics of these systems. This was accomplished by using MD simulations of the four selective substrate-free mutants, namely, E234A, N305A, G412A, and F415A, for which significant changes in catalysis were observed. In each case, three replicates of 25 ns MD simulations were conducted. To assess if each system has reached an equilibrated state, the rmsd of each frame of the MD simulation trajectory was computed from their respective initial coordinates for all four mutants (Figure 2). The rmsd values were observed to fluctuate within 2.0 Å during the production period (final 20 ns) of the simulations, indicating stability for each system. Also, the quality of the simulations was assessed by computing the rms fluctuation of each amino acid from the time-averaged structure using the last 20 ns of MD simulation data. The rms fluctuations for each replica, as well as the replica-averaged fluctuations of the WT and the four ProRS variants, are reported in Figure 8. Analysis of these rms fluctuation data indicates that the backbone flexibilities are quite reproducible for each of these protein systems, with only a propagated uncertainty of 0.2-0.3 Å for the three replica simulations (Figure 8). These results indicate that all simulations have reached equilibrated states.

Next, we probed the impact of these mutations on the collective dynamics. For each system, PCA was conducted using the combined 60 ns trajectory (last 20 ns of three replicates) and the first three clusters representing the predominant collective dynamics were extracted. The rms fluctuations of  $C_{\alpha}$  atoms were computed from their respective average structures, normalized, and averaged over the three clusters. The impact of mutation of on-pathway residues on the protein flexibility was examined by computing the difference of these cluster-averaged rms fluctuations between the WT and a



**Figure 8.** Root-mean-square fluctuations of individual amino acids for WT ProRS and four variants. In each stacked plot, the rms fluctuations of  $C_{\alpha}$  atoms calculated from the time-averaged structures over the last 20 ns of MD trajectories are shown. Fluctuations for the three replicas are separated by 2 Å and color-coded for the sake of clarity: red for replica 1, blue for replica 2, and green for replica 3. In each case, the bottom plot (purple) represents the replica-averaged rms fluctuations. The calculated propagated uncertainties are 0.3 Å for WT, 0.2 Å for E234A, 0.2 Å for N305A, 0.3 Å for G412A, and 0.3 Å for F415A.

specific mutant (Figure 9). Analysis of the results indicates that the flexibility of the protein backbone was altered by varied extents because of the alanine substitutions at these sites. The overall flexibility of the catalytic-domain residues, including the PBL (residues 199–206), was found to be considerably impacted by the alanine substitution of N305 (Figure 9), which is located in the editing domain and has been observed to have significantly reduced proline activation and aminoacylation activities. The fluctuations of the editing-domain residues were also impacted in all four mutants, with the most significant changes observed for N305A, G412A, and F415A. A noticeable alteration in editing activity was also observed for these mutants, indicating that coupled thermal motions are important for Ec ProRS function.

The PBL is critical for substrate binding and catalysis. Therefore, to examine the impact of these point mutations on the collective dynamics of the PBL, we conducted PCA of WT Ec ProRS and the same four Ec ProRS variants (F415A, G412A, E234A, and N305A) using the last 20 ns of MD simulation data for the  $C_{\alpha}$  atoms of the PBL. The root-meansquare projections (rmsps)16 of the PBL for the WT and mutant proteins are shown in Figure 10. In this study, the rmsp (averaged over three replicate simulations) represents the collective displacement of the loop along principal components 1-10 (i.e., PC1-PC10, respectively), which are  $C_{\alpha}$  eigenvectors. The PCA shows that each mutation had an impact on the collective dynamics of the PBL. The displacement of the PBL along PC1 is significantly altered for all the mutant proteins compared to the WT enzyme. However, the error in PBL displacement along PC1 for E234A is large, making it difficult to ascertain the precise effect of this mutation on the PBL dynamics, and further studies are required. Noticeable changes

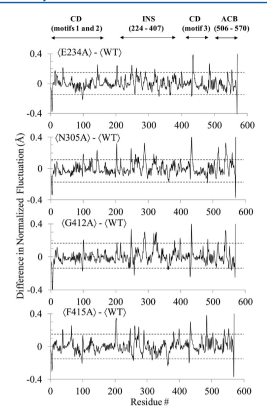
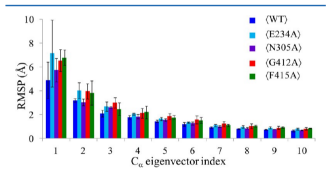


Figure 9. Changes in the normalized rms fluctuations of  $C_{\alpha}$  atoms observed in the collective dynamics of the four mutants with respect to WT Ec ProRS. The angular bracket indicates that the rms fluctuations are averaged over the first three clusters representing the predominant collective protein motions. The propagated uncertainties for each of these plots are within 0.15 Å and are shown with two parallel dotted lines. Abbreviations: CD, catalytic (aminoacylation) domain; INS, insertion domain; ACB, anticodon binding domain.



**Figure 10.** Analysis of root-mean-square projections (rmsp) of the PBL  $C_{\alpha}$  atoms for WT ProRS and the four variant proteins. The angular bracket indicates that the rmsp values are averaged over the three replica simulations. These average rmsps and standard deviations for eigenvectors 1–10 are shown.

in PBL displacement were also observed for PC2 and PC3. Therefore, this analysis indicates that mutation of the pathway residues could impact PBL dynamics and potentially alter substrate binding.

## DISCUSSION

Protein Dynamics and Site-to-Site Communication. A protein's internal dynamics is critical for many important biochemical processes ranging from catalysis to allostery. Various studies have demonstrated that coupling of dynamics

between domains is a prerequisite to the coordination of biological events occurring in distant sites. 17-20,46-48 It has been reported that correlated backbone motion provides pathways for transfer of structural and dynamics information.<sup>36</sup> Alternatively, long-range intraprotein communication can be transmitted by correlated side chain fluctuations.<sup>35</sup> These studies suggest that both local interactions and global dynamics are critical for long-range communication. Moreover, evolutionary studies of myosin motor protein and other proteins systems, including G protein-coupled receptors and hemoglobin, have demonstrated that conserved residues are key contributors to allosteric communication. 21,22,49 Recently. nuclear magnetic resonance experiments have provided information about the dynamic processes through which the KIX domain of the CREB binding protein communicates allosteric information. 50 This study also revealed that the information is transmitted through an evolutionarily conserved network of residues. Similarly, we showed that coupled-domain motions are mediated by conserved and evolutionarily coupled residues in Tt LeuRS. 20 Recent experimental 42,51 and computational<sup>20,44,52</sup> studies of other AARS systems have also revealed the existence of residue-residue interaction networks that promote interdomain communication.

An earlier study using principal component analysis of the trajectory of MD simulations of prokaryotic-like ProRSs demonstrated that the INS is engaged in coupled motion with various structural elements of the aminoacylation domain, including the PBL. 16 The collective dynamics of the PBL was found to be altered by the deletion of the INS or point mutation at the INS-aminoacylation-domain junction (vide infra). 16 Because the PBL undergoes a large-scale conformational transition upon substrate binding, 15 we sought to understand the molecular mechanism by which the PBL dynamics is modulated by distant protein segments. The STCA method was specifically applied to identify pre-existing residue-residue interactions between the INS and aminoacylation domains of Ec ProRS that could modulate PBL dynamics. Four probable pathways of communication between the aminoacylation domain (C443/R450) and INS (K279) (Table 2 and Figure 4) were identified. Residues involved in these pathways were found to be predominantly conserved and polar (Table 2). In addition, the majority of pathway residues are noncatalytic, suggesting that these residues are conserved in the ProRS family to maintain functional dynamics.

Enzymatic Function and Mutation. The impact of the alanine substitution of pathway residues on amino acid activation is significant (Table 3). Earlier studies showed that mutation of two separate pathway residues, E218 (path IV) and C443 (paths I and II), resulted in activation efficiencies significantly lower than those of WT ProRS. 16,38 Interestingly, mutation of residues E234 and N305, which are located farther from the aminoacylation active site, also impacts amino acid activation. For example, a 50-fold decrease in amino acid activation efficiency was observed for the N305A variant compared to that of the WT enzyme. In general, the mutations that resulted in decreased amino acid activation efficiency also resulted in similar reductions in aminoacylation efficiency, as well as in pre- and post-transfer editing activities, which strongly suggests that the predicted pathways of interdomain communication are important for substrate binding and catalysis. In one case, a catalytic-domain mutant (F415A) severely defective in amino acid activation, aminoacylation, and pre-transfer editing, which are all believed to occur in the

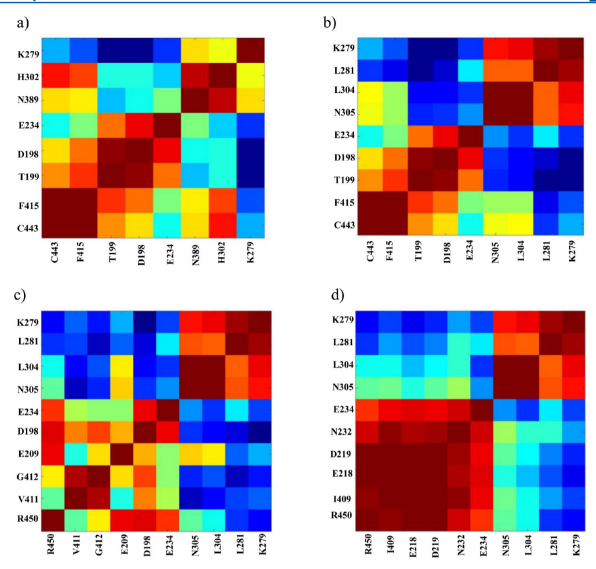


Figure 11. Coupling of thermal motions between residues in various pathways. Cross-correlations in fluctuation between residue pairs in (a) path I, (b) path II, (c) path III, and (d) path IV, extracted from the C matrix of the MD simulation. A strongly correlated motion between residues is colored red, whereas strongly anticorrelated motions are colored blue.

aminoacylation active site, was more active than WT ProRS in post-transfer editing, an activity that resides in the INS domain. This suggests that F415 is coupled to the INS. Moreover, mutations of residues at the INS—aminoacylation domain interface (N305A and E234A) have a greater impact on overall enzyme function, consistent with interdomain communication occurring between the INS and aminoacylation domain through their interface region.

Amino acid activation results for the N305A/G412A double mutant ( $\Delta\Delta G > 3.7$ , based on the level of detection) also support coupling between pathway residues even though they are located in distant domains. This mutant is totally inactive in amino acid activation, an effect that is more severe than predicted on the basis of the single mutations ( $\Delta\Delta G = 2.5$  kcal/mol + 0.31 kcal/mol = 2.81 kcal/mol). Moreover, the aminoacylation activity of this double mutant was completely abolished. Note that this double mutant is active in hydrolyzing Ala-tRNA<sup>Pro</sup>; only a 2-fold reduction in post-transfer editing activity was observed. This study also revealed cross-talk between the predicted pathways. Free energy analysis for the H302A (path I)/G412A (path III) double mutant for the

aminoacylation reaction indicates subadditivity suggesting coupling between H302 and G412, which implies communication between pathways. Close scrutiny of the predicted pathways showed that they are intertwined through residue E234, a component of all four of the predicted pathways. This observation suggests that E234, which is located in a bottleneck position of the  $\beta$ -strand connecting the catalytic and INS domains, may serve as a key hinge in site-to-site communication

Coupled Dynamics between the Two Functional Sites. If the identified residue networks facilitate coupled-domain dynamics, then a mutation along these predicted paths is expected to have an observable impact on the protein dynamics. This hypothesis was supported by the rms fluctuation analysis of the WT and mutant ProRS variants, which demonstrates that alanine substitution of on-pathway residues has a considerable impact on the flexibility of various domains of Ec ProRS (Figure 9). In addition, the collective dynamics of the PBL residues were also impacted by these mutations (Figure 10). Thus, noncatalytic residues present in the communication pathways are indeed important for

propagating substrate-induced conformational transitions from the active site pocket to the catalytically important PBL. Taken together, the *in silico* mutational study supports the role of the predicted interaction networks in maintaining distant coupleddomain dynamics.

In addition, analysis of cross-correlation coefficients between adjacent pairs of residues in each pathway revealed an interesting pattern of dynamic correlations (Figure 11). Adjacent residues in a pathway are primarily engaged in correlated motion; however, two distinct clusters of residues are observed. Residues within each cluster are engaged in strong correlated motion about the hinge residue (E234) between the INS and the aminoacylation domain. This observation is significant as it portrays a unique feature of coupled-domain dynamics in Ec ProRS; although correlated motions exist within each cluster, the coupling of these motions ultimately results in anticorrelated dynamics between the two domains (Figure S1 of the Supporting Information). The existence of anticorrelated motion between the INS and the aminoacylation domain apparently modulates the PBL conformational transition from the open to closed state that is required for substrate binding and for protecting the Pro-AMP from hydrolysis.

Proteins are dynamical in nature; mutation of any residue will have some impact on protein dynamics and may affect its function. Although mutation of catalytic residues is expected to impact an enzyme's function, in this study, we found that sitedirected changes of noncatalytic residues that are part of key communication pathways also affect enzyme function. Most of these residues in the predicted pathways not only impact the function of the domain in which they are located but also affect the distant domain with which they are dynamically coupled. For example, alanine substitution of F415 of the aminoacylation domain has a significant impact on the editing reaction. Similarly, the N305A mutation (INS domain residue) exhibited significantly reduced efficiencies in proline activation and aminoacylation of tRNA Pro. On the other hand, residues that do not belong to the predicted communication pathways are expected to have little impact on the function of a distant domain. Previous experimental results showed that alanine substitution of several residues of the INS domain (viz., T257, H369, and D386) resulted in an only minor change in aminoacylation activity (≤2.5-fold decrease) and proline activation remained unaffected by these mutations.<sup>7</sup> These three residues are not members of the predicted pathways. Analysis of the dynamic cross-correlation matrix indeed revealed low average correlation coefficients (-0.15 to -0.29) of these residues with respect to those of residues in the predicted pathways. In summary, coupled motions between INS and the aminoacylation domain are maintained by some defined residue-residue interaction networks and play important roles in both aminoacylation and editing functions.

## CONCLUSIONS

There are two proposed models<sup>53</sup> for long-range site-to-site communication: the "induced-fit" model in which a substrate-induced conformational change is propagated through a single pathway of residue—residue interactions and the "population-shift" model according to which a perturbation at a distant site alters the conformational equilibrium through multiple "preexisting" pathways of residue—residue interactions. Separate experimental and computational studies have suggested that allosteric signal propagation is mediated by a network of coupled residues and could be regulated by enthalpic

(conformational) and/or entropic (dynamic) changes.<sup>53–57</sup> In this study, an effort was made to identify the preexisting network of residue—residue interactions that could facilitate substrate-induced conformational changes from one site to another in Ec ProRS through correlated motion.

It is well established that the structural and dynamic information of a protein is encoded in its primary sequence. Therefore, the substrate-free structure can be used to extract information regarding dynamically coupled residues. Moreover, amino acid residues that are important for structure, function, and dynamics should be conserved or have co-evolved resulting in a functionally more efficient enzyme. Therefore, integration of evolutionary and dynamic information is essential for identifying residues important for maintaining coupled dynamics and, thereby, function of proteins. The STCA method takes into account both dynamic and evolutionary information in predicting preexisting pathways of interdomain communication in multidomain proteins like Ec ProRS.

Results from the STCA analysis suggest that Ec ProRS employs multiple pathways of communication between the INS and aminoacylation domains to modulate the catalytically important PBL dynamics. A significant number of residues participating in interdomain communication are evolutionarily conserved. Altering these conserved residues has a considerable impact on enzyme function. Moreover, in silico mutation followed by principal component analyses suggested that residues on the predicted pathways of communication alter protein dynamics. Taken together, this study has identified residues that are likely to be involved in maintaining the coupled dynamics between functional domains of Ec ProRS. Moreover, this study supports the notion that to facilitate longrange site-to-site communication, multidomain proteins like Ec ProRS use parallel paths of residue-residue interactions. Additional mutational studies are underway to explore the relative importance of these pathways.

# ASSOCIATED CONTENT

# S Supporting Information

Dynamic cross-correlation matrix of WT Ec ProRS obtained from the PCA-based cluster analysis, a normalized plot of  $\Delta G_i^{\rm stat}$  versus residue number obtained from the statistical coupling analysis, and a table containing the list of primers used in this study. This material is available free of charge via the Internet at http://pubs.acs.org.

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#### Notes

The authors declare no competing financial interest.

## ABBREVIATIONS

AARS, aminoacyl-tRNA synthetase; Ec, E. coli; INS, insertion domain; MD, molecular dynamics; PBL, proline-binding loop; PCA, principal component analysis; Pro-AMP, prolyl-adenylate; ProRS, prolyl-tRNA synthetase; rms, root-mean-square; rmsd, root-mean-square deviation; rmsp, root-mean-square projection; SCA, statistical coupling analysis; STCA, statistical thermal coupling analysis; Tt LeuRS, T. thermophilus leucyl-tRNA synthetase; WT, wild-type.

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