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Transfer RNA Bound to MnmH Protein Is Enriched with Geranylated tRNA – A Possible Intermediate in Its Selenation?

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Abstract

The wobble nucleoside 5-methylaminomethyl-2-thio-uridine (mnm⁵⁵²U) is present in bacterial tRNAs specific for Lys and Glu and 5-carboxymethylaminomethyl-2-thio-uridine (cmnm⁵⁵²U) in tRNA specific for Gln. The sulfur of (c)mnm⁵⁵²U may be exchanged by selenium (Se) – a reaction catalyzed by the selenophosphate-dependent tRNA 2-selenouridine synthase encoded by the mnmH (ybbB, selU, sufY) gene. The MnmH protein has a rhodanese domain containing one catalytic Cys (C97) and a P-loop domain containing a Walker A motif, which is a potential nucleotide binding site. We have earlier isolated a mutant of Salmonella enterica, serovar Typhimurium with an alteration in the rhodanese domain of the MnmH protein (G67E) mediating the formation of modified nucleosides having a geranyl (ge)-group (C₁₀H₁₇-fragment) attached to the s² group of mnm⁵⁵²U and of cmnm⁵⁵²U in tRNA. To further characterize the structural requirements to increase the geranylation activity, we here report the analysis of 39 independently isolated mutants catalyzing the formation of mnm⁵ges²U. All these mutants have amino acid substitutions in the rhodanese domain demonstrating that this domain is pivotal to increase the geranylation activity. The wild type form of MnmH⁺ also possesses geranyltransferase activity in vitro although only a small amount of the geranyl derivatives of (c)mnm⁵⁵²U is detected in vivo. The selenation activity in vivo has an absolute requirement for the catalytic Cys97 in the rhodanese domain whereas the geranylation activity does not. Clearly, MnmH has two distinct enzymatic activities for which the rhodanese domain is pivotal. An intact Walker motif in the P-loop domain is required for the geranylation activity implying that it is the binding site for geranylpyrophosphate (GePP), which is the donor molecule in vitro in the geranyltransfer reaction. Purified MnmH from wild type and from the MnmH(G67E) mutant have bound tRNA, which is enriched with geranylated tRNA. This in conjunction with earlier published data, suggests that this bound geranylated tRNA may be an intermediate in the selenation of the tRNA.
Introduction

Transfer RNA plays a pivotal role in translating the information residing in mRNA into proteins. The different species of tRNA are about 75–90 nucleotides in length and of all nucleic acids present in the cell, tRNA contains the highest proportion of modified nucleosides. These are derivatives of the four normal nucleosides adenosine (A), guanosine (G), uridine (U) and cytidine (C). At present more than 100 different modified nucleosides in tRNA from different organisms are characterized (http://mods.rna.albany.edu/ or http://modomics.genesilico.pl/). The synthesis of the modified nucleosides is catalyzed by highly specific enzymes using precursor tRNA molecules as substrates. Position 34 (the wobble nucleoside) and position 37 (the nucleoside adjacent and 3’ of the anticodon) in tRNAs from all organisms are not only frequently modified (61% and 75%, respectively) but a large variety of different modified nucleosides are also present in these two positions [1]. These modified nucleosides in tRNA are important for a proper and efficient reading of the mRNA and lack of some of them results in severe physiological defects and/or no viability (Reviewed in [1]). Although several of these modified nucleosides have different chemical structures, are present in different tRNAs, and at different positions in the tRNA, many of them are important for keeping the ribosome in the correct reading frame [2].

The modified nucleosides (c)mnm5s2U34 in bacteria and mcm5s2U34 in eukaryotes are present in the wobble position of Gln-, Lys- and Glu-tRNA and presence of these modifications in tRNA is essential for viability in yeast [3] and in bacteria (unpublished observation). Deficiency of (c)mnm5s2U as well as hypermodification of tRNA (see below) in bacteria induces error in the reading frame maintenance, which is phenotypically manifested by ability to suppress +1 frameshift mutations. Lack of either the side chain in position 5 or the sulfur at position 2 induces frameshifts as well as addition of a geranyl group (C10H17-fragment) to the sulfur at position 2 of (c)mnm5s2U does [2, 4]. This geranylation reaction is catalyzed by the MnmH protein and the geranylation at the s2-group is increased by a mutation in the mnmH gene (earlier denoted sufY, ybbB or selU) resulting in an amino acid substitution G67E of MnmH [4]. Such +1 frameshift suppressors were first isolated by Riddle and Roth [5, 6] and later by Jäger et al [7]. The first isolated mnmH mutations are genetically dominant mediating the addition of a geranylgroup to the sulfur of cmnm5s2U of tRNAGln [4]. This was later confirmed by Dumelin et al [8] and these authors further showed that a geranyl group is also added to mcm5s2U in tRNAlys5s2U34 and in tRNAGlu5s2U [8]. G67E alteration of the MnmH increases the intrinsic low geranylation activity of the wild type MnmH [4, 8]. Earlier the MnmH (SelU, SufY, YbbB) protein was also shown to replace the sulfur of (c)mnm5s2Uys by selenium [9]. Clearly, the MnmH protein has two distinct enzymatic activities—a geranylation and a selenation activity.

The synthesis of cmnm5s2U is complex and requires eleven different proteins of which two (MnmC and MnmH) have two enzymatic activities (Summarized in Fig 1; Reviewed recently in [1, 10]). The sulfur pathway depends on the IscS protein, which mobilizes the sulfur of cysteine which is then relayed to the wobble uridine in tRNA to form the (c)mnm5s2U [11]. The two pathways—synthesis of the (c)mnm5-side chain and the thiolation at position 2—occur independently of each other; i.e lack of one modification does not influence the synthesis of the other. At a certain level and depending on the availability of selenium and geranylpyrophosphate (GePP) a fraction of the (c)mnm5s2U is selenated or geranylated. Two alternative metabolic pathways may be envisioned—the MnmH either selenates or geranylates the s2-position of cmnm5s2U (Alternative I) or the geranylated (c)mnm5s2U (abbreviated (c)mnm5ges2U; “ge-” is the abbreviation used for the geranyl group) is an intermediate in the selenation reaction creating the final product (c)mnm5Se2U (Alternative II). In Alternative I the MnmH protein might
be a moonlighting protein provided that both these distinct activities resides in the same domain (one criteria for a moonlighting protein [12, 13]). In Alternative II the MnmH protein has two distinct activities operating in a linear biosynthetic pathway with the end product (c) mmn5Se2U. The MnmC protein also has two activities (an oxidase and a methyltransferase activity) operating in a linear biosynthetic pathway but these activities reside in two distinct domains.

The MnmH protein is composed of two domains—a rhodanese domain and a P-loop domain (Fig 2). Rhodaneses are ubiquitous proteins that catalyze the transfer of a sulfur atom.
from thiosulfate (rhodanese) or 3-mercaptopyruvate to e.g. cyanide in vitro (Reviewed in [14, 15]). The rhodaneses contain a conserved Cys (C97 in MnmH) that is absolutely required for their catalysis. In the single-domain rhodaneses this domain comprises the entire protein (e.g. GlpE) but such a domain can also be fused to other protein domains; e.g. the ThiI protein contains a rhodanese, a P-loop, and a THUMP domain and it catalyses both the biosynthesis of the thiazole ring of thiamine and the modified nucleoside 4-thiouridine (s4U) [16]. Whereas the two domains of bacterial MnmH are encoded by a single gene, the archaeal MnmH protein is encoded by two separated genes of which the gene encoding the P-loop is immediately upstream of the gene encoding the rhodanese domain [17]. Such a gene organization will result in the synthesis of two separate proteins and both of them are required to obtain selenation activity [17]. In both bacterial and archaeal orthologues of MnmH there is a conserved amino acid insertion of about 33 amino acids (36 amino acids in Salmonella enterica serovar Typhimurium, yellow in Fig 2), which is not present in single-domain rhodaneses like GlpE [9]. The rhodanese signature motifs (CH2A and CH2B), the conserved active center (C97-XX-G) in rhodaneses, and the P-loop are shown. Position G67 is also shown since the first mutant form characterized of MnmH has an amino acid substitution (G67E) at this position.

![Domain organization of the MnmH protein from S. enterica](Modified from [9]). MnmH consists of two domains—a rhodanese domain with its Cys97 in the active center (Red indicates amino acids which has been altered by a site specific alterations) and a Walker A motif GCTGSGKT (a Walker A motif is G-x(4)-GKT [18, 19]), which is a binding site for ATP and GTP. A conserved 36 amino acid insertion present in the rhodanese domain of MnmH orthologues is not present in single-domain rhodaneses like GlpE [9]. The rhodanese signature motifs (CH2A and CH2B), the conserved active center (C97-XX-G) in rhodaneses, and the P-loop are shown. Position G67 is also shown since the first mutant form characterized of MnmH has an amino acid substitution (G67E) at this position.

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Geranylation of tRNA

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Geranylation of tRNA
selenation activities [9]. One possibility would be that selenation and geranylation occur in parallel catalysed by MnmH (Alternative I, Fig 1). We further show that the purified protein from various mutant forms of MnmH induces a strong binding of the chaperon GroEL whereas the wild type version of MnmH binds GroEL much less. Both the wild type form and all mutant forms of MnmH have tRNA strongly bound to them. Interestingly, the enzyme bound tRNA is enriched with the geranylated form of tRNA suggesting a possible role of geranylated derivative of (c)mnm5s2U as being an intermediate in the replacement of the sulfur by selenium (Alternative II, Fig 1).

**Materials and Method**

**Bacteria and growth conditions**

The bacterial strains used were derivatives of *Salmonella enterica* serovar Typhimurium or *Escherichia coli* K12 (S1 Table). As rich medium Luria-Bertani (LB) was used [20] (C:\GetARef\Refs\Refsmanus\FS_mutant_2011.ref #2; C:\GetARef\Refs\Refsmanus\Kristina-Gln05.ref #38; The minimal solid medium was made from the basal medium C:\GetARef\Refs\Refsmanus\FS_mutant_2011.ref #4; Vogel & Bonner [21] (C:\GetARef\Refs\Refsmanus\Kristina-Gln05.ref #39; with 15g of agar per liter and supplemented with 0.2% glucose and required amino acids and/or vitamins [22] (C:\GetARef\Refs\Refsmanus\FS_mutant_2011.ref #5; C:\GetARef\Refs\Refsmanus\Kristina-Gln05.ref #40; 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*) [23] (C:\GetARef\Refs\Refsmanus\FS_mutant_2011.ref #20; was performed as previously described [22] (C:\GetARef\Refs\Refsmanus\Kristina-Gln05.ref #40; 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.

**Random mutagenization of the codon for G67 in MnmH**

Plasmid pUST210 (pCL1921 harbouring the *mnmH* gene and a spectinomycin resistant marker) was amplified with two primers where nucleotides 199–201 (encoding G67) were scrambled creating codons for different amino acids. The resulting plasmids were transformed into strain JM109 supercompetent cells (Promega) and plated on agar plates containing LB and spectinomycin (100 μg/ml). 300 transformants were collected and used for preparing a pool of plasmids which was thereafter introduced into strain GT6874 (∆*mnmH*, *hisO1242, hisC3737* (a +1 frameshift mutation in the *hisC* gene)). The transformation mixture was plated on medium E agar plates containing 0.2% glucose and spectinomycin (400 μg/ml) to select His⁺ Sp⁺ transformants. Thirty six such colonies were purified by restreaking and the *mnmH* gene on the plasmid was sequenced. To move the mutations to the chromosome a fragment with a tetracycline resistance marker was placed in the chromosomal located *mnmH* gene [24]. A fragment containing a mutation was amplified from the corresponding plasmid and this fragment was used in a linear transformation to replace the tetracycline resistant (Tc⁺) marker by selecting tetracycline sensitive (Tc⁻) clones [25, 26]. The mutated codon presence on the chromosome was confirmed by sequencing.
Mutagenesis to obtain independently isolated mutants with an altered MnmH protein

Mutagenesis of the strain GT5688 (insertion zbb-2523::Tn10dTc close to mnmH) was performed by inducing the expression of DinB [27, 28] from plasmid pSMP24 (Cb-R) [29]. GetARef\Refs\Refsmanus\Kristina-Gln05.ref #44; which harbor the dinB expressed from an ara-promoter\C:\GetARef\Refs\Refsmanus\Kristina-Gln05.ref #43; C:\GetARef\Refs\Refsmanus\Kristina-Gln05.ref #42; An over-night culture of strain GT5688 was diluted 2x106-fold and inoculated into 500 μl of LB+100 μg carbenicillin (Cb)/ml +0.08% L-arabinose. After 24 hours of growth at 37°C phage P22 was added to make a phage stock. The phage lysates were used to infect strain GT7321 (mnmH+, hisD10122) containing a +1 frameshift mutation in the hisD gene. Following selection for tetracycline resistant (TcR) colonies, His+ clones were selected among them. After a backcross the mnmH gene was sequenced to identify the mutation causing the His+ phenotype.

To mutagenize with hydroxylamine a phage P22 lysate prepared on strain GT5688 (insertion zbb-2523::Tn10dTc close to mnmH) was treated with hydroxylamine as described [30] C:\GetARef\Refs\Jocke\proM.ref #26; until approximately 0.1 per cent infectious phage P22 particles remained. This lysate was used to transduce strain GT7321 (hisD10122; a +1 frameshift mutation in the hisD gene) and following selection for TcR clones, His+ clones were selected.

Nitrosoguanidine (NG) induced mutations were obtained by placing a crystal of NG on a lawn of the donor strain GT5688 (insertion zbb-2523::Tn10dTc close to mnmH). Around the NG crystal a ring of growing bacteria emerged that contains bacteria with mutations induced by NG. Bacteria were scraped off from several fractions of the bacterial ring and suspended in 1 ml of LB and suspended cells were allowed to grow for 1–2 hours before phage P22 was added to make a lysate. As above the phage lysate was used to infect His strain harboring the +1 frameshift mutation hisD10122. TcR clones were selected and among them His+ clones were selected. To avoid siblings only one mutant from each phage stock was saved.

Amino acid alterations of Cys96, Cys97, or K155 of MnmH

To create amino acid alterations at defined positions, a fragment bearing the mutation was transformed into a strain with a Tn10dTc tetracycline resistance cassette in the mnmH gene. Tetracycline sensitive recombinants were selected on Bochner plates [25], purified, and tested for loss of TcR. The presence of the desired mutation was confirmed by sequencing of TcS clones. The alteration of K155 to A was done by placing a KmR cassette after the mnmH gene, and then amplifying a fragment with a primer harbouring the mutation as one primer and a primer binding in the cassette as the other primer. This fragment containing the mutation and KmR was then transformed into LT2 (wt) and GT1484 (mnmH204(G67E)), recombinants were selected on kanamycin plates and KmR clones were sequenced.

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 4x10^8 cells/ml. Cells were lysed and total RNA was prepared [31] and dissolved in 2 ml buffer R200 (10 mM Tris-HCl, 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 2.0 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. Transfer RNA
was digested to nucleosides by nuclease P1 followed by treatment with bacterial alkaline phosphatase at pH 8.3 [32]. The hydrolysate was analyzed as described earlier [33] using a Supelcosil C-18 column (Supelco) or Develosil C-30 (Phenomenex) with a Waters Alliance HPLC system. The degradation method used results in nucleosides except if a complex nucleoside is present which usually gives a dinucleotide as a result. The complex geranylated derivatives are therefore present in the hydrolysate as a dinucleotide (NpN). Since geranylation of tRNA is occurring only on (c)mnm5s2U containing tRNAs, i.e. Gln, Lys and Glu specific tRNAs, there will always be a U35 next to the geranylated (c)mnm5s2U34. Therefore, the degradation by enzyme P1 results in a dinucleotide of the type ges2UpU. The separation of the various geranylated derivatives are shown in S1 Fig. The various ges2-derivatives were identified by their characteristic spectra and by MS analysis of their molecular weights ([4] and data not shown).

Preparations of membranes
Membranes were prepared essentially as described by Wai et al [34] from 50 ml culture grown over night in LB Broth. The cells were washed with 10 mM Tris-HCl, pH 8.0, suspended in 1 ml of the same buffer and sonicated. The cell debris was pelleted by centrifugation at 5000 x g for 10 min and the resulting supernatant was centrifuged 30 min at 10000 x g. The sediment was used as the total membrane fraction. This fraction was separated into inner and outer membranes by adding 100 μl of 2% N-lauryl sarcosyl in 10 mM Tris buffer pH 8.0 and the mixture was incubated at room temperature for 30 min. After centrifugation for 30 min at 15000 x g the supernatant, which is the inner membrane fraction, was dialysed against 1000 ml 10 mM Tris-HCl, pH 8.0. The outer membrane in the pellet was suspended in 350 μl 10 mM Tris-HCl, pH 8.0 buffer and dialysed against the same buffer. Both outer and inner membrane fractions were extracted twice with phenol-chloroform-isoamylalcohol solution (25:24:1). The water phase, which contains RNA, was concentrated and loaded on a 12.5% urea PAGE gel in 0.5 M Tris-Borat-EDTA buffer. The gel was run for 1h at 200V and was blotted to Zetaprobe (Biorad) in a wet blot apparatus using the same buffer. As positive control total tRNA from strain GT8277 (ΔmnmH) was used. The tRNA was detected by Northern hybridization essentially as described earlier [35] using oligonucleotides labelled with [γ32P]-triphosphate (Perkin-Elmer). The oligonucleotides were complementary to nucleotides at positions 25–50 of tRNAGln cmnm5s2UUG and at positions 25–49 of tRNAPro cmnm5s2UGG. Northern blots were visualized by Phosphor-Imager analysis (S3 Fig).

Purification of FLAG-proteins
The Flag-tagged proteins, expressed in strain LMG194, were purified essentially as described by Moukadiri et al [36] except that the induced cells were grown over night at 30°C. The respective mnmH genes were amplified by PCR using FLAG-specific mnmH primers and cloned into the expression vector (pBAD TOPO TA) to generate the three FLAG-MnmH derivatives. The proteins were purified by standard procedure and routinely stored at -20°C in the presence of 15% glycerol.

Determining the modification pattern in tRNA bound to purified MnmH proteins
Fractions of purified protein were pooled, mixed with equal volume of phenol, and the nucleic acid was precipitated from the water phase by 2.5 volumes of ethanol with 1% KAc added. The precipitate was dissolved in water and once more the tRNA was extracted with phenol and the
tRNA was precipitated with 2.5 volume of ethanol, washed with 70% ethanol, dissolved in 100 μl MQ-water, degraded into nucleosides as described above, and the degraded products were analysed by HPLC. The modification pattern was compared to digested total tRNA from *E. coli*.

**Results**

The size and not the chemical features of amino acids at position 67 mediates increased geranylation activity of tRNA *in vivo*

Four independent +1 frameshift suppressor mutants were obtained earlier [5–7], which all had alterations at position 67 of MnmH located in the conserved 36 amino acid insertion in the rhodanese domain, which is found in many MnmH orthologues (Fig 2). Although the alterations were either a basic (Arg) or an acid amino acid (Glu), it still suggested that features at this position might be critical to obtain an increased level of geranylation of tRNA. To get more information about this aspect, we randomly scrambled the three nucleotides of codon 67 of *mmnH* on a plasmid. To analyze the geranylation activity of these altered MnmH proteins under the same condition as the original mutants, we transferred the mutated *mmnH* genes to the chromosome. We obtained nine new alterations at position 67 of MnmH inducing a +1 frameshift and concomitantly increasing the synthesis of (c)mnm5ges2U34 in tRNA (Table 1). The chemical properties of the amino acid side chain were all different (acid, basic hydrophobic, hydrophilic) but a common feature of the alterations was that the side chains have a larger van der Waals volume [37] than glycine, which is the amino acid in position 67 of the wild type MnmH (Table 1). Therefore, it appears that it is a space alteration in the vicinity of position 67 that is critical to generate an increased level of (c)mnm5ges2U34 in tRNA and thereby increases the endogenous weak activity of wild type MnmH to synthesize (c)mnm5ges2U34 in tRNA. We also considered that the 36 amino acid insertion might act as a negative element blocking an activity to

Table 1. Size and not the chemical feature of the amino acid side chain at position 67 is critical for increased geranylation activity of MnmH. ND, not detected. Relevant genotype: Besides indicated alterations of MnmH at position 67 encoded from the relevant *mmnH* gene located on the chromosome, all strains contain zbb-2558::Tn10dTc (close to *mmnH*), hisO1242, hisD10122 (CCC-CAA; +1 frameshift mutation), zdd-2532::Cat. Frameshifting (FS) of the indicated strains were monitored as growth of single cells on plates without histidine after 4 days at 37°C (+++, large colonies; ++, medium size colonies, +, small size colonies and (+), very small colonies). Wild type shows no growth under the same conditions. The frameshift phenotype is caused by shortage of Gln-tRNA\textsubscript{Gln}\textsubscript{cmnm5s2UUG} (cmnm5s2U) due to the formation of cmnm5gesU34 of this tRNA. This inhibition of the glutaminylation reaction causes lower levels of charged non-geranylated Gln-tRNA\textsubscript{Gln}\textsubscript{cmnm5s2UUG} which in turn induces the frameshifting activity [10]. Geranylated compounds were determined by HPLC analysis of tRNA from the indicated strains.

| Strain | aa at pos.67 | Chemistry of aa side chain | V | Total ges^2-derivatives (x10^3) | FS |
|--------|--------------|---------------------------|---|-------------------------------|----|
| GT7321 | G (wt)       | Hydrophobic               | 63.8| ND                           | -  |
| GT8216 | E            | Acidic                    | 140.8| 17 ± 8.2                     | +  |
| GT8187 | I            | Aliphatic/hydrophobic     | 164.9| 13 ± 6.7                     | +++|
| GT8188 | T            | Weak acidic               | 120.0| 19 ± 4.3                     | +++|
| GT8189 | L            | Aliphatic/hydrophobic.    | 164.6| 15 ± 1.5                     | ++ |
| GT8190 | R            | Strongly basic            | 192.8| 11 ± 0.9                     | ++ |
| GT8191 | N            | -                         | 127.5| 10 ± 3.2                     | (+) |
| GT8192 | C            | Hydrophobic/acidic        | 103.5| 16 ± 2.5                     | +  |
| GT8193 | H            | Aromatic/weak basic       | 159.3| 18 ± 6.7                     | +  |
| GT8194 | V            | Aliphatic/hydrophobic     | 139.1| 15 ± 1.4                     | +  |
| GT8195 | M            | Hydrophobic               | 167.7| 23 ± 1.0                     | +  |
generate (c)mnm5ges2U34. However, an MnmH protein with these 36 amino acids deleted, did not mediate any formation of (c)mnm5ges2U34 (data not shown).

Altering in or close to the conserved 36 amino acid insertion in the rhodanese domain increases the level of (c)mnm5ges2U34 in tRNA

We have isolated earlier one mutant with increased geranylation activity following a genome wide selection of frameshift suppressors [7], and three mutants were isolated 1970 as +1 frameshift suppressors by Riddle and Roth [5, 6] (Table 2). Thus, four independently isolated mutants have been characterized and all were located in the rhodanese domain suggesting that this domain is pivotal for increasing the geranylation activity. To further characterize the requirement for increased geranylation activity, we randomly mutagenized the region containing the mnmH gene and selected clones with increased +1 frameshift activity and accordingly an increased geranylation of tRNA\(^{Gln}_{\text{mmn5ges2U34}}\). (For details see Materials and Methods). Since the selection procedure used required enhanced ability to suppress a +1 frameshift mutation, we did not expect any mutants with a decreased geranylation activity. In total we characterized 39 independently isolated mnmH mutants, of which 35 were obtained by the above described localized mutagenesis. Of the latter, four were double mutants (two alterations at positions 69 and 66 in conjunction with alteration in position 67 and one (position 54) in conjunction with an alteration R79W). Thirty five of the 39 mutants had alterations in or in the vicinity of the conserved 36 amino acid insertion in the rhodanese domain (Fig 3). None of the mutants had

### Table 2. Characterization of 39 independent isolated mutants with increased frameshifting activity caused by increased geranylation activity following localized mutagenesis of the mnmH gene or by whole genome selection for increased frameshifting activity. A. Mutants were isolated following localized mutagenesis as described in M-M. All alterations in MnmH were obtained by the mutagen hydroxylamine except the alterations A63P and one of the two L66P alterations which were obtained using overexpression of dinB and the mutagen nitrosoguanine, respectively. In addition three mutants (One G67E and two G67R) obtained by Riddle and Roth [6] and one mutant (G67R) obtained by Jäger et al [7] are also included in the table. All these mutants have increased +1 frameshifting activity (data not shown), which is a caused by increased geranylation of tRNA. We did not observe a simple correlation between ability to suppress a +1 frameshift mutation to the level of cmnm5ges2UpU, which might not be expect, since induction of frameshifting is complex although partly dependent on a decreased level of charged Gln-tRNAGln\(^{cmnm5ges2UUG}\) [4]. B. Total ges\(^-\)derivatives consist of the level of cmnm5ges2U and mnm5ges2U determined as described in S1 Fig. Standard deviations are given as +/- . C. Not done. The level of the geranylation in these double mutants was not determined since they all had an alteration, which as a single alteration give increased geranylation activity. They all had increased frameshifting activity. D. Values are from Table 1. E. Following independent isolated mutants with G67R alteration in MnmH are present in strains GT945 and GT946 [6]. In addition, but not included in this table, are strains GT8190 (G67R) and GT8194 (G67V), which were obtained by scrambling the codon in position 67 (Table 1).

| Alterations in MnmH | No. of mutants | Total ges\(^-\)derivatives/ \(\Psi (x10^{-3})\) |
|---------------------|---------------|------------------------------------------|
| Wild (B8typeGT7321) | -             | <1                                       |
| G67Ed               | 2             | 17 ± 8.2                                 |
| R129C               | 2             | 9 ± 4.7                                  |
| G67E, R69Q          | 1             | Not done\(^{c}\)                         |
| R79W                | 21            | 16 ± 3.5                                 |
| A63P                | 1             | 15 ± 6.5                                 |
| L66P                | 2             | 17 ± 4.8                                 |
| A82T                | 1             | 14 ± 6.9                                 |
| R79W, C54C          | 1             | Not done\(^{c}\)                         |
| G67R, L66L          | 2             | Not done\(^{c}\)                         |
| G67R\(^{b}\)        | 5             | 11 ± 0.9                                 |
| G67V\(^{d}\)        | 1             | 15 ± 1.4                                 |

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Fig 3. Distribution of the 39 independent amino acid substitutions in MnmH that generate increased geranylation activity. Thirty-five (35) of these had amino acid substitutions in or in close proximity to the conserved 36 amino acid insertion in the rhodanese domain present in MnmH orthologues. X at position 67 indicates the various alterations were obtained in this position as shown in Table 1.

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an alteration in the P-loop domain, suggesting that potential alterations in this domain are not able to increase the geranylation activity. The results suggest that important features to increase the geranylation activity reside in the rhodanese domain.

Of the amino acid substitutions obtained in MnmH increasing the level of geranylation of tRNA, three were in 100% conserved positions (A63, G67 and R129), one in a 86% conserved position (L66), one in a 71% conserved position (R79), and one in a 50% conserved position [9]. Thus, alterations that increase the geranylation activity were all in the conserved or semi-conserved positions of the rhodanese domain.

High level of geranylated $tRNA^{Gln}_{cmnm5s2UUU}$, $tRNA^{Lys}_{mmn5s2UUU}$, and $tRNA^{Glu}_{mmn5s2UC}$ does not grossly affect cellular growth

The three tRNAs having the (c)mnm5s2U derivatives are about 50–70% geranylated in a strain having the MnmH(G67E) amino acid substitution (Table 3). One would then expect a reduction in growth unless the geranylated tRNA is able to efficiently decode the message. The charging level of $tRNA^{Gln}_{cmnm5s2UUU}$ is reduced to 26% as compared to the wild type level of 86% [4], a reduction that also results in increased frameshifting [4]. However, the level of charging of $tRNA^{Lys}_{mmn5s2UUU}$ and $tRNA^{Glu}_{mmn5s2UC}$ is not affected [8]. Still, the geranylation of $tRNA^{Glu}_{mmn5s2UC}$ induces +1 and -1 frameshifting and the efficiency of reading the glutamic acid codon GAA is reduced by 50% due to the geranylation of $tRNA^{Glu}_{mmn5s2UC}$ [8]. All these observed phenotypes would suggest that the geranylation may reduce the growth. We therefore tested the influence on growth of three strains [GT8277 $(\Delta mnmH)$, GT5688 $(mnmH^+)$], and strain GT5687 $(mnmH204 (G67E))$ with no, low level, and high level of geranylated tRNA, respectively. However, there was no difference in growth, measured as size of single cell colonies, between these strains on three different growth media (Rich medium, glucose and acetate minimal medium) and at three different temperatures (30, 37 and 41°C). Also, no difference in the specific growth rate constant was observed between these strains in rich liquid medium at 37°C (S2 Fig).

Table 3. The altered MnmH (G67E) mediates high level of geranylation of the tRNAs specific for Gln, Lys and Glu. A. The indicated levels of geranylated derivatives were calculated by assuming that the reduced level of (c)mnm5s2U has been converted to the corresponding geranylated derivative. B. Analysis of the purified species $tRNA^{Gln}_{cmnm5s2UUU}$. C. HPLC analysis of bulk tRNA from strain GT6680 $(\Delta mnmH)$ revealed no geranylated tRNA (unpublished observation). D. The level of geranylated derivatives from an E. coli $(\Delta mnmH)$ strain containing a plasmid harboring an mnmH (G67E) allele.

| tRNA $^{\text{gln}}_{\text{cmnm5s2UUG}}$ b) | % ges$^2$-derivatives of total (c) mnm5s2U $^a$ | % ges$^2$-derivatives of total (c) mnm5s2U $^a$ | % total ges$^2$-derivatives of total (c) mnm5s2U $^a$ |
|------------------------------------------|---------------------------------|---------------------------------|---------------------------------|
| $\Delta mnmH$                           |                                 |                                 |                                 |
| tRNA $^{\text{gln}}_{\text{cmnm5s2UUG}}$ b) | < 3                             | Not done                        | 70                              |
| In tRNA Gln, Lys, and Glu.              | 8                               | Not detected $^c$               | 50                              |
| In tRNA Gln, Lys and Glu.              | 3–7                            | Not detected $^c$               | 45 $^d$                         |

Table 3. The altered MnmH (G67E) mediates high level of geranylation of the tRNAs specific for Gln, Lys and Glu. A. The indicated levels of geranylated derivatives were calculated by assuming that the reduced level of (c)mnm5s2U has been converted to the corresponding geranylated derivative. B. Analysis of the purified species $tRNA^{Gln}_{cmnm5s2UUU}$. C. HPLC analysis of bulk tRNA from strain GT6680 $(\Delta mnmH)$ revealed no geranylated tRNA (unpublished observation). D. The level of geranylated derivatives from an E. coli $(\Delta mnmH)$ strain containing a plasmid harboring an mnmH (G67E) allele.

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The hydrophobic geranyl group does not cause an enrichment of geranylated tRNA in the outer or inner membranes

The envelope of a gram negative bacteria consists of an inner and an outer membrane with a peptidoglycan cell wall and a periplasmic space in between. Whereas the outer membrane consists of lipopolysaccharides and proteins the inner membrane consists of a bilayer with a hydrophobic layer of fatty acids. A human tRNA$^{Sec}$ containing the prenylated modified nucleoside 6-isopentenyl-adenosine (i6A) has an anticodon loop that is highly hydrophobic. Such
tRNA is retained in the hydrophobic HeLa cell membrane demonstrating that some tRNAs may be located in another cellular compartment than the cytoplasm [38]. The prenylated geranyl group of (c)mnm\textsubscript{5}ges\textsubscript{2}U, which is also highly hydrophobic, might mediate insertion of a geranylated tRNA into the hydrophobic lipid bilayer similar to the insertion of tRNA\textsubscript{Sec} into the human membrane. Thus, geranylation of tRNA might thereby reduce the concentration of such tRNA in the cytoplasm and hence reduce the amount of ternary complex available for translation. Shortage of ternary complex may induce frameshifting by inducing a pause of ribosome waiting for the ternary complex to bind to the ribosomal A-site and such an event will induce a frameshifting event by the peptidyl-tRNA \cite{2, 7, 39} as is observed for geranylated Gln-tRNA \cite{4} and geranylated Lys-tRNA \cite{8}. Moreover, such shortage of the ternary complex may also reduce the rate of translation as was observed for GAA(Glu) codon \cite{8}. Thus, it was of interest to investigate if the hydrophobic geranyl group would mediate an enrichment of geranylated tRNA to the membranes and, if so, explain some of the above mentioned phenotypes. We therefore prepared outer and inner membranes and monitored the presence of tRNA\textsubscript{Gln}\textsubscript{cmnm\textsubscript{5}ges\textsubscript{2}UUG}, which potentially might be geranylated, and tRNA\textsubscript{Pro}\textsubscript{GGG}, which is not a substrate for geranylation, in the membrane fractions. We used \textsuperscript{32}P-labelled oligonucleotides specific for these tRNAs as probes in a Northern blot analysis of membranes of three strains (GT8277 (\textit{ΔmnmH}), GT5688 (\textit{mnmH\textsuperscript{+}}), and strain GT5687 (\textit{mnmH204(G67E)}) with no, low level, and high level of geranylated tRNA, respectively. No positive signal for tRNA\textsubscript{Gln}\textsubscript{cmnm\textsubscript{5}ges\textsubscript{2}UUG} or for tRNA\textsubscript{Pro}\textsubscript{GGG} was observed in the outer membrane fraction, whereas in the inner membrane fraction signals for tRNA\textsubscript{Gln}\textsubscript{cmnm\textsubscript{5}ges\textsubscript{2}UUG} and tRNA\textsubscript{Pro}\textsubscript{GGG} were observed in all three strains (S3 Fig). Thus, apparently at least these two tRNAs were associated with the inner membrane. However, the signal for the presence of tRNA\textsubscript{Gln}\textsubscript{cmnm\textsubscript{5}ges\textsubscript{2}UUG} was similar in these three strains demonstrating that geranylation of the tRNA did not enrich the association of this tRNA with the inner membrane. We conclude that geranylation of tRNA does not induce an enrichment of tRNA associated with the membranes.

An intact Walker motif in the P-loop domain, but not an intact catalytic cysteine in the rhodanese domain, is required for the geranylation activity

As pointed out above, the rhodanese domain is critical for the enhancement of the geranylation activity, since all alterations of MnmH obtaining this feature resided in this domain. Therefore, the catalytic cysteine (C97) in this domain might be required for this activity in a way similar to the importance of it in selenation \cite{9}. The functionality of C97 in geranylation was tested by constructing amino acid substitutions C97A and C97S. Although the non-conserved cysteine at position 96 is not required for selenation \cite{9}, we also tested its relevance for the geranylation reaction due to its close location to the catalytic cysteine in the rhodanese domain. The geranyl group is transferred \textit{in vitro} from geranylpyrophosphate (GePP) by MnmH to the tRNA (see below). The P-loop domain contains a Walker motif (GCTGSK\textsubscript{155}T, Fig 2), which is a potential nucleotide binding site and therefore a potential binding site for GePP. We therefore tested the relevance of K155 in this sequence for geranylation by altering K155 to A155. These alterations were combined with the G67E amino acid substitution in MnmH, which mediates increased geranylation activity as compared to wild type MnmH. Analysis of mutants having the corresponding mutations on the chromosome demonstrated that the Walker motif in the P-loop domain must be intact to obtain geranylation activity whereas altering C97 or C96 reduced the level of geranylation but did not abolish it (Table 4). To increase the geranylation activity we placed the \textit{mnmH} gene and its derivatives on a plasmid. In this case the K155A alteration still reduced substantially the activity of the MnmH\textsuperscript{+} as well as of the MnmH(G67E) derivative (Table 4). The alterations of
C97 (either to an Ala or to a Ser) or C96S reduced the activity but much less than the K155A amino acid substitution. Thus, the C97 or C96 is not absolutely required for geranylation although a structural change in the neighborhood of them caused by changing the Cys to either Ser or Ala reduced the activity. We conclude that the catalytic cysteine in the rhodanese domain is not required for geranylation in sharp contrast to the absolute requirement of C97 for selena- 

The accumulation of geranylated derivatives in vivo is partly sensitive to the presence or absence of the (c)mnm5-side chain of the target nucleoside (c)mnm5s2U34

The first step in the synthesis of the cmnm5-side chain is catalyzed by the MnmEG complex. In the presence of Gly the cmnm5-side chain is first synthesized and this side chain is thereafter cleaved by the MnmC(ø) activity to form the nm5-side chain. When instead ammonium ion is present the MnmEG complex synthesizes directly the nm5-side chain. To generate the cmnm5-side chain the nm5-side chain formed by either of these reactions is methylated by MnmC(m) (Fig 1). In the wild type, the cmnm5-side chain is present only in the tRNAcmnm5s2UUG, whereas the tRNAcmnm5s2UUC and tRNAcmnm5s2UUC contain the cmnm5-side chain. In the presence of Gly, deletion of the mnmC gene (ΔmnmC) results in the accumulation of the cmnm5-side chain in the Gln-, Lys, and Glu-tRNAs. Deleting the mnmE gene results in no synthesis of a side chain and s2U accumulates in all these three tRNAs. To test the influence of the (c)mnm5-group on the geranylation activity, we used strains with the mnmH204 (G67E) mutation in conjunction with either ΔmnmE or ΔmnmC deletion and monitored thereby the level of geranylation of tRNA with no side or the cmnm5-side chain, respectively, at position 5 of the target nucleoside (c)mnm5s2U. Table 5 shows that in all strains with different side chains at position 5 of the target nucleoside, the level of ges2U-derivatives were reduced by about 50% suggesting that the geranylation activity is only partly sensitive to the length or absence of the (c)mnm5-side chain.

Purified MnmH(G67E) and MnmH(G67E, K155A) but not MnmH+ bind the chaperon GroEL

To characterize the geranylation further, we purified the FLAG-tagged MnmH proteins expressed from three different plasmids–the wild type MnmH+, the MnmH(G67E) mutant,
and the double mutant MnmH(G67E, K155A). Fig 4 shows that all preparations contain, besides the MnmH proteins, GroEL and tRNA (the GroEL was identified by N-terminal amino acid sequencing). The mutant forms contain substantial amount of GroEL whereas only a weak band of GroEL was observed in the wild type preparation. GroEL is a chaperon suggesting that the aberrant proteins MnmH(G67E) and MnmH(G67E, K155A) require this chaperon to be folded properly. Presence of such aberrant proteins might also induce synthesis of GroEL but this was not the case, since the level of GroEL as monitored by GroEL specific antibodies was the same in the strain overproducing these proteins as in wild type strain without any plasmid (data not shown). We also tried to purify other mutant forms of MnmH but such preparations had degraded MnmH and very low amount of intact peptide in conjunction with large amount of GroEL (data not shown). We judged that these preparations were not suitable to test the in vitro geranylation activity.

The purified wild type MnmH and the MnmH(G67E) mutants but not the double mutant MnmH(G67E K155A) transfer in vitro a geranyl group from geranylpyrophosphate to tRNA

To characterize further the geranylation activity of some of the altered MnmH, we tested purified MnmH proteins for their geranylation activity in vitro. Transfer RNA was purified from strain GT6680 (ΔmnmH), since such tRNA does not contain any (c)mnm5ges2U derivatives and such ges2-deficient tRNA should be optimal as an acceptor for a potential transfer of a geranyl (ge) group from the geranyl group donor GePP to tRNA. Since an intact P-loop is required in vivo to generate (c)mnm5ges2U34, the double mutant (G67E, K155A) should not possess any geranylation activity. We also expected no or only weak activity for the wild type form of MnmH and very low amount of intact peptide in conjunction with large amount of GroEL (data not shown). We judged that these preparations were not suitable to test the in vitro geranylation activity.
no geranylated tRNA at zero time. Apparently, the tRNA bound to this enzyme (Fig 4) is not geranylated.

The purified MnmH proteins have a bound tRNA population that is enriched by the geranylated form of the tRNA

In the wild type MnmH preparation a strong band of tRNA (Fig 4) was observed consistent with the observation by Wolfe et al. [9], C:\GetARef\Refs\Refsmanus\MnmH-2014.ref #15; who estimated that two moles of tRNAs were bound per mole of wild type MnmH\textsuperscript{+} [9]. Transfer RNA was also bound to the MnmH(G67E) and MnmH (G67E,K155A) mutant forms of MnmH (Fig 4). As mentioned above we also noticed that monitoring geranylation activity in vitro, samples taken at zero time of incubation had a substantial level of geranylated tRNA (see Fig 5). This might indicate that the purified enzymes had geranylated tRNA bound to it. To test this, we prepared tRNA from enzyme preparation of wild type MnmH\textsuperscript{+} as well as from some of the mutant forms of MnmH and analyzed the modification pattern of the enzyme bound tRNA. Modified nucleosides (Cm, m\textsuperscript{1}G, and ms\textsuperscript{2}i\textsuperscript{6}A) not present in tRNAs specific for Gln-, Lys, or Glu-tRNA, which are the only substrates for geranylation, were underrepresented...
compared to bulk tRNA (indicated by arrows in Fig 6). On the other hand modified nucleosides present in the Gln-Lys- and Glu- tRNAs (t6A, m2A, (c)mnm5ges2U) were overrepresented. Similar results were obtained when tRNA bound to MnmH(G67E) enzyme was analyzed (data not shown). We conclude that tRNAs bound to the enzymes were enriched with the geranylated derivatives. Apparently, the MnmH proteins have strong affinity for such tRNA (Fig 6).

Discussion

The first alteration in MnmH was discovered in a mutant mediating the ability to suppress +1 frameshift mutation and this induced phenotype is due to geranylation of tRNA_{Gln}^{GlnGluGluGlu} [4] alteration is within the 36 amino acid insertion in the rhodanese domain conserved among bacterial and archaeal MnmH orthologues (Fig 2). To obtain more information what other changes in the MnmH protein would induce a similar phenotype we isolated several independent mutants with increased suppressor activity and accordingly an increased level of geranylation of the tRNA. Interestingly, the alterations in position 67, in which the first isolated mutants were altered, suggest that it is not the chemical features but the size of the amino acid that is important for the induced phenotype (Table 1). However, isolating randomly induced mutations in the mnmH gene we found alterations outside this conserved 36 amino acid insertion but they were all in the rhodanese domain (Table 2, Fig 3). Still, the alterations were in the vicinity of this 36 amino acid insertion and all were close to the region where the catalytic Cys97 of the rhodanese domain is located. Thus, the rhodanese domain is important for the geranylation activity. This is also the case for the selenation activity, since the latter has an

Fig 5. In vitro geranylation of ΔmnmH tRNA, which is devoid of geranylated (c)mnm5s2U derivatives. MnmH (G67E) (dashed line), MnmH+ (solid line) and double mutant MnmH (G67E, K155A) (narrow dashed line) proteins were used as enzyme source. Corrected for the presence of GroEL in the different enzyme preparations, the specific activities (min⁻¹, μg MnmH protein⁻¹) were 0.12 x 10⁻³ (wild type), 0.40 x 10⁻³ (G67E), and 0 (G67E, K155A). The reaction was stopped at indicated times by phenol extraction and the geranylated compounds in the reaction mixture were determined by HPLC after digestion with P1 as described in Material and Methods. Note, that the zero time points for MnmH (G67E) and MnmH+ already show the presence of geranylated compound(s) whereas this is not the case for the double mutant MnmH (G67E, K155A).

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absolute requirement for an intact Cys97 whereas the geranylation activity does not (Table 3). The two activities are mechanistically different although the features of the rhodanese domain are important for both. We also show that the Walker motif in the P-loop domain is required for geranylation implying that this sequence might be important for the binding of GePP and this would argue against that MnmH is a moonlighting protein. Even though this sequence is outside the rhodanese domain the importance of an intact P-loop domain points towards a role in the geranylation reaction may be through binding of GePP. Wolfe et al [9] suggested that the P-loop domain binds tRNA and thus is required for both activities. This suggestion is supported by analysis of a MnmH orthologue in Archaea. In these organisms the two domains of MnmH orthologue are encoded by two genes of which the P-loop domain is encoded by a gene upstream of the gene encoded the rhodanese domain. Still, both half of the MnmH protein have to be present to obtain selenation activity and tRNA binding [17].

The MnmH (G67E) mutant was discovered as being +1 frameshift suppressor [4]. Since this alteration of MnmH also resulted in reduced charging of tRNAGln\textsubscript{cmnm5s2UUG} this suppressor phenotype is consistent with the "peptidyl slippage model" [7, 38]. This model suggests that a shortage of an aminoacylated tRNA reduces the rate of entry to the ribosomal A-site by the corresponding ternary complex (aa-tRNA·GTP·elongation factor Tu (EF-Tu)). Such a shortage will induce a ribosomal pause and thereby increases the probability of a slippage one nucleotide forward by the peptidyl-tRNA. Thus, the fact that geranylation of the tRNA\textsubscript{Gln\textsubscript{cmnm5s2UUG}} reduces the charging of tRNA\textsubscript{Gln\textsubscript{cmnm5s2UUG}} explains the frameshift suppressor phenotype of the
MnmH (G67E) mutant [4]. According to Dumelin et al [8] geranylation of tRNA^{lys}_{mnm5ges2UUU} is fully charged but still induces +1 frameshift errors, which is not consistent with the proposed peptidyl-tRNA slippage model. However, a possible explanation of this frameshifting phenotype according to the peptidyl-tRNA slippage model, would be that a fraction of the tRNA^{lys}_{mnm5ges2UUU} is not available for translation. If so, it would decrease the concentration of active ternary complex containing Lys-tRNA^{lys}_{mnm5ges2UUU} and thereby inducing a ribosomal pause and thus frameshifting according to this model. One way to reduce the available concentration of tRNA^{lys}_{mnm5ges2UUU} for translation would be that the geranylated tRNA is present in another compartment than the cytoplasm where translation occurs. We therefore tested the possibility that the geranylated tRNA was associated with the hydrophobic membrane. However, we did not find any evidence for an enrichment of geranylated tRNA in the membrane ruling out this possibility to explain the frameshifting phenotype at lysine codons. Another model to explain the frameshifting phenotype induced by the geranylated and fully charged tRNA^{lys}_{mnm5ges2UUU} would be that the ternary complex containing geranylated lys-tRNA^{lys}_{mnm5ges2UUU} has a reduced decoding activity of the lysine codon AAA and thereby inducing a peptidyl-tRNA slippage. Indeed, geranylated 2-thiothymidine base pairs poorly to A but fairly well to G compared to the non geranylated derivative [40]. Accordingly, the translation rate of Glu codon GAA is more reduced by geranylation of tRNA^{glu}_{mnm5ges2UUC} than its reading of Glu codon GAG [8]. Therefore, geranylation of tRNA^{lys}_{mnm5ges2UUU} may also cause a slow decoding of AAA in the A-site, which would in turn induce a frameshift event according to the peptidyl slippage model. Alternatively, the ternary complex containing the geranylated tRNA^{lys}_{mnm5ges2UUU} is inactive and this will reduce the concentration of a functional ternary complex containing non-geranylated tRNA^{lys}_{mnm5ges2UUU} and thereby reducing the reading of the AAA codon. This will induce a pause and thereby a frameshifting event in the P-site. In summary, there are several ways to explain the frameshifting activity induced by fully charged geranylated tRNA^{lys}_{mnm5ges2UUU} to be consistent with the peptidyl-tRNA slippage model. Still, a different mechanistic explanation not in line with the peptidyl-tRNA slippage model cannot be ruled out.

Chemically synthesized mnm^{5}ges^{2}U serves as a starting molecule to generate the selenium derivative mnm^{5}Se^{2}U by adding only sodium hydroselenide (NaSeH) in an ethanol solution [41]. Thus, a spontaneous chemical conversion occurs to generate mnm^{5}Se^{2}U from mnm^{5}ges^{2}U. This would suggest that a linear conversion of mnm^{5}s^{2}U → mnm^{5}ges^{2}U → mnm^{5}Se^{2}U (Fig 1, alternative II) is operating to replace sulfur of mnm^{5}s^{2}U by selenium and that mnm^{5}ges^{2}U serves as an intermediate in the selenation reaction. If a linear reaction mechanism is operating (Fig 1, Alternative II) inhibition of the selenation reaction would result in an increased level of the potential intermediate geranylated tRNA. In a selD mutant selenation is blocked due to the lack of the selenium donor (SePO_{4}), still the level of geranylation is about the same as in a selD^{+} strain ([4], data not shown). Moreover, a purified MnmH protein catalyzes the formation of the selenium derivative mnm^{5}Se^{2}U in vitro, without the presence of geranylpyrophosphate (GePP), which is the normal donor of a geranyl group [9]. These results would argue against any involvement of a geranylated intermediate in the selenation reaction. However, the selenium concentration in the growth medium might be very low, why a block in the synthesis of the selenium donor SePO_{4} by deleting the selD gene may not in a significant way further reduce the concentration of the selenium donor. This may explain why we did not observe any change in geranylation caused by lack of SelD even if a linear reaction mechanism is operating (Fig 1, Alternative II). Moreover, as shown in Fig 6 purified MnmH enzyme is enriched with geranylated tRNA, why such tRNA was likely present also in the purified enzyme preparations used by Wolfe et al [9]. Therefore, the geranylated tRNA bound to their purified
MnmH enzymes may serve as an intermediate for selenation if a linear process according to Fig 1 (Alternative II) is operating explaining why GePP is not required in their selenation in vitro [9]. Wolfe et al [9] also added tRNA prepared from a ΔmnmH strain and such tRNA contains no geranylated or selenated derivative. Omitting such tRNA in the reaction mixture, the selenation is reduced by 52% as compared when it is present and increasing the amount of ΔmnmH tRNA also increases the extent of selenation. Thus, in some way ΔmnmH tRNA added to the reaction mixture stimulates selenation although no GePP was added [9]. This would suggest a more complex mechanism than a linear process as outlined in Fig 1 alternative II. However, since the MnmH protein has a strong affinity for geranylated tRNA (Fig 6), the purified MnmH enzyme used in selenation in vitro may also contain the substrate GePP, which would allow geranylation of the added ΔmnmH tRNA which then serves as an intermediate in selenation. The chemical mechanism of geranylation and selenation is likely to be different. Still, the rhodanese domain is important for both reactions, suggesting that although mechanistically different, both reactions may occur on this domain perhaps without the intermediate geranylated tRNA leaving the enzyme. These results are consistent with a geranylated tRNA as intermediate in selenation. The generation of (c)mnm5ges2U34 may result in a leaving group, which facilitates the exchange of the ges2-group by selenium. Clearly, more detailed studies are required to elucidate the details of the mechanism of selenation.

Supporting Information
S1 Table. Strains and plasmids.
(DOCX)

S1 Fig. HPLC.
(EPS)

S2 Fig. Growth.
(EPS)

S3 Fig. Northern.
(EPS)

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Author Contributions
Conceived and designed the experiments: GRB. Performed the experiments: GJ PC. Analyzed the data: GJ PC. Contributed reagents/materials/analysis tools: GRB GJ. Wrote the paper: GRB GJ.

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