Fine-tuning Interaction between Aminoacyl-tRNA Synthetase and tRNA for Efficient Synthesis of Proteins Containing Unnatural Amino Acids

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Fine-tuning Interaction between Aminoacyl-tRNA Synthetase and tRNA for Efficient Synthesis of Proteins Containing Unnatural Amino Acids

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Supporting Information

ABSTRACT: By using a directed evolution approach, we have identified aminoacyl-tRNA synthetase variants with significantly enhanced activity for the incorporation of unnatural amino acids into proteins in response to the amber nonsense codon in bacteria. We demonstrated that the optimization of anticodon recognition of tRNA by aminoacyl-tRNA synthetase led to improved incorporation efficiency that is unnatural amino acid-specific. The findings will facilitate the creation of an optimized system for the genetic incorporation of unnatural amino acids in bacteria.

KEYWORDS: genetic code expansion, anticodon recognition, unnatural amino acid, aminoacyl-tRNA synthetase engineering, amber suppression

Orthogonal tRNA-aminoacyl-tRNA synthetase pairs are widely used for the site-specific incorporation of nearly 80 unnatural amino acids (unAAs) in Escherichia coli, Saccharomyces cerevisiae, plant, and mammalian cells in response to unique nonsense (e.g., amber) and frameshift (e.g., quadruplet) codons. This methodology enables site-specific introductions of unique chemical or physical probes into proteins, which could facilitate the study of protein structure and function as well as the investigation of biological processes.\(^1\)\(^6\)\(^8\) Recently, nonsense codon suppression-mediated regulation of biological events were also explored in synthetic biology applications.\(^9\)\(^11\) Although unAAs are typically incorporated into proteins in response to nonsense codon with good efficiency and excellent fidelity, further system optimization to increase the incorporation efficiency is still highly desirable. This is because inefficient incorporation not only results in a low yield of the desired unAA-containing protein but also leads to an increased accumulation of truncated protein products that may negatively affect the function of host cells.

An engineered Methanocaldococcus jannaschii amber suppressor tyrosyl-tRNA (MjTyrRS) and tyrosyl-tRNA-synthetase (MjTyrRS) pair is the most extensively used system for the evolution of aminoacyl-tRNA synthetase (aaRS) variants that incorporate unAAs with aromatic functional groups in E. coli. In a previous effort, optimization of the interaction between MjTyrRS and the E. coli native translational machinery, such as EF-Tu,\(^1\) was explored in order to improve the incorporation efficiency. Here, we report an approach that focuses on fine-tuning anticodon recognition of MjTyrRS by MjTyrRS variants, which led to additional improvement in unAA incorporation efficiency beyond the EF-Tu strategy. The observed improvement likely resulted from more efficient aminoclylation of MjTyrRS\(^1\) by the evolved MjTyrRS mutants.

The evolved MjTyrRS variant catalyzes the aminoclylation reaction of MjTyrRS\(^1\) with a tyrosine analogue (an unAA) at the expense of an ATP. The catalytic efficiency of this reaction, which contributes to the overall efficiency of unAA incorporation, is dictated by the substrates (MjTyrRS\(^1\) and unAA) recognition of the enzyme. Previous work successfully altered the specificity of MjTyrRS toward unAAs by focusing on changing the enzyme’s amino acid recognition pocket. While the anticodon of the tRNA was changed from GUA (Figure 1) into CUA (MjtRNAGUA\(^1\)) to enable amber suppression with unAA, the substrate promiscuity of the MjTyrRS toward MjtRNAGUA\(^1\) is unnatural amino acid-specific. The findings will facilitate the creation of an optimized system for the genetic incorporation of unnatural amino acids in bacteria.

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recognition of the mutated MjtRNA_Tyr by MjTyrRS variants. We used a directed evolution approach to identify MjTyrRS mutants that led to significantly higher amber suppression efficiency than both the parent MjTyrRS variants and the reported MjTyrRS mutant.\textsuperscript{13,14} While other beneficial factors may be picked up as well in our cell growth-based selection process, the major contributor to the observed improvement is likely due to better recognition of MjtRNA_Tyr by MjTyrRS mutants. Our results also showed that the mutations for each unAA-specific MjTyrRS variant are different, which further strikes the notion that the catalytic efficiency of MjTyrRS is determined by recognitions of both substrates, the tRNA and the amino acid. We finally demonstrated that the MjTyrRS mutants obtained in this study, when paired with a previously evolved MjtRNA_Tyr (MjtRNA_Tyr-Nap1) that has optimized interaction with E. coli EF-Tu,\textsuperscript{12} further improved amber suppression efficiency.

Examining the X-ray crystal structure of the MjtRNA_Tyr-MjTyrRS complex\textsuperscript{13} reveals that the anticodon of MjtRNA_Tyr is recognized by the C-terminal domain of MjTyrRS (Figure 1). Residues Phe261 and His283 engage in stacking interaction with the base of G34 in MjtRNA_Tyr. Residue Asp286 forms two hydrogen bonds with N1 and N2 of G34. Asp286 is well conserved among the archaeal and eukaryotic TyrRSs. It was reported that mutation of Asp286 into alanine led to a 10-fold reduction of the aminoacylation rate of MjtRNA_Tyr by MjTyrRS.\textsuperscript{15} On the other hand, the Asp286Arg mutation resulted in a more efficient aminoacylation of MjtRNA_Tyr (an amber suppressor tRNA with G34C mutation). In addition to Phe261, His283, and Asp286, residue Met285 is in the close proximity to G34 and may provide additional interaction to fine-tune the anticodon recognition by MjTyrRS. We envisaged that a directed evolution approach involving mutagenesis of above residues within the anticodon recognition pocket of MjTyrRS could optimize the interaction between MjTyrRS and MjtRNA_Tyr and therefore improve the overall efficiency of unAA incorporation in response to amber nonsense codon.

To test the hypothesis, we first examined an MjTyrRS variant, AcPheRS (referred as AcPheRS-wt hereafter),\textsuperscript{16} that was evolved previously for the incorporation of 3-acetyl-L-phenylalanine (AcPhe, Figure 2A) in response to amber nonsense codon. We created an AcPheRS library in which residues Phe261, His283, Met285, and Asp286 were completely randomized. Overlapping polymerase chain reaction (PCR) was performed with synthetic oligonucleotide primers in which the randomized residues were encoded as NNK (N = A, C, T, or G; K = T or G) to generate a library with a theoretical diversity of $1.05 \times 10^6$. The quality of the library (>99% coverage) was validated by DNA sequencing. The resulting AcPheRS library was subjected to a positive selection to identify functional AcPheRS variants followed by a negative selection to remove AcPheRS variants that could charge MjtRNA_Tyr with natural amino acid as previously described.\textsuperscript{16} Briefly, the positive selection is based on resistance to chloramphenicol (Cm), which is conferred by the suppression of an amber mutation at a permissive site (Asp112) in the chloramphenicol acetyltransferase-encoding gene in the presence of MjtRNA_Tyr-AcPhe, and functional AcPheRS mutants. The negative selection uses the toxic barnase gene with amber mutations at permissive sites (Gln2TAG and Asp44TAG) and was carried out in the absence of AcPhe. The surviving AcPheRS variants from two positive and one negative rounds of selection were subsequently screened for chloramphenicol resistance level in the presence and absence of AcPhe. A few clones that survived on 150 μg/mL chloramphenicol in the presence of AcPhe and did not grow on 75 μg/mL chloramphenicol in the absence of AcPhe were identified. Among these clones, AcPheRS-8G and AcPheRS-12B displayed the fastest growth rate in the presence of chloramphenicol and the brightest GFP fluorescence (The selection plasmid, pREP,\textsuperscript{17} contains a T7 RNA polymerase gene with amber mutation at permissive site. The synthesis of full-length T7 RNA polymerase with amber suppression drives the expression of a green fluorescent protein). Another clone, AcPheRS-2B, showed higher amber suppression efficiency than that of AcPheRS-wt, but the efficiency is lower than that of AcPheRS-8G and AcPheRS-12B. We next examined the relative protein expression level of AcPheRS-wt and AcPheRS-8G (Supporting Information Figure S3) by Western blot. We did not detect any notable difference between the two, suggesting that the observed improvement in AcPhe incorporation was not a result of higher expression level of the evolved AcPheRS-8G.
mutant. We also conducted cell growth experiments and observed similar growth rates of strains harboring different AcPheRS variants (Supporting Information Figure S4A). It is therefore unlikely that the observed improvement was due to lower toxicity of the evolved AcPheRS mutants.

DNA sequencing results revealed mutation convergence at positions Phe261 and Asp286 of all three hits (Figure 2B). The Asp286 residue was mutated into neutral residues (Asn, Gln, and Leu) that have similar side chain size to that of the Asp residue in AcPheRS-wt. Residue Phe261 in all three hits was mutated into nonaromatic amino acids, which apparently reduce the stacking interaction between the synthetase and the pyrimidine base of C34 in MjtRNA\text{\textsubscript{ACUA}}. This observation indicates that the favorable stacking interaction in MjTyrRS-MjtRNA\text{\textsubscript{ACUA}} may not be essential for AcPheRS-MjtRNA\text{\textsubscript{ACUA}}, when the G34C mutation leads to the replacement of a purine base with a pyrimidine base in MjtRNA\text{\textsubscript{ACUA}}. In addition to mutations at positions Phe261 and Asp286, Met285 was mutated to smaller amino acids (Figure 2B), which is inconsistent with the notion that cytosine is a smaller base than guanine and a larger amino acid may be needed to restore the lost interaction.\textsuperscript{13,18} The possible explanation is that the amino acid residue at position 285 does not directly interact with the nucleotide but rather affect anticodon recognition through fine-tuning the local conformation of the anticodon recognition region of AcPheRS.

To determine the efficiency and the fidelity of AcPhe incorporation into proteins in E. coli, an amber mutation (TAG) was introduced at position Asp149 in a C-terminal His-tagged GFP variant (GFP149TAG). Protein expression experiments using the two most promising hits (AcPheRS-8G and AcPheRS-12B) as well as two controls (AcPheRS-wt and AcPheRS-D286R) were carried out in LB medium supplemented with and without 1 mM AcPhe. Fluorescence analysis of E. coli cultures showed that significant amount of full-length GFP protein was produced only in the presence of AcPhe for the two evolved AcPheRS variants (Figure 2C). This result indicates that the evolved AcPheRS mutants are not cross-active with any endogenous amino acids in E. coli. The incorporation fidelity of the evolved AcPheRS mutants is comparable to that of the AcPheRS-wt. Fluorescence intensities of GFP also showed that the evolved AcPheRS-8G and AcPheRS-12B mutants had significantly higher amber suppression efficiency (Figure 2C) than that of AcPheRS-wt and AcPheRS-D286R, an AcPheRS variant with the previously reported beneficial mutation (Asp286Arg).\textsuperscript{13,14} Comparing to the rational designed AcPheRS-D286R mutant, better recognition of MjtRNA\text{\textsubscript{ACUA}} by AcPheRS-8G and AcPheRS-12B was likely achieved by exploiting a much larger conformational space using the directed evolution approach. Between the two best hits, AcPheRS-8G displayed better reproducibility and lower background in AcPhe incorporation (Figure 2C). We, therefore, focused on the AcPheRS-8G hit in following studies.

We next investigated if the beneficial mutations in the evolved AcPheRS-8G could be functionally transferred to other unAA-specific MjTyrRS variants to achieve general improvements in the incorporation efficiency. We reasoned that the beneficial mutations within the anticodon recognition region are away from the amino acid-binding pocket of MjTyrRS variants and might not affect the unAA recognition by MjTyrRSs. To this end, we focused on two other MjTyrRS variants, BpaRS\textsuperscript{5} and sTyrRS\textsuperscript{21}, which were evolved previously to recognize p-benzoyl-L-phenylalanine (Bpa, Figure 3A) and sulfotyrosine (sTyr, Figure 4A), respectively. The Bpa is a useful cross-linking amino acid for the study of protein–protein interactions. The sTyr, which is a product of post-translational modification,\textsuperscript{21} is found in many secreted and membrane-bound proteins. The direct and more efficient incorporation of sTyr into proteins is useful for the investigation of its biological functions.\textsuperscript{22} Side chain structures of these two unAAs are significantly different from that of AcPhe. We constructed
BpaRS-8G and sTyrRS-8G, each of which contained beneficial mutations (Phe261Gly, His283Leu, Met285Val, and Asp286Gln) from AcPheRS-8G. Based on the expression of GFP149TAG (Supporting Information Figure S1), no expected improvements were observed. Two possible explanations could be proposed from the above result: (1) the observed improvement of unAA incorporation via the optimization of anticodon recognition by MjTyrRS is an isolated case for AcPheRS, and (2) the anticodon and unAA recognition by MjTyrRS are mutually dependent.

To investigate above possibilities, we examined if BpaRS and sTyrRS variants with improved incorporation efficiency can be obtained through the evolution of anticodon recognition pocket. Using the same approach in AcPheRS evolution, we created a BpaRS library and a sTyrRS library, where residues Phe261, His283, Met285, and Asp286 were randomized in each library. After consecutive rounds of positive and negative selections, three BpaRS hits (BpaRS-8E, BpaRS-11D, and BpaRS-11H) and two sTyrRS hits (sTyrRS-2A and sTyrRS-5A) were identified to display higher amber suppression efficiency over their parents (Figures 3 and 4). Among these, BpaRS-8E and sTyrRS-5A are the best ones for the incorporation of Bpa and sTyr, respectively. It is worth of noticing that the degree of chloramphenicol resistance (amber suppression at position 112 of chloramphenicol acetyltransferase) does not correlate very well with the intensity of GFP fluorescence (amber suppression at position 149 of GFP). While cells containing BpaRS hits can survive much higher concentrations of chloramphenicol (>250 μg/mL) than sTyrRS hits (75 μg/mL), greater fluorescence intensities were observed for sTyrRS hits (~6000–8000 au/OD600, and Figure 4C) relative to those of the BpaRS hits (~1200–1600 au/OD600) (Figure 3C). In addition, the fold improvement of sTyrRS hits over the sTyrRS-wt is apparently greater when the chloramphenicol resistance level was used to evaluate the improvement. The observation may be due to the structure/function changes of reporter proteins caused by the incorporation of a given unAA with unique physical and chemical properties. Nevertheless, the general trend of improvement is obvious regardless which data set is used for evaluation.

As shown in Figure 3B, BpaRS-11D and BpaRS-11H have converged protein sequence, which is different from that of BpaRS-8E. The two sTyrRS hits, sTyrRS-2A and sTyrRS-5A, have different sequences (Figures 4B). None of the BpaRS and sTyrRS hits has the same mutation combinations as the AcPheRS hits. The most significant difference was observed at position 286 of BpaRS hits. While all the AcPheRS and sTyrRS hits contain a neutral amino acid residue at position 286, the BpaRS hits either retained a negatively charged Asp residue or changed into a positively charged Arg residue. On the other hand, all the evolved BpaRS, sTyrRS, and AcPheRS variants have mutations at either positions 261 or 283 or both, which apparently led to reduced stacking interactions with the C34 of MjTyrRS

Figure 5. GFP expression with AcPheRS and MjtRNACUA variants. (A) GFP fluorescence assays of AcPhe incorporation by using different combinations of AcPheRS and MjtRNACUA variants. Fluorescence intensity was normalized to cell growth; (B) GFP expression yield and SDS-PAGE analysis.
that harbors MjtRNA_{UA}^{Tyr} and a chloramphenicol acetyltransferase-encoding gene with an amber mutation at position Asp112. Transforms were cultivated in LB media containing kanamycin and tetracycline. After 12 h of cultivation, cells were harvested. Based on calculation, a certain number of cells (>4.6× the size of the library) were plated on LB agar containing kanamycin (to maintain the pBK-MjTyrRS plasmid), tetracycline (to maintain the pREP plasmid), 1 mM unAA (e.g., AcPhe), and chloramphenicol (varied from 50 to 250 μg/mL). The selection plates were incubated at 37 °C for 24 h. Survived cells were pooled and the pBK-MjTyrRS plasmids were isolated.

**Negative Selection Assay.** *E. coli* GeneHogs was cotransformed with plasmids pNEG^{27} (containing MjtRNA_{UA}^{Tyr} and a barnase-encoding gene with two amber mutations at permissive sites, Gln2 and Asp44) and pBK-MjTyr RS plasmids isolated from the positive selection. Transforms were plated on LB agar containing ampicillin (to maintain the plasmid), kanamycin, and 0.2% l-arabinose (to activate the transcription of mutant barnase gene). The selection plates were incubated at 37 °C for 12 h. Survived cells were then pooled and the pBK-MjTyrRS plasmids were isolated.

**Hit Verification.** Selected numbers of single colonies from the last round of positive selection were screened by replication onto plates with varied concentrations of chloramphenicol (34, 50, 75, 100, 150, or 250 μg/mL) in the presence and absence of the appropriate unAA. Only the ones that grew in the presence of unAA and did not grow in the absence of unAA were selected for further evaluation.

**Fluorescence Analysis of Bacterial Culture.** *E. coli* GeneHogs strain harboring plasmids pBK-MjTyrRS variant and pLei-GFP-N149TAG^{12} was cultured in 5 mL LB media containing kanamycin and chloramphenicol at 37 °C. The protein expression was induced at the OD_{600 nm} of 0.6 by additions of IPTG (0.1 mM) and appropriate unAA (1 mM). Following cultivation at 37 °C for an additional 16 h, 1 mL of cell culture was collected, washed, resuspended in 1 mL of potassium phosphate buffer (50 mM, pH 7.4). The processed cell suspensions were directly used for fluorescence and cell density measurements using a Synergy H1 Hybrid plate reader (BioTek Instruments). The fluorescence of GFP (GFP_{UV}) was monitored at λ_{exc} = 390 nm and λ_{em} = 510 nm. The cell density was estimated by measuring the absorbance at 600 nm. Values of fluorescence intensity were normalized to cell growth. Reported data are the average of two or more measurements with standard deviations.

**Protein Expression and Purification.** Similar cell cultivation procedure for fluorescence analysis was applied to preparing 25 mL of *E. coli* culture for protein purification. Cells were collected by centrifugation at 5000 g and 4 °C for 15 min. Harvested cells were resuspended in lysis buffer containing potassium phosphate (20 mM, pH 7.4), NaCl (300 mM), and imidazole (10 mM). Cells were subsequently disrupted by sonication. Cellular debris was removed by centrifugation (21 000 g, 30 min, 4 °C). The cell-free lysate was applied to Ni Sepharose 6 Fast Flow resin (GE Healthcare). Protein purification followed manufacturer’s instructions. Protein concentrations were determined by Bradford assay (Bio-Rad). Purified proteins were analyzed by SDS-PAGE.
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I. Primers

1. For library construction.

   **pBK-TyrRS-Nde1-F1**
   TTGAGGAATCCCATATGGACGA

   **TyrRS-Lib1-R1**
   CAATTCCTTATTTTTAATAA

   **TyrRS-Lib1-F2**
   TTATTTAAAAATAAGGAATTGNNKCCANNKNNKTTAAAAATGCTGTAAGCTGAAG

   **TyrRS-Lib1-R2**
   GCAGACGCCTTTATCCGGCCTG

   **TyrRS-F261-R**
   TTTTCTGGCCTTTTTATGGT

   **TyrRS-F261-F**
   ACCATAAAAAGGCCAGAAAAANNKGGGTGGAGATTTGACAGTAAATA

   **pBK-wtAcF-D286R-F2**
   AAAAATAAGGAATTGCATCCAATTCGCTTTAAAAATGCTG

2. For mutation transfer

   TTGAGGAATCCCATATGGACGA
   CTCCATCCCTCCCAACAGC
   GCTGTGGAGGGATGGAG
   GCGAACGCCTTTATCCGGCCTG
**Figure S1.** GFP fluorescence assays of cells expressing BpaRS and sTyrRS variants that contain anticodon recognition region mutations from AcPheRS-8G.

(A) The structures of $p$-benzoyl-L-phenylalanine (Bpa) and sulfotyrosine (sTyr); (B) Fluorescence readings of *E. coli* GeneHogs cells expressing BpaRS and sTyrRS variants, each co-expressed with MjtRNA$_{CUA}^{\text{Tyr}}$, in the presence of 1 mM Bpa and 10 mM sTyr, respectively. Fluorescence intensity was normalized to cell growth.
Figure S2. GFP fluorescence assays of AzPhe incorporation by AcPheRS-wt and AcPheRS-8G. Fluorescence readings of *E. coli* GeneHogs cells expressing AcPheRS-wt and AcPheRS-8G, each co-expressed with MjtRNA<sub>CUA</sub><sup>Tyr</sup>, in the presence or the absence of 1 mM AzPhe. Fluorescence intensity was normalized to cell growth.
Figure S3. Western blot analysis of protein expression level. Lane 1, marker; lane 2, AcPheRS-8G; lane 3, AcPheRS-8G; lane 4, AcPheRS-wt; lane 5, AcPheRS-wt. The AcPheRS-8G and AcPheRS-wt were sub-cloned and expressed as C-terminus His6 fusion proteins. Each sample lane was normalized to the same number of cells based on OD600nm measurement. Protein expression experiments were conducted as duplicates for each protein. The protein bands were visualized immunochemically using anti-His antibody and HRP-labeled secondary antibody. No significant differences in protein expression level were observed between AcPheRS-8G and AcPheRS-wt.
Figure S4. Growth rate of *E. coli* cells that contain different *Mj*TyrRS variants. Data are the average of three experiments.

Note. Similar cell growth rates of strains harboring different AcPheRS variants were observed. It is therefore unlikely that the evolved aminoacyl-tRNA synthetase mutants mis-aminoacylate endogenous tRNAs at detectable higher level than their parent aminoacyl-tRNA synthetases. Otherwise, stronger cell growth inhibition would be observed in the presence of the evolved aminoacyl-tRNA synthetase mutants.