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Pyrrolysine and Selenocysteine Use Dissimilar Decoding Strategies

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Abstract: Selenocysteine (Sec) and pyrrolysine (Pyl) are known as the 21st and 22nd amino acids in protein. Both are encoded by codons that normally function as stop signals. Sec specification by UGA codons requires the presence of a *cis*-acting selenocysteine insertion sequence (SECIS) element. Similarly, it is thought that Pyl is inserted by UAG codons with the help of a putative pyrrolysine insertion sequence (PYLIS) element. Herein, we analyzed the occurrence of Pyl-utilizing organisms, Pyl-associated genes, and Pyl-containing proteins. The Pyl trait is restricted to several microbes, and only one organism has both Pyl and Sec. We found that methanogenic archaea that utilize Pyl have few genes that contain in-frame UAG codons, and many of these are followed with nearby UAA or UGA codons. In addition, unambiguous UAG stop signals could not be identified. This bias was not observed in Sec-utilizing organisms and non-Pyl-utilizing archaea, as well as with other stop codons. These observations as well as analyses of the coding potential of UAG codons, overlapping genes, and release factor sequences suggest that UAG is not a typical stop signal in Pyl-utilizing archaea. On the other hand, searches for conserved Pyl-containing proteins revealed only four protein families, including methylamine methyltransferases and transposases. Only methylamine methyltransferases matched the Pyl trait and had conserved Pyl, suggesting that this amino acid is used primarily by these enzymes. These findings are best explained by a model wherein UAG codons may have ambiguous meaning and Pyl insertion can effectively compete with translation termination for UAG codons obviating the need for a specific PYLIS structure. Thus, Sec and Pyl follow dissimilar decoding and evolutionary strategies.

Abbreviations: Pyl, pyrrolysine; Sec, selenocysteine; MtmB, monomethylamine methyltransferase; MtbB, dimethylamine methyltransferase; MttB, trimethylamine methyltransferase; *pylT*, tRNA^{Pyl} gene; PylS, pyrrolysyl-tRNA synthetase; SECIS, selenocysteine insertion sequence; PYLIS, pyrrolysine insertion sequence; ORF, open reading frame; UTR, untranslated region; RF1, class I release factor; RF2, release factor 2; *selA*, Sec synthase gene; SelB, Sec-specific elongation factor; EFSec, eukaryotic Sec-specific elongation factor; *selC*, tRNA^{Sec} gene; SelD, selenophosphate synthetase; SBP2, SECIS-binding protein 2; nt, nucleotide(s).

INTRODUCTION

Pyrrolysine (Pyl) has recently been identified in the active site of monomethylamine methyltransferase (MtmB) from *Methanosarcina barkeri*, and sequences encoding Pyl-containing homologs of this protein were found in several other methanogenic archaea, including *Methanosarcina acetivorans*, *Methanosarcina mazei*, and *Methanosarcina thermophila* (1–3). Methylamine methyltransferase genes from these organisms contain in-frame UAG codons, which do not

halt translation, but encode Pyl. Following this discovery, additional Pyl-containing methyltransferases have been identified in *Methanosarcina*, and to date three classes of Pyl-containing methylamine methyltransferase genes are known: *mtmB*, dimethylamine methyltransferase (*mtbB*), and trimethylamine methyltransferase (*mttB*) (1, 2). Some *Methanosarcina* contain several paralogs of each methyltransferase family. Using this information, various genome sequences were scanned for genes encoding homologous Pyl-containing proteins. This search identified an *mttB* homolog in a Gram-positive bacterium *Desulfotobacterium hafniense* (2). More recently, an Antarctic archaeon, *Methanococcoides burtoni*, has also been reported to utilize Pyl (4). In contrast, no Pyl-containing methyltransferases have been reported in eukaryotes. It is also not known whether the utilization of Pyl is restricted to methyltransferases or other Pyl-containing proteins exist.

Although the mechanism of Pyl biosynthesis and incorporation into protein is not fully understood, the presence of a *Methanosarcina* tRNA^{Pyl} gene (*pylT*) with the CUA anticodon and of class II aminoacyl-tRNA synthetase gene (*pylS*) argued for cotranslational incorporation of Pyl (2). A recent study suggested that *pylT* and *pylS* are the only foreign genes necessary for translating UAG as Pyl in *Escherichia coli*, when the cells are supplemented with exogenous Pyl (5). In addition, it was reported that PylS could activate and ligate Pyl directly onto tRNA^{Pyl} (5, 6) and that tRNA^{Pyl} is directly recognized by the standard elongation factor EF-Tu (7). Analysis of the genomic context of *pylT* and *pylS* identified *pylB*, *pylC*, and *pylD*, which were suggested to participate in Pyl biosynthesis or insertion into protein (2). *pylT*, *-S*, *-B*, *-C*, and *-D* genes constitute a Pyl gene cluster (or Pyl operon), and *pylT* and *pylS* genes are considered as the Pyl utilization signature.

Because Pyl is inserted in response to a codon that in most organisms functions as a terminator, there are three distinct possibilities for how Pyl insertion can be achieved: (i) redefinition of a subset of UAG stop codons by a *cis*-acting mRNA signal to encode Pyl; (ii) reassignment of all UAG codons to encode Pyl; and (iii) ambiguous meaning of UAG codons, e.g. a competition between read-through and termination such that a fraction of ribosomes translating the UAG codon incorporate Pyl, whereas the rest support termination (8). However, the attention of researchers has previously focused on the first possibility, because of the analogy between Pyl and sele-

nocysteine (Sec) (9). Both Pyl and Sec are encoded by “termination” codons and are the only known additions to the pool of 20 universal, directly encoded, amino acids. Therefore, Sec and Pyl are known as the 21st and 22nd amino acids.

The mechanism of Sec insertion is known in much detail (10–13). Incorporation of Sec requires the presence of selenocysteine insertion sequence (SECIS) element, a hairpin structure residing in 3'-untranslated regions (3'-UTRs) of selenoprotein mRNAs in eukaryota and archaea, or immediately downstream of Sec UGA codons in eubacteria (13–15). SECIS is essential for Sec insertion, whereas in its absence UGA serves as terminator (16). Several attempts have been made to search for analogous stem-loop structures in mRNAs encoding Pyl-containing proteins. A putative secondary structure was predicted 5–6 nucleotides downstream of the Pyl-encoding UAG codon in *mtmB* mRNAs and designated as pyrrolysine insertion sequence (PYLIS) element (9, 17). This predicted structure has not been tested experimentally for functional relevance.

Identification of genes encoding Sec- and Pyl-containing proteins in genomic sequences is challenging, because standard annotation tools interpret UGA and UAG as stop signals. For example, most methylamine methyltransferases in *Methanosarcina* are incorrectly annotated. At present, no tools are available for prediction of Pyl-containing proteins, and previous *in silico* approaches were limited to manual analyses and BLAST searches (2). In the case of Sec, tools have been developed and successfully used to identify selenoprotein genes by searching for SECIS elements (18–20) and Sec/Cys pairs in homologous sequences (21, 22).

In this study, we used bioinformatics approaches to analyze Pyl-utilizing organisms and Pyl-containing proteins, and to examine possible mechanisms of Pyl insertion. Our data suggest that indiscriminate Pyl insertion at UAG may be tolerated in Pyl-utilizing archaea and that Pyl decoding processes are different from those of Sec.

EXPERIMENTAL PROCEDURES

Sequence Databases and Resources—260 completely sequenced prokaryotic genomes were downloaded from the NCBI ftp server (<http://ftp.ncbi.nih.gov/genomes/Bacteria>). To analyze incompletely sequenced genomes, we used partial genomic sequences (contigs) from the NCBI data base of microbial genomes as well as a non-redundant nucleotide data base. Both web-based and local Blast programs (23) were used for sequence analysis (available at <http://ftp.ncbi.nih.gov/blast> and <http://www.ncbi.nlm.nih.gov/BLAST>).

Identification of Pyl Gene Cluster Homologs and Known Pyl-containing Proteins—*pylT* and *pylS* sequences from *M. barkeri* (accession number AY064401 [GenBank]) were used as queries to search genomic databases for possible homologs with an *e* value below 0.01. Candidate tRNA^{Pyl} sequences were further analyzed to identify structural features associated with known tRNA^{Pyl}, such as a 6-bp acceptor stem and a base between the D and acceptor stems (2). Other genes in the Pyl gene cluster (*pylB*, *-C*, and *-D*) were similarly analyzed by comparative sequence analyses. We further examined whether these genes were organized in clusters.

A tblastn program with default parameters was used to search for Pyl-containing methylamine methyltransferases in different organisms. Open reading frames (ORFs) and conservation of UAG-flanking regions were then examined manually. Multiple alignments and phylogenetic trees were generated with ClustalW (24).

Analysis of Candidate PYLIS Elements in Methylamine Methyltransferase Genes—Sequences either downstream of in-frame UAG codons or in the putative 3'-UTR of methylamine methyltransferase gene mRNAs were analyzed manually to search for possible conserved structures and sequence features within these structures. RNA secondary structures were predicted with RNAfold 1.4, which is a part of the Vienna RNA package (available at [http://www.tbi.univie.ac.at/~ivo/RNA/\(25\)](http://www.tbi.univie.ac.at/~ivo/RNA/(25))).

Analyses of UAG Codon Function—To characterize functions of UAG codons, a homology-based approach was developed and used to analyze UAG-flanking regions in four Pyl-utilizing organisms, *M. acetivorans*, *M. maei*,

M. burtonii, and *D. hafniense*. This procedure was implemented using simple Perl scripts (available upon request). First, genes terminating with UAG were extracted from the original annotation files and extended until the next non-UAG stop signal (UAA/UGA). ORFs translated from the elongated genes were analyzed by tblastn against non-redundant and microbial genome databases. We also screened for conservation of UAG codons in nucleotide sequences and of UAG-flanking regions in protein sequences. This procedure assigned each UAG codon to one of three categories as follows: (i) A UAG was interpreted as a terminator if an elongated sequence was sufficiently long (>30 nucleotides), and all of its homologs had a true stop signal (either a non-UAG terminator in Pyl-utilizing organisms or any termination signal in other organisms) that corresponded to the UAG codon. (ii) A UAG was interpreted as a candidate Pyl codon if an elongation was >30 nt, and >50% homologs extended beyond the UAG and terminated near the termination site of the elongated sequence. All identified sequences were then analyzed for conservation of UAG in Pyl-utilizing organisms with blastn and blastp. (iii) A UAG was not assigned a function if the two situations discussed above could not be satisfied (for example, if we observed short elongations beyond UAG codons, lack of sequence similarity between homologs in regions flanking UAG, or a small number of homologs extending beyond the UAG).

A non-Pyl-utilizing archaeon, *Methanococcus jannaschii*, was also analyzed using the same approach. It served as the control in searches involving Pyl-utilizing archaea.

Analysis of Overlaps between Elongated UAG-containing Genes and Downstream Genes—Overlaps between genes are common in prokaryotic genomes (26). To examine how extensions of UAG-containing genes relate to the extent of the overlap, we analyzed overlapping genes before and after sequence elongation downstream of predicted stop codons in *M. acetivorans* and *M. maei*. A simple Perl script was developed for this analysis (available upon request). We first identified overlapping genes in the original genome annotations, determined the number of overlaps in each genome, and measured overlap lengths. The longest overlap in a genome was defined as an overlap threshold. We then extended genes terminated at UAG until the next non-UAG stop signal using the approach described above and repeated the overlap analysis procedure. We reasoned that if no significant increase in the number of genes whose overlap was longer than the threshold would be observed, the situation would be consistent with the use of UAG as either a terminator or a Pyl codon. However, if the sequence extension procedure generated many genes with large overlaps with the downstream genes, the situation would be consistent with the use of UAG codon as terminator. In addition, the gene overlaps involving genes terminated at UAA and UGA codons were analyzed using the same strategy (e.g. before and after extension). These served as controls.

Identification of Genes Associated with Pyl Utilization—All predicted ORFs in *M. acetivorans* and *M. maei* genomes were searched for exclusive occurrence in genomes that utilize Pyl. The tblastn program was used to search these sequences against 260 completely sequenced prokaryotic genomes, non-redundant nucleotide data base and unfinished microbial contigs with an *e* value below 0.05. A simple script was developed to parse the tblastn output and examine presence/absence of homologs in analyzed genomes. A pairwise alignment tool, b12seq, was then used with an *e* value cutoff set to 0.001 to cluster protein sequences into different families. The occurrence of these proteins in *D. hafniense* was then analyzed.

RESULTS AND DISCUSSION

Distribution of Pyl-utilizing Organisms and Pyl-containing Proteins

Available completely and incompletely sequenced prokaryotic genomes were screened for tRNA^{Pyl} (*pylT*) and pyrrolysyl-tRNA synthetase (*pylS*) sequences, and their patterns of occurrence were compared with those of other Pyl genes (*pylB*, *-C*, and *-D*). We found that the products of *pylB* (biotin synthase homolog) and *pylC* (carbamoyl-phosphate synthetase homolog) have close homologs in a wide variety of organisms. In contrast, *pylS*, *pylT*, and *pylD* (nucleoside-diphosphate sugar epimerase homolog) are specific for methanogenic archaea and *D. hafniense* (Figure 1). In *D. hafniense*, *pylSn* and *pylSc* encode the N- and C-terminal parts of PylS (2), and small overlaps occur between

pylT and *pylSc* (4 nt) and between the *pylB* and *pylC* genes (52 nt). *pylS*, *pylT*, and *pylD* always cluster with *pylB* and *pylC*, and the overall Pyl gene cluster has identical mutual organization of these sequences, except that *D. hafniense* *pylSn* is located at the end of the cluster (Figure 1). Thus, the five Pyl genes define the Pyl gene cluster, but only *pylT* and *pylS* (and perhaps *pylD*) sequences can be used as the signature for the Pyl trait.

A search of completely and incompletely sequenced prokaryotic genomes for Pyl genes revealed only six organisms that could utilize Pyl, including four members of *Methanosarcina* genera, *M. burtonii*, and *D. hafniense*. *Methanosarcina* species and *M. burtonii* belong to *Methanosarcinales*, suggesting that Pyl is encoded by a UAG codon in a restricted group of phylogenetically related organisms that occupy a specific environmental niche. High conservation of the Pyl gene cluster and the small number of organisms that utilize Pyl suggest its relatively recent origin.

In the 6 Pyl-utilizing organisms, a total of 29 Pyl-containing methylamine methyltransferase genes was identified (Table I). They are distributed in three enzyme families that do not share significant sequence similarity. Figure 2 shows the occurrence of these genes in genomes and contigs. Only *mtmB* genes cluster with the Pyl operon genes (in three *Methanosarcina* organisms). In *M. mazei*, two distant duplicate *mtmB* genes are present. In *M. barkeri*, two duplicate *mtmB1* genes cluster together and are on the opposite strands with the Pyl cluster.

Further analyses of the three methylamine methyltransferase protein families revealed conservation of Pyl in MtmB and MtbB (*i.e.* no MtmB and MtbB homologs were detected, in which Pyl is replaced with other residues or in which the Pyl-encoding UAG codon is replaced with a non-UAG stop signal). On the other hand, multiple MttB homologs were detected, in which Pyl is not conserved and replaced with various amino acids (Figure 3). This situation is in contrast to Sec, which is highly conserved. In addition, most selenoproteins have homologs, in which Sec is replaced with cysteine (Cys). In fact, the Sec/Cys pair in homologous sequences is a feature that is used for identification of selenoproteins in genomic databases (21, 22).

Phylogenetic analyses of the three methylamine methyltransferase families as well as of the *pylT* and *pylS* genes typically placed *D. hafniense* genes as outliers (Figure 4). In Pyl-utilizing archaea, all *mtmB*, *mtbB*, and *mttB* genes encode Pyl-containing proteins. Conversely, *D. hafniense* possessed *mttB* genes encoding proteins with and without Pyl. MttB homologs that did not have Pyl were broadly distributed in other bacteria.

Although the use of Pyl appears to be prevalent in methanogenic archaea, from our data it could not be established with certainty whether the Pyl trait evolved in these organisms or in bacteria. Both Pyl operon sequences and *mttB* have optimal codon usage in organisms in which they are present (data not shown), arguing against a recent (traceable) lateral transfer of the Pyl trait between methanogenic archaea and *D. hafniense*.

Analyses of Candidate PYLIS Elements

By analogy to SECIS, a stem-loop structure in selenoprotein mRNAs that reprograms a small fraction of UGA codons in a genome to serve in Sec insertion (11, 13), the occurrence of putative PYLIS structures was proposed (9, 17). However, we found that the RNA structures downstream of UAG codons in several *mtbB* and *mttB* mRNAs that were recently suggested as putative PYLIS elements (17) are dissimilar and do not occur in *mtmB*. Manual analyses of sequences downstream of UAG codons, as well as sequences in UTRs revealed no obvious common structure shared by members of all three methylamine methyltransferase families. A possibility remains that structures downstream of UAG codons inhibit termination.

Functions of UAG Codons in Pyl-utilizing Organisms

Because UGA has a dual function in Sec-utilizing organisms (Sec insertion and translation termination), there is a possibility that UAG, in a similar fashion, also serves two functions (Pyl insertion and translation termination) in Pyl-utilizing organisms. If PYLIS is absent, what might be a mechanism for discriminating between the two functions of UAG codons? An alternative possibility would be a reassignment of all UAG codons from stop to Pyl. To address these possibilities, we analyzed the distribution of the three stop codons in Pyl-utilizing organisms (Table II). In *D. hafniense*, the percentage of the genes predicted to terminate at UAG is 22.5%, which is similar to the proportion of genes predicted to terminate at UGA (28.4%). The extensive utilization of UAG in this bacterium suggests that this codon likely has a dual role in protein synthesis.

In contrast, the proportion of genes that are predicted to terminate at UAG in Pyl-containing archaea is <5.0%. For example, only 126 (including all incorrectly annotated methylamine methyltransferases) of 3,371 genes are predicted to use UAG terminator in *M. mazei*. This value is much lower than the proportion of UAA or UGA terminators, suggesting that UAG might be functionally distinct from UAA and UGA in these archaea.

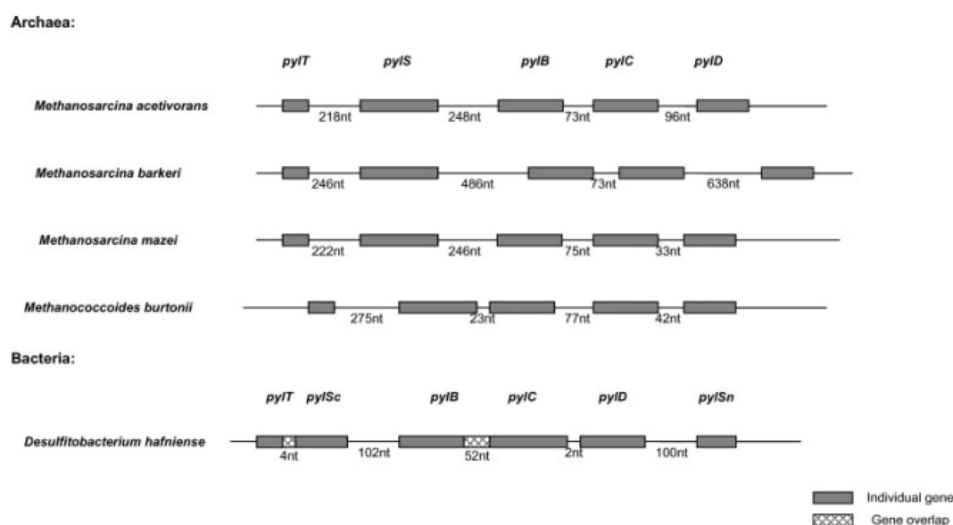


FIGURE 1. **Pyl gene cluster (operon).** The Pyl gene cluster includes five neighboring genes: *pylT*, *pylS*, *pylB*, *pylC*, and *pylD*. In archaeal genomes, these genes show identical organization. In *D. hafniense*, two genes (*pylSn* and *pylSc*) code for the N- and C-terminal parts of PylS, and there are small overlaps between *pylT* and *pylSc* and between the *pylB* and *pylC* genes as shown.

TABLE I. Distribution of Pyl-containing methylamine methyltransferase genes in Pyl-utilizing organisms

Domain of life	Organism	MMA methyltransferase (<i>mtmB</i>)	DMA methyltransferase (<i>mtbB</i>)	TMA methyltransferase (<i>mttB</i>)
Archaea	<i>M. acetivorans</i>	<i>mtmB1</i> , <i>mtmB2</i>	<i>mtbB1</i> , <i>mtbB2</i> , <i>mtbB3</i>	<i>mttB1</i> , <i>mttB2</i>
	<i>M. barkeri</i>	<i>mtmB1</i> (2), <i>mtmB2</i>	<i>mtbB1</i> , <i>mtbB2</i> , <i>mtbB3</i>	<i>mttB</i>
	<i>M. mazei</i>	<i>mtmB</i> (2)	<i>mtbB1</i> , <i>mtbB2</i> , <i>mtbB3</i>	<i>mttB1</i> , <i>mttB2</i>
	<i>M. thermophila</i>			<i>mttB</i>
	<i>M. burtonii</i>	<i>mtmB1</i> , <i>mtmB2</i>	<i>mtbB1</i> , <i>mtbB2</i>	<i>mttB1</i> , <i>mttB2</i>
Bacteria				
	<i>D. hafniense</i>			<i>mttB</i>

We further used a homology-based search strategy to examine the coding potential of UAG codons as an additional test to characterize the function of these codons in archaea. Our approach was to extend the reading frames of all genes predicted to contain UAG codons until the next non-UAG stop signal. For each elongated UAG-containing gene, the tblastn program was used to identify candidate homologs in other organisms and to examine the conservation of UAG-flanking regions within translated sequences. We reasoned that if a sequence is sufficiently extended (>30 nt) beyond the UAG, and all of its homologs in other organisms have true stop signals that corresponded to the UAG codon (that is, only the sequence upstream of UAG is conserved, whereas sequence similarity is absent downstream of UAG), the UAG should be a terminator (Figure 5A). Using this strategy, we could reliably identify UGA stop signals in Sec-utilizing organisms and distinguish them from UGA codons for Sec (data not shown).

Conversely, if the extension is long (>30 nt), and most candidate homologs extend beyond the UAG to end near (or after) the site corresponding to the non-UAG stop codon in the elongated sequence, the UAG codon in the sequence of interest is considered a Pyl codon candidate. In addition to testing UAG function, this strategy could also be

used for identification, in Pyl-encoding organisms, of candidate Pyl-containing proteins. To avoid the possibility of dealing with a sequencing error or a pseudogene, we required the presence of sequences encoding a candidate Pyl-containing protein in two or more genomes of Pyl-utilizing organisms (Figure 5B). In this case, methylamine methyltransferases served as true positives, because they can be extended beyond their Pyl UAG codons, share homology with other proteins in sequences downstream of their UAG codons, and occur in at least 4 Pyl-utilizing organisms as Pyl-containing forms (Figure 3).

In other situations (see Figure 5C for specific examples), we could not distinguish between Pyl-encoding functions and stop signals. However, if the UAG was followed with a nearby stop codon, either Pyl insertion or translation termination could presumably be tolerated.

Surprisingly, among all the genes with predicted in-frame UAG codons in the Pyl-utilizing archaea, we could not detect a single unambiguous candidate containing UAG as its terminator (Table III). Instead, we found that in most genes either UAG codons are followed with additional nearby non-UAG stop signals (40.3% of UAGs), or UAG-containing genes can be extended to generate conserved sequence alignments downstream of UAGs (24.3%). The former situations cannot

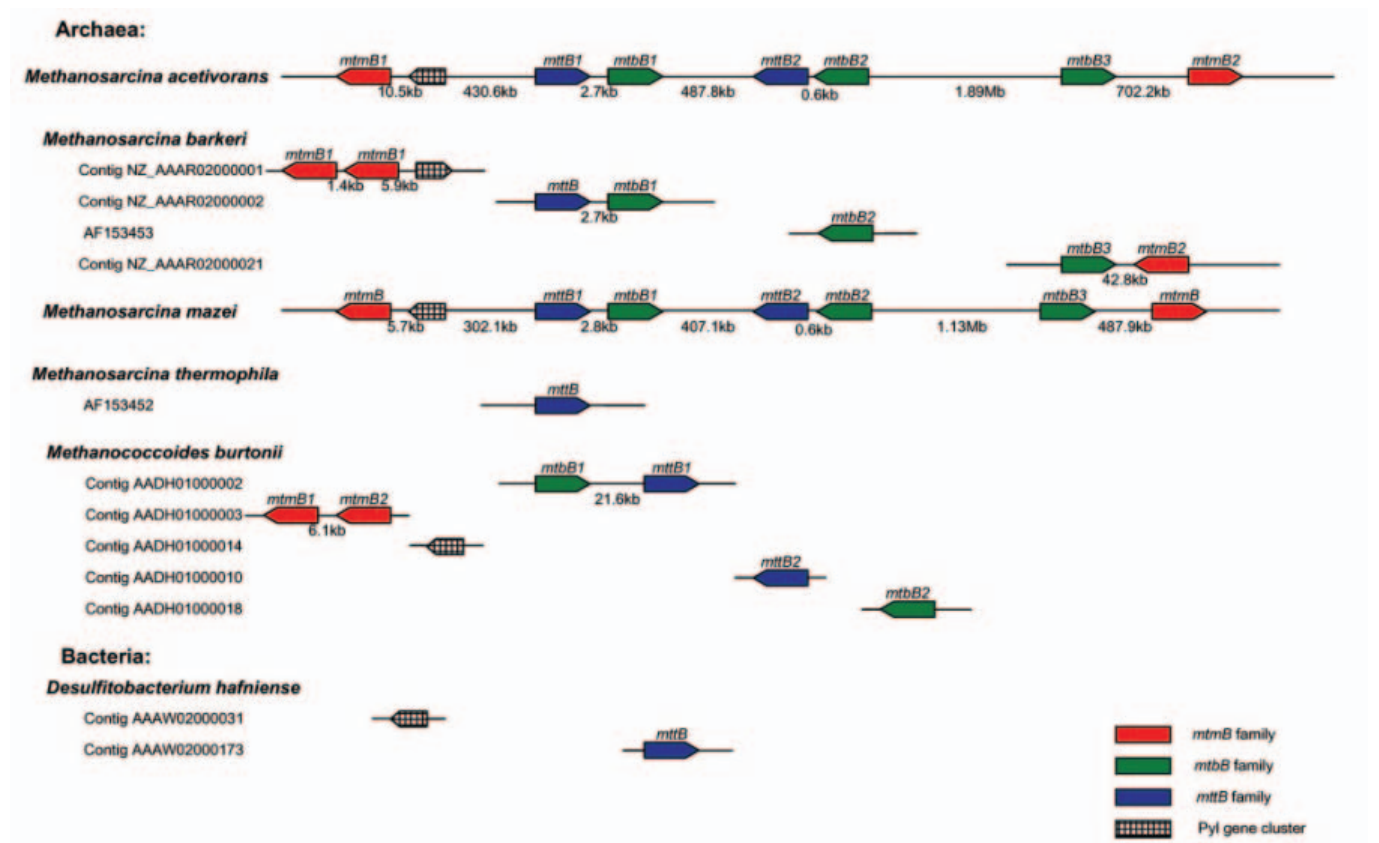


FIGURE 2. Occurrence of methylamine methyltransferase genes in Pyl-utilizing organisms. The *mtmB*, *mtbB*, and *mttB* genes and the Pyl gene clusters are shown by the indicated color scheme in the completed genomes of *M. acetivorans* and *M. mazei* and in contigs containing these genes in other Pyl-utilizing organisms. Coding direction and distances between the genes are indicated.

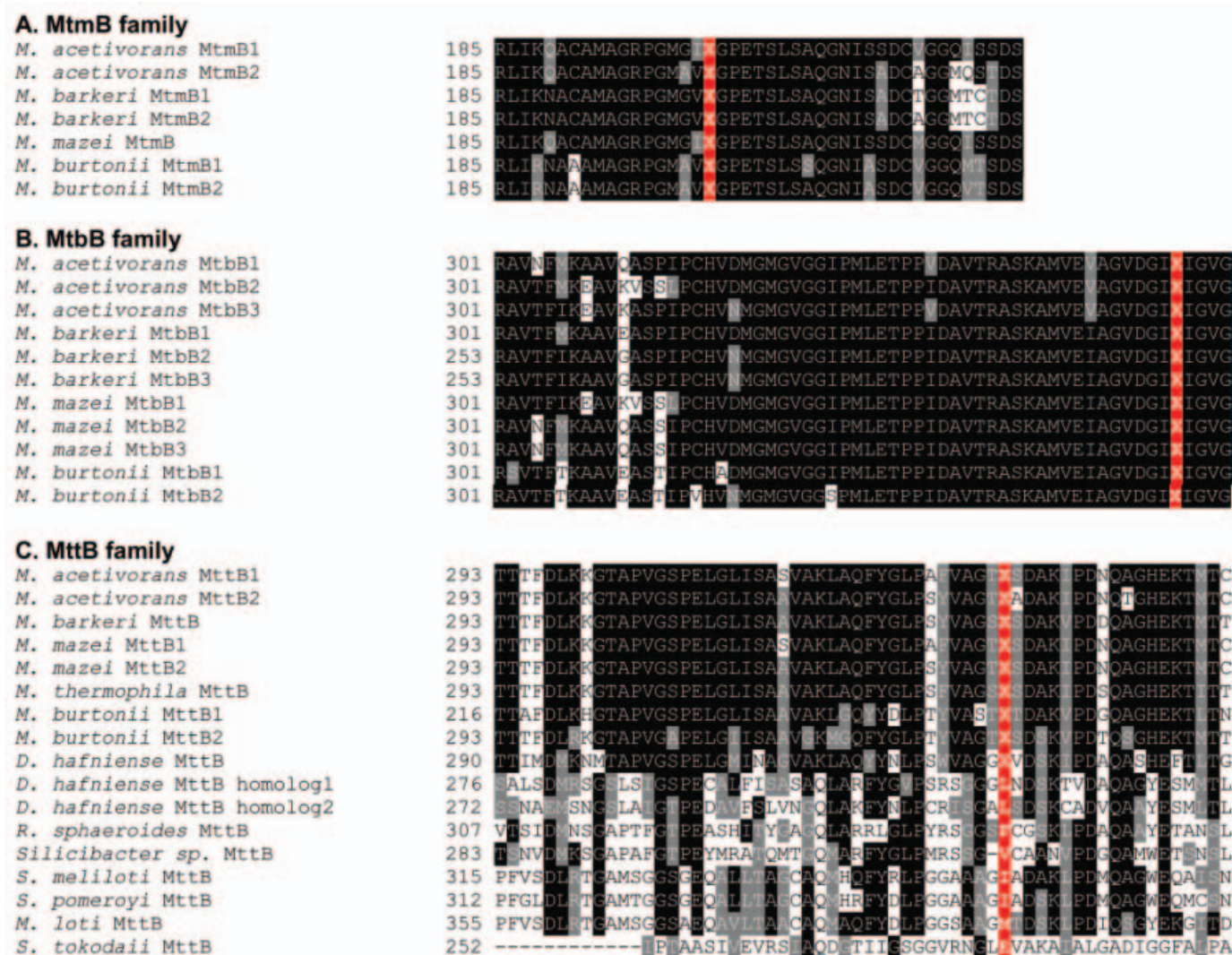


FIGURE 3. Multiple alignments of Pyl-flanking regions in methylamine methyltransferase families (MtmB, MtbB, and MttB). Pyl is shown by X, and its location in the alignment is highlighted in red. MttB homologs that lack Pyl are also shown.

distinguish between termination and Pyl insertion, whereas the latter correspond to candidate Pyl-containing proteins.

Interestingly, these searches revealed only four protein families, including three known methylamine methyltransferase families and a family of transposases, as candidate Pyl-containing proteins encoded in at least two genomes. UAG-containing forms of transposases were identified in *M. acetivorans* (4 sequences) and *M. mazei* (16 sequences), and all had a single UAG codon at the same position. These data suggest that transposases are a family of novel Pyl-containing proteins (Figure 5B).

However, only methylamine methyltransferases were found in three or more genomes, and these enzymes provided the best match to the Pyl trait. This observation suggests that Pyl utilization was conserved during evolution for its use by methylamine methyltransferases. At least in MtmB, Pyl is located at the enzyme active site, where it was suggested to be directly involved in catalysis serving as a strong electrophile (3). It cannot be excluded that additional, species-specific proteins that use catalytic Pyl residue occur in organisms capable of Pyl insertion. However, our searches suggest that the Pyl trait is associated exclusively with methylamine methyltransferases and not with any other family of Pyl-containing proteins.

The bias against unambiguous assignment of a fraction of the occurrences of a codeword as terminator that we observed in Pyl-utilizing archaea, was not seen in either Sec-utilizing organisms (data not shown) or the Pyl-decoding bacterium *D. hafniense*. In *D. hafniense*, we detected a number of "true" UAG stop signals (221 hits, 19.6% of all UAG codons). In addition, 33 *D. hafniense* sequences, including one incorrectly annotated *mttB* gene, have a predicted in-frame UAG codon. However, except for *mttB*, these are present in single copies and so are likely false positives. Overall, these findings are consistent with the hypothesis that UAG has a dual function in *D. hafniense*, but no evidence for such a dual function was observed in archaea.

Using the same approach, we analyzed a non-Pyl-utilizing archaeon, *Methanococcus jannaschii*, in which UAG is known to have unambiguous stop codon meaning. Like *Methanosarcina*, *M. jannaschii* has a small number of UAG codons (UAG: UGA:UAA = 164:222:1343). However, we detected 52 genes (31.7% of all UAGs), in which these UAGs could be classified as true stop signals (Figure 5A). Only 5 genes showed sequence homology downstream of their UAG, but these were represented by single UAG-containing sequences. Thus, there is a clear difference in the use of UAG codons between Pyl-utilizing and non-Pyl-utilizing archaea (Table III).

TABLE II. Distribution of annotated UAA, UAG, and UGA codons in the genomes of Pyl-utilizing organisms

Organism	Genome size (nt)	Annotated genes	Stop codon		
			UAG	UAA	UGA
Archaea					
<i>M. acetivorans</i>	5,751,492	4,540	224 (4.9%)	2,217 (48.8%)	2,099 (46.3%)
<i>M. mazei</i>	4,096,345	3,371	126 (3.7%)	1,800 (53.4%)	1,445 (42.9%)
<i>M. burtonii</i>	~2.6 M (unfinished)	2,782	131 (4.7%)	1,264 (45.4%)	1,383 (49.9%)
Bacterium					
<i>D. hafniense</i>	~6.1 M (unfinished)	4,999	1127 (22.5%)	2,427 (48.5%)	1,422 (28.4%)

Partially overlapping genes are common in prokaryotic genomes (26, 27), however, most overlaps are short. We reasoned that if UAG codons function as terminators, extensions of these sequences downstream of UAGs would result in some overlaps that are abnormally long. In contrast, if UAG codons function in Pyl insertion, the extended sequences would not result in overlaps or would produce mostly short overlaps with the downstream genes.

To examine these possibilities, we analyzed overlaps involving previously annotated genes and genes whose ORFs were extended beyond the annotated stop signals for all three stop codons in *M. acetivorans* and *M. mazei* (Table IV). The annotated genes showed a maximum of a 98 nucleotide overlap in *M. acetivorans* and a 241 nucleotide overlap in *M. mazei*, and these values were set as thresholds to evaluate the overlaps obtained after the genes were extended to the next stop signal.

We found that, among all UAG-extended genes, 27 *M. acetivorans* and 43 *M. mazei* genes overlap with downstream genes that are in different reading frames (genes that overlapped in the same reading frame were excluded as these produced gene fusions rather than overlaps and could correspond to Pyl-containing proteins). However, in *M. mazei*, the lengths of all overlaps are below the threshold (241 nt). In *M. acetivorans*, only 3 genes have overlaps longer than the threshold (98 nucleotides), but these (including the longest overlap of 172 nucleotides) are still shorter than the *M. mazei* threshold. Thus, extension of UAG-containing genes until the next stop signal does not modify the overall composition of the overlapping genes.

In contrast, extension of UAA- and UGA-containing genes generated many genes that overlap above the thresholds (e.g. 117 genes for UAA and 195 genes for UGA in *M. acetivorans*), with some overlaps being above 1000 nt (the longest overlap is 1934 nt for UGA in *M. acetivorans*, Table IV). Thus, both the number of abnormal gene extensions and the longest observed overlaps involving UGA- and UAA-containing sequences are much higher than those of the UAG-containing sequences, suggesting that the usage of UAA and UGA is functionally distinct from that of UAG in Pyl-utilizing archaea and that read-through of UAG codons should be tolerated better than read-through of UAA or UGA stop codons.

Analysis of Class I Release Factors in Archaea

Most eukaryotes and archaea possess a single class I release factor (RF1) responsible for the recognition of all stop codons in mRNA (28, 29). Structural studies of human RF1 suggest that its N-terminal domain interacts with mRNA during stop codon recognition (30). Numerous studies demonstrated that the sequences near the highly conserved NIKS motif in this domain are critical for stop codon recognition (31–33). If UAG codons were reassigned to code for Pyl in Pyl-containing archaea, their release factors would have specific changes responsible for the alteration of their specificity. We have searched for release factors in all sequenced *Euryarchaeota*. Unexpectedly, we found that two Pyl-utilizing archaea, *M. acetivorans* and *M. barkeri*, encode two non-identical RF1s. Figure 6 shows a phylogenetic tree of RF1s from *Euryarchaeota*. All RF1s from Pyl-containing archaea have specific amino acid changes in the area surrounding the NIKS motif (Figure 7). Interestingly, in organisms with two release factors, additional

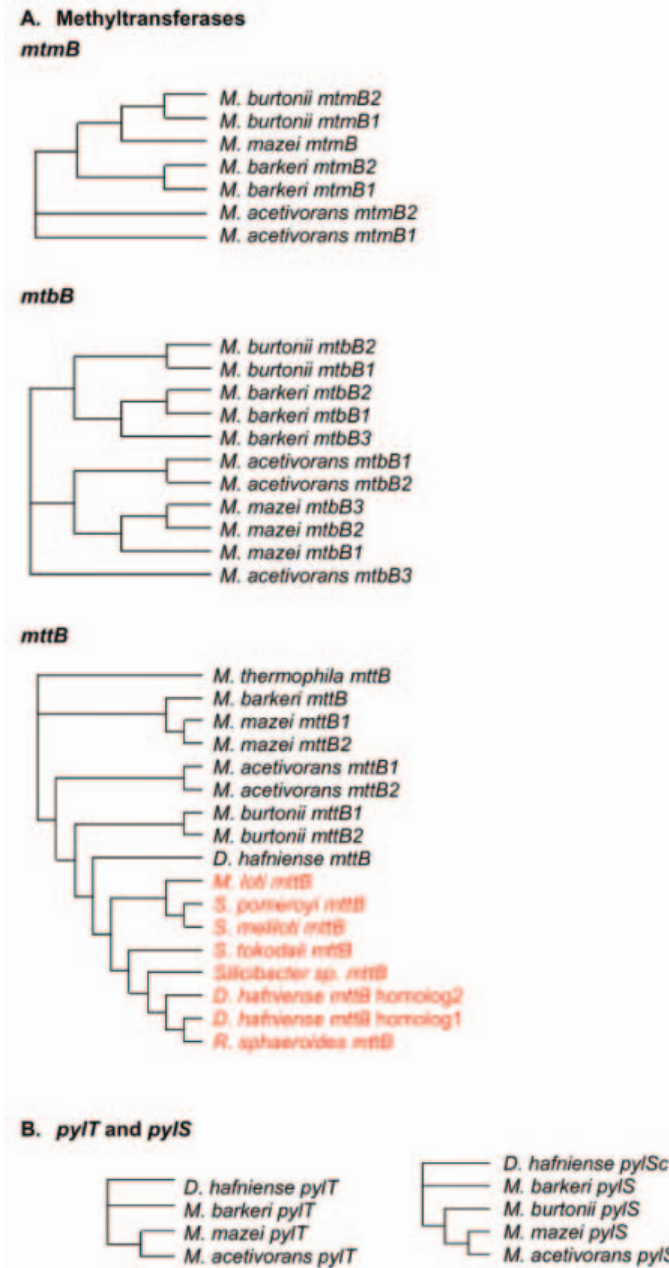


FIGURE 4. Phylogenetic analysis of methyltransferases, pylT and pylS. A, phylogenetic analysis of methyltransferase genes. mttB homologs that do not encode Pyl are shown in red. B, phylogenetic analysis of pylT and pylS genes.

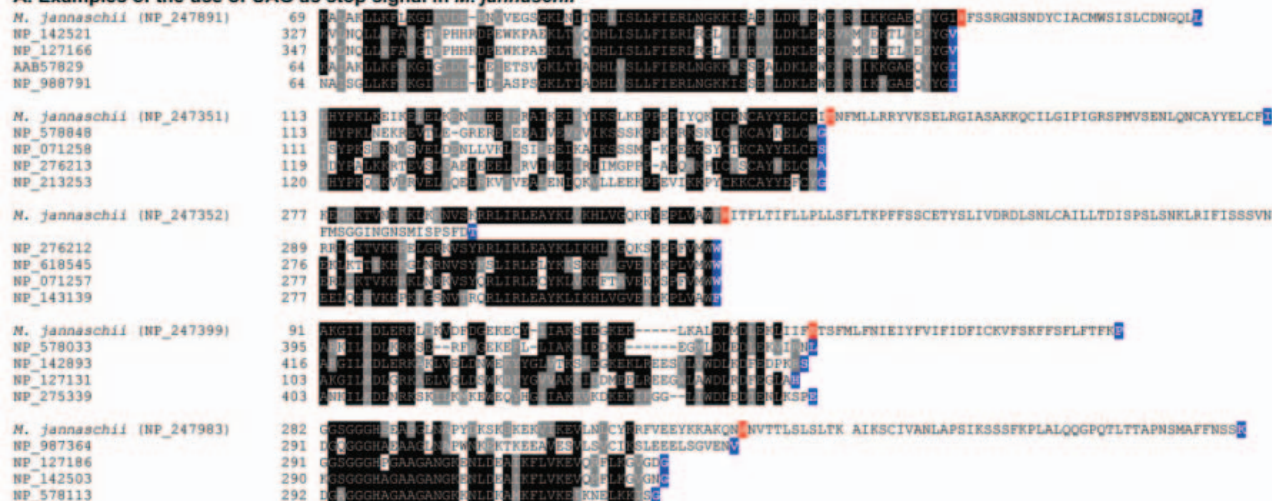
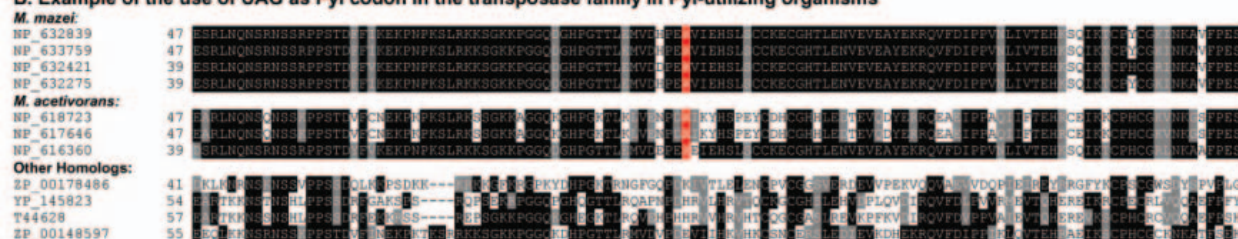
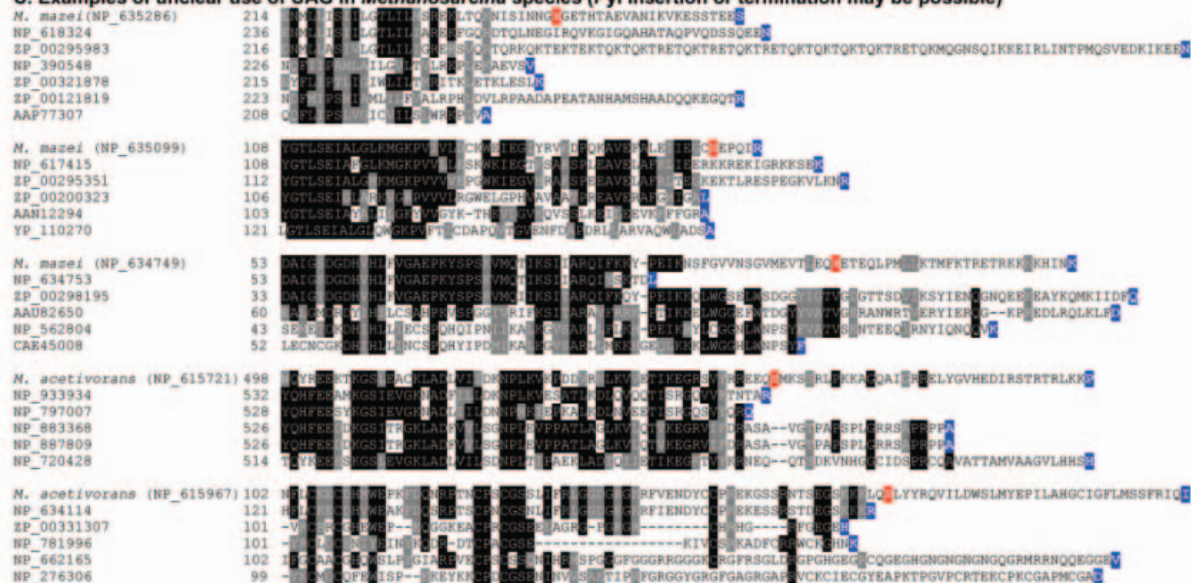
A. Examples of the use of UAG as stop signal in *M. jannaschii***B. Example of the use of UAG as Pyl codon in the transposase family in Pyl-utilizing organisms****C. Examples of unclear use of UAG in *Methanosarcina* species (Pyl insertion or termination may be possible)**

FIGURE 5. Multiple sequence alignments of *M. jannaschii* and *Methanosarcina* proteins containing in-frame UAG codons. X indicates hypothetical translation of UAG codons, and its location in the alignment is highlighted in red. C-terminal amino acid residues in proteins and hypothetical extensions with UAG read-through are highlighted in blue. Accession numbers for proteins are also shown. A, alignments involving *M. jannaschii* proteins in which UAG serves as terminator. Conserved regions were found only upstream of UAG codons. In addition, all homologs had a true stop signal (either a non-UAG terminator in Pyl-utilizing organisms or any of the three terminators in other organisms) that corresponded to UAG codons in query sequences. B, alignment of transposases, a newly identified family of Pyl-containing proteins. C, alignments of several proteins from *Methanosarcina*, in which the function of UAG codons is unclear.

RF1s contain amino acid substitutions in the NIKS motif itself. These data suggest that RF1s from Pyl-containing archaea may have indeed evolved to change their stop codon recognition specificity.

Several organisms are known, in which stop codons are reassigned to sense codons (34, 35). Although in bacteria (in which there are two

semi-specific class I release factors) stop codon reassignment involves loss of one release factor, in eukaryota stop codon reassignment involves changes in the specificity of release factors. For example, in the ciliate *Euplotes*, UGA was reassigned to encode Cys (36), and it has been shown that its RF1 does not recognize UGA stop codons (37).

TABLE III. Analysis of UAG-flanking regions in archaeal genomes

Organism	Number of UAG-utilizing genes	Distribution of genes containing in-frame UAG codons					
		Distance between UAG and the next terminator is ≤30 nt (unclear function of UAG)	True stop signals (no conservation of sequences downstream of UAG)	Distance between UAG and the next terminator is >30 nt			Other genes (unclear function of UAG)
				Candidate Pyl-containing genes			
				Candidate Pyl-containing genes represented by single sequences (conservation of sequences downstream of UAG > 50%)	Candidate Pyl-containing genes (at least two UAG-containing forms)	Candidate Pyl-containing genes (at least three UAG-containing forms)	
Pyl-utilizing archaea							
<i>M. acetivorans</i>	224	96 (42.9%)	0 (0.0%)	35 (15.6%, 7 methyltransferases + 4 transposases + 24 other genes)	11 (7 methyltransferases + 4 transposases)	7 (7 methyltransferases)	103 (46.0%)
<i>M. mazei</i>	126	56 (44.4%)	0 (0.0%)	39 (31.0%, 7 methyltransferases + 16 transposases + 7 other genes)	23 (7 methyltransferases + 16 transposases)	7 (7 methyltransferases)	31 (24.6%)
<i>M. burtonii</i>	131	42 (32.1%)	0 (0.0%)	43 (32.8%, 6 methyltransferases + 37 other genes)	6 (6 methyltransferases)	6 (6 methyltransferases)	46 (35.1%)
Total	481	194 (40.3%)	0 (0.0%)	117 (24.3%)	40 (8.3%)	20 (4.2%)	180 (37.4%)
Archaea that do not utilize Pyl							
<i>M. jannaschii</i>	164	80 (48.8%)	52 (31.7%)	5 (3.0%)	0 (0.0%)	0 (0.0%)	26 (11.0%)
Pyl-utilizing bacteria							
<i>D. hafniense</i>	1127	390 (34.6%)	221 (19.6%)	33 (2.9%, including <i>mttB</i>)	1 (<i>mttB</i>)	1 (<i>mttB</i>)	483 (42.9%)

TABLE IV. Analysis of overlapping genes in Pyl-utilizing archaea.

Organism	Codon	Number of genes ^a	Overlap threshold (nt) ^b	Overlapped genes (prior to extension) ^c	Overlapped genes (after extension) ^d	Gene fusion caused by extension ^e	Genes that overlap beyond threshold ^f	Longest overlap (nt)
<i>M. acetivorans</i>	UAG	224	98	15 (6.7%)	42 (18.8%)	11 (4.9%)	3 (1.3%)	172
	UAA	2217		123 (5.5%)	495 (22.3%)	132 (6.0%)	117 (5.3%)	464
	UGA	2099		332 (15.8)	643 (30.6%)	102 (4.9%)	195 (9.3%)	1934
<i>M. mazei</i>	UAG	126	241	5 (4.0%)	48 (38.1%)	30 (23.8%)	0 (0.0%)	434
	UAA	1800		86 (4.8%)	390 (21.7%)	105 (5.8%)	8 (0.4%)	1081
	UGA	1445		197 (13.6%)	400 (27.7%)	66 (4.6%)	28 (1.9%)	

^a Number of predicted genes containing corresponding in-frame stop codons.

^b Longest overlap between two genes in the genome.

^c Number of genes that overlap in the genome.

^d Number of genes that overlap considering read-through function of predicted stop codons.

^e Number of genes that become fused with downstream genes (*i.e.*, both genes are in the same frame and are separated by stop codon).

^f Number of genes that overlap beyond the threshold for the organism considering read-through function of stop codon.

Interestingly, there are also two non-identical RFIs in *Euplotes* (38, 39). An attractive hypothesis is that codon reassignment involves extensive mutational alteration of release factors, including duplication of their corresponding genes. If so, the fact that there are two release factors in some Pyl-encoding archaea can be also used to support the proposition that UAG function might be partially or fully reassigned from stop to Pyl.

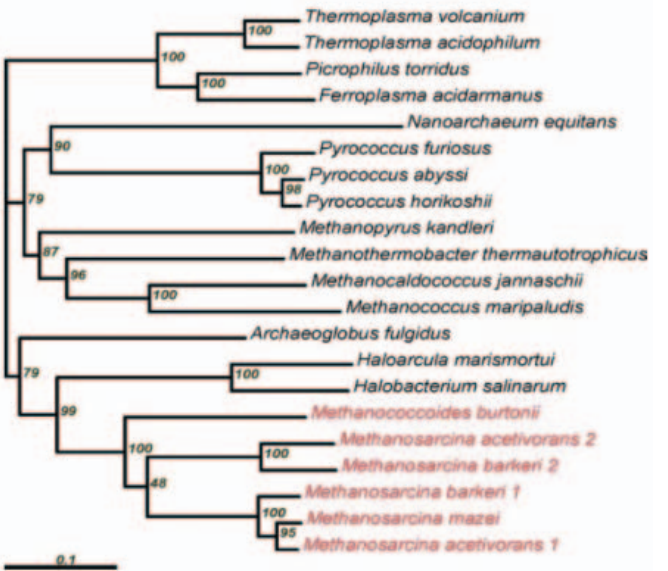


FIGURE 6. Phylogenetic analysis of RF1s from *Euryarchaeota*. Names of Pyl-utilizing organisms are highlighted in red.

Putative Mechanisms for Discrimination between Pyl Decoding and Stop Signal Functions of UAG Codons

The several lines of evidence described above imply that, at least in Pyl-decoding archaea, UAG codons do not function as standard terminators. Instead, there appears to be either a complete reassignment of UAG codons from stop to Pyl, or a competition between Pyl insertion and translation termination that favors Pyl insertion.

Reassignment of codon meaning has been documented for genetic codes of several prokaryotic and eukaryotic organisms, and it is highly common in genetic codes of organelles (34, 40, 41). Most frequently stop rather than sense codons are reassigned. This is probably because it is less deleterious for a cell, because stop codon usage is lower than the usage of sense codons. In addition, terminators generally do not correspond to crucial parts of the protein as is the case with many sense codons. In case of UAG, a complete reassignment of this terminator to Pyl codon does not seem to result in adverse functional consequences as there only a few UAG codons in Pyl-utilizing archaea and many of these would simply be slightly extended at the C terminus if UAG codes for Pyl.

The second possibility is that the meaning of UAG is ambiguous and specifies both termination and Pyl, making UAG the polysemous codon in methanogenic archaea. There is one known example of such ambiguous meaning of a codon. In several members of the *Candida* species a standard leucine (Leu) CUG codon is translated as serine (Ser). tRNA^{Ser}_{Cag}, which has 1-methylguanosine at position 37, can be charged with Leu both *in vitro* and *in vivo* and incorporate both Ser and Leu at CUG codons in these organisms (42, 43). A recent genetic exercise to manipulate the editing activity of *E. coli* isoleucyl-tRNA synthetase has generated a strain of *E. coli* where Cys is misincorporated at isoleucine (Ile) codons (44). This results in generation of “statistical proteins” where individual members of a protein pool have either Cys

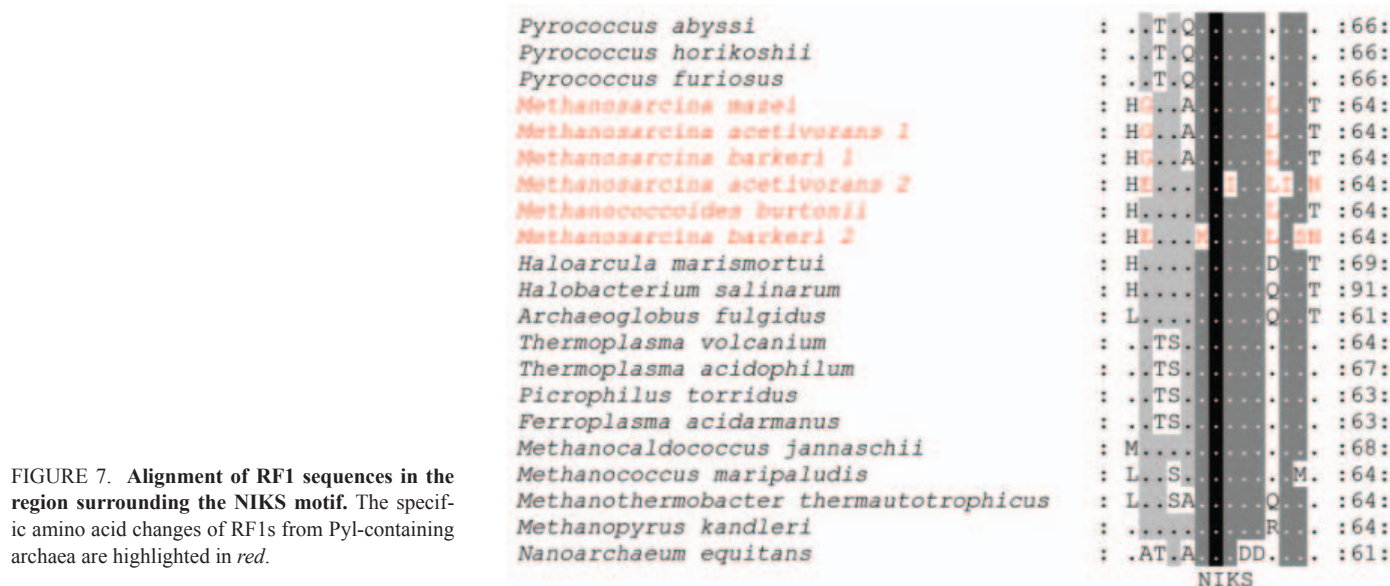


FIGURE 7. Alignment of RF1 sequences in the region surrounding the NIKS motif. The specific amino acid changes of RF1s from Pyl-containing archaea are highlighted in red.

or Ile in the same positions. It has been shown that generation of heterogeneous proteins could have a growth yield advantage and be beneficial at some steps of evolution (42–44).

An attractive possibility is that the competition between termination and Pyl insertion for UAG codons is influenced by physiological stimuli, e.g. this is a regulated competition or a partial reassignment of UAG codons. Considering that only one new family of candidate Pyl-containing proteins (transposases) could be identified, and it occurred only in two archaea, methylamine methyltransferases emerged as the key enzymes that utilize Pyl. MtmB and MtbB families of methylamine methyltransferases strictly conserve Pyl, whereas the occurrence of Pyl-containing *mttB* methylamine methyltransferases strictly matches that of the Pyl trait. Thus, it seems likely that Pyl insertion is maintained, because it is required for use by these enzymes. It is possible that the efficiency with which UAG is decoded as Pyl depends on specific environmental conditions, such as growth on methylamines (Figure 8). In this case, methylamines may activate Pyl biosynthesis and expression of methylamine methyltransferase genes. Under these conditions, Pyl insertion may out-compete translation termination at many or all UAG codons. However, because methylamine methyltransferases are extremely abundant enzymes (1–3), Pyl will primarily serve the methyltransferase UAG codons.

It is also possible that the rate with which Pyl is inserted at UAG codons is gene-specific, e.g. it depends on strength or leakiness of the UAG codon as a terminator. It is known that stop signals are extended elements with upstream and downstream sequences contributing to efficiency of decoding, terminating, and read-through functions (45–47). These stop-codon-flanking sequences were reported to cross-link with release factors and influence termination efficiency (48, 49). Thus, the context of UAG codons may be an important feature that influences the outcome of UAG decoding. However, the small number of true Pyl-containing proteins (e.g. methyltransferases) was insufficient for us to make definitive conclusions regarding the presence of sequence signals than flank UAG and discriminate between the two coding functions of UAG codons.

If metabolites influence competition between Pyl insertion and termination, in the absence of methylamines, Pyl might not be synthesized, and termination at UAG codons may prevail, because there would be no Pyl-tRNA^{Pyl} to compete with termination. If so, many UAG codons could serve as stop signals under certain growth conditions (e.g. no methylamines), but will insert Pyl when methylamines are present. This hypothesis is also consistent with the identical pat-

terns of occurrence of methylamine methyltransferase genes (*mtmB* and *mtbB*) and the Pyl cluster.

In any case, the data suggest that there appears to be no need for a highly specific PYLIS element in archaeal Pyl-containing mRNAs. For example, if UAG codons are primarily decoded as Pyl, there might be only a small fraction of truncated forms of methyltransferases due to competition with termination. In addition, insertion of Pyl at other UAG codons should be well tolerated, because this would result in only slight increases in protein masses. At the same time, the small number of UAG-containing genes in Pyl-decoding archaea suggests that Pyl is not a common amino acid in protein in these organisms. Presumably, either utilization of Pyl could affect functions of some proteins, or the flux through the Pyl biosynthetic pathway could not satisfy the demand for this amino acid, if it is to be commonly used. Further research is needed to address these hypotheses.

Searches for Genes Associated with the Pyl Trait

The availability of multiple genomic sequences of Pyl-utilizing organisms provides an opportunity to identify genes associated with the Pyl trait by analyzing patterns of gene occurrence. Our strategy

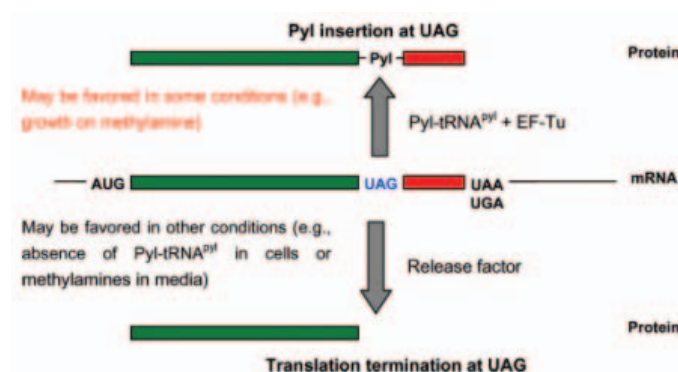


FIGURE 8. A model of Pyl insertion and termination of translation at UAG codons in Pyl-utilizing archaea. Decoding UAG codons as Pyl may be predominant under conditions that require Pyl (e.g. growth on methylamines), whereas in other situations, UAG codons may serve as terminators. Such competition between Pyl insertion and translation termination may occur in many or all UAG-containing genes, or it may be heavily shifted toward exclusive use of UAG as Pyl codon.

TABLE V. Identification of proteins associated with the Pyl trait

Organism	Annotated proteins	Proteins present in Pyl-utilizing archaea	Proteins present in Pyl-utilizing archaea and <i>D. hafniense</i>
<i>M. acetivorans</i>	4,540	226	8
<i>M. mazei</i>	3,371	188	7
Total	7,911	414 (168 protein families/groups)	15 (9 protein families/groups)

TABLE VI. Comparison of Sec and Pyl biosynthesis and insertion

Feature	Selenocysteine	Pyrrolysine
Distribution		
Archaea	<i>M. jannaschii</i> , <i>M. kandleri</i> , and <i>M. maripaludis</i>	<i>Methanosarcina</i> , <i>M. burtonii</i>
Bacteria	Widespread	<i>D. hafniense</i>
Eukaryotes	Widespread	Not found
Protein families	Diverse families (26 eukaryotic and 25 prokaryotic families)	Methylamine methyltransferases (MtmB, MtbB, MttB), transposases
Biosynthetic pathway		
Approach	Sec synthesized on tRNA ^{Sec} , which first aminoacylated with Ser	Pyl synthesized prior to aminoacylation to tRNA ^{Pyl}
Synthesis	Ser-tRNA ^{Sec} → Sec-tRNA ^{Sec}	Pyl + tRNA ^{Pyl} → Pyl-tRNA ^{Pyl}
Key enzymes	SelA and SelD (SPS)	PylS, PylB, PylC, and PylD
Non-canonical tRNA	tRNA ^{Sec}	tRNA ^{Pyl}
Decoding requirements		
Codon	UGA	UAG
Elongation factor	SelB (EFSec)	EF-Tu
cis-Element in mRNA	SECIS element	Not found
trans-Element	SelB or EFSec/SBP2	Unknown
Other		
Location	Active sites of various selenoproteins	Active site of MtmB
Function	Sec is a strong nucleophile and serves redox function	Electrophile in MtmB
Homologs	Conserved Cys-containing homologs	No other conserved amino acid found in homologs

was to test each of the annotated genes present in *M. acetivorans* and *M. mazei* (a total of 7911 genes) for their possible exclusive presence in Pyl-utilizing organisms (Table V). We first analyzed the presence of these genes in Pyl-utilizing archaea and absence in other archaea. This search identified 414 genes, which belonged to 168 protein families and included *mtmB*, *mtbB*, *pylS*, and *pylD*. These sequences were then analyzed for their occurrence in *D. hafniense*. Interestingly, only 15 proteins, which belonged to 9 families, were identified, including PylS, PylD, and a family represented by NP_632172 [GenBank] in *M. mazei*, which belonged to COG2043 (uncharacterized protein conserved in archaea). Other protein families were represented by hypothetical proteins with no known function or conserved motifs, including NP_618552 [GenBank] (*M. acetivorans*), NP_615464 [GenBank] (*M. acetivorans*), NP_617162 [GenBank] (*M. acetivorans*), NP_618396 [GenBank] (*M. acetivorans*), NP_632241 [GenBank] (*M. mazei*), and NP_633144 [GenBank] (*M. mazei*). The identified sequences may contain proteins associated with the Pyl trait. The possible functions may include roles in Pyl biosynthesis, insertion, or degradation, as well as functions associated with regulation and utilization of Pyl and Pyl-containing proteins. Since 4 out of the 15 proteins were true positives (PylS and PylD in each genome), our approach is clearly capable of identifying Pyl-associated genes. The availability of additional genomes with the Pyl trait will help in further characterization of the identified genes.

Parallels and Differences in Sec and Pyl Insertion Systems

A hypothesis of a parallel between Sec and Pyl insertion systems was suggested (3, 9, 50) based on similarities at various steps in the two translation pathways. However, our analyses described above argue against this possibility. We further analyzed and compared the known features of Pyl and Sec biosynthesis and insertion (Table VI).

Distribution of Sec and Pyl Traits—Sec is used by approximately a quarter of prokaryotic organisms, which have been characterized by genome sequencing, and by many eukaryotic organisms, with the exception of yeasts and higher plants (1, 13, 21). At least 26 eukaryotic and 25 prokaryotic selenoprotein families are known; most are present in proteins with distinct structures and functions (20, 21). This relatively widespread use of Sec contrasts with the rare occurrence of Pyl. We compared the Sec and Pyl decoding traits and found that only one organism, *D. hafniense*, utilizes both rare amino acids. Searches against all known selenoproteins identified only one selenoprotein, formate dehydrogenase α subunit, in *D. hafniense* (data not shown). Formate dehydrogenases contain one Sec, which is present in the active site, coordinates molybdenum, and is directly involved in oxidation of formate to carbon dioxide (51, 52). Thus, it appears that *D. hafniense* has only one Sec residue and one Pyl residue in its set of proteins. Whereas the use of Pyl and Sec is limited in *D. hafniense*, this organism appears to use all 22 natural amino acids currently known. The use of both Sec and Pyl by this organism is clearly supported by the presence of corresponding biosynthetic and insertion machineries.

Comparison of Sec and Pyl Biosynthetic Pathways—One distinctive feature of Sec biosynthesis is that its synthesis occurs on tRNA^{Sec} (the *selC* gene product) (53, 54). The tRNA^{Sec} has features that distinguish it from canonical tRNAs, including its length (typically 90 nucleotides; more than any other tRNAs), few post-transcriptional modifications, an unusually long variable arm, and the presence of 13 nucleotides in the acceptor and T Ψ C stems (13, 55, 56). Serine is initially acylated to tRNA^{Sec} by seryl-tRNA synthetase and then the *selA* gene product, Sec synthase, converts Ser-tRNA^{Sec} to Sec-tRNA^{Sec} in a two-step reaction (54). The selenophosphate donor for this reaction is synthesized by the *selD* gene product, selenophosphate synthetase (54,

57). Although the biosynthesis of Pyl has not been completely characterized, *pylB*, *-C*, and *-D* genes are candidates to play a role in Pyl synthesis (5, 58–60). Recent data suggest that, like the 20 common amino acids, Pyl is synthesized prior to its attachment to tRNA^{Pyl} (5, 6). The *pylS* gene product is dedicated to acylating Pyl to tRNA^{Pyl}. Direct charging of Pyl onto its tRNA contrasts with the tRNA-based pathway in Sec biosynthesis. In addition, like some canonical tRNAs, tRNA^{Pyl} lacks the variable stem.

Comparison of Sec and Pyl Insertion Pathways—The mechanism of Sec insertion in response to UGA has been most thoroughly elucidated in *E. coli* (53, 61, 62). Sec-tRNA^{Sec} forms a complex with the Sec-specific elongation factor SelB and GTP, and subsequently binds the SECIS element within ribosome-bound mRNAs (10). The resulting quaternary complex directs the insertion of Sec at in-frame UGA codons (54, 63, 64). The mechanism of Sec insertion in archaea and eukaryotes differs from that in bacteria in two aspects: (i) in bacteria, the SECIS element is located immediately downstream of the in-frame UGA codon, but it is present in 3'-UTRs in archaea and eukaryotes; (ii) SelB in bacteria binds both Sec-tRNA^{Sec} and SECIS, whereas in eukaryotes (and possibly archaea), SelB (EFSec) binds Sec-tRNA^{Sec} and is associated with SECIS via SECIS-binding protein 2 (SBP2) (13, 65, 66). Release factor RF2 can recognize UGA efficiently if any component of the Sec incorporation machinery is missing (45).

Understanding the mechanism of Pyl insertion has lagged behind that of Sec. As discussed above, we neither could identify true UAG stop codons nor find conserved stem-loop structures (PYLIS) immediately downstream of the in-frame UAG codons or in UTRs of methyltransferase genes in methanogenic archaea. Our data, viewed as a whole, argue for differences between Sec and Pyl insertion pathways.

CONCLUSIONS

The nature of the processes that extend the universal genetic code is still being debated. In recent years, many non-standard amino acids in proteins have been identified (67–69), but almost all are formed by post-translational modifications. Sec and Pyl are the only two exceptions identified to date. Both are encoded by canonical stop codons using specific tRNAs. These properties may have evolutionary significance in regard to the extension of the genetic code (8, 50).

In this study, various genomic sequences were scanned to identify and scrutinize Pyl-utilizing organisms and Pyl-containing proteins. Analysis of the distribution of stop codons revealed that UAG is a rare codon in Pyl-encoding archaea, but common in the bacterium *D. hafniense*. It appears that in *D. hafniense* UAG codons must serve two functions, stop and Pyl insertion. However, having only a single Pyl-containing protein in this organism precludes computational searches for possible *cis*-elements that modify the function of the UAG codon. In contrast, in Pyl-utilizing archaea, the lack of unambiguous candidates for UAG stop codons, the relatively large proportion of genes that could only slightly be extended beyond UAG and the acceptable overlaps for extended UAG-containing genes suggest that UAG codons might be decoded as Pyl in many or all UAG-containing genes. These features also suggest that insertion of Pyl could be well tolerated in these organisms, even though the number of proteins that require Pyl for their function is small and might be limited to methylamine methyltransferases. Thus, although both Sec and Pyl insertion may compete with termination, there is a difference between decoding strategies employed by Pyl and Sec, with Pyl resembling the standard amino acids and Sec having its own biosynthetic and decoding mechanisms.

Our findings are also consistent with analyses of release factor sequences in archaea as well as with the unsuccessful search for PYLIS elements in methylamine methyltransferase genes. If some discrimination between Pyl decoding and stop signal functions of UAG codons is needed, it could potentially be provided by the context of the UAG codons or structural elements that inhibit termination thus allowing Pyl

insertion. Alternatively, UAG codons could serve as Pyl codons under conditions that stimulate Pyl synthesis (e.g. during growth of cells on methylamines) by out-competing the weak terminator function of UAG. However, in situations where methyltransferase genes are not required, Pyl biosynthesis may be inhibited resulting in depletion of the pool of Pyl-tRNA^{Pyl}, which would favor termination at UAG codons. For most genes, Pyl insertion or termination may be nearly equivalent, as gene products would only slightly differ in C-terminal regions. Combined with the fact that only a small number of genes utilize UAG codons in archaea, this change in UAG coding function could be well tolerated by these organisms.

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