Crystal structure of Pyrococcus horikoshii tryptophanyl-tRNA synthetase and structure-based phylogenetic analysis suggest an archaean origin of tryptophanyl-tRNA synthetase

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ABSTRACT

The ancient and ubiquitous aminoacyl-tRNA synthetases constitute a valuable model system for studying early evolutionary events. So far, the evolutionary relationship of tryptophanyl- and tyrosyl-tRNA synthetase (TrpRS and TyrRS) remains controversial. As TrpRS and TyrRS share low sequence homology but high structural similarity, a structure-based method would be advantageous for phylogenetic analysis of the enzymes. Here, we present the first crystal structure of an archaean TrpRS, the structure of Pyrococcus horikoshii TrpRS (pTrpRS) in complex with tryptophanyl-5'AMP (TrpAMP) at 3.0Å resolution which demonstrates more similarities to its eukaryotic counterparts. With the pTrpRS structure, we perform a more complete structure-based phylogenetic study of TrpRS and TyrRS, which for the first time includes representatives from all three domains of life. Individually, each enzyme shows a similar evolutionary profile as observed in the sequence-based phylogenetic studies. However, TyrRSs from Archaea/Eucarya cluster with TrpRSs rather than their bacterial counterparts, and the root of TrpRS locates in the archaean branch of TyrRS, indicating the archaean origin of TrpRS. Moreover, the short distance between TrpRS and archaean TyrRS and that between bacterial and archaean TrpRS, together with the wide distribution of TrpRS, suggest that the emergence of TrpRS and subsequent acquisition by Bacteria occurred at early stages of evolution.

INTRODUCTION

Aminoacyl-tRNA synthetases (aaRSs) are a family of enzymes that play a vital role in maintaining the fidelity of transferring the genetic information from mRNA to protein in protein synthesis (1). They catalyze the aminoacylation reaction by first activating amino acids to form aminoacyl-AMPs and then attaching the activated amino acids to the 3'-end of their cognate tRNAs to form aminoacyl-tRNAs. These aminoacyl-tRNAs further recognize the trinucleotide codons of mRNA through the complementary anticodons on tRNAs during the protein synthesis. Since the amino acid sequence of a protein determines its structure and further function(s), errors in the aminoacylation reaction that result in incorrect incorporation of amino acids during protein synthesis can lead to serious consequences. Due to their importance, aaRSs have been suggested to be the first group of enzymes to evolve from the ancient ‘RNA world’ to the present ‘protein world’. Being ancient and ubiquitous, aaRSs are good candidates for studying early evolutionary events and hence have been subjects of intense interest (2).

aaRSs form two mutually exclusive classes with different structural architectures and aminoacylation mechanisms which are suggested to evolve from two distinct ancestors as a result of convergent evolution (3,4). Each of the two classes of aaRSs encompasses three subclasses

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The authors wish it to be known that, in their opinion, the first two authors should be regarded as joint First Authors.

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and the members of each subclass are more closely related to each other than to others in the same class (5). So far, the evolutionary relationship between TrpRS and TyrRS, the only two members of subclass Ic, remains controversial. TrpRS and TyrRS are found to be closely related as crystal structures of Bacillus stearothermophilus TrpRS and TyrRS significantly resemble each other despite their low sequence similarity (6,7). Moreover, some mutants of Bacillus subtilis TrpRS lost the discrimination against Tyr, suggesting a close evolutionary relationship between TrpRS and TyrRS (8). In 1996, Ribas de Pouplana et al. (9) investigated the phylogenetic relationship of TrpRS and TyrRS based on a multiple sequence alignment of 16 bacterial and eukaryotic sequences available at that time and found that the two types of enzymes were clustered according to their bacterial or eukaryotic nature but not amino acid specificity. However, when Brown et al. (10) conducted a similar analysis with 32 sequences from a broader range of taxa in 1997, a contradictory conclusion was drawn as TrpRSs and TyrRSs form separate clades on the basis of amino acid specificity, which was considered to be attributed to the inclusion of sequences from more species especially those from Archaea. One year later, Diaz-Lazcoz et al. (11) performed a similar study with all available sequences of aaRSs including 49 TrpRS and TyrRS sequences. The pyramidal classification of the sequences implies that the archaeal/eukaryotic TyrRSs resemble more to the archaeal/eukaryotic TrpRS than to their bacterial counterparts, which is in accord with part of the observations by Ribas de Pouplana et al. (9). On the other hand, their results also partially support the notion by Brown et al. as bacterial TrpRSs were grouped with archaeal/eukaryotic TrpRSs in the pyramidal classification and the constructed combined phylogenetic trees of TrpRS and TyrRS seemed to have more similarities with those generated by Brown et al. (10). The subsequent work by Woese et al. showed that the full canonical pattern holds for either enzyme, although the phylogenetic relationship between TrpRS and TyrRS was not assessed (2). Later, Yang et al. suggested that the amino acid specificities of TrpRS and TyrRS were established in the early stages of bacterial evolution (12).

All the aforementioned studies were based on the multiple sequence alignment method. However, our analysis of the TrpRS and TyrRS sequences using NCBI BLAST shows that the sequence identities between TrpRSs and TyrRSs and those between eukaryotic TrpRSs/TyrRSs and their bacterial counterparts are under 20% which is below the twilight zone threshold (13), and hence the sequence alignment method is less reliable for phylogenetic study of these enzymes. It is well known that for homologous proteins 3D structures and structural features are more conserved than primary sequences, and thus protein structures also provide useful evolutionary information especially when the sequence homology is low (14–16). In 2003, O’Donoghue and Luthey-Schulten (17) investigated the evolutionary paths of aaRSs with a structural alignment method. Specifically, the analysis based on the unweighted pair group method with arithmetic averages (UPGMA) algorithm showed that TrpRS and TyrRS conform to the canonical pattern. However, in the UPGMA tree, the branching between Homo sapiens TyrRS (hTyrRS) (representing the archaeal type) and bacterial TrpRS is short, and the phylogenetic tree based on the neighbor-joining (NJ) algorithm controversially grouped hTyrRS with bacterial TrpRS. The ambiguity was suggested to be attributed to the lack of crystal structures of archaeal/eukaryotic TrpRSs (17).

During the past several years, with the rapid development of structural biology and structural genomics, increasingly more crystal structures of aaRSs have been determined, including those of TrpRSs from Eucarya (12,18) and more TyrRSs from Archaea (19). However, by the time this study was initiated, no structure of TrpRS from Archaea was available. Thus, we were motivated to solve the crystal structure of Pyrococcus horikoshii TrpRS (pTrpRS) and employ the structural information for further evolutionary study. Here, we report the crystal structure of pTrpRS in complex with tryptophanyl-5’ AMP (TrpAMP), the first archaeal TrpRS structure, and present the results of a more complete structure-based phylogenetic analysis of TrpRS and TyrRS which for the first time includes structures from species representing all three domains of life. Our data suggest the origination of TrpRS from archaeal TyrRS and the subsequent horizontal transfer of TrpRS from Archaea to Bacteria, providing new insights into the evolutionary paths of TrpRS and TyrRS.

MATERIALS AND METHODS

Cloning, expression and purification

The P. horikoshii genomic DNA was used as the template to amplify the trpSI gene. The gene fragment was inserted into the NcoI and SacI restriction sites of the pET28a expression vector (Novagen) which adds a hexahistidine tag at the C-terminus of the protein product. Escherichia coli strain BL21 (DE3) was transformed with the plasmid and when OD600 of the transformed cells reached 0.6–0.8, 1 mM IPTG was added to induce protein expression at 37°C for 4 h. The cells were harvested, followed by sonication in the lysis buffer (50 mM NaH2PO4/Na2HPO4, pH 8.0, 300 mM NaCl, 5 mM 2-mercaptoethanol and 1 mM PMSF). The supernatant fraction of the lysate was applied onto a Ni-affinity column (Qiagen) pre-equilibrated with the lysis buffer, and then the column was loaded in turn with the washing and elution buffers (the lysis buffer supplemented with 30 and 200 mM imidazole, respectively). The eluted fractions were dialyzed against the dialysis buffer (20 mM bicine buffer, pH 9.0, 2 mM MgCl2, 5 mM 2-mercaptoethanol and 1 mM PMSF) and concentrated to 20 mg/ml for crystallization screening.

Crystallization, data collection, structure determination and refinement

Prior to crystallization screening the purified protein was heated at 70°C for 10 min followed by centrifugation.
at 4°C. The supernatant was collected and 2 μl of the protein solution was mixed with 2 μl of the crystallization solution to form crystallization drops. The crystallization process was carried out at 20°C using hanging drop vapor diffusion method and crystals grew from drops containing the protein solution and the crystallization solution [0.1 M sodium citrate, pH 5.2, 1.6 M (NH₄)₂SO₄ and 10 mM MnCl₂] supplemented with 0.5 μl of 20 mM Trp and 0.5 μl of 100 mM ATP.

X-ray diffraction data were collected from a flash-cooled crystal at synchrotron beamline BL17A of Photon Factory, Japan and processed using the HKL2000 software package (20). The structure of pTrpRS was solved by the molecular replacement method with program PHASER (21) [implemented in the CCP4 suite (22)] using the structure of hTrpRS in complex with TrpAMP [PDB code 2QUJ, (23)] as the search model. There are two monomers in an asymmetric unit forming a homodimer and the monomers were refined independently. In the initial difference Fourier maps, there are evident electron densities corresponding to a bound TrpAMP at each active site (Figure 1B). This is consistent with the previous results that in the presence of Trp and ATP, the Trp activation reaction took place in the crystallization solution, leading to the formation of TrpAMP (23). The initial structure refinement was carried out with the program CNS (24) following the standard protocols and the model building was performed manually with the help of programs COOT (25) and O (26). The final structure refinement was performed using the maximum likelihood algorithm implemented in the program REFMAC5 (27). Throughout the refinement, a free R-factor monitor calculated with 5% of randomly chosen reflections and a bulk solvent correction

Figure 1. Structure of pTrpRS in complex with TrpAMP. (A) Overall structure of the pTrpRS–TrpAMP complex. There are two pTrpRS–TrpAMP complexes in an asymmetric unit forming a homodimer. For clarity, only one monomer is shown with the N-terminal domain in yellow, the RF catalytic domain in cyan and the C-terminal domain in green. The four characteristic motifs, namely the N-terminal β-hairpin, AIDQ motif, HIGH motif and KMSAS loop are marked and colored in orange, purple, red and violet, respectively. The bound TrpAMP molecules are shown in ball-and-stick models. (B) A representative SIGMAA-weighted 2Fo-Fc composite omit map (1σ contour level) for the bound TrpAMP. (C) Structural comparisons of pTrpRS with hTrpRS (left panel) and bTrpRS (right panel) based on a superposition of the core region of the RF domain (equivalent to residues 74–229 of pTrpRS). The structures of the pTrpRS–TrpAMP, hTrpRS–TrpAMP (PDB code 2QUJ) and bTrpRS–TrpAMP (PDB code 1I6M) complexes are shown in green, orange and magenta, respectively. (D) A stereoview showing interactions of the bound TrpAMP (yellow) with the surrounding residues (green) at the active site. The hydrogen-bonding interactions are indicated with thin dashed lines. (E) Sequence alignment of the insertion region of TrpRSs from different species representing all three domains Bacteria, Archaea and Eucarya. The insertion region (equivalent to residues 204–209 of pTrpRS) is labeled and marked with a red line. The sequence alignment is generated by ESPript (47) with the secondary structures of bTrpRS and hTrpRS at the top and bottom of the alignment, respectively. The invariant residues are highlighted in shaded red boxes while the conserved ones in open red boxes.
were applied. The summary of statistics of the diffraction data and the structure refinement is listed in Table 1.

Structural alignment and phylogenetic analysis

Among the TrpRS and TyrRS structures available in the RCSB Protein Data Bank, there are multiple structures for a given enzyme in which the enzyme is in apo form or in complexes with various ligands. The previous structural studies have shown that both TrpRS and TyrRS undergo conformational changes upon ligand binding (6,7,19,23,28–33). For example, upon tryptophan binding, hTrpRS transforms from an open conformation of the unliganded form to a semi-closed conformation with the closure of the AIDQ motif and the KMSAS loop toward the active site followed by the rotation of the N- and C-terminal domains towards the Rossmann fold (RF) catalytic domain (23). In the presence of TrpAMP or TrpNH₂O (a tryptophan analog) and ATP, the enzyme takes a closed conformation mainly with a further movement of the KMSAS loop toward the active site when compared with the semi-closed conformation (23). For bacterial *Bacillus stearothermophilus* TrpRS (bTrpRS), the enzyme converts from an open conformation in the unliganded state or in complex with Trp or ATP alone to a closed conformation in the pre-transition (in complex with TrpNH₂O and ATP) and post-transition (in complex with AQP) states and further to a distinct closed conformation in the product state (in complex with TrpAMP) (6,28–31). For TyrRS, upon ligand binding, the overall structures of the enzymes remain similar and the conformational changes occur mainly at the KMSKS loop which rearranges during the reaction (19,32,33). For instance, depending on the binding of ligands, the KFGKT loop of the bacterial TyrRSs (equivalent to the...
KMSKS signature of TrpRSs) may take an open conformation (in the tyrosine-bound complexes), a semi-open conformation (in the TyrAMP/TyrAMS-bound complexes) or a closed conformation (in the presence of tyrosinol and ATP) (19,32,33). To ensure equal weight of the species in the structural alignment to minimize the effect of conformational differences on the phylogenetic analysis, one structure of TrpRS/TyrRS was selected for each species and in particular, for a given enzyme that has multiple structures, the structure that is the most similar to that of the pTrpRS–TrpAMP complex is selected. In total, 6 TrpRS structures and 10 TyrRS structures were chosen in which the enzyme is alloyed with either TrpAMP or Trp/Tyr. The chosen structures in complexes with TrpAMP from B. stearothermophilus (PDB code 1I6K), Mycoplasma pneumoniae (PDB code 2YY5), H. sapiens (PDB code 2QUJ) and P. horikoshii (reported herein), and with tryptophan from Deinococcus radiodurans (PDB code 1YI8) and Thermotoga maritima (PDB code 2G36). The chosen structures of TyrRSs include those of TyrRSs in complexes with TyrAMP or TyrAMS from B. stearothermophilus (PDB code 3TSI), E. coli (PDB code 1VBM), Saccharomyces cerevisiae (PDB code 2DLC) and human mitochondria (PDB code 2PID), with tyrosine or tyrosinol from P. horikoshii (PDB code 2CYC), Methanococcus jannaschii (PDB code 1J1U), Archeoglobus fulgidus (PDB code 2CYB), Thermus thermophilus (PDB code 1H3F) and H. sapiens (PDB code 1Q11), and with a Tyr analog from Staphylococcus aureus (PDB code 1J1J).

Selection of appropriate region(s) is a key step for reliable structure-based phylogenetic analyses of aaRSs. Discrepancies may arise from incompleteness or dispersion of the crystal structures due to deletion, invisibility and/or significant conformational differences of the N- and C-termini or flexible regions, and from possible posterior evolution of the anticodon recognition domain and the N-terminal domain which are less conserved than the catalytic domain. Therefore, in our structural alignment, we selected only the core region of the conserved RF domain (corresponding to residues 71–257 of pTrpRS) which is structurally conserved and adopts similar conformations in all of the selected structures. The region is almost identical to that used in the sequence-based phylogenetic study by Brown et al. (10) (equivalent to residues 71–257 of pTrpRS) except that the KMSKS loop which adopts different conformations depending on the binding of different ligands (see above) was not included in our study. The sequence identities of this region between TrpRSs and TyrRSs remain below the twilight zone threshold (13), again underscoring the advantage of the utilization of the structural alignment method in our study. Similarly, only the RF domains were aligned in the structure-based phylogenetic study of class I aaRSs by O’Donoghue and Luthey-Schulten (17) although the exact region was not specified.

The structural alignment was carried out using the multiple structural alignment program STAMP integrated in the molecular visualization program VMD, version 1.8.5 (34) with the parameters npass = 2, scanscore = 6 and scanslide = 5. Protein homology was assessed with the statistical similarity measure $Q_H$ which was adapted by O’Donoghue et al. (17) on the basis of the original measure $Q$ (35) to include the effects of the gaps on the aligned portion. $Q_H$ ranges from 0 to 1, where $Q_H = 1$ means that the two proteins are identical. A distance matrix of pairwise structure dissimilarity value (1–$Q_H$) was generated (Table 2) and used as input for phylogenetic analyses with the UPGMA method using software MultiSeq in VMD (36), and with the NJ algorithm and the minimum evolution (ME) method, respectively, using program Mega4 (37).

RESULTS

Structure of the pTrpRS–TrpAMP complex

The crystal structure of the pTrpRS–TrpAMP complex was determined at 3.0 Å resolution with an $R$-factor of 23.8% and a free $R$-factor of 25.8% (Table 1). The asymmetric unit contains two pTrpRS molecules each bound with a TrpAMP at the active site (Figure 1A). The pTrpRS consists of three typical domains: an N-terminal domain (residues 1–68), an RF catalytic domain (residues 69–246 and 358–386) and a C-terminal $\alpha$-helical domain (residues 247–357) [hereafter the nomenclature of the secondary structures of pTrpRS is after Yang et al. (38)]. Most of the residues are well defined with good electron density except the N-terminal three residues and residues 281–300.
Table 2. Distance (1-OH) matrix for UPGMA, NJ and ME dendrogram of subclass Ic aaRSs

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*Designation of the enzymes is the same as in Figure 2.

Superposition of the structure of the pTrpRS–TrpAMP complex to the corresponding structures of eukaryotic hTrpRS (23) and bacterial bTrpRS (29) yields root mean square deviations of 1.4 Å and 2.6 Å, respectively, for all Ca atoms (Figure 1C), indicating that the overall structure of pTrpRS resembles more to hTrpRS than bTrpRS, which is consistent with its higher sequence similarity with hTrpRS than with bTrpRS (44% versus 23%). In addition, the archaeal pTrpRS has an N-terminal domain which is approximately as long as that of the T1 form hTrpRS (39), whereas this domain is absent in bTrpRS. This domain contains a β-hairpin (residues 5–13) (hereafter the residues of aaRSs from P. horikoshii, H. sapiens and B. stearothermophilus are indicated with superscripted letters P, H and B, respectively) and the equivalent β-hairpin of hTrpRS has been shown to be important to the ATP binding (23,38). Within the β-hairpin a salt bridge between Lys66 and Glu111P (equivalent to Phe84H and Thr89H) appears to stabilize the region in the extreme environment of high temperature.

TrpAMP is bound at the active site which is composed by key structural elements including the KMSAS, HIGH and AIDQ motifs. The amino acid specificity of pTrpRS is mainly determined by residues Gln111P and Tyr77P (equivalent to Gln194H and Tyr159H, respectively). The nitrogen of the indole ring of the tryptophanyl moiety forms hydrogen bonds with the side-chain carboxyl oxygen of Gln111P (3.2 Å) and the hydroxyl oxygen of Tyr77P (2.8 Å) (Figure 1D). Additionally, the hydroxyl of the indole nitrogen is stabilized by the phenol group of Tyr77P via π–π stacking interaction. The amino group of the tryptophanyl moiety is bound by Glu116P (equivalent to Glu199H) via a salt bridge and Gln198P (equivalent to Gln284H) via a hydrogen bond (3.0 Å) through their side-chains, while the carbonyl group interacts with the side-chain nitrogen of Lys117P (equivalent to Lys200H) (3.1 Å).

The binding mode of the tryptophanyl moiety by archaeal pTrpRS is quite similar to that by eukaryotic hTrpRS, and together the archaeal and eukaryotic TrpRSs show great differences from bacterial bTrpRS. In the bTrpRS–TrpAMP complex, the indole nitrogen of the tryptophanyl moiety is recognized by Asp132B via a hydrogen bond and Phe5B via hydrophobic interaction (29). In addition, the amino group and the carbonyl group are bound by Tyr125B and Gln9P, respectively. Among the residues of bTrpRS participating in the Trp binding, only Phe5B has similar physicochemical properties as its equivalent in archaeal/eukaryotic TrpRSs (Tyr77P/Tyr159H) although the hydrogen-bonding interaction between the indole nitrogen and Tyr77P/Tyr159H is absent in the bTrpRS complex. In addition, the interaction between the amino group and Gln198H/Gln284H observed in the pTrpRS/hTrpRS complexes is also missing in the bTrpRS complex.

For the AMP group, the N6 atom of the adenosine moiety of TrpAMP is recognized by the main-chain carbonyl groups of Val244B and Met254B of the KMSAS motif. The ribose group is positioned by the AIDQ motif with the 2’-hydroxyl group interacting with the side-chain carboxylate of Asp216B and both the 2’-hydroxyl and 3’-hydroxyl groups interacting with the main-chain nitrogen of Ala214B. Considering that the AIDQ motif is highly conserved in Archaea/Eucarya and the corresponding GEDQ motif of bTrpRS binds to AMP with equivalent interactions (29), the binding mode of AMP seems to be conserved in all TrpRSs. As for the α-phosphate, Lys195B of the KMSKS loop in bTrpRS binds to it; however, due to the lack of a lysine at the equivalent position, the moiety is bound by the side-chain amino of Arg80P/Arg162H and the main-chain amide of Gly81P/Gly163H on a distal strand β5.

Although the overall structure and the TrpAMP binding mode of pTrpRS are more similar to those of hTrpRS than to bTrpRS, the dimer interface of pTrpRS resembles more to that of bTrpRS. The dimer interface of pTrpRS buries 1877 Å² (or 10%) solvent-accessible surface of each monomer and involves five α-helices (α8 and α10-α13) of the RF domain and the C-terminus. Compared with pTrpRS and bTrpRS, hTrpRS has an insertion (residues 290–305) in the RF domain (Figure 1E) that forms η4 and ζ14 and blocks the extension of the C-terminus to the dimerization interface. The equivalent regions of pTrpRS (residues 204–209) and bTrpRS (residues 135–139) are much shorter and the C-terminus...
can extend towards the other subunit, which substantially increases the dimer interface (1877 Å² in pTrpRS/bTrpRS versus 1501 Å² in hTrpRS). It is also noteworthy that, compared with hTrpRS or bTrpRS, pTrpRS contains four additional pairs of inter-molecular salt bridges at the dimer interface formed between Glu163P and Lys130P and between Glu185P and Lys178P, respectively. As the formation of more salt bridges is one of the features of thermo-stable proteins, these additional salt bridges may contribute to the hyperthermostability of pTrpRS.

**Structural alignment of TrpRS and TyrRS**

Protein structures are more conserved than sequences and thus contain evolutionary information. Comparative studies of different structures of several enzymes have been carried out to investigate their evolutionary paths (17,40). Previously, the evolutionary relationship between TrpRS and TyrRS was examined using a structural alignment method by O’Donoghue et al. (17). In their work, different algorithms rendered conflicting results about the branching of archaeal/eukaryotic TyrRS: in the UPGMA tree, archaeal/eukaryotic TyrRS (represented by *H. sapiens* TyrRS) clustered with its bacterial counterparts, while in the NJ tree it grouped with the enzymes with specificity to tryptophan. The discrepancy was attributed to the unavailability of a crystal structure of TrpRS from Archaea/Eukarya at that time, which could cause errors arising from ‘attractor effects’ between long uninterrupted branches (41). With the structure of pTrpRS presented here, we can now perform a more complete study to get a more reliable picture of the evolutionary history of TrpRS and TyrRS.

To prevent discrepancies arising from incompleteness or dispersion of the crystal structures and to avoid bias caused by posterior evolution of certain regions, we selected the core region of the RF domain for the structural alignment (for details see ‘Materials and Methods’ section). In our study, this region can be accurately aligned and the gaps and insertions can be unambiguously identified (Figure 2A). In contrast, in a previous study that was based on sequence alignment of a similar region alone without utilization of the structural information, certain residues/regions were not properly aligned (10). For example, a structurally conserved Gln residue corresponding to Gln101P on β6 of pTrpRS (Figure 2A) which participates in the interactions of β6 with several other structural elements of the RF domain, is incorrectly aligned in the sequence-based study (10). In addition, a part of an α-helix in the archaeal/eukaryotic TyrRSs (corresponding to residues Pro86 to Leu92 of human TyrRS) which should be aligned to its equivalent helix (corresponding to residues Thr93 to Glu99 of *E. coli* TyrRS) (Figure 2A), was mistakenly aligned to a loop in the bacterial TyrRSs (corresponding to residues Ala86 to Asn92 of *E. coli* TyrRS) (10). Thus, the missing of an equivalent loop in the archaeal/eukaryotic TyrRSs (indicated by a gap in Figure 2A) was not detected. In a later phylogenetic study based on multiple sequence alignment, the structural information was applied to adjust the sequence alignment of *B. stearothermophilus* TrpRS and TyrRS and subsequently all other sequences (11). The utilization of the structural information obviously improved the quality and reliability of the multiple sequence alignment in that the aforementioned Gln residue became properly aligned. However, the gap was still undistinguished, which is possibly limited by the availability of only two crystal structures of TrpRS and TyrRS (those from *B. stearothermophilus*) at that time. Intriguingly, the application of different sequence alignment methods (10,11) in the two phylogenetic studies yielded inconsistent results about the classification of TrpRSs and TyrRSs, with the latter in agreement with our result (see ‘Discussion’ section later), indicating that whether structural information is considered and introduced in the alignments may account for the divergence in the positioning of archaeal/eukaryotic TyrRSs in the evolutionary trees.

**Structure-based phylogenetic analysis of TyrRS and TrpRS**

With the inclusion of the pTrpRS structure and the other recently reported TrpRS and TyrRS structures, the structure-based evolutionary trees were generated with the UPGMA and NJ algorithms which have been used by O’Donoghue and Luthey-Schulten (17), and additionally the ME algorithm which is commonly used for distance-based analyses. Although the UPGMA algorithm assumes the molecular clock hypothesis, the phylogenetic trees calculated with the UPGMA, ME and NJ algorithms, respectively, show a congruent topology (Figure 2B). The root of the TrpRS tree separates the archaeal/eukaryotic TyrRSs and their bacterial counterparts. The archaeal/eukaryotic TyrRSs group with TrpRSs without ambiguity, which is in accord with the observations by both Ribas de Pouplana et al. (9) and Diaz-Lazcoz et al. (11) that the archaeal and eukaryotic TyrRSs resemble more to TrpRSs than their bacterial counterparts (see ‘Discussion’ section later).

When examined individually, TrpRSs conform to the full canonical pattern (Figure 2B). According to Woese et al. (2), at least six distinct subgroups can be identified within the bacterial genre (2). Here, only four structures of TrpRSs from bacterial species are available and included in the analysis, representing four subgroups of the bacterial genre. In our evolutionary trees, TrpRSs from *B. stearothermophilus* [closely related to *B. subtilis* whose sequence was included in the work by Woese et al. (2)] and *M. pneumoniae* cluster together while TrpRS from *T. maritima* is more closely related to that from *D. radiodurans*, consistent with the evolutionary relationships of the subtypes revealed by Woese et al. (2). Although the bacterial TrpRSs show great divergence, they form a group distinct from the archaeal and eukaryote TrpRSs represented by the enzymes from *P. horikoshii* and *H. sapiens*, respectively, suggesting that the division between bacterial TrpRSs and archaeal/eukaryotic TrpRSs occurred at an early stage of bacteria evolution.

For TyrRSs, the generated evolutionary trees essentially have the same topology as the previously reported sequence-based trees (2,42), which all strongly support
the distinct separation of Bacteria, Archaea and Eucarya (Figure 2B). In the bacterial genre, TyrRSs are divided into two far related subtypes, namely TyrRS and TyrRZ, with TyrRSs from *B. stearothermophilus*, *E. coli* and *S. aureus* belonging to the TyrRS subgroup and that from *T. thermophilus* belonging to the other. These results support the notion that bacterial TyrRSs can be divided into TyrRS and TyrRZ, which was originally indicated by the sequence-based phylogenetic studies (43,44). Additionally, human mitochondrial TyrRS is more similar to the bacterial enzymes of the first subtype. TyrRSs from *A. fulgidus* and *M. jannaschii* cluster with eukaryotic TyrRSs, while that from *P. horikoshii* forms a separate branch, although they are all from Euryarchaeota. Considering that *P. horikoshii* TyrRS clusters with TyrRSs from plants in the sequence-based trees (2), our results suggest that TyrRSs are partially intermixed in the archaeal and eukaryotic genre. To verify this notion, further study is needed to include more structures of eukaryotic TyrRSs especially those from plants.

**DISCUSSION**

As the sequence identity between TrpRS and TyrRS is below the twilight zone threshold (13), the connections of the two enzymes suggested by the previous sequence-based phylogenetic studies are controversial. For proteins...
sharing low sequence identity, 3D structures are better than primary sequences for modeling of protein evolution (15–17). In particular, despite the low sequence homology, TyrRS and TrpRS share high structure similarity, supporting the use of the structural alignment method for phylogenetic studies of these enzymes. However, due to the limitation of available crystal structures of TrpRS and TyrRS, the structure-based study of aaRSs by O’Donoghue and Luthey-Schulten (17) yielded conflicting results and thus was unsuccessful to give a conclusive answer regarding the evolutionary relationship between TrpRS and TyrRS (17). In our study, we demonstrate the advantage of the structural alignment method over the sequence alignment method by showing that the residues are correctly aligned and the insertions and gaps are unambiguously identified (Figure 2A).
In addition, we achieve consistent results as shown by the congruent topology of the generated phylogenetic trees (Figure 2B). Therefore, our results are more accurate and valuable to discern the evolution history of the two enzymes. This strategy can also be applied to examine the relationships between other aaRSs as the sequence identities of aaRSs with different specificities are typically below the twilight zone threshold (17).

Analysis of a set of paralogs of aaRSs has suggested that generally aaRSs form monophyletic groups regarding their amino acid specificities, implying that the enzymes appeared prior to the separation of the three kingdoms (2). However, several exceptions exist: asparaginyl-tRNA synthetase (AsnRS) and glutaminyl-tRNA synthetase (GlnRS) are suggested to arise from the archaean genre of aspartyl-tRNA synthetase (AspRS) (2,45) and the eukaryotic lineage of glutamyl-tRNA synthetase (GluRS) (2,46), respectively. In the structural dendograms presented in Figure 2B, the root of TrpRS is located in the archaean branch of TyrRS, making the TyrRS group paraphyletic and thus breaking the monophyletic rule. These results indicate that similar to AsnRS, TrpRS originates from an archaean lineage. Despite the same archaean origin, the evolutionary path of TrpRS exhibits some differences from that of AsnRS. The branching between archaean AsnRS and archaean AspRS is relatively long, suggesting that the appearance of archaean AsnRS and the subsequent acquisition of bacterial AsnRS occurred at late stages, which explains the absence of AsnRS in most archaean and some of the bacterial taxa (2). In contrast, the relatively short distance between TrpRS and archaean TyrRS and that between bacterial TrpRS and archaean TrpRS suggest that TrpRS had already emerged right after the division between Bacteria and Archaea and soon horizontally transferred to the bacterial genre, which is also supported by the fact that TrpRS is widely distributed in all three kingdoms. The transfer of TrpRS from Archaea into Bacteria is consistent with the notion by Woese et al. (2) that the horizontal gene transfer of aaRSs between Archaea and Bacteria appears to be asymmetric as the synthetases were transferred only from Archaea to Bacteria, but not the reverse. On the other hand, the close occurrences of the two events (appearance of TrpRS and the acquisition by the bacterial genre) might be another reason for the divergence of the positioning of archaean TrpRS in the previous evolutionary studies (9–12,17).

Collectively, our data imply that before the division of Bacteria and Archaea, the ancestor TyrRS had existed, whereas no aminoacyl synthetase ancestor solely with a Trp specificity was present. After the division between Bacteria and Archaea, the ancestor of TyrRSs diverged to the archaean version and the bacterial version, and soon TrpRSs evolved from the archaean lineage of TyrRS probably through gene duplication, followed by the early acquisition of TrpRSs by Bacteria through horizontal gene transfer.

Protein Data Bank accession code

The structure of trytophanyl-tRNA synthetase from *P. horikoshii* in complex with TrpAMP has been deposited.
with the RCSB Protein Data Bank under accession code 3JXE.

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