Identification of the yeast gene encoding the tRNA m¹G methyltransferase responsible for modification at position 9

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ABSTRACT

Methylation of tRNA at the N-1 position of guanosine to form m¹G occurs widely in nature. It occurs at position 37 in tRNAs from all three kingdoms, and the methyltransferase that catalyzes this reaction is known from previous work of others to be critically important for cell growth in *Escherichia coli* and the yeast *Saccharomyces cerevisiae*. m¹G is also widely found at position 9 in eukaryotic tRNAs, but the corresponding methyltransferase was unknown. We have used a biochemical genomics approach with a collection of purified yeast GST-ORF fusion proteins to show that m¹G₉ formation of yeast tRNA^{GIy} is associated with ORF YOL093w, named *TRM10*. Extracts lacking Trm10p have undetectable levels of m¹G₉ methyltransferase activity but retain normal m¹G₃₇ methyltransferase activity. Yeast Trm10p purified from *E. coli* quantitatively modifies the G₉ position of tRNA^{GIy} in an *S*-adenosylmethionine-dependent fashion. Trm10p is responsible in vivo for most if not all m¹G₉ modification of tRNAs, based on two results: tRNA^{GIy} purified from a *trm10-\Delta/trm10-\Delta* strain is lacking detectable m¹G; and a primer extension block occurring at m¹G₉ is removed in *trm10-\Delta/trm10-\Delta/trm10-\Delta* strains. Trm10p bears no detectable resemblance to the yeast m¹G₃₇ methyltransferase, Trm5p, or its orthologs. Trm10p homologs are found widely in eukaryotes and many archaea, with multiple homologs in several metazoans, including at least three in humans.

Keywords: Saccharomyces cerevisiae; S-adenosylmethionine; TRM10; YOL093w; 1-methylguanosine

INTRODUCTION

tRNA molecules are extensively modified after transcription. In addition to 5'- and 3'-end maturation, CCA addition, and splicing of introns, all tRNAs contain a number of nucleoside modifications (Bjork 1995; Grosjean et al. 1997; Hopper and Phizicky 2003). In *Saccharomyces cerevisiae*, 25 different modifications of base or sugar moieties have been identified within the 34 tRNA species that have been characterized (Sprinzl et al. 1998), and every tRNA molecule has some subset of these modifications, altering nucleoside identity at an average of 11 positions. Some modifications such as i⁶A or m₂²G occur only at a single position in one or more tRNAs, whereas other modifications such as Ψ

occur at a number of different positions in tRNAs (Sprinzl et al. 1998). The identity and position of many of these modifications are highly conserved; however, despite their widespread conservation, the role that these modifications play in the cell is not fully understood.

One modification of interest is methylation of guanosine (G) at the N-1 position of the base (m¹G; Fig. 1A). This modification occurs at two locations in tRNAs. m¹G is found at position 37 in several tRNAs from all three major kingdoms. It is also found at position 9 in several cytoplasmic tRNAs from all eukaryotes that have sequenced tRNAs, in mitochondrial tRNAs from several animal species, in two vertebrate viral tRNAs, and in tRNA from at least one archaeal species (Sprinzl et al. 1998). In the yeast *S. cerevisiae*, m¹G is found at position 9 in 10 tRNAs and at position 37 in 8 tRNAs, with one species, tRNA^{Pro}, having m¹G at both positions. The methyltransferase proteins responsible for m¹G₃₇ formation are critical in both bacteria and yeast; *Salmonella typhimurium trmD* mutants and *S. cerevisiae*

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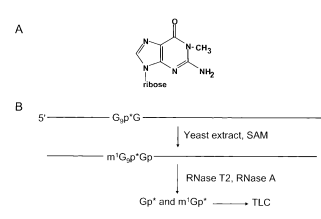


FIGURE 1. (*A*) N1-Methylguanosine. (*B*) Assay scheme to detect $m^{1}G$ formation in tRNA^{Gly} uniquely labeled at position 9 (G₉*Gly). This assay results in the production of either Gp* if the substrate remains unmodified, or $m^{1}Gp^{*}$ if it is modified.

trm5 mutants are each severely compromised for growth (Bjork et al. 2001).

The goal of this work was to identify the gene and protein responsible for m^1G_9 formation in *S. cerevisiae* tRNA. It is known that the yeast m^1G_{37} methyltransferase plays no role in m^1G_9 formation in yeast (Bjork et al. 2001). Furthermore, unlike the case with many other modification enzymes, there is no obvious candidate gene in the yeast database with homology to the known m^1G_{37} methyltransferases (Bjork et al. 2001) that might be responsible for m^1G_9 methylation of tRNAs. Using a biochemical genomics ap-

proach, we show here that the protein encoded by ORF YOL093w catalyzes tRNA m¹G methyltransferase activity in vitro at position 9 of tRNA substrates, and that this protein (Trm10p) is responsible for m¹G formation at position 9 in yeast tRNAs in vivo.

RESULTS

To determine the protein responsible for catalyzing the modification of tRNA at G_9 to m¹G, we first developed a sensitive assay using yeast tRNA^{Gly}, a tRNA that is known to contain m¹G₉ in vivo (Sprinzl et al. 1998). This tRNA was synthesized in vitro and then uniquely labeled at the phosphate immediately 3' of G₉, by directing cleavage at this position with RNase H, labeling the 5' nucleotide of the resulting 3' RNA product, and reforming the tRNA by ligation, as described in Materials and Methods (Yu 1999). After incubation of G_{o}^{*} Gly tRNA with a source of m¹ G_{o} methyltransferase activity (crude yeast

extracts or purified protein) in the presence of the methyl donor *S*-adenosylmethionine (SAM), RNA was treated with a combination of RNase T2 and RNase A, and nucleoside 3'-monophosphate products were resolved on cellulose TLC plates (Fig. 1B). Because G_9 was the only labeled nucleotide in this substrate, modification of the tRNA at this position is easily detected by looking for a change in mobility of the labeled spot. As shown in Figure 2A (lanes a and b), treatment of tRNA with yeast crude extract results in the quantitative conversion of Gp* to a modified product that migrates significantly more slowly in the solvent system. This modification was presumed to be m¹Gp*, which will be demonstrated below.

Identification of a yeast ORF associated with an activity that modifies position 9 of tRNA^{Gly}

To identify the protein responsible for this modification activity, we screened a library of purified yeast GST-ORF fusion proteins for the activity, as has been done for a number of other activities (Martzen et al. 1999; Alexandrov et al. 2002; Hazbun and Fields 2002; Xing et al. 2002). This library is an ordered collection of 6144 yeast strains each expressing a predicted yeast open reading frame (ORF) as a GST-ORF fusion protein. For ease of assaying more than 6000 proteins, the library of strains is grown in pools derived from each microtiter plate, such that after purification each pool contains a mixture of 96 individual GST-ORF fusion proteins. After activity is observed in one pool, fur-

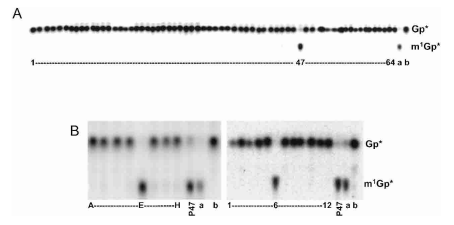


FIGURE 2. Identification of a yeast ORF associated with m^1G_9 methyltransferase activity. Reaction mixtures containing G_9^* Gly tRNA and protein as indicated were incubated in methyltransferase buffer at 30°C for 4 h, and then RNA was digested with nucleases to produce 3'-phosphorylated nucleotides, which were resolved by thin layer chromatography. (*A*) Assay of a genomic collection of pools of purified yeast GST-ORF fusion proteins for G_9 methyltransferase activity. Substrate G_9^* Gly tRNA was incubated with 64 pools of purified GST-ORF fusion proteins, each derived from 96 yeast strains of a library of strains expressing individual GST-ORF fusion proteins, as indicated. (Lane *a*) *Saccharomyces cerevisiae* crude extract; (lane *b*) no extract. (*B*) Assay of subpools from pool 47 for tRNA G_9 methyltransferase activity. (*First* panel) Substrate G_9^* Gly tRNA was incubated with GST-ORFs derived from the strains in rows *A*–*H* from plate 47, as indicated. (Lane *P47*) plate 47; (lane *a*) crude extract; and (lane *b*) no extract. (*Second* panel) Substrate was incubated with GST-ORFs from columns 1–12 of plate 47. (Lanes *P47*,*a*,*b*,) Same as previous panel.

ther deconvolution leads to the identification of the individual fusion protein responsible for the activity. When the GST-ORF pools were screened for the modification activity that altered G₉ of tRNA^{Gly}, a positive signal was observed in pool 47 (Fig. 2A). To determine the individual ORF responsible for this modification, we assayed 8 subpools of 12 proteins comprising each row (A-H) of plate 47 and 12 subpools of 8 proteins comprising each column (1-12) of plate 47. Activity was detected in row E and column 6 (Fig. 2B). To confirm the assignment of the activity to the strain at position E6, the corresponding strain (MRM4470) was grown, and extracts were analyzed for activity. Crude extracts made from this strain overexpress tRNA m¹G₉ methyltransferase activity at least fivefold relative to wild-type strains (data not shown), demonstrating that the corresponding ORF, YOL093w, is the limiting factor required for activity. The protein encoded by YOL093w has no previously identified function in yeast; we have assigned it the name TRM10 because, as we show below, its protein product is the tRNA m¹G₉ methyltransferase.

Trm10p is necessary for modification of position 9 of tRNA^{Gly} and has no effect on modification of position 37

To determine if Trm10p is required for modification at position 9, we prepared extracts from the homozygous $trm10-\Delta/trm10-\Delta$ strain and its isogenic $TRM10^+/TRM10^+$ parent strain, and compared their activities with the tRNA^{Gly} substrate (G₉*Gly) used above. Extracts from wild-type cells were active with the G₉*Gly substrate when diluted as much as 125-fold, whereas G₉ modification activity was undetectable in the $trm10-\Delta/trm10-\Delta$ strain even at the highest concentration of extract tested (Fig. 3A). This is consistent with the identification activity.

Because yeast tRNA also contains m¹G at position 37 of some tRNA species, we investigated the site specificity of the

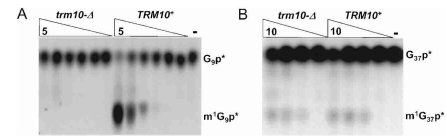


FIGURE 3. Assay of a $trm10-\Delta/trm10-\Delta$ strain for methyltransferase activity at positions 9 and 37. Extracts were prepared from isogenic wild-type $(TRM10^+/TRM10^+)$ and $trm10-\Delta/trm10-\Delta$ strains, and assayed for methyltransferase activites. (A) m¹G₉ methyltransferase activity. Assays contained G₉*Gly tRNA and decreasing concentrations of crude extracts (5 mg/mL to 1.6 µg/mL by factors of 5) made from wild-type and $trm10-\Delta/trm10-\Delta$ yeast strains and were carried out at 30°C for 1 h. (B) m¹G₃₇ methyltransferase activity. Assays contained G₃₇*Leu tRNA with decreasing concentrations of crude extracts from wild-type and $trm10-\Delta/trm10-\Delta$ yeast strains (2.5 mg/mL to 2.5 µg/mL by factors of 10) and were carried out at 30°C for 5 h.

reaction using a tRNA_{CAA}^{Leu} substrate normally modified by Trm5p to m¹G at position 37. Although our in vitro assay for Trm5p activity uses a similar uniquely labeled substrate (G₃₇*Leu) and similar conditions, we consistently observe much lower levels of Trm5 methyltransferase activity than we observe with Trm10p and its substrate. Nonetheless, m¹G₃₇ methyltransferase activity can be observed by using longer incubation times (Fig. 3B). Assay of the wildtype and $trm10-\Delta/trm10-\Delta$ extracts with the G₃₇*Leu substrate indicates no difference in m¹G₃₇ methyltransferase activity between them, as expected if the modification activity of Trm10p is specific for position 9 (Fig. 3B). We note that because the modification at G₃₇ of tRNA_{CAA} leu is known to be m¹G (Bjork et al. 2001), the identical migration of the products observed with the two different substrates is consistent with the modification at position 9 also being m¹G.

Trm10p is sufficient for m¹G₉ modification in vitro

To determine if Trm10p could catalyze modification of tRNA without additional S. cerevisiae proteins, the ORF was cloned into a pET14b-based vector for production of the protein in Escherichia coli. This plasmid, pJEJ12-3, expresses soluble His₆-Trm10 fusion protein when introduced into BL21(DE3)pLysS cells and induced for expression, as observed by SDS-PAGE of extracts (Fig. 4A, lanes 1,2). The prominent new band in lane 2 has a molecular mass consistent with the predicted molecular mass of 35.5 kD for the His₆-Trm10 fusion protein. Assays of extracts demonstrated a >1000-fold increase in G₉*Gly modification activity from cells expressing His₆-Trm10p compared with cells with vector alone (Fig. 4B). As shown in Figure 4B, there was no background m¹G_o methyltransferase activity in extracts made from cells containing the control vector; this is as expected because E. coli cells do not contain an m¹G₉ modification in their tRNAs.

The His₆-Trm10 fusion protein was purified from ex-

tracts with immobilized metal-ion affinity chromatography (IMAC; Fig. 4A, lanes 5,6). The purified protein is nearly homogeneous, has an increased specific activity of about twofold over the original extract (Fig. 4A, lanes 4-6), and is obtained at 30% yield (data not shown), indicating that Trm10p is sufficient for catalysis in vitro in the absence of other interacting partners. The activity of the purified protein from E. coli is completely dependent on the addition of 0.5 mM SAM in the assay, although this dependence is only apparent at low enzyme concentrations, where the enzyme is required to undergo multiple turnovers during the course of the reaction

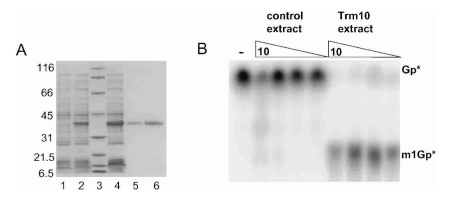


FIGURE 4. Overexpression and purification of Trm10p from *Escherichia coli.* (*A*) SDS-PAGE gel of Trm10p overproduction and purification. (Lane 1) Crude extract from control strain with vector only (40 μ g); (lane 2) crude extract from strain with plasmid pJEJ12-3 expressing His₆-Trm10p (40 μ g); (lane 3) broad range MW markers; (lane 4) crude extract used for purification, from strain containing pJEJ12-3 (40 μ g); (lanes 5,6) purified Trm10p (4 and 8 μ g, respectively). (*B*) Assay of *E. coli* extracts for m¹G₉ methyltransferase activity. Assays containing G9*Gly tRNA and decreasing concentrations of either the control extract (panel *A*, lane 1) or Trm10p-expressing extract (panel *A*, lane 2) were performed at 30°C for 1 h.

(data not shown). This indicates that some amount of bound SAM copurifies with Trm10p and therefore that Trm10p is able to bind SAM with some degree of affinity.

m¹G is the product of Trm10p activity

To show that the product of Trm10p activity is m¹G, we modified in vitro transcribed tRNA^{GIy} with Trm10p purified from E. coli in a reaction mixture containing 10 µM tRNA and 10 µM Trm10p, and then analyzed nucleosides from the product tRNA by HPLC (Fig. 5A, bottom two traces). Unmodified tRNA^{Gly} contains four main peaks for the four expected nucleosides C, U, G, and A, whereas the Trm10p-modified tRNA contains an additional peak that comigrates with and has a spectrum identical to an m¹G nucleoside standard (Fig. 5A, upper trace). This evidence demonstrates that m¹G is the product of the Trm10p reaction. Quantification of the amount of m¹G indicates 1.1 mole of m¹G/mole tRNA based on the areas of the peaks observed before and after reaction with Trm10p, consistent with modification of a single nucleotide in tRNA^{Gly} to m¹G (Table 1).

TRM10 is responsible for m¹G₉ modification of tRNA^{Gly} in vivo

To determine if Trm10p is responsible for formation of m^1G_9 of tRNA^{Gly} in vivo, we purified tRNA^{Gly} from both wild-type and *trm10-\Delta/trm10-\Delta* cells, and analyzed the nucleosides by the same HPLC analysis. The peak for m^1G is evident in tRNA^{Gly} derived from the wild-type parent (Fig. 5B, upper trace) and absent in the tRNA^{Gly} derived from the *trm10-\Delta/trm10-\Delta* strain (Fig. 5B, lower trace). Other modified nucleosides known to be present in

tRNA^{Gly} (m⁵C and Cm) were identified based on retention time and spectrum of the respective peaks and are labeled in Figure 5B. Quantitation of the amount of modified nucleosides apparent in these tRNAs indicates that <1% m¹G remains in tRNAGly isolated from the $trm10-\Delta/trm10-\Delta$ cells, whereas all other nucleotide modifications remain essentially unchanged, except for dihydrouridine levels, which are slightly altered between the two samples but well within the variation observed because of its proximity to pseudouridine and lack of a clear absorption peak (Table 1). For tRNA from both wild-type and *trm10-* Δ strains, we reproducibly observe a higher than expected amount of m⁵C; this probably reflects an additional m⁵C modification in tRNAGly that has not

been previously reported. Thus, Trm10p activity is the only significant in vivo source of m¹G in tRNA^{Gly}, and its absence has little effect on other modifications.

TRM10 is responsible for m¹G₉ modification of other tRNAs in vivo

To examine the in vivo effect of Trm10p on m¹G modification of other tRNA species, we have used a primer extension assay. The presence of m¹G in a nucleotide sequence should result in generation of a primer extension block at the position immediately before an m¹G, because the N-1 methyl group of the m¹G residue impairs its ability to form the normal Watson-Crick base pair necessary for continued extension by reverse transcriptase (RT). Because primers used in this assay are specific for an individual tRNA species, this assay allows for rapid analysis of the modification status of multiple tRNAs in a single preparation of bulk RNA. We tested this method by probing tRNA^{Gly} from bulk preparations of RNA, using a primer that could hybridize to it at a region ending in the D-loop (shown in bold in Fig. 6A). With tRNA from wild-type cells, a block to primer extension was observed at position G₁₀ (corresponding to C in the sequencing ladder shown in Fig. 6A), because the presence of m^1G_9 has prevented RT from reading past the modified nucleotide. However, with RNA from the trm10- $\Delta/trm10-\Delta$ strain, this primer is extended through G_o to the 5'-end of the tRNA, as expected because of the presence of an unmodified G₉ in tRNA^{Gly} from this strain (Fig. 5B; Table 1).

The primer extension assay described above was used to analyze m^1G_9 content in 8 of the 9 other *S. cerevisiae* tRNA species known to contain m^1G_9 : tRNA^{Trp}, tRNA^{Pro}, tRNA^{Val}, tRNA^{Ile}, tRNA^{Ile}, tRNA^{ICG}, and both tRNA_{UCU}

А

 $\begin{array}{c} C \\ M^{5}C \\ C \\ M^{5}C \\ C \\ M^{5}C \\ M^{5}C \\ M^{1}G \\ M^{1}G$

FIGURE 5. Identification of m¹G as the product of Trm10p activity in vitro and in vivo. (*A*) Trm10 protein modifies tRNA^{Gly} to produce material that matches m¹G. (*Upper* trace) m¹G chemical standard. (*Middle* trace) In vitro transcribed tRNA^{Gly} treated with buffer and digested to nucleosides. (*Lower* trace) In vitro transcribed tRNA^{Gly} treated with Trm10p and digested to nucleosides. All samples were analyzed by HPLC as described in Materials and Methods. (*B*) Comparison of nucleosides of tRNA^{Gly} from wild-type and *trm10-*Δ/ *trm10-*Δ strains. tRNA^{Gly} was purified from wild-type and *trm10* mutant strains, and its nucleosides were prepared and resolved on HPLC as described above.

species. Primer extension assays with tRNA^{Ala} have been repeatedly unsuccessful for unknown reasons. With the eight tRNAs for which we were able to see primer extension products, the results were the same as that observed for tRNA^{Gly}. For example, primer extension of tRNA^{Val}, tRNA^{Trp}, and tRNA_{ICG}^{Arg} from wild-type cells is blocked before reaching the 5'-end of the RNA, at a position consistent with the presence of m¹G₉ in these tRNA species (Fig. 6B). This primer extension block is completely absent in tRNA from *trm10-* Δ /*trm10-* Δ cells, strongly implying a quantitative loss of m¹G₉ in these cells (Fig. 6B). The same results were observed with the other five tRNA species examined (data not shown). In contrast, control primer extension assays with primers targeting tRNAs that do not

TABLE 1.	Quantification	of nucleoside	modifications	to tRNA ^{Gly}
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Modified nucleoside	Number expected per mole of tRNA ^a	In vitro product of Trm10p ^b	tRNA ^{Gly} from wild type ^b	$tRNA^{Gly}$ from $trm10 - \Delta^{b}$
m ¹ G	1	1.1	0.98	< 0.01
m⁵C	1	_	1.7	1.8
Cm	1		1.0	1.0
DHU	2		0.92	1.4
pseudoU	4	—	3.9	3.9

^a(Sprinzl et al. 1998).

^bCalculated from areas under peaks observed upon HPLC elution of nucleosides, as described in Materials and Methods.

contain m¹G₉ exhibit only the primer extension stop at the 5'-end of the tRNA in RNA derived from either the wildtype or the *trm10-* Δ /*trm10-* Δ strain (Fig. 6B, tRNA_{UUU}^{Lys}). For tRNA_{ICG}^{Arg} (Fig. 6B) and a few of the other tRNA species (tRNA_i^{Met}, tRNA^{IIe}, and both tRNA_{UCU}^{Arg} species; data not shown), the block appeared as more of a stutter rather than as the block at a single site observed with tRNA^{Gly} (Fig. 6A), tRNA^{Val}, tRNA^{Trp} (Fig. 6B), and tRNA^{Pro} (data not shown). This may reflect a sequencespecific variability in the way AMV-RT deals with the presence of m¹G in various sequences; variation in the response of RT to sequences that act as blocks to primer extension has been observed previously (Lorsch et al. 1995). These results demonstrate that TRM10 is required for the catalysis of at least 9 of the 10 m¹G modifications observed at position 9 in yeast. It is likely that Trm10p is also responsible for modification of G₉ of tRNA^{Ala}.

Specificity of Trm10p activity in vivo

The primer extension assay has also been used to confirm that Trm10p is not involved in modification of G_{37} in vivo. Primers were used to target two tRNA species known to be substrates for Trm5p: tRNA_{UAG}^{Leu}, which contains m¹G₃₇, and tRNA^{Phe}, which contains Y₃₇ (wybutosine), the synthesis of which is initiated by formation of m¹G₃₇ by Trm5p (Bjork et al. 2001). Primer extension assays demonstrated that the extent of m¹G₃₇ modification is unchanged in strains lacking Trm10p (Fig. 6C). This result was confirmed by HPLC analysis of nucleosides generated from another tRNA known to contain m¹G₃₇, tRNA_{CAA}^{Leu}; the m¹G content of this tRNA is similar when purified from wild-type or *trm10-*Δ/*trm10-*Δ cells (data not shown).

Effects of loss of m¹G₉ on cell viability

As we have shown above, Trm10p is the major, if not the only, m¹G₉ tRNA methyltransferase acting in *S. cerevisiae* cells. However, *trm10-\Delta/trm10-\Delta* strains do not exhibit any obvious growth defect when compared with their isogenic *TRM10⁺/TRM10⁺* parent strain in rich media and in minimal media at several different temperatures (data not shown). In contrast, a *trm5-\Delta* strain grows extremely slowly (Bjork et al. 2001).

The Trm10 methyltransferase family is widely conserved, but is not homologous to the yeast $m^{1}G_{37}$ methyltransferase, Trm5

Iterative BLAST searches show that Trm10p is a member of a family of proteins found widely in all free-living eukaryotes (but not in the obligate parasite microsporidia, *Encephalitozoon cuniculi*) and half of all archaeal species sequenced (Fig. 7) but not in any Eubacteria. Indeed, this family of proteins has been described in both the PFAM

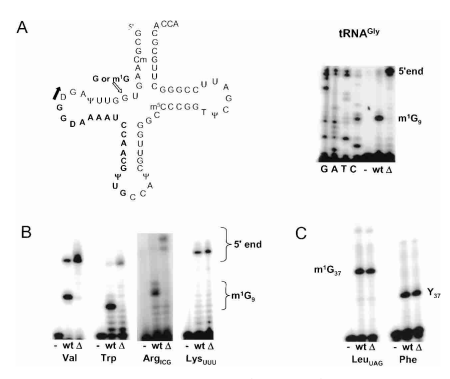


FIGURE 6. Deletion of *TRM10* results in loss of m¹G₉, but not m¹G₃₇ in vivo. (*A*) Establishment of primer extension assay to assess m¹G modification. (*Left* panel) Cloverleaf structure of tRNA^{Gly}. Nucleotides in bold indicate the position of the primer used for primer extension analysis of the 5'-end of the tRNA. (*Right* panel) Primer was used with in vitro transcribed tRNA^{Gly} and AMV Reverse Transcriptase to generate a sequencing ladder (lanes *G,A,T*, and *C*), and alone (–) or with RNA derived from wild type (wt) or *trm10-*Δ/*trm10-*Δ (Δ) strains to analyze for the presence of m¹G₉ in tRNA^{Gly} from each population of RNA. (*B*) Primer extension to assess modification status of tRNA^{Val}, tRNA^{Trp}, tRNA_{ICG}^{Arg}, and tRNA_{UUU}^{Lys} at position G₉. Primer extensions were performed with either (–) primer alone, (wt) wild-type RNA, or (Δ) *trm10-*Δ/*trm10-*Δ RNA. (*C*) Primer extension to assess modification status of tRNA_{UAG}.

database (DUF 425) and the COGs (Cluster of Orthologous Groups, COG2419) database as being conserved, albeit with no ascribed function. Unexpectedly, the database searches also revealed the presence of multiple *TRM10*-like homologs in each of several metazoan genomes. For instance, the human and mouse genomes harbor at least three homologs, whereas mosquito, fruit fly, and nematode genomes possess at least two. The occurrence of multiple Trm10p family members in metazoans is unexpected because Trm10p is responsible for most, if not all, of the m¹G₉ tRNA methylation observed in yeast (Fig. 6).

The presence of the archaeal and multiple metazoan homologs prompted a phylogenetic reconstruction of the evolution of this family, presented in Figure 8. In this phylogeny, the archaeal and eukaryotic homologs clearly group separately from one another. In addition, it is apparent that at least one of the lineages found in metazoans (human #3 in Fig. 8) appears to branch separately from those that group the human homologs #1 and #2 with yeast Trm10p. This implies two possibilities: either this lineage of proteins predates the human–yeast split; or it has rapidly evolved, causing its phylogenetic placement to be artifactual. Com-

parison of the human-mouse orthologs for each of lineages 1, 2, and 3 indicates that all three pairs are evolving under similar evolutionary constraints, supporting the argument against artifactual placement in the tree. Therefore, we conclude that there were at least two separate lineages of Trm10-like proteins in eukaryotes, only one of which has been retained in protists, fungi, and plants, whereas metazoans continue to harbor both. One of the Caenorhabditis elegans homologs branches even earlier than the two metazoan lineages, but this may be caused by rapid evolution rather than an ancient divergence, as is the case for a number of C. elegans genes (Blaxter et al. 1998).

Remarkably, database searches, pairwise BLAST, and comparisons of multiple alignments do not identify even any remote homology between Trm10p and Trm5p, the *S. cerevisiae* m^1G_{37} methyltransferase (Bjork et al. 2001; see Materials and Methods); thus, these two proteins are likely unrelated, despite their conserved biochemical activity (see also Discussion). We note also that Trm10p was not identified in a search for SAM-dependent methyltransferases in *S. cerevisiae* based on known SAM binding motifs (Niewmierzycka and Clarke 1999), and that none of our

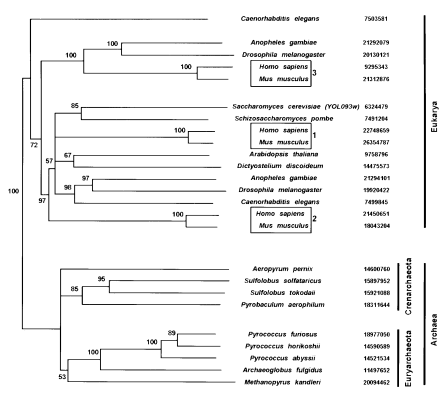
searches revealed any similarity with known methyltransferases. Because Trm10p requires SAM, this indicates that Trm10p may contain a more subtle or a different SAMbinding sequence.

DISCUSSION

We have shown that *S. cerevisiae* Trm10p, encoded by YOL093w, catalyzes the methylation of G residues to m¹G in tRNA^{Gly} at position 9. Using a substrate that is uniquely labeled with ³²P at the junction between residues 9 and 10, *S*-adenosylmethionine as a methyl donor, and assays of a genomic array of GST-fusion proteins, we have identified the corresponding protein and gene (Fig. 2). Cloning of this gene and subsequent overproduction of Trm10p activity in *E. coli* demonstrates that Trm10p alone is sufficient for catalysis of m¹G₉ methylation in vitro (Fig. 4). The product of the Trm10p reaction is m¹G, based on comparison of the TLC mobility of the RNase T2 digestion product m¹G p with that produced by modification of G₃₇ of tRNA^{Leu} to m¹G in extracts (Fig. 3), and on comparison of the HPLC mobility

MKKEADORKKAERDERKVAERRVERRKERENTHIIYGLGHTSLFLRIYDTTINHWONNRLTRAMOFAOKWVLDCS KVKKAROIKKEMKAAAREEAKNIKLLETTEEDKOKNKLERLWDRNMDIAMGWKGAOAMOFGOPLVFDMA KMCURORWEENKAKYNAERRVKKRELEHERSAKIOEKIDRGEEVPOELIREPRINVNOTDSGIETIDDCS KLIKOROWEEORELRKOKREKRKERKKELRHERSAKIOENSDGHDK.KRVRDVWHSTLRLIDCS KLIKOROWEEORELRKOKREKRKERKKORREAKIOEKRTGPSRKELKKRVRDVWHSTLRLIDCS KORLAEFAELRKLEREREKKROERREKKORREAKELGLPKRSEEKEKKIERKROLADGGKSGLSVAIDD GERKLKEWEETLANATEERLKLIERERKKORRREAKELGLPKRSEEKEKKIERKROLADGGKSGLSVAIDD GERKLKEWEETLANATEERLKLIESRKSDRRENNFENDVOTDSGELONOD GERKLKEWEETLANATEERLKLIESRKSDRRENNFENDFOONSSKELGEN VORKORHWEKTVAAKSKREGERERKKORRENSERREND SORTLECOMENTIERIEND SORTLEETLANATEERLEKKVEEREREKKORRENDEGERTOONSSKELGEN	YDEH, NNREATYAAK QLMMCFAENRMNDEPPDHTYCNTQMD SRCMQSAQRYIPTMHNPEPPINLHSKC. FTEL YENYK K KELONTVSQLLESEGWRRNVDPFHITTCNLKID GALHREVKRYGEKMD. KLLLTSTEKS. HVDL FDELWDKKLEVSLSNQVTRAKSANRRANHPAEIKVAPPEKLKOFFTTKNTNYENMHFKELPDDKIMFGDEH FDELWDKELKTK. FDELWDKKEHKQOCRTAKENRALHPVOFTTSHGGGLKKMDENDKGW. VNWKDIH IKPEHTSEL FDHLWVLKDIKKEHKQOCLRITTINRRSPQFGNKHFGIRR NGHIHESFKKN EGMENWH. VQTTFDKGHTDI FAHL SESEISSLVQQIMYCKAVNGRSTSPCHMLTGVQG KMSTELDKL. PGFEKWFIEKESRCTIEAMA FAHL SESEISSLVQQIMYCKAVNGRSTSPCHMLTGVQG KMSTELDKL	<pre>P R 0 NLVYLTPHCRED_VTYN . PDDITIVGANVD TMNNEPLSLAKAKRLG LRMARLPLDRTLOWGSG F P K 0 NLVYLTADS PNWTTFR . HDKVYVTGSFVD KSM0PGTSLAKAKRLN LATECLPLDKYLOWEIG I S KD K I VYLTADS PN WTTFR . HDK VYTGSFVD KNN YKELCLKKAG K KLN LATECLPLDKYLOWEIG I KKE D LIYLTSDS PN L KELD . ES KAYVTGGLVD KNN YKELCLKKAG K KG I PHACLPLDKYLOWEIG F F S C V VLT C ES D R VLDKLQ . PGCTYVIG GLVD KNN YKELCLKKAG K KG I PHACLPLD KYLLWEIG F F S C V VLT C ES D R VLDKLQ . PGCTYVIG GLVD KNN F KELCLKKAG K KG I TTARLPLD KYLLOWEIG F F S C V VLT C ES D R VLDKLQ . PGCTYVIG GLVD KNN F KG WF C G S C G C C S C F F A K KNN S F F I S C V VLT C ES D R VLDKLQ . PK HI YI I G C V V C F S A T X A S A T X A S C S C C S C F C C S C F V N N S F C C S C F C F C C F C C F C C F C C C C</pre>	<pre>SG KSLTLNOMINIMIDLKK TGDWDTALKHVPRRKVVONEFOPREKDH</pre>	FIGURE 7. Alignment of Trm10p homologs. The various Trm10 homologs were identified using iterative BLAST searches, and aligned using CLUSTALX. A representative alignment is shaded to a 50% consensus using MacBoxShade, and indicates residues that are similar (gray background) or identical across all species (black background). The numbers adjacent to the name of each species are the accession numbers for the proteins used. The numbers in parentheses before and after the alignment indicate the numbers of amino acids from individual sequences not included in the alignment for clarity of presentation.
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FIGURE 8. Phylogeny of the Trm10p homologs. A phylogeny of the various Trm10p homologs from eukaryotes and archaea was constructed using the neighbor-joining method, rooted on the archaeal homologs. Bootstrap analysis was carried out using the PAUP* version 4.0b10, and bootstrap support (percentage of 1000 trials where a certain grouping was supported) is indicated at the respective nodes. The tree is collapsed to a 50% bootstrap support. The source species of each protein is listed beside the appropriate line, together with the corresponding accession number.

and the spectrum of m^1G derived from Trm10p-modified tRNA^{Gly} with a commercial source of m^1G (Fig. 5).

Our data also show that Trm10p is specific for G_9 of tRNA substrates. Extracts from $trm10-\Delta/trm10-\Delta$ strains lack m¹G₉ methyltransferase activity but have m¹G₃₇ methyltransferase activity, as expected if Trm10p catalyzes modification of position 9, but not at position 37 (Fig. 3). In addition, because only 1 mole of m¹G is produced during modification of tRNA^{Gly} transcript in vitro (Table 1), because the modified site is known to be position 9 with the G₉*Gly substrate (Fig. 2), and because this is the only position known to be modified to m¹G in tRNA^{Gly}, we conclude that Trm10p primarily modifies G₉ in vitro.

Finally, we have shown that Trm10p is required in vivo for methylation of G_9 in 9 of the 10 yeast tRNAs known to have this modification, based on two results. First, a comparison of tRNA^{Gly} purified from wild-type and *trm10-* Δ cells shows that m¹G modification is dependent on Trm10p (Fig. 5). Second, RT pauses at position 10 in primer extension assays of 9 tRNAs species known to contain m¹G₉, but not if the tRNAs were isolated from an otherwise isogenic *trm10-* Δ /*trm10-* Δ strain (Fig. 6).

There is a clear demarcation of position specificity for m¹G modification by Trm10p at G₉, as shown here, and by Trm5p at G₃₇ of substrate tRNAs (Bjork et al. 2001). This strict site specificity for what is essentially the same chemical reaction is similar to that observed with S. cerevisiae Tad1p and Tad2p/Tad3p, which are responsible for formation of I₃₇ and I₃₄, respectively (Gerber et al. 1998; Gerber and Keller 1999), and with Pus4p and Pus6p, which are responsible for formation of Ψ_{55} and Ψ_{31} , respectively (Becker et al. 1997; Ansmant et al. 2001). Other tRNA modifications that occur at more than one position are catalyzed by proteins with region-specific modification activity, such as Pus3p, which forms Ψ_{38} and Ψ_{39} (Lecointe et al. 1998), and Trm7p, which 2'-O-methylates at positions 32 and 34 (Tsai et al. 2002). In contrast, some proteins exhibit promiscuous modification activity at multiple sites, such as Trm4p, which catalyzes m⁵C formation at positions 34, 40, 48, and 49 (Motorin and Grosjean 1999), and Pus1p, which catalyzes Ψ formation at up to eight widely different positions in tRNA substrates (Simos et al. 1996; Motorin et al. 1998).

The mechanistic basis for the site specificity and catalysis of m¹G modification by Trm5p and Trm10p is an in-

triguing question that remains to be addressed by further study. A search for homologs of each of these two S. cerevisiae proteins by BLAST yields a distinct family of enzymes for each one (Fig. 7; data not shown). Independent alignments of these two predicted families of m¹G methyltransferases yield core sequences of conserved amino acids within each group; however, none of these blocks of conserved sequence are shared between the two groups of enzymes. It is possible that Trm5p and Trm10p share a common chemical mechanism for m¹G methylation but that the amino acids in the active site are hard to identify. Alternatively, the two enzymes may catalyze m¹G modification in completely distinct ways. Further analysis of the mechanism of these two enzymes will help to address this question and may elucidate how Trm5p and Trm10p each recognize their different positions and tRNA substrates for m¹G methylation.

Another intriguing specificity question is how Trm10p is able to distinguish between tRNA species to be modified and those to remain unmodified at position 9. Every eukaryotic tRNA sequenced to date contains a purine at position 9, with the exception of two tRNA^{His} species, one from Drosophila and one from sheep. In S. cerevisiae position 9 is occupied most often by G (in 23 out of 34 tRNAs sequenced); however, the mere presence of a G at position 9 is not the determining factor for m¹G formation. In S. cerevisiae there are 10 tRNA species that contain m¹G₉ (tRNA^{Gly}, tRNA^{Trp}, tRNA^{Pro}, tRNA^{Val}, tRNA_{ICG}^{Arg}, both tRNA_{UCU}^{Arg} species, tRNA^{IIe}, tRNA_i^{Met}, and tRNA^{Ala}), but there are 12 other tRNA species that contain a G at position 9 that remains unmodified in vivo (Sprinzl et al. 1998). There is no obvious primary sequence similarity in the region of position 9 that would explain m¹G modification at this site. One possibility is that $m^{1}G_{9}$ is formed either in response to, or to participate in, a common structural feature. An interaction involving the residue at position 9 has been observed in the crystal structure of tRNA^{Phe}, in which a base-triple has been observed between an A at position 9 and the U₁₂-A₂₃ base pair via a reverse-Hoogsteen interaction between the non-Watson-Crick faces (N7 and N6) of A₉ and A₂₃ (Kim et al. 1974). An investigation of the sequences of the 22 G₉-containing elongator tRNAs from S. cerevisiae yields no sequence pattern for the base pair at position 12-23 that covaries with the modification status of G_9 that would indicate that m^1G_9 participates in a similar structural feature (Sprinzl et al. 1998). However, in eukaryotic initiator tRNAs position 9 is overwhelmingly found to be a G in eukaryotes (17 out of 18 tRNAs sequenced) and, moreover, that G is always modified to m¹G (Sprinzl et al. 1998; Marck and Grosjean 2002). Interestingly, the base pair at position 12–23 in the $m^{1}G_{9}$ -containing tRNA_i^{Met} species is invariably a G₁₂-C₂₃ base pair. Although the electron density of the m^1G_9 in the crystal structure of S. cerevisiae tRNAi^{Met} is diffuse and therefore a specific interaction cannot be proven, it does appear to be in the correct position and orientation to interact with the G12-C23 base pair in an analogous interaction to that observed between A₉-A₂₃-U₁₂ in tRNA^{Phe} (Basavappa and Sigler 1991). If such an interaction does occur, it may be that m¹G₉, although present in other tRNA species, plays its most important role in the correct structure or function of eukaryotic initiator tRNAs.

Trm10p is a highly conserved protein that is widely found in eukaryotes (Figs. 7, 8), consistent with the widespread occurrence of m^1G_9 in eukaryotic tRNAs. The finding of multiple paralogous proteins in some metazoans is unexpected, and raises the question of their origin and function. The fact that these multiple homologs are no more related to each other than to other members of the Trm10p family indicates that the duplication occurred prior to the divergence of eukaryotic species, implying that organisms such as yeast have lost the duplicated members after their divergence. It seems likely that at least one of the different paralogs in humans and other organisms is responsible for G_9 methylation of tRNAs. The other paralogs may have altered substrate specificity, contributing to m¹G modification of specific subclasses of tRNAs or other RNAs, or they may have altered tissue specificity; alternatively, they may have evolved other functions. The presence of related archaeal homologs implies that they too may be m^1G_9 methyltranferases, because at least one archaeal organism, *Halobacterium cutirubrum*, has tRNA with this modification (Gu et al. 1983), although whether *H. cutirubrum* has a Trm10p homolog cannot be ascertained at this point owing to lack of sequence information.

We note that the pattern of m¹G₉ modification in tRNAs from other eukaryotes is also not obvious, either by the subset of G_9 residues that are modified to m^1G_9 , or by the identity of the tRNA species that are modified to $m^{1}G_{9}$ in each organism. For example, the yeast tRNAs for Ala, Gly, Ile, Pro, Arg, Val, Trp, and initiator tRNA all contain $m^{1}G_{9}$, whereas in humans only Asn, Gln, and initiator tRNAs have been shown to have this modification, and at least one sequenced tRNA^{Gly} species has an unmodified G at position 9. All told, at least six other tRNAs (those for Asp, Glu, Met, Asn, Gln, and Thr) are modified with $m^{1}G_{0}$ in some eukaryotes, but not in yeast. This indicates that the enzymes from different eukaryotes have evolved ways to distinguish between different sets of tRNAs to be modified, and may provide an explanation for why there are multiple paralogs of Trm10p in some metazoans. Cloning and analysis of other eukaryotic Trm10p homologs, particularly from species with multiple paralogs, will help to address these issues of specificity.

It is noteworthy that the extended superfamily of Trm5p homologs includes proteins that modify substrates other than tRNA, including rRNA and DNA. The finding of other methyltransferases in the Trm5 protein family, although still not identifying Trm10p, underscores their evolutionary unrelatedness. Although the Trm10 family of proteins is smaller, our phylogenetic analysis indicates that there may have been functional diversification in this family of proteins as well; further analysis of the multiple homologs present in metazoans will be required to elucidate any diverse functions.

The cellular role of m¹G₉ modification is unclear at present, despite the evolutionary conservation of this modification and the conservation of the protein (Figs. 7, 8). Cells lacking Trm10p have no obvious growth defects under standard growth conditions in rich or minimal media. For a number of other mutants of tRNA modification enzymes, dramatic phenotypes are only manifested in combination with other mutations that alter tRNA modification or tRNA levels (Grosshans et al. 2001; Johansson and Bystrom 2002). Further tests of this type may need to be done to clarify the role of Trm10 protein. The question of why this modification (and other modifications) is so widely conserved in eukarvotes is an important one because tRNA molecules are among the most abundant nucleic acids in the cell, and therefore a large amount of cellular energy is being spent on production of modifications. From our estimation of the number of mature tRNA molecules produced during a generation of growth in *S. cerevisiae*, methylation at position 9 alone requires the consumption of ~720,000 molecules of SAM per generation.

MATERIALS AND METHODS

Preparation of specifically labeled tRNA substrates

tRNA^{Gly} specifically labeled at the phosphate following guanosine 9 (G₉*Gly) was prepared as follows. In vitro transcribed tRNA^{Gly} (1 nmole) was incubated with Rnase H (a gift from Y.-T. Yu, University of Rochester Medical Center) and a chimeric 2'-Omethylated/deoxy oligonucleotide (HHMI/Keck Center, Yale) to direct cleavage 3' of G₉ (Yu 1999). After gel purification of the resulting 3' half-molecule, 100 pmole was treated with alkaline phosphatase (Roche), and then rephosphorylated with T4 Polynucleotide kinase (PNK; Roche) in the presence of $[\gamma^{-32}P]ATP$ (ICN, 7000 Ci/mmole) to yield a labeled, 5'-phosphorylated 3' half-molecule (beginning at G_{10}) that was gel-purified. The labeled RNA (40-50 pmole) was ligated with a synthetic RNA comprising the first nine nucleotides of tRNA^{Gly} (100-200 pmole; Dharmacon Research), using a DNA oligonucleotide (100-200 pmole) designed to bridge the desired ligation site, 80 U of T4 DNA ligase (USB), and incubation in buffer containing 1× T4 DNA ligase reaction buffer (USB) supplemented with an additional 1 mM ATP at 37°C for 4-12 h. Ligated RNA was purified by PAGE. Typical ligation efficiencies were 50%–70% of labeled 3' molecule ligated.

 G_{37} *Leu substrate was produced in essentially the same way, except that the chimeric oligonucleotide used in the RNase H reaction directed cleavage 3' of G_{37} in in vitro transcribed mature tRNA_{CAA}^{Leu}, and the 5' half-molecule used for the subsequent ligation was PAGE-purified from the same RNase H reaction.

Primer extension assays were performed on the purified 3' fragments of RNA produced by RNase H to demonstrate that cleavage occurred at the correct position for each RNA. Ligated reaction products were analyzed by RNase T2 and P1 nuclease cleavage reactions to determine that the correct junction sequence was obtained after ligation.

Methyltransferase assay conditions

m¹G methyltransferase activity was assayed for both G₉*Gly and G₃₇*Leu tRNA substrates, in 10-µL reaction mixtures containing 50 mM Tris-HCl (pH 8.0), 2.5 mM MgCl₂, 1 mM dithiothreitol (DTT), 50 mM ammonium acetate, 0.05 mM ethylenediaminetetraacetic acid (EDTA), 1 mM spermidine, 0.5 mM S-adenosylmethionine (SAM), 5000-10,000 cpm of the corresponding specifically labeled tRNA, and a source of m¹G methyltransferase activity (crude extract or purified protein). Reactions were incubated at 30°C for 1-8 h as indicated, and then stopped by the addition of 90 μL of 0.5 M Tris-HCl (pH 8.0) containing 20 μg of carrier tRNA, followed by phenol extraction and ethanol precipitation of the RNA. RNA was resuspended in a 4-µL solution containing 20 mM ammonium acetate (pH 5.2), 1 mM EDTA, 10 µg of RNase A, and 1 U of RNase T2, and incubated at 50°C for 1 h, and then digestion products were applied to cellulose TLC plates (EM Science) and resolved in buffer containing saturated ammonium sulfate:isopropanol:water (80:18:2).

GST-ORF fusion proteins

The pools and subpools of purified GST-ORF fusion proteins were prepared as previously described (Martzen et al. 1999).

Strains and plasmids

The homozygous diploid strain deleted for YOL093w (*trm10-* Δ / *trm10-* Δ) and its isogenic parent strain (BY4743) were obtained from Invitrogen. Plasmid pJEJ12-3 (N-terminal His₆-Trm10p) was constructed by PCR amplification of the yeast *TRM10* ORF, followed by insertion of the DNA into a pET-derived vector specifying the N-terminal sequence Met-Ala-His₆ before the first Met codon (gift of E. Grayhack, University of Rochester School of Medicine), and sequencing of the resulting insert.

Growth and preparation of crude yeast extracts

To prepare crude extracts, 250 mL of each strain was grown from single colonies in YPD media to an OD_{600} of 2, and cells were harvested, resuspended in 1 mL of extraction buffer containing 50 mM Tris-HCl (pH 7.5), 1 mM EDTA, 4 mM MgCl₂, 5 mM DTT, 10% glycerol, 1 M NaCl, with 2.5 µg/mL leupeptin and 2.5 µg/mL pepstatin, supplemented with 4 mM AEBSF and sheared with a bead-beater containing zirconium beads. Extracts were clarified by centrifugation; protein concentration was determined by BioRad protein assay, and samples were quick-frozen and stored at $-80^{\circ}C$.

Purification of Trm10p

Plasmid pJEJ12-3 (N-terminal His₆-Trm10p) was freshly transformed into *E. coli* BL21(DE3) pLysS cells (Novagen), and an individual transformant was grown at 37°C in 500 mL of LB containing 100 µg/mL ampicillin to OD₆₀₀ = 0.4, and induced for 5 h by the addition of 1 mM isopropyl β-D-thiogalactopyranoside (IPTG). Purification of His₆-Trm10p was performed essentially as previously described (Steiger et al. 2001), using TALON resin (Clontech) for immobilized metal-ion affinity chromatography (IMAC), except that the protein bound to the TALON resin was eluted using a gradient of 0–100 mM imidazole. Trm10p eluted as a single peak at ~80–90 mM imidazole and was dialyzed into buffer containing 20 mM Tris-HCl (pH 7.5), 4 mM MgCl₂, 2 mM EDTA, 55 mM NaCl, 1 mM DTT, and 50% glycerol for storage at ~20°C. The purified protein (6.9 mg/mL) was estimated to be >95% pure on SDS-PAGE by Coomassie staining.

Preparation of small-molecular-weight RNA from yeast

To prepare RNA, the wild-type parent strain BY4743 (*TRM10⁺*/ *TRM10⁺*) and the *trm10-* Δ /*trm10-* Δ strain were grown in 125 mL of YPD to an OD₆₀₀ of 2–3. Cells were harvested, and RNA was extracted with hot phenol, precipitated twice with ethanol, and resuspended in 1 mL of ddH₂O essentially as described previously (Rubin 1975) and stored at –80°C. The concentration of the final purified RNA was calculated by assuming A₂₆₀ of 40 µg/mL RNA = 1.

Purification of tRNA^{Gly} from yeast

tRNA^{Gly} was purified from total yeast cellular RNA using a 5'biotinylated oligonucleotide (Integrated DNA Technologies) complementary to the 3'-end of tRNA^{Gly}. Binding of the biotinylated oligonucleotide to streptavidin (SA) magnetic particles (Roche) was performed according to the manufacturer's instructions. The tRNA^{Gly} was purified from 1 mg of total cellular RNA using 2 mg of bound beads according to the method of Tsurui et al. (1994). The eluted tRNA was centrifuged at 13,000 rpm for 10 min to remove any residual beads and desalted/concentrated by centrifugation using a Centricon YM-10 (Amicon) with successive concentration and dilution into ddH₂O. The concentration of purified tRNA was determined using A₂₆₀; ~10–15 µg of purified tRNA was recovered per milligram of total cellular RNA. The same procedure was used to individually purify tRNA_{CAA}^{Leu} from wildtype and *trm10-*Δ-derived RNAs using a biotinylated oligonucleotide complementary to the 3'-end of this tRNA.

HPLC analysis of nucleosides

Purified tRNA^{Gly} isolated from yeast (1 µg) or in vitro transcribed tRNA^{Gly} (20 µg) incubated with Trm10p or buffer was treated with 4 µg of P1 nuclease in buffer containing 30 mM sodium acetate (pH 5.2) and 0.2 mM ZnCl₂ at 37°C for 16 h, and then with 8 U of calf intestinal alkaline phosphatase (Roche) in 1× alkaline phosphatase reaction buffer at 37°C for 3 h. The resulting nucleosides were resolved by HPLC (Waters Alliance Model 2690, equipped with Waters 996 photodiode array detector) on a reverse phase C18 column (supelcosil LC-18-T, 25 cm × 4.6 µm; Supelco, Inc.) essentially as described (Gehrke and Kuo 1989), and individual spectra of the nucleosides were used to confirm the assignments. N-1 methylguanosine (m¹G) was obtained from R.I. Chemical and analyzed in the same way.

Quantification of modified nucleosides was carried out by measuring the area under each nucleoside peak at its known maximum absorbance. The areas of the four main nucleosides (C, U, G, and A) were used to determine the total number of moles of tRNA in each injection by dividing the area observed by the known extinction coefficient for each nucleoside and then normalizing for the expected number of that nucleoside in the tRNA being analyzed. The moles of modified nucleotide in each injection were then determined by dividing the area by the known extinction coefficient for each modified nucleoside; this number was compared with the total moles of tRNA in the sample to obtain the ratio of modified nucleoside per tRNA.

Primer extension assays

Primer extension experiments were carried out using 15–20 nt primers (Integrated DNA Technologies) that were 5'-end-labeled with T4 PNK and $[\gamma-^{32}P]$ ATP (ICN, 7000 Ci/mmole), followed by centrifugation through a MicroBioSpin 6 column (BioRad) to remove unincorporated ATP. In a 5-µL reaction containing 50 mM Tris-HCl (pH 8.3), 30 mM NaCl, and 10 mM DTT, 1 pmole of each primer was annealed with 2 µg of total RNA (isolated as described above) by heating to 95°C for 3 min followed by slow cooling to 37°C. Then 2 µL of annealed RNA/primer was added to a 5-µL reaction containing 0.1–0.4 mM each dNTP (G, A,T, and C) and 0.7 U of AMV-Reverse transcriptase (Promega, high concentration) in 1× AMV-RT reaction buffer (Promega). Sequencing reactions additionally contained 0.2 mM of each dNTP (G, A, T, or C) as indicated. After 5 min at room temperature, extensions

were allowed to continue at 37°C for 30 min–1 h and then stopped by the addition of 7 μ L of formamide containing 0.2 mg/mL carrier RNA, 0.1% bromophenol blue, and 0.1% xylene cyanol. Reactions were run on 10% polyacrylamide, 4 M urea gels and visualized using the PhosphorImager (Molecular Dynamics).

Bioinformatics and phylogenetic analysis

Using the yeast Trm10p as the query sequence, PSI-BLAST searches were carried out (Altschul et al. 1997). These led to the identification of archaeal homologs in the second iteration, which were also used as queries in subsequent PSI-BLAST analyses to obtain a comprehensive list of Trm10p homologs. To ascertain if there was any homology between the m¹G₉ and m¹G₃₇ methyltransferases, we used the Blocks database (http://blocks.fhcrc.org; Henikoff et al. 2000); preconstructed blocks corresponding to the m¹G₃₇ methyltransferases (IPB002905) were used as queries to search both the list of individual Trm10p homologs using PSI-BLAST, or consensus blocks constructed from the Trm10p homologs, using a LAMA search (Pietrokovski 1996). In neither case was any significant homology found. The Trm10p homologs were aligned using CLUSTALX (Thompson et al. 1997); the alignment was used to generate a phylogeny using the neighbor-joining program (Saitou and Nei 1987), and bootstrap analysis was carried out in the PAUP* suite (Swofford 2002).

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