Alanyl-tRNA synthetase genes of *Vanderwaltozyma polyspora* arose from duplication of a dual-functional predecessor of mitochondrial origin

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Received June 14, 2011; Revised August 17, 2011; Accepted August 22, 2011

ABSTRACT

In eukaryotes, the cytoplasmic and mitochondrial forms of a given aminoacyl-tRNA synthetase (aaRS) are typically encoded by two orthologous nuclear genes, one of eukaryotic origin and the other of mitochondrial origin. We herein report a novel scenario of aaRS evolution in yeast. While all other yeast species studied possess a single nuclear gene encoding both forms of alanyl-tRNA synthetase (AlaRS), *Vanderwaltozyma polyspora*, a yeast species descended from the same whole-genome duplication event as *Saccharomyces cerevisiae*, contains two distinct nuclear AlaRS genes, one specifying the cytoplasmic form and the other its mitochondrial counterpart. The protein sequences of these two isoforms are very similar to each other. The isoforms are actively expressed in vivo and are exclusively localized in their respective cellular compartments. Despite the presence of a promising AUG initiator candidate, the gene encoding the mitochondrial form is actually initiated from upstream non-AUG codons.

A phylogenetic analysis further revealed that all yeast AlaRS genes, including those in *V. polyspora*, are of mitochondrial origin. These findings underscore the possibility that contemporary AlaRS genes in *V. polyspora* arose relatively recently from duplication of a dual-functional predecessor of mitochondrial origin.

INTRODUCTION

Faithful decoding of mRNA into protein depends on accurate aminoacylation of tRNA by aminoacyl-tRNA synthetases (aaRSs) and specific codon/anticodon base pairing. AaRSs are a group of structurally diverse enzymes responsible for protein translation, each of which couples a specific amino acid to its cognate tRNA. The resultant aminoacyl-tRNAs are then delivered by elongation factor (EF)-1 to ribosomes for protein synthesis. In prokaryotes, there are typically 20 aaRSs, one for each amino acid (1–3). In eukaryotes, protein synthesis occurs in the cytoplasm and organelles, such as mitochondria and chloroplasts (4). Thus, eukaryotes, such as yeast, commonly have two genes that encode distinct sets of proteins for each aminoacylation activity, one localized in the cytoplasm and the other in mitochondria. Each set aminoacylates the tRNAs within its respective cellular compartment, and is sequestered from isoacceptors confined to other cellular compartments.

Cytoplasmic and mitochondrial forms of an aaRS are commonly encoded by two nuclear genes of distinct evolutionary origins in eukaryotes (5). Typically, the gene that encodes the cytoplasmic form is of eukaryotic (or archaean) origin, while the gene that encodes its mitochondrial counterpart is of prokaryotic origin. However, in some cases, a single nuclear gene encodes both forms of a given aaRS through alternative initiation of translation, while its orthologue was lost during evolution. Examples of this type include yeast *ALA1* [which encodes alanyl-tRNA synthetase (AlaRS)] (6), *GRS1* (which encodes glycyl-tRNA synthetase) (7), *HTS1* (which encodes histidyl-tRNA synthetase) (8) and *VAS1* (which encodes valyl-tRNA synthetase) (9). Since isoforms target different cellular compartments, they cannot be substituted for one another in vivo. A similar phenotype was observed for genes encoding cytoplasmic and mitochondrial forms of *Arabidopsis thaliana* AlaRS, threonyl-tRNA synthetase and valyl-tRNA synthetase (10). With the exception of a few controversial incidents caused by genetic recombinations or horizontal gene transfer, the evolutionary scenarios of eukaryotic aaRSs largely fit in one of the two paradigms described above. Nevertheless, it was recently reported that two paralogous valyl-tRNA synthetase genes of mitochondrial origin coexist in the fission yeast *Schizosaccharomyces pombe*,...
one that encodes the cytoplasmic form and the other its mitochondrial counterpart (11).

Many examples of non-AUG initiation were reported in higher eukaryotes, where cellular and viral mRNAs initiate from codons that differ from AUG by one nucleotide, such as ACG, UUG and AUU (12). However, only a few cases were found to date in low eukaryotes such as yeast. ALA1 and GR5 are among the first genes shown to use naturally occurring non-AUG triplets as an alternative translation initiator in the model yeast S. cerevisiae (6,7). While sequence context has little effect on the translation-  
al efficiency of AUG initiation in yeast (13–15), translation initiation from a non-AUG codon is profoundly affected (by up to 32-fold) by nucleotides at relative positions –3 to –1, especially –3. AARuug (R denotes A or G; uug denotes a non-AUG initiation codon) appears to represent the most favorable sequence context (16,17). An interesting feature of the expression of ALA1 is that this gene contains two consecutive ACG initiator codons for its mitochondrial form (6). Consecutive ACG codons possess stronger initiation activity than does a single ACG codon (18). This feature of remedial initiation from repeated non-AUG initiator codons per se may represent a novel mechanism to improve the efficiency of translation initiation. Examples of non-AUG initiation were also recently reported in other yeast species (11,19).

As aaRSs are one of the most ancient groups of enzymes that continue to function in all free-living organisms, they are often targeted for taxonomic and phylogenetic studies (20–22). In this report, we explored the functions and historical relationships of AlaRSs from 13 different yeast species. It is our hope that results obtained in this study will advance our understanding of the biochemical properties of the eukaryotic aaRS gene, and provide new insights into its evolutionary footprints. Results presented herein show that while all other yeast species studied possess a single AlaRS gene with dual functions, Vanderwartzyma polyspora, a close relative of S. cerevisiae, contains two distinct AlaRS genes that respectively encode the cytoplasmic and mitochondrial forms. Unexpectedly, all yeast AlaRS genes are of the mitochondrial type, irrespective of the cellular compartments where the protein products are active. These and other results imply that AlaRS genes in V. polyspora underwent a major genetic recombination at least twice. First, the mitochondrion-type gene gained cytoplasmic function and became a dual-functional gene, while its orthologue was lost. Later, the dual-functional gene of mitochondrial origin was duplicated into two copies, each retaining a single function (cytoplasmic or mitochondrial).

MATERIALS AND METHODS

Construction of plasmids

Cloning of ALA1 genes of S. cerevisiae and Candida albicans was previously described (6,23). To clone the ALA1 gene of V. polyspora (VpALA1) into pRS315-2xFLAG (a low-copy-number yeast shuttle vector carrying a LEU2 marker and a 2xFLAG sequence), a pair of gene-specific primers was used to amplify the gene via a polymerase chain reaction (PCR) using yeast genomic DNA as the template. The forward primer with an EagI site is located 300 bp upstream of the first ATG codon of the open reading frame (ORF), while the reverse primer with an XhoI site is located immediately upstream of the stop codon. The ~3.3-kb PCR-amplified DNA fragment was subsequently digested with EagI and XhoI prior to cloning into pRS315. Cloning of VpALA2 and SpALA1 into pRS315 followed a similar protocol. Note that these clones were expressed under the control of their respective native promoters.

Cloning of SpALA1 into pADH (a high-copy-number yeast shuttle vector with a LEU2 marker, an ADH promoter preceding the multiple cloning sites, and a 2xFLAG sequence following the multiple cloning sites) (24) followed a similar protocol, except that the forward primer was annealed to a sequence 12 bp upstream of the first ATG codon. Note that this clone was expressed under the control of a constitutive ADH promoter. Construction of initiator mutants of VpALA1 and VpALA2 followed a strategy described earlier (7). Mutagenesis was carried out following a protocol provided by the manufacturer (Strategene, La Jolla, CA, USA). Green fluorescence protein (GFP) assays followed a protocol described elsewhere (25).

Complementation assays for cytoplasmic AlaRS activity

The yeast ALA1 knockout strain, TRY11 (MATa, his3Δ200, leu2Δ1, lys2-801, trp1Δ101, ura3Δ52 and ala1Δ::TRP1) was maintained by a plasmid carrying the wild-type ALA1 gene and a URA3 marker (26). Complementation assays of cytoplasmic AlaRS activity were carried out by introducing a test plasmid carrying the gene of interest and a LEU2 marker into TRY11, and the ability of the transformants to grow in the presence of 5-FOA was determined. Starting from a cell density of 4.0 A600, cell cultures were 5-fold serially diluted, and 10-μl aliquots of each dilution were spotted onto designated plates containing 5-FOA. Plates were incubated at 30°C for 3–5 days. The transformants evicted the maintenance plasmid with a URA3 marker in the presence of 5-FOA, and thus could not grow on the selection medium unless a functional cytoplasmic AlaRS was encoded by the test plasmid.

Complementation assays for mitochondrial AlaRS activity

TRY11 was co-transformed with a test plasmid and a second maintenance plasmid (carrying a HIS3 marker) that only expresses the cytoplasmic form of AlaRS (due to a mutation in the non-AUG initiator codon). In the presence of 5-FOA, the first maintenance plasmid (carrying a URA3 marker) was evicted from the co-transformants, while the second maintenance plasmid was retained. Thus, all co-transformants survived 5-FOA selection due to the presence of the cytoplasmic AlaRS derived from the second maintenance plasmid. The mitochondrial phenotypes of the co-transformants were further tested on YPG plates at 30°C, with results
documented on Day 3 following plating. Because a yeast cell cannot survive on glycerol without functional mitochondria, the co-transformants did not grow on YPG plates unless a functional mitochondrial AlaRS was generated from the plasmid.

Reverse transcription—PCR

To determine the relative levels of endogenous \( ALA1 \) and \( ALA2 \) mRNAs in \( V. polyspora \), a semiquantitative reverse transcription (RT)–PCR experiment was carried out following the protocols provided by the manufacturer (Invitrogen, Carlsbad, CA, USA). Total RNA was first isolated from the transformant, and then treated with DNase to remove contaminating DNA. Aliquots (~1 μg) of the RNA were then reverse-transcribed into single-stranded complementary (c)DNA using an oligo-dT primer. After RNase H treatment, single-stranded cDNA products were amplified by a PCR using a pair of gene-specific primers. The forward and reverse primers (GSP1 and GSP2) for \( VpALA1 \) contained sequences complementary to nucleotides 1936–1961 (5'-CG TAATGTCGAAGAAGTCTGTAACC-3') and nucleotides 2379–2405 (5'-TATTAGACAAATGGATAGTT GACC-3') of \( VpALA1 \), respectively. The forward and reverse primers (GSP3 and G-SP4) for \( VpALA2 \) contained sequences complementary to nucleotides 1966–1992 (5'-TGATGTCAGTAAAGAACCTGTC-3') and nucleotides 2320–2346 (5'-GTCGAGAATGGTGATGAGT TCAG-3') of \( VpALA2 \), respectively. To obtain more-reliable data, four different cycle numbers of PCR were carried out for each cDNA preparation as indicated in the figure.

A quantitative real-time RT–PCR experiment was subsequently used to obtain more-accurate data. Two sets of primers, GSP5/GSP6 and GSP7/GSP8, were respectively used to amplify the cDNA fragments of \( VpALA1 \) and \( VpALA2 \). GSP5 and GSP6 are complementary to nucleotides +1998 to +2022 and +2111 to +2135 of \( VpALA1 \), respectively, while GSP7 and GSP8 are complementary to nucleotides +2004 to +2028 and +2016 to +2040 of \( VpALA2 \), respectively. Quantitative data were obtained from three independent experiments and averaged.

Western blot analysis

Protein expression patterns of the \( ALA1 \) and \( ALA2 \) constructs were determined by a chemiluminescence-based western blot analysis. INVSc1 (\( MATa \), \( his3A1 \), \( leu2 \), \( trp1-289 \), \( ura3-52 \); \( MATa \), \( his3A1 \), \( leu2 \), \( trp1-289 \), \( ura3-52 \) (Invitrogen) was first transformed with the constructs of interest, and total protein extracts were prepared from the transformants. Aliquots of protein extracts (40 μg) were loaded onto a mini gel (of 8 x 10 cm) containing 10% polyacrylamide and electrophoresed at 100 V for 1–2 h. Following electrophoresis, the resolved proteins were transferred using a semi-dry transfer device to a polyvinyliene fluoride (PVDF) membrane in buffer containing 30 mM glycine, 48 mM Tris base (pH 8.3), 0.037% sodium dodecysulfate (SDS) and 20% methanol. The membrane was probed with an anti-FLAG tag antibody (Sigma) followed by an HRP-conjugated goat anti-mouse IgG antibody (Invitrogen), and then exposed to X-ray film following the addition of appropriate substrates.

RESULTS

Vanderwaltozyma polyspora possesses two distinct nuclear AlaRS genes

Previous studies showed that the \( ALA1 \) gene of \( S. cerevisiae \) encodes both mitochondrial and cytoplasmic forms of AlaRS (6, 23). We wondered whether a similar feature was conserved in the AlaRS genes of other yeast species. Pursuant to this objective, 14 different yeast AlaRS genes were retrieved from the EMBL database, and the predicted amino acid sequences were analyzed. Except for \( V. polyspora \), which contained two distinct nuclear AlaRS genes (designated herein as \( VpALA1 \) and \( VpALA2 \)), all yeast species studied possessed a single AlaRS gene (designated herein as \( ALA1 \)) (Figure 1). As aminoaacylation activities are required for protein synthesis in both the cytoplasm and mitochondria, the single AlaRS gene was expected to provide both cytoplasmic and mitochondrial functions in these yeast species. Indeed, two in-frame initiator candidates were found in each of the ORFs (Figure 1A). Despite only one ATG initiator codon being found in \( ALA1 \) genes of \( Ashbya gossypii \), \( C. glabrata \) and \( Kluyveromyces lactis \), a promising non-ATG initiator candidate was mapped in the leader sequence of these genes (Figure 1A). In agreement with this tentative assignment, the N-terminal pre-sequences deduced from these genes were all rich in positively charged and hydroxylated amino acid residues but were devoid of acidic residues, characteristic of a mitochondrial targeting signal.

Despite \( V. polyspora \) containing two distinct nuclear AlaRS genes, neither of the protein forms deduced from the ORFs of these two genes (starting from the ATG1 codon) possessed a canonical N-terminal mitochondrial targeting signal. Sequence alignment further indicated that the first ATG codons of these two genes were both located at a position comparable to that of the ATG initiator codon for the cytoplasmic form of ScAlaRS (AlaRS of \( S. cerevisiae \)) (Figure 1A). A closer look at the 5'-end sequences of these two genes suggested that ATG1 was the only possible initiator codon for \( VpALA1 \), while \( VpALA2 \) contained several promising non-ATG initiator candidates 5' to ATG1, such as TTG(-28), ACG(-23) and TTG(-15) (Figure 1B). The protein forms deduced from the ORFs of these two genes considerably resembled each other (~66% identity), with \( VpAlaRS1 \) (AlaRS1 of \( V. polyspora \)) having a higher similarity to ScAlaRS (83% identity) (Figure 1C).

The top three base pairs in the acceptor stem are highly conserved in all yeast tRNAs\(^{Ala}\)

Previous studies showed that the major identity elements of tRNA\(^{Ala}\) reside in its acceptor stem. In particular, a G3:U70 base pair is conserved in the acceptor stems of all known tRNAs\(^{Ala}\) from prokaryotes, archaea, eu- karyote cytoplasm and chloroplasts (27, 28). Comparison of acceptor stems of tRNAs\(^{Ala}\) from various yeast species
indicated that G3:U70, G1:C72 and G2:C71 are strictly preferred G4:C69, U5:A68, U6:A67 and A7:U66. While the mitochondrial counterparts moderately preferred G4:C69, U5:A68, U6:A67 and A7:U66. This preference for particular base pairs in the bottom part of the acceptor stem suggests that while VpALA1 is exclusively initiated from upstream non-ATG codons.

Vanderwaltozyma polyspora ALA1 and ALA2 encode the cytoplasmic and mitochondrial forms of AlaRS, respectively

To further investigate the functional potential of these AlaRS genes, several representative sequences were cloned in pRS315 (a low-copy-number yeast shuttle vector), and their cross-species complementation activities were assayed in TRY11, an ala1 strain of S. cerevisiae. Consistent with previous reports, CaALA1 and ScALA1 rescued both the cytoplasmic and mitochondrial defects of the knockout strain (6,23). Transformants bearing either of these two constructs grew robustly on 5-FOA and YPG plates (Figure 3A–C, nos 1 and 2). In contrast, SpALA1, when cloned in the same vector pRS315, rescued only the mitochondrial defect of the knockout strain (Figure 3A–C, no. 3), presumably due to poor gene expression by its native promoter (Figure 3D). A dual-functional phenotype was observed for this gene when it was cloned into pADH, a high-copy number yeast shuttle vector with a constitutive ADH promoter (Figure 3A–C, no. 4). Thus, SpALA1 can recognize both cytoplasmic and mitochondrial tRNA\textsubscript{Ala} isoacceptors of S. cerevisiae. In contrast to the dual-functional nature of these genes, AlaRS genes from V. polyspora appeared to have single functions. VpALA1 and VpALA2, respectively, rescued the cytoplasmic and mitochondrial defects of TRY11, suggesting that these two genes respectively provide cytoplasmic and mitochondrial functions (Figure 3B and C, nos 5 and 6). The protein expression patterns of these genes were subsequently determined by western blotting. As shown in Figure 3D, all genes, except for SpALA1, were well expressed from their native promoters in S. cerevisiae. It appeared that the native promoter of SpALA1 cannot efficiently be recognized by the transcription apparatus of S. cerevisiae. In addition, the protein expression level of VpALA1 was much lower than that of VpALA2.

Vanderwaltozyma polyspora ALA2 is exclusively initiated from upstream non-ATG codons

As mentioned above, no N-terminal mitochondrial targeting signal was predicted for the ATG1-initiated protein sequence of VpALA2. How is this protein form targeted to mitochondria for its functioning? To verify whether this gene is actually initiated from ATG1, this codon was subsequently mutated, and the complementation activity of the resultant construct was tested. As a reference, the first ATG codon of VpALA1 was also mutated and tested. As expected, a mutation at ATG1 of VpALA1 drastically impaired its cytoplasmic activity. However, a mutation at ATG1 of VpALA2 had little effect on its mitochondrial activity (Figure 4A–C, nos 2 and 4). This result suggests that VpALA1 is initiated from ATG1, while VpALA2 is initiated elsewhere.

To obtain more clues into solving this mystery, upstream in-frame non-ATG codons that fit the following three criteria were preferentially selected as initiator
candidates and tested for their initiating activity. First, they had to differ from ATG by only one nucleotide. Second, they had to bear an ‘A’ nucleotide at relative position 3. Lastly, they had to possess the potential to initiate translation of a mitochondrial targeting signal devoid of acidic amino acid residues. The only candidate that perfectly matched all of these criteria was ACG.

Unexpectedly, a mutation at this candidate slightly reduced, but not eliminated, the mitochondrial activity of VpALA2 (Figure 4A–C, no. 5), suggesting the presence of an alternative initiation site. To gain further insights, a less likely candidate, TTG, that matched some of the criteria described above was also tested for its initiation activity. While a mutation at TTG alone had little effect on the mitochondrial activity of VpALA2, simultaneous mutations at ACG and TTG drastically impaired its mitochondrial activity (Figure 4A–C, nos. 6–7). Note that TTG contained a ‘C’ at relative position 3. Mainly for this reason, TTG was not selected as one of the primary targets in the first round. So, despite the presence of a promising ATG initiator candidate, VpALA2 was exclusively initiated from upstream non-ATG codons, with ACG being the major initiation site and TTG being the minor initiation site.

To verify the cellular localization of these two proteins in vivo, a DNA sequence encoding the GFP was inserted in-frame at the 3'-ends of these two genes, resulting in VpALA1-GFP and VpALA2-GFP. As genetic tools for V. polyspora are not yet fully developed, we herein used an S. cerevisiae strain, INVScl, as the platform for the localization assay. The GFP fusion constructs were individually transformed into INVScl, and the subcellular localization of these GFP fusion proteins in the transformants was examined under fluorescence microscopy. As shown in Figure 4D, VpAlaRS2-GFP exclusively existed in mitochondria (d–f), while VpAlaRS1-GFP was evenly distributed in the cytosol (a–c). This result provides further support to the notion that VpALA1 and VpALA2 encode the cytoplasmic and mitochondrial forms of AlaRS, respectively.

Both ALA1 and ALA2 are actively expressed in Vanderwaltozyma polyspora

To investigate whether VpALA1 and VpALA2 are actively expressed in their native host, V. polyspora, relative levels of their specific mRNAs were determined by a semiquantitative RT-PCR. Two sets of gene-specific primers were designed, one set complementary to VpALA1 and the other to VpALA2. The expected sizes of the PCR-amplified products were 469 and 380 base pairs for VpALA1 and VpALA2, respectively. As shown in Figure 5A, similar levels of VpALA1 and VpALA2 DNA fragments were amplified by their respective primers using the genomic DNA of V. polyspora as a template (upper panel), suggesting that these two sets of primers were annealed to their respective target genes with similar efficiencies under the conditions used. The same sets of primers were subsequently used to amplify the cDNA of these two genes, which had been reverse-transcribed from a total RNA preparation of V. polyspora using oligo-dT as the primer. Figure 5A shows that similar levels of VpALA1 and VpALA2 DNA fragments were amplified using the reverse-transcribed cDNA as a template (lower panel), suggesting that both genes are actively expressed in V. polyspora and are transcribed at comparable efficiencies. Further determination using a quantitative (q)RT–PCR shows that VpALA1 had a transcriptional efficiency two-fold higher than that of VpALA2 (Figure 5B). Note that these two sets of primers were annealed to their respective target genes.
with an almost identical efficiency using the genomic DNA of V. polyspora as a template.

To test the substrate specificities of VpAlaRS1 and VpAlaRS2, recombinant AlaRS-His6 enzymes were purified from yeast transformants harboring appropriate plasmids by Ni-NTA column chromatography to homogeneity, and in vitro-transcribed tRNAn Ala (nuclear-encoded cytosolic tRNA Ala) and tRNAm Ala (mitochondrion-encoded tRNA Ala) of V. polyspora were used as the substrates for the aminoacylation reactions. Figure 6 shows that VpAlaRS1 and VpAlaRS2 had a similar tRNA specificity. This result reinforces the hypothesis that localization of these two enzymes determines their functions in vivo.

Both ALA1 and ALA2 of Vanderwaltozyma polyspora are of mitochondrial origin

As V. polyspora is so far the only known yeast species that possesses two distinct AlaRS genes (Figure 1), possible historical relationships between VpAlaRS1, VpAlaRS2, and their yeast homologues were analyzed using the neighbor-joining method. To better illustrate this phylogeny, representative AlaRS sequences from all three of the major domains of life (Bacteria, Archaea, and Eukarya) were also subjected to the analysis. Bias in the alignments of these sequences was eliminated by reducing them to their core active sites. This portion comprises only 53% of the sequence of V. polyspora AlaRS2. As shown in Figure 7, except for AlaRS sequences from mitochondrion-deficient eukaryotes Giardia lamblia and Trichomonas vaginalis, all eukaryotic AlaRS sequences, including those from S. cerevisiae, C. albicans, Y. lipolytica, V. polyspora, A. thaliana, Drosophila melanogaster (cytoplasmic and mitochondrial isoforms) and Homo sapiens (cytoplasmic and mitochondrial isoforms), clustered within a monophyletic branch that showed a higher similarity to the bacterial branch than to the archaean branch (Figure 7). This finding suggests that all AlaRS sequences from mitochondrion-containing
eukaryotes, including those from yeasts, are of the mitochondrial type. Thus, irrespective of the cellular compartments in which their protein products are active, \( VpALA1 \) and \( VpALA2 \) actually arose from the same ancestor. Moreover, it appears that loss of the AlaRS gene of eukaryotic origin in mitochondrion-containing eukaryotes occurred long before the events of yeast speciation.

DISCUSSION

As aaRSs are likely to be one of the most ancient families of proteins, they are often targets of choice for phylogenetic analyses. Results presented herein showed that while all other yeasts contained a single AlaRS gene with dual functions, \( V.\ polyspora \) possessed two distinct AlaRS genes (\( VpALA1 \) and \( VpALA2 \)) (Figures 1 and 3). \( VpALA1 \) and \( VpALA2 \) can substitute for the cytoplasmic and mitochondrial functions of \( S.\ cerevisiae\ ALA1 \), respectively (Figures 3 and 4). Strangely enough, the protein sequence deduced from the ORF of \( VpALA2 \) (starting from ATG1) lacks a cleavable N-terminal mitochondrial targeting signal as predicted by a PSORTII analysis (29). In fact, the ATG1 codon of this gene is located at a position equivalent to that of the ATG initiator codon for the cytoplasmic form of \( S.\ cerevisiae\ AlaRS \) (Figure 1). Further investigation revealed that \( VpALA2 \) is exclusively initiated from upstream non-ATG codons (Figure 4). This unexpected finding may account for the observation that while this gene is transcribed at an efficiency equivalent to that of its counterpart (i.e. \( VpALA1 \)) (Figure 5), it has a much lower protein expression level (Figure 3). Also unexpected is the finding that one of the non-ATG initiator codons of \( VpALA2 \) contains a less favorable nucleotide, C, at relative position -3 (Figures 1 and 4). Unlike most other examples of alternative initiation of translation (30–33), the downstream ATG initiator candidate of \( VpALA2 \) appears to be silent in this case.

The specificity of an aminoacylation reaction is generally accomplished by direct recognition of tRNA by its cognate aaRS. In some instances, recognition depends mainly on the anticodon loop, while in others it relies more on sequences or structures in the acceptor stem (34). In the instance of tRNA\(^{Ala}\), the primary identity determinant is a G3:U70 base pair, which is strictly conserved in the acceptor stems of all known tRNA\(^{Ala}\) molecules from prokaryotes, archaea, eukaryote cytoplasm and chloroplasts (27,28). This strong preference for a single determinant may provide a functional basis for the high feasibility and high efficiency of cross-species and cross-compartmental complementation (Figures 3 and 4) (6). Amazingly, even a human AlaRS gene can complement an \( ALA1 \) knockout strain of \( S.\ cerevisiae \) (26). On the other hand, not all eukaryotic tRNA\(^{Ala}\)s bear a G3:U70 base pair in their acceptor stems. One such example was found in \( Drosophila\ melanogaster \), where the cytoplasmic tRNA\(^{Ala}\) has a canonical G3:U70 base pair in its acceptor stem, but the GU base pair has been translocated to position 2:71 in its mitochondrial...
isoacceptor. As a result, the mitochondrial tRNA\(^{\text{Ala}}\) (with a G2:U71 base pair) can only be recognized by its mitochondrial cognate enzyme (35). Further investigation revealed that the ability to recognize the translocated GU base pair is conferred by an insertion peptide of 27 amino acid residues embedded inside the mitochondrial enzyme (35). Despite \(V.\) polyspora also possessing two distinct AlaRS genes, no similar insertion peptide was found in their protein products, and all of their cognate tRNAs contained a G3:U70 base pair (Figure 2).

An earlier phylogeny proposed that the existing AlaRS gene in mitochondrion-containing eukaryotes was horizontally transferred to the nucleus from an ancestral mitochondrion and therefore should be regarded as being of mitochondrial origin (36,37). However, the AlaRS gene in the amitochondriate protists \(G.\) lamblia and \(T.\) vaginalis appears to have evolved from a different origin. In fact, the AlaRS sequences retrieved from these two eukaryotic organisms were clustered into a sister group that grossly deviated from the major eukaryotic branch and exhibited higher affinity to the archaeal branch instead (36) (Figure 7). Conceivably, the evolutionary history of AlaRS in eukaryotes is far more sophisticated than anticipated. In this regard, our findings reported herein are of particular interest, as they show that all yeast AlaRS genes, including those in \(V.\) polyspora, are of mitochondrial origin, regardless of the cellular compartments to which their protein products are targeted (Figure 7). As all other yeast species studied contain a single dual-functional AlaRS gene (Figures 1–3), it is likely that contemporary AlaRS genes in \(V.\) polyspora arose from duplication of such a gene. Notably, only one copy of the AlaRS gene was retained in the genomes of \(S.\) cerevisiae and \(C.\) glabrata after the sorting-out process (Figures 1 and 7). Moreover, it was shown that only a single copy of the valyl-tRNA synthetase gene was retained in yeast species descended from the aforementioned event (11). Thus, the evolutionary footprints in relation to genome duplication and reduction appear to be independent for each aminoacylation activity, even in the same organism.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

**FUNDING**

The National Science Council (Taipei, Taiwan; NSC97-2311-B-008-002-MY3, NSC97-2311-B-008-003-MY3 to C.C.W.); Armed Forces Taoyuan General Hospital (Lungtan, Taiwan; to C.Y.K., C.C.W.).
Funding for open access charge: National Science Council (Taiwan).

Conflict of interest statement. None declared.

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