

### RESEARCH PAPER

OPEN ACCESS Check for updates



# Anticodon sequence determines the impact of mistranslating tRNA<sup>Ala</sup> variants

Ecaterina Cozma 📭, Megha Rao 📭, Madison Dusick 📭, Julie Genereaux, Ricard A. Rodriguez-Mias 🕞, Judit Villén 6, Christopher J. Brandl 6, and Matthew D. Berg 6

<sup>a</sup>Department of Biochemistry, The University of Western Ontario, London, Ontario, Canada; <sup>b</sup>Department of Genome Sciences, University of Washington, Seattle, WA, USA

#### **ABSTRACT**

Transfer RNAs (tRNAs) maintain translation fidelity through accurate charging by their cognate aminoacyl-tRNA synthetase and codon:anticodon base pairing with the mRNA at the ribosome. Mistranslation occurs when an amino acid not specified by the genetic message is incorporated into proteins and has applications in biotechnology, therapeutics and is relevant to disease. Since the alanyl-tRNA synthetase uniquely recognizes a G3:U70 base pair in tRNA Ala and the anticodon plays no role in charging, tRNA Ala variants with anticodon mutations have the potential to mis-incorporate alanine. Here, we characterize the impact of the 60 non-alanine tRNA anticodon variants on the growth of *Saccharomyces cerevisiae*. Overall, 36 tRNA anticodon variants decreased growth in single- or multi-copy. Mass spectrometry analysis of the cellular proteome revealed that 52 of 57 anticodon variants, not decoding alanine or stop codons, induced mistranslation when on single-copy plasmids. Variants with G/C-rich anticodons resulted in larger growth deficits than A/U-rich variants. In most instances, synonymous anticodon variants impact growth differently, with anticodons containing U at base 34 being the least impactful. For anticodons generating the same amino acid substitution, reduced growth generally correlated with the abundance of detected mistranslation events. Differences in decoding specificity, even between synonymous anticodons, resulted in each tRNAAla variant mistranslating unique sets of peptides and proteins. We suggest that these differences in decoding specificity are also important in determining the impact of tRNA Ala anticodon variants.

#### **ARTICLE HISTORY**

Revised 8 June 2023 Accepted 31 August 2023

#### **KEYWORDS**

Mistranslation: tRNAAla; genetic code; expanded decoding; tRNA biology

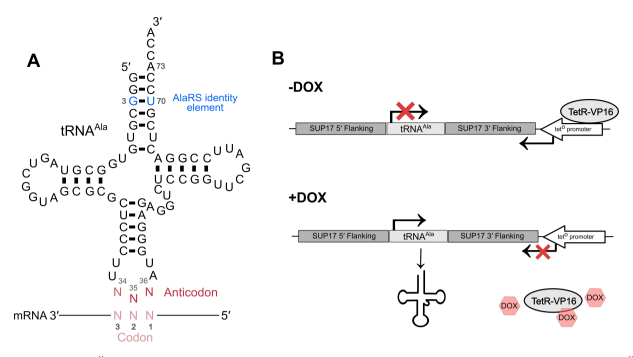
### Introduction

Transfer RNAs (tRNAs) are adaptor molecules that decode the genetic message contained within mRNA into protein sequence during translation. Translation fidelity and proteome accuracy are maintained by tRNAs at two steps. First, tRNAs are recognized by their aminoacyl-tRNA synthetase (aaRS) and charged with the cognate amino acid. Second, accurate base pairing between the tRNA anticodon and mRNA codon during decoding at the ribosome ensures that the correct amino acid is incorporated into the growing polypeptide chain [1-3]. Errors at either step lead to mistranslation, the incorporation of an amino acid not specified by the genetic message. Mis-incorporation events occur naturally in cells with estimated frequencies of 10<sup>-5</sup> to 10<sup>-3</sup> depending on the codon [4-7]. In addition, mistranslation increases in response to environmental conditions [8-10] and in the presence of tRNA variants [11-14].

Aminoacyl-tRNA synthetases recognize their cognate tRNAs through individual bases or structural motifs within the tRNA called identity elements [15-17]. For most tRNAs, the anticodon is the main identity element, providing a direct link between the genetic code and amino acid being incorporated. This is not the case for tRNA Ala and tRNA Ser, which are recognized through identity elements outside of the anticodon [18-21]. Specifically, the alanyl-tRNA synthetase (AlaRS) recognizes a conserved G3:U70 base pair in the tRNA Ala acceptor stem (Figure 1A) [18,19]. Seryl-tRNA synthetase recognizes a uniquely long variable arm 3' of the anticodon stem in tRNA Ser [20,21]. Identity elements for leucylation of tRNA<sup>Leu</sup> are species specific but are also largely independent of the anticodon [22-24]. Consequently, with one or two mutations to the anticodon tRNA Ala, tRNA Ser and tRNA Leu are theoretically capable of mis-incorporating alanine, serine or leucine, respectively, at non-cognate codons [13,14,25,26].

When introduced into cells, tRNA variants capable of mistranslation give rise to a heterogeneous mixture of molecules arising from the same genetic template but differing at a subset of amino acid positions. Carl Woese described these as statistical proteins which can provide cells with a wider range of functions than would be achievable from a homogeneous proteome [27]. As further indicated by Woese, statistical proteins form the basis of a powerful strategy for searching protein phase space to find novel protein activities [28].

Mistranslating tRNA variants have implications for health and disease. Santos et al. demonstrated that mistranslating tRNAs induce cell transformation and result in faster growing tumours in mice [12]. Flies and nematodes expressing



**Figure 1.** Schematic of tRNA<sup>Ala</sup> containing a degenerate anticodon and the tetracycline inducible tRNA expression system. (A) The structure of tRNA<sup>Ala</sup> with a degenerate anticodon. Bases colored in blue represent the G3:U70 base pair required for recognition and charging by AlaRS. The anticodon is shown in red. (B) tRNA<sup>Ala</sup> in the tetracycline inducible system used to regulate tRNA<sup>Ala</sup> anticodon variant expression flanked by up- and downstream *SUP17* sequence. In the absence of doxycycline, the tet<sup>O</sup> promoter is bound by the TetR-VP16 transcriptional activator which represses tRNA expression by driving RNA polymerase II expression across the tRNA gene. In the presence of doxycycline, TetR-VP16 binds doxycycline and dissociates from the promoter allowing the tRNA to be transcribed by RNA polymerase III.

mistranslating tRNAs show morphological defects [29,30]. Mistranslating flies also have locomotive defects similar to those seen for flies expressing alleles associated with neurodegeneration [29,31-33]. Consistent with a possible role in disease, tRNA variants with the potential to mistranslate are found in human populations [34,35]. Lant et al. demonstrated that in cell models of neurodegeneration a naturally occurring mistranslating tRNA variant slowed protein aggregate formation and inhibited the clearance of protein aggregates [36]. Further to their impact on health and disease, mistranslating tRNAs have potential in gene therapy as first demonstrated for β-thalassemia by Temple et al. [37]. Improvements in methods to introduce genes and RNA into cells have resulted in a renewed exploration of suppressor tRNA therapies being developed to treat diseases arising from premature termination codons, such as Duchenne muscular dystrophy and cystic fibrosis [38,39]. Lastly, mistranslating tRNA variants provide novel tools for determining the effects of thousands of missense mutations simultaneously [40], enabling the functional interpretation of missense mutations at a genome-wide level for personalized medicine applications.

The diverse applications and disease implications of mistranslating tRNAs led us to engineer all possible anticodon variants of *Saccharomyces cerevisiae* tRNA<sup>Ala</sup> and measure their impact on growth and ability to induce mistranslation. We hypothesized that the non-alanine anticodon variants would impact cells differently, as a result of making different amino acid substitutions. Indeed, differences in growth inhibition and mistranslation were observed for both non-synonymous and synonymous anticodons when found in the context of tRNA<sup>Ala</sup>. Impact was generally greatest for G/C-

rich anticodons and less for anticodons with U at base 34. The abundance of mistranslated peptides caused by the tRNA Ala variant, the specific proteins and positions mistranslated and the type of amino acid substitution contributed to the extent cell growth was affected by each variant.

## Materials and methods

## Yeast strains and growth

Wild-type haploid yeast strains are derivatives of BY4742 ( $MAT\alpha$   $his3\Delta1$   $leu2\Delta0$   $lys2\Delta0$   $ura3\Delta0$ ) [41]. The haploid strain CY8652 ( $MAT\alpha$   $his3\Delta1$   $leu2\Delta0$   $lys2\Delta0$   $ura3\Delta0$   $tTA^*-URA3$ ) containing the tet 'off' activator ( $tTA^*$ ) marked with URA3 was derived from R1158 [42] after crossing with BY4741 and sporulation.

Yeast strains were grown at 30°C in yeast peptone medium or in synthetic medium supplemented with nitrogenous bases and amino acids containing 2% glucose. Transformations were performed using the lithium acetate method as previously described [43].

## **Growth assays**

Inducible  $tRNA^{Ala}$  anticodon variants. Yeast strain CY8652 constitutively expressing the TetR-VP16 protein and containing a LEU2 plasmid expressing either a control  $tRNA^{Ala}_{GGC(Ala)}$  decoding alanine codons or the anticodon variants were grown to saturation in medium lacking uracil and leucine without doxycycline. Strains were diluted to  $OD_{600}$  of 0.1 in the same medium containing 0, 0.01 or 1.0

µg/mL of doxycycline and grown for 24 hours at 30°C with agitation. OD<sub>600</sub> was measured every 15 minutes for 24 hours in a BioTek Epoch 2 microplate spectrophotometer. Doubling time was calculated using the R package 'growthcurver' [44] and was normalized to the wild-type strain grown at the respective doxycycline concentration. For two variants, CGG(Pro) and CAG(Leu), which did not double at least twice in medium containing 1.0 µg/mL of doxycycline, we used area under the curve relative to the area under the curve of the control GGC(Ala) to determine relative growth. Each strain was assayed in triplicate.

Multicopy tRNA Ala anticodon variant derivatives. Yeast strain BY4742 was transformed with 1.0 µg of plasmid expressing the control tRNA Ala CGC(Ala) variant or anticodon variant tRNAs and plated onto minimal medium lacking uracil. Plates were imaged after 2 days of growth and quantified using the ImageJ [45] 'Watershed' package.

### **DNA** constructs

The construct encoding the tRNA Ala variants was synthesized as a GeneString fragment (ThermoFisher) containing a random mix of each nucleotide at anticodon positions 34, 35 and 36 and the tRNAAla gene was flanked by approximately 300 base pairs of up- and downstream sequence from SUP17. The synthetic gene fragment was cloned as a HindIII/ NotI fragment into pCB4699, a LEU2 centromeric plasmid (YCplac111) containing a tetracycline regulated promoter downstream of the tRNA gene as previously described [46]. For variants that were not selected out of the randomized anticodon pool, constructs expressing these tRNAAla variants were engineered via two-step PCR reactions. To reduce the number of primers needed, mismatched primer sets were used to generate two variants from a single PCR reaction. PCR fragments were subcloned into pGEM-Teasy (Promega) and then moved as HindIII/NotI fragments into pCB4699. The sequences of the 64 tRNA Ala anticodon variants generated and plasmid numbers are listed in Table S1. The cloning approach and primers used to generate each variant (if applicable) are listed in Table S2.

Multicopy plasmids expressing tRNAAla variants were made by cloning HindIII/EcoRI fragments containing each tRNA variant into YEplac195. Plasmid numbers are listed in Table S3.

## **Mass spectrometry**

Strains were grown overnight, in quadruplicate, in medium lacking leucine and uracil. Cultures were diluted to OD<sub>600</sub> of 0.01 in the same medium containing 1.0 µg/mL doxycycline except for the strain expressing tRNA Ala CGG(Pro) which was diluted to OD<sub>600</sub> of 0.1. Cultures were harvested by washing once in 1× yeast nitrogen base and frozen in liquid nitrogen when they reached an OD<sub>600</sub> between 0.8 and 1.0, except for tRNA Ala GCC(Glv) which due to its slow growth was harvested 36 hours after dilution at  $OD_{600} \sim 0.4$ .

Cells were lysed by bead-beating with 0.5 mm glass beads at 4°C in urea lysis buffer (8 M urea, 50 mM Tris pH 8.2, 75 mM NaCl). Lysates were cleared by centrifugation at 21,000 × g for 10 minutes at 4°C, and protein concentration was determined by bicinchoninic acid assay (Pierce, ThermoFisher Scientific). Proteins were reduced with 5 mM dithiothreitol for 30 minutes at 55°C, alkylated with 15 mM iodoacetamide for 30 minutes at room temperature in the dark, and the alkylation was quenched with an additional 5 mM dithiothreitol for 30 minutes at room temperature. For each sample, 50 µg of protein was diluted fourfold with 50 mM Tris pH 8.9 and digested for 4 hours at 37°C with 1.0 µg LysC (Wako Chemicals). Digestions were acidified to pH 2 with trifluoro-acetic acid and desalted over Empore C18 stage tips [47].

For strains containing the variants with UUU(Lys) and CUU(Lys) anticodons and the corresponding control strain containing tRNA Ala GGC(Ala) samples were lysed as above in 4 M guanidine hydrochloride, 100 mM triethylammonium bicarbonate pH 8.0. Samples were prepared as above, except after quenching the alkylation reaction, 100 µg of protein was acetylated with 200 µg of Sulfo-NHS-acetate (Pierce, ThermoFisher Scientific) for 1 hour at room temperature. Robotic purification and digestion of proteins into peptides was performed on the Kingfisher Flex (ThermoFisher Scientific) using trypsin and the R2-P1 method [48]. Digestions were acidified and desalted as above. Peptides were resuspended in 200 mM EPPS pH 9.5 with 5% hydroxylamine and incubated for 5 hours at room temperature to reduce acetylation on serine, threonine and tyrosine residues before peptides were desalted again.

Peptide samples were resuspended in 4% acetonitrile, 3% formic acid and subjected to liquid chromatography coupled to tandem mass spectrometry on a tribrid quadrupole Orbitrap mass spectrometer (Orbitrap Eclipse; ThermoFisher Scientific). Samples were loaded onto a 100  $\mu$ m ID  $\times$  3 cm precolumn packed with Reprosil C18 3 µm beads (Dr. Maisch GmbH) and separated by reverse-phase chromatography on a 100 µm ID × 30 cm analytical column packed with Reprosil C18 1.9 µm beads (Dr. Maisch GmbH) housed in a column heater set at 50°C. Peptides were separated using a gradient of 7-50% acetonitrile in 0.125% formic acid delivered at 400 nL/ min over 95 minutes, with a total 120-minute acquisition time. The mass spectrometer was operated in datadependent acquisition mode with a defined cycle time of 3 seconds. For each cycle, one full mass spectrometry scan was acquired from 350 to 1200 m/z at 120,000 resolution with a fill target of 3E6 ions and automated calculation of injection time. The most abundant ions from the full MS scan were selected for fragmentation using 1.6 m/z precursor isolation window and beam-type collisional-activation dissociation (HCD) with 30% normalized collision energy. MS/MS spectra were acquired at 30,000 resolution by setting the AGC target to standard and injection time to automated mode. Fragmented precursors were dynamically excluded from selection for 60 seconds.

MS/MS spectra were searched against the S. cerevisiae database (downloaded sequence from Saccharomyces Genome Database resource in 2014) using Comet (release 2015.01) [49]. The precursor mass tolerance was set to 20 ppm. Constant modification of cysteine carbamidomethylation (57.0125 Da) and variable modification of methionine oxidation (15.9949 Da) was used for all searches.

For samples containing the UUU(Lys) and CUU(Lys) anticodon variants and the corresponding control samples, variable modification of lysine acetylation (42.0105 Da) was included. A variable modification of each amino acid to alanine was used for the respective searches. A maximum of three of each variable modification was allowed per peptide. Search results were filtered to a 0.05% false discovery rate at the peptide spectrum match level using Percolator [50]. Peptide intensity was determined using in-house implementation of the moFF algorithm [51] to extract MS1 intensity. To determine if variants were inducing mistranslation, the proportion of identified mistranslated peptides was calculated from the number of unique mistranslated peptides for which the non-mistranslated sibling peptide was also observed. The mistranslated proportion is defined as the number of unique mistranslated peptides, where alanine had been mis-incorporated, divided by the number of total peptides containing the wild-type amino acid and expressed as a percentage. At the codon level, the decoding specificity of each tRNA Ala variant was determined using a custom Perl script (Supplemental File 1) to map codons back to mistranslated residues. Only peptides with one possible substitution event were considered to allow for accurate localization of the mistranslated residue.

The raw mass spectrometry data have been deposited to the ProteomeXchange Consortium via the PRIDE [52] partner repository with the dataset identifier PXD038242, and an annotated list of the file names can be found in Table S4.

## Factors used to correlate with growth impact of each variant

The following factors were correlated using multiple linear regression with the relative growth impact of each variant as the dependent variable: the mistranslation abundance calculated from the mass spectrometry data and the difference of the replaced amino acid relative to alanine for the following five properties: volume [53], electron-ion interaction potential [54], hydrophobicity [55],  $\alpha$ -helix propensity [55] and  $\pi$ -helix propensity [56].

#### Statistical analyses

Statistical analyses were performed using R studio 1.4.1717. Example code can be found in Supplemental File 2. Welch's t-test was used to compare doubling time or area under the curve of strains containing tRNA Ala anticodon variants to the control tRNA Ala GGC(Ala) strain and corrected for multiple tests using the Benjamini-Hochberg method. Wilcoxon rank sum test was used to compare median colony size of strains transformed with tRNA Ala anticodon variants on multicopy plasmids relative to the control tRNA Ala CGC(Ala) and corrected for multiple tests using the Bonferroni method. Welch's *t*-test was used to compare the proportion of mistranslated peptides detected measured for strains containing tRNAAla anticodon variants to the control tRNA Ala GGC(Ala) strain and corrected for multiple tests using the Benjamini-Hochberg method. For the multiple linear regression, independent variables with p-values less than 0.05 after Bonferroni multiple test

correction were considered to contribute to the impact each tRNAAla anticodon variant had on growth. Association between growth impact and estimated mistranslation frequency was determined using Spearman's correlation.

### **Results**

## Engineering and characterizing the growth impact of tRNAAla anticodon variants

To engineer tRNA Ala anticodon variants with the potential to mis-incorporate alanine at non-alanine codons, we cloned tRNA<sup>Ala</sup> with each of the 64 possible anticodon sequences into a tetracycline inducible system [46] to regulate tRNA expression and control mistranslation-associated toxicity (Figure 1, Table S1). In the absence of the tetracycline analog doxycycline, tRNA expression is repressed. When doxycycline is added to the growth medium, the tRNA variant is expressed in a titratable manner. Since we previously found that regulation of this system was improved with tRNA Ser SUP17 flanking sequence [46], the tRNAAla encoding gene was positioned precisely into the SUP17 locus (Figure S1). As such, all anticodons were examined in the same context and of note, the anticodon should not play a role in the stability or expression of a tRNA [13].

To assess the impact of the tRNA Ala variants on yeast growth, plasmids containing one of the 64 tRNAAla anticodon variants were transformed into a yeast strain constitutively expressing TetR-VP16 (CY8652) [46]. Transformants were isolated and their growth characterized in liquid minimal media containing 0, 0.01 or 1.0 μg/mL doxycycline. Growth curves were performed in biological triplicate for each strain at 30°C over 24 hours. Figure 2A shows representative growth curves for strains containing plasmids with the control  $tRNA^{Ala}_{\ GGC(Ala)}$  or  $tRNA^{Ala}$  variants with CAU(Met), GAA(Phe) and AGG(Pro) anticodons as examples of those having minimal, intermediate and high impact on growth, respectively. As expected, impact on growth was less severe at lower doxycycline concentrations for tRNA Ala variants with GAA(Phe) and AGG(Pro) anticodons. Growth of strains with the control tRNAAla<sub>GGC(Ala)</sub> or the variant with a CAU(Met) anticodon was unchanged at all doxycycline concentrations. Representative curves for all variants are shown in Figure S2.

The relative growth of each strain containing one of the 64 possible tRNA Ala variants in medium with 1.0 μg/mL doxycycline is shown in Figure 2B, and relative growth at all three doxycycline concentrations is shown in Figure S3 (raw data can be found in Table S5). Of the 60 variants with non-alanine anticodons, 25 significantly decreased growth compared to the control tRNA Ala GGC(Ala) (Table S6). We observed a range of relative growth rates indicating that tRNAAla variants with anticodons decoding the 19 non-alanine amino acids have diverse impacts. For example, G/C-rich variants with CGG(Pro), GAC(Val), GCC(Gly) and CAG(Leu) anti-codons resulted in the greatest growth reduction. Anticodon variants that caused no reduction in growth included the A/U-rich variants with AAA(Phe), AAU(Ile), UAU(Ile), UUU(Lys) and AUU(Asn) anticodons as well as variants with anticodons decoding the three stop codons.

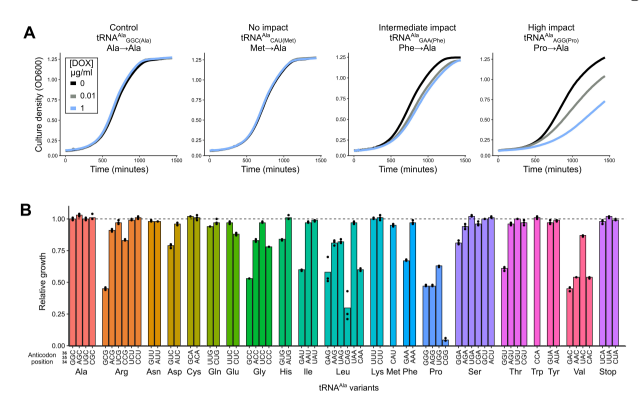


Figure 2. Impact of tRNA<sup>Ala</sup> anticodon variants on yeast growth. (A) Representative growth curves of yeast strain CY8652 containing a plasmid with the control tRNA<sup>Ala</sup><sub>GGC(Ala)</sub>, a tRNA<sup>Ala</sup><sub>CAU(Met)</sub> variant with no growth impact, a tRNA<sup>Ala</sup><sub>GAA(Phe)</sub> variant with intermediate growth impact or a tRNA<sup>Ala</sup><sub>AGG(Pro)</sub> variant with a severe growth impact. Black, grey and blue lines represent strains grown in media containing 0, 0.01 and 1.0 µg/mL doxycycline, respectively. (B) Yeast strain CY8652 containing plasmids with the control tRNA<sup>Ala</sup> GGC(Ala) or one of the tRNA<sup>Ala</sup> anticodon variants were grown for 48 hours at 30°C in medium lacking uracil and leucine, diluted to an OD600 of 0.1 in the same medium with 1.0 µg/mL doxycycline and grown for 24 hours at 30°C with agitation. OD600 was measured at 15-minute intervals and doubling time was quantified. Relative growth of tRNA Ala anticodon variants was calculated by multiplying the inverse of each variant doubling time by the average doubling time of the tRNA Ala GGC(Ala) control (for raw data and statistical comparisons see Table S5 and S6). For strains containing CGG(Pro) and CAG(Leu) variants area under the curve was used to determine relative growth. Each point represents one biological replicate (n = 3).

The minimal impact of anti-codons variants decoding stop codons may reflect factors other than A/U richness such as the strong competition from release factors, the requirement for only one stop codon per gene, the presence of redundant stop codons downstream of many genes and the ability of many proteins to tolerate a C-terminal extension [57]. Interestingly, different growth impacts were observed for variants decoding synonymous codons. For example, the variant with a GGU(Thr) anticodon reduced growth to 61% of the control tRNA Ala, whereas the other three threonine decoding variants had a minimal effect.

To analyse the importance of anticodon G/C content on the impact of tRNA Ala variants, we plotted the relative growth of strains containing each variant on Grosjean and Westhof's alternative representation of the genetic code [58], which places 'strong' G/C-rich anticodons at the top of the circle plot and 'weak' A/U-rich anticodons at the bottom as determined by the Turner binding energy of the base pairing between the first and second positions of the codon with bases 35 and 36 of the anticodon. As shown in Figure 3A, 'strong' anticodons generally resulted in a greater reduction in growth compared to 'weak' anticodons.

Next, we evaluated how the identity of tRNA base 34, the first position of the anticodon, impacts growth. Figure 3B shows the relative growth of strains containing synonymous anticodon variants decoding the same amino acid and differing only at base 34. In four-box sets, variants with U at position 34 had less impact on growth as compared to their

A, G or C counterparts. Similarly, two-box variants with A at position 34 had less impact on growth than their G-containing counterparts. Consistently, G or C at position 34 had the greatest impact on growth in each isodecoder set.

Of the 57 possible anticodons not decoding alanine or stop codons, genes encoding 18 of these anticodons are not found in S. cerevisiae. In Figure 3B, non-native and native anti-codons are represented by open and filled points, respectively. For the three- and four-box tRNA variants including those decoding Ser, Arg and Leu codons, the most impactful anti-codons is non-native in seven of eight cases (the exception being the variant with the GCC(Gly) anticodon). This pattern does not hold in any of the three cases for the two-box tRNAs where it can be evaluated.

## Increasing tRNAAla anticodon variant copy number reveals growth impact

In single copy, 35 non-alanine variants did not statistically impact growth when compared to the control strain express-ing tRNA Ala GGC(Ala). Since each tRNA Ala anticodon variant competes with the native tRNA species for decoding, we cloned a subset of the variants into a multicopy plasmid to determine if increasing copy number would produce a growth phenotype. The wild-type BY4742 strain was transformed with the multicopy plasmids and transformant colony size quantified relative to a control

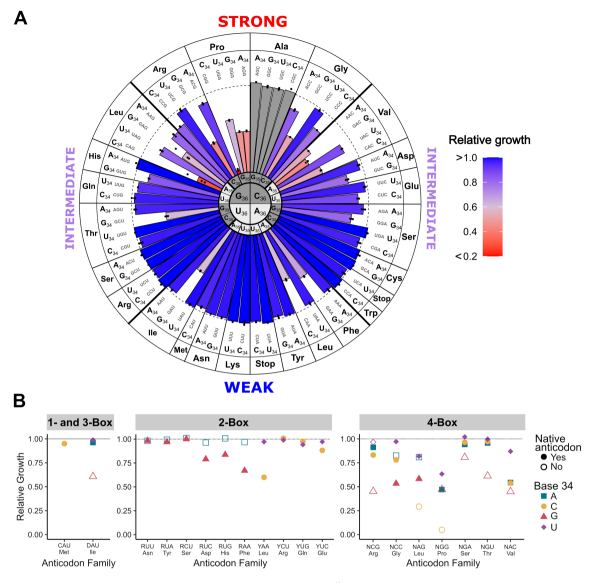


Figure 3. Anticodon sequence affects growth impact. (A) Relative growth for each tRNA<sup>Ala</sup> variant as in Figure 2B plotted for each anticodon on Grosjean and Westhof's alternative representation of the genetic code [58] where 'strong' codon:anticodon pairs are at the top of the circle plot and 'weak' pairs are at the bottom. The length of each bar is proportional to the relative growth of each variant. (B) Relative growth for mistranslating tRNA<sup>Ala</sup> anticodon variants decoding synonymous anticodons differing at base 34. Variants are grouped by number of different possible anticodons decoding a specific amino acid (1-, 2-, 3- or 4-box) with the same base at positions 35 and 36. Amino acid identity is listed below the anticodon. Native and non-native anticodon sequences in yeast are depicted with filled and empty shapes, respectively. D: adenine, quanine or uracil; R: purine; Y: pyrimidine; N: any base.

transformation of tRNA Ala CGC(Ala) on a multicopy plasmid (Figure 4, Figure S4, Table S7). We chose to quantify colony size after transformation to reduce the number of passages and selection pressure of deleterious plasmids towards lower copy. As a control to demonstrate that overexpressing a mistranslating variant exacerbates growth impact, we also analysed the GAA(Phe) anticodon variant that resulted in slow growth on a single-copy plasmid. When this variant was expressed on a multicopy plasmid, no transformants were obtained. Of the 17 low impact variants tested, 12 significantly decreased median colony size compared to the control when introduced on a multicopy plasmid. Four of these led to mean colony size 50% less than the control (variants with UCC(Gly), CAU(Met), AGU(Thr), AAA(Phe) anticodons) and one variant produced transformants that were too small to quantify (variant with

CUG(Gln) anticodon). These variants likely mistranslate in single copy, but at a low frequency that does not lead to a growth phenotype. In contrast, five tRNA<sup>Ala</sup> variants showed no statistical impact on colony size, even in high copy number (variants with GCA(Cys), UUU(Lys), CCA(Trp), AUA(Tyr) and CUA(Stop) anticodons).

# Estimating mistranslation frequency of each tRNA<sup>Ala</sup> anticodon variants

To evaluate the relationship between reduced growth caused by the tRNA  $^{\rm Ala}$  variants and their ability to induce mistranslation, we analysed the cellular proteome by mass spectrometry after growth of each strain in medium containing 1.0 µg/mL doxycycline and isolation of cellular protein in denaturing conditions. On average, we identified ~3000 unique proteins

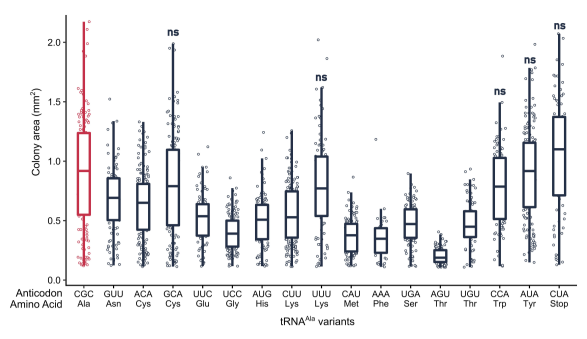


Figure 4. Overexpressing some but not all tRNA<sup>Ala</sup> variants reduce growth. Strain BY4742 was transformed with 1.0  $\mu$ g of a *URA3* multicopy plasmid containing each tRNA<sup>Ala</sup> anticodon variant, plated on medium lacking uracil and grown at 30°C. Transformants were imaged after 48 hours (raw images are in Figure S4) and colony area was quantified using the ImageJ 'Watershed' package. Each point represents one colony and horizontal bars at the center of each boxplot represent median colony size. Median colony size of each variant was compared to the control tRNA<sup>Ala</sup><sub>CGC(Ala)</sub> (shown in red) using a Wilcoxon rank sum test with Bonferroni correction to determine significance (see Table S7 for *p*-values). Variants lead to a statistically significant decrease in growth (p < 0.01) unless denoted by 'ns'.

per sample (Figure S5A). To determine if mistranslation was occurring, we calculated the fraction of unique mistranslated peptides relative to the total number of peptides containing the wild-type amino acid (Figure 5A, Table S8). Of the 57 non-alanine anticodon variants measured, 52 resulted in statistically elevated levels of mistranslated peptides compared to the control strain (variants decoding stop codons were not analysed). We identified mistranslated peptides mapping to up to 566 unique proteins depending on the strain (Figure S5B). Mistranslation events were more commonly found for more abundant wild-type peptides (Figure S5C), likely reflecting the ease of identifying the fractional occurrence of mistranslation in more abundant proteins. Similar to the effect these variants had on growth, the proportion of mistranslated peptides detected spanned a wide range. For example, we detected 12.8%, 11.9% and 11.6% of peptides mistranslated for variants UGG(Pro), GAC(Val) and GUG(His) anticodons, respectively, whereas we detected 0.12% of peptides mistranslated for the variant with an AAA(Phe) anticodon. It is important to recognize that these values are estimates, as the measured proportion of peptides mistranslated reflects specific properties of the mistranslated versus wild-type peptides, such as differences in protein turnover and the ability to detect individual peptides.

We hypothesized that the impact each variant has on growth is influenced by the abundance of mistranslated proteins present in the cell under steady-state conditions, which takes into account how frequently mistranslation occurs and the number of sites mistranslated throughout the proteome. Therefore, we calculated the summed MS1 intensity of detected mistranslated peptides normalized to the summed MS1 intensity of all peptides and plotted this against relative growth (Figure S6). There was a weak correlation, where

variants that had a greater overall abundance of mistranslated peptides tended to have more impact on growth (linear  $r^2 = 0.20$ , Spearman's rank correlation  $p = 1.81 \times 10^{-9}$ ).

Since the type of amino acid substitution would also be expected to influence the impact on growth, we plotted the estimated abundance of mistranslated peptides versus relative growth separately for the eight non-alanine isoacceptor sets that have more than two anticodons (Figure 5B). In general, the estimated abundance of mistranslated peptides increased, the impact on growth was more severe. Negative correlation coefficients greater than 0.9 were seen for 5 of the 8 cases. Indicative of some substitutions being better tolerated than others, the slope of the line reflecting the relationship between estimated total mistranslation and growth differed between amino acid families. The arginine to alanine and serine to alanine substitution plots had the greatest and least slopes, respectively. The presence of additional factors impacting growth in an anti-codons specific fashion was apparent from the variants with anticodons decoding glycine, leucine and valine codons, where reduced growth caused by one or more variants did not correlate well with the summed MS1 intensity of the mistranslated peptides. In contrast to the other variants, the plot for variants mis-incorporating alanine at proline codons showed that NGG(Pro) anticodon variants causing less mistranslation had a greater impact on growth.

The amino acid-specific differences in the slopes of the mistranslation abundance versus relative growth plots support that the chemical properties of the substituted amino acid relative to alanine are a factor in determining the impact of the mistranslation event on growth. To determine if we could attribute this relationship to a specific factor, we performed multiple linear regression using the difference between each replaced amino acid and alanine for five factors identified by

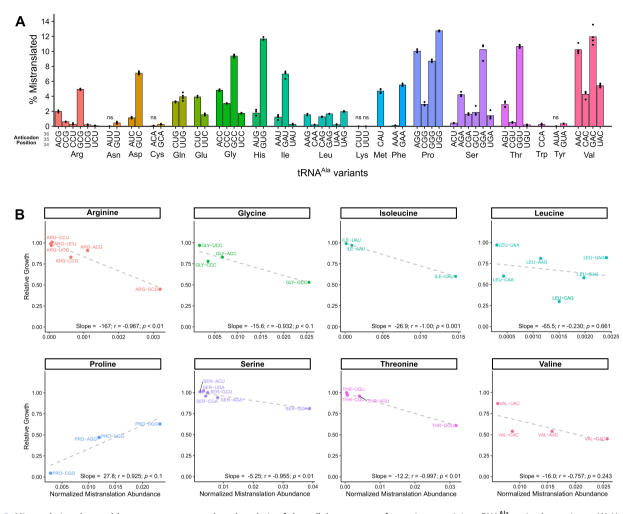


Figure 5. Mistranslation detected by mass spectrometry-based analysis of the cellular proteome for strains containing tRNA<sup>Ala</sup> anticodon variants. (A) Yeast strain CY8652 containing a tRNA<sup>Ala</sup> anticodon variant were diluted into medium containing 1.0 μg/mL doxycycline to induce tRNA expression and harvested at an OD<sub>600</sub> of ~ 1.0. Cells were lysed in a denaturing buffer and mass spectrometry analysis of the cellular proteome was performed. The proportion of peptides detected as mistranslated, expressed as a percentage, was calculated for each variant from the number of unique peptides where alanine mis-incorporation was observed relative to the number of unique peptides where the wild-type amino acid was observed. Each point represents one biological replicate (n ≥ 3). Variants have statistically elevated levels of mistranslated peptides relative to the control strain containing tRNA<sup>Ala</sup><sub>GGC</sub> (Welch's t-test; Benjamini-Hochberg corrected p < 0.01) unless denoted by 'ns'. (B) Summed MS1 intensity of mistranslated peptides normalized to the summed MS1 intensities of all detected peptides (normalized mistranslation abundance) plotted against the relative growth for each tRNA<sup>Ala</sup> variant for amino acid families with 3 or more anticodons. The grey dashed line represents the linear correlation between the two variables. Note the x-axis is scaled differently for each amino acid plot and does not always start at zero.

Then *et al.* that distinguish the proteinogenic amino acids [59]: (1) volume [53]; (2) electron-ion interaction potential [54]; (3) hydrophobicity [55]; (4)  $\alpha$ -helix propensity [55] and (5)  $\pi$ -helix propensity [56]. When these factors were included, the correlation with growth improved (linear  $r^2 = 0.44$ ,  $p = 3.25 \times 10^{-5}$ ) as compared to when we only considered mistranslation abundance. Volume, electron-ion interaction potential and  $\pi$ -helix propensity significantly contributed to the improved correlation, with electron-ion interaction potential having the largest impact followed by  $\pi$ -helix propensity and volume (Table S9).

# Synonymous anticodons target different proteins and positions

For the variants that substitute glycine, valine and leucine with alanine, reduced growth caused by one or more variants did not correlate well with the summed MS1 intensity of the mistranslated peptides. The tRNA<sup>Ala</sup> variants with

leucine anticodons resulted in mistranslation amounts that were the least correlated with their effect on growth (Figure 5B). The UAG(Leu) anticodon variant induced the greatest amount of mistranslation within this set and reduced growth to ~80% of the control strain, whereas the CAG(Leu) variant induced an intermediate amount of mistranslation but reduced growth to 30%. We hypothesized that the different impact on growth might be explained by codon-specific mistranslation by the tRNAAla variants, resulting in the mistranslation of different positions within different proteins. Figure 6A shows codons that were mistranslated for the four tRNAAla variants with NAG leucine anticodons. The decoding specificity, which we define as the set of synonymous codons where we observe mistranslation, for variants with UAG(Leu) and CAG(Leu) anticodons was distinct from each other and from the variants with AAG(Leu) and GAG(Leu) anticodons, which were similar. In line with the differences in decoding specificity, there was minimal overlap in the mistranslated peptides

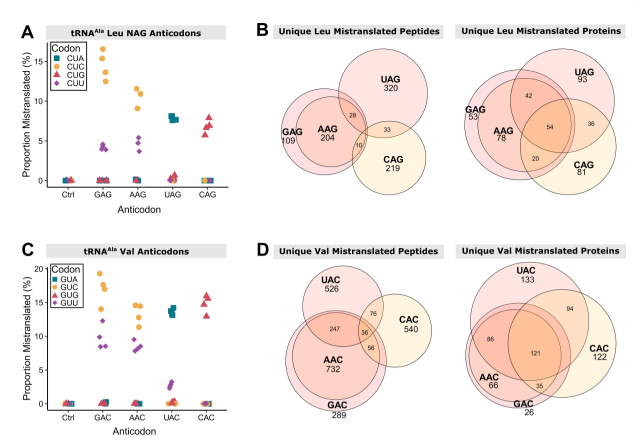


Figure 6. tRNA<sup>Ala</sup> variants with leucine NAG anticodons or valine NAC anticodons mistranslate at unique codons leading to different subsets of the proteome experiencing mistranslation. (A) Codon specific proportion of unique mistranslated peptides for the control strain containing tRNA<sup>Ala</sup> GGC(Ala) and four tRNA<sup>Ali</sup> containing synonymous anticodons decoding leucine codons. Proportion of mistranslated peptides at each codon was calculated from mass spectrometry data as the number of unique mistranslated peptides divided by the number of wild-type peptides where the leucine residue is coded for by the codon indicated. Only peptides containing one leucine residue were considered. Each point represents one biological replicate ( $n \ge 3$ ). (B) Venn diagrams showing the overlap between the unique peptides (left) and proteins (right) where mistranslation was detected by mass spectrometry in strains containing tRNAAla variants that have one of four synonymous anticodons decoding leucine codons. (C) Codon specific proportion of unique mistranslated peptides for the control strain containing tRNA<sup>Ala</sup><sub>GGC(Ala)</sub> and four tRNA<sup>Ala</sup> variants containing synonymous anticodons decoding valine codons as in (A). (D) Venn diagrams showing the overlap between the unique peptides (left) and proteins (right) where mistranslation was detected by mass spectrometry in strains containing tRNAAla variants that have one of four synonymous anticodons decoding valine codons.

and proteins identified in strains containing the variants with UAG(Leu) and CAG(Leu) anticodons (Figure 6B). In contrast, variants with AAG(Leu) and GAG(Leu) anticodons mistranslated relatively similar sets of peptides and proteins and their growth impacts were proportional to their estimated mistranslation abundance.

The same analysis was done for the set of tRNA Ala variants substituting alanine for valine. The estimated total mistranslation by variants with UAC(Val) and CAC(Val) anticodons was similar but these variants decreased growth to 87% and 54% of the control strain, respectively. The decoding specificity of the variants with UAC(Val) and CAC(Val) anticodons was distinct from each other and from the variants with AAC(Val) and GAC(Val) anticodons, which were similar (Figure 6C). Again, in line with the differences in decoding specificity, there was minimal overlap in the mistranslated peptides and proteins identified in strains containing the variants with UAC(Val) and CAC(Val) anticodons (Figure 6D), while variants with AAC(Val) and GAC(Val) anticodons mistranslated relatively similar sets of peptides and proteins. These results suggest that differences in decoding

specificity could contribute to the impact on growth of synonymous anticodon variants.

# Decoding specificity of tRNA<sup>Ala</sup> variants

The mass spectrometry data identifying sites of mistranslation allowed us to determine the codons mistranslated by each variant. Figure 7 shows the decoding specificity for tRNA Ala variants with statistically elevated mistranslation frequencies (raw data in Table S10, Figure S7). In 10 of 13 cases, G at position 34 resulted in mistranslation of codons with C and U at the third codon position. Interestingly, in these cases, with the exception of GCC(Gly), a greater proportion of mistranslated peptides were identified with codons ending in a 3' C rather than a 3' U. Variants with A at position 34 have the potential to be modified to I34 (inosine) and decode codons ending with U, C and A [60-62]. Of the nine A34containing variants, seven mistranslated codons ending in U and C, with the exceptions being ACC(Gly) and AUG(His). This suggests that these variants have expanded decoding consistent with inosine modification; however, we only detected mistranslation at codons ending in A for the

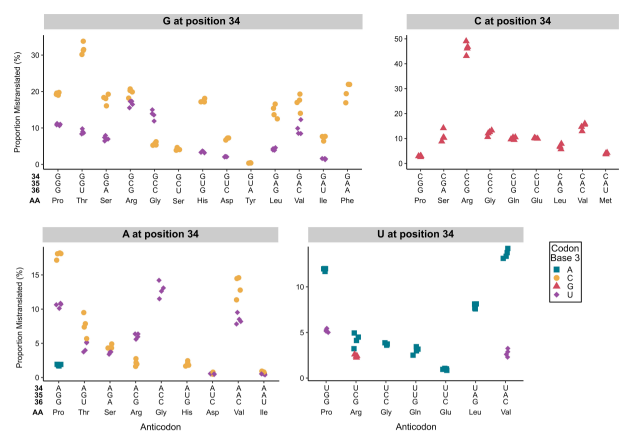


Figure 7. Profile of mistranslation events detected at synonymous codons for tRNA<sup>Ala</sup> variants with G, A, C or U at base 34. For each tRNA<sup>Ala</sup> anticodon variant, the proportion of mistranslated peptides identified at each synonymous codon was calculated from the number of unique peptides with a mistranslation event at that codon relative to the number of unique peptides containing a wild-type residue at that specific codon. In all cases, positions 35 and 36 formed Watson-Crick pairs with positions 2 and 1 of the codon, respectively. To confidently localize the mistranslation event and identify the mistranslated codon, only peptides containing a single target amino acid were used in this analysis. Only mistranslation that is statistically above the frequency measured for the same codon in the wild-type strain is shown (Welch's *t*-test; Benjamini-Hochberg corrected p < 0.01). Each point represents one biological replicate ( $n \ge 3$ ). Note the scale of the y-axis is different for each plot.

AGG(Pro) variant. Variants with U at position 34 predominantly mistranslated codons with 3' A, with U:G decoding only detected for the UCG(Arg) variant. Interestingly, U:U decoding, indicative of superwobble, was detected for UGG(Pro) and UAC(Val) variants. Mistranslation by tRNA<sup>Ala</sup> variants with C at position 34 was limited to codons ending in G.

## **Discussion**

Our study has characterized the 60 non-alanine tRNA<sup>Ala</sup> anti-codons variants for their mistranslation potential and impact on the growth of. *cerevisiae*. Twenty-five of the 60 variants reduced growth when present in single copy. Slow growth due to some but not all remaining variants was observed when their copy number was increased. As the translation machinery, including tRNAs, is conserved throughout evolution, these results suggest that any potential impact of tRNA<sup>Ala</sup> anticodon variants on human health or disease could depend on the specific sequence of the anticodon. Mistranslation was observed for 52 of 57 non-alanine variants, indicating that mistranslation does not always lead to reduced growth. The 52 variants that do mistranslate will have applications in synthetic biology and could potentially

be used as therapeutics. Furthermore, the fact that these tRNA<sup>Ala</sup> variants can be introduced into cells without causing cell death has implications for the adaptability and evolution of the genetic code (see for example [63,64]).

# Factors influencing the impact of $tRNA^{Ala}$ variants on cell growth

There are multiple factors that interface to influence the extent to which a mistranslating tRNA impacts cell growth. The first to consider is the overall level of mistranslation. The amount of mistranslation experienced by a cell is dependent on the frequency of mistranslation, the number of codons that are mistranslated and the expression level of proteins containing the effected codons. Likewise, the frequency of mistranslation is dependent on multiple factors including the abundance of native, wild-type isodecoder tRNAs competing with the mistranslating tRNA variant and factors related to the efficiency of the tRNA variant in the translation process. Kinetic aspects of translation are relatively uniform for native amino-acyl-tRNA despite the diversity in amino acid structure and tRNA sequence [65-68]. At first glance, this would suggest that all tRNA Ala anticodon variants should mistranslate at similar levels and therefore mistranslation frequency would only depend on

the number of competing native tRNAs. However, as noted by the Uhlenbeck lab [65,69], to obtain translational uniformity each tRNA has evolved sequence or modification differences to tune their activity to a similar level. Our results support this argument since when taken out of their native context, the tRNA Ala anticodon variants do not elicit an equivalent level of mistranslation or impact on growth. This is most easily seen for 2-box anticodons where there is only one native isodecoder in S. cerevisiae (Phe, Asn, Asp, His, Tyr and Cys), and thus the competition with native tRNAs is the same for both synonymous anticodons. For all six of these amino acid families, the two tRNA Ala variants have different estimated mistranslation frequencies and different impacts on growth (Figure S8). In all of these cases, the variant containing a G at position 34 had a more severe impact than the A-containing counterpart. In fact, throughout the entire set of variants, G/C-rich anticodons had a greater impact on growth compared to the A/U-rich variants. It is possible that tRNA modifications play a role in this effect by tuning translation efficiency. For example, native tRNAs with A/U-rich anticodons are often modified at base 37 to N<sup>6</sup>-isopentenyladenosine or N<sup>6</sup>-threonylcarbamoyladenosine to stabilize base pairing interactions [70-73]. These modifications likely do not occur for tRNAAla, which has a G/C-rich anticodon, thus potentially reducing the activity of A/U-rich anticodon variants. Consistent with this, Pernod et al. found limited codon:anticodon interactions between A/U-rich codons in an in vitro system [74]. We note that we cannot exclude the possibility that subtle changes in tRNA Ala structure arise as the result of non-cognate anticodon sequences that could impact alanylation by AlaRS or translational efficiency.

A second principal factor determining the impact of each mistranslating tRNA Ala variant is the specific set of proteins and peptides that are mistranslated. Even small changes in expression or function of certain proteins can impact growth [75]. Since amino acid composition varies from protein to protein, each tRNA Ala anticodon variant has the potential to contribute to slow growth by decreasing the functional amount of one or more distinct proteins. The ability of variants to impact different cellular pathways is supported by our previous finding where two tRNAs mistranslating at similar frequencies, one mis-incorporating alanine at proline codons and the other serine at arginine codons, had unique sets of genetic interactions [76]. Furthermore, because codon usage is not random, synonymous anticodons target different proteins and thus impact growth to different extents as demonstrated by tRNA Ala variants decoding the variants with UAG(Leu) and CAG(Leu) anticodons. Also related to mistranslation contributing to changes in protein expression is the ability of specific mistranslation events to influence translational elongation and regulate protein folding. Reduced translation rate at the 5' end of genes, which minimizes abortive synthesis, and ribosome stalling, which can promote protein folding, are achieved by the selection of specific codons (reviewed in [77-79]). Proline codons specifically have a significant role since peptide bond formation involving proline as a donor or acceptor is slow and stretches of two or more prolines lead to ribosome stalling. Therefore, mistranslation events replacing proline have the potential to alter protein expression. In this regard, it is interesting that tRNA Ala variants with proline anticodons are, as a group, the most impactful on growth and the only set where a positive correlation between the amount of mistranslation and growth was observed.

Related to the importance of codon usage regulating protein expression, Gamble et al. identified 17 codon pairs in yeast that negatively impact translation [80]. These codon pairs tend to be conserved throughout yeasts [81], suggesting they have a functional role. When non-native tRNAs with anticodons cognate to one of the codons in the inhibitory pair are introduced into cells, the inhibitory effect of the pair on translation is alleviated [80]. Likewise, introducing the non-native mistranslating tRNAAla variants might affect the regulation of gene expression achieved through inhibitory codon pairs. Interestingly, 3 of the 17 codon pairs contain the proline CCG codon, providing another possible explanation for why the CGG(Pro) anticodon variant had the largest impact on growth while having relatively low levels of mistranslation.

We also considered the nature of the amino acid substitution as a determinant of the impact of each tRNA<sup>Ala</sup> variant on growth. The methyl side chain of alanine is small and hydrophobic. Mis-incorporating alanine for amino acids with distinct chemical or functional properties might be expected to have a greater impact. Supporting this argument, the correlation between the impact on growth and abundance of mistranslated peptides improved when the anticodons were separated by amino acid substitution. When we considered five properties that differentiate the 20 amino acids [59], differences in volume, electron-ion interaction potential and  $\pi$ -helix propensity of the replaced amino acid relative to alanine contributed to the impact each substitution had on growth.

In this discussion, we have restricted our consideration to the roles of tRNAs in translation. However, many reports are continuing to expand the known roles for tRNAs in other processes. For example, tRNA Gln CUG is important for signalling in response to poor nitrogen sources [82], and the anticodon is required to impart this function [83]. We cannot exclude the possibility that these non-translational functions contribute to the growth impairment of some of the tRNA Ala variants.

## Similarities and differences with mistranslating tRNA Ser variants

Previously, Zimmerman et al. investigated the impact of tRNA<sup>Ser</sup> variants with all 58 non-serine anticodons on the growth of yeast [13]. While their data was acquired in pooled format, making direct comparison with our tRNA are results difficult, their findings of anticodon specific differences including those between synonymous anticodons are consistent with our tRNAAla results.

We previously analysed some individual tRNA Ser variants [14] and find that in general, tRNA Ser anticodon variants are more toxic than their tRNAAla counterparts. Mistranslation by a tRNA Ala or tRNA Ser variant would be reduced if its aminoacylation by AlaRS or SerRS, respectively, was in competition with the anticodon-cognate



aaRS. This neutralizing mechanism would be most likely for the 11 aaRSs that use the full anticodon as a primary identity element for aminoacylation (Asn, Asp, Cys, Gln, His, Ile, Lys, Met, Phe, Thr and Trp) [17] and for native anticodon sequences. Anticodon cognate aminoacylation may account for differences between tRNAAla tRNA Ser variants since the unique variable arm of tRNA Ser is likely to negate interaction with the anticodon cognate aaRSs.

## **Decoding specificity of anticodon variants**

In addition to decoding their exact Watson-Crick base pairing codons, expanded decoding by tRNAs occurs through G:U wobble pairing and modification of base 34 [84]. Our dataset allowed us to investigate the in vivo decoding specificities for mistranslating tRNAAla anticodon variants. For most variants with G, C or U at position 34, the Watson-Crick cognate codon was mistranslated the most frequently. Variants with A34 mistranslated codons ending in C and U and to a lesser extent A consistent with A34 being modified to inosine [60,85]. The exception was the ACC(Glv) anticodon variant, which is not native in yeast. Interestingly, we only detected mistranslation at codons ending in A for the AGG(Pro) variant. The need for mechanisms to regulate decoding by variants modified to have I34 may reflect that for two-box tRNA families, I:A decoding would result in translation across the box. It is possible that I:A decoding is minimized in non-cognate circumstances to prevent across the box mistranslation should the A34 arise through mutation of native two-box tRNAs. Consistent with this, we did not observe across the box alanine mis-incorporation for two-box anticodon variants with A34.

Anticodon variants with G or U at position 34 also exhibited expanded decoding. For 10 of 13 variants with G at position 34, codons ending in U were mistranslated. Conversely, only the UCG(Arg) variant decoded codons ending in G. This agrees with previous reports that G:U (anticodon:codon) pairs are more favourable than U:G pairs, unless U is modified [58,74,86,87]. For two variants with U at position 34 (UGG(Pro) and UAC(Val) anti-codons), we observed decoding of codons ending in U. This suggests 'superwobble' where anticodons with U at position 34 decode codons ending in all four nucleotides [88] and is consistent with the ability of S. cerevisiae tRNA Pro UGG to decode all four proline codons [89].

## **Acknowledgments**

We would like to thank Josh Isaacson and Josephine Davey-Young for critically reading the manuscript and Patrick O'Donoghue, Tallulah Andrews and members of the Villén lab for insightful discussions.

### **Disclosure statement**

No potential conflict of interest was reported by the author(s).

## **Funding**

This work was supported from the Natural Sciences and Engineering Research Council of Canada [RGPIN-2015-04394 to C.J.B.] and generous donations from Graham Wright and James Robertson to M.D.B. Mass spectrometry work was supported by a research grant from the Keck Foundation, National Institutes of Health grants RM1HG010461, R35 GM119536 and associated instrumentation supplement (to J.V.).

### **Author contributions**

E.C., C.J.B. and M.D.B. conceived and developed the project. E.C. and J. G. cloned the tRNA variants. E.C. conducted the growth experiments, analysed the data, identified factors correlating with mistranslation abundance and growth impact and created Figures 1-4 and associated supplemental figures. M.D.B. conducted the mass spectrometry experiments with assistance from R.A.R.-M., analysed the data and created Figures 5-7 and associated supplemental figures. M.R. constructed multicopy variants. M. R. and M.D. participated in data analysis. C.J.B. supervised the work and provided funding. J.V. supervised the mass spectrometry work and provided funding. M.D.B. supervised the project. E.C., C.J.B. and M.D.B. wrote the manuscript, and all authors edited the manuscript.

## Data availability statement

Raw images and data are available in the supplemental information. The raw mass spectrometry data have been deposited to the ProteomeXchange Consortium via the PRIDE [52] partner repository with the dataset identifier PXD038242, and an annotated list of the file names can be found in Supplemental Table S4.

### **ORCID**

Ecaterina Cozma Dhttp://orcid.org/0000-0003-4446-8302 Megha Rao (b) http://orcid.org/0000-0002-2096-1322 Madison Dusick (b) http://orcid.org/0000-0002-1958-8671 Ricard A. Rodriguez-Mias http://orcid.org/0000-0001-7960-6387 Judit Villén (b) http://orcid.org/0000-0002-1005-1739 Christopher J. Brandl http://orcid.org/0000-0001-8015-9668 Matthew D. Berg (b) http://orcid.org/0000-0002-7924-9241

### References

- [1] Chapeville F, Lipmann F, von Ehrenstein G, et al. On the role of soluble ribonucleic acid in coding for amino acids. Proc Natl Acad Sci. 1962;48:1086-1092. doi: 10.1073/pnas.48.6.1086
- [2] Ogle JM, Murphy FV, Tarry MJ, et al. Selection of tRNA by the ribosome requires a transition from an open to a closed form. Cell. 2002;111(5):721-732. doi: 10.1016/S0092-8674(02)01086-3
- [3] Loveland AB, Demo G, Grigorieff N, et al. Ensemble cryo-EM elucidates the mechanism of translation fidelity. Nature. 2017;546 (7656):113-117. doi: 10.1038/nature22397
- [4] Loftfield RB, Vanderjagt D. The frequency of errors in protein biosynthesis. Biochem J. 1972;128(5):1353-1356. doi: 10.1042/ bi1281353
- [5] Stansfield I, Jones KM, Herbert P, et al. Missense translation errors in Saccharomyces cerevisiae. J Mol Biol. 1998;282 (1):13-24. doi: 10.1006/jmbi.1998.1976
- Joshi K, Bhatt MJ, Farabaugh PJ. Codon-specific effects of tRNA anticodon loop modifications on translational misreading errors in the yeast Saccharomyces cerevisiae. Nucleic Acids Res. 2018;46 (19):10331-10339. doi: 10.1093/nar/gky664
- [7] Mordret E, Dahan O, Asraf O, et al. Systematic detection of amino acid substitutions in proteomes reveals mechanistic basis of ribosome errors and selection for translation fidelity. Mol Cell. 2019;75(3):427-441.e5. doi: 10.1016/j.molcel.2019.06.041
- [8] Gomes AC, Miranda I, Silva RM, et al. A genetic code alteration generates a proteome of high diversity in the human pathogen

- Candida albicans. Genome Biol. 2007;8(10):R206. doi: 10.1186/gb-2007-8-10-r206
- [9] Netzer N, Goodenbour JM, David A, et al. Innate immune and chemically triggered oxidative stress modifies translational Nature. 2009;462(7272):522-526. doi: 10.1038/ nature08576
- [10] Wiltrout E, Goodenbour JM, Fréchin M, et al. Misacylation of tRNA with methionine in Saccharomyces cerevisiae. Nucleic Acids Res. 2012;40(20):10494-10506. doi: 10.1093/nar/gks805
- [11] Lant JT, Berg MD, Sze DHW, et al. Visualizing tRNA-dependent mistranslation in human cells. RNA Biol. 2018;15(4-5):567-575. doi: 10.1080/15476286.2017.1379645
- [12] Santos M, Pereira PM, Varanda AS, et al. Codon misreading tRNAs promote tumor growth in mice. RNA 2018;15:773-786. doi: 10.1080/15476286.2018.1454244
- [13] Zimmerman SM, Kon Y, Hauke AC, et al. Conditional accumulation of toxic tRNAs to cause amino acid misincorporation. Nucleic Acids Res. 2018;46(15):7831-7843. doi: 10.1093/nar/gkv623
- [14] Berg MD, Zhu Y, Genereaux J, et al. Modulating mistranslation potential of tRNA Ser in Saccharomyces cerevisiae. Genetics. 2019;213(3):849-863. doi: 10.1534/genetics.119.302525
- [15] Rich A, RajBhandary UL. Transfer RNA: molecular structure, sequence, and properties. Annu Rev Biochem. (1):805-860. doi: 10.1146/annurev.bi.45.070176.004105
- [16] de Duve C. The second genetic code. Nature. 1988;333 (6169):117-118. doi: 10.1038/333117a0
- [17] Giegé R, Eriani G. The tRNA identity landscape for aminoacylation and beyond. Nucleic Acids Res. 2023;51(4):1528-1570. doi: 10.1093/nar/gkad007
- [18] Hou YM, Schimmel P. A simple structural feature is a major determinant of the identity of a transfer RNA. Nature. 1988;333 (6169):140-145. doi: 10.1038/333140a0
- [19] Hou YM, Schimmel P. Evidence that a major determinant for the identity of a transfer RNA is conserved in evolution. Biochemistry. 1989;28(17):6800-6804. doi: 10.1021/bi00443a003
- [20] Asahara H, Himeno H, Tamura K, et al. Escherichia coli seryl-tRNA synthetase recognizes tRNA Ser by its characteristics tertiary structure. J Mol Biol. 1994;236(3):738-748. doi: 10.1006/ jmbi.1994.1186
- [21] Biou V, Yaremchuk A, Tukalo M, et al. The 2.9 Å crystal structure of T. thermophilus seryl-tRNA synthetase complexed with tRNA<sup>Ser</sup>. Science. 1994;263(5152):1404-1410. doi: 10.1126/ science.8128220
- [22] Asahara H, Himeno H, Tamura K, et al. Discrimination among E. coli tRNAs with a long variable arm. Nucleic Acids Symp Ser. 1993;29:207-208.
- [23] Soma A, Kumagai R, Nishikawa K, et al. The anticodon loop is a major identity determinant of Saccharomyces cerevisiae tRNA Leu. J Mol Biol. 1996;263(5):707-714. doi: 10.1006/jmbi. 1996.0610
- [24] Breitschopf K, Achsel T, Busch K, et al. Identity elements of human tRNA<sup>Leu</sup>: structural requirements for converting human tRNA Ser into a leucine acceptor in vitro. Nucleic Acids Res. 1995;23(18):3633-3637. doi: 10.1093/nar/23.18.3633
- [25] Berg MD, Hoffman KS, Genereaux J, et al. Evolving mistranslating tRNAs through a phenotypically ambivalent intermediate in Saccharomyces cerevisiae. Genetics. 2017;206(4):1865-1879. doi: 10.1534/genetics.117.203232
- [26] Geslain R, Cubells L, Bori-Sanz T, et al. Chimeric tRNAs as tools to induce proteome damage and identify components of stress responses. Nucleic Acids Res. 2010;38(5):e30. doi: 10.1093/nar/ gkp1083
- [27] Woese CR. On the evolution of the genetic code. Proc Natl Acad Sci U S A. 1965;54(6):1546-1552. doi: 10.1073/pnas.54.6.1546
- [28] Woese CR. A new biology for a new century. Microbiol Mol Biol Rev. 2004;68(2):173-186. doi: 10.1128/MMBR.68.2.173-186.2004
- [29] Isaacson JR, Berg MD, Charles B, et al. A novel mistranslating tRNA model in Drosophila melanogaster has diverse, sexually dimorphic effects. G3 Bethesda Md. 2022;12(5):jkac035. doi: 10. 1093/g3journal/jkac035

- [30] Reverendo M, Soares AR, Pereira PM, et al. TRNA mutations that affect decoding fidelity deregulate development and the proteostasis network in zebrafish. RNA Biol. 2014;11(9):1199-1213. doi: 10.4161/rna.32199
- [31] Feany MB, Bender WW. A Drosophila model of Parkinson's disease. Nature. 2000;404(6776):394-398. doi: 10.1038/35006074
- Song L, He Y, Ou J, et al. Auxilin underlies progressive locomotor deficits and dopaminergic neuron loss in a Drosophila model of Parkinson's disease. Cell Rep. 2017;18(5):1132-1143. doi: 10.1016/ i.celrep.2017.01.005
- [33] Aggarwal A, Reichert H, VijayRaghavan K. A locomotor assay reveals deficits in heterozygous Parkinson's disease model and proprioceptive mutants in adult Drosophila. Proc Natl Acad Sci U S A. 2019;116(49):24830–24839. doi: 10.1073/pnas.1807456116
- [34] Berg MD, Giguere DJ, Dron JS, et al. Targeted sequencing reveals expanded genetic diversity of human transfer RNAs. RNA Biol. 2019;16(11):1574-1585. doi: 10.1080/15476286.2019.1646079
- [35] Parisien M, Wang X, Pan T. Diversity of human tRNA genes from the 1000-genomes project. RNA Biol. 2013;10(12):1853-1867. doi: 10.4161/rna.27361
- [36] Lant JT, Kiri R, Duennwald ML, et al. Formation and persistence of polyglutamine aggregates in mistranslating cells. Nucleic Acids Res. 2021;49(20):11883-11899. doi: 10.1093/nar/gkab898
- [37] Temple GF, Dozy AM, Roy KL, et al. Construction of a functional human suppressor tRNA gene: an approach to gene therapy for βthalassaemia. Nature. 1982;296(5857):537-540. doi: 10.1038/ 296537a0
- [38] Ko W, Porter JJ, Sipple MT, et al. Efficient suppression of endogenous CFTR nonsense mutations using anticodon-engineered transfer RNAs. Mol Ther Nucleic Acids. 2022;28:685-701. doi: 10.1016/j.omtn.2022.04.033
- [39] Porter JJ, Heil CS, Lueck JD. Therapeutic promise of engineered nonsense suppressor tRNAs. Wiley Interdiscip Rev RNA. 2021;12 (4):e1641. doi: 10.1002/wrna.1641
- [40] Rodriguez-Mias RA, Hess KN, Ruiz BY, et al. Proteome-wide identification of amino acid substitutions deleterious for protein function. bioRxiv. 2022 April;pre-print: not peer-reviewed. doi: 10.1101/2022.04.06.487405:9
- [41] Brachmann CB, Davies A, Cost GJ, et al. Designer deletion strains derived from Saccharomyces cerevisiae S288C: a useful set of strains and plasmids for PCR-mediated gene disruption and other applications. Yeast. 1998;14(2):115-132. doi: 10.1002/ (SICI)1097-0061(19980130)14:2<115:AID-YEA204>3.0.CO;2-2
- [42] Hughes TR, Marton MJ, Jones AR, et al. Functional discovery via a compendium of expression profiles. Cell. 2000;102(1):109-126. doi: 10.1016/S0092-8674(00)00015-5
- [43] Berg MD, Genereaux J, Zhu Y, et al. Acceptor stem differences contribute to species-specific use of yeast and human tRNA ser. Genes. 2018;9(12):E612. doi: 10.3390/genes9120612
- [44] Sprouffske K, Wagner A. Growthcurver: an R package for obtaining interpretable metrics from microbial growth curves. BMC Bioinf. 2016;17(1):172. doi: 10.1186/s12859-016-1016-7
- Schneider CA, Rasband WS, Eliceiri KW. NIH image to ImageJ: 25 years of image analysis. Nat Methods. 2012;9(7):671-675. doi: 10.1038/nmeth.2089
- [46] Berg MD, Isaacson JR, Cozma E, et al. Regulating expression of mistranslating tRNAs by readthrough RNA polymerase II transcription. ACS Synth Biol. 2021;10(11):3177-3189. doi: 10. 1021/acssynbio.1c00461
- [47] Rappsilber J, Mann M, Ishihama Y. Protocol for micro-purification, enrichment, pre-fractionation and storage of peptides for proteomics using StageTips. Nat Protoc. 2007;2 (8):1896-1906. doi: 10.1038/nprot.2007.261
- Leutert M, Rodríguez-Mias RA, Fukuda NK, et al. R2-P2 rapidrobotic phosphoproteomics enables multidimensional cell signaling studies. Mol Syst Biol. 2019;15(12):e9021. doi: 10.15252/msb. 20199021
- [49] Eng JK, Jahan TA, Hoopmann MR. Comet: an open-source MS/ MS sequence database search tool. Proteomics. 2013;13(1):22-24. doi: 10.1002/pmic.201200439

- [50] Käll L, Canterbury JD, Weston J, et al. Semi-supervised learning for peptide identification from shotgun proteomics datasets. Nat Methods. 2007;4(11):923-925. doi: 10.1038/nmeth1113
- [51] Argentini A, Goeminne LJE, Verheggen K, et al. moFF: a robust and automated approach to extract peptide ion intensities. Nat Methods. 2016;13(12):964-966. doi: 10.1038/nmeth.4075
- Perez-Riverol Y, Bai J, Bandla C, et al. The PRIDE database resources in 2022: a hub for mass spectrometry-based proteomics evidences. Nucleic Acids Res. 2022;50(D1):D543-52. doi: 10.1093/ nar/gkab1038
- [53] Grantham R. Amino acid difference formula to help explain protein evolution. Science. 1974;185(4154):862-864. doi: 10. 1126/science.185.4154.862
- [54] Veljkovic V, Cosic I, Dimitrijevic LD. Is it possible to analyze DNA and protein sequences by the methods of digital signal processing? IEEE Trans Biomed Eng. 1985;BME-32(5):337-341. doi: 10.1109/TBME.1985.325549
- [55] Prabhakaran M. The distribution of physical, chemical and conformational properties in signal and nascent peptides. Biochem J. 1990;269(3):691-696. doi: 10.1042/bj2690691
- [56] Fodje MN, Al-Karadaghi S. Occurrence, conformational features and amino acid propensities for the  $\pi$ -helix. Protein Eng Des Sel. 2002;15(5):353-358. doi: 10.1093/protein/15.5.353
- Huh W-K, Falvo JV, Gerke LC, et al. Global analysis of protein localization in budding yeast. Nature. 2003;425(6959):686-691. doi: 10.1038/nature02026
- Grosjean H, Westhof E. An integrated, structure- and energy-based view of the genetic code. Nucleic Acids Res. 2016;44(17):8020-8040. doi: 10.1093/nar/gkw608
- [59] Then A, Mácha K, Ibrahim B, et al. A novel method for achieving an optimal classification of the proteinogenic amino acids. Sci Rep. 2020;10(1):15321. doi: 10.1038/s41598-020-72174-5
- [60] Crick FH. Codon—anticodon pairing: the wobble hypothesis. J Mol Biol. 1966;19(2):548-555. doi: 10.1016/S0022-2836(66)80022-0
- [61] Torres AG, Piñeyro D, Filonava L, et al. A-to-I editing on tRNAs: biochemical, biological and evolutionary implications. FEBS Lett. 2014;588(23):4279-4286. doi: 10.1016/j.febslet.2014.09.025
- [62] Senger B, Auxilien S, Englisch U, et al. The modified wobble base inosine in yeast tRNA le is a positive determinant for aminoacylation by isoleucyl-tRNA synthetase. Biochemistry. 1997;36 (27):8269-8275. doi: 10.1021/bi970206l
- [63] Yona AH, Bloom-Ackermann Z, Frumkin I, et al. tRNA genes rapidly change in evolution to meet novel translational demands. Elife. 2013;2:e01339. doi: 10.7554/eLife.01339
- [64] Koonin EV, Novozhilov AS. Origin and evolution of the genetic code: the universal enigma. IUBMB Life. 2009;61(2):99-111. doi: 10.1002/iub.146
- [65] Ledoux S, Uhlenbeck OC. Different aa-tRNAs are selected uniformly on the ribosome. Mol Cell. 2008;31(1):114-123. doi: 10. 1016/j.molcel.2008.04.026
- [66] Fahlman RP, Dale T, Uhlenbeck OC. Uniform binding of aminoacylated transfer RNAs to the ribosomal A and P sites. Mol Cell. 2004;16(5):799-805. doi: 10.1016/j.molcel.2004.10.030
- [67] Wohlgemuth I, Pohl C, Rodnina MV. Optimization of speed and accuracy of decoding in translation. EMBO J. 2010;29 (21):3701-3709. doi: 10.1038/emboj.2010.229
- [68] Kothe U, Rodnina MV. Codon reading by  $tRNA^{Ala}$  with modified uridine in the wobble position. Mol Cell. 2007;25(1):167-174. doi: 10.1016/j.molcel.2006.11.014
- Olejniczak M, Uhlenbeck O. tRNA residues that have coevolved with their anticodon to ensure uniform and accurate codon recognition. Biochimie. 2006;88(8):943-950. doi: 10.1016/j.biochi.2006.06.005
- Boccaletto P, Machnicka MA, Purta E, et al. MODOMICS: a database of RNA modification pathways. 2017 update. Nucleic Acids Res. 2018;46(D1):D303-7. doi: 10.1093/nar/gkx1030
- [71] Yarian C, Townsend H, Czestkowski W, et al. Accurate translation of the genetic code depends on tRNA modified nucleosides.

- J Biol Chem. 2002;277(19):16391-16395. doi: 10.1074/jbc. M200253200
- [72] Konevega AL, Soboleva NG, Makhno VI, et al. Purine bases at position 37 of tRNA stabilize codon-anticodon interaction in the ribosomal a site by stacking and Mg<sup>2+</sup>-dependent interactions. RNA. 2004;10(1):90-101. doi: 10.1261/rna.5142404
- [73] Jenner LB, Demeshkina N, Yusupova G, et al. Structural aspects of messenger RNA reading frame maintenance by the ribosome. Nat Struct Mol Biol. 2010;17(5):555-560. doi: 10.1038/nsmb.1790
- [74] Pernod K, Schaeffer L, Chicher J, et al. The nature of the purine at position 34 in tRNAs of 4-codon boxes is correlated with nucleotides at positions 32 and 38 to maintain decoding fidelity. Nucleic Acids Res. 2020;48(11):6170-6183. doi: 10.1093/nar/gkaa221
- [75] Breslow DK, Cameron DM, Collins SR, et al. A comprehensive strategy enabling high-resolution functional analysis of the yeast genome. Nat Methods. 2008;5(8):711-718. doi: 10.1038/nmeth. 1234
- [76] Berg MD, Zhu Y, Ruiz BY, et al. The amino acid substitution affects cellular response to mistranslation. G3 Bethesda. 2021;11 (10):jkab218. doi: 10.1093/g3journal/jkab218
- Stein KC, Frydman J. The stop-and-go traffic regulating protein biogenesis: how translation kinetics controls proteostasis. J Biol Chem. 2019;294(6):2076-2084. doi: 10.1074/jbc.REV118.002814
- Knight JRP, Garland G, Pöyry T, et al. Control of translation elongation in health and disease. Dis Model Mech. 2020;13(3): dmm043208. doi: 10.1242/dmm.043208
- [79] Melnikov S, Mailliot J, Rigger L, et al. Molecular insights into protein synthesis with proline residues. EMBO Rep. 2016;17 (12):1776-1784. doi: 10.15252/embr.201642943
- Gamble CE, Brule CE, Dean KM, et al. Adjacent codons act in concert to modulate translation efficiency in yeast. Cell. 2016;166 (3):679-690. doi: 10.1016/j.cell.2016.05.070
- [81] Ghoneim DH, Zhang X, Brule CE, et al. Conservation of location of several specific inhibitory codon pairs in the Saccharomyces sensu stricto yeasts reveals translational selection. Nucleic Acids Res. 2019;47(3):1164-1177. doi: 10.1093/nar/gky1262
- [82] Murray LE, Rowley N, Dawes IW, et al. A yeast glutamine tRNA signals nitrogen status for regulation of dimorphic growth and sporulation. Proc Natl Acad Sci U S A. 1998;95(15):8619-8624. doi: 10.1073/pnas.95.15.8619
- [83] Tate JJ, Rai R, Cooper TG. Nitrogen starvation and TorC1 inhibition differentially affect nuclear localization of the Gln3 and Gat1 transcription factors through the rare glutamine tRNA<sub>CUG</sub> in Saccharomyces cerevisiae. Genetics. 2015;199(2):455-474. doi: 10. 1534/genetics.114.173831
- [84] Agris PF, Eruysal ER, Narendran A, et al. Celebrating wobble decoding: half a century and still much is new. RNA Biol. 2018;15(4-5):537-553. doi: 10.1080/15476286.2017.1356562
- [85] Basilio C, Wahba AJ, Lengyel P, et al. Synthetic polynucleotides and the amino acid code. Proc Natl Acad Sci U S A. 1962;48 (4):613-616. doi: 10.1073/pnas.48.4.613
- [86] Johansson MJO, Esberg A, Huang B, et al. Eukaryotic wobble uridine modifications promote a functionally redundant decoding system. Mol Cell Biol. 2008;28(10):3301-3312. doi: 10.1128/MCB.
- [87] Rozov A, Westhof E, Yusupov M, et al. The ribosome prohibits the G•U wobble geometry at the first position of the codonanticodon helix. Nucleic Acids Res. 2016;44:6434-6441. doi: 10. 1093/nar/gkw431
- [88] Rogalski M, Karcher D, Bock R. Superwobbling facilitates translation with reduced tRNA sets. Nat Struct Mol Biol. 2008;15 (2):192-198. doi: 10.1038/nsmb.1370
- [89] Bloom-Ackermann Z, Navon S, Gingold H, et al. A comprehensive tRNA deletion library unravels the genetic architecture of the tRNA pool. PLoS Genet. 2014;10(1): e1004084. doi: 10.1371/journal.pgen.1004084