

Amino-Acid-Dependent Shift in tRNA Synthetase Editing Mechanisms

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Supporting Information

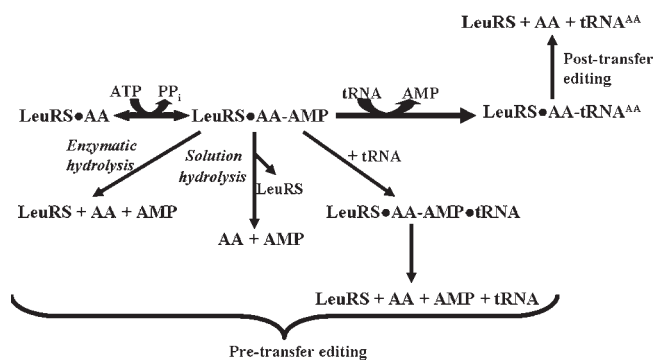
ABSTRACT: Many aminoacyl-tRNA synthetases prevent mistranslation by relying upon proofreading activities at multiple stages of the aminoacylation reaction. In leucyl-tRNA synthetase (LeuRS), editing activities that precede or are subsequent to tRNA charging have been identified. Although both are operational, either the pre- or post-transfer editing activity can predominate. Yeast cytoplasmic LeuRS (ycLeuRS) misactivates structurally similar noncognate amino acids including isoleucine and methionine. We show that ycLeuRS has a robust post-transfer editing activity that efficiently clears tRNA^{Leu} mischarged with isoleucine. In comparison, the enzyme's post-transfer hydrolytic activity against tRNA^{Leu} mischarged with methionine is weak. Rather, methionyl-adenylate is cleared robustly via an enzyme-mediated pre-transfer editing activity. We hypothesize that, similar to *E. coli* LeuRS, ycLeuRS has coexisting functional pre- and post-transfer editing activities. In the case of ycLeuRS, a shift between the two editing pathways is triggered by the identity of the noncognate amino acid.

Translational machinery relies on aminoacyl-tRNA synthetases (AARSs) to attach correct amino acids to cognate tRNAs.¹ To ensure fidelity in protein synthesis, about half of the AARSs, including leucyl-tRNA synthetase (LeuRS), have evolved editing mechanisms against structurally similar noncognate amino acids to clear mistakes that originate in the synthetic active site.² The absence of these editing mechanisms results in statistical mutations in the proteome.³

Amino acid editing may occur either before (pre-transfer)^{4,5} or after (post-transfer)⁶ the activated amino acid is transferred to the tRNA.⁷ Thus, pre-transfer editing targets the misactivated aminoacyl-adenylate intermediate (AA-AMP) for hydrolysis, while post-transfer editing cleaves mis-aminoacylated tRNA (AA-tRNA^{AA}; Scheme 1) to clear the AARSs mistakes.^{2,8} For LeuRS, the post-transfer editing active site is housed in a discrete insertion domain, called connective polypeptide 1 (CP1), that is separated from the aminoacylation active site by ~35 Å.⁹

Yeast cytoplasmic LeuRS (ycLeuRS) has been proposed to have relatively poor initial substrate discrimination¹⁰ and misactivates structurally similar or isosteric standard and nonstandard amino acids such as isoleucine, methionine, norvaline, and homocysteine.¹⁰ An early report by Englisch et al. also suggested that ycLeuRS edited predominantly by a pre-transfer editing mechanism, while *E. coli* LeuRS clears mistakes exclusively by a post-transfer editing mechanism.¹⁰ In *E. coli* LeuRS, we have shown that the CP1 editing domain that is responsible for post-transfer editing masks a pre-transfer editing activity that is associated

Scheme 1. Pre- and Post-transfer Amino Acid Editing Pathways



with the canonical aminoacylation core.^{11,12} In ycLeuRS, we hypothesize that both pre- and post-transfer editing mechanisms coexist and that one predominates, similar to *E. coli* LeuRS. Here, we determined that a shift between the two AARS fidelity mechanisms can be dependent on the identity of the standard amino acids that threaten LeuRS fidelity. A partition of fidelity mechanisms has also been reported for the human cytoplasmic LeuRS (hcLeuRS) against nonstandard biosynthetic intermediates norvaline and α -aminobutyrate.¹³ This multi-step strategy for quality control could be akin to the evolution of auxiliary editing domains such as ybak to clear mischarged Cys-tRNA^{Pro} in *trans*, in addition to Ala-tRNA^{Pro} that is produced and then cleared by *cis* editing activity of prolyl-tRNA synthetase.^{14,15}

We cloned the gene for yeast cytoplasmic tRNA^{Leu} (yctRNA^{Leu}_{CAA}) into the pTrc-99 vector¹⁶ for overexpression in *E. coli*. The yctRNA^{Leu} was extracted with phenol/tris saturated solution, followed by denaturing gel purification. ycLeuRS, with a six-histidine tag, was purified via affinity purification.¹⁷ Using *in vitro* deacylation assays that incorporated purified yctRNA^{Leu} mischarged with [³H]-isoleucine, we determined that ycLeuRS has a robust post-transfer editing mechanism that clears mischarged Ile-tRNA^{Leu} (Figure 1A). In comparison, this yeast enzyme exhibited significantly reduced hydrolytic activity against [³H]-Met-tRNA^{Leu} (Figure 1A). This amino-acid-dependent difference in post-transfer editing was surprising in that it contrasts with other LeuRSs that effectively deacylate tRNA^{Leu} mischarged with either isoleucine or methionine.^{13,18–21}

Despite its weak post-transfer editing activity, we did not observe accumulation of mischarged Met-tRNA^{Leu} for wild-type ycLeuRS (Figure 1B), as compared to an editing defective mutant

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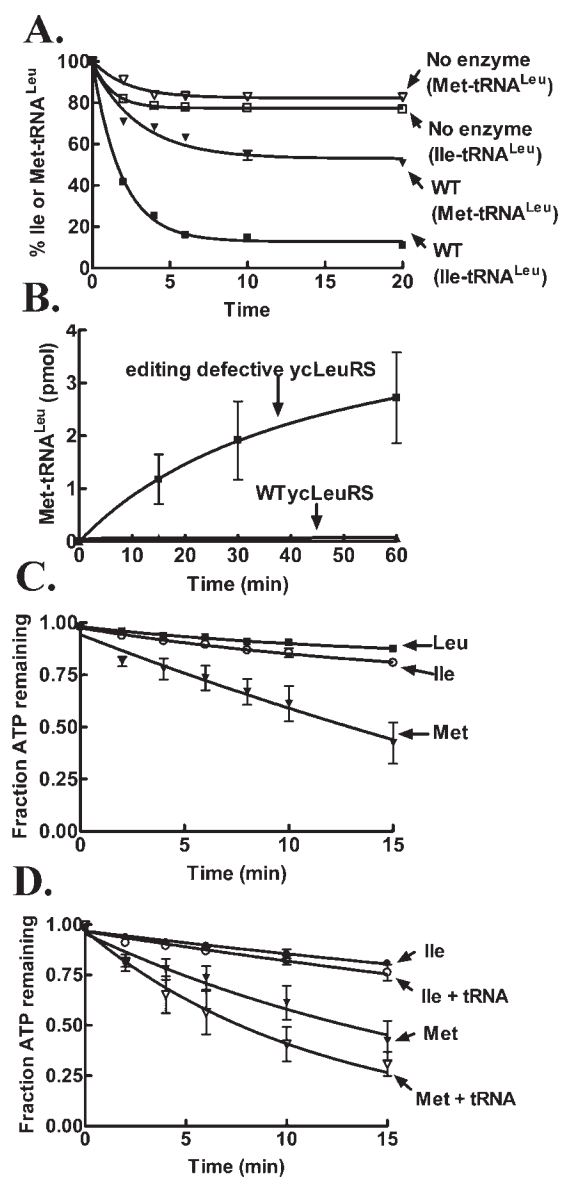


Figure 1. *In vitro* enzymatic activities of ycLeuRS. Deacylation of [^3H]-Ile-tRNA^{Leu} and [^3H]-Met-tRNA^{Leu} by 1 μM ycLeuRS (A). Mis-aminoacylation of yctRNA^{Leu} with 25 μM [^3H]-methionine (422 $\mu\text{Ci}/\text{mL}$) by 1 μM LeuRS (B). ATP hydrolysis in the absence (C) or presence (D) of tRNA. The reactions contained 250 μM ATP (600 $\mu\text{Ci}/\text{mL}$), amino acids (2.5 mM leucine or 100 mM isoleucine or methionine), and 1 μM ycLeuRS (C). Assays are described in detail in the Supporting Information.

(D419A) that we previously characterized.¹⁷ In ycLeuRS, mutation of the universal Asp419 to alanine disrupts overall editing (pre- and post-transfer).¹⁷ Thus, the mutant enzyme mischarges yctRNA^{Leu} with noncognate amino acids. It remains unclear how this single site impacts both pre- and post-transfer editing. Because of this striking difference in deacylation activities, we hypothesized that methionine that is misactivated by ycLeuRS is more efficiently cleared by an alternate fidelity pathway to post-transfer editing.

Consistent with previous activation measurements of other noncognate amino acids by ycLeuRS,¹⁰ we measured a poor discrimination factor of 1/72 for methionine (data not shown).

Table 1. Rate Constants (min^{-1}) for tRNA-Independent AARS Pre-transfer Editing Activities of Standard Amino Acids^a

	enzyme-dependent	non-enzymatic	fold difference	ref
	AMP formation	hydrolysis of AA-AMP		
<u>LeuRS</u> Met	2.0	0.100	20	this work
<u>IleRS</u> Val	3.2	0.120	30	23
<u>GlnRS</u> Gln ^b	1.5	0.042	35	24
<u>ProRS</u> Ala	27.2	0.113	240	22, 25
<u>ValRS</u> Thr	22.2	0.070	300	23

^a For enzymatic AMP formation, k_{obs} was reported for all systems, except IleRS and ValRS activation of valine and threonine, respectively, for which k_{cat} was reported. For non-enzymatic solution hydrolysis, k_{obs} was reported. ^b The GlnRS-Gln system requires presence of tRNA^{Gln}.

Since ycLeuRS lacks a robust post-transfer editing activity to clear Met-tRNA^{Leu}, we predicted that the methionyl adenylate intermediate might be cleared by pre-transfer editing. Pre-transfer editing is typically characterized by increased consumption of ATP in the presence of noncognate amino acids.^{13,22,23} We analyzed both ATP hydrolysis and AMP formation using TLC-based assays²⁴ that utilize [α - ^{32}P]-ATP to visualize separated ATP, AMP, and AA-AMP. In the absence of tRNA, ATP hydrolysis for methionine was stimulated relative to cognate leucine (Figure 1C), yielding an accumulation of 25 μM of AMP with $k_{\text{obs}} = 2.0 \pm 0.3 \text{ min}^{-1}$ for AMP formation (Table 1). By comparison, isoleucine-dependent AMP formation was only slightly elevated, with $k_{\text{obs}} = 0.7 \pm 0.1 \text{ min}^{-1}$ in the absence of tRNA.

Similar increases in AMP formation have been measured for other AARSs in the presence of noncognate amino acids as an indicator for tRNA independent pre-transfer editing activity (Table 1). The addition of tRNA enhanced ATP hydrolysis for methionine but failed to significantly stimulate isoleucine-dependent ATP hydrolysis (Figure 1D), suggesting an increase in overall editing against methionine. This is consistent with early reports¹⁰ where tRNA addition was reported to weakly stimulate pre-transfer editing of noncognate amino acids by ycLeuRS.¹⁰

Hydrolysis of AA-AMP can be catalyzed by the enzyme or depend on a selective release mechanism.^{22,25} The latter relies on enzyme ejection of adenylate intermediate into the aqueous environment for hydrolysis of noncognate AA-AMP (Scheme 1).^{2,8,22} To distinguish between these two possibilities for Met-AMP hydrolysis, we performed a chase assay²³ using reaction conditions that were identical to the AMP formation assays described above. Enzyme-synthesized AA-AMP accumulated for 10 min, followed by addition of a large molar excess of non-radioactive ATP (25 mM) to displace AA-AMP from the synthetic active site. Enzyme-independent hydrolysis of methionyl-adenylate in solution occurred at a rate of $0.1 \pm 0.001 \text{ min}^{-1}$ (Figure 2), which is 20-fold slower than the rate of methionine-dependent AMP formation by ycLeuRS (2.0 min^{-1}) (Table 1). This is consistent with other AARSs that have been proposed to edit by tRNA-independent pre-transfer editing mechanisms (Table 1). We estimate that enzyme-associated hydrolysis of the methionyl-adenylate would account then for $\sim 95\%$ of the tRNA-independent pre-transfer editing activity of ycLeuRS against methionine.

Adenylate hydrolysis by the homologous IleRS has been proposed to involve its translocation from the synthetic site to the CP1 domain editing site in a tRNA-dependent manner.²⁶

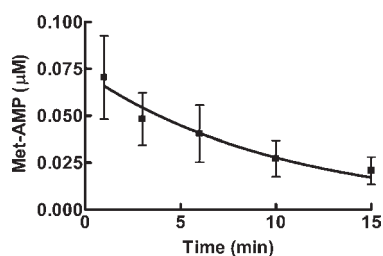


Figure 2. Non-enzymatic hydrolysis of methionyl-adenylate. TLC-based ATP chase assays include 251 μM ATP (1800 $\mu\text{Ci}/\text{mL}$), 100 mM methionine, and 5 μM ycLeuRS. AA-AMP was chased from the enzyme active site into solution by adding 100-fold excess of non-radioactive 25 mM ATP. Assays are described in the Supporting Information.

In contrast, for *E. coli* LeuRS, a robust pre-transfer editing activity was stimulated only when the CP1 domain was deleted from the enzyme¹¹ or when a mutation at CP1 domain-based Ala293 was introduced.¹² Similarly, pre-transfer editing activity is activated in ProRS²⁵ and ThrRS²⁷ mutants under circumstances where post-transfer activity was selectively inactivated. Pre-transfer editing activities that are tRNA independent have also been reported for SerRS,²⁸ which lacks a specialized editing domain, as well as in GlnRS,²⁴ which apparently does not require editing activity. In addition, MetRS clears homocysteine in a pre-transfer editing cyclization mechanism to produce thiolactone in the synthetic active site.²⁹

Segregation of amino acids to different editing pathways has been reported for hcLeuRS, which edits biosynthetic intermediates (α -amino butyrate and norvaline) under *in vitro* conditions via pre- and post-transfer hydrolysis, respectively.¹³ A tRNA-independent pre-transfer editing activity for *Aquifex aeolicus* LeuRS has also been identified to clear norvalyl-adenylate.³⁰ Our results demonstrate that pre- and post-transfer editing co-exist in ycLeuRS, as also found previously for *E. coli* LeuRS.^{11,12} Methionine-stimulated ATP hydrolysis and AMP formation in the absence of tRNA as well as ATP chase experiments support that quality control for ycLeuRS is highly dependent on pre-transfer editing under *in vitro* conditions. It is possible that the inclusion of tRNA shifts the editing mechanism preference and increases overall global editing. The same conditions that incorporate isoleucine rather than methionine suggest that the ycLeuRS relies on post-transfer editing for isoleucine clearance. Although most *in vitro* enzyme experiments fail to recapitulate the dynamic cellular environment, direct *in vitro* comparison for ycLeuRS suggests that methionine and isoleucine partition for clearance between different editing pathways. It remains possible that ycLeuRS relies on a combination of editing mechanisms, but our results support that the preference of editing pathways can be dependent on substrate identity.

We hypothesize that these two fidelity mechanisms in ycLeuRS are not redundant but are adapted to accommodate diverse specificities of the noncognate amino acids that challenge LeuRS fidelity in the cell.^{3,20} Under this scenario, ycLeuRS pre- and post-transfer editing sites would have evolved in a complementary way, similar to AARSs that are dependent on independent auxiliary editing domains. Thus, in the case of ycLeuRS, we propose that pre- and post-transfer editing pathways partition in a way that is dependent on the chemical structure of the non-cognate amino acid to ensure that all errors are efficiently targeted for clearance by ycLeuRS. This is critical to the cell since disruptions

or absence of editing activities in LeuRS can cause mistranslation and cell death.^{3,20}

■ ASSOCIATED CONTENT

S Supporting Information. Details on all materials and methods. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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