# Coevolution of an aminoacyl-tRNA synthetase with its tRNA substrates

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Contributed by Dieter Söll, September 23, 2003

Glutamyl-tRNA synthetases (GluRSs) occur in two types, the discriminating and the nondiscriminating enzymes. They differ in their choice of substrates and use either tRNAGlu or both tRNAGlu and tRNAGIn. Although most organisms encode only one GluRS, a number of bacteria encode two different GluRS proteins; yet, the tRNA specificity of these enzymes and the reason for such gene duplications are unknown. A database search revealed duplicated GluRS genes in >20 bacterial species, suggesting that this phenomenon is not unusual in the bacterial domain. To determine the tRNA preferences of GluRS, we chose the duplicated enzyme sets from Helicobacter pylori and Acidithiobacillus ferrooxidans. H. pylori contains one tRNAGlu and one tRNAGln species, whereas A. ferrooxidans possesses two of each. We show that the duplicated GluRS proteins are enzyme pairs with complementary tRNA specificities. The H. pylori GluRS1 acylated only tRNAGlu, whereas GluRS2 was specific solely for tRNAGIn. The A. ferrooxidans GluRS2 preferentially charged tRNA<sup>GIn</sup><sub>UUG</sub>. Conversely, A. ferrooxidans GluRS1 glutamylated both tRNAGlu isoacceptors and the tRNAGIN species. These three tRNA species have two structural elements in common, the augmented D-helix and a deletion of nucleotide 47. It appears that the discriminating or nondiscriminating natures of different GluRS enzymes have been derived by the coevolution of protein and tRNA structure. The coexistence of the two GluRS enzymes in one organism may lay the groundwork for the acquisition of the canonical glutaminyl-tRNA synthetase by lateral gene transfer from eukaryotes.

Faithful protein biosynthesis relies on the correct attachment of amino acids to their corresponding (cognate) tRNAs catalyzed by the aminoacyl-tRNA synthetases (AARSs) (1). Given the presence of 20 canonical amino acids, one would expect each organism to contain at least 20 AARSs to ensure high accuracy during aminoacylation of the complete set of tRNAs. However, this is true only for eukarya and some bacteria. Most of the other bacterial organisms (2, 3), all known archaea (4, 5), and eukaryotic organelles (6) lack the AARS specific for glutamine. In addition, a canonical asparaginyl-tRNA synthetase is missing from many archaeal and bacterial genomes (7), whereas a canonical cysteinyl-tRNA synthetase is not distinguishable in some methanogenic archaea (8).

In addition to direct acylation of the cognate tRNA by AARSs, an indirect pathway was found to be responsible for the formation of both Gln-tRNA<sup>Gln</sup> and Asn-tRNA<sup>Asn</sup>. In these cases, formation of the mischarged product (Glu-tRNA<sup>Gln</sup> and Asp-tRNA<sup>Asn</sup>, respectively) is a required intermediate in the synthesis of the cognate amino acid–tRNA pairs. To accomplish this, the indirect pathway to Gln-tRNA<sup>Gln</sup> formation uses a nondiscriminating (ND) form of glutamyl-tRNA synthetase (GluRS) that is able to efficiently glutamylate both tRNA<sup>Glu</sup> and tRNA<sup>Gln</sup> (9). In the next step, a tRNA-dependent amidotransferase, Glu-tRNA<sup>Gln</sup> amidotransferase, amidates Glu-tRNA<sup>Gln</sup> to the correctly charged Gln-tRNA<sup>Gln</sup>, which is then used as a substrate for protein synthesis (reviewed in ref. 10).

In bacteria, the ND Bacillus subtilis GluRS has been extensively studied (9). Due to the lack of a canonical glutaminyltRNA synthetase (GlnRS) in this organism, the ND-GluRS is an essential enzyme in Gln-tRNA formation as it generates GlutRNA<sup>Gln</sup>. This product is then converted to Gln-tRNA<sup>Gln</sup> by Glu-tRNA<sup>Gln</sup> amidotransferase (10-12). However, the mechanism of how the ND-GluRS recognizes two different tRNA substrates is not known, and the tRNA identity set for such an enzyme has never been determined. In contrast, organisms with a canonical GlnRS enzyme possess a discriminating GluRS (D-GluRS), which recognizes only tRNA<sup>Glu</sup>. The tRNA identity elements of a D-GluRS enzyme were reported for E. coli (13). Major identity determinants for this enzyme are located in the augmented D-helix (indicated in Fig. 1), which is formed by nucleotides in the D-stem and D-loop, as well as the bases connecting the acceptor helix and D-stem, and some bases in the extra arm (13). In vitro mutagenesis of Thermus thermophilus D-GluRS expanded its tRNA specificity to the recognition of tRNA<sup>Glu</sup> with a glutamine-specific anticodon (14). However, this work did not afford real insight into a ND-GluRS, because the tRNA specificity was not changed.

In addition to the lack of some synthetases (see above), whole genome analysis also revealed several organisms containing more than one AARS for certain amino acids. In many cases, the function of such a redundancy is unknown; however, in some organisms, the roles of the duplicated AARSs have been explained (15). A second aspartyl-tRNA synthetase in Deinococcus radiodurans, for instance, is shown to be involved in tRNAdependent asparagine biosynthesis (7). In E. coli, two genes for lysyl-tRNA synthetase (LysRS) exist; one LysRS gene (lysS) is constitutively expressed, and the other one is heat-inducible (lysU) (16, 17). Similarly, two paralogous gltX genes (encoding GluRS) exist in some proteobacteria and a scattering of other bacterial groups (18); however, the function of this duplication is not clear. There are indications that the two GluRSs in Helicobacter pylori preferentially glutamylate either tRNA<sup>Glu</sup> or tRNA<sup>Gln</sup>.\*\* To investigate further the duplicated gltX genes, we chose the Acidithiobacillus ferrooxidans and H. pylori systems and examined the substrate preferences of the two GluRS enzymes that coexist in these organisms.

### **Materials and Methods**

**Oligonucleotides, DNA Sequencing, and Radiochemicals.** Oligonucleotides were synthesized, and DNAs were sequenced by the Keck Foundation Biotechnology Resource Laboratory at Yale University. Uniformly labeled [14C]Glu (254 mCi/mmol) and

Abbreviations: AARS, aminoacyl-tRNA synthetase; ND, nondiscriminating; GluRS, glutamyl-tRNA synthetase; GlnRS, glutaminyl-tRNA synthetase.

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<sup>\*\*</sup>Hendrickson, T. L., Skouloubris, S., de Reuse, H. & Labigne, A. (2002) Abstr. Papers Am. Chem. Soc. 224, 121.

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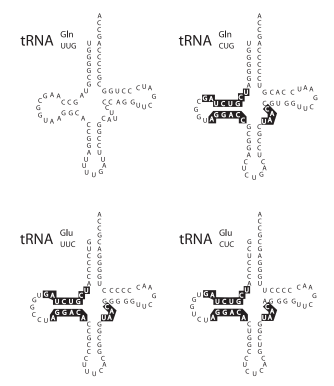


Fig. 1. Cloverleaf representation of A. ferrooxidans tRNAGlu and tRNAGln species. The bases that form part of the augmented D-helix (13), in both isoacceptors of tRNA Glu and in tRNA Gln, are shown in white letters and black

[ $^{14}$ C]Gln (244 mCi/mmol) and [ $\gamma$ - $^{32}$ P]ATP (6,000 Ci/mmol) were from Amersham Pharmacia Biosciences.

Cloning of gltX Genes from H. pylori and A. ferrooxidans. The H. pylori genes (HP0476, gltX1; HP0643, gltX2) were obtained by PCR from the genomic DNA (strain 26695) by using the Expand High Fidelity PCR System (Roche Molecular Biochemicals). Oligonucleotide primers introduced an NdeI site at the 5' end and a SapI site at the 3' end of each gene. The PCR products, digested with NdeI and SapI, were cloned into the pTYB1 vector (New England BioLabs) digested with the same enzymes, to form the plasmids pHPgltX1 and -X2. The identity of the clones was determined by DNA sequencing.

Analysis (http://tigrblast.tigr.org/ufmg/index.cgi?database = a\_ferrooxidans seq) of the incomplete A. ferrooxidans genome sequence revealed two gltX genes. They were cloned into the pCR2.1-TOPO vector (Invitrogen) after PCR amplification of genomic A. ferrooxidans DNA (ATCC 23270) by using oligonucleotides that introduced an EcoRI site on both ends of each gene. After sequence confirmation, the genes were cloned into the EcoRI site of pGEX-2T (Amersham Biosciences) to yield the plasmids pAFgltX1 and -X2.

Expression and Purification of GluRSs. The plasmids pHPgltX1 and -X2 were transformed into Escherichia coli BL21(DE3) (Invitrogen). The expression of both genes and the purification and cleavage of the intein fusion proteins were done as described (New England BioLabs). For the overexpression of A. ferrooxidans genes and purification of the GST-fusion protein, we followed the previously described protocols (19).

Preparation of Unfractionated tRNA. The A. ferrooxidans tRNA<sup>Gln</sup> genes (sequence from the TIGR database), were constructed by synthesis of the corresponding oligonucleotides and subsequent

ligation into the pKK223-3 vector (Amersham Biosciences) and digested with EcoRI and HindIII enzymes, generating the plasmids pJSQUUG and pJSQCUG. These were transformed into E. coli DH5 $\alpha$ . Purification of unfractionated tRNA containing the specified tRNA gene transcript was carried out as described (20). Comparison of aminoacylation reactions by using E. coli GlnRS and unfractionated tRNA obtained from the cells, transformed with pJSQUUG, pJSQCUG, or empty vector, showed that the A. ferrooxidans  $tRNA_{UUG}^{Gin}$  and  $tRNA_{CUG}^{Gin}$  isoacceptors comprised between 10% and 20% of the total tRNA.

To obtain mature unfractionated tRNA from both organisms, H. pylori cells (strain 43526) were grown (21), and the unfractionated tRNA was prepared as described (20). A. ferrooxidans (strain 19859) was grown in a bioreactor in 9-K medium [0.4 g/liter (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 0.4 g/l MgSO<sub>4</sub>(7H<sub>2</sub>O), 0.056 g/l K<sub>2</sub>HPO<sub>4</sub>(3H<sub>2</sub>O)/5 g/l FeSO<sub>4</sub>(7H<sub>2</sub>O) adjusted to pH 1.6 with concentrated H<sub>2</sub>SO<sub>4</sub>, and sterilized by filtration. The ferrous ion was regenerated by 0.5 A of current]. Total tRNA extracted from 0.5 g of A. ferrooxidans cells was purified by the Qiagen RNA/DNA kit as described by the manufacturer (Qiagen, Valencia, CA).

Purification of tRNAGIn and tRNAGIu by Affinity Chromatography. The A. ferrooxidans tRNA $_{\rm UUG}^{\rm Gln}$  and tRNA $_{\rm CUG}^{\rm Gln}$  (see above) and E. coli tRNA $_{\rm CUG}^{\rm Gln}$  and tRNA $_{\rm CUG}^{\rm Gln}$  were purified from unfractionated tRNA by affinity chromatography on immobilized T. thermophilus EF-Tu (22) with the following specifications. Unfractionated *E*. coli tRNA was aminoacylated with E. coli GlnRS or GluRS. The Gln-tRNAGln or Glu-tRNAGlu was separated from the uncharged tRNA by the formation of a ternary complex with the EF-Tu-GTP immobilized on a Ni-NTA-agarose column. After the elution of Gln-tRNAGln or Glu-tRNAGlu, the tRNA was deacylated for 15 min at 65°C in 0.1 mM borate-KOH, pH 9.0. The pure tRNA<sup>Gln</sup> was not contaminated with tRNA<sup>Glu</sup> and vice versa, as judged by aminoacylation with pure E. coli GluRS or GlnRS, respectively.

**Aminoacylation Assays with GluRS1 and -RS2.** *In vitro* acylations with H. pylori and A. ferrooxidans GluRS enzymes was carried out at 37°C in an 80-μl reaction mixture containing 100 mM Hepes-KOH, pH 7.2; 30 mM KCl; 12 mM MgCl<sub>2</sub>; 5 mM ATP; 2 mM DTT; 25  $\mu$ M [14C]Glu plus 75  $\mu$ M unlabeled glutamate and 40–60 μg of unfractionated tRNA or 0.7–1.3 μg of pure tRNA. The reactions were started by adding the enzyme (100-150 nM). Aliquots (15  $\mu$ l) were taken at different times, spotted on 3MM filter paper discs, and washed twice with 5% trichloroacetic acid. For acidic gel separations (see below), unfractionated H. pylori or A. ferrooxidans tRNA was charged in the presence of 0.1 mM unlabeled glutamate.

Acid Urea Gel Electrophoresis of tRNA and Aminoacyl-tRNA. This method (23) allows the separation of charged from uncharged tRNA due to a difference in electrophoretic mobility between the two species. Hybridization of a sequence-specific probe permits the determination of the identity of the tRNA on the gel. Unfractionated tRNA (80  $\mu$ g) from *H. pylori* or *A. ferrooxidans* was glutamylated for 15 min at 37°C with the homologous synthetases (see above), and half of the reaction was deacylated for 15 min at 65°C with 0.1 mM borate-KOH, pH 9.0. After extraction with phenol (saturated with 0.3 M sodium acetate, pH 5.0/10 mM EDTA), the charged and uncharged tRNAs were recovered by ethanol precipitation. They were dissolved (at a final concentration of 5  $\mu$ g/ $\mu$ l) in sample buffer (0.1 M sodium acetate, pH 5.0/8 M urea/0.05% bromophenol blue/0.05% xylene cyanol) The RNA samples (10  $\mu$ g) were loaded on a 9.5% polyacrylamide gel ( $50 \times 20$  cm, 0.4 mm thick) containing 7 M urea and 0.1 M sodium acetate, pH 5.0 and run at 4°C, 600 V in 0.1 M sodium acetate, pH 5.0, for 40 h.

Detection of the tRNAs was performed by Northern blotting.

Table 1. Organisms with two gltX genes

Organism	Bacterial class	tRNA <sup>Glu</sup> species	tRNA <sup>GIn</sup> species	Gat CAB	gluX	GlnRS
Anaplasma marginale	$\alpha$ -Proteobacteria	1	1	+	_	
Anaplasma phagocytophilum	$\alpha$ -Proteobacteria	1	1	+	_	_
Brucella melitensis	$\alpha$ -Proteobacteria	2	2	+	+	_
Brucella suis	$\alpha$ -Proteobacteria	2	2	+	+	_
Ehrlichia chaffeensis	$\alpha$ -Proteobacteria	1	1	+	_	_
Magnetospirillum magnetotacticum	$\alpha$ -Proteobacteria	2	2	+	+	_
Mesorhizobium loti	$\alpha$ -Proteobacteria	2	2	+	+	_
Neorickettsia sennetsu	$\alpha$ -Proteobacteria	1	1	+	_	_
N. aromaticivorans	$\alpha$ -Proteobacteria	2	2	+	+	_
Rhodobacter sphaeroides	$\alpha$ -Proteobacteria	1	1	+	+	_
Rhodospirillum rubrum	$\alpha$ -Proteobacteria	2	2	+	+	_
Rickettsia conorii	$\alpha$ -Proteobacteria	1	1	+	_	_
Rickettsia prowazekii	$\alpha$ -Proteobacteria	1	1	+	_	_
Silicibacter pomeroyi	$\alpha$ -Proteobacteria	1	1	+	+	_
Wolbachia sp.	$\alpha$ -Proteobacteria	1	1	+	_	_
A. ferrooxidans	$\gamma$ -Proteobacteria	2	2	+	+	_
Coxiella burnetii	γ-Proteobacteria	1	1	+	_	_
Methylococcus capsulatus	γ-Proteobacteria	2	2	+	+	_
C. jejuni	$\varepsilon$ -Proteobacteria	1	1	+	_	_
H. pylori sp.	$\varepsilon$ -Proteobacteria	1	1	+	_	_
Helicobacter hepaticus	arepsilon-Proteobacteria	1	1	+	_	-
D. hafniense	Clostridia	2	1	+	_	+
Thermoanaerobacter tengcongensis	Clostridia	2	2	+	_	-
T. maritima	Thermotogae	2	2	+	-	-

gluX, a truncated gltX gene (see text); GatCAB, Glu-tRNA<sup>Gln</sup> amidotransferase; + or -, presence or absence, respectively.

For this purpose, the portion of the gel containing the tRNAs was electroblotted onto a Hybond-*N* + membrane (Amersham Bioscience) by using a Hoefer Electroblot apparatus (Amersham Bioscience) at 10 V for 10 min and then at 30 V for 90 min with 10 mM Tris acetate, pH 7.8/5 mM sodium acetate/0.5 mM Na-EDTA as transfer buffer. The membranes were then baked at 72°C for 2 h. The tRNAs were detected by hybridization with a 5′-<sup>32</sup>P-labeled oligodeoxyribonucleotide probe. For *H. pylori* tRNA<sup>Gln</sup>, the probe complementary to nucleotides 1–21. *H. pylori* has two slightly different tRNA<sup>Glu</sup><sub>GUC</sub> genes; however, by Northern blot, we detected the product of one gene as the main tRNA<sup>Glu</sup> (data not shown); the probe used to detect that tRNA was complementary to nucleotides 1–21 (5′-TAACCACTA-GATGAAGGAGCC-3′).

The *A. ferrooxidans* genome sequence reveals two  $tRNA^{Gln}$  isoacceptors,  $tRNA^{Gln}_{UUG}$  and  $tRNA^{Gln}_{CUG}$  (see Fig. 1). The probes were complementary to nucleotides 13–37 and 53–75 for  $tRNA^{Gln}_{UUG}$  and to nucleotides 14–37 for  $tRNA^{Gln}_{CUG}$ . In the case of  $tRNA^{Glu}_{Glu}$ , we also found two isoacceptors that were detected by using the probes complementary to nucleotides 15–39 for each tRNA.

### **Results**

Selection of the Experimental Systems. The presence of two paralogous gltX genes has been previously noticed in a few bacterial genomes (18). To survey the presence of duplicated GluRS genes or pieces thereof, we performed TBLASTN searches (24) of all currently available bacterial genomes. As a result, gltX duplications were found in the genomes of >20 organisms (Table 1). With the exception of two members of the Clostridiales and the deep-rooted  $Thermotoga\ maritima$ , all organisms with duplicated gltX genes are proteobacteria.

Apart from duplications of the entire gltX gene, many genomes were found to contain gluX (yadB in  $E.\ coli$ ), a truncated form of gltX encoding a GluRS protein that lacks the enzyme's entire C-terminal anticodon-binding domain ( $\approx 35\%$  of the total protein) and has no ascribed function; it was suggested to be a

pseudogene (25) and shown to be not essential in *E. coli* (ref. 26, J.C.S., unpublished results). The relatively wide distribution (from *E. coli* to *D. radiodurans* and *Corynebacterium*) of the genes encoding these precisely truncated GluRS fragments suggests that these proteins may have a yet unknown function. Nonetheless, GluX was not analyzed further.

Class I lysyl-tRNA synthetase (LysRS) is structurally very related (27) and displays sequence homology to GluRS near its C terminus (COG 0008). Its rare occurrence in bacteria is confined largely to  $\alpha$ -proteobacteria. It is interesting to note that the entire set of  $\alpha$ -proteobacteria with two gltX genes (Table 1) is identical to that containing a class I LysRS. Whether this is a functionally significant correlation remains to be seen.

Phylogenetic analysis of GluRS sequences in a general tree showed a segregation of the duplicated GluRS proteins into two groups (Fig. 2). The first enzyme, GluRS1, branched with discriminating GluRS of *E. coli* and other proteobacteria. The second enzyme, GluRS2, appeared quite diverged but was in some cases closer to the ND-GluRSs, e.g., *B. subtilis* GluRS. Therefore, we assumed that GluRS1 is a canonical discriminating synthetase, whereas GluRS2 is the ND enzyme capable of forming Glu-tRNA<sup>Gln</sup>, the tRNA substrate in the transamidation pathway of Gln-tRNA formation (reviewed in ref. 10). The key enzyme in this pathway, the Glu-tRNA<sup>Gln</sup> amidotransferase (encoded by *gatCAB*) (10–12), is present in all of the organisms with two *gltX* genes (Table 1). A relationship, mediated by a common partner, has been established between *H. pylori gltX2* and *gatA* in a genomic protein–protein interaction map (29).

To complement the enzyme analysis, we proceeded to examine the tRNA<sup>Glu</sup> and tRNA<sup>Gln</sup> species in these organisms. Our survey showed that some of the organisms possess only one isoacceptor of each tRNA<sup>Glu</sup> and tRNA<sup>Gln</sup> (e.g., *H. pylori* and *Campylobacter jejuni*), whereas others have two (e.g., *A. ferrooxidans* and *Novosphingobium aromaticivorans*). Closer examination brought better insight: in most cases where a genome contains two tRNA<sup>Gln</sup> isoacceptors, they are dissimilar and

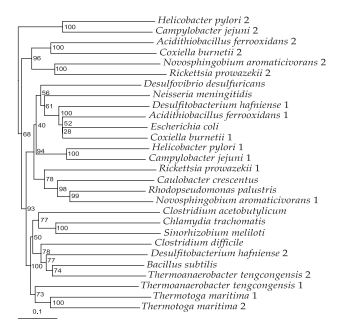


Fig. 2. Phylogeny of bacterial-type GluRS proteins. Amino acid sequences were aligned by using the CLUSTALW program (Version 1.82) (28). Initial analyses were performed with GluRS proteins from numerous bacterial organisms, and later the same was confirmed with a set of representative GluRSs. Phylogeny was inferred by using the neighbor-joining method to create and evaluate 1,000 resampled alignments. Bootstrap percentages are given for each branch. (Bar = 10 aa replacements per 100 positions.)

structurally distinguished by the presence or absence of an augmented D-helix (13). The tRNA<sup>Glu</sup> and tRNA<sup>Gln</sup> species of A. ferroxidans are depicted in Fig. 1; the tRNA<sub>CUG</sub> has an augmented D-helix, whereas the tRNA<sub>UUG</sub> lacks this element; in addition it has a different transfer of the control of the con addition it has a different acceptor stem sequence. Thus, the A. ferrooxidans tRNA<sub>CUG</sub> may be a substrate for the E. coli-type GluRS (GluRS1). Together, these findings suggest that the two GluRS proteins may bind structurally different tRNA substrates. Therefore, we chose A. ferrooxidans and H. pylori, organisms with duplicated gltX genes, to examine the possible reason for the presence of two GluRS proteins by determining the tRNA substrate preferences of these enzymes.

# The Two H. pylori GluRSs Possess Complementary tRNA Specificities. The gltX1 and gltX2 genes from H. pylori and A. ferrooxidans were cloned into expression vectors to encode intein or GST fusion proteins, respectively. After expression in E. coli and affinity chromatography, the native H. pylori enzymes were generated by intein cleavage, whereas the A. ferrooxidans GluRS enzymes were used as GST fusion proteins. Preliminary experiments showed that GluRS1 and -RS2 from both organisms efficiently charge total E. coli tRNA with glutamate but, as expected, not glutamine (data not shown). We then proceeded to determine

the enzymes' tRNA specificities. Because H. pylori possess only one tRNA<sup>Glu</sup> and one tRNA<sup>Gln</sup> species, we analyzed this system first. Because H. pylori tRNAs are very similar in sequence to E. coli tRNAs (>80% sequence identity for both species), we proceeded to purify tRNA Glu and tRNAGIn from unfractionated E. coli tRNA by EF-Tu affinity chromatography (22). The resulting pure E. coli tRNA species were then acylated with glutamate by the H. pylori GluRS1 and -RS2 enzymes. GluRS1 charged tRNA<sup>Glu</sup> well, whereas GluRS2 did not recognize this tRNA (Fig. 3A). In contrast, the result with tRNA<sup>Gln</sup> was completely reversed, because this tRNA was charged exclusively by GluRS2 (Fig. 3B). To ascertain the validity of these results obtained in charging the heterologous E.

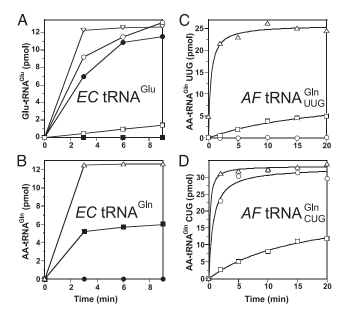


Fig. 3. Aminoacylation of tRNA<sup>Glu</sup> and tRNA<sup>Gln</sup> from *E. coli* and *A. ferrooxi*dans. The reaction conditions were as indicated in Materials and Methods by using the pure tRNAs. (A) E. coli tRNAGlu. (B) E. coli tRNAGln. (C) A. ferrooxidans tRNAGIN (D) A. ferrooxidans tRNAGIN. The enzymes used were E. coli GluRS  $(\nabla)$  and GlnRS  $(\Delta)$ , H. pylori GluRS1  $(\bullet)$  and GluRS2  $(\blacksquare)$ , and A. ferrooxidans GluRS1 (o) and GluRS2 ( $\square$ ). Glutamic acid was used with the GluRS, and glutamine was used with GlnRS.

coli tRNA, we also analyzed glutamylated mature H. pylori tRNA samples by Northern blot of acid/urea gels (23). Glu-tRNAs generated by H. pylori GluRS1 and -RS2 and the control Gln-tRNA formed by E. coli GlnRS were used. After running the gel, Northern blot analysis was performed by using a specific probe for each tRNA of interest. Glu-tRNA Gln was formed only in the presence of GluRS2 (Fig. 4A, lane 3), whereas GlutRNAGlu was formed only in the presence of the GluRS1 (Fig. 4A, lane 6). These results confirmed that GluRS1 and -RS2 recognize the homologous tRNAGlu and tRNAGln, respectively (Fig. 4A, compare lanes 2 and 3 to lanes 6 and 8). As expected, Gln-tRNA<sup>Gln</sup>, made by E. coli GlnRS, moved more slowly than Glu-tRNA<sup>Gln</sup> (Fig. 4A, compare lanes 1 and 3), which suggests that this method could also be used to provide a clue to the amino acid identity of the reaction products.

# tRNA Specificity of GluRS Is Linked to Idiosyncratic tRNA Features.

The A. ferrooxidans GluRS system is expected to be more complex than that of H. pylori, because there are two tRNAGlu and two tRNAGin isoacceptors (Fig. 1). Although these four tRNAs have different anticodons, both tRNAGlu isoacceptors are uniform in their secondary structure and similar to E. coli and *H. pylori* tRNA<sup>Glu</sup>, one of the tRNA<sup>Gln</sup> isoacceptors shows a striking divergence in its structure and is in some aspects more similar to tRNA<sup>Glu</sup> than tRNA<sup>Gln</sup> (Fig. 1). We expressed the A. ferrooxidans tRNA genes in E. coli and purified them by EF-Tu chromatography (22). The aminoacylation results show clearly that tRNA<sub>UUG</sub> was not charged by A. ferrooxidans GluRS1 but moderately by GluRS2 and well by E. coli GlnRS (Fig. 3C). †† However, tRNA<sub>CUG</sub> was charged well by GluRS1 and E. coli GlnRS but less well by GluRS2 (Fig. 3D). As expected, only A. ferrooxidans GluRS1 could charge E. coli tRNAGlu (Fig. 3A).

 $<sup>^{\</sup>dagger\dagger}$ A. ferrooxidans GluRS2 does not reach the same plateau as  $\it E.~coli$  GlnRS (Fig. 3 $\it C$ ). This may be due to the lack of a nucleotide modification in the heterologously produced tRNAGING or to the presence of a supernumerary one introduced by E. coli. This is plausible, because mature A. ferrooxidans tRNAGIn is fully charged by GluRS2 (Fig. 4B, lane 5).

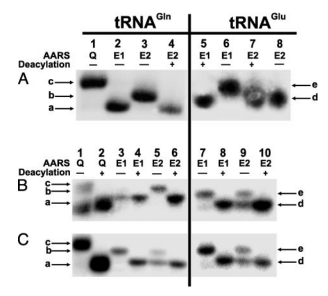


Fig. 4. Northern blot analysis of *H. pylori* and *A. ferrooxidans* tRNA after *in vitro* acylation by the different GluRS enzymes. The blots were probed with <sup>32</sup>P-labeled oligonucleotide complementary to the anticodon stem or plus acceptor stem (see *Materials and Methods*). Blots correspond to the detection of tRNA<sup>Gln</sup> (*Left*) and to tRNA<sup>Glu</sup> (*Right*). (*A*) Unfractionated tRNA from *H. pylori*. (*B*) Unfractionated tRNA from *A. ferrooxidans* probed for tRNA<sup>Gln</sup><sub>UUC</sub> or tRNA<sup>Glu</sup><sub>GUC</sub>. (C) Unfractionated tRNA from *A. ferrooxidans* probed for tRNA<sup>Gln</sup><sub>GuC</sub> tRNA<sup>Glu</sup><sub>GUC</sub>. Lane 1, charging of unfractionated tRNA with glutamine by *E. coli* GlnRS. The enzymes used are: E1, GluRS1; E2, GluRS2; Q, *E. coli* GlnRS. The arrows indicate positions of AA-tRNA: a, uncharged tRNA<sup>Gln</sup>; b, Glu-tRNA<sup>Gln</sup>; c, Gln-tRNA<sup>Gln</sup>; d, uncharged tRNA<sup>Glu</sup>.

These tRNA specificities were confirmed by glutamylation of mature unfractionated A. ferrooxidans tRNA followed by acidic gel analysis, as described above for H. pylori. The A. ferrooxidans GluRS1 substrates are both tRNA<sup>Glu</sup> isoacceptors (Fig. 4 B and C, lane 7) and tRNA<sup>Gln</sup><sub>CUG</sub> (Fig. 4C, lane 3), but not tRNA<sup>Gln</sup><sub>Glu</sub> (Fig. 4B, lane 3). On the other hand, the A. ferrooxidans GluRS2 recognized primarily tRNA<sup>Gln</sup><sub>GluG</sub> (Fig. 4B, lane 5) but also exhibited some charging of the other tRNA<sup>Gln</sup> isoacceptor and tRNA<sup>Glu</sup> (Figs. 4B, lane 9, and C, lanes 5 and 9), thus retaining some characteristics of the ND-GluRS enzymes. That GluRS1 charged exclusively tRNAs with augmented D-helix structure strongly suggests this is the major identity element for this group of enzymes

Altogether, these results imply the existence of a definite relationship between GluRS types and the specific structure of their tRNA substrates.

## Discussion

RNA Sequence Suggests Structural Differences in Bacterial tRNA<sup>Gln</sup> Species. Inspection of the tRNA<sup>Glu</sup> and tRNA<sup>Gln</sup> sequences in the Genomic tRNA Database (http://rna.wustl.edu/tRNAdb) revealed that most archaeal and eukaryal tRNA<sup>Glu</sup> and tRNA<sup>Gln</sup> species (with the exception of *Saccharomyces cerevisiae*) possess sequences containing the augmented D-helix element. Thus, this RNA structural motif may be a relatively old element contributing to the function of these tRNAs. However, restricted to bacteria, structurally dissimilar (with respect to the augmented D-helix) tRNA<sup>Glu</sup> and tRNA<sup>Gln</sup> species, are frequently observed. In addition, tRNA<sup>Gln</sup> isoacceptors exist that differ in the presence or lack of augmented D-helix (e.g., *H. pylori*, one species with a nonaugmented D-helix; *A. ferrooxidans*, two tRNA<sup>Gln</sup> species, one with and one without augmented D-helix; *Caulobacter crescentus*, one species with an augmented D-helix).

Thus, it is reasonable to assume the existence of different GluRS types as being related to this unusual tRNA diversity.

Bacterial gltX Gene Duplications Occurred Several Times. Phylogenetic analyses suggest a common origin of the GluRS and GlnRS family of AARSs (25, 30, 31). It is widely accepted that GlnRS probably evolved within the eukaryal ancestor through duplication of a *gltX* gene and later spread by horizontal gene transfer to the bacterial domain (25). Thus, it is plausible that a similar gltX gene duplication event may have occurred in some bacterial species and gave rise to a new subgroup of genes encoding enzymes with the tRNA specificity of GlnRS. Phylogenetic analysis shows that the tRNAGIn-specific GluRS (GluRS2) described here arose from the ancestral gene duplication within bacterial GluRSs (ref. 30; this work), possibly in  $\alpha$ -proteobacteria. A standard ND-GluRS may have preceded the GluRS2type enzyme, because A. ferrooxidans GluRS2 retains traces of tRNA<sup>Glu</sup>-binding capability. Such a gene duplication must have taken place several times in bacterial evolution, because GluRS2 enzymes in proteobacteria, in Thermoanaerobacter tengcongensis and in Thermotoga maritima, for instance, are clearly not of the same origin. This gene duplication in *T. maritima* is a much more recent event, and its two GluRSs group together in the phylogenetic tree (Fig. 2).

GluRS Duplication, a Strategy to Enhance Fidelity of Aminoacylation? The consistent occurrence of structurally different tRNA<sup>Glu</sup> and tRNA<sup>Gln</sup> sequences in organisms with a *gltX1* and a *gltX2* gene suggests that the separation of tRNA specificities in the corresponding GluRS enzymes may have provided higher fidelity of aminoacylation. It is reasonable to assume that use of specialized enzymes like GluRS1, GluRS2, or GlnRS would be favorable in terms of aminoacylation accuracy, compared with the broader specificity of GluRS-ND enzymes.

**GluRS2, a New Type of AARSs.** The essential ND-GluRS (e.g., *B. subtilis* GluRS) found in many organisms is a synthetase specific for one amino acid (Glu) and two acceptor RNA families (tRNA<sup>Gln</sup> and tRNA<sup>Glu</sup>). This simple view of a single type of ND-GluRS among bacteria is not universally correct. Our results uncovered at least two additional types of GluRS. The first is still a ND-GluRS that, in addition to tRNA<sup>Glu</sup>, recognizes only the subset of the tRNA<sup>Gln</sup> isoacceptor having an augmented D-helix (e.g., *A. ferrooxidans* GluRS1). The second enzyme is unique among AARSs; it is a new "mischarging" synthetase with an amino acid specificity (Glu) different from its single tRNA specificity (tRNA<sup>Gln</sup>). This GluRS (e.g., *H. pylori* GluRS2) is an enzyme halfway between the discriminating GluRS and GlnRS enzymes, each of which recognizes only (and correctly) their cognate amino acid/tRNA pair.

**GluRS2** and **GlnRS** Evolved Independently. It is tempting to speculate that GluRS2 might be on its way to developing into a true bacterial-origin GlnRS that has not yet evolved a glutaminebinding capacity. However, all known bacterial GlnRS proteins show the highest similarity to eukaryal GlnRSs (25) and are apparently not descendants of the tRNAGin-specific GluRS2 proteins discussed here. Furthermore, examination of the still incomplete Desulfitobacterium hafniense genome revealed (Table 1) the coexistence of ORFs for GluRS1 and -RS2 proteins with the predicted complementary tRNA specificities and for a canonical GlnRS. This implies that the GluRS2 and GlnRS enzymes evolved independently. Possibly the coexistence of GluRS1 and -RS2 may have prepared the organism for the acquisition of a canonical GlnRS by horizontal gene transfer from eukarya and subsequent loss of the nonorthologous gltX2 gene. This might have been an easier scenario than evolving a new amino acid specificity (Glu to Gln) in gltX2. These assump-

Diversity of Bacterial GluRS Enzymes. Based on currently available information, there appear to be three groups of GluRS covering most of the examples investigated. Many bacteria [e.g., Sinorhizobium meliloti (3) or Chlamydia trachomatis (20)] have a single ND-GluRS and tRNAGln isoacceptor without the augmented D-helix. The GluRS2-type enzyme is adapted to recognize only tRNA<sup>Gln</sup> lacking the augmented D-helix, a tRNA species also recognized by the canonical GlnRS. Last, there is GluRS1, an enzyme on its way to becoming a discriminating GluRS and preferentially recognizing tRNA species with the augmented D-helix. Therefore, the discriminating or ND natures of GluRS may have formed by coevolution of protein and tRNA structure.

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Considering all of the data, it is clear that the GluRS/GlnRS family of enzymes shows exquisite diversity of gene arrangement and pathways, which might be linked to a relatively young evolution of canonical GlnRS activity.

**Note Added in Proof.** During the course of publication, a similar study with comparable results was published (32).

We are grateful to Jesse Rinehart for assistance with the gel-shift experiment; to Klaus Melchers (ALTANA Pharma, Konstanz, Germany) for the gift of H. pylori DNA; to David Graham and Michael Ibba for critical questions; and to Milind Deshpande and Jolene Ehler for assistance in growing H. pylori cells at the Center for Biocatalysis at the University of Iowa. This work was supported by grants from the National Institute of General Medical Sciences (to L.D. and D.S.), the Department of Energy (to D.S.), the National Aeronautics and Space Administration (to D.S.), the U.S. Department of Agriculture (to L.D.), and Fondo Nacional de Desarrollo Científico y Tecnológico Chile (Grant 1020087 to O.O.).

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