

Genetics and Molecular Biology | Full-Length Text

Alternate routes to mnm⁵ s ²U synthesis in Gram-positive bacteria

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ABSTRACT The wobble bases of tRNAs that decode split codons are often heavily modified. In bacteria, tRNA^{Glu, Gln, Asp} contains a variety of xnm⁵s²U derivatives. The synthesis pathway for these modifications is complex and fully elucidated only in a handful of organisms, including the Gram-negative *Escherichia coli* K12 model. Despite the ubiquitous presence of mnm⁵s²U modification, genomic analysis shows the absence of *mnmC* orthologous genes, suggesting the occurrence of alternate biosynthetic schemes for the conversion of cmnm⁵s²U to mnm⁵s²U. Using a combination of comparative genomics and genetic studies, a member of the YtqA subgroup of the radical Sam superfamily was found to be involved in the synthesis of mnm⁵ s ²U in both *Bacillus subtilis* and *Streptococcus mutans*. This protein, renamed MnmL, is encoded in an operon with the recently discovered MnmM methylase involved in the methylation of the pathway intermediate nm⁵s²U into mnm⁵s²U in *B. subtilis*. Analysis of tRNA modifications of both *S. mutans* and *Streptococcus pneumoniae* shows that growth conditions and genetic backgrounds influence the ratios of pathway intermediates owing to regulatory loops that are not yet understood. The MnmLM pathway is widespread along the bacterial tree, with some phyla, such as Bacilli, relying exclusively on these two enzymes. Although mechanistic details of these newly discovered components are not fully resolved, the occurrence of fusion proteins, alternate arrangements of biosynthetic components, and loss of biosynthetic branches provide examples of biosynthetic diversity to retain a conserved tRNA modification in Nature.

IMPORTANCE The xnm⁵s²U modifications found in several tRNAs at the wobble base position are widespread in bacteria where they have an important role in decoding efficiency and accuracy. This work identifies a novel enzyme (MnmL) that is a member of a subgroup of the very versatile radical SAM superfamily and is involved in the synthesis of mnm⁵s²U in several Gram-positive bacteria, including human pathogens. This is another novel example of a non-orthologous displacement in the field of tRNA modification synthesis, showing how different solutions evolve to retain U34 tRNA modifications.

KEYWORDS rSAM, sequence similarity network, comparative genomics, mnm⁵s²U, tRNA modification, YtqA, YtqB, MnmM, MnmL

P ost-transcriptional modifications of the first position of the anticodon of tRNA molecules (position 34, also called "wobble" position, Fig. 1A) are critical for accurate decoding of mRNAs [\(1\)](#page-19-0). These modifications are often complex and can expand or decrease the wobble capacity that allows one given tRNA molecule to decode codons differing just by their third base [\(2\)](#page-19-0). For example, bacterial tRNAs that decode NNA/NNG codons, particularly in split codon boxes, generally harbor xnm⁵(s^2)U derivatives where nm⁵ stands for aminomethylene [\(3\)](#page-19-0). The presence of the xnm⁵ side chain increases pairing to G-ending codons, while the s^2 group may be the major contributor to restricting pairing with U- and C-ending codons. In addition, some tRNAs carry ribose

Editor Tina M. Henkin, Ohio State University, Columbus, Ohio, USA

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The authors declare no conflict of interest.

[See the funding table on p. 19.](#page-18-0)

Received 27 December 2023 **Accepted** 9 March 2024 **Published** 29 March 2024

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FIG 1 Validated and predicted xnm⁵(s²)U34 synthesis pathways in the two main model bacteria. (A) Wobble position of tRNAs. (B) Validated xnm⁵(s²)U34 synthesis enzymes for *E. coli* K12. (C) Validated xnm⁵(s²)U34 synthesis enzymes and pathway holes for *B. subtilis 168, S. mutans,* and *S. pneumoniae*. Abbreviations: 5-carboxymethylaminomethyl-2-thiouridine (cmnm 5 s²U); 5-carboxymethylaminomethyl-2′-O-methyluridine (cmnm 5 Um); 5-aminomethyl-2-thiouridine (nm 5 s²U); 5-methylaminomethyluridine (mnm⁵U); 5-methylaminomethyl-2-thiouridine(mnm⁵s²U).

methylation at the 2'-OH and/or the mnm⁵ base modification, which may induce restrictive wobbling [\(4\)](#page-19-0). The enzymes involved in introducing $x n m^{5}(s^{2})U$ derivatives are well characterized in the model Gram-negative *E. coli* K12 (Fig. 1B) (see reference [\(5\)](#page-19-0) for review). The insertions of the s^2 and xnm⁵ moieties involve two separate biosynthetic branches, both requiring multiple enzymes [\(4, 5\)](#page-19-0). Structural analysis of the *E. coli* MnmA-tRNA complex suggests that a partially modified xnm⁵ intermediate species may not fit into the active site, meaning that thiolation of C2 precedes modification of C5 at U34 of tRNA [\(6\)](#page-19-0).

The first step in the synthesis of the xnm⁵ moiety is catalyzed by a heterodimeric 5-carboxymethylaminomethyluridine-tRNA synthase complex MnmGE that can use glycine or ammonium as substrates to produce the cmnm⁵U or nm⁵U derivatives, respectively [\(7, 8\)](#page-19-0) (Fig. 1B). The cmnm⁵U derivative can be subsequently converted into mnm⁵U by the bifunctional enzyme MnmC which contains two independent catalytic domains. The C-terminal domain of MnmC, also known as MnmC/MnmC1 or MnmC(o), promotes the FAD-dependent oxidoreduction of cmnm⁵s²U34 to form nm⁵U. Whereas the N-terminal MnmC domain, MnmD/MnmC2 or MnmC(m), harbors the tRNA 5-(aminomethyl)−2-thiouridylate-methyltransferase (EC:2.1.1.61 domain), which catalyzes the methylation of nm⁵s²U to mnm⁵s²U using S-Adenosyl-L-Methionine (SAM) as the methyl donor [\(9\)](#page-19-0) (Fig. 1B). Some organisms such as *Aquifex aeolicus* only encode a stand-alone MnmC2 protein and lack the MnmC1 ortholog as their MnmGE enzymes only use ammonium to directly produce nm⁵(s²)U [\(10\)](#page-19-0). Other bacteria, such as Myco*plasma capricolum,* only encode the MnmEG genes and no MnmC1 or MnmC2 genes and harbor cmnm⁵(s²)U in their tRNAs [\(11\)](#page-19-0).

Environmental and nutritional conditions affect the relative levels of tRNA modifi-cations, including xnm⁵(s²)U derivatives [\(12\)](#page-19-0), leading to varying translation efficiency and cellular viability. In general, media, growth phase, and tRNA identity influence the proportions of the different routes, leading to different xnm⁵(s^2)U derivatives [\(13\)](#page-19-0). It is also known that additional enzymes are involved in ribose methylation (TrmL)[\(14\)](#page-19-0) or in producing tRNA^{Lys}UUU and tRNA^{Glu} UUC with U34 modified with selenium instead of sulfur (MnmH) [see reference [\(4\)](#page-19-0) for review]. The MnmH enzyme is also involved in the geranylation of the same tRNAs in *E. coli* and *Salmonella typhimurium* [\(4\)](#page-19-0). The presence of xnm⁵(s²)U modifications is critical for fitness as it has been shown in *E. coli* that mutations in *mnmE* and *mnmA* are synthetic lethal [\(15, 16\)](#page-19-0). Mutations in *mnmE*, *mnmG,* and *mnmA* confer a wide range of phenotypes, including defects in virulence in diverse bacteria, including species from Gram-positive and Gram-negative groups [\(3, 17\)](#page-19-0). Their orthologs are also important for modification of mitochondrial tRNAs and their absence leads to mitochondrial dysfunction and disease [\(3, 18\)](#page-19-0).

E. coli K12 is one of the rare organisms where all the tRNA modification genes have been identified [\(19\)](#page-19-0). Even in the well-studied Gram-positive model organism *Bacillus* subtilis sp subtilis strain 168, the functional assignment of genes involved in mnm⁵s²U synthesis remains incomplete [\(20\)](#page-19-0). While the accumulation of cmnm⁵s²U was anticipated, given the presence of phylogenetically conserved *mnmEG* orthologs in this organism, the presence of mnm⁵ s ²U was initially unexpected since *B. subtilis* lacks *mnmC*-like gene or sequences coding for MnmC1 and MnmC2 domains. This intriguing finding suggested that a non-orthologous replacement must have occurred for the last steps of the pathway (Fig. 1C) [\(21\)](#page-19-0). The existence of alternate biosynthetic schemes for tRNA thiolation in this organism is further supported through the occurrence of an abbreviated pathway for s^2U tRNA modification involving only two enzymes YrvO-MnmA, bypassing the Tus sulfur relay system required in *E. coli* (Fig. 1C) [\(6\)](#page-19-0).

Foundational studies from the Armengod group showed that cell extracts of *B. subtilis* were competent to promote mnm⁵s²U synthesis from precursor cmnm⁵s²U tRNA isolated from *E. coli ΔmnmC* strain) [\(21\)](#page-19-0). Interestingly, these reactions also formed the nm⁵s²U intermediate, which was consumed in reactions supplemented with S-adenosyl-methionine (SAM), but not FAD. More recently, YtqB (MnmM) was identified as the enzyme responsible for the conversion of nm^5s^2U into mnm⁵s²U, performing an MnmC2-like reaction [\(22\)](#page-19-0). This assignment was attributed to the depletion of mnm⁵s²U accompanied by the accumulation of nm⁵s²U tRNA in the *B. subtilis ΔmnmM* strain. Trans-complementation of not only *B. subtilis mnmM* but also *Staphylococcus aureus* and *Arabidopsis* thaliana mnmM orthologs restored mnm⁵s²U levels [\(22\)](#page-19-0). Furthermore, according to Cho and collaborators, *B. subtilis mnmM* expression in *E. coli ΔmnmC* shows detectable levels of mnm⁵s²U but at levels significantly lower than those of the wild type. While MnmM is competent in catalyzing methylation of nm⁵s²U to yield mnm⁵s²U, it remains unanswered if additional components participate in this biosynthetic scheme in *B. subtilis* and other species lacking MnmC. For instance, although experimentally demonstrated in crude extracts, the identity of an MnmC1-like enzyme catalyzing the conversion of cmnm⁵s²U into nm⁵s²U tRNA is unknown.

Comparative genomic methods combining phylogenetic distribution, gene fusion, and physical clustering analyses have been powerful tools in identifying many missing tRNA modification genes, particularly for complex modification pathways that require many enzymes [\(23–27\)](#page-19-0). Now that many tRNA modification genes have been identified in different organisms, it is clear that different evolutionary solutions have emerged to catalyze the insertion of the same modifications [\(28\)](#page-19-0). In this work, we used comparative genomic methods combined with genetic studies to better understand the last steps in mnm⁵s²U synthesis in microbes lacking MnmC and identified the versatile radical SAM superfamily family protein YtqA as a key player. These combined approaches reveal the occurrence of alternate pathways leading to hypermodification of U34 tRNA^{Glu, Gln, Lys} in bacterial species.

RESULTS

Phylogenetic distribution analysis of the mnm⁵ s ²U pathway suggests MnmC1 has been replaced by non-orthologous enzymes in a number of firmicutes

Analysis of the distribution of the known mnm⁵s²U34 synthesis protein family members in the bacterial genomes covered in the COG database revealed a patchy distribution for the latter part of the pathway (Fig. 2; Fig. S1; Data S1 and S2). Of the 968 organisms that encode the three proteins required for cmnm⁵s²U biosynthesis—MnmA (COG0482/ K00566), MnmE (COG0486/K03650), and MnmG (COG0445/K03495)—22% (216/968) encode bifunctional MnmC1/MnmC2 (or MnmC, COG4121/K15461) proteins like *E. coli,* while ~24% (232/968) encode only MnmC2 proteins without MnmC1 domains like *A. aeolicus*. Finally, over half (520/968) of the analyzed organisms lack both MnmC1 and MnmC2 homologs. Thus, the occurrence of MnmC1 is sparse and matches the distribution of MnmC1 and MnmC2 sequences recently reported by Cho et al. [\(22\)](#page-19-0) for a smaller data set. The absence of MmnC1 can be, however, associated with distinct biochemical scenarios. First, in the subset of organisms that only use the ammonia pathway to directly synthesize nm⁵s²U, no MnmC1 is required, and the subsequent methylation to mnm⁵s²U can be performed by MnmC2 family proteins, like in *A. aeolicus* [\(10\)](#page-19-0) or possibly by

FIG 2 Phylogenetic distribution analysis of MnmC1, MnmC2, and YtqA/YhcC proteins. Percentages for each group represent the number of organisms that encode a given protein group [MnmC1/MnmC2 fusions (MmmC1C2), MnmC1 alone (MnmC1), MnmC2 alone (MnmC2), and YtqA/YhcC] out of the 968 bacterial genomes that encode MmnAGE analyzed (Table S1; Data S2, S3, and S4). In addition, the percentage of genomes where the *ytqA* and *mnmM* genes are physically clustered on the chromosome (YtqA-MnmM clustered) is reported (Data S5 and S6). The pixel width of branches reflects the relative intensity of corresponding heatmaps and their dominant (highest value) color of those same heatmaps (0.0–1.0 heatmap range corresponds to 1–3 px branch width). Heatmap per branch with the highest intensity across all five heatmap rows was assigned to the branch color to highlight taxonomic patterns. Only Deferribacteres possess a combined color reflecting the paired 1.0 values for both MnmC2 and YtqA/YhcC heatmaps (orange and green of equal intensity combine to make brown). The Firmicute branch is colored red.

YtqB/MnmM proteins [\(22\)](#page-19-0). Second, some organisms such as *Lactococcus lactis* sub lactis Il1403 [\(29\)](#page-19-0) or *M. capricolum* [\(11\)](#page-19-0) harbor only cmnm⁵s²U and, therefore, do not require any enzymes beyond MnmEG. Third, in species that accumulate both cmnm⁵s²U and mnm⁵ s ²U, as in the case of *B. subtilis* and possibly in many other Firmicutes [\(30, 31\)](#page-20-0) (Fig. 2; Fig. S1), a non-orthologous replacement of MnmC1 must be present and is yet to be identified.

Comparative genomic analyses led to the identification of a radical SAM family subgroup as a candidate for the missing mnm⁵ s ²U synthesis enzyme in Firmicutes

Protein fusions and physical clustering analysis are powerful methods to identify missing genes [\(32, 33\)](#page-20-0) and both led to the identification of a subgroup of the YtqA/YhcC family as a strong candidate for an enzyme involved in the synthesis of $nm⁵s²U$ in Firmicutes. A domain analysis of the COG4121 family, which encompasses both MnmC1 and MnmC2 domains, across the 968 bacterial genomes of the COG Database [\(34\)](#page-20-0) that also encode the three MnmA, MnmG, and MnmE proteins, led to the identification of at least four other types of MnmC2 fusions, in addition to the well-characterized MnmC1/MnmC2 examples: fusions with the queuine tRNA-ribosyltransferase (TGT) (COG0343, PF01702); the DNA modification methylase YhdJ (PF01555, COG0863); the menaquinone biosynthesis MqnA superfamily subgroup MqnD (PF02621, cd13634); and with a radical SAM family domain of the YtqA/YhcC subgroup (COG1242, PF16199, cd01335, PF04055) (Data S3; Fig. S1). The YtqA name comes from the subfamily member found in *B. subtilis* (BSU30480) and the YhcC name comes from the subfamily member found in *E. coli* (b3211). Three examples of YtqA/MnmC2 fusion proteins were identified in the genomes present in the COG database and 70 more in the InterPro database (Fig. 3A; Fig. S1; Data S4A). Genomic synteny analyses show that *ytqA* is often in an operon or in a close physical neighborhood with the mnmM gene encoding the recently identified nm⁵s²U methyltransferase, including in the model Gram-positive *B. subtilis* where the two genes are co-expressed (Fig. 3B and 4B; Fig. S2) [\(35\)](#page-20-0). Both YtqA/YhcC and MnmM are part of superfamilies that are currently not well separated in most protein family databases (see below and discussion), but when analyzing the distribution of *ytqA-mnmM* clusters (Fig. 2; Data S5), these are prevalent in organisms that are missing MnmC1 and MnmC2 such as the Firmicutes and the deltaproteobacteria (Fig. 2; Data S6 and S7).

YhcC/YtqA sequences display signature motifs of members of the radical SAM superfamily of enzymes, known to facilitate a wide range of chemistries. First, these sequences were located in the sequence similarity network (SSN) analysis of the whole radical SAM family generated by the Gerlt group using the searchable web resource RadicalSAM.org [\(36\)](#page-20-0). The YtqA/YhcC subgroup was identified as part of the Elongator Protein-like mega cluster 2–4.1 (Fig. 4A). Proteins of this subgroup are members of the KEGG orthology group K07139 and the InterPro family IPR005911 (Protein YhcC-like). YtqA unique functional motifs and genomic synteny of its coding sequence make it a candidate for replacing MnmC1 in many organisms. Proteins within the YhcC sub-group would not be isofunctional since the *E. coli* genome encodes for YhcC, and MnmC1 is the only protein catalyzing the cmnm⁵s²U to nm⁵s²U reaction in this organism [\(9\)](#page-19-0). We, therefore, constructed SSNs of 10,896 IPR005911 proteins, which allowed differentiation of the *E. coli* YhcC subgroup (group four in Fig. 4B) from the *B. subtilis* YtqA subgroup (group one in Fig. 4B). Mapping of additional data such as gene neighborhood, fusion, and gene presence/absence data onto this network suggests that in addition to YtqA proteins from subgroup 1, homologs from subgroups 3, 5, 7, and 8 are most likely linked to mnm⁵s²U synthesis. Collectively, bioinformatic analysis showing the physical clustering, gene fusion, and taxonomic distribution data strongly suggests a role of the YtqA protein subfamily in Firmicutes in mnm⁵s²U synthesis in tRNA.

FIG 3 Physical clustering and domain fusion link the YtqA subgroup with mnm⁵s²U synthesis. (A) Examples of fusions between proteins involved in the last steps of mnm^ss²U synthesis. Protein domains are color-coded MnmC1 (dark green), MnmC2 (light green), MnmM (light gray), and YtqA (dark gray). The number of examples of such fusions present in the InterPro database is given between brackets; see Data S4A for the complete list of YtqA/MnmM, MnmC1/MnmM, and YtqA/MnmC2 fusions. (B) Examples of physically clustered *ytqA* (dark gray) and *mnmM* (light gray) in Gram-positive species. Data extracted from the SSN analysis performed for Fig. 4.

Genetic analysis in *S. mutans* **confirms the involvement of YtqA in mnm⁵ s ²U synthesis**

To validate the predicted role of YtqA in mm^5s^2 U synthesis, we constructed deletion mutants of the *ytqA and mnmM genes* in *Streptococcus mutans*. Like in *B. subtilis*, the two genes are in an operon in *S. mutans* (Fig. 3). A non-polar deletion of the *ytqA* (*smu_1699* c) was constructed by inserting an Erythromycin Resistance (Ery^R) cassette lacking a terminator. The same cassette was used to delete the *mnmM* (*ytqB*, *smu_1697* c) gene as well as to delete both genes. As expected, tRNA extracted from a *Streptococcus mutans* strain containing a deletion of the *ytqA-mnmM* genes (*smu_1699 c and smu_1697 c*) lacks mnm⁵s²U and accumulates increased relative levels of nm⁵s²U and cmnm⁵s²U intermediates (Fig. 5). Inactivation of only *mnmM* confirms the involvement of the methyltransferase in mnm⁵s²U formation and also showed an accumulation of nm⁵s²U, compatible with its proposed role in catalyzing the final methylation step in mnm⁵ s ²U. Inactivation of *ytqA* led to higher levels of cmnm⁵s²U and no nm⁵U accumulation, a phenotype that supports the involvement of *ytqA* in this pathway. Notably, the distinct phenotypes of *ytqA-mnmM* and *ytqA* deletions demonstrate the absence of polar effects on the *ytqA* knockout strain.

Genetic analysis of *B. subtilis* **YtqA shows its involvement in mnm⁵ s 2U pathway**

The genomic co-localization of *ytqA* and *mnmM* observed in *S. mutans* is also observed in *B. subtilis* [\(20, 22\)](#page-19-0). Recent functional studies in *B. subtilis* showed that the inactivation of mnmM led to a loss of mnm⁵s²U [\(22\)](#page-19-0). Two research groups participating in this study independently analyzed the relative levels of tRNA modifications in *B. subtilis ytqA* and

FIG 4 Sequence similarity network of the YtqA/YhcC family. (A) The radical SAM domains that are fused to MnmC2 are members of the Megacluster-2-4-1 (in black), on RadicalSAM.org a subgroup of Megacluster-2–4 ((Elongator protein-like); (B) YtqA/YhcC SSN overview generated using InterPro HMM signature identifier IPR005911 (EFI-EST; AS100, repnode 55). Different types of data were mapped to the subset of nodes (black outlines) present in the benchmark genomes (968 representative genomes that encode MnmAGE proteins): (1) Presence/absence data of MnmC1, MnmC2, and MnmC1/C2 fusions (Data S3); (2) YtqA/MnmC2 fusions data (Data S3); (3) physical clustering data (Data S5). Dark blue nodes were found to be without Gene Cluster data in KEGG. For complete mapping, see Data S8.

mnmM mutants and have not been able to reproduce these results. Collectively, our analyses showed the absence of mnm⁵s²U modification in the DytqA strain and the presence of this modification in the *ΔmnmM* strain from LB cultures isolated at mid and late lag phase (OD 0.5–1) (Fig. 6; Fig. S3). The results reported here are in contrast with the recent Cho et al. report on the analysis of *mnmM* knockout strain using tRNA samples isolated from overnight cultures that showed no detectable levels of mnm⁵s²U [\(22\)](#page-19-0) but in agreement with earlier analysis of *B. subtilis* IC6779 strain containing *mnmM* inactivation that retained detectable levels of the fully modified base [\(21\)](#page-19-0). In our hands, tRNA samples extracted from *B*. *subtilis* cultured beyond log phase (OD = or > 2) led to inconsistent results (Fig. S3B), likely associated with cells entering sporulation. Remarkedly, tRNA isolated from *B. subtilis ΔytqA* showed high levels of the pathway intermediate nm⁵s²U that is not detectable in tRNA extracted from the wild-type strain. Therefore, ytqA inactivation resulted in the accumulation of nm^5s^2U and no detection of mnm $5s^2U$ (20, [22\)](#page-19-0). Overall, these analyses showed the involvement of ytqA and mnmM in mnm⁵s²U synthesis. It is worth noting that cells lacking YtqA or MnmM do not display any growth phenotype when compared to the wild type, suggesting that the intermediate c_{mm} ⁵s²U modification may already provide the proper conformation of s²U34 to guarantee fidelity in base pairing with cognate tRNAs.

The ability of *B. subtilis* YtqA and MnmM to functionally replace *E. coli* MnmC was then assessed through heterologous trans-complementation experiments as a probe for

FIG 5 *S. mutans* YtqA and MnmM are involved in mnm5s2U tRNA modification. *S. mutans* wild type, and *ytqA*, *mnmM*, and *ytqA-mnmM* deletion strains were cultured in a BHI medium. tRNA samples were processed through the VDCL2 method and analyzed through LC-MS method 2, as described in the Materials and Methods section. For each sample, 600 ng of digested tRNA was analyzed through LC-MS, and two technical replicates were analyzed. The reported averages and standard deviation were calculated based on data acquired from three independent growth experiments. The signal of each analyte was confirmed by both qualifier and quantifier transitions and reported as the fragment mass of each modification, as indicated. Relative levels of each modification were determined by normalizing the area of mass abundance associated with each modification to the mass abundance of dihydrouridine in the same sample. The star mark denotes the absence of modification or levels below the detection limit.

MnmC1 and MnmC2 equivalent reactions. As previously reported, *E. coli DmnmC* accumulates cmn⁵s²U and completely lacks mnm⁵s²U [\(21\)](#page-19-0). Co-expression of *B. subtilis* YtqA-MnmM restored the synthesis of mnm⁵s²U to 1/3 of wild-type levels, confirming their involvement in this pathway (Fig. 6B). Expression of *B. subtilis* MnmM alone also resulted in mnm⁵ s ²U formation, albeit at much lower levels (~5% of *E. coli* MG1655 levels) and similar to residual levels reported by Cho et al. [\(22\)](#page-19-0).

Altered pathway for the mnm⁵ s ²U synthesis in *S. pneumoniae*

Genomic analysis of YtqA and MnmM orthologs showed the presence of an altered pathway for mnm⁵ s ²U in some Gram-positive such as *Streptococcus pneumoniae* D39. This organism harbors truncated copies of *ytqA* coding the N-terminal and C-terminal sequences of YtqA (*spd_0565* and *spd_0566*) upstream of *mnmM (spd_0567*) (Fig. 3; Fig. S4), raising the question of whether these species represent evolutionary representatives in the process of losing the last step of this pathway. In fact, analysis of modified nucleosides from S. pneumoniae showed much-reduced levels of mnm⁵s²U (approximately 10-fold lower than *B. subtilis* or *E. coli*, Fig. 6) and confirmed the involvement of *spd_565* to *spd_567* gene region in this pathway (Fig. 7). Interestingly, a spontaneous nonsense mutation within *mnmM* generating a Q152stop was identified within the unencapsulated *S. pneumoniae* D39 genetic background (IU1824) (Fig. S4). Analysis of modified nucleosides showed that this truncation near the C-terminal end of the protein affects MnmM activity and results in loss of mnm⁵s²U synthesis like the deletion of the whole *mnmM* gene (IU19966) (Fig. 7). These phenotypes were complemented by repairing the mutation (strain IU19835) or expressing the *mnmM* ectopically under a zinc inducible promoter. The lack of an apparent growth defect upon elimination of mnm⁵s²U was also observed in S. pneumoniae, reinforcing the proposal that the cmnm⁵s²U modification is sufficient to fulfill functional requirements. Nevertheless, out of 242 complete *S. pneumoniae* genomes in the BV-BRC database at the time of the analysis (version 3.32.13a), all harbored truncated copies of *ytqA (e*xamples given in Fig. S5), suggesting this truncation must have occurred just before the *S. pneumoniae* speciation. Phylogenetic analysis indicates that the closest species with a common ancestor, *Streptococcus mitis* [\(37\)](#page-20-0) has a full copy of the gene (Fig. S5). To investigate the possible impacts of the truncated *spd_0567*(*mnmM)(*Q152stop) on pneumococcal physiology, we performed the following experiments and detected no differences between IU1824

FIG 6 B. subtilis mnm⁵s²U pathway involves YtqA and MnmM. (A) The inactivation of *ytqA* and *mnnM* genes affects the relative accumulation of mnm⁵s²U and pathway intermediates. Modified nucleosides were extracted from *B. subtilis* wild type, *DytqA,* and *DmnmM* strains were cultured in LB medium until OD of 1. (B) *B. subtilis* YtqA-MnmM complements defects associated with *E. coli* D*mnmC* inactivation. *E. coli* wild-type MG1655 *and* D*mnmC* strain transformed cells with empty plasmid (pBAD), pDS358 (*pBAD-ytqA-mnmM*), pDS359 (*pBAD-ytqA*), or pDS365 (*pBAD-mnmM*) were cultured in LB ampicillin and arabinose until OD of 1. tRNA samples were isolated using the PDS method and analyzed using LC-MS method 1. Relative levels of each modification were determined by normalizing the mass abundance associated with each modification to the mass abundance of dihydrouridine in the same sample. The reported averages were calculated based on data acquired from three independent growth experiments. The star mark denotes the absence of modification or levels below the detection limit.

(*spd_0567*(*mnmM*)(Q152stop) and IU19835 (*mnmM*⁺) (data not shown). The conditions tested were growth in BHI broth at 30°C, 37°C, 42°C, and in BHI broth containing 0.3 M NaCl at 37°C. There were no differences between cell widths and lengths of cells grown at 37°C, or survival rates after 30-min treatment with 20 mM hydrogen peroxide. In addition, antibiotic disk-diffusion assays showed no differences in the diameter of the zones of inhibition using discs containing fosfomycin, cefotaxime, penicillin, and tetracycline between IU1824 (*spd_0567* (*mnmM*) (Q152stop) and IU19835 (*mnmM*⁺).

DISCUSSION

Bioinformatic analysis shows the occurrence of diverse biochemical strategies for hypermodification of U34 in tRNA^{Glu, Gln, Lys} in bacteria. Our results from the *S. mutans* and *S. pneumoniae ΔmnmM* strains confirm the functional assignment of Cho et al. [\(22\)](#page-19-0) that MnmM is required in the SAM-dependent methylation reaction nm^5s^2U to form mnm⁵ s ²U. Our analysis from *B. subtilis ΔmnmM,* however, did not show the same requirement, suggesting that, in this organism, other methylases may provide functional replacement under certain growth conditions. Previous analysis of single knockout strains of several methylases in *B. subtilis*, including MnmM also failed to show inactiva-tion of mnm⁵s²U synthesis [\(21\)](#page-19-0). We believe that the differences in the observed phenotypes of the *mnmM* knockout strain between ours and other studies may be associated with the functionality of the NH₄⁺ and glycine-dependent routes with varying growth conditions and stages. The NH₄⁺ route could be the major route to mnm⁵s²U in the stationary phase (conditions used in the Cho et al. analyses). Hence, MnmM would be the

FIG 7 S. pneumoniae (Sp) MnmM is involved in mnm⁵s²U tRNA modification. tRNA isolated from Sp cultures using MW methods were digested and analyzed through LC-MS Method 1. Modified nucleosides were purified tRNA from Sp strains: (1) Sp IU19835: D39 Δcps WT (*spd_0565⁺ , spd_0566⁺ , spd_0567⁺*) (black), (2) Sp IU19964: Δspd_0565-0566 markerless (*ΔytqA*) (dark gray), (3) Sp IU19966: *Δspd_0567* markerless (*ΔmnmM*) (light gray), (4) Sp IU19968: Δ*spd_0565-0567* markerless (*ΔytqAmnmM*)(*white*), (5) Sp IU19974: Δ*spd_0567* markerless (*ΔmnmM*) ΔbgaA-Pzn-*mnmM* grown in Zn, (complementation of *ΔmnmM*) (gray stripped), and (6) Sp IU1824: original WT strain (IU1824) that has spd_0567 (Q152stop) mutation (black stripped). *spd_0565 and spd_0566* encode the N-terminal and C-terminal of YtqA and *spd_0567* encodes MnmM. Relative levels of each modification were determined by normalizing the mass abundance associated with each modification to the mass abundance of dihydrouridine in the same sample. The reported averages were calculated based on data acquired from three independent growth experiments. The star mark denotes the absence of modification or levels below the detection limit.

major player in these conditions. However, the glycine route may be the major route in exponential growth and another enzyme or pathway would be required to synthesize mnm⁵s²U from cmnm⁵s²U. Physical clustering and gene fusion data point to YtqA as a candidate to complete this pathway in *B. subtilis*. Indeed, the *B. subtilis ΔytqA* mutant is devoid of mnm⁵s²U and accumulates nm⁵s²U, supporting the occurrence of the NH₄+ pathway in this organism. This phenotype cannot be due to a polar effect on *mnmM,* as we found that deleting this gene did not lead to $nm⁵s²U$ accumulation and retained mnm⁵ s ²U modification. In addition, co-expression of *ytqA-mmnM* allowed partial complementation of the *E. coli ΔmnmC* mnm⁵ s ²U deficiency, whereas expression of mnmM alone only led to minimal mnm⁵s²U accumulation. The expression of ytqA alone did not increase nm⁵U nor mnm⁵U levels in the same *mnmC* strain compatible with the model that thiolation of C2 of U34 precedes the installation of the C5 modification route (data not shown).

The exact role of YtqA is still not fully understood and will require further biochemical and genetic characterization. Nevertheless, experimental evidence provided in this study shows the involvement of YtqA in the biosynthesis of $\text{mnm}^5\text{s}^2\text{U}$; therefore, we propose renaming this enzyme as MnmL (Fig. 8). Our current model is that MnmL-MnmM can promote the conversion of cmnm⁵s²U into mnm⁵s²U. MnmL (YtqA) is classified as a member of the radical SAM enzyme superfamily, known to catalyze diverse chemical transformations triggered by the generation of a 5′-deoxyadenosyl radical (dAdo●) from SAM [\(38\)](#page-20-0). This dAdo \bullet could abstract an H-atom from cmnm⁵s²U, causing oxidative decarboxylation by forming an xnm⁵s²U intermediate that could further react to yield mnm⁵s²U. Potential sites for the dAdo^o-catalyzed H-atom abstraction include the glycine alpha carbon, the amide nitrogen, or the $5U$ -methylene carbon (Fig. S6). Experimental determination of the site of H-atom abstraction would require isotope labeling studies. However, the viability of this chemistry can be probed using density functional theory (DFT) geometry optimizations to predict the outcome of H-atom abstraction from each of these sites within the cmnm⁵s²U substrate—these results are summarized in Table S2. For example, H-atom abstraction from the amide nitrogen promotes oxidative decarboxylation and the formation of a terminal imine that could be hydrolyzed to give nm⁵s²U. This chemistry is very similar to that catalyzed by radical SAM enzymes HydG and ThiH, which abstract hydrogen from the amino group of an exogenous L-tyrosine, leading to the formation of *p*-cresol and dehydroglycine [\(39\)](#page-20-0). However, the observation made in the present work that both YtqA and MnmM are required to produce the maximal

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FIG 8 Diversity in the cmnm⁵s'U/mnm⁵s'U biosynthetic routes in bacteria. (A) Pathways utilizing MnmL(YtqA) and MnmM show varying complexity. (B) Bacteria utilize alternate enzymes or domains to complete the synthesis of mnm⁵s²U. The pathways in organisms in bold have been experimentally validated (see main text for references), and those marked with * and + have been predicted based on phylogenetic distribution and gene fusion data, respectively.

amount of mnm⁵s²U suggests some interplay between the two enzymes. Sequence homology (cf. Fig. 4) suggests that MnmL (YtqA) possesses a $(\beta/\alpha)_{6}$ three-quarter TIM barrel fold, like 75% of all radical SAM enzymes [\(40\)](#page-20-0). Often, an additional protein subunit occupies the gap in the TIM barrel and is integral in the catalyzed chemistry, as is the case in the tRNA-modifying MiaB [\(41\)](#page-20-0). MnmL is a quite small radical SAM enzyme (sequence length ≈330 amino acids), and no additional subunit is predicted by AlphaFold {https://alphafold.ebi.ac.uk/entry/O35008}. Therefore, it is possible that tRNA binding may mediate the formation of a ternary complex with MnmM, which enables the conversion of cmnm⁵s²U to mnm⁵s²U. This model is compatible with the observation that no nm⁵ s ²U is accumulated in *S. mutans*, *B. subtilis*, and *S. pneumoniae* wild-type strains. The requirement of a MnmL-MnmM-tRNA ternary complex for completion of the two consecutive steps also helps to explain the lack of mnm⁵s²U when MnmL (YtqA) is absent and the accumulation of nm^5s^2U which is likely derived from the NH $_4^+$ pathway. In this model, it is possible that another yet-unidentified methylase(s) can provide a functional replacement when MnmM is absent in *B. subtilis*. Nevertheless, biochemical validation of these proposed biosynthetic schemes will be tested in subsequent studies.

The occurrence of alternate pathways for the synthesis of mnm⁵s²U cannot be simply explained by the existence of an optimized single enzyme catalyzing two consecutive reactions as in the case of MnmC or by the recruitment of a proposed MnmL-MnmM complex. In fact, nature has found multiple combinations to promote hypermodification of U34 tRNA involving both C2 and C5 positions, which guarantees fidelity in translation. Distinct pathways have been ascribed for s²U in prokaryotes, as well as distinct cytosolic and mitochondrial eukaryotic systems [\(6,](#page-19-0) [42, 43\)](#page-20-0). As discussed in the introduction, for modifications in the C5 position, some species do not synthesize mnm⁵s²U and only accumulate cmnm⁵s²U with apparent no metabolic penalties. At the same time, other species have bypassed the cmnm⁵s²U intermediate by utilizing only the ammonia route through the direct formation of nm⁵s²U intermediate and an MnmC2 enzyme to

complete the pathway, as in the case of *A. aeolicus* [\(10\)](#page-19-0). Phylogenetic distribution and fusion analyses [\(Fig. 2](#page-3-0) [and 3;](#page-5-0) Fig. S1 and S8 of Cho et al. [\(22\)](#page-19-0)) revealed the patchwork of enzymatic solutions that have evolved in bacteria for the synthesis of the x^5s^2U34 modifications (summarized Fig. 8). Some of these solutions have been experimentally validated (in bold in Fig. 8), while others are predicted from the fusions analysis (annotated with $a + sign$ in Fig. 8) or from phylogenetic distribution data (annotated with a * sign in Fig. 8). For example, *Chlamydiae trachomatis* encodes an MnmM but no MnmL(YtqA) nor MnmC1 (Table S6 of Cho et al. [\(22\)](#page-19-0) and Data S4B). We predict that, like A. aoelicus, this organism only uses the ammonium route to produce nm⁵s²U but then uses MnmM (instead of MnmC2 in *A. aoelicus*) to perform the last step (Fig. 8). Similarly, the MnmL/MnmC2 and MnmC1/MnmM fusions (Data S4A) suggest these two combinations of enzymes can be used to convert cmnm ${}^{5}s^{2}U$ into nm ${}^{5}s^{2}U$ in addition to the known *E. coli* MnmC1/nmC2 combination. These predictions await experimental validation as one cannot rule out that other yet-unidentified enzymes are present in some organisms. Also, in organisms that only harbor MnmAGE, it is impossible to predict without experimental data if the glycine route or the $NH₄$ route is used alone or in combination, even if to date no organisms seem to stop at the $nm⁵s²U$ intermediate. It is also clear that the genes encoding enzymes involved in the late steps of the mnm⁵s²U synthesis pathway can be quite easily lost in specific clades, as shown here in the *S. pneumoniae* clade.

Functional assignment solely based on sequence similarities is not a suitable approach for computational prediction of the mnm⁵s²U pathways. Both YtqA/MnmL and YtqB/MnmM proteins are members of protein superfamilies that have many paralog subgroups and one must be very cautious in transferring functional annotations from the experimentally validated members to homologs to avoid mis-annotation of paralogs. The experimentally validated *B. subtills* MnmM is a member of the SAM-dependent methyltransferase superfamily (IPR029063) that contains hundreds of families and cannot be used to propagate the annotation. A more specific profile has been recently created, which places MnmM in the methyltransferase MnmM-like family (IPR010719), and it has been used to propagate the annotation in Interpro and Uniprot databases and experimentally validates for several members of the family [\(22\)](#page-19-0). For the transfer of the MnmL/YtqA functional annotation, one can confidently propagate within the proteins in Group 1 of the SSN generated in Fig. 4. Proteins of subgroups 3, 7, and 8 also have a high probability of isofunctionality with proteins of subgroups 1 based on fusion and physical clustering (Fig. 4). For all other subgroups, experimental testing is required before any annotation transfer can occur. One subgroup that is most certainly not isofunctional with MnmL/YtqA is group 4, which contains the *E. coli* YhcC protein. As discussed above, there is no cmnm⁵s²U to mnm⁵s²U transformation activity when the *mnmC* gene of *E. coli* that encodes the MnmC1/MnmC2 fusion is deleted. In addition, physical clustering data of the larger YtqA/YhcC family never links members of *E. coli* YhcC subgroups with any protein of the nm⁵s²U pathway (Data S5). The genes that cluster most frequently with genes encoding proteins of the YhcC subgroup (group 4 in Fig. 4) are genes involved in anaerobic/aerobic growth conditions and oxidative stress resistance such as *gltB*, *gtlD*, *arcB*, *ptsN,* and *elbB* (Data S5). It has also been shown that *yhcC* expression is regulated by the anaerobic transition regulator FNR [\(44\)](#page-20-0). At this stage, further studies will be required to elucidate the function of members of the YhcC subgroup.

Overall, this study shows the diversity of pathways involving the synthesis of xnm⁵s²U tRNA in bacteria. Genomic analysis guided the computational assignment of alternate pathways and led to the identification of MnmL/YtqA as a component of this pathway. The occurrence of fusion proteins containing alternate arrangements of MnmC1/MnmL and MnmC2/MnmM supports the proposed role of MnmL in catalyzing the conversion of cmnm⁵s²U into nm⁵ s²U in a reaction that is coupled to the final methylation by MnmM. Experimental validation of such computational assignments was completed for both *S. mutans* and *B. subtilis*. Interestingly, analysis of *S. pneumoniae-modified tRNA* suggests that the arrangement of MnmL as two polypeptides spanning the N- and C-

halves of the enzyme leads to loss of function. Thus, *S. pneu*m*oniae* likely represents an evolutionary intermediate in route to losing this branch of the pathway. The accumulation of cmnm⁵s²U appears to compensate for the deficiency of the fully modified mnm⁵s²U, as strains lacking this modification do not display a growth phenotype or any other metabolic defects. Validation of functional assignments for alternate biosynthetic pathways awaits experimental demonstration. Nevertheless, the examples reported in this study provide direction for these investigations.

MATERIALS AND METHODS

General bioinformatic resources

For literature and sequence retrievals, the resources at NCBI (https:// [www.ncbi.nlm.nih.gov/\) \(45\), UniProt \(https://www.uniprot.org\) \(46\), and BV-BRC](https://www.ncbi.nlm.nih.gov/) [\(https://www.bv-brc.org\)](https://www.bv-brc.org) [\(47\)](#page-20-0) were routinely used. PaperBlast was used to find published papers on members of a given family (papers.genomics.lbl.gov/) [\(48\)](#page-20-0). SSNs were generated with the Enzyme Function Initiative (EFI) suite of web tools (https://efi.igb.illi[nois.edu/\) \(49\). SSNs were visualized using Cytoscape \(50\). Species phylogenetic trees](https://efi.igb.illinois.edu/) were usually constructed using phyloT v2 (data version 2023.2). Both species and protein phylogenetic trees were visualized and annotated using iTOL v6 (version 6.8.1; [https://itol.embl.de\)](https://itol.embl.de) [\(51\)](#page-20-0). The tree building and visualization tools of BV-BRC were also used. Absence-presence data were retrieved from the Database of Clustered Orthologous Genes (COG Db) [\(https://ftp.ncbi.nih.gov/pub/COG/COG2020/data/,](https://ftp.ncbi.nih.gov/pub/COG/COG2020/data/) accessed 22 August 2022) [\(34\)](#page-20-0). Physical clustering data were retrieved from the KEGG Database (*via* the protein sequence entry option, "Gene Cluster"). Metabolic pathway and gene symbol/family name annotations were retrieved from KEGG (release 108.0; https:// [www.kegg.jp/kegg/\)\(52\) and UniProtKB \(release 2023_04; https://www.uniprot.org\) \(46\).](https://www.kegg.jp/kegg/) Gizmogene [\(http://www.gizmogene.com/\)](http://www.gizmogene.com/) was used to create gene cluster figures. SubtiWiki was used to explore coexpression data in *B. subtilis* [\(53\)](#page-20-0). Protein sequences were aligned using MultiAlin [\(54\)](#page-20-0).

Absence-presence data extraction, benchmark genome filtration for encoding of MnmAGE homologs, and MnmC1-/MnmC2fusion analysis

The MnmAGE families (COG0482, COG0445, and COG0486, respectively) absence-presence data were extracted from the Database of Clustered Orthologous Genes (COG) [\(https://ftp.ncbi.nih.gov/pub/COG/COG2020/data/,](https://ftp.ncbi.nih.gov/pub/COG/COG2020/data/) accessed 22 August 2022) [\(34\)](#page-20-0). This set was used to create a benchmark set of 968 bacterial genomes encoding MnmAGE homologs (Data S1) that was used to then analyze the presence/absence of MnmC protein family members (COG4121). Because COG does not distinguish between the different subgroups of MnmC (i.e., MnmC1, MnmC2, MnmC1/C2 fusion), CDD Search (batch search, standard view) was used to determine MnmC1 domain-containing proteins, MnmC2 domain-containing proteins, MnmC1/C2 fusions, and other fusions with MnmC2 [\(https://www.ncbi.nlm.nih.gov/Structure/bwrpsb/bwrpsb.cgi,](https://www.ncbi.nlm.nih.gov/Structure/bwrpsb/bwrpsb.cgi) accessed 22 August 2022) [\(55\)](#page-20-0). The specific hits per CDD Search results provided the labels of sequences used in data tables and figures (Data S3). Batch CDD search results also supplied respective sequence lengths.

Physical clustering analyses

Using the COG database for the 968 bacterial benchmark genomes filtered for MnmAGE, YtqA family sequences were acquired (ordered locus names per assembly) (Data S4). For each identified YtqA homolog, the KEGG Database "Gene Cluster" protein entry feature was used to extract physical clustering data. Each set of genes per "Gene Cluster" varied in number, determined by KEGG. These data were expanded into a tabular format per respective cluster YtqA sequence (UniProt Accession ID) (Data S5). All gene cluster proteins were mapped to their respective UniProt Accession IDs and investigated

for methyltransferase (i.e., MnmM-/YtqB-like) domains using three different methods, one BLAST-driven and two HMM-driven (Data S5C through E): (1) traditional BLASTp (release, ncbi-blast-2.14.1+-win64; e-value = 0.0001; UniProtKB derived FASTA sequences as inputs) [\(56\)](#page-20-0); (2) CDD sequence search (output format, Standard; RefSeq identifiers as inputs) [\(57\)](#page-20-0); and (3) InterProScan sequence search (UniProtKB derived FASTA sequences as inputs) [\(58\)](#page-20-0). All MnmM-/YtqB-like absence-presence data were managed and curated, and the outputs of the three different retrieval methods were merged using Microsoft Office Excel (Data S5C and 5D). The same was true for the YtqA-containing gene cluster data and subsequent data transformations required for calculating absence-presence proportions/percentages across various genome subgroups within phylum-level defined taxonomic groups (taxonomic groups derived from those included in the COG absencepresence data; representative taxonomic group tax IDs were manually curated) (Data S6). These mapped percentages would be the source data mapped to the phylogenetic trees of Fig. 2. Trees were generated, mapped, and edited using phyloT v2 (https://phylot.bio[byte.de/index.cgi\) and iToL \(51\). UniProt IDs for all genes of each cluster were mapped.](https://phylot.biobyte.de/index.cgi) Metabolic categories were assigned to each cluster member using KEGG and UniProt pathway category assignments (Data S8). Gene symbol/family names were assigned per their respective annotations (Data S8). Using this data, annotations were subsequently mapped to a tree using iTOL or to SSNs using Cytoscape.

Sequence similarity and gene neighborhood networks

SSNs were generated with the Enzyme Function Initiative (EFI) suite of web tools [\(https://efi.igb.illinois.edu/\)](https://efi.igb.illinois.edu/) [\(49\)](#page-20-0). SSNs were visualized using Cytoscape [\(50\)](#page-20-0). A network was generated using the InterPro identifier IPR005911 (Protein YhcC-like), the taxonomy restricted to include only bacterial sequences. Sequences annotated as fragments by UniProt were excluded. Three preliminary networks were generated for alignment scores (AS) of 90, 97, and finally 100 ("repnode" percentages of 95, 95, and 55, respectively). AS100, repnode 55 was selected for figure generation. After acquiring UniProt accession IDs for all sequences possible, fusions and absence-presence data for MnmC and YtqA (see preceding subsection) were mapped to each preliminary network and then to the final colorized network (Fig. 4). The AS100 network was colorized by cluster and [with all singletons and 2-node clusters removed \(see Data S9 at https://doi.org/10.6084/](https://doi.org/10.6084/m9.figshare.25403692.v1) m9.figshare.25403692.v1).

Density functional theory (geometry optimizations)

Density functional theory (DFT) calculations were carried out using Orca 5.0 [\(59\)](#page-20-0). The set of cmnm⁵s²U molecular models was generated with a variety of protonation states and sites for the initial H-atom abstraction (summarized in Fig. S6) using Chem3D (PerkinElmer). The ribose moiety and the rest of the tRNA were approximated with a methyl group, but all other atoms of the 2-thiouracil were kept as shown in Fig. S6A. Geometry optimization was done in Orca with the BP86 functional [\(60, 61\)](#page-20-0) and def2-SVP basis set [\(62\)](#page-20-0) with def2/J auxiliary functions [\(63\)](#page-20-0) and using the conductor-like polarizable continuum model for water (dielectric constant = 80.4 , refractive index = 1.33) [\(64\)](#page-20-0).

Bacterial strains and growth conditions

All bacterial strains and plasmids used in this work are listed in Table S3. Strains of *S. mutans* were routinely cultured statically at 37°C in 5% CO₂ in BHI medium (BD Biosciences) with 10 µg/mL erythromycin (Sigma) when appropriate. Strains of *B. subtilis* were routinely cultured with shaking at 200 rpm in LB media (Fisher) at 37°C with 50 mg/mL of kanamycin (Sigma-Aldrich) or 10 µg/mL erythromycin when appropriate. *E. coli* strains were cultured with shaking at 200 rpm on LB agar at 37°C supplemented with 100 µg/mL ampicillin (Acros) or 50 µg/mL kanamycin when appropriate. *S. pneumoniae* cells were cultured statically in BHI medium at 37°C in an atmosphere of 5% CO₂. S. *pneumoniae* were strains grown on plates containing trypticase soy agar II (modified;

Becton-Dickinson), and 5% (vol/vol) defibrinated sheep blood (TSAII-BA). Plates were incubated at 37 $^{\circ}$ C in an atmosphere of 5% CO₂. TSAII-BA plates for selections contained antibiotics at concentrations described previously [\(65\)](#page-20-0).

Genetic construction methods

E. coli

The *B. subtilis ytqA-mnmM* gene region was amplified from *B. subtilis* CU1065 genomic DNA prepared using a DNA extraction kit (QuickExtract, Epicentre). PCR amplification was performed using primers YtqA_F and YtqB_R (Table S3) and the resulting fragment was digested with NcoI and BamHI prior to ligation into NcoI/BglII sites of pBAD, the resulting plasmid was named pDS358. The coding sequence for *ytqA* and *ytqB* was amplified with restriction sites NcoI and XhoI/BamHI and NcoI and BamHI from pDS358 using primers YtqA_NcoI_F and YtqA_XhoIBamHI_R and YtqB_NcoI_F and YtgB_BamHI_R, respectively. The resulting amplicons were digested and ligated into pBAD to create pDB359 and pDS365, respectively. The sequences of these constructs were verified through DNA sequence (Azenta). pBAD expression plasmids containing *ytqA-mnmM*, *ytqA*, and *mnmM* were transformed into *E. coli* ∆*mnmC* and cultured in 500 mL of LB medium containing ampicillin (100 µg/mL) and arabinose (2 mg/mL) up to OD600 nm of ~1.0. Cells were harvested by centrifugation and stored at −20°C prior to further analysis.

S. mutans

Mutations in *S. mutans* UA159 were constructed by a PCR ligation mutagenesis method [\(66\)](#page-20-0)). Different sets of primers (listed in Table S4) were used to amplify regions directly upstream and downstream from the gene targeted for deletion and to introduce complementary ligation sites to a non-polar erythromycin resistance cassette. The upstream, downstream, and resistance markers were digested and ligated into a linear piece of DNA, which was subsequently used to transform *S. mutans* cells in the presence of competence stimulating peptide [\(67\)](#page-20-0), and double-cross-over recombination events were selected by plating on BHI agar supplemented with 10 $\mu q/mL$ erythromycin. The correct deletion of the gene was confirmed by PCR with external primers and amplicon length comparison.

S. pneumoniae

Streptococcus pneumoniae strains used in this study are listed in Table S3. Strains were derived from unencapsulated strain IU1824 (D39 Δ*cps rpsL1*), which were derived from the encapsulated serotype-2 D39W progenitor strain IU1690 [\(68\)](#page-21-0). Compared with the progenitor strain IU1690, a spontaneous mutation (Q150STOP) was found in *spd_0567* (*mnmM*) during the construction of IU1824 [\(69\)](#page-21-0). The *mnmM* (Q150STOP) mutation in IU1824 was repaired to *mnmM⁺* in strain IU19835. No growth or microscopic differences were observed between IU1824 and IU19835 (spd_0567⁺) in BHI broth. Strains containing antibiotic markers were constructed by transformation of CSP1-induced competent pneumococcal cells with linear DNA amplicons synthesized by overlapping fusion PCR [\(65\)](#page-20-0). Strains containing markerless alleles in native chromosomal loci were constructed using allele replacement via the P_c-[kan-rpsL⁺] [\(70\)](#page-21-0). Primers used to synthesize different amplicons are listed in Table S4. TT1519 and TT1522 are outside forward primers and TT1520 is the outside reverse primer for strains IU19964, IU19966, and IU19968. Fusion primers for IU19964, IU19966, and IU19968 are TT1618/TT1619, TT1623/TT1624, and TT1621/TT1622, respectively. D39 genomic DNA was used as a template. For the construction of IU19974, the 5′ fragment was obtained using IU11077 [\(69\)](#page-21-0) which contains the 5′ fragment of Δ*bgaA::kan*-t1t2-PZn and primers TT657 and TT1577. Middle fragment and 3′ fragments were obtained using D39 gDNA as templates and primer pairs TT1578/TT1579 and TT1580/CS121. Mutant constructs were confirmed by PCR and DNA sequencing of chromosomal regions corresponding to the amplicon regions used

for transformation. Ectopic expression of *mnmM* was achieved with a P_{Zn} zinc-inducible promoter in the ectopic *bgaA* site. 0.2 mM Zn^{2+} and 0.02 mM Mn^{2+} were added to BHI broth for inducing conditions. Mn^{2+} was added with Zn^{2+} to prevent zinc toxicity [\(71\)](#page-21-0).

B. subtilis

The *B. subtilis mnmM* and *ytqA* mutant were obtained from the Bacillus Genetic Stock Center [\(https://bgsc.org/index.php\)](https://bgsc.org/index.php) and verified by PCR using Oligos BSU_ytqB_VF and BSU_ytqB_VR or BSU_ytqA_VF and BSU_ytqA_VR (see Table S2).

Growth, cell width and length measurements, hydrogen peroxide, and antibiotic sensitivity tests of *Streptococcus pneumoniae*

Growth in BHI broth containing 0.3 M NaCl was performed by the addition of 0.21 M NaCl to BHI broth, which contained 0.09 M NaCl. Cell width and length measurements were performed as described in reference [\(65\)](#page-20-0). Hydrogen peroxide sensitivity assays were performed on bacterial cells grown exponentially in BHI to OD_{620} ~0.2 as descri-bed in reference [\(72\)](#page-21-0). Cells were treated with 20 mM H_2O_2 for 30 min, samples were serially diluted in physiological saline solution [0.9% (wt/vol) NaCl] at room temperature and 5 μL were spotted on TSAII-BA plates. Antibiotic disk-diffusion assays were carried out with the antibiotic Sensi-Disc (Becton Dickinson Pty Ltd., Fosfomycin; cat# 231709, Cefotaxime; cat# 231606, Tetracycline; cat# 230998, and penicillin; cat# 230918) as described in [\(69\)](#page-21-0).

Bulk tRNA extraction

Samples of bacterial culture were used to prepare bulk tRNA extracts for *E. coli*, *S. mutans*, *S. pneumoniae*, and *B. subtilis*. As the LC-MS analyses were done by three laboratories and utilized different methods to prepare tRNA samples were used. In the PDS method, 0.5 L cultures of *E. coli* and *B. subtilis* were harvested by centrifugation, and the cell pellets were treated as previously described to obtain the bulk tRNA samples [\(73\)](#page-21-0) that were stored at −20°C until LC-MS analysis. In the VDCL1 method*, B. subtilis* cell pellets were prepared by diluting overnight (16-18 h) liquid cultures to an initial OD_{600nm} of 0.02, allowed to grow at 37°C with shaking at 200 rpm until a final OD_{600nm} of ~1.0 or another specified value as indicated, and 5 mL of cells were centrifuged. For *B. subtilis* samples, each cell pellet was re-suspended in a solution containing 10 mg/mL lysozyme and sodium acetate (pH5.3, 50 mM sodium acetate) and transferred to a 1.5 mL microcentrifuge tube followed by an incubation at 37°C for 1 h. The resulting lysate was then used to isolate bulk tRNA as previously described [\(74\)](#page-21-0).

S. mutans tRNA samples were prepared using the VDCL2 method. Using this method, each sample was prepared by first diluting overnight (16–18 h) liquid cultures to an initial OD_{600nm} of 0.02 and allowed to grow statically at 37°C in 5% CO₂ until a final OD600nm of ~0.6 was achieved, and 5 mL were centrifuged. Each *S. mutans* cell pellet was re-suspended in a solution of 5 mg/mL lysozyme (lysozyme from chicken egg white, Sigma Aldrich—6876) and TE Buffer (pH 8, 10 mM Tris, 1 mM EDTA) and transferred to a microcentrifuge tube. Each sample was then incubated on wet ice for 30 min. After the cold incubation, a solution of 10% SDS in water was added to each sample tube to reach a final concentration of 1% SDS, followed by an incubation at 95°C for 5 min. The resulting *S. mutans* lysate was then used to isolate bulk tRNA as previously described [\(74\)](#page-21-0).

S. pneumoniae tRNA samples were prepared using the MW method. Each sample processed by the MW was prepared by diluting overnight cultures (12–15 h) to an initial OD_{620nm} 0.005 and allowed to grow until an OD_{620nm} of 0.1-0.3 was achieved. For induction of mmm in the merodiploid strain IU19974, 0.2 mM Zn^{2+} , and 0.02 mM Mn^{2+} were added to BHI broth during day growth. For each tRNA preparation, two 40 mL of culture grown to OD₆₂₀ \approx 0.19 to 0.22 were centrifuged at 16,000 x *q* for 7 min at 4°C. Each pellet was added with 100 µL of lytic buffer (1.2% triton, 2 mM EDTA, 20 mM Tris, pH 8.3), vortexed, and followed by the addition of 500 µL of Trizol. The bacteria Trizol mixtures from the two pellets were combined and added to a 2 mL

tube containing lysing matrix B (MP biomedical, 116911-500), and processed with a FastPrep-24 machine (MP biomedical) three times for 40 s at setting 6 with a 24 \times 2 holder at 4°C. The tubes were centrifuged at 16,000 x *g* for 7 min at 4°C, and the upper layers (~900 µL) were transferred to new microfuge tubes and incubated for 5 min at room temperature. Chloroform (450 µL) was added to the transferred sample, vortexed for 10 s, and incubated for 5 min at room temperature, followed by centrifugation at 16,000 x *g* for 10 min at 4°C. Approximately 450 µL of the upper phase was transferred to microfuge tubes, added with 0.5 x volume (225 µL) of 100% room temperature ethanol, and inverted 10 times to mix. The mixture was added to a PureLink miRNA Isolation Kit Spin Columns, followed by centrifugation at 1 min at 12,000 x *g*. The flow through (-600μ) was transferred to a new microfuge tube and mixed with 700 μ L of 100% ethanol. The mixture was added in two runs to a new PureLink miRNA Isolation Kit Spin Column, centrifuged for 1 min at 12,000 x g, and the flow-through discarded. Wash buffer W5 from the PureLink miRNA Isolation Kit (500 µL) was added to the column and centrifuged for 1 min at 12,000 x *g*. The wash step was performed three times, followed by a final centrifugation of 3 min. The spin column was transferred to a new microfuge tube, and 62 µL of RNase-free water was added and incubated for 1 min. The sample was eluted by centrifugation for 1 min at 21,000 \times g. Samples were quantified with a NanoDrop spectrophotometer, showing typical 260/280 readings of ~2 and 260/230 readings of 2.0 to 2.2. Typical yields from this protocol were 40 to 70 µg from 80 mL of cell culture collected at OD₆₂₀ ≈0.19 to 0.22. The purified samples were stored at −80°C until further analysis.

Mass spectrometric analysis of modified ribonucleosides in tRNA from *B. subtilis***,** *E. coli***, and** *S. pneumoniae*

LC-MS Method #1

tRNA modifications were analyzed by two chromatography-coupled mass spectrometry methods (LC-MS). In LC-MS Method #1, total RNA (40 µg) was heat-denatured at 100°C for 5 min, chilled on ice for 5 min, and allowed to come to room temperature. The RNA was digested in 20 mM TMA-acetate pH 5.3 containing 5 µL P1 nuclease, 10 µL alkaline phosphatase (RSAP), and 20 µL CutSmart buffer. The reaction was incubated at 37°C for 1 h, centrifuged at 12,800 \times *g* for 10 min, and the supernatant was spiked with 5 µL of methanol-formic acid (80:4) before mass spectrometry analysis. Digested total RNA samples were analyzed by HPLC using a Polaris C-18 column (Agilent), coupled with a mass spectrometer (Orbitrap) in positive ion mode, as described in Edwards et al. [\(75\)](#page-21-0). Optima LC-MS grade solvents were 0.1% formic acid in water (solvent A) and 0.1% formic acid in methanol (solvent B). The column was pre-equilibrated with 2% Solvent B for 15 column volumes at 300 µL min−1. Tune file parameters used during runs were as follows: Voltage (kV): 4.01; Sheath Gas Flow Rate (arb): 47.00; Auxiliary Gas Flow Rate (arb): 30.00; Sweep Gas Flow Rate (arb): 0.00; Capillary Voltage (V): 2.00; Capillary Temperature (C): 350; and Tube Lens Voltage (V): 49.89. Digested nucleosides were separated and detected over the course of a 39 min run using the gradient of solvent B as follows: 0– 4 min, hold at 2% B (vol/vol); 4–25 min, increase B from 2% to 100%; 25–33 min, hold at 100% B; and 33–39 min switch from 100% B to 2% B to allow column re-equilibration at 2% B. Standards of dihydrouridine (Dalton) and pseudouridine (Barry & Associates) were run as controls for normalizing data as they are present in lower abundance, similar to the modifications analyzed in this study (76) . The desired analytes D ($[M + H]$ ⁺:247.0925), nm⁵s²U ([M + H]⁺: 290.0805), cmnm⁵s²U ([M + H]⁺: 348.086), mnm⁵s²U ([M + H]⁺:304.096), where the masses were determined within 5 ppm accuracy.

LC-MS Method #2

tRNA from each sample (*B. subtilis:* 1 µg; *S. mutans* 2 µg) was hydrolyzed in a 50 µL digestion cocktail containing 12.5 U (*B. subtilis*) or 8 U (*S. mutans*) benzonase, 5 U CIAP (calf intestinal alkaline phosphatase), 0.15 U PDE I (phosphodiesterase I),

0.1 mM deferoxamine, 0.1 mM BHT (butylated hydroxytoluene), 5 ng coformycin, 50 nM $[^{15}N]_{5}$ -deoxyadenosine ([¹⁵N]-dA; internal standard), 2.5 mM MgCl₂, and 5 mM Tris-Hcl buffer pH 8.0. The digestion mixture was incubated at 37°C for 6 h. After digestion, all samples were analyzed by chromatography-coupled triple-quadrupole mass spectrometry (LC-MS/MS). For each sample, 200 ng (*B. subtilis*) or 600 ng (*S. mutans*) of hydrolysate was injected in two technical replicates. Using synthetic standards, HPLC retention times of RNA modifications were confirmed on a Waters Acuity BEH C18 column (50 \times 2.1 mm inner diameter, 1.7 µm particle size) coupled to an Agilent 1290 HPLC system and an Agilent 6495 triple-quad mass spectrometer. The Agilent sample vial insert was used. The HPLC system was operated at 25°C and a flow rate of 0.3 mL/min in a gradient (Table S5) with Buffer A (0.02% formic acid in double distilled water) and Buffer B (0.02% formic acid in 70% acetonitrile). The HPLC column was coupled to an Agilent 6495 triple quad mass spectrometer with an electrospray ionization source in positive mode with the following parameters: Dry gas temperature, 200°C; gas flow, 11 L/min; nebulizer, 20 psi; sheath gas temperature, 300°C; sheath gas flow, 12 L/min; capillary voltage, 3,000 V; and nozzle voltage, 0 V. Multiple reaction monitoring (MRM) mode was used for detection of product ions derived from the precursor ions for all the RNA modifications with instrument parameters including the collision energy (CE) optimized for maximal sensitivity for the modification. Based on synthetic standards (synthesized as noted) with optimized collision energies, the following transitions and retention times were monitored: cmnm⁵s²U [\(77\)](#page-21-0), m/z 348 \rightarrow 141, 2.3 min; mnm⁵s²U (Biosynth), m/z 304 \rightarrow 172, 1.6 min; nm⁵s²U (Granlen), m/z 290 \rightarrow 158, 1.2 min; D, and m/z 115, 0.864 min. Signal intensities for each ribonucleoside were normalized by dividing by the sum of the UV signal intensities of the four canonical ribonucleosides recorded with an in-line UV spectrophotometer at 260 nm.

ACKNOWLEDGMENTS

We would like to thank Chelsey Bomar for identifying the first YtqA_MnmD fusion during the MCB6318 Spring 2022 class and Scott Dawson, Brystol Habermacher, and Mira Baki for assistance in the cloning of *B. subtilis* genes.

This work was funded by the National Institutes of Health (R01 GM70641 to V.D.C.L., R35 GM131767 to M.E.W., and ES026856, ES024615, and ES031529 to Thomas Begley), the National Research Foundation of Singapore through the Singapore-MIT Alliance for Research and Technology Antimicrobial Resistance Interdisciplinary Research Group (P.C.D.S.), and the National Science Foundation (MCB-1716535 to P.C.D.S.).

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ADDITIONAL FILES

The following material is available [online.](https://doi.org/10.1128/jb.00452-23)

Supplemental Material

Data S1 (JB00452-23-s0001.xlsx). List of the 968 organisms in the COG database that encode the three proteins MnmA, MnmG and MnmE and distribution of the COG4121(MnmC) in those genomes.

Data S2 (JB00452-23-s0002.xlsx). Summary of MnmC1/MnmC2/MnmC1C2 distributions across bacterial orders in the benchmark set of 968 genomes.

Data S3 (JB00452-23-s0003.xlsx). Domain analysis of MnmC1/MnmC2/MnmC1C2 (via CDD Search) in homologs present in benchmark set of 968 genomes.

Data S4 (JB00452-23-s0004.xlsx). (a) List of YtqA/mmM, MnmC1/MnmM and YtqA/ MnmC2 fusion proteins present in InterPro; (b) YtqA/YhcC family proteins found in the benchmark set of 968 genomes.

Data S5 (JB00452-23-s0005.xlsx). YtqA/YhcC gene clusters (a) all, (b) those with mnmM.

Data S6 (JB00452-23-s0006.xlsx). MnmC1/MnmC2/MnmC1C2, and physically clustered YtqA-MnmM occurrence data across bacterial orders.

Data S7 (JB00452-23-s0007.xlsx). Differential YtqA gene cluster, MnmC1/MnmC2/ MnmC1C2 fusion analysis data summary tables.

Data S8 (JB00452-23-s0008.xlsx). Complete YtqA/YhcC gene cluster, MnmC1/MnmC2/ MnmC1C2 fusion analysis data mapping to SSN figure.

Supplemental material (JB00452-23-s0009.pdf). Tables S1 to S6, Fig. S1 to S7, and legends for Data S1 to S8.

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