#### **ORIGINAL ARTICLE**



# **Evolution of Eukaryal and Archaeal Pseudouridine Synthase Pus10**

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#### Abstract

In archaea, pseudouridine ( $\Psi$ ) synthase Pus10 modifies uridine (U) to  $\Psi$  at positions 54 and 55 of tRNA. In contrast, Pus10 is not found in bacteria, where modifications at those two positions are carried out by TrmA (U54 to m<sup>5</sup>U54) and TruB (U55 to  $\Psi$ 55). Many eukaryotes have an apparent redundancy; their genomes contain orthologs of archaeal Pus10 and bacterial TrmA and TruB. Although eukaryal Pus10 genes share a conserved catalytic domain with archaeal Pus10 genes, their biological roles are not clear for the two reasons. First, experimental evidence suggests that human Pus10 participates in apoptosis induced by the tumor necrosis factor-related apoptosis-inducing ligand. Whether the function of human Pus10 is in place or in addition to of  $\Psi$  synthesis in tRNA is unknown. Second, Pus10 is found in earlier evolutionary branches of fungi (such as chytrid *Batrachochytrium*) but is absent in all dikaryon fungi surveyed (Ascomycetes and Basidiomycetes). We did a comprehensive analysis of sequenced genomes and found that orthologs of Pus10, TrmA, and TruB were present in all the animals, plants, and protozoa surveyed. This indicates that the common eukaryotic ancestor possesses all the three genes. Next, we examined 116 archaeal and eukaryotic Pus10 protein sequences to find that Pus10 existed as a single copy gene in all the surveyed genomes despite ancestral whole genome duplications had occurred. This indicates a possible deleterious gene dosage effect. Our results suggest that functional redundancy result in gene loss or neofunctionalization in different evolutionary lineages.

Keywords Phylogeny · Protein evolution · Subfunctionalization · Pseudogene · Orthologs

# Introduction

During protein synthesis, transfer RNA (tRNA) requires modifications at specific nucleotides to ensure the correct incorporation of amino acids to a growing peptide chain. A superfamily of pseudouridine synthases families can

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isomerize uridine (U) to pseudouridine ( $\Psi$ ) at multiple positions in RNA post-transcriptionally (Kaya and Ofengand 2003). In this superfamily, members of TruA, TruB, and TruD families modify tRNA; members of the RluA and RsuA families primarily involve in uridine modification of rRNA; and members of Pus10 can modify tRNA at position U54 to  $\Psi$  (Gurha and Gupta 2008).

Both TruB (of bacterial origin) and Pus10 (of archaeal origin) can modify U to  $\Psi$  at tRNA position 55 in vitro (Blaby et al. 2011; Gurha and Gupta 2008; Roovers et al. 2006). In bacteria, TrmA methylates tRNA at position U54 to m<sup>5</sup>U54 or ribo-T54 (ribothymidine) (Ny and Björk 1980). These three enzymes—TruB, TrmA, and Pus10—all have eukaryal orthologs and thus, have potential overlapped roles in modification of uridines at position 54 (by TrmA and Pus10) and 55 (by TruB and Pus10) of tRNAs. Indeed, when crystal structures of Pus family members were compared, overlapping structural similarities of an aspartate residue as the catalytic amino acid in the active pocket were revealed and confirmed. The structure is preserved in all the eukaryal orthologs studied (Foster et al. 2000; Hoang 2004; Hoang

et al. 2006; McCleverty et al. 2007; Pan et al. 2003; Sivaraman et al. 2002). Moreover, mutational analysis of Pus family members identified additional four amino acids and structural features—forefinger loop (FFL) and thumb loop—to be important for substrate recognition and binding (Chan and Huang 2009; Conrad et al. 1999; Gurha and Gupta 2008; Hamilton et al. 2005; Hur and Stroud 2007; Joardar et al. 2013; Kamalampeta et al. 2013; Spedaliere et al. 2000).

In this paper, the molecular evolution of Pus10 and resolution of functional redundancy between orthologs of Pus10, TruB, and TrmA in different eukaryotic lineages are described. Lineage-specific alterations in key Pus10 sequence features are shown: length of the FFL and substitution of amino acids in the catalytic region and thumb loop. Different evolutionary hypotheses are proposed to explain how neofunctionalization, subfunctionalization, and gene loss have given different lineage-specific outcomes.

# **Materials and Methods**

#### **Collection of Datasets**

For the identification of Pus10-like proteins, only the species with whole-sequenced and annotated genomes were retrieved from the National Center for Biotechnology Information (NCBI), the DOE Joint Genome Initiative (JGI), the Solanaceae Genomic Network (SGN), and the Ensembl genome browser (version 62) (Bombarely et al. 2011; Hubbard et al. 2009; McGinnis and Madden 2004). Sequence alignments were performed using the basic local alignment tool Protein BLAST (BLASTP; using default settings) in NCBI. The full length Pus10 sequences from Homo sapiens (NP\_653310, PDB: 2V9K), Arabidopsis thaliana (NP\_173466), Methanocaldococcus jannaschii (NP\_247004), and Pyrococcus furiosus (NP\_578868.1) were used as a query. Similarity Sequences with BLASTP score of 40 or better and with > 25% coverage were initially retained. However, those that did not harbor the essential Asp (aspartic acid) in the conserved core catalytic sequence GREDXD (Joardar et al. 2013) or were found to be significantly better aligned to another known Pus protein (e.g., Cbf5, Pus4) were excluded from later analysis. These were likely not functional orthologs of Pus10. Accession numbers for all the proteins obtained are given in Supplementary Table 1.

#### **Protein Alignment and Phylogenetic Analysis**

Orthologous Pus10 protein sequences of 116 species were collected from Eukarya and Archaea. A weakly aligned

Pus10 homolog was found for Nanoarchaeum but was excluded from the downstream phylogenetic analysis due to extreme divergence of the sequence and absence of key Pus10 sequence features. The N-terminal region of Pus10 showed high divergence specific to each taxonomic domain and kingdom, and resulted in very low or no bootstrap supported values. Thus, protein alignment and phylogenetic analyses were conducted using only the conserved C-terminal catalytic domain (e.g., Gly286-Asp528 in human Pus10). In this study, amino acid positioning of Pus10 refers to the corresponding sequence position in relation to M. jannaschii. Isolation of the C-terminal domain was done by multiple sequence alignment using ClustalX (version 2.0.12) of whole protein sequences followed by removing the N-terminus region (e.g., residues 1-286 in human Pus10) (Larkin et al. 2007; McCleverty et al. 2007). Maximum likelihood analyses of all taxa with respect to the C-terminal of orthologous Pus10 were calculated with the Whelan and Goldman (WAG + I + G)empirical substitution model (Whelan and Goldman 2001). The web server RAxML Black box (http://phylobench. vital-it.ch/raxml-bb/) was used for maximum likelihood analyses (Stamatakis 2006). Newick files were uploaded into FigTree (version 1.3.1) (http://tree.bio.ed.ac.uk/software/figtree) and the following settings were altered. Line weight: 2x, cladogram was chosen as displayable transform version, while node labels were chosen to display bootstrap values. Further features were selected: node bars and scale bar (altered to line weight of two). Tree branches were colored to emphasize different kingdoms and the tree graphic was exported as a jpeg format.

#### **Annotation of Phylogenetic Trees**

The molecular phylogeny of the C-terminal Pus10 was annotated using the Interactive Tree Of Life (version 2.0) (iTOL; http://itol.embl.de/index.shtml) (Letunic and Bork 2011, 2016) featuring the key amino acids in the active pocket and structural features relevant for recognition and binding of tRNA substrates. Substitutions in the key Pus10 features were mapped onto a maximum likelihood tree and color coded. A separate species tree with 131 taxa (including those without detectable PUS10) was generated to illustrate the presence/absence pattern of Pus10 in relation to species ancestry and to determine potential gain/ loss events (Supplementary Table 1). A reduced version with of the species tree was generated by acquiring taxonomy numbers from NCBI uploaded onto iTOL. The resulting species tree was color coded with respect to kingdom, except for phylum amoebozoa. Furthermore, presence/ absence of Pus10 within clades was emphasized (Letunic and Bork 2011) (refer to illustration shown in Fig. 5).

# Homology Modeling and Structural Superimposition

A homology model of the protein structure of *M. jannaschii* Pus10 was calculated using 3D-JIGSAW (version 2.0; http:// bmm.cancerresearchuk.org/~3djigsaw/) (Bates et al. 2001). The crystal structure of *H. sapiens* (PDB: 2V9K, *B* factor 38.6, resolution: 2.0 Å) was used as a template (McCleverty et al. 2007). Both the crystal structures of *H. sapiens* Pus10 and the homology-modeled protein structure of *M. jannaschii* were uploaded onto the Swiss-PDB Viewer (version 4.1) and superimposed via the 'generate structural alignment tool' (Guex et al. 2009). Pus10 of *H. sapiens* and *M. jannaschii* are displayed as ribbon and each backbone of the 3D-structure was color coded. Conserved amino acids in all pseudouridine synthases were individually displayed, color coded, and labeled (refer to Fig. 3).

#### **Results**

# Characterization of Key Conserved Structural Features of Pus10

Members of the Pseudouridine Synthase superfamily have a similar 3D structure and a conserved catalytic Asp (Mueller and Ferré-D'Amaré 2009). Other accessory domains with binding capacity and features (e.g., FFL and thumb loop) are conserved within sub-families and can distinguish between different members (Fig. 1). Crystal structure analysis of RluA and TruA identified FFL and thumb loop in the catalytic site as the 'pinch mechanism' holding the substrate in place (Hoang et al. 2006; Hoang and Ferré-D'Amaré 2001). Only three families (TruA, RluA, and Pus10) have both the FFL and thumb loop, whereas other families have either loop individually (Fig. 1).

The THUMP domain is unique to Pus10 within the pseudouridine synthase superfamily (Fig. 2). It is structurally similar to the THUMP domain of bacterial ThiI and present at the N-terminus, and likely involved in substrate recognition and binding (Aravind and Koonin 2001). Modification of the N-terminal THUMP domain in Pus10 was observed



Fig. 1 Domain distribution of pseudouridine synthases. Pseudouridine synthase superfamily contains (six) families and their substrates. Domains are represented as boxes, while loops are represented as ovals. Catalytic site and thumb loop are represented by their consensus sequences. Conserved catalytic aspartate (D) across all pseudouridine synthases is marked in bold, X indicates any amino acid. Due to the focus of this paper, Pus10 and tRNA55 are in bold. TruD family=TruD, TruA family=TruA, TruB family=TruB, RsuA family=RluB and RsuA, RluA family=RluC and RluA. Members of RluA and RsuA families differ in their structural features. RluA is lacking two extensions compared to its family member RluC. RsuA is lacking the thumb loop and the C-terminal extension compared to its member RluB (Mueller and Ferré-D'Amaré 2009)

Fig. 2 THUMP domain modification of Pus10 in different lineages. Blue colored helices and arrows emphasize structural elements belonging to typical THUMP domain and are observed in archaeal Pus10. Three insertions were observed in multiple sequence alignment across 116 taxa and are indicated with arrows. + Means presence of an insertion; – means no insertion. (Color figure online)



 Table 1
 Structural alignment with THUMP domain via DaliLite

Species	Lineage	Z-score
H. sapiens	Animal	3.7
D. melanogaster	Animal	3.7
C. elegans	Animal	2.8
H. volcanii	Archaea	3.3
M. jannaschii	Archaea	2.8
P. furiosus	Archaea	2.5
R. oryzae	Fungi	4.3
N. bombycis	Fungi	2.5
A. thaliana	Plant	3.0
P. patens	Plant	2.6

throughout different eukaryotic lineages, and extensively so in plants and animals (Fig. 2). Eukaryotic Pus10 THUMP domains contain large insertions to prevent detection by routine search of hidden Markov models (HMM) for a THUMP domain (when using PFAM PF02926 as template). But, all archaeal Pus10 THUMP domains tested fit the PFAM model.

Structural pairwise alignment of bacterial ThiI (PDB: 2C5S) was made to the homology-modeled THUMP domain across eukaryal and archaeal lineages. The structural comparison analysis by the DaliLite (version 3.3) (Holm et al. 2008; Holm and Rosenstrom 2010) showed an overall high similarity Z-score (Table 1). The Z-scores of PUS10 from *P. furiosus* and *H. sapiens* are very high, 2.5 and 3.7, respectively. These scores indicate that although observed deletions in *P. furiosus* and insertions in *H. sapiens* could theoretically have affected the folding of the protein, this region still remains structurally similar to ThiI THUMP.

In Pus10 of *H. sapiens*, the zinc-binding site consists of four cysteine (Cys) residues at positions 21, 24, 109, and 112, and is thought to be involved in the maintenance of the native N-terminal structure (McCleverty et al. 2007). Multiple sequence alignment revealed the presence of

the four Cys residues in nearly all Pus10 orthologs of Archaea and Eukarya. The exceptions were found in the fungi lineage. Here, all four Cys were present in early fungi Mucoromycotina and in several Chytridiomycota, but entirely absent in Microsporidia. Furthermore, the first Cys pair (Cys21 and 24) was absent in one chytrid fungus (*Allomyces macrogynus*). Interestingly, the latter was encountered in the following invertebrates: *Branchiostoma floridae*, *Daphnia pulex*, and *Caenorhabditis elegans*. Ala (alanine) substitutions of the second pair, C106/109A in *M. jannaschii* (equivalent to C109/C112 of human Pus10) showed a reduced  $\Psi$ 54 modification, but  $\Psi$ 55 modification was unaffected (Joardar et al. 2013). This suggests Pus10 proteins may retain partial ( $\Psi$ 55) function despite the absence of the second Cys pair.

In the catalytic region, five amino acids (Asp275, Tyr339, Ile412, Lys413, Leu440 in M. jannaschii) were determined to be conserved throughout all pseudouridine synthase families (Hamma and Ferré-D'Amaré 2006; McCleverty et al. 2007). Superimposition of a homology-modeled structure of M. jannaschii Pus10 onto that of H. sapiens Pus 10 revealed the same orientation and 3D positioning of the conserved amino acids present in the active pocket. This was observed throughout all 116 organisms surveyed (Fig. 3) with the exceptions of B. floridae (Cephalochordata), D. pulex (Brachiopoda), Manihot esculenta, and Solanum lycopersicum. A conserved consensus sequence of the catalytic site in the Pus10 orthologs centered on the catalytic Asp275 and five surrounding residues 'GREDVD' (Fig. 1). Substitutions within this catalytic site have been observed throughout Archaea and Eukaryota (Fig. 4). In addition to the active pocket, two loops were experimentally confirmed to be of value for the pseudouridylation of tRNAs at positions U55 and U54. The FFL proved to contribute to modify U55 over U54. Therefore, the length of the loop was crucial, not the identities of the amino acids within the loop. The opposite was observed for the thumb loop, where the amino acid Fig. 3 Substrate-specific amino acids conserved throughout all pseudouridine synthases. a superimposition of human Pus10 (gray) and of M. jannaschii (blue) generated via Swiss-PDB viewer version 4.0.1 (Guex and Peitsch 1997). FFL and thumb loop (indicated with black arrows) are located in the C-terminal part of the protein. **b** Back view of the C-terminus with close up focusing on the catalytic core, which shows the catalytic Asp275 (white), Leu440 (orange), Tyr339 (blue), Ile412 (green), and Lys413 (pink). (Color figure online)



identity (His and Arg) is of importance for pseudouridylation (Joardar et al. 2013).

# Presence, Absence, and Copy Number of Pus10 Orthologs in Whole Genomes

Organisms with fully sequenced genomes were surveyed by BLASTP to identify Pus10 orthologs. In the identification of Pus10 orthologs, several eukaryotic and a few archaeal species were found to lack Pus10 and were placed on a species tree (Fig. 5). Four nodes of Pus10 loss were identified (Fig. 5). In fungi, all dikaryotes (Ascomycota and Basidiomycota) lacked Pus10 but had TruB orthologs. The most parsimonious answer would be a single evolutionary event in which Pus10 was lost in the common ancestor of all Dikaryon fungi. Other fungal taxa, including Chytrids and Microsporidia, contain Pus10 orthologs (Fig. 5). Most of Archaea had Pus10 but only two genera lacked functional Pus10 genes (Fig. 5). In Sulfolobus tokodaii, S. acidocaldarius, S. solfataricus, their most similar sequences (to Pus10) all lack the catalytic aspartate residue. The entire conserved 'GREDVD' sequence surrounding the catalytic aspartate residue is significantly altered (i.e., "PYSEPSDVR" in S. acidocaldarius). Although most of the residues upstream of this site are somewhat conserved. There is less conservation of the protein downstream of the catalytic site. Many of the other structural features thought to be needed to perform pseudouridylation are also missing or significantly divergent from those conserved between Archaea and Eukaryotes. The Pus10 sequence of Nanoarchaeum showed high overall divergence, with numerous substitutions throughout the sequence and resulted in low bootstrap values upon phylogenetic analysis. For these reasons, these two genera of Archaea were excluded from phylogenetic analysis. In four Archaeal orders and a few members within Desulfurococcales and Halobacteriales, shorter FFLs were observed. However, the majority of Archaea species in this study had Pus10 orthologs with completely intact Pus10 structural features (Fig. 4).

A surprising finding in all the genomes surveyed was the presence of only a single Pus10 gene per genome, despite the numerous genome duplications that have occurred in the evolution of animals, fungi, and especially in plants (where



**Fig. 4** Summary of the Pus10 characteristics of the C-terminal across Eukarya and Archaea. Here we want to emphasize on the substitutions between clades and chose taxa (97 out of 111) that differ in their Pus10 characteristic. The maximum likelihood tree was calculated via RAxML black box, model WAG+G+I. The annotation was generated via iTOL. Forefinger-loop features are represented in a single-value bar chart to emphasize on the length differences: Forefinger loop=15=insertion (longer), 10=normal length based on *H. sapiens*, 5=partial deletion (shorter). Presence/absence/substitution of

genes frequently duplicate and subfunctionalize) (Koonin and Wolf 2010; Proost et al. 2011). This finding might point to a potential dosage effect or other harmful consequence for higher copy numbers of Pus10.

amino acid is represented with a color gradient: presence of amino acid=50, substitution=25, absence=0. The catalytic region: Y339 (yellow), I412 (orange), K413 (purple), and L440 (light blue). Catalytic site including the catalytic Asp (D) (green). Thumb loop (focus on His and Arg presence; blue). All substitutions are displayed. Pus10 key features are displayed as seen in multiple sequence alignment. Species labels were color coded based on their belonging to archaea (light green) and eukaryote (light blue). (Color figure online)

# **Pus10 in Archaea**

A maximum likelihood tree was calculated for the C-terminal domain of the Pus10 in 44 archaeal species (Supplementary Fig. 1). The tree comprises 4 clades where the outer most clade is represented by members of the



Thermococcales: Cenarchaeum symbiosum and Nitrosopumilus maritimus (Thaumarchaeota). These are incorporated into the Methanococcales clade and show long branches, indicating high divergence and possible false placement due to long branch attraction or parallelism (Anderson and Swofford 2004). This sequence convergence indicates some possible alteration of the Pus10 catalytic function. When comparing key features of Pus10 protein sequences—FFL, catalytic 'GRED[V/X]D,' and thumb loop with His and Arg-of Thaumarchaeota to other Archaea and Eukaryota, Pus10 of both C. symbiosum and N. maritimus show the same FFL size as that of M. jannaschii. This could explain the incorporation of C. symbiosum and N. maritimus into the clade of Methanococcales. However, the sequence of their catalytic site differs considerably. In addition, the thumb loop lacks a conserved histidine residue. Multiple sequence alignment of Pyrobaculum arsenaticum, P. islandicum, and Thermoproteus neutrophilus also show modification in all structural features for Pus10.

The most important differences in Pus10 features were observed in Methanococcales and Thermococcales. Methanobacteriales, Halobacteriales, and Methanococcales have insertions in the FFL, creating a longer loop than the other clades in Archaea (Fig. 4). Moreover, the Thermococcales clade, within this insert-containing group, had further changes in the length of the FFL and substitutions (His376Asn and Arg377Ser) in the thumb loop.

#### Pus10 in Fungi

Of the 50 fungi with fully sequenced genomes, a total of nine species were found with orthologous Pus10 sequences (Supplementary Fig. 2). All nine were found in the earliest diverging families of fungi, the Unikaryonidae and Nosematidae (Microsporidia), Mucoraceae (Zygomycota), and Batrachochytrium (Chytridiomycota). The related lineages Ascomycota (as yeasts and sac fungi) and Basidiomycota (as mushroom fungi) did not have Pus10 orthologs in any of the 41 sequenced genomes surveyed, although other pseudouridine synthase genes (e.g., Pus4, Cbf5) were found (Fig. 5). This suggested that Pus10 was secondarily lost in the common ancestor of these fungal species.

Common sequence features among the Pus10 sequences of fungi included the presence of ten amino acid deletions in the 11th  $\alpha$ -helix and 13th  $\alpha$ -helix C-terminal to the catalytic site (Supplementary Fig. 2) (McCleverty et al. 2007). Microsporidia are also missing the first three secondary structures ( $\alpha 4$ ,  $\alpha 6$ ,  $\beta 1$ ) which are critical to the THUMP domain (Fig. 2). All early fungi species contain the same sequence length of the FFL, and three conserved amino acids were present in these Pus10 FFLs ('PxxxxGxxxxxS').

The catalytic site of Pus10 in *R. oryzae* and *M. circinelloides* showed conversion of the key second Asp 277 to Asn. Other minor substitutions in the catalytic site have also been observed (Fig. 4). The thumb loop in all nine fungi species is highly conserved in its sequence, except for in *R. oryzae* and *M. circinelloides* (Gln, instead of His). This changes the composition of the thumb loop from a basic amino acid to an acidic amino acid and could affect the binding of tRNA (Fig. 4). A TrmA ortholog was present in *B. dendrobatidis* but not in any other species of Microsporidian or Zygomycota (Table 2). Interestingly, TruB (Pus4) orthologs are not present in the earlier fungi lineages (Microsporidia, Chytridiomycota and Zygomycota) but are present in Ascomycota (*S. cerevisiae*).

#### **Pus10 in Plants**

*Arabidopsis thaliana* has a total of 24 pseudouridine synthase-like proteins. Most are paralogous copies of TruA, indicating expansion of this family in plants. Only one copy of Pus10 and TruD was detected in *A. thaliana*, and in 18 other plant genomes.

Multiple sequence alignment showed the incorporation of Ile (isoleucine), instead of Val (valine), in the catalytic site of 'GREDID' in plants. Substitutions were observed in the following species: 'GREDLD' in *Cucumis sativus*; 'GREDMD' in the moss Physcomitrella patens and green algae Ostreococcus tauri; 'GREDAD' in Chlamydomonas reinhardtii (Fig. 4). A highly modified THUMP domain was present in most plants. Nicotiana tabacum and Solanum lycopersicum (Solanaceae), Carica papaya, and Oryza sativa all lack  $\alpha 4$  and  $\beta 1$  of the THUMP domain. In addition, N. tabacum and S. lycopersicum are also missing  $\alpha 6$  and β2. Sorghum bicolor Pus10 uncharacteristically lacks the catalytic Asp and several amino acids of the catalytic site (Fig. 4), indicating it is a non-functioning protein, however other grasses all seem to have a functional Pus10 catalytic site.

The molecular phylogenetic analysis of Pus10 has the same topology as the phylogenetic tree of species, with the exception of *Ectocarpus siliculosus* (Heterokontophyta; brown algae) (Fig. 4). This resulted in a split of the two green algae representatives—*C. reinhardtii* and *O. tauri*, Chlorophyta—indicating possible genome fusion in the brown algae (Fig. 4). All plant genomes contain TruB and TrmA orthologs (N-terminus of TruB in *C. reinhardtii*).

Most species within the plant kingdom have functional Pus10; but *C. reinhardtii* and *C. papaya* have modifications in the catalytic site that may have an effect on its function.

# Pus10 in Animals and the Importance of THUMP Domain

THUMP is a RNA-binding domain present in 17 protein architectures and found in over 3587 proteins according to the PFAM database. Many of these proteins have different biological roles. For example, S-adenosyl methionine-dependent methyltransferase, rhodanese-like proteins, FtsJ-like methyltransferase, SpoU rRNA methylase family, and cytidine zinc-binding region. In this study, three major insertions which potentially disrupt or significantly alter the functional THUMP domain in Pus10 were identified (Fig. 2). All eukaryotic Pus10 THUMP domains were larger than archaeal ones and included numerous lineage-specific insertions between conserved portions of the THUMP-motif (Fig. 2); they make identification of the THUMP domain by sequence alignment difficult. The first insertion was observed on the N-terminal side of the THUMP domain and is found in all taxa except fungi and Archaea. A second insertion within the THUMP domain was observed to be present in animals and in some plants. The third insertion is found shortly before the FFL; it is conserved in higher eukaryotes and in several Protista (Paramecium tetraurelia and Trichoplax adhaerens). However, the insertions were absent in Archaea (Fig. 2). In this study, 42 animal species representing 32 different orders, a few animal-specific modifications in the THUMP domain and key Pus10 structure features (conserved residues in the catalytic site, FFL, and thumb loop) were observed (Fig. 5). The C-terminal catalytic site is highly conserved in all animals with some minor differences in earlier lineages. A maximum likelihood tree showed that the Protista, Tricoplacia, and Dictyostelia, followed by representatives of Nematoda, generally show the most diverse sequences alterations of key Pus10 features (Supplementary Fig. 3). Nematodes, C. elegans and Pristionchus pacificus, have a shorter FFL and the substitution of His to Lys in the thumb loop; same observations were made for the Diptera clade. However, Tribolium castaneum

Table 2Presence/absence ofTruB, TrmA, and Pus10

Species	TruB	Pus10	TrmA
Homo sapiens	NP_631908.1	NP_653310.2	NP_892029.2
Saccharomyces cerevisiae	EGA73247	ND	EDN59959
Zygomycota	ND	Yes	ND
Batrachochytrium dendrobatidis	ND	EGF78467	EGF83624.1
Microsporidians	ND	Yes	ND
Escherichia coli	TruB	ND	TrmA

ND not detected by BLAST in the genome, Yes present in all taxa sampled

had 'GREDFD' in the catalytic motif/core instead of I412 with respect to *M. jannaschii* (refer to Fig. 5). Although there is no experimental evidence, these modifications to the THUMP domain may potentially alter or weaken non-specific Pus10 RNA binding and potentially alter the range of substrate RNA.

In H. sapiens, Pus10 is also known as Downstream Of Bid (DOBI). It has been shown to be a downstream interactor of the Bid (BH3 interacting-domain death agonist) protein signal involved in TRAIL-induced apoptosis and release of cytochrome C from mitochondria (Aza-Blanc et al. 2003; Jana et al. 2017). TRAIL is tumor necrosis factor-related apoptosis-inducing ligand and TNF-related apoptosis-inducing ligand. Mammalian Pus10 is recognized and cleaved by caspase-3 or caspase-8, which may create the functional unit for TRAIL signaling (Park et al. 2009). To analyze the ancestry of Pus10 involvement in TRAIL-induced apoptosis and tRNA pseudouridylation, 17 animal species were tested for the presence of (a) intact orthologs of redundant functioning pathways (TruB, TrmA) and (b) essential interacting components of TRAIL-induced apoptosis, caspase-3, and caspase-8 (Table 3). TrmA as well as caspase-3 and caspase-8 genome were not detectable in D. pulex, suggesting TRAIL-induced apoptosis is absent and Pus10 is likely to perform pseudouridylation of U54 (Table 3). Caspase-3 and caspase-8 orthologs were found in *T. adhaerens*, a multicellular organism belonging to the phylum Placozoa which lacks the presence of organs (Table 3). This, together with the absence of caspase-3 and caspase-8 orthologs in all fungi and plants, indicates that TRAIL-induced apoptosis could have begun as early as the last common ancestor of animals with *T. adhaerens*, at the base of the metazoan clade, approximately 635My ago.

# Discussion

#### Evolutionary Resolution of Functional Redundancy in Pus10 and TruB

Pus10 is the most recently identified member of the pseudouridine synthase family and it has been experimentally shown to be crucial for tRNA pseudouridylation in Archaea (Gurha and Gupta 2008; Mueller and Ferré-D'Amaré 2009; Roovers et al. 2006). While there is significant similarity between eukaryal (*H. sapiens*) and archaeal (*M. jannaschii*) Pus10 orthologs, no remotely similar protein sequences were found in Bacteria. On the other hand, TrmA and TruB are absent in Archaea, but are present in Bacteria and Eukarya. This suggests that Pus10 must have originated in Archaea

Species	TruB	TrmA (3BT7)	Caspase-3	Caspase-8
Homo sapiens	NP_631908.1	NP_892029.2	NP_116786.1	NP_001219.2
Gallus gallus	XP_421776.1	XP_415080.2	NP_990056.1	NP_989923.1
Taeniopygia guttata	XP_002186667.1	XP_002187418.1	XP_002191294.1	XP_002190143.1
Anolis carolinensis	XP_003223551.1	XP_003226338.1	ND	ND
Danio rerio	NP_001122159.1	NP_956223.1	NP_571952.1	NP_571585.2
Daphnia pulex	346,482	ND	ND	ND
Branchiostoma floridae	XP_002607561.1	XP_002611903.1	XP_002610410.1	XP_002605135.1
Ciona intestinalis	XP_002130935.1	XP_002130635.1	XP_002131300.2	XP_002122848.1
Lottia gigantea	139,266	121,926	ND	ND
Paramecium tetraurelia	XP_001430104.1	XP_001442539.1	ND	ND
Trichoplax adhaerens	XP_002108027.1	XP_002115265.1	XP_002114146.1	XP_002114150.1
Dictyostelium discoideum	XP_640508.1	XP_629224.1	ND	ND
Drosophila melanogaster	NP_525120.1	NP_649083.2	NP_524551.2	NP_476974.1
Apis mellifera	XP_397244.4	XP_396538.2	XP_395697.2	XP_395697.2
Tribolium castaneum	XP_967291.1	XP_973242.1	XP_967501.2	XP_967501.2
Caenorhabditis elegans	NP_499370.1	NP_503353.2	ND	NP_001022453.1
Mucor circinelloides	113,436	111,833	ND	ND
Ostreococcus tauri	XP_003080670.1	XP_003078170.1	ND	ND
Physcomitrella patens	XP_001773683.1	XP_001766898.1	ND	ND
Oryza sativa	NP_001054428.2	EEC70100.1	ND	ND
Arabidopsis thaliana	NP_196950.2	NP_188767.2	ND	ND
Ricinus communis	XP_002521488.1	XP_002530289.1	ND	ND
Populus trichocarpa	EEF11588.1	XP_002314271.1	ND	ND

Number accession number in GenBank, ND not detected by BLAST of genome

**Table 3**Presence/absence ofTruB, TrmA, caspase-3, andcaspase-8

and is required to modify tRNA at both positions U55 and U54. The coexistence of Pus10, with parallel tRNA modifying genes TrmA and TruB, impacts the function of Pus10 in Eukarya. This situation may have also occurred in the observed species of Sulfolobus which have genes similar to bacterial TruB and TrmA, possibly acquired through horizontal gene transfer (HGT). In this case, the presence of both of these genes might have lead to the loss of key amino acids involved in the pseudouridine synthase function, but not the complete loss of the Pus10 protein. In another Archaea, P. furiosus, in which HGT has introduced bacterial RumA to modify tRNA position U54 (but not U55), may have resulted in a more specific loss of Pus10 function for position U54 only. In general, after a duplication event, an organism is less likely to maintain more than one copy of a gene involved in tRNA processing and this coincides with our study of Pus10 across all species examined. Redundancy of Pus10 with nonhomologous proteins performing similar functions may have led to its functional diversification.

The most important differences in terms of potential for revealing the mechanism of function in Pus10 were observed in Methanococcales and Thermococcales (refer to Fig. 4). Structural comparison of P. furiosus to M. jannaschii showed these modifications are very likely relevant for hook formation and substrate binding, which may be related to the loss of in vivo modification of U54 without affecting U55 modification (Joardar et al. 2013). Several in vitrogenerated mutants of M. jannaschii Pus10 at these residues lost Ψ54 activity, but not activity of Ψ55. P. furiosus Pus10 does have a weak in vitro ability to pseudouridylate at tRNA position 54, whereas M. jannaschii has a robust one (Gurha and Gupta 2008). An in silico mutagenesis of His376 to Asn and Arg377 to Ser in M. jannaschii shows that this change in the thumb loop should increase the gap width between both loops (Joardar et al. 2013). A wider gap generated by changes to the thumb loop, and/or shortening of the FFL, could contribute to a reduction in the tight pinching of the tRNA substrate that might be required for modification of U54.

In the case of *P. furiosus*, a bacterial rRNA methyltransferase (in the subfamily RumA of Pus superfamily) that is known to be acquired through HGT, is responsible for U54 modification in tRNA (Urbonavičius et al. 2008). Thus, in *P. furiosus*, the presence of RumA likely resulted in the partial relaxation of selection pressure and subsequent loss of Pus10  $\Psi$ 54 synthase ability, whereas its  $\Psi$ 55 activity is retained. These differences in sequence across Archaea may indicate the extent of possible subfunctionalization and uncoupling of  $\Psi$ 55 and  $\Psi$ 54 activity of Pus10 for the order Thermococcales.

Based on our findings, it is likely that only TruB and TrmA orthologs remain functionally unchanged. Pus4 (TruB ortholog) converts U55 to  $\Psi$ 55, and Trm2 (TrmA ortholog) modifies U54 to ribo-T54 in tRNAs of eukarvotes (Becker et al. 1997; Nordlund et al. 2000; Nurse et al. 1995). A different pseudouridine synthase, Cbf5, performs pseudouridylation of rRNA in a guide-RNAdependent manner in both eukaryotes and Archaea. However, in eukaryotes, Cbf5 has also neofunctionalized to play a role during mitosis by binding to microtubules and centromeres (Blaby et al. 2011; Jiang et al. 1993; Rashid et al. 2006). Thus, like Cbf5, Pus10 may have neofunctionalized to signal TRAIL-induced apoptosis in humans, if not all animals either in place of or in addition to the pseudouridylation function. Pus10 protein is nuclear localized, and translocates to the mitochondria upon cleavage by caspase-3, which in turn amplifies caspase-3 activity, creating a positive feedback loop that has a central role in apoptosis (Jana et al. 2017). However, some mammalian tRNAs have been shown to have  $\Psi$  at position 54 instead of ribo-T. This indicates the existence of a protein to provide  $\Psi$  synthase activity for U54 (Roe and Tsen 1977). The obvious candidate is Pus10 (Gurha and Gupta 2008). Thus, we hypothesize that in some eukaryotic lineages Pus10 and TrmA orthologs may both have subfunctionalized, partitioning U54 modification activity, each with its own specific subset of tRNA substrates. Furthermore, we observed in the multiple sequence alignment of all 131 taxa, three insertions in the THUMP domain, which could alter RNA recognition. Interestingly, a few human mitochondrial tRNA<sup>LEU</sup> do contain  $\Psi$  at position 55 (Helm 2006). However, the presence of TruB in H. sapiens does not leave out the possibility that TruB performs the aforementioned pseudouridylation rather than Pus10 (Table 2). While it is clear that mammalian Pus10 is part of TRAIL-apoptosis it remains unclear if it continues tRNA pseudouridylation at position 54 in the Mammalian clade, though it is the most obvious candidate. If so, it may still be capable to perform tRNA pseudouridylation at position 55 as it may be impossible to uncouple this reaction from position 54 pseudouridylation.

A RsuA-like  $\Psi$  synthase protein is known to be a suppressor of *var2* mutant phenotype and plays a role in variegation, an apoptosis of the chloroplast, in *A. thaliana* (Yu et al. 2008). This suggested a potential TRAIL-like apoptosis pathway might exist in plants. However, no sequences like caspase-3 were found. Plants contain 3 metacaspases, which are involved in the so-called 'deathosome,' to regulate programmed cell death (Coll et al. 2010; Vercammen 2004; Vercammen et al. 2007). Metacaspases are also found in the kingdoms of Protozoa, fungi, and plants. Plants contain type I and II metacaspases, but not animal-like caspase-4 and metacaspase-9 (AtMC4, AtMC9) recognize positively charged amino acids such as Arg or Lys in the motifs of substrate peptides with the amino acids FR, GRR, GKR, and VRPR and cleave after Arg or Lys (Vercammen

2004). Future experimental work will be needed to determine whether Pus10 plays a part of either a mitochondrial or plastid apoptotic-like machinery in plants.

#### **Molecular Evolution of Pus10 in Eukaryota**

In Eukarya, the common ancestor likely had three proteins from Bacteria and Archaea through genome fusion (Koonin 2010) that modified residues U54 and U55 of tRNA, which created a functional redundancy. Over time the modification and loss of Archaeal-origin Pus10 interacts with that of bacterial-origin TruB and TrmA orthologs. Surprisingly, unlike most redundancy created through gene duplication, this redundancy was not resolved through gene loss in most lineages (Sémon and Wolfe 2007). Individual eukaryotic lineages were examined and revealed lineage-specific differences in the fate of these three proteins in the fungi, plants, and animals.

A surprising finding in all the sequenced genomes of this study was the presence of only a single Pus10 gene per genome, despite the numerous genome duplications that have occurred in the evolution of animals, fungi, and especially in plants (where genes frequently duplicate and subfunctionalize) (Koonin and Wolf 2010; Proost et al. 2011). This finding might point to a potential dosage effect or other harmful consequence for higher copy numbers of Pus10.

In Fungi, TruB orthologs are not present in the earlier fungi lineages (Microsporidia, Chytridiomycota, and Zygomycota) but are present in Ascomycota. The most parsimonious solution is that the three early-branching fungal lineages have lost both TruB and TrmA, but have retained Pus10 (Table 2), while in Dikaryon fungi (Ascomycota, Basidiomycota), the opposite order of gene loss is observed, TruB and TrmA are retained and Pus10 is lost. Thus, it seems that these two groups of fungi have undergone the opposite selection of the enzymes Pus10, TruB, and TrmA that can modify uridine residues at positions 54 and 55 of tRNA. The pattern of gain and loss here clearly points to the random elimination of redundant enzymes for tRNA U54 and U55 modification that could be due to the loss of selection pressure. Given the phylogenetic analysis, it is unlikely that this pattern was the result of HGTs (refer to Fig. 5). This simple explanation for resolving functional redundancy through random gene loss is however not the case in plants and animals which retained seemingly functional orthologs of all three genes.

A recent duplication event in angiosperms should have resulted in at least two copies of Pus10 and in the 11 eudicot genomes surveyed (Proost et al. 2011). However, it has been shown that the extra paralogous genes coding for signaling and metabolism (e.g., protein kinases) are often maintained in plants while paralogous genes involved in DNA repair and modification are typically thinned to a limited number through rapid gene loss (Thomas et al. 2006). Gene ontology analysis of a tetraploid genome which revealed the preference of keeping or losing duplicated genes is frequently based on their biological role. Indeed, duplicated genes which are responsible for tRNA processing were less likely to be maintained in the genome.

Ala (alanine) substitutions in *M. jannaschii* differentially reduced  $\Psi$ 54 activity while retaining most of wild-type  $\Psi$ 55 activity (Joardar et al. 2013). However, a triple mutant in *M. jannaschii* (IIe412A/Lys413A/Leu440A) showed no pseudouridine synthase activity at either site. These two species, along with *Sorghum bicolor* that is lacking a catalytic Asp, indicate that Pus10 activity is lost in some plant species. However, the overall presence of conserved, seemingly functional Pus10 in all other plant species in this study would indicate a positive selection pressure for some function of this enzyme.

A RsuA-like pseudouridine synthase protein is known to be a suppressor of *var2* mutant phenotype and plays a role in variegation, an apoptosis of the chloroplast, in A. thaliana (Yu et al. 2008). This suggested a potential TRAILlike apoptosis pathway might exist in plants. However, no sequences like caspase-3 and caspase-8 were found. Plants contain 3 metacaspases, which are involved in the so-called 'deathosome,' to regulate programmed cell death (Coll et al. 2010; Vercammen 2004; Vercammen et al. 2007). Metacaspases are also found in the kingdoms of Protozoa, fungi, and plants. Plants contain type I and II metacaspases, but not animal-like caspases. Metacaspase-4 and metacaspase-9 (AtMC4, AtMC9) recognize positively charged amino acid such as Arg or Lys in the motifs of substrate peptides with the amino acids FR, GRR, GKR, and VRPR and cleave after Arg or Lys (Vercammen 2004). Future experimental work will be needed to determine whether Pus10 plays a part of either a mitochondrial or plastid apoptotic-like machinery in plants.

# Conclusion

This study provides evidence of Pus10 subfunctionalization in Thermococcales (Archaea) and possibly in the Chytridiomycota. In both cases, TrmA/Trm2-like enzymes are present in the genome, indicating an alternate route for modification of U54 that co-occurs with mutations that reduce or eliminate Pus10 U54 modification. Apparently, methylation of U54 of tRNA to ribo-T54 by proteins acquired through either HGT or descent can lead to functional diversity in  $\Psi$ 54 synthase activity of Pus10. A gap in the catalytic site of Pus10 in *S. bicolor* (plant) was noticed which could indicate that Pus10 is on the verge to become a pseudogene in some plant lineages. Acknowledgements This work was supported by NIH Grant GM55045 to R.G.

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