An unmodified wobble uridine in tRNAs specific for Glutamine, Lysine, and Glutamic acid from *Salmonella enterica* Serovar Typhimurium results in nonviability—Due to increased missense errors?

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**Abstract**

In the wobble position of tRNAs specific for Gln, Lys, and Glu a universally conserved 5-methylene-2-thiouridine derivative (xm\(^5\)s\(^2\)U34, x denotes any of several chemical substituents and 34 denotes the wobble position) is present, which is 5-(carboxy)methylamino-methyl-2-thiouridine ((c)mnm\(^5\)s\(^2\)U34) in Bacteria and 5-methylcarboxymethyl-2-thiouridine (mcm\(^5\)s\(^2\)U34) in Eukarya. Here we show that mutants of the bacterium *Salmonella enterica* Serovar Typhimurium LT2 lacking either the s\(^2\)- or the (c)mnm\(^5\)-group of (c)mnm\(^5\)s\(^2\)U34 grow poorly especially at low temperature and do not grow at all at 15˚C in both rich and glucose minimal media. A double mutant of *S. enterica* lacking both the s\(^2\)- and the (c)mnm\(^5\)-groups, and that thus has an unmodified uridine as wobble nucleoside, is nonviable at different temperatures. Overexpression of tRNA\(^{\text{Gln}}\)\(_{\text{cmnm-s2UUG}}\) lacking either the s\(^2\)- or the (c)mnm\(^5\)-group and of tRNA\(^{\text{Lys}}\)\(_{\text{mnm-s2UUU}}\) lacking the s\(^2\)-group exaggerated the reduced growth induced by the modification deficiency, whereas overexpression of tRNA\(^{\text{Lys}}\)\(_{\text{mnm-s2UUU}}\) lacking the mnm\(^5\)-group did not. From these results we suggest that the primary function of cmnm\(^5\)s\(^2\)U34 in bacterial tRNA\(^{\text{Gln}}\)\(_{\text{cmnm-s2UUG}}\) and mmns\(^5\)s\(^2\)U34 in tRNA\(^{\text{Lys}}\)\(_{\text{mnm-s2UUU}}\) is to prevent missense errors, but the mnm\(^5\)-group of tRNA\(^{\text{Lys}}\)\(_{\text{mmms-s2UUU}}\) does not. However, other translational errors causing the growth defect cannot be excluded. These results are in contrast to what is found in yeast, since overexpression of the corresponding hypomodified yeast tRNAs instead counteracts the modification deficient induced phenotypes. Accordingly, it was suggested that the primary function of cmnm\(^5\)s\(^2\)U34 in these yeast tRNAs is to improve cognate codon reading rather than prevents missense errors. Thus, although the xm\(^5\)s\(^2\)U34 derivatives are universally conserved, their major functional impact on bacterial and eukaryotic tRNAs may be different.
Introduction

Transfer RNA from all organisms contains modified nucleosides, which are derivatives of the four major nucleosides adenosine (A), guanosine (G), cytidine (C), and uridine (U). More than 100 different modified nucleosides have been characterized (http://mods.rna.albany.edu or http://modomics.genesilico.pl). Although their presence in tRNA is scattered over the entire molecule, there are two positions in the tRNA that are more frequently modified than others—about 50% of nucleosides in position 34 (the wobble position) and about 80% in position 37 (next to and 3’ of the anticodon) are modified [1, 2]. Not only are these two positions frequently modified but they also contain many chemically different modifications, which are important for an efficient and accurate decoding of the mRNA (Reviewed in [1]). These modifications alter the chemical properties of the base but also the adoption of syn or anti conformations as well as inducing different tautomeric forms [3]. Unmodified U is present rarely in this position of tRNAs from free living organisms [2]. Apparently, a modified U in the wobble position of tRNAs of most cellular organisms is required for efficient decoding with required fidelity.

Modified wobble U’s are classified in two groups based on their chemical structures. One group consists of 5-hydroxyuridine derivatives with a chemical group attached at position 5 of the uracil base via an ether linkage (xoU34-derivatives) whereas the other group consists of 5-methyluridine derivatives (xmU34-derivatives) with a methylene carbon attached to the C5 atom of uracil (x denotes any of several different chemical substituents). The xoU34-derivatives are found in tRNAs reading family codon boxes and the modification expands wobble capacity of uridine to read 3 to 4 codons in such a codon box. The xmU34-derivatives are present in tRNAs reading mixed codon boxes and decode codons ending with A or G. They may also have a sulfur bound to position 2 of uracil (xmS2U34-derivatives; reviewed in [1]). These wobble derivatives (xmS2U34) are universally conserved in tRNAs specific for Gln, Lys, and Glu and, accordingly, are present in the three phylogenetic domains Bacteria, Eukarya and Archaea and also in organelles such as mitochondria and chloroplasts. The 5-(carboxy)methylaminomethyl-2-thiouridine ((c)mnmS2U34) derivative of these modifications are present as wobble nucleosides in bacterial tRNAs specific for Gln, Lys, and Glu whereas the wobble nucleoside in the eukaryotic tRNA counterparts is 5-methylcarboxymethyl-2-thiouridine (mcmS2U34) (Fig 1). Such derivatives were thought to restrict the wobble capacity of U and thereby improving the recognition of purine-ending codons and prevent misreading of the near-cognate codons ending in pyrimidines [4] (cf. Fig 2). The synthesis of the latter in yeast requires the activity of several genes, of which ELP3 is required for the early step(s) in the synthesis of the mcmS2-side chain and TUC1 is required for last step in the thiolation at position 2 [5, 6]. The yeast double mutant elp3, tuc1, which thus contains an unmodified wobble uridine in tRNAs specific for Gln, Lys, and Glu, is nonviable but can be rescued by overexpression of the corresponding hypomodified tRNAs. The latter result suggests that the primary function of mcmS2U34 is to improve the efficiency of decoding cognate codons rather than to prevent missense errors [5].

In bacteria the functional aspect of (c)mnmS2U34 is more complex. Although it was thought that the function of these modifications was to prevent misreading, Hagervall et al [7] showed that this is not the case for tRNA^{Lys}_{Gln^{mnmS2U34}}, since lack of either the mnmS2- or the S2-group reduces misreading of the near-cognate Asn-codons (AAU/C, see Fig 2). Moreover, Manickam et al [8] confirm these results for the lack of the mnmS2- side chain and further show that such hypomodified tRNA^{Lys}_{Gln^{mnmS2U34}} also decreases other missense errors (Arg codons AGA/G; see Fig 2). However, lack of mnmS2-side chain in tRNA^{Glu}_{Glu^{mnmS2U34}} increases missense errors of Gly (GGA) and Asp (Glu/C) codons [8]. Thus, the same modification (mnmS2-side
chain) in tRNA_{Lys}^{mnm}s^2U or in tRNA_{Glu}^{mnm}s^2U influences the frequency of missense errors differently—increasing (tRNA_{Lys}^{mnm}s^2U) or decreasing (tRNA_{Glu}^{mnm}s^2U). Therefore, the functional impact of the modification is sensitive to the structural context it is part of, which has also been shown for the modified nucleosides queosine (Q34) [8] and the uridine-5-oxyacetic acid (cmo^5U34) [9].

Transfer RNA specific of Gln, Lys, and Glu are the only tRNA species in bacteria that contain (c)mnm^5s^2U34 as wobble nucleoside. Whereas tRNA_{Gln}^{cmmn}s^2U34 contains a mixture of cmnm^5s^2U (80%) and mnm^5s^2U (20%) [10–12], tRNA_{Lys}^{mmn}s^2U34 and tRNA_{Glu}^{mmn}s^2U34 contain mnm^5s^2U34. The formation of (c)mnm^5s^2U34 requires the activity of several enzymes as summarized in Fig 3. A predicted minimal translation apparatus requires the mnmA, mnmE, and mnmG genes but not the mnmC gene to synthesize cmnm^5s^2U34, which suggests an important role of this wobble modification for a proper translation [13]. The s^2- and the (c)mnm^5-group of (c)mnm^5s^2U34 are pivotal for reading frame maintenance and lack of either of them induces ability to suppress frameshift mutations [14]. We have characterized several frameshift suppressor mutants lacking this modified nucleoside and these mutants are our tools in studying the function of the (c)mnm^5s^2U34 wobble nucleoside [15]. Mutations in mnmA, tusB, or tusE block the sulfur relay pathway whereas mutations in mnmE or mnmG (gidA) block the synthesis of the (c)mnm^5-side chain and accordingly such mutants lack (c)mnm^5s^2U34 (Fig 3, Table 1).

This paper focuses on the functional aspect of the wobble nucleoside (c)mnm^5s^2U34 in bacteria. We show here that lack of either the s^2- or the (c)mnm^5-group induces poor growth and cold sensitivity, that a double mutant having an unmodified U34 as wobble nucleoside is not viable, and that overexpression of tRNA_{Gln}^{cmmn}s^2U34 or tRNA_{Lys}^{mmn}s^2U34 lacking the s^2-group reduces growth considerably suggesting that such hypomodified tRNAs increase missense error. Overexpression of tRNA_{Gln}^{Gln} lacking the cmnm^5-group reduces growth although much less than s^2-deficiency, whereas lack of the mnm^5-group of overexpressed tRNA_{Lys}^{Lys} does not.
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![Codon table and the anticodon loop of tRNAs specific for Gln, Lys, and Glu which contain cmnm^5s^2U (Gln) or mnm^5s^2U (Lys and Glu) as wobble nucleoside in bacterial tRNAs.](https://doi.org/10.1371/journal.pone.0175092.g002)

**Fig 2.** Codon table and the anticodon loop of tRNAs specific for Gln, Lys, and Glu which contain cmnm^5s^2U (Gln) or mnm^5s^2U (Lys and Glu) as wobble nucleoside in bacterial tRNAs. Above the third column (second codon base A) the anticodon stem and loop of the tRNAs containing the (c)mnm^5s^2U in position 34 is shown (denoted in red cmnm^5s^2U). Position 36 of the anticodon is color coded with Green G36, position 32 of tRNA Gln red U36, and position 33 of tRNA Lys and Glu blue C36. Similar color code is denoted for the modified nucleosides present in position 37 in the corresponding tRNAs. N denotes G, A or C, respectively, for first nucleoside in the relevant codons read by these tRNAs. Note that these tRNAs are rich in U, which is a poor stacker [24] making the anticodon very flexible especially if the modifications is absent and this is especially true of tRNA^Lys^Gln. In the codon table the letters outside the box, to the left, above, and to the right indicate the first, second, and third position of the codon. Circles connected by a line, or a single circle, represent one tRNA species. A filled circle indicates the capacity of that tRNA to base pair with the indicated codon, either by Watson-Crick or by wobble according to the revised wobble hypothesis [1]. Red and yellow circles indicate tRNAs that are sequenced at the RNA level while green circles represent tRNAs for which only a partial tRNA sequence is available. A red or green circle indicates efficient base pairing while a yellow circle indicates a restricted wobble. An open (white) circle is a base pairing that is not according to the revised wobble hypothesis. However, data in vivo from mutants where only this tRNA is left to decode all codons in the codon box, suggest that the tRNA in fact is able to read that codon [9, 25]. Data are compiled from Sprinzl data base (http://trna db.bioinf.uni-leipzig.de/) and Modomics data base (http://modomics. genesilico.pl/). Mutations in mmmE or mmmG genes result in no formation of the (c)mnm^5s^2U present in tRNAs specific for Glu, Lys and Glu but also cmnm^5s^2U in tRNA^Glu^ for Arg and Gly. Mutations in mmmA result in no thiolation of cmnm^5s^2U34.  

a) The modifications in position 34 of tRNA^Glu^ from S. enterica are cmnm^5s^2U (80%) and mnm^5s^2U (20%).  

b) The majority of this tRNA^Glu^ contains mnm^5U34 but there is also a small amount of cmnm^5U34.

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Materials and methods

Bacteria and growth conditions

The bacterial strains used were derivatives of *Salmonella enterica* serovar Typhimurium LT2 or *Escherichia coli* K12 (S1 Table, Supporting materials). As rich medium Luria-
Bertani (LB) was used [16]. The minimal solid medium was made from Vogel & Bonner basal salt medium [17]. When necessary antibiotics were added at following concentrations: carbenicillin 100 μg ml⁻¹, kanamycin 50 μg ml⁻¹, and chloramphenicol 25 μg ml⁻¹.

### Genetic procedures

Transduction with phage P22 HT105/1 (int-201) [19] was performed as previously described [18]. DNA sequencing was performed on plasmid DNA or PCR products following the manual of Applied Biosystems ABI PRISM Cycle Sequencing Ready Reaction Kit Big Dye™ v.1.1 or by LightRun™ sequencing at GATC Biotech, Cologna, Germany. The inactivation of mnmA and mnmE genes was done according to Datsenko & Wanner [20] and the resulting insertions were confirmed by PCR.

### Analysis of modified nucleosides in tRNA

Bacterial strains were grown over night in medium LB, diluted 100 times in 100 ml of the same medium and grown at 37°C to a cell density of about 4x10⁸ cells/ml. Cells were lysed and total RNA was prepared [21] and dissolved in 2 ml buffer R200 (10 mM Tris-H₃PO₄ (pH 6.3), 15% ethanol, 200 mM KCl) and applied to a Nucleobond® AX500 column (Macherey-Nagel GmbH & Co., Düren, Germany), pre-equilibrated with the same buffer. The column was washed once with 6 ml R200 and once with 2 ml R650 (same composition as R200, except for 650 mM KCl instead of 200 mM KCl). Finally, tRNA was eluted with 7 ml R650, precipitated by 0.7 volumes cold isopropanol, washed twice with 70% ethanol and dissolved in water.

### Table 1. Level of (c)mnm⁵s²U in tRNAs of various strains used in this study.

| Strain      | Relevant genotype | s²C/Ψ 254 nm | mnm⁵s²U/Ψ # 254 nm | s²U/Ψ 254 nm | (c)mnm⁵s²U/s⁴U 314 nm | s²U/s⁴U 314 nm |
|-------------|------------------|--------------|--------------------|--------------|----------------------|---------------|
| GT7321      | mnmA⁺, mnmE⁺     | 0.10         | 0.042              | <0.001       | 0.047                | <0.001        |
| GT8173      | mnmA16<cat       | 0.12         | <0.001             | <0.001       | <0.003               | <0.002        |
| GT8176      | mnmE17<kan       | 0.12         | <0.003             | 0.048        | <0.002               | 0.019         |
| GT8177      | pmmnA⁺/mnmA16<cat | 0.12         | 0.054              | <0.001       | 0.046                | <0.002        |
| GT7440      | mnmA3(G24D)      | 0.12         | <0.001             | d            | <0.001               | D             |
| GT7453      | tusB27(Q31stop)  | 0.12         | <0.001             | d            | <0.001               | D             |
| GT7432      | tusE30(K128stop) | 0.11         | <0.003             | d            | <0.002               | D             |
| GT7436      | mnmE13(codon 241–271 deleted) | 0.14 | <0.001 | d | <0.001 | D |
| GT7478      | mnmG1(gidA1, 58 nt deletion from A402) | 0.13 | <0.002 | d | <0.001 | D |

a) The cmnm5s2U peak is hidden under contaminating material, so it is not possible to determine the area.
b) This level is likely to be an overestimate due to poor separation from especially G. (See Fig 4C). The level is about 100% of the average level of mnm⁵s²U in the two wild type strains GT7321 and GT8177.
c) Strain GT8177 contains a plasmid from the Saka collection, [45], which harbors the wild type allele of mnmA⁺ gene and the mnmA16<cat insertion on the chromosome.
d) These analysis were made using a Supelco C-18 column, which does not separate s²U from s⁴U. However, this analysis determines excellently the level of mnm⁵s²U at 254 nm and both cmnm⁵s²U and mnm⁵s²U at 314 nm.

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Transfer RNA was digested to nucleosides by nuclease P1 followed by treatment with bacterial alkaline phosphatase at pH 8.3 [22]. C:\GetAREf\Refs\Jocke\cmo5U.ref #91; The hydrolysate was analyzed as described earlier [23] using a Supelcosil C-18 column (Supelco) or Develosil C-30 (Phenomenex) with a Waters Alliance HPLC system.

**Results**

The *mmmA* or *mmmE* mutants have undetectable level of (c)mnm$^5$s$^2$U34 in tRNA

To test the viability of a mutant containing mutations in both *mmmA* and *mmmE*, selectable markers linked to the inactivation of these genes were required. We therefore inserted in the *mmmA* gene a chloramphenicol (*cat*) resistant element and in the *mmmE* gene a kanamycin (*Km*) resistant element (denoted *mmmA*16<i>&gt;*cat* and *mmmE*17<i>&gt;*Km*, respectively), which completely destroy the synthesis of the corresponding enzymes and block the synthesis of (c)mnm$^5$s$^2$U34. In an *mmmA*16<i>&gt;*cat*, *mmmE*17<i>&gt;*Km* double mutant tRNAs specific for Gln, Lys and Glu will therefore contain an unmodified wobble uridine (U34). These resistance elements should not induce any polar effect on the expression on downstream genes, since both genes are transcribed as single cistrons (http://regulondb.cgg.unam.mx/).

The biosynthesis of the cmnm$^5$- and the s$^2$-groups are suggested to be independent of each other [27–29]. Accordingly, a deletion of the *mmmE* or the *mmmG* gene should result in tRNAs specific for Gln, Lys and Glu containing s$^2$U34 instead of (c)mnm$^5$s$^2$U (Fig 3). If the thiolation is not dependent of the presence of the (c)mnm$^5$-side chain, a similar level of moles of s$^2$U should accumulate as the level of (c)mnm$^5$s$^2$U34 present in the wild type. Indeed, s$^2$U is observed in the *mmmE*17<i>&gt;*Km* mutant but at a level of 40% (at 314 nm) and 100% (at 254 nm) of the level of (c)mnm$^5$s$^2$U in the wild type (assuming the same extension coefficient constants for (c)mnm$^5$s$^2$U and s$^2$U; Table 1, Fig 4). Note, that s$^2$U is not is well separated from s$^2$U at 314 nm and G at 254 nm (Fig 4C and 4E) making the analysis not optimal. However, analysis of tRNA of an *mmmE* mutant of *E. coli* s$^2$U is also about 50% of the level of cmnm$^5$s$^2$U in the wild type at analysis conditions where s$^2$U is well separated from s$^2$U [30]. On the other hand determinations using radioactive labelling have found higher level of s$^2$U in relation to the level of (c)mnm$^5$s$^2$U (80% by Hargervall et al [11]). Although the *mmmE* mutant used has s$^2$U in its tRNA we cannot rule out that the level might not reach the expected level even though thiolation, as has been suggested, is not dependent on the presence of the cmnm$^5$-side chain.

A mutation in the *mmmA*, *tusB* or *tusE* genes should result in the presence of cmnm$^5$U34 in the Gln- and cmnm$^5$U34 in Lys- and Glu-tRNAs (cf Fig 3). However, in the HPLC analysis used the cmnm$^5$U is not separated from the major nucleoside C. However, we have earlier analysed the distribution of modified nucleosides in an *mmmA* mutant and such a mutant contains cmnm$^5$U34 instead of cmnm$^5$s$^2$U34 [7]. Moreover, *mmmA*, *tusB* or *tusE* deletion mutants contain cmnm$^5$U34 in tRNA instead of cmnm$^5$s$^2$U34 [31]. Since the (c)mnm$^5$s$^2$U is not detected in tRNA of the *mmmA*, *tusB* and *tusE* mutants used (Table 1), it is likely that they instead have (c)mnm$^5$U in their tRNAs specific to Gln-, Lys-, and Glu.

Lack of the s$^2$- or (c)mnm$^5$-group of (c)mnm$^5$s$^2$U as in the *mmmA* and *mmmE* mutants, respectively, results in a severe growth reduction especially at low temperature

As described in the introduction yeast has the chemically related wobble nucleoside cmcm$^5$s$^2$U in the corresponding tRNA species in which (c)mnm$^5$s$^2$U34 is present in *S. enterica* and *E. coli* (Fig 1). The Elp3p catalyses the first step in the synthesis of the cmcm$^5$-group and the Tuc1p the
last step in synthesis of the s\(^2\)-group [5, 6]. A double mutant elp3, tuc1 is nonviable demonstrating that an unmodified U34 in the Gln, Lys and Glu-tRNA does not support viability in yeast [5]. However, lowering the temperature allows such a double mutant to grow, although poorly. Assuming that a bacterial mnmA, mnmE double mutant is also viable at lower temperature, we monitored the growth of the mnmA\(^+\) and mnmE\(^+\) single mutants at several temperatures and growth media. However, both single mutants grew very poorly at all conditions tested, especially at low temperature, and not at all at 15˚C [on rich (mnmA and E mutants) and minimal medium (mnmA mutant)] (Table 2). The reduction of growth at 37˚C is similar to the growth behaviour of mnmA, tusB and tusE mutants described by Suzuki and collaborators [31].

**Overexpression of Gln or Lys specific tRNAs lacking the 2-thio- or the (c)mnm\(^5\)- group of (c)mnm\(^5\)s\(^2\)U34 causes severe growth reduction**

In a yeast elp3, tuc1 double mutant tRNAs specific for Gln, Lys, and Glu lack cmm\(^5\)s\(^2\)U34 in their wobble positions and have instead an unmodified U34. Such a double mutant is not viable, but overexpression of hypomodified versions of these tRNAs rescues the double mutant elp3, tuc1 [5]. A corresponding bacterial mnmA, mnmE double mutant, which should also have an unmodified wobble U34 in the corresponding tRNAs, would therefore be potentially nonviable. Since the x\(^m\)s\(^2\)U34 derivatives are universally conserved their function might be similar so that overexpression of bacterial hypomodified tRNAs might, as in yeast, rescue a
Table 2. The mmmA and mnmE mutants are cold sensitive on rich and minimal glucose media.

|                      | LAL (relative colony size) | Glucose+His (relative colony size) |
|----------------------|---------------------------|-----------------------------------|
|                      | GT7321 (wt)               | GT8176 (mnmE17<->Km)              | GT8176 (mnmA16<->cat) | GT8173 (mnmA16<->cat) | GT8173 (mnmE17<->Km) |
| Temp 42.5°C          | 1.0 (1.7)                 | 0.7                               | <0.1                  | 41 (3)                | 1.0 (1.8)           |
| 37°C                 | 1.0 (1.2)                 | 0.75                              | 0.5                   | 37 (3)                | 1.0 (2.0)           |
| 30°C                 | 1.0 (3.2)                 | 0.16                              | no sc                 | 30 (3)                | 1.0 (1.5)           |
| RT                   | 1.0 (1.1)                 | 0.45                              | no sc                 | RT (3)                | 1.0 (0.5)           |
| 15°C                 | 1.0 (1.0)                 | <0.1                              | no sc < 0.1 (14 d)    | 15 (3)                | 1.0 (0.9)           |

Values are given as colony size (in mm; average of 10 isolated colonies) relative to that of the size of the wild type following the same time of incubation. The colony size of the wild type is given within parenthesis. <0.1, colonies less than 0.1 mm; no sc: no single cell colonies was observed and growth was observed only on primary and secondary streaks. The size of the colony was determined under a magnifying glass and its diameter was determined using a slide calliper. (LAL-plates, rich media, Glucose+His–plates with minimal media)
a) scored after one day  
b) scored after two days  
c) scored after three days  
d) scored after four days and  
e) scored after six days. When the time of incubation of the plates was longer than that for the wild type, the number of days are indicated within parenthesis.

Potential nonviable double mutant mmmA, mnmE. Overexpression of a hypomodified tRNA (e.g. hypomodified tRNA\textsubscript{Gln}\textsuperscript{cmnm\textsubscript{5}\textsubscript{2}UUG}) in the recipient cell might therefore facilitate the construction of a viable mutant having both mmmA and mnmE mutations on the chromosome and a rescuing plasmid encoding e.g. hypomodified tRNA\textsubscript{Lys}\textsuperscript{cmnm\textsubscript{5}\textsubscript{2}UUG}. We therefore first monitored how overexpression of a hypomodified tRNA\textsubscript{Gln}\textsuperscript{cmnm\textsubscript{5}\textsubscript{2}UUG} or tRNA\textsubscript{Lys}\textsuperscript{cmnm\textsubscript{5}\textsubscript{2}UUG} influenced growth of the mmmA3 and mnmE13 mutants.

**Overexpression of tRNA\textsubscript{Gln}\textsuperscript{cmnm\textsubscript{5}\textsubscript{2}UUG} lacking the s\textsuperscript{2}-group and possessing only the cmnm\textsuperscript{5}-group of cmnm\textsuperscript{5}\textsuperscript{2}T34.** Strains having a mutation in mmmA, tusB or tusE genes lack the s\textsuperscript{2}-group of (c)mmm\textsubscript{5}\textsuperscript{2}U34 of tRNA\textsubscript{Gln}\textsuperscript{cmnm\textsubscript{5}\textsubscript{2}UUG} and such a tRNA has only the cmnm\textsuperscript{5}-group (See above). Plasmid pUST313 harbors the metT operon consisting of genes glnU and glnW encoding the (c)mmm\textsuperscript{5}\textsuperscript{2}T34 containing tRNA\textsubscript{Gln}\textsuperscript{cmnm\textsubscript{5}\textsubscript{2}UUG} and genes glnV and glnX encoding the C34 containing tRNA\textsubscript{Gln}\textsuperscript{CUG}. This operon also contains the metT/U genes encoding tRNA\textsubscript{Met}\textsuperscript{cmnm\textsubscript{5}\textsubscript{2}UUG} and the leuW gene encoding tRNA\textsubscript{Leu}\textsuperscript{cmnm\textsubscript{5}\textsubscript{2}UUG} (Fig 5). Note, no other full sized gene is present on this plasmid. We introduced this plasmid into mmmA3, tusB27 and tusE30 mutants as well as into the wild type strain. Overexpression of tRNA\textsubscript{Gln}\textsuperscript{cmnm\textsubscript{5}\textsubscript{2}UUG} from plasmid pUST313 in the wild type influenced the growth only to a minor degree (Table 3). Slow growth as such was not sensitive to the overexpression of fully modified tRNA\textsubscript{Gln}\textsuperscript{cmnm\textsubscript{5}\textsubscript{2}UUG} as tested by introducing the same plasmid into a mutant having a similar growth rate as an mmmA mutant (data not shown). However, plasmid pUST313 severely reduced the growth of the three independent mutants mmmA3, tusB27 and tusE30 all lacking the s\textsuperscript{2}-group of (c)mmm\textsuperscript{5}\textsuperscript{2}T34 in tRNA\textsubscript{Gln}\textsuperscript{cmnm\textsubscript{5}\textsubscript{2}UUG}. Since the results were the same in the three independent s\textsuperscript{2}-deficient mutants, it was the lack of the s\textsuperscript{2}-group of (c)mmm\textsuperscript{5}\textsuperscript{2}U in tRNA\textsubscript{Gln}\textsuperscript{cmnm\textsubscript{5}\textsubscript{2}UUG}, that caused the reduced growth and no other aberrations in the cell. We obtained a few large colonies among the small colonies when introducing plasmid pUST313 into the mmmA3 mutant. Sequencing the metT operon of the plasmid present in such a large spontaneously occurring colony revealed that a recombination had occurred between the plasmid encoded metT and metU genes resulting in the loss of genes glnU and glnW, which code for tRNA\textsubscript{Gln}\textsuperscript{cmnm\textsubscript{5}\textsubscript{2}UUG} (Fig 5). The resulting plasmid, pUST314,
Fig 5. Gene organization of plasmids pUST313 and pUST314. Plasmid pUST313 harbors the genes glnU and glnW (red) encoding the (c)mnm5s2U34 containing Gln-tRNA (tRNA<sup>Gln</sup><sub>cmnm5s2U34</sub>) and genes glnV and glnX (blue) encoding the C34 containing tRNA<sup>Gln</sup><sub>cUG</sub>. Plasmid pUST314 is a spontaneous plasmid containing glnU and glnW genes encoding tRNA<sup>Gln</sup><sub>cmnm5s2U34</sub> and leuW (black). Note, only part of the asnB and STM0672 genes are present (shaded gray) suggesting that these genes do not express any full sized protein. Plasmid p815 (not shown in the figure but used in some experiments) contains the valU operon, which consists of genes valU, valX, valY and lysV. The latter is the structural gene for tRNA<sup>Met</sup><sub>ac4CAU</sub>

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therefore lacks these genes and have only metT/U gene encoding tRNA<sup>Met</sup><sub>ac4CAU</sub> and glnV and glnX genes encoding the C34 containing tRNA<sup>Gln</sup><sub>cUG</sub>. This plasmid caused only a 50% growth reduction of the mmmA3 and the tusB27 mutants compared to the almost complete inhibition of growth caused by the tRNA<sup>Gln</sup><sub>cmnm5s2U34</sub> expressing plasmid pUST313. Thus, a substantial part of the growth reduction is caused by overexpression of tRNA<sup>Gln</sup><sub>cmnm5s2U34</sub> (Table 3). Plasmid pUST314 also reduced growth suggesting that overexpression of tRNA<sup>Gln</sup><sub>cmnm5s2U34</sub> and of tRNA<sup>Gln</sup><sub>cUG</sub> had some growth reducing effect in strains lacking the s<sup>2</sup>-group of (c)mnm5s<sup>2</sup>U34 but not at all in the wild type strain. In summary, overexpression of s<sup>2</sup>U-deficient tRNA<sup>Gln</sup><sub>cmnm5s2U34</sub> in three independent mutants (mmmA3, tusB27 and tusE30) induced a severe growth reduction that was not observed if this tRNA was not overexpressed.

On minimal medium we also observed a severe growth reduction by overexpression of tRNA<sup>Gln</sup><sub>cmnm5s2U34</sub> in all mutants lacking the s<sup>2</sup>-group of (c)mnm5s<sup>2</sup>U34 but not in the wild type control. A similar growth reduction was also observed with plasmid pUST314, which does not encode tRNA<sup>Gln</sup><sub>cmnm5s2U34</sub> but overexpressed tRNA<sup>Met</sup><sub>ac4CAU</sub> and tRNA<sup>Gln</sup><sub>cUG</sub>. Therefore, we cannot attribute this growth reduction to the overexpression of tRNA<sup>Gln</sup><sub>cmnm5s2U34</sub> but to the s<sup>2</sup>-deficiency of (c)mnm5s<sup>2</sup>U34 in the cell, since a similar growth reduction was not observed in the wild type strain (Table 3). Apparently, overexpression of the tRNAs from these two plasmids (+/- expression of tRNA<sup>Gln</sup><sub>cmnm5s2U34</sub>) induces a severe growth reduction. Since this was observed on minimal medium, an unbalanced tRNA population due to the overexpression of the tRNAs encoded by these plasmids might induce in s<sup>2</sup>-deficient strains aberrations of translation of several genes involved in the synthesis of metabolites resulting in multiple auxotrophy.

Overexpression of tRNA<sup>Gln</sup><sub>cmnm5s2U34</sub> lacking the cmmn<sup>5</sup>-group and possessing only the s<sup>2</sup>-group of (c)mnm5s<sup>2</sup>U34. Plasmid pUST313 was introduced to the mmmE13 and mmmG1 mutants, which both lack the cmmn<sup>5</sup>-group of cmmn5s<sup>2</sup>U34 in their tRNA<sup>Gln</sup><sub>cmnm5s2U34</sub>, and have instead s<sup>2</sup>U34 as wobble nucleoside. This caused also a growth reduction on rich medium although not to the same extent as when this plasmid was introduced into cells lacking the s<sup>2</sup>-group of (c)mnm5s<sup>2</sup>U34 (Table 3). Since a similar result was obtained in two different mutants (mmmE13 or mmmG1) inducing a similar mmm<sup>5</sup>-group deficiency, the growth reduction is
Table 3. Overexpression of hypomodified tRNA\(^{Gln, Lys, Glu}_{\text{Gln, Lys, Glu}}\) severely reduced cellular growth.

| Relevant genotype on chromosome | Wobble nucleoside in tRNA\(^{Gln, Lys, Glu}_{\text{Gln, Lys, Glu}}\) | Plasmid (Relevant genotype) | Growth on LAL-plates. (Rel. colony size (1 d)) | Growth on His-glucose-plates. (Rel. colony size (2 d)) |
|---------------------------------|-----------------------------------------------|-----------------------------|-----------------------------------------------|--------------------------------------------------|
| Wild type                       | (c)mnm\(^5\)s\(^2\)U34                       | No plasmid                  | 0.9 1.0                                       | 1.0 1.0                                          |
|                                 | pUST312 (vector)                             | 1.0 1.0                     | 1.0 1.0                                       |                                                  |
|                                 | pUST313 (glnU,X\(^X\))                      | 0.8 0.8                     | 0.6 0.8                                       |                                                  |
|                                 | pUST314 (ΔglnU,X\(^X\))                     | 1.0                         | 0.9                                           |                                                  |
|                                 | p815 (lysV\(^X\))                           | 0.9                         | 1.0                                           |                                                  |
| mmmA3                           | (c)mnm\(^5\)s\(^2\)-group                    | No plasmid                  | 1.1                                           | 1.1                                              |
|                                 | pUST312 (vector)                             | 1.0 1.0                     | 1.0 1.0                                       |                                                  |
|                                 | pUST313 (glnU,X\(^X\))                      | No sc; <0.1                 | <0.1; <0.1                                    |                                                  |
|                                 | pUST314 (ΔglnU,X\(^X\))                     | 0.6;                         | <0.1                                          |                                                  |
|                                 | p815 (lysV\(^X\))                           | 0.2                         | 0.1                                           |                                                  |
| tusB27                          | (c)mnm\(^5\)s\(^2\)-group                    | No plasmid                  | 0.9                                           | 1.0                                              |
|                                 | pUST312 (vector)                             | 1.0 1.0                     | 1.0                                            |                                                  |
|                                 | pUST313 (glnU,X\(^X\))                      | No sc; No sc; No sc         | <0.1; No sc                                   |                                                  |
|                                 | pUST314 (ΔglnU,X\(^X\))                     | 0.5;                         | 0.1                                           |                                                  |
|                                 | p815 (lysV\(^X\))                           | 0.2                         | 0.1                                           |                                                  |
| tusE30                          | (c)mnm\(^5\)s\(^2\)-group                    | No plasmid                  | 0.9                                           | 1.1                                              |
|                                 | pUST312 (vector)                             | 1.0                          | 1.0                                           |                                                  |
|                                 | pUST313 (glnU,X\(^X\))                      | No sc; No sc; No sc         | <0.1; No sc                                   |                                                  |
|                                 | pUST314 (ΔglnU,X\(^X\))                     | 0.2;                         | 0.1                                           |                                                  |
|                                 | p815 (lysV\(^X\))                           |                             |                                               |                                                  |
| mmmE13                          | s\(^2\)U34(lacks the (c)mnm\(^5\)-group)    | No plasmid                  | 0.9                                           | 1.0                                              |
|                                 | pUST312 (vector)                             | 1.0                          | 1.0                                           |                                                  |
|                                 | pUST313 (glnU,X\(^X\))                      | No sc; No sc; No sc         | 0.2; No sc                                    |                                                  |
|                                 | pUST314 (ΔglnU,X\(^X\))                     | 0.2;                         | 0.1                                           |                                                  |
|                                 | p815 (lysV\(^X\))                           |                             |                                               |                                                  |
| mmmG1                           | s\(^2\)U34(lacks the (c)mnm\(^5\)-group)    | No plasmid                  | 0.9                                           | 1.1                                              |
|                                 | pUST312 (vector)                             | 1.0 1.0                     | 1.0 1.0                                       |                                                  |
|                                 | pUST313 (glnU,X\(^X\))                      | 0.5; 0.4                     | 0.4 0.2                                       |                                                  |
|                                 | pUST314 (ΔglnU,X\(^X\))                     | ND                           | ND                                            |                                                  |
|                                 | p815 (lysV\(^X\))                           | 1.2                         | 1.0                                           |                                                  |

Indicated strains were grown at 37˚C in the rich media LB overnight and then streaked out on LAL agar plates or on glucose medium containing histidine, since the host strain GT7321 requires histidine for growth. No sc, denotes that no single colonies were observed and growth only on primary and/or secondary outstreaks were observed. Numbers are the average of 10 similar sized colonies after indicated time of incubation. When two numbers are present the experiment was repeated once. The size of the colonies was determined as described in Table 2. ND = not done

a) Plasmid pUST313 carry the metT operon which contains the tRNA genes: metT-leuW-glnU-glnW-metU-glnV-glnX. The genes glnU and glnW encodes glutamine tRNA having (c)mnm\(^5\)s\(^2\)U34 as wobble nucleoside whereas glnV and glnX encodes glutamine tRNA having C34 as wobble nucleoside.

b) Plasmid pUST314 carry a part of the metT operon due to a spontaneous recombination between the metT and metU genes. It therefore lacks leuW-glnU-glnW genes and contains only the metT/glnV-glnX. The glnV and glnX genes encodes gln-tRNA having C34 as wobble nucleoside. Thus this plasmid lacks genes for the gln-tRNA havning (c)mnm\(^5\)s\(^2\)-group as wobble nucleoside.

c) Plasmid p815 contains the valU- operon which contains the valU-valX-valY and lysVtRNA genes. The lysV gene encodes lys-tRNA having mnm\(^5\)s\(^2\)U34 as wobble nucleoside.

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likely due to the hypomodified tRNA\textsubscript{Gln}\textsubscript{mnmA16} and not to some other aberrations in these cells. There was no or very little growth reduction caused by plasmid pUST314, which lacks the genes encoding tRNA\textsubscript{Gln}\textsubscript{mnmE17}. Thus, the decreased growth observed is caused by overexpression of hypomodified tRNA\textsubscript{Gln}\textsubscript{mnmA16}. Note, that the observed growth reduction on minimal medium was larger than on rich medium and it was dependent on overexpression of tRNA\textsubscript{Gln}\textsubscript{mnmA16}. Thus, overexpression of tRNA\textsubscript{Gln}\textsubscript{mnmA16} lacking the cmnm\textsuperscript{5}-group causes a substantial growth reduction on rich medium and even a more severe growth reduction on minimal medium. We suggest that tRNA\textsubscript{Gln}\textsubscript{mnmA16} lacking the cmnm\textsuperscript{5}-group induces missense errors. Interestingly, tRNA\textsubscript{Gln}\textsubscript{mnmE17} lacking the cmnm\textsuperscript{5}-group also increases misreading of GGA (Gly), GAU (Asp) and GAC (Asp) codons [8].

**Overexpression of tRNA\textsubscript{Lys} having only the cmnm\textsuperscript{5}- or the s\textsuperscript{2}-group of cmnm\textsuperscript{5}s\textsuperscript{2}U34.**

Plasmid p815 contains the val\textsubscript{U} operon, which consists of the genes val\textsubscript{U}, val\textsubscript{X}, val\textsubscript{Y}, and lys\textsubscript{V}, the latter of which is the structural gene for tRNA\textsubscript{Lys}. Mutations mnmA3 or mnmE13 influence only the structure of the tRNA\textsubscript{Lys} and overexpression of these tRNAs, including tRNA\textsubscript{Lys}\textsubscript{mnmA3} and tRNA\textsubscript{Lys}\textsubscript{mnmE13} resulted in a severe growth reduction of strains lacking the s\textsuperscript{2}-group of cmnm\textsuperscript{5}s\textsuperscript{2}U34, suggesting that overexpressing thiol-deficient tRNA\textsubscript{Lys} may induce translational errors, such as missense errors (Table 3). However, Hagervall et al [7] showed that s\textsuperscript{2}-deficiency of cmnm\textsuperscript{5}s\textsuperscript{2}U34 in tRNA\textsubscript{Lys}\textsubscript{mnmA3} results in less missense errors in reading the AAU/C (Asn) codons. If our results reflect increased missense errors, they may be due to other such errors than those monitored by Hagervall et al [7]. If so, the missense errors induced by s\textsuperscript{2}-deficiency of cmnm\textsuperscript{5}s\textsuperscript{2}U34 in tRNA\textsubscript{Lys}\textsubscript{mnmA3} may be codon specific, which has been observed earlier [8]. However, such a growth reduction was not observed when the overexpressed tRNA\textsubscript{Lys}\textsubscript{mnmA3} was cmnm\textsuperscript{5}-deficient, suggesting that this deficiency did not induce any increased translational errors such as missense error. This result is indeed consistent with earlier reports showing that cmnm\textsuperscript{5}-deficient tRNA\textsubscript{Lys}\textsubscript{mnmA3} results in less missense errors in decoding AGA/G (Arg) or AAU/C (Asn) codons [7, 8]. These results strengthen our suggestion that the growth reduction observed by overexpressing hypomodified tRNAs may be correlated to increased missense errors.

The combined results above show that overexpressed s\textsuperscript{2}-deficient tRNA\textsubscript{Gln}\textsubscript{mnmA16} and tRNA\textsubscript{Lys}\textsubscript{mnmA3} imposed a considerable growth reduction suggesting that such hypomodified tRNAs increase missense error. Overexpression of tRNA\textsubscript{Gln}\textsubscript{mnmA16} lacking the cmnm\textsuperscript{5}-group reduced growth although much less than s\textsuperscript{2}-deficiency, whereas lack of the cmnm\textsuperscript{5} group of overexpressed tRNA\textsubscript{Lys}\textsubscript{mnmA3} did not.

**Presence of the wobble (c)cmnm\textsuperscript{5}s\textsuperscript{2}U34 in tRNAs specific for Gln, Lys, and Glu is essential for viability of S. enterica.**

To construct a double mutant mmmA, mmmE we wanted to combine strains harboring both the selectable mmmA16\textsuperscript{<}\textsuperscript{-}\textsuperscript{>cat (Cm\textsuperscript{R}) and mmmE17\textsuperscript{<}\textsuperscript{-}\textsuperscript{>Km (Km\textsuperscript{R}) mutations and such a double mutant should have tRNAs specific for Gln, Lys, and Glu with an unmodified wobble uridine (U34). We grew the mmmA16\textsuperscript{<}\textsuperscript{-}\textsuperscript{>cat mutant in rich medium at 37°C and mixed the cells with the transducing phage P22 grown on the mmmE17\textsuperscript{<}\textsuperscript{-}\textsuperscript{>Km. We then spread the mixture on rich agar plates containing kanamycin to select for the mmmE17\textsuperscript{<}\textsuperscript{-}\textsuperscript{>Km mutation and thereby creating the wanted double mutant mmmA16\textsuperscript{<}\textsuperscript{-}\textsuperscript{>cat, mmmE17\textsuperscript{<}\textsuperscript{-}\textsuperscript{>Km. The Km\textsuperscript{R} transductants were selected at several temperatures to monitor if a double mutant might be viable at another temperature than 37°C. The donor strain GT8176 (STM3453-2550::Tn10dTc) also contains a selectable (tetracycline resistant, Tc\textsuperscript{R}) mutation known not to induce any growth defect and this mutation was used to monitor the efficiency of transduction. As recipient we also used strain GT8173 (pmmmA\textsuperscript{+} mmmA16\textsuperscript{<}\textsuperscript{-}\textsuperscript{>cat), which contains the same mmmA16\textsuperscript{<}\textsuperscript{-}\textsuperscript{>cat mutation on the chromosome and the complementing plasmid pmmmA\textsuperscript{+} containing the wild type allele
of the mnmA gene. Such strain should behave as the wild type unless the mutation mnmA16<>cat introduces some unknown phenotype, which is not related to the mutation in mnmA gene and therefore not complemented by the pmmnA+ plasmid.

Table 4 shows that no double mutant mnmA16<>cat, mnmE17<>Km was obtained at any temperature clearly demonstrating that such a double mutant is not viable. This result cannot be due to an inefficient transduction, since the transfer of the STM3453-2550::Tn10dTc marker was similar to the transfer frequency of the mnmE17<>Km mutation to the wild type (Compare the number of TcR colonies to the number of KmR colonies in the wild type). Moreover, the transduction frequency of mnmE17<>Km was similar when strain GT8177 (pmmnA+/ mnmA16<>cat) was used as recipient. Note also, that we waited several days in order to allow a very slow growing double mutant to appear but still no such mutant was recovered. Moreover, we did not obtain any external suppressor mutant with mutation counteracting the lack of modification of U34. We conclude that cells having an unmodified wobble uridine of tRNAs specific for Gln, Lys and Glu are not viable on rich medium at temperatures between room temperature (about 20˚C) and 42˚C.

The inability to obtain any transductants at 15˚C when selecting TcR transductants (STM3453-2550::Tn10dTc) shows that the recipient mnmA16<>cat mutant is not able to grow at such low temperature consistent with the results shown in Table 2.

### Discussion

We [5] and others [32]) have shown that the major function of mcm5 s2U34 in yeast tRNAs is to improve the efficiency of the cognate anticodon-codon interaction rather than preventing missense errors, since overexpression of hypomodified tRNAs specific for Gln, Lys or Glu counteracts the phenotypes induced by deficiency of either s2, mcm5- or both modifications. However, overexpression of bacterial tRNA\textsuperscript{Gln}\textsubscript{mcm5s2UUG} did not counteract the slow growth

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**Table 4.** The double mutant mnmA, mnmE is nonviable on rich medium at 15 to 37˚C.

| Temp (˚C) | Donor (Relevant genotype) | Recipient (Relevant genotype) | Tc\textsuperscript{R} No. of colonies (days) | Km\textsuperscript{R} No. of colonies (days) |
|-----------|---------------------------|-------------------------------|-------------------------------------------|-------------------------------------------|
| 37        | GT8176(mnmE17<>Km, STM3453-2550::Tn10dTc) | GT7321(mnmA+mnmE\textsuperscript{+}) | 976 (3) | 691 (3) |
| 37        | " | GT8177 (pmmnA+/ mnmA16<>cat) | 463 (3) | 393 (3) |
| 37        | " | GT8173 (mnmA16<>cat) | 915 (3) | 0 (7); 2\textsuperscript{a} |
| 30        | " | GT7321(mnmA+mnmE\textsuperscript{+}) | 584 (3) | 308 (5) |
| 30        | " | GT8173 (mnmA16<>cat) | 628 (3) | 0 (29) |
| RT        | " | GT7321(mnmA+mnmE\textsuperscript{+}) | 521 (4) | 191 (14) |
| RT        | " | GT8173 (mnmA16<>cat) | 482 (6) | 0 (35) |
| 15        | " | GT7321(mnmA+mnmE\textsuperscript{+}) | 155 (10) | 28 (10) |
| 15        | " | GT8177 (pmmnA+/ mnmA16<>cat) | 224 (10) | 80 (10) |
| 15        | " | GT8173 (mnmA16<>cat) | 0 (44) | 0 (44) |

Recipient cells were grown in rich medium at 37˚C. Following transduction using phage P22 grown on indicated donor strains, selection was made on rich plates containing either Km (50 ug/ml) or Tc (20ug/ml). Selection for Tc\textsuperscript{R} colonies shows the efficiency of transduction. The plates were incubated at indicated temperatures for indicated days (in parenthesis) when the number of colonies was scored. Plasmid pmmnA\textsuperscript{+} is the pNTR-SD mnmA\textsuperscript{+} plasmid harbouring the wild type allele of the mnmA\textsuperscript{+} gene from E. coli[45].

\textsuperscript{a} Two small colonies were observed after seven days but they were not analysed further due to their instability and poor growth.

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caused by $s^2$- or $cmnm^5$- deficiency but rather exaggerated the reduced growth of these mutants (Table 3). This growth reduction is caused by the $s^2$- or $cmnm^5$- deficiency and not to some other cellular aberrations, since the same result was obtained by three independent mutants ($mmmA3$, $tusB27$ and $tusE30$) defective in the thiolation and two independent mutants ($mmmA13$ and $mmmG1$) defective in the synthesis of the side chain (Table 3). Moreover, this severe growth reduction by overexpression of tRNA$^{\text{Gln}}_{cmnm5,6s2UUG}$ is due to the surplus of this particular tRNA, since a spontaneous faster growing mutant derivative was obtained that have a plasmid derivative lacking genes $glnU/W$, which both encode this tRNA (Fig 5). These results suggest that the major impact of hypomodified tRNA$^{\text{Gln}}_{cmnm5,6s2UUG}$ caused increased translational errors, such as missense errors, rather than reduced efficiency of reading the cognate codons as was suggested for yeast tRNA lacking $mcm^5s^4U34$ [5]. If the function of (c)$mmm^5s^2U34$ were to improve the efficiency of cognate codon reading, increased concentration of hypomodified tRNA should rather counteract the reduced growth of the $mmmA3$ and $mmmA13$ mutants than to exaggerate it. Thus, although the $xm^5s^2U34$ wobble modifications are universally conserved, their functional impacts may be different, at least in yeast and in S. enterica. A difference in translation accuracy between yeast and E. coli has been noted earlier, since missense errors are about 10-fold lower in yeast compared to E. coli. Yeast might have evolved mechanism(s) not present in bacteria to reduce missense errors [33].

An unmodified wobble uridine is rarely present in any cytosolic tRNAs [2]. In organelles, like mitochondria, unmodified uridines are present in tRNAs reading family boxes but not in tRNAs specific for Gln, Lys, and Glu, which always have an $xm^5s^2U34$ derivative. It was thought that the presence of this modification in the wobble position of Gln, Lys and Glu tRNAs would prevent missense errors [4]. The phenotypes of the $mmmA3$ and $mmmA17$ mutants show that these modifications are pivotal for cellular growth (Table 2) and, indeed, a double mutant $mmmA16tusB27$, $mmmA17tusE30$, which has an unmodified U34 as wobble nucleoside, did not grow at several different temperatures (Table 4). These results are consistent with an earlier report that showed that a similar double mutant of E. coli is not able to grow [34]. Clearly, the functional impact of (c)$mmm^5s^2U34$ is of the outmost importance and explain why such a modification is always present in these kind of tRNAs and predicted to be present in a minimal translation apparatus [13]. Based on the results presented in this paper we suggest that the major reason for inability to support growth with an unmodified uridine in the wobble position of tRNAs specific for Gln, Lys, and Glu is increased missense errors.

It has been suggested that the $s^2$- and $xm^5$- groups of $xm^5s^2U$ derivatives present in the wobble position would prevent missense errors [4, 35]. This kind of modification is present in split codon boxes (Fig 2) and thus should prevent decoding the near-cognate pyrimidine ending codons in such boxes and thereby missense errors. Although the three tRNAs specific for Gln, Lys and Glu all have the $xm^5s^2U34$ wobble modification they still are part of distinct structural context. The tRNA$^{\text{Lys}}_{mmmA5s2U34}$ has a very flexible anticodon loop due to the presence of several U (a poor stacker) including U36, whereas Gln has G36 and Glu has C36. Moreover, tRNA$^{\text{Lys}}_{mmmA5s2U34}$ has the large hydrophobic cyclic N6-threonylcarbamoyladenosine (ctA) in position 37 and both tRNA$^{\text{Gln}}_{mmmA5s2U34}$ and tRNA$^{\text{Glu}}_{mmmA5s2U34}$ have the much smaller methylated derivative m$^2$A. The ctA improves the stacking interaction with the base above (A38) and below (A36) of the anticodon loop but it also makes a cross strand stack to the first base (A) of the codon and thereby stabilizes the weak A-U36 base pairing in the first position [36, 37]. This strong functional impact on the stability of U-rich anticodon loops may explain why this modification is essential for viability [38]. The functional impact of m$^2$A, which is present in tRNA$^{\text{Gln}}_{mmmA5s2U34}$ and tRNA$^{\text{Lys}}_{mmmA5s2U34}$, is likely to be different and should have a lesser functional impact than ctA and accordingly it is not essential for viability [39]. Thus, the structural
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context for these \textit{\textsuperscript{5}s}U wobble modifications is different in these three tRNAs not only because differences in the primary sequence of the anticodon loop but also because of differences in the modification pattern and in particular the nature of the modifications at position 37 (ctA in tRNA\textsuperscript{Gln} and m\textsuperscript{2}A in tRNA\textsuperscript{Glu} and tRNA\textsuperscript{Lys}, see Fig 2). These features may influence whether the functional impact by (c)mnm\textsuperscript{5}sU34 is primarily to prevent missense errors or to improve the efficiency of cognate reading. The frequency of missense errors also depends on the efficiency of the competing near-cognate tRNAs [40]. Such a consideration may explain that missense errors for hypomodified tRNA\textsuperscript{Gln} are decreased [7, 8] (Table 3), whereas they are increased for tRNA\textsuperscript{Glu} (Table 3) and tRNA\textsuperscript{Lys} [8]. Note also, that when a specific missense error is monitored it is codon dependent but if growth reduction caused by overexpression of a specific hypomodified tRNA monitors missense errors it will be the total outcome of many possible such errors of several near- and/or non-cognate codons. Therefore, one would not expect identical results using two such different methods to monitor translational errors. Still, in the case of the functional impact of the mnm\textsuperscript{5}-side chain on tRNA\textsuperscript{Lys} the two methods gave consistent results (Table 3 and [7, 8]).

We have suggested that the reduced growth linked to the overexpression of hypomodified tRNA\textsuperscript{Gln} or tRNA\textsuperscript{Lys} is mainly due to increased missense errors, although we cannot exclude other translational errors, such as errors in the reading frame maintenance, misfolding of nascent proteins, or ribosomal drop off. The latter probably does not explain the observed growth reduction, since overexpression of tRNA would counteract such errors. Deficiency of either s\textsuperscript{2}- or (c)mnm\textsuperscript{5}-group of (c)mnm\textsuperscript{5}sU34 induces frameshifts by the peptidylslippage mechanism (reviewed in [1]) and according to this model overexpression of the tRNA reading the next codon downstream of the frameshifting site decreases frameshifting. Still, Pande et al [41] noticed that overexpression of a few specific yeast tRNAs increases frameshifts probably by an out-of-frame binding, which pulls the ribosome into the shifted frame. The increase in frameshifting was modest and no major effect on growth was reported. Overexpression of hypomodified tRNA specific for Gln, Lys and Glu in yeast counteracts the phenotypes induced by modification deficiency and does not induce any growth defect [5, 32]. Moreover, as in bacteria overexpression of fully modified yeast tRNA does not cause any reduction in growth. Note also that one of the most efficient frameshift suppressors (sufA6) does not in any major way influence cellular growth (unpublished observation). Taken together, increased frameshifting as an explanation to the severe growth reduction observed upon overexpression of hypomodified tRNAs is not likely. On the other hand the growth reduction may cause an aberrant folding of several proteins. The nascent polypeptide is folded on the ribosome and a change of the polypeptide synthesis rate may influence the folding of proteins [42]. The speed with which the mRNA is decoded is correlated to the concentration of cognate tRNAs [43, 44]. Overexpression of an unmodified tRNA may change the rate of polypeptide synthesis and thereby be influencing the folding of the nascent peptides resulting in an accumulation of misfolded proteins. Such aberrant proteins may in turn induce a growth defect. Still, overexpression of fully modified tRNA did not influence the growth (Table 3), a condition which also should change the rate of polypeptide synthesis due to increased concentration of some tRNAs. Therefore, if the observed reduction of growth is caused by misfolding of proteins, it would be specific for modification deficiency. Overexpression of normal or hypomodified yeast tRNAs specific for Gln, Lys and Glu does not reduce cellular growth but rather counteracts the modification deficient induced phenotype(s). Thus in yeast overexpression of hypomodified tRNAs does not induce frameshifts or aberrant folding of proteins to such a degree that it influences growth, although both kinds of translational errors may well be...
operating also in this organism [5, 32]. From these considerations we favor missense errors causing the observed growth reduction, since theoretical considerations have suggested that the function of the $\text{xm}^5\text{s}^2\text{U34}$ is to prevent such errors. Moreover, such errors have been observed experimentally [8] although in some experiments reduced missense errors by hypomodified tRNA was noticed [7, 8]. These experiments monitored effects on specific codons why it is not excluded that misreading of other codons may be operating. Overexpression of $\text{tRNA}^{\text{Lys}}_{\text{mnm}^5\text{ssUU}}$ lacking the $\text{mnm}^5$-group did not reduce growth suggesting no increase missense errors consistent with results obtained earlier [7, 8]. Thus, monitoring growth may well be relevant to estimate missense errors although not as specific as a direct measure of missense errors. Even if we cannot rule out other translational errors, we still found it likely that the growth reduction observed is caused mainly by increased missense errors. This suggestion is in line with the fact that the $\text{xm}^5\text{s}^2\text{U34}$ wobble modification have been shown theoretically and experimentally to influence the accuracy of translation besides its effect on the efficiency of cognate codon reading. Therefore, the primary function of the $\text{xm}^5\text{s}^2\text{U34}$ modifications may be different in yeast and in bacteria.

Supporting information
S1 Table. Strains and plasmids.

(DOCX)

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