

Discrimination of Cognate and Noncognate Substrates at the Active Site of Class I Lysyl-tRNA Synthetase[†]

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ABSTRACT: The aminoacyl-tRNA synthetases are divided into two unrelated structural classes, with lysyl-tRNA synthetase (LysRS) being the only enzyme represented in both classes. On the basis of the structure of L-lysine complexed with *Pyrococcus horikoshii* class I LysRS (LysRS1) and homology to glutamyl-tRNA synthetase (GluRS), residues implicated in amino acid recognition and noncognate substrate discrimination were systematically replaced in *Borrelia burgdorferi* LysRS1. The catalytic efficiency of steady-state aminoacylation ($k_{\text{cat}}/K_{\text{M}}$) with lysine by LysRS1 variants fell by 1–4 orders of magnitude compared to that of the wild type. Disruption of putative hydrogen bonding interactions through replacement of G29, T31, and Y269 caused up to 1500-fold reductions in $k_{\text{cat}}/K_{\text{M}}$, similar to changes previously observed for comparable variants of class II LysRS (LysRS2). Replacements of W220 and H242, both of which are implicated in hydrophobic interactions with the side chain of lysine, resulted in more dramatic changes with up to 40000-fold reductions in $k_{\text{cat}}/K_{\text{M}}$ observed. This indicates that the more compact LysRS1 active site employs both electrostatic and hydrophobic interactions during lysine discrimination, explaining the ability of LysRS1 to discriminate against noncognate substrates accepted by LysRS2. Several of the LysRS1 variants were found to be more specific than the wild type with respect to noncognate amino acid recognition but less efficient in cognate aminoacylation. This indicates that LysRS1 compromises between efficient catalysis and substrate discrimination, in contrast to LysRS2 which is considerably more effective in catalysis but is less specific than its class I counterpart.

Aminoacyl-tRNAs (aa-tRNAs)¹ are synthesized when an amino acid is esterified to the 3'-end of a transfer RNA (tRNA) (1, 2). After being synthesized, aa-tRNAs are screened for their correct pairing by elongation factor TU (3) and taken to the ribosome where they base pair with the complementary mRNA and participate in protein synthesis. Correctly aminoacylated tRNAs are essential for the faithful translation of mRNA into the encoded polypeptide sequence. The aa-tRNAs are synthesized primarily by the aminoacyl-tRNA synthetases (aaRSs) (4). The accuracy of aa-tRNA synthesis is ensured by the extremely high substrate specificity of the aaRSs, and this is further enhanced in some synthetases by the existence of editing mechanisms directed against noncognate substrates (5). The 20 aaRS proteins, as for example found in *Escherichia coli*, are divided into two mutually exclusive structural groups, comprised of 10

members each, termed class I and class II (6–8). The assignment of an aaRS specific for a particular amino acid to one structural class or the other is almost completely conserved. The only widespread exception to this rule is the lysyl-tRNA synthetases (LysRSs). These are class I enzymes (LysRS1) in certain bacteria and archaea but are otherwise members of class II (LysRS2) (9–11). The class I LysRS is found in a number of pathogenic bacteria (e.g., *Borrelia*, *Treponema*, and *Rickettsia* species) and is fundamentally different from the human class II enzyme, suggesting it may be a useful antimicrobial target (12, 13).

The existence of both class I- and class II-type LysRSs indicates that exactly the same activity (lysylation of tRNA^{Lys}) has been achieved in two completely different structural frameworks by convergent evolution. The continued retention of both forms of LysRS prompted investigation of possible biochemical differences between the two enzymes. In vitro studies revealed variation in the recognition of both the anticodon and acceptor stem of tRNA^{Lys} by LysRS1 and LysRS2 (14–16), but these differences were found to be less significant in vivo (17). In contrast, structural differences between their active sites lead to significantly divergent patterns of noncognate amino acid discrimination by LysRS1 and LysRS2 both in vitro and in vivo (18). The lysine-binding pocket appears more open in LysRS2 than in LysRS1, and as a result, lysine analogues such as S-(2-aminoethyl)-L-cysteine (AEC) are efficient substrates for the class II, but not the class I, enzyme in vitro. LysRS1 imparts resistance to growth inhibition by AEC as a result of the

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¹ Abbreviations: aa-tRNA, aminoacyl-tRNA; aaRS, aminoacyl-tRNA synthetase; AEC, S-(2-aminoethyl)-L-cysteine; LysRS, lysyl-tRNA synthetase.

difference in noncognate amino acid recognition compared to LysRS2, illustrating how LysRS distribution can contribute to quality control during protein synthesis.

High-resolution crystal structures are available for both LysRS1 and LysRS2 in complexes with lysine (19, 20). Comparison of the LysRS1 and LysRS2 substrate complexes revealed that while the mechanisms of recognition of the R group of L-lysine rely on similar arrangements of amino acids in each binding pocket, the active sites are different. On the basis of the structure of L-lysine complexed with *E. coli* LysRS2 (*lysS*), residues implicated in amino acid recognition and discrimination were systematically replaced, and the resulting variants were characterized (21). This revealed that LysRS2 predominantly recognizes lysine via hydrogen bonding interactions with a series of charged residues in the active site. Manipulation of the amino acid binding site allowed an up to 4-fold improvement in AEC discrimination by LysRS2 but did not rival the highly effective discrimination of noncognate substrates by LysRS1. These findings suggest fundamentally different roles in substrate discrimination for the conserved aromatic residues found in the two LysRS active sites. To compare substrate selection and discrimination at the two different LysRS active sites, we have now investigated the role of each residue implicated in lysine binding by LysRS1. Comparisons to previous studies indicate that while both enzymes employ a network of hydrogen bonding interactions to bind lysine, LysRS1 employs an additional network of hydrophobic interactions absent in LysRS2. The impact of this extended substrate binding network on noncognate amino acid recognition suggests that it may directly account for the substrate specificity of LysRS1 being greater than that of LysRS2.

EXPERIMENTAL PROCEDURES

Bacterial Strains and Plasmids. *Borrelia burgdorferi* *lysK*-encoded LysRS1 cloned into the pET15b vector was used as a template for development of LysRS1 variants. Sets of two complementary primers with 27 nucleotides were designed to encode each desired point mutation. The LysRS1 variants used here were G29A, T31S, T31G, E43N, G43I, W220L, W220A, W220Y, H242A, H242L, H242W, Y269F, and Y269S. PCRs using *Pfu* turbo DNA polymerase (Stratagene) were performed according to the manufacturer's procedure. PCR products and the pET15b vector were digested with *Nde*I and *Bam*HI (New England BioLabs), then ligated, and transformed into DH5 α cells. Point mutations were confirmed by sequencing each gene completely. In vitro transcribed *B. burgdorferi* tRNA^{Lys} (UUU anticodon) was prepared and purified as described previously (18).

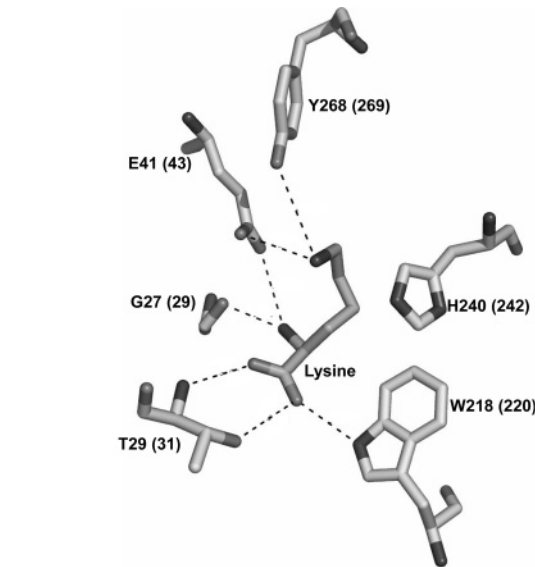
Lysyl-tRNA Synthetase Purification. The *B. burgdorferi* *lysK*-encoded LysRS1 and mutants cloned into the pET15b vector were expressed in *E. coli* BL21(DE3) cells as described previously (18). Cells were harvested by centrifugation and washed in column buffer [20 mM Tris-HCl (pH 8), 300 mM NaCl, 30 mM imidazole, 1 mM MgCl₂, and 10% glycerol]. Cells were resuspended in column buffer supplemented with protease inhibitor (Hoffman-La Roche), passed through a French pressure cell, and then centrifuged at 25000g for 30 min. The resulting supernatant was loaded onto a nickel affinity bead column (QIAGEN). Protein was eluted from the nickel affinity column in a buffer of 50 mM

Tris-HCl (pH 8), 1 mM MgCl₂, 300 mM NaCl, 250 mM imidazole, 10% glycerol, and 10 mM 2-mercaptoethanol. The fractions containing LysRS1 were pooled and then concentrated by ultrafiltration using an Amicon Ultra-15 unit (Millipore). The concentrated LysRS1 was further loaded onto a Superose 6 column (Amersham Pharmacia Biotech) and eluted in a buffer of 50 mM Hepes (pH 7.2), 25 mM KCl, 10 mM MgCl₂, 10% glycerol, and 5 mM 2-mercaptoethanol. The fractions containing LysRS1 were again concentrated by ultrafiltration and stored at -80 °C. The concentration of LysRS1 was determined by the Bradford dye binding procedure using the Bio-Rad protein assay system.

Aminoacylation Assays. Aminoacylation was performed at 37 °C in 100 mM Hepes (pH 7.2), 25 mM KCl, 10 mM MgCl₂, 5 mM DTT, 5 mM ATP, 16 μ M tRNA^{Lys}, and 50 nM to 5 μ M LysRS1. For the determination of K_M for L-Lys, all the concentrations were fixed except for that of L-[³H]-Lys (L-[U-³H]lysine, 91.0 Ci/mmol, Amersham Corp.), which was added at concentrations between 0.2- and 5-fold greater than the K_M . The same procedure was followed for tRNA and ATP K_M determinations, in which tRNA or ATP was added at concentrations between 0.2- and 5-fold greater than the K_M . Aliquots (9 or 18 μ L) were taken every 30 s to 15 min and spotted onto 3 mm filter disks presoaked in 5% TCA (w/v) containing 0.5% (w/v) L-[¹²C]Lys. Sample disks were washed, and radioactivity was counted as described previously (18). The steady-state kinetic parameters that are given represent the average of at least two independent experiments where values deviated by no more than 10% between individual determinations.

K_i Determination. To determine the K_i for S-(2-aminoethyl)-L-cysteine (AEC), at least five different concentrations of the analogue were first screened in the aminoacylation reaction for each variant under standard conditions with K_M concentrations of L-[¹⁴C]Lys for each LysRS1 variant. Analogue concentrations at which the initial rate of aminoacylation was decreased by 20–50% when compared with that of the reaction without analogue were then established, and these levels were used for K_i determinations. The K_i values presented represent the average of at least two independent experiments where values deviated by no more than 10% between individual determinations.

tRNA Aminoacylation with Noncognate Amino Acids. Synthesis of the tRNA^{Lys} [³²P]pA76 transcript was performed as described previously (22). Briefly, the CCA-3' end of the tRNA was first removed by treatment of 20 μ M tRNA transcript with 73 μ g/mL *Crotalus atrox* venom (Sigma) in a buffer containing 40 mM Na-Gly (pH 9.0) and 10 mM magnesium acetate. The mix was incubated for 2 h at 21 °C, phenol/chloroform extracted, ethanol precipitated, and finally desalted by gel filtration through a Sephadex G25 column (Pharmacia). The CCA-3' end of the tRNA was reconstituted and radiolabeled by incubation for 10 min at 37 °C with 0.5 μ M snake venom-treated tRNA, in 50 mM Na-Gly (pH 9.0), 10 mM MgCl₂, 10 μ M CTP, 9 μ M ATP, 1 μ M [α -³²P]ATP, and 3 μ g/mL *E. coli* tRNA-terminal nucleotidyltransferase. The reaction was stopped by addition of 1 volume of phenol, and the mixture was gel filtered twice through a G25 column. As described previously (23), the aminoacylation reaction was performed in 10 μ L of aminoacylation medium (see above) containing 5 mM cold



<i>B. burgdorferi</i>	LysRS1:	28	SGITPSGTVHIGNFRE
<i>P. horikoshii</i>	LysRS1:	26	SGITPSGYVHVGNFRE
<i>T. thermophilus</i>	GluRS:	6	IAPSP TGDP HVGT AYI
<i>B. burgdorferi</i>	LysRS1:	220	WRIDWPMRWKYEKVDFEPAGK
<i>P. horikoshii</i>	LysRS1:	218	WRVDWPMRWSHFGVDFEPAGK
<i>T. thermophilus</i>	GluRS:	187	YHLANVVDHLM GV TDVIRAE
<i>B. burgdorferi</i>	LysRS1:	241	DHSSGGSFDTSKNIVK-IFQGSPPVTFQY
<i>P. horikoshii</i>	LysRS1:	239	DHLVAGSSYDTGKEIIKEVYKGEAPLSLMY
<i>T. thermophilus</i>	GluRS:	208	EWLV---STPIHVLVLYRA- FGWEAP --RFY

FIGURE 1: Active site of *P. horikoshii* LysRS1. (A) L-Lysine in the active site of *P. horikoshii* LysRS1 (adapted from ref 20). Hydrogen bonds are shown as dashed lines, and the numbering of the corresponding positions in *B. burgdorferi* LysRS1 is indicated in parentheses. (B) Alignment of amino acid binding site residues from the *B. burgdorferi* and *P. horikoshii* class I LysRSs and *T. thermophilus* GluRS. Positions conserved in at least two of the sequences are shown in bold; those replaced in this study are denoted with an asterisk.

amino acids, 5 μ M transcript, and a trace of radiolabeled tRNA. After incubation for 15 min, an aliquot was removed and incubated for 30 min at room temperature with P1 RNase. The liberated [α - 32 P]AMP and aminoacyl-[α - 32 P]-AMP were separated by TLC on PEI-cellulose and visualized as described previously (23).

RESULTS

Selection and Characterization of LysRS1 Variants. The structure of the active site of *Pyrococcus horikoshii* LysRS1 implicates six residues in lysine binding (Figure 1A). Of these amino acids, W220 (*B. burgdorferi* LysRS1 numbering) is conserved in all 61 known class I LysRS proteins, while G29 and Y269 vary in only one known example each (Ala in *Nanoarchaeum equitans* and Phe in uncultured *Archaeon GZfos32E7*). H242, E43, and T31 are also well-conserved, as illustrated by the conservation of all of the corresponding residues between the *B. burgdorferi* and *P. horikoshii* proteins (Figure 1B). *B. burgdorferi* LysRS1 variants were made with the aim of testing the role of each lysine-binding residue in the active site and investigating conservation of the synonymous residues in the closely related glutamyl-tRNA synthetase (GluRS) active site (Figure 1B). The G29A

Table 1: Steady-State Aminoacylation Kinetics of Wild-Type and Variant *B. burgdorferi* LysRS1 with Lysine

LysRS	K_M (mM)	k_{cat} (s $^{-1}$)	k_{cat}/K_M (R) ^a
wild-type	0.230 \pm 0.02	0.34 \pm 0.009	1
G29A	6.3 \pm 0.50	0.45 \pm 0.1	20
T31G	4.4 \pm 0.7	0.0046 \pm 0.0003	1400
T31S	0.76 \pm 0.07	0.53 \pm 0.03	2
E43Q	Nd ^b		
E43I	Nd ^b		
W220A	0.330 \pm 0.04	0.00075 \pm 0.00004	650
W220L	3.3 \pm 0.6	0.00017 \pm 0.00001	29000
W220Y	4.7 \pm 0.4	0.13 \pm 0.05	50
H242A	3.3 \pm 0.9	0.00012 \pm 0.00001	40000
H242L	0.18 \pm 0.01	0.29 \pm 0.008	0.9
H242W	Nd ^b		
Y269F	0.40 \pm 0.04	0.099 \pm 0.003	6
Y269S	2.8 \pm 0.3	0.0026 \pm 0.0001	1600

^a Relative decrease compared to that of the wild type. ^b No detectable activity.

replacement was intended to disrupt hydrogen bonding with the α -amino group of lysine. T31S and T31G were intended to displace and disrupt, respectively, hydrogen bonding with the α -carboxyl group, E43Q and E43I to disrupt electrostatic interaction with the ϵ -amino group, and Y269F and Y269S to disrupt hydrogen bonding with the ϵ -amino group.

Previous studies and comparisons of the active sites of the other subclass Ib aaRSs, glutamyl-tRNA synthetase (GlnRS) and GluRS, revealed residues that participate in tRNA-dependent amino acid recognition. Structural (24) and biochemical (25) studies have shown that residues Y211 and F233 of *E. coli* GlnRS stack against the terminal adenosine of tRNA^{Gln} during formation of the glutamine-binding site. A conserved role in other class Ib aaRSs for the residue equivalent to Y211 of GlnRS is suggested by the strict conservation of the analogous position as Tyr in GluRS. Multiple-sequence alignments with GluRS show that the analogous position in all known class I LysRS sequences is instead occupied by Trp (W220 in *B. burgdorferi*). Positions analogous to F233 of GlnRS are also fairly well conserved, being His or Trp in GluRS and His or Leu (rarely Ile) in LysRS1 (H242 in *B. burgdorferi*). Therefore, the W220Y and H242W modifications were designed to determine whether protein–RNA stacking interactions affect lysine binding, which would be expected to result in corresponding changes in the kinetic parameters for both lysine and tRNA.

The steady-state kinetic parameters for lysine during aminoacylation were measured for the wild type and LysRS1 variants (Table 1). The catalytic efficiency (k_{cat}/K_M) for lysine compared to that of the wild type decreased significantly for most variants, the only exceptions being H242L and T31S which both showed almost no change and Y269F that displayed an only 6-fold drop in efficiency. No aminoacylation was detected using either the filter binding or 3'-labeling assay for E43Q or E43I (Table 1 and Figure 2), indicating that replacements at this position abolish activity. H242W was also inactive in filter binding assays but exhibited slight activity in alternative assays (Figure 2), indicating that this replacement severely reduces aminoacylation efficiency. The H242A replacement also strongly impacted efficiency (40000-fold reduction), while H242L caused almost no change. W220A had little effect on K_M but dramatically reduced k_{cat} (450-fold decrease), while W220Y increased K_M 20-fold but reduced k_{cat} only 3-fold.

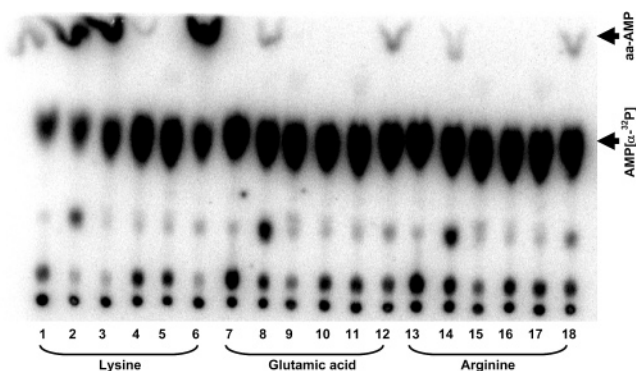


FIGURE 2: TLC analysis of tRNA^{Lys} aminoacylation with lysine and noncognate amino acids by wild-type and variant *B. burgdorferi* LysRS1: lanes 1–6, G29A, T31S, W220Y, H242W, E43I, and wild-type LysRS1, respectively, with lysine; lanes 7–12, like lanes 1–6 except with glutamic acid instead of lysine; and lanes 13–18, like lanes 1–6 except with arginine instead of lysine.

Table 2: Steady-State Aminoacylation Kinetics of Wild-Type and Variant *B. burgdorferi* LysRS1 with tRNA^{Lys}_{UUU}

LysRS	K_M (μ M)	k_{cat} (s^{-1})	k_{cat}/K_M (R) ^a
wild-type	2.0 \pm 0.1	0.33 \pm 0.005	1
G29A	5.1 \pm 0.3	0.59 \pm 0.015	1.4
T31G	2.1 \pm 0.1	0.0050 \pm 0.0001	50
T31S	2.1 \pm 0.2	0.68 \pm 0.02	0.5
E43Q	Nd ^b		
E43I	Nd ^b		
W220A	2.5 \pm 0.2	0.00056 \pm 0.00002	740
W220L	1.1 \pm 0.1	0.00012 \pm 0.000003	1500
W220Y	1.2 \pm 0.1	0.23 \pm 0.006	0.9
H242A	1.0 \pm 0.2	0.00013 \pm 0.000008	1300
H242L	1.3 \pm 0.05	0.29 \pm 0.004	0.7
H242W	Nd ^b		
Y269F	0.83 \pm 0.1	0.14 \pm 0.004	1
Y269S	1.1 \pm 0.09	0.0023 \pm 0.00006	79

^a Relative decrease compared to that of the wild type. ^b No detectable activity.

Y269F had a much weaker effect on both K_M and k_{cat} than Y269S. Y269F increased K_M 2-fold and decreased k_{cat} 3-fold, while Y269S increased K_M 10-fold and reduced k_{cat} 130-fold. Similarly, T31S had a much weaker effect on both K_M and k_{cat} than T31G. T31S slightly increased both K_M and k_{cat} (3- and 2-fold, respectively), while T31G increased K_M 20-fold and decreased k_{cat} 70-fold.

LysRS1, together with arginyl-tRNA synthetase (ArgRS), GlnRS, and GluRS, forms a small subgroup of aaRSs that requires the presence of tRNA for amino acid activation. To investigate whether the observed effects of the various replacements on lysine binding were the indirect results of changes in tRNA binding, the kinetic parameters for tRNA^{Lys} were determined (Table 2). No significant changes compared to the wild type were seen in the K_M values for tRNA, and the changes in k_{cat} were very similar to those observed for lysine. Comparison of LysRS1 to the closely related GluRS active site suggested that T31 and Y269 might also participate directly in ATP binding. The aminoacylation kinetic parameters for variants containing replacements of T31 and Y269 showed only relatively modest changes in the K_M for ATP (2–7-fold decreases, Table 3), suggesting that the main role of these residues is in lysine binding.

Substrate Discrimination by LysRS1 Variants. Previous studies indicated that LysRS1 and LysRS2 are markedly different in their abilities to discriminate lysine analogues

Table 3: Steady-State Aminoacylation Kinetics of Wild-Type and Variant *B. burgdorferi* LysRS1 with ATP

LysRS	K_M (mM)	k_{cat} (s^{-1})	k_{cat}/K_M (R) ^a
wild-type	1.3 \pm 0.4	0.57 \pm 0.09	1
T31G	9.5 \pm 2	0.0011 \pm 0.0002	3800
T31S	1.7 \pm 0.3	1.7 \pm 0.2	0.4
Y269F	1.6 \pm 0.3	0.017 \pm 0.002	40
Y269S	6.9 \pm 0.4	0.00051 \pm 0.00006	5900

^a Relative decrease compared to that of the wild type.

Table 4: Kinetic Parameters for the Inhibition of Steady-State Aminoacylation by Wild-Type and Variant *B. burgdorferi* LysRS1 in the Presence of AEC

LysRS	K_i (mM)	k_{cat} (s^{-1})	K_i/K_i^{wt}	K_M/K_M^{wt}
wild-type	3.6 \pm 0.2	0.32 \pm 0.006	1	1
W220A	19 \pm 2	0.00082 \pm 0.00003	5	1
W220L	11 \pm 1	0.00024 \pm 0.000003	3	14
W220Y	17 \pm 1	0.15 \pm 0.003	5	20
H242A	8.5 \pm 1	0.00016 \pm 0.000005	2	14
H242L	2.9 \pm 0.3	0.36 \pm 0.01	1	1
G29A	11.8 \pm 2	0.40 \pm 0.02	3	30

(18). For example, the competitive inhibitor AEC has a K_i for the aminoacylation reaction that is several 100-fold lower for LysRS2 than for LysRS1. Structural modeling suggested that this difference was in part due to steric hindrance by H242 and W220 in LysRS1, as the comparable region of the LysRS2 active site appears more open. To investigate the structural basis of lysine analogue discrimination by LysRS1, the K_i values for AEC of H242 and W220 variants were determined (Table 4). The G29A variant was also investigated in light of its stronger discrimination of canonical amino acids (see below). For all the LysRS1 variants that were tested, the k_{cat} did not change significantly in the presence of AEC, indicating that it acted as a competitive inhibitor. With the exception of LysRS1 H232L, the K_i as compared to that of the wild type increased for all the variants. In comparison to that of the wild type, the changes in apparent AEC (K_i/K_i^{wt}) and lysine (K_M/K_M^{wt}) binding showed that, with the exception of W220A and H242L, the various replacements were considerably more detrimental to cognate than noncognate substrate binding. To further investigate the relationship between cognate amino acid recognition and the discrimination of noncognate substrates, we tested the ability of LysRS1 variants to activate canonical amino acids other than lysine.

Aminoacylation of Canonical Amino Acids by LysRS1 Variants. Structural studies have previously shown that *P. horikoshii* LysRS1 bears a striking similarity to *Thermus thermophilus* GluRS that extends throughout the whole structure and includes the active site (20). Detailed comparisons of amino acid binding based on the *P. horikoshii* LysRS1 structure suggest that *B. burgdorferi* LysRS1 residues W220, H242, T31, G29, and E43 most closely correspond to Y187, W209, S9, A7, and I21, respectively, in *T. thermophilus* GluRS. The corresponding LysRS1 variants, for which the identities of the appropriate residues were changed to those observed in GluRS, were then tested for their ability to utilize the canonical noncognate amino acids glutamic acid and arginine (Figure 2). As previously observed (18), wild-type LysRS1 is able to aminoacylate tRNA^{Lys} with both glutamic acid and arginine, albeit at substantially lower levels than with lysine. However, of the

LysRS1 variants that were tested, only T31S retained the ability to utilize glutamic acid and arginine, with all the other variants exhibiting no such activity.

DISCUSSION

Specific Recognition of Lysine in the LysRS1 Active Site. While comparisons to our previous study of *E. coli* LysRS2 clearly identify residues with similar roles in the class I and II enzymes, it is also evident that the two residues characteristic of the more closed LysRS1 active site are functionally specific for the class I protein. H242 is positioned very close to bound lysine in the structure and plays an important role in hydrophobic interaction with the substrate. When the potential for this interaction was lost in LysRS1 H242A, this was accompanied by a 40000-fold drop in catalytic efficiency, while a conservative substitution that maintained hydrophobicity (H242L) had no effect. The almost complete loss of activity for H242W likely results from the larger indole moiety limiting the access of lysine to the active site. The proposed role of H242 in contributing to hydrophobic interactions while limiting access to the active site correlates with data from LysRS1 sequence alignments, where this position is conserved as either His or Leu and rarely Ile (there are 45 examples of His, 15 of Leu, and one of Ile, in the 61 known aligned LysRS1 sequences). Initial examination of the active site of LysRS1 suggested similar roles in lysine binding for H242 and W220, but biochemical analyses indicate that they differ in function. The W220L and W220Y replacements both significantly increased the K_M for lysine, but W220A had no effect, suggesting that direct interactions with the substrate are not as important at this position as at H242. The changes at W220 may instead affect the position of H242, which is only 3.2 Å away, thereby perturbing both direct interactions with lysine and stacking with A76 and C75 of the tRNA (Figure 3A). The stacking array is also important for the correct position and orientation of A76 of the tRNA into the active site (in GluRS of *T. thermophilus*, the W220 corresponds to Y187). Disruption of the correct positioning of H242 in the W220A and W220L variants severely impaired the k_{cat} , while W220Y resulted in almost no changes in catalytic efficiency. The predicted effect of W220 substitutions would be to impair tRNA binding since the correct positioning of the stacking array would be compromised. However, no variation in K_M for tRNA was observed for modifications of W220 which suggests this residue is not involved in the same direct interactions with A76 as the corresponding Y187 in GluRS. Nevertheless, further studies are now needed to determine the extent to which interactions between W220 and H242 and tRNA^{Lys} might contribute to amino acid recognition and catalysis through local conformational adjustments of the active site.

The crystal structure of *P. horikoshii* LysRS1 suggests that Y268 may be too far from the ϵ -amino group of lysine to allow strong interaction with it through hydrogen bonding. Instead, comparison to other tRNA-dependent class I aaRSs indicates that Y268 is in a hydrophobic pocket and is able to bind the adenine portion of ATP through stacking interactions (Figure 3B). Superposition of GluRS, ArgRS, and LysRS1, using the α backbones to search for similar structural motifs, shows that in ArgRS the same position is occupied by the conserved residue F402 which can also stack against ATP (Figure 3B). The decreases in catalytic ef-

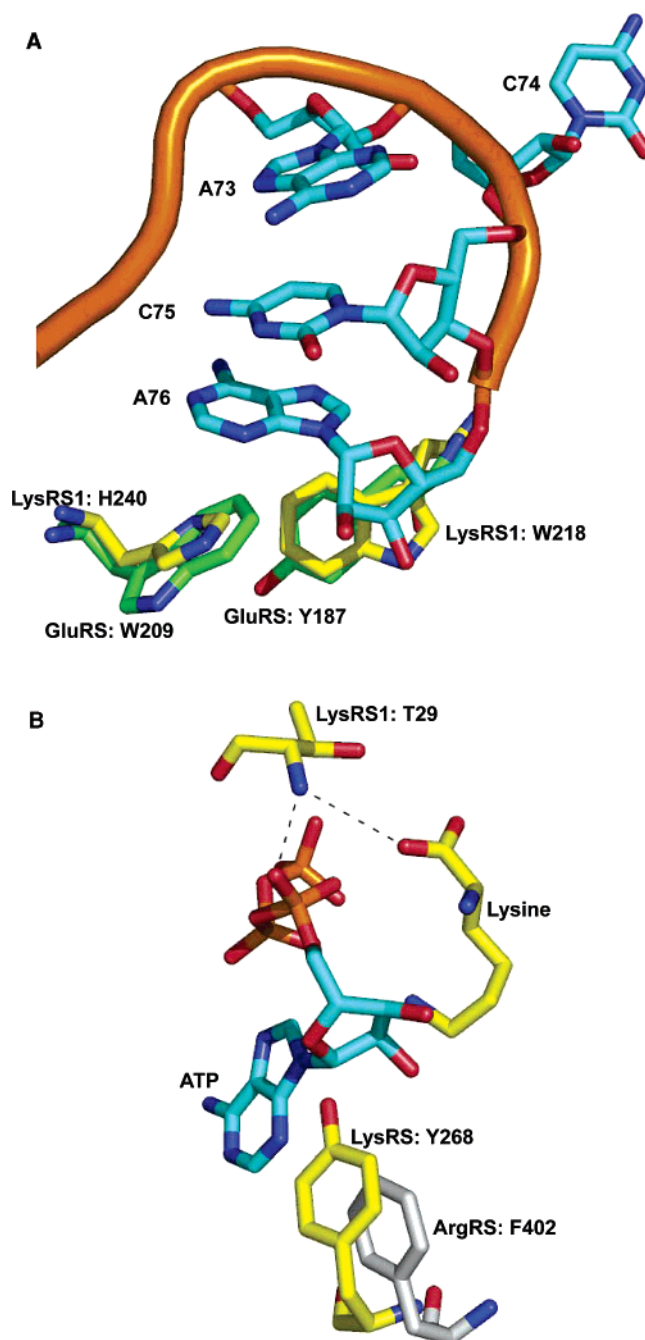


FIGURE 3: (A) Detail of the superposition of the structures of *P. horikoshii* LysRS1 (31) and the *T. thermophilus* GluRS-tRNA^{Glu} complex (26). The 3'-end of tRNA^{Glu} is colored cyan with the phosphate backbone in orange, GluRS active site residues Tyr187 and Trp209 in green, and the corresponding LysRS1 residues Tyr218 and His240 in yellow. (B) Detail of the superposition of the structures of the active sites of *P. horikoshii* LysRS1, the GluRS-ATP complex of *T. thermophilus* (26), and yeast ArgRS (27). LysRS1 residues are colored yellow, GluRS residues cyan, and ArgRS residues gray. The positions of substrates, L-lysine for LysRS1 and L-glutamic acid-ATP for GluRS, are also indicated. Structure superpositions were made using SARF2 (28) to search for similar structural motifs based on α backbones and subsequently visualized with PyMOL (29).

iciency for Y269S LysRS1 (1600-fold for lysine and 5900-fold for ATP) confirmed that the aromatic ring of Tyr is important for interaction with ATP. In agreement with a role in stacking at this position, Y269F showed almost no reduction in K_M for any of the substrates and only a small

drop in k_{cat} . Thus, rather than Y269 being directly involved in lysine binding, this residue is instead involved in the correct positioning of ATP by stacking with the adenosine moiety. This contrasts with the most analogous residue in LysRS2, Y280, which instead interacts directly with lysine.

Kinetic analyses indicated that the role of G29 in LysRS1 involves hydrogen bonding with the α -amino group of lysine. The increase in K_M with almost no change in k_{cat} for lysine indicated that G29 in LysRS1 performs the same function as G216 in LysRS2 of *E. coli*. Another prediction, that T31 is involved in hydrogen bond interactions with the α -carboxylate group of lysine, was confirmed by the T31G replacement which caused a decrease in the specific activity of this enzyme, while T31S exhibited almost the same kinetic parameters as the wild type. The docking model (Figure 3B) indicated that T31 also forms a hydrogen bond with the oxygen atom from the α -phosphate of ATP. The corresponding position in *T. thermophilus* GluRS, S9, is involved in hydrogen bonding with the oxygen atom from the α -phosphate of ATP. The kinetics for ATP indicated that T31G affected the correct positioning and possible orientation of ATP in the active site. The T31S substitution had no effect on the kinetic parameters for lysine or ATP, indicating that the conservation of H-bonding capacity at this position among the class Ib aaRSs is important for interaction with ATP.

Substrate Specificity Determinants in LysRS1. The enhanced discrimination of lysine analogues and noncognate amino acids by LysRS1 was predicted to be based on the compactness of its active site when compared to that of LysRS2 (18). In our previous model, we proposed that H242 and W220 would sterically hinder AEC binding in the active site of LysRS1. Substitutions of H242, W220, and G29 showed that any changes to the active site that compromise lysine binding also affect AEC, as also seen for similar replacements in LysRS2 (21). W220A is the only replacement to show almost no effect on the K_M for lysine but a large increase in the K_i for AEC, suggesting that the absence of the bulky indole moiety promotes closure of the active site thereby further hindering the binding of AEC.

Investigation of the effect of synonymous residues conserved within GluRS and LysRS1 on the discrimination of noncognate amino acids indicated that the active site is fine-tuned to compromise between efficiency and specificity. Almost all the replacements severely reduced efficiency (k_{cat}/K_M for lysine) but increased specificity as they lost the ability to charge tRNA^{Lys} with glutamic acid and arginine. Both glutamic acid and arginine are used by the wild-type enzyme, but only the variant T31S, a conserved replacement with little loss in lysine efficiency, could also charge the noncognate substrates. Any attempts to replace residues involved in lysine binding and convert the LysRS1 active site into a form more similar to GluRS caused severe decreases in the level of lysine binding and a complete loss of glutamic acid and arginine binding. Thus, despite the residual structural and mechanistic similarities between LysRS1 and GluRS, both have diverged sufficiently to maximize binding and selectivity for cognate amino acids while excluding noncognate amino acids within very similar active site architectures.

Divergent Mechanisms of Substrate Discrimination in LysRS1 and LysRS2. The evolution of two unrelated structures to perform the same reaction is so far limited to LysRS

among the aaRSs. We previously proposed that the active site of LysRS2 is more open and, therefore, able to accommodate lysine analogues more easily than LysRS1. These findings corroborate this idea and further suggest that the binding network in LysRS1 is different from that in LysRS2. The kinetic data for W220 and Y269 variants indicate these two residues do not play important roles in lysine binding but rather form the active site scaffold for interactions with ATP and tRNA. In the LysRS2 structure, lysine binding is mainly based on a complex hydrogen bonding network and is much more robust (19), as also indicated by biochemical studies (21). The usage of electrostatic interactions and hydrogen bonds in LysRS2 leads to tight lysine binding (2.6 μM) when compared to that in LysRS1 (225 μM), and since the lysine binding network is more extensive, any single mutation in the LysRS2 active site does not cause as dramatic a change in the catalytic efficiency as observed in LysRS1 variants. There is only one LysRS2 replacement (E278D) which causes a reduction of more than 1000-fold in k_{cat}/K_M for lysine compared to the wild type, and changes in the active site at the other seven residues interacting directly with lysine lower the catalytic efficiency considerably less. The crystal structure of *P. horikoshii* LysRS1 identifies only four residues (E43, H241, T31, and G29) that could play important roles in lysine binding. Since fewer residues in the active site of LysRS1 are directly involved in binding and discrimination of lysine, any nonconservative replacement of these residues can cause severe decreases in catalytic efficiency. The lysine binding network in LysRS1 takes advantage of the specific size and positioning of the substrate in the active site with two distinct sides: polar hydrogen bond interactions with the functional groups of lysine positioned on one side and hydrophobic interactions with the lysine backbone on the other. This dual interaction contributes to the difference in sensitivity toward the discrimination of lysine analogues by excluding larger analogues such as AEC. The same mechanism is not observed in LysRS2, which efficiently screens for hydrogen bond interactions within a larger active site, thereby allowing inhibition by AEC.

Attempts to explain the evolutionary role of two unrelated LysRSs belonging to class I and class II initially focused on identity elements of the tRNA, since this is the largest and potentially most diverse molecule that the two LysRSs must discriminate (14). However, the subtle differences in tRNA^{Lys} discrimination in vitro between LysRS1 and LysRS2, with a G-U wobble in the second base pair of the acceptor stem being an anti-determinant for LysRS2, were not significant in vivo (17). Subsequent analyses of lysine analogues demonstrated in vivo and in vitro that the evolutionary pressure to retain two forms of LysRS likely arose to prevent infiltration of the genetic code by these near-cognate amino acids. The mechanisms of lysine recognition and discrimination by both enzymes are sufficiently different that it was not possible to convert the active site of LysRS2, which is more catalytically efficient, into a more selective site as found in LysRS1 (21). These earlier findings, together with the data presented here, clearly indicate that the continued evolutionary retention of the less efficient class I form of LysRS is a result of the higher substrate specificity offered by the class I active site compared to that of LysRS2.

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