Role of Pseudouridine Formation by Deg1 for Functionality of Two Glutamine Isoacceptor tRNAs

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Abstract: Loss of Deg1/Pus3 and concomitant elimination of pseudouridine in tRNA at positions 38 and 39 (ψ38/39) was shown to specifically impair the function of tRNA\textsuperscript{Gln_{UG}} under conditions of temperature-induced down-regulation of wobble uridine thiolation in budding yeast and is linked to intellectual disability in humans. To further characterize the differential importance of the frequent ψ38/39 modification for tRNAs in yeast, we analyzed the in vivo function of non-sense suppressor tRNAs \textsuperscript{SUP4} and \textsuperscript{sup70-65} in the absence of the modifier. In the tRNA\textsuperscript{Gln_{A}} variant \textsuperscript{SUP4}, UAA read-through is enabled due to an anticodon mutation (UψA), whereas \textsuperscript{sup70-65} is a mutant form of tRNA\textsuperscript{Gln_{CUG}} (SUP70) that mediates UAG decoding due to a mutation of the anticodon-loop closing base pair (G31:C39 to A31:C39). While \textsuperscript{SUP4} function is unaltered in deg1/pus3 mutants, the ability of \textsuperscript{sup70-65} to mediate non-sense suppression and to complement a genomic deletion of the essential SUP70 gene is severely compromised. These results and the differential suppression of growth defects in deg1 mutants by multi-copy SUP70 or tQ(UUG) are consistent with the interpretation that ψ38 is most important for tRNA\textsuperscript{Gln_{UG}} function under heat stress but becomes crucial for tRNA\textsuperscript{Gln_{CUG}} as well when the anticodon loop is destabilized by the \textsuperscript{sup70-65} mutation. Thus, ψ38/39 may protect the anticodon loop configuration from disturbances by loss of other modifications or base changes.

Keywords: tRNA modification; PUS3/DEG1; translation; 5-methoxycarbonylmethyl-2-thiouridine; non-sense suppression; sup70-65

1. Introduction

In addition to standard nucleosides, tRNA is known to contain a large variety of modified nucleosides, formed by the addition of chemical groups or isomerization. Pseudouridine (ψ) is an isomer of uridine (5-ribosyl-uracil) and represents the initially discovered and most abundant modified nucleoside present in all three domains of life [1,2]. The model organism \textit{Saccharomyces cerevisiae} has ten characterized pseudouridine synthases (Pus1–9 and Cbf5) that modify cytoplasmic and mitochondrial tRNAs and/or other types of RNA, including tRNA, small nuclear RNA (snRNA) and mRNA [2]. Despite its ubiquitous presence in tRNA, ψ appears to affect tRNA function rather subtly, since it is not generally required for cell viability, as demonstrated by the non-essential nature of all of the PUS genes in the yeast system [3]. However, the absence of PUS3/DEG1 in yeast uniquely causes slow growth, in particular at elevated temperature [4]. The general effects of ψ on tRNA function are likely mediated by increasing the rigidity of the sugar phosphate backbone and base stacking [2,5]. In humans, homozygous mutation of the DEG1 orthologue PUS3 is correlated with intellectual disability, suggesting a significant contribution to tRNA functioning not only in yeast but also in humans [6].

In yeast, Deg1 dependent formation of ψ occurs in a majority of tRNAs either at position 38 or 39 [4,7]. Slow growth of yeast mutants lacking Deg1 is strongly aggravated if parts of the...
wobble uridine modification 5-methoxy-carbonylmethyl-2-thiouridine (mcm\(^{5}\)s\(^{2}\)U) are simultaneously absent [8,9]. The mcm\(^{5}\) component of the mcm\(^{5}\)s\(^{2}\)U modification is formed by a pathway requiring the Elongator complex and a variety of accessory and potentially regulatory proteins [10–19]. The separate Urm1 pathway mediates the transfer of sulfur to the carbon 2 of the uracil base. This requires the sulfur carrier and ubiquitin-like modifier protein Urm1 and a number of additional proteins operating upstream (Nfs1, Tum1, Uba4) and downstream (Ncs2, Ncs6) of Urm1 resulting in sulfur flow from cysteine to mcm\(^{5}\)U [11,20–25]. The absence of either part alone (mcm\(^{5}\)U or s\(^{2}\)U) induces shared pleiotropic phenotypes, including growth defects at elevated temperature and sensitivity to a number of exogenous stressors including the target of rapamycin complex 1 (TORC1)-inhibiting drug rapamycin [26,27]. On a functional level, partial loss of mcm\(^{5}\)s\(^{2}\)U reduces binding of tRNA to the ribosomal A-site and in turn causes reading frame slippage [28,29]. Combined loss of mcm\(^{5}\) and s\(^{2}\)U increases the severity of these phenotypes and causes a defect in protein homeostasis resulting in the accumulation of protein aggregates [30–32]. Similarly, combination of Elongator or Urm1 pathway mutations with a deg1 deletion aggravates growth defects and induces protein aggregation, which is further linked to severe problems in cytoskeleton organization, cell polarity and nuclear segregation [9].

Pleiotropic phenotypes of mutants lacking ψ\(^{38}/39\) together with mcm\(^{5}\)s\(^{2}\)U can be specifically rescued by overexpressing tRNA\(^{GlnUUG}\), which is reasonable since this tRNA alone carries both modifications [8,9]. In addition to tRNA\(^{GlnUUG}\), there is an isoacceptor tRNA with the anticodon sequence CUG that is thought to decode the G-ending codon for Gln. Genetic data from yeast suggests that the U34-containing isoacceptor mainly decodes the A-ending codon and only inefficiently reads the G-ending codon [33]. Because of this, the single tQ(CUG) gene has been found to be essential for yeast cell viability [33]. Except for the wobble nucleoside and a variable base (A/G) in position 42, tRNA\(^{GlnCUG}\) contains an anticodon stem loop (ASL) identical with that of tRNA\(^{GlnUUG}\) and is also modified by Deg1 at position U38 [34]. Interestingly, however, growth defects of deg1 single mutants were found to be suppressible by overexpression of tRNA\(^{GlnUUG}\), but not tRNA\(^{GlnCUG}\). This raises the possibility of differential functional dependencies of the two Gln isoacceptor tRNAs on the presence of ψ\(^{38}\). In this study, we investigated potential reasons for such divergent effects of one and the same modification on two highly similar tRNA species.

2. Results and Discussion

2.1. Phenotypic Rescue of deg1 Mutants by tRNA Overexpression

tRNA\(^{GlnUUG}\) functionality has been shown to depend on ψ\(^{38}\), in particular under conditions of heat stress [8]. Since the tRNA\(^{GlnCUG}\) isoacceptor harbours an ASL almost identical to that of tRNA\(^{GlnUUG}\), we first investigated whether there are indeed differential dependencies of the two Gln isoacceptors on the presence of ψ\(^{38}\). The thermosensitive growth phenotype of deg1 mutants was found to be specifically suppressible by overexpression of tRNA\(^{GlnUUG}\) but not tRNA\(^{GlnCUG}\) or any other tRNA carrying U38 or U39, which is modified by Deg1 to ψ [8]. We confirm this result and find a clear suppression of deg1-induced thermosensitivity by multicopy tRNA\(^{GlnUUG}\) but not tRNA\(^{GlnCUG}\) or other tester tRNAs (Figure 1A). However, rescue by multicopy tRNA\(^{GlnUUG}\) is clearly incomplete as growth at 39 °C is not fully restored to wild type levels (Figure 1A). Therefore, we checked whether co-overexpression of the isoacceptor tRNA\(^{GlnCUG}\) together with tRNA\(^{GlnUUG}\) improves growth at elevated temperature (39 °C) compared to overexpression of tRNA\(^{GlnUUG}\) alone. We find that under these conditions, growth of deg1 mutants already overexpressing tRNA\(^{GlnUUG}\) is not further improved by additional overexpression of tRNA\(^{GlnCUG}\) (Figure 1B). These results suggest that under heat stress conditions, growth defect of yeast cells lacking ψ\(^{38}\) results almost exclusively from malfunction of tRNA\(^{GlnUUG}\) rather than tRNA\(^{GlnCUG}\), despite containing almost identical ASLs.
why tRNA remains unaffected by the temperature effect on wobble uridine thiolation. tRNA\textsubscript{GlnCUG} the presence of an additional functional impairment of the former tRNA due to wobble uridine \(\psi\) to be essential for yeast viability \([33]\), indicating that this C34-containing tRNA is indeed indispensable for translation as compared to the minor tRNA\textsubscript{GlnCUG}. It should also be taken into account that copy numbers of the two Gln isoacceptors in yeast differ significantly due to the number of the respective tRNA genes (nine copies of tRNA\textsubscript{GlnUUG} versus one copy of tRNA\textsubscript{GlnCUG} (\textit{Saccharomyces cerevisiae} genome database)). This correlates with a biased codon usage and may impact on the efficiency of phenotypic suppression by tRNA overexpression. Based on this consideration, phenotypic effects of the \textit{DEG1} deletion may be rescuable by overexpression of tRNA\textsubscript{GlnUUG}, since this is the major Gln isoacceptor and consequently more important for overall translation as compared to the minor tRNA\textsubscript{GlnCUG}. However, the single tRNA\textsubscript{GlnCUG} gene is in fact known to be essential for yeast viability \([33]\), indicating that this C34-containing tRNA is indeed indispensable for translation, despite its lower frequency of utilization compared to the one containing mcm\textsuperscript{5}s\textsuperscript{2}U34. Hence, the differential dependency of both isoacceptors on \(\psi38\) as indicated by differential phenotypic suppression of the \textit{deg1} phenotype is presumably, at least in part, due to the difference at the wobble base. Interestingly, tRNA\textsubscript{GlnUUG} is known to harbour a hypomodified wobble uridine under mild heat stress conditions, which results from a destabilization of the \textit{IRM1} pathway and in turn suppresses the tRNA thiolation reaction that converts mcm\textsuperscript{5}U to mcm\textsuperscript{5}s\textsuperscript{2}U \([8,25,35-37]\). Thus, an explanation why tRNA\textsubscript{GlnUUG} depends more than tRNA\textsubscript{GlnCUG} on \(\psi38\) under heat stress conditions might be the presence of an additional functional impairment of the former tRNA due to wobble uridine hypomodification. tRNA\textsubscript{GlnCUG} carries a different unmodified wobble base (C34) and therefore remains unaffected by the temperature effect on wobble uridine thiolation.

Figure 1. Rescue of \textit{deg1} growth defect by tRNA overexpression. (A) Single tRNA overexpression. Indicated tRNA species were expressed from multicopy plasmids and in \textit{deg1} mutants and serial dilutions spotted on yeast peptone dextrose (YPD) plates that were incubated at 30 °C or 39 °C. In wild type cells, some tRNA species selected for overexpression carry 5-methoxy-carbonylmethyl-2-thiouridine (mcm\textsuperscript{5}s\textsuperscript{2}U), 5-carbamoylmethyluridine (mcm\textsuperscript{5}U), 5-methoxy-carbonylmethyluridine (mcm\textsuperscript{5}s\textsuperscript{2}U), pseudouridine (\(\psi\)) 38/39 or combinations thereof, as indicated by (+). (B) Combined overexpression of tRNA\textsubscript{GlnUUG} and tRNA\textsubscript{GlnCUG}. Cells carrying either a multicopy construct for overexpression of tRNA\textsubscript{GlnUUG} (tQ(UUG)), tRNA\textsubscript{GlnCUG} (tQ(CUG)) or both were assayed as in (A) for growth at 30, 37 and 39 °C. WT: wild type.
2.2. Genetic Interactions of DEG1 under TORC1 Inhibition

In support of a specific requirement for ψ38 in tRNA^{Gln}_{UUG} in the event of wobble base hypomodification, strong synthetic growth defects were observed upon combining the deg1 mutation with mutations in Elongator or Urm1 pathway genes, indicative of functional crosstalk between the modifications at position 34 and 38 in tRNA^{Gln}_{UUG} [8,9,38]. Elongator and Urm1 pathway mutants are further known to exhibit shared phenotypes, including translational inaccuracy and sensitivity to TORC1 inhibiting agents such as caffeine or rapamycin and these are synergistically increased in double mutants lacking both modification activities [9,30–32]. To check whether negative genetic interactions between DEG1, Elongator and URM1 can be extended to conditions of TORC1 inhibition, we scored rapamycin sensitivity of deg1, urm1 and elp3 single and all possible double mutant combinations. As shown in Figure 2, there is a clear rapamycin sensitivity in all three single mutants and this sensitivity is strongly aggravated in all the double mutants, with stronger effects seen in elp3 deg1 and urm1 deg1, where U34 and U38/39 are hypomodified. Thus, there is an overlapping phenotype in deg1, elp3 and urm1 mutants and all affected modifications (mcm5^s/2U, ψ38/39) appear to mediate rapamycin tolerance independently of each other, possibly by maintaining a related aspect of tRNA function. In particular, the observed rapamycin sensitivity of deg1 single mutants in the absence of heat stress suggests that ψ38 alone becomes important for tRNA function under at least one stress condition that is different from heat. Since there is strong negative genetic interaction on rapamycin between DEG1 and URM1/Elongator, the latter of which affect tRNA^{Gln}_{UUG} but not tRNA^{Gln}_{CUG}, we assume that these effects are caused by specific malfunction of tRNA^{Gln}_{UUG}.

![Figure 2](image)

**Figure 2.** Interactions between DEG1 and genes coding for Elongator (ELP3) or Urm1 (URM1) pathway components under TORC1 inhibition. Indicated single and double mutants were spotted on drug-free yeast peptone dextrose (YPD) plates which were incubated at 30 °C or 37 °C or YPD medium containing 5 nM of the TORC1 inhibiting drug rapamycin.

2.3. Replacement of tQ(CUG) by the sup70-65 Allele in Haploid Yeast

The finding that tRNA^{Gln}_{UUG} lacking ψ38 becomes vulnerable to disturbances of the natural anticodon loop configuration (loss of U34 modification) suggested that tRNA^{Gln}_{CUG} devoid of ψ38 might also be sensitized to destabilization of the ASL. However, as C34 is unmodified and no modification except ψ38 is known for the ASL of tRNA^{Gln}_{CUG} [7,34], we utilized a variant of tRNA^{Gln}_{CUG} with a disturbance of the ASL due to a sequence change (G31A). This mutation, known as the tQ(CUG) allele sup70-65, replaces the Watson–Crick base pair G31:C39 normally closing the anticodon loop of tRNA^{Gln}_{CUG} by a mismatch (A31:C39) and reduces stability and charging efficiency of the mutant tRNA [39–41]. Despite these strong effects on the tRNA product of the essential single-copy gene tQ(CUG), diploid yeast cells carrying the homozygous sup70-65 mutation are viable [39–41] and may provide an opportunity to analyze the effects of ψ38 removal on tRNA^{Gln}_{CUG} in the presence of a destabilized ASL. We utilized a haploid strain (S288C) lacking the single genomic
copy of tQ(CUG) and carrying the wild type tQ(CUG) gene on a plasmid that is counter-selectable by 5-fluoro-orotate (5-FOA) [31,40]. This strain was transformed with a second centromeric plasmid carrying the sup70-65 allele of the tQ(CUG) gene. Subsequently, the former plasmid was removed by growing cells on 5-FOA medium, resulting in a strain carrying sup70-65 as the sole genetic source of tQ(CUG) (Figure 3A). The viability of this strain suggests that even in haploid S288C cells, the mutant form of tQ(CUG) is active to maintain sufficient levels of translation. However, we find that in comparison to wild type, a strong thermosensitivity is caused by the mutation, while no significant growth defect was observed at 30 °C (Figure 3B). This finding possibly indicates that the destabilized form of tRNAGlnCUG (sup70-65) is sufficiently active in translation at normal, but not at elevated temperatures. The absence of defects in growth or CAG decoding in diploid sup70-65 cells was already documented [39]. However, more recent work demonstrated a reduced expression of reporter genes with tandem CAG codons and several natural CAG-codon-containing mRNAs [40,41]. Hence, a translational defect is already present in sup70-65 cells at 30 °C and this may be aggravated at elevated temperatures, as evidenced by the observed growth defect (Figure 3B). It is noteworthy that tRNAGlnCUG can be functionally replaced by the inefficient CAG decoder tRNAGlnUUG if the latter is present at elevated levels and carries the full mcm5s2U modification [33]. In such strain, CAG codons are exclusively decoded by tRNAGlnUUG, involving U-G wobble and the mcm5s2U modified wobble base. In particular, the absence of s2U in tRNAGlnUUG entirely prevents the suppression of lethal effects of a tQ(CUG) deletion by multi-copy tQ(UUUG) [33]. Thus, the strong growth defect of haploid sup70-65 cells at elevated temperatures may also indicate that tRNAGlnUUG contributes, to some extent, to CAG (Gln) decoding in this mutant, since tRNAGlnCUG is functionally impaired. If so, elevated temperature would repress thiolation and in turn prevent the engagement of tRNAGlnUUG in decoding of the G-ending Gln codon (CAG).

Figure 3. Role of ψ38 for the sup70-65 allele of the tQ(CUG) gene. (A) Experimental design for the construction of a haploid sup70-65 strain and plasmid shuffle. A strain carrying a genomic deletion of the essential single-copy tQ(CUG) is kept alive by plasmid pAK1 (tQ(CUG)-URA3). A second plasmid is introduced carrying the mutant sup70-65 allele of tQ(CUG) and pAK01 is eliminated from the strain by growth on 5-fluoro-orotate (5-FOA) media; (B) Drop-dilution assay as in Figure 1 with WT, deg1 and the haploid strain carrying sup70-65 as the sole genetic source of tQ(CUG); (C) Plasmid shuffle assay to check for the ability to lose pAK01 (tQ(CUG)-URA3) in the presence (right) or absence (left) of an additional deg1 deletion.

To test whether sup70-65 is still able to replace wild type tQ(CUG) when ψ38 is removed, we deleted the DEGI gene in a strain carrying a genomic deletion of tQ(CUG) and containing the wild type tQ(CUG) gene on the counter-selectable plasmid. Then, the ability to lose the tQ(CUG) plasmid in
the presence of the sup70-65 plasmid was assayed in comparison to the strain lacking the deg1 mutation. We find that colony formation on 5-FOA medium, which counter-selects the tQ(CUG) wild type allele, is severely reduced in the strain lacking DEG1 (Figure 3C). This result supports the interpretation that ψ38 becomes critical for function of tRNA<sup>Gln</sup><sub>CUG</sub> in the presence of a destabilized ASL.

2.4. Comparison of Non-Sense Suppression by sup70-65 and SUP4 in the Presence and Absence of ψ38/39

Since sup70-65 is known as an amber suppressor [39,40], we analyzed how loss of ψ38 influences the efficiency of UAG stop codon read-through by sup70-65. If ψ38 is indeed crucial for tRNA<sup>Gln</sup><sub>CUG</sub> function in the presence of a destabilized ASL, then stop codon read-through by sup70-65 could be impaired by the deg1 mutation. As a comparison, we scored the efficiency of SUP4, a distinct non-sense suppressor tRNA, derived from tRNA<sup>Tyr</sup><sub>GψA</sub> [10]. SUP4 can read-through ochre (UAA) stop codons due to a base change at the wobble position and was already shown to crucially depend on the presence of the Elongator-mediated mcm<sup>5</sup>U modification (mcm<sup>5</sup>UψA) [10]. Since tRNA<sup>Tyr</sup><sub>GψA</sub> harbours a DEG1 dependent ψ in position 39 [7], SUP4 could in principle also be affected by the deg1 mutation. To test this, we utilized a W303-1B-derived strain carrying SUP4 and ade2-1, an allele of ADE2 causing adenine auxotrophy due to an ochre mutation [10]. Non-sense suppression of the ade2-1 transcript is known to cause adenine prototrophy and change of colony color from red (no read-through) to white (read-through) [10]. In this background, we deleted the DEG1 gene and scored adenine auxotrophy as well as color colony in direct comparison with an elp3 mutant that is known to affect SUP4-mediated read-through. While the presence of SUP4 causes the expected change in color and adenine prototrophy, we find that this indeed depends on a functional ELP3 gene but not on DEG1 (Figure 4A). Since this assay might be too insensitive to detect smaller reductions in SUP4-mediated read-through and to confirm the obtained result with a premature ochre codon in another reporter gene, we utilized a quantitative assay relying on a lacZ reporter with an engineered ochre codon [33,42]. Again, read-through in this reporter was strongly enhanced by the presence of SUP4 and suppressed by the elp3 mutation, yet not by deg1 (Figure 4B). Further, even application of mild temperature stress (37 °C) to cells carrying the reporter constructs did not significantly reduce read-through levels of the deg1 mutant as compared to the wild type SUP4 strain under the same conditions (Figure 4B). Thus, Deg1-mediated ψ39 in itself does not detectably impact on the function of SUP4 in non-sense suppression in two different read-through contexts. This result is consistent with the absence of suppression of temperature sensitive growth of the deg1 mutant by multi-copy tRNA<sup>Tyr</sup><sub>GψA</sub> (Figure 1A) and supports the conclusion that ψ38 formed by Deg1 in distinct tRNA species may have different functional consequences.

For sup70-65-mediated non-sense suppression, we utilized strain W303-1B, which carries the trp1-1 allele that contains a premature amber (UAG) codon and is efficiently suppressible by sup70-65 when the mutant tRNA gene is present on a multi-copy plasmid [40]. We deleted the DEG1 gene in W303-1B and introduced sup70-65 on a multi-copy construct. As a control, we used the parental DEG1 wild type strain and introduced sup70-65 on both, single and multi-copy vectors [40]. While there is no major difference in growth on tryptophan-supplemented minimal media, the presence of the sup70-65 gene in both single and multi-copy conferred an ability to grow on tryptophan-free media specifically to the DEG1 wild type. As expected, multi-copy expression of sup70-65 was required for induction of efficient growth in the absence of tryptophan, due to relatively inefficient amber suppression by sup70-65 via first base wobble (U1:G36/A2:U35/G3:C34) [40]. In the absence of the suppressor tRNA, no growth on tryptophan-free medium was observed and, most importantly, growth was also absent in the deg1 mutant with the multi-copy sup70-65 construct causing efficient read-through in the DEG1 wild type strain. Thus, indeed sup70-65-mediated non-sense suppression critically depends on the presence of ψ38 formation by Deg1.
3. Materials and Methods

3.1. Strains, Plasmids and General Methods

Yeast strains used are listed in Table 1. Standard methods were used in this study for yeast growth [43]. Yeast was routinely grown in complex yeast peptone dextrose (YPD) medium. To select transformants or maintain plasmids, either synthetic complete (SC) medium lacking specific nutrients or YPD medium containing 200 μg·mL⁻¹ G418 was used. Deletion of DEG1 involved PCR-mediated synthesis of deletion cassettes using previously described oligonucleotides [9] and template plasmids pUG6 and pUG27 [44]. Transformation was done using the PEG/lithium acetate method [45]. Correct genomic replacement of the DEG1 gene was verified by PCR using oligonucleotides described [9]. Multicopy plasmids for overexpression of tRNA^Glu^\text{UUC}, tRNA^Lys^\text{UUU},
tRNA<sup>Arg</sup><sub>UCU</sub>, tRNA<sup>Leu</sup><sub>UAA</sub>, tRNA<sup>Tyr</sup><sub>GUA</sub> and tRNA<sup>Gly</sup><sub>GCC</sub> were described earlier [13] as well as the construct for tRNA<sup>Gln</sup><sub>UUG</sub> [46]. Wild type tRNA<sup>Gln</sup><sub>CUG</sub> overexpression used 2 µ plasmid pSUP70 [40]. pAK01 is a single copy (CEN/ARS) plasmid carrying the wild type tQ(CUG) gene as well as URA3 as a counter-selectable marker [40]. Counter-selection was done using synthetic complete medium containing uracil and 5-FOA at 1 mg·mL<sup>−1</sup>. Plasmids pSUP70-65-2 µ and pSUP70-65-CEN carry the sup70-65 allele of tQ(CUG) and either the 2 µ origin or CEN-ARS, respectively [40].

### Table 1. Strains used in this study.

| Strain                        | Genotype                           | Reference/Source               |
|-------------------------------|------------------------------------|--------------------------------|
| Saccharomyces cerevisiae      | BY4741 MATα, his3Δ, leu2Δ, met15Δ, ura3Δ | Euroscaf, Frankfurt            |
| S. cerevisiae elp3            | BY4741 elp3ΔKanMX4                 | Euroscaf, Frankfurt            |
| S. cerevisiae urm1            | BY4741 urm1ΔKanMX4                 | Euroscaf, Frankfurt            |
| S. cerevisiae deg1            | BY4741 deg1ΔKanMX4                 | Euroscaf, Frankfurt            |
| S. cerevisiae elp3 urm1       | BY4741 elp3ΔKanMX4 urm1ΔHIS3       | [31]                           |
| S. cerevisiae deg1 urm1       | BY4741 urm1ΔKanMX4 deg1ΔSpHIS5     | [9]                            |
| S. cerevisiae deg1 elp3       | BY4741 elp3ΔKanMX4 deg1ΔSpHIS5     | [9]                            |
| S. cerevisiae sup70           | BY4741 pAK01 tQ(CUG)AKLEU2         | [31]                           |
| S. cerevisiae sup70-65-pAK01  | BY4741 pAK01 tQ(CUG)AKLEU2 pSUP70-65-CEN | this work            |
| S. cerevisiae W303-1B         | MATα leu2-3,112 trp1-1 can1-100 ura3-1 ade2-1 his3-11,15 | [47]                           |
| S. cerevisiae RK207           | W303-1B deg1ΔSpHIS5                | this work                      |
| S. cerevisiae RK273           | W303-1B deg1ΔloxP                  | this work                      |
| S. cerevisiae UMY2893         | MATα SUP4 leu2-3,112 trp1-1 can1-100 ura3-1 ade2-1 his3-11,15 | [10]                           |
| S. cerevisiae UMY2916         | UMY2893 elp3ΔKanMX4               | [10]                           |
| S. cerevisiae RK208           | UMY2893 deg1ΔSpHIS5               | this work                      |

### 3.2. Phenotypic Assays

Yeast strains were either grown on YPD or appropriate SC solid medium to select for the presence of plasmids for 24 to 36 h. Cells were recovered from the plates and resuspended in sterile water. Cell densities were measured using absorbance at 600 nm and dilutions prepared from this suspension with final OD<sub>600nm</sub> values of 0.15, 0.015, 0.0015 and 0.00015. These were spotted on either drug-free solid YPD plates or YPD plates supplemented with sterile rapamycin at a final concentration of 5 nM. Subsequently, plates were incubated at different temperatures for 36–72 h and photographed. The longer incubation time (72 h) was used for phenotypic assay with deg1 elp3 double mutants. For qualitative read-through of ade2-1 and trp1-1 alleles, serial dilution were spotted on SC lacking adenine or tryptophan respectively and incubated for 48 to 60 h at 30 °C.

### 3.3. Quantitative Read-through Assay

Cells carrying pUKC815 (wild type lacZ) or pUKC817 (UAA ochre insertion) [42] were grown at 30 °C or 37 °C in liquid SC medium lacking uracil to OD<sub>600</sub> of 2–3 and harvested by centrifugation. Cells were washed and resuspended in Z-buffer (60 mM Na<sub>2</sub>HPO<sub>4</sub>, 40 mM NaH<sub>2</sub>PO<sub>4</sub>, 10 mM KCl, 50 mM 2-mercaptoethanol, pH 7). Cell density was measured by absorbance at 600 nm. To 500 µL aliquots, two drops of 0.01% sodium dodecyl sulfate (SDS) solution and chloroform were added and samples mixed on a vortex for 30 s each. Following incubation at 37 °C for 5 min, 100 µL of 4 mg·mL<sup>−1</sup> ortho-nitrophenyl-β-galactoside dissolved in Z-buffer was added. Reactions were stopped by the addition of 250 µL 1 M Na<sub>2</sub>CO<sub>3</sub> and absorbance at 420 nm measured. Activity units were calculated by employing Miller’s formula [48]. Relative read-through efficiency (%) was calculated by dividing the beta galactosidase activity measured with the pUKC817 construct by the one measured with the pUKC815 construct [33,42]. For each strain, at least three independent cultures were measured with both constructs.
4. Conclusions

The emerging picture from the presented work and previous studies places Deg1-mediated ψ38 in a role to maintain the function of both Gln isoacceptor tRNAs. However, the presence of ψ38 becomes critical for tRNA function only in the presence of an additional disturbance of the ASL structure. In one case (tRNA^{Gln}_{UUG}), a strong growth defect indicative of a significant loss of function is observed specifically when deg1 mutants are shifted to conditions known to downregulate the formation of a distinct modification (mcm^{5}s^{2}U) (Figures 1 and 2). In another case (tRNA^{Gln}_{CUC}) where this modification is absent per se, the introduction of a destabilization of the ASL can substitute for a hypomodified wobble base and sensitize this tRNA as well for the negative effects of the deg1 mutation (Figures 3 and 4). It appears possible that other tRNA species such as tRNA^{Tyr}_{GψA} do not lose function in the absence of ψ38/39 due to the existence of additional modifications that may stabilize the ASL. For example, tRNA^{Tyr}_{GψA}-derived suppressor tRNAs are known to carry N^{6}-isopentenyl-adenosine (i^{6}A) in position 37 [7]. The presence of i^{6}A has been shown to support efficient ochre read-through [49] and therefore may protect the ASL from destabilization. We assume that multiple modifications of the ASL provide multiple layers of structural protection, which routinely are not ruptured in single tRNA modification mutants and this functional redundancy may explain the paradox situation of missing phenotypes for many of the evolutionary conserved tRNA modification genes.

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