Observing the fate of tRNA and its modifications by nucleic acid isotope labeling mass spectrometry: NAIL-MS

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ABSTRACT
RNA in yeast, especially rRNA and tRNA are heavily modified to fulfill their function in protein translation. Using biosynthetic stable isotope labeled internal standards we quantified 12 modified nucleosides in tRNA from S. cerevisiae over 24 hours. We observed different quantities of modified nucleosides in dependence of the growth phase. To elucidate the underlying mechanism of the observed tRNA modification profile adaptation, it is necessary to distinguish the pre-existing tRNA pool and its modifications from newly-synthesized tRNAs. By combination of 2 differentially isotope labeled media we developed NAIL-MS, nucleic acid isotope labeling coupled mass spectrometry. During the yeast growth cycle we observe dilution of pre-existing tRNAs by newly-synthesized tRNAs by the growing number of cells. tRNA was found to be highly stable with only little degradation over the observed period. The method was further used to quantify the levels of modified nucleosides in the original and new tRNA pools. By addition of deuterium-labeled methionine, we could observe the incorporation of new methyl marks on pre-existing tRNAs. For 2'-O-methylcytidine (Cm) we observed a global increase in log phase. We identified extensive 2'-OH-cytidine methylation of the pre-existing tRNAs and the new tRNAs which masks an actual decrease of pre-existing Cm. In contrast, global 5-methylcytidine (m5C) levels decreased during growth due to a drop in m5C quantities in the original tRNA pool.

The NAIL-MS data suggests different mechanisms for tRNA modification adaptation depending on the individual modification observed. With this new tool it is possible to follow the fate of methylated RNAs during growth and potentially compare the impact of different stress conditions on the epitranscriptome.

Introduction

The central dogma of molecular biology states that DNA is the storage of the genetic code, which is transcribed into (messenger) RNA, which is translated into proteins by another (transfer) RNA. This fundamental life process is dominated by nucleic acids, which are composed of the 4 canonical nucleosides cytidine, guanosine, adenosine and thymidine in DNA and uridine in RNA, respectively. The sequence of these building blocks defines the genetic code which contains all information necessary for each organism. While DNA is the storage of the genetic code, RNAs can be found as regulators and effectors of all essential life processes. Especially tRNA (tRNA) and rRNA (rRNA) are key players in protein translation and therefore highly abundant. To fulfill their function, both tRNA and rRNA are extensively modified and across all domains of life more than 150 modified RNA nucleosides have been identified.1 In recent years, the term epitranscriptomics has emerged to describe the modification level and profile of RNAs. The modification is introduced by dedicated RNA modifying enzymes which are referred to as RNA writers. Common RNA writers include methyltransferases, thiotransferases and pseudouridine-synthases and each writer has a specific substrate RNA. While modifications in tRNA and rRNA have been known for decades, modifications in mRNA have only recently been described.2,3 Alongside mRNA writer enzymes, so-called RNA erasers have been identified which actively demethylate the methylated adenosine in mRNA.

Like all organisms analyzed to date, S. cerevisiae tRNAs are heavily modified as recently reviewed.4 Over 25 different modified nucleosides have been identified in these tRNAs, of which 12 are mostly S-adenosyl methionine methylated derivatives of the canonical nucleosides. The quantities of tRNA modifications are known to adapt during cell growth5 and during stress.6 Furthermore they are essential for protein homeostasis and cell morphogenesis.7 The dynamic adaptation of tRNA modifications prompts to ask the question how tRNA modification levels are changed from a mechanistic viewpoint. One possibility is an addition of modifications by tRNA writers to the already existing tRNA pool and the newly synthesized tRNA which results in increased tRNA modification levels. A decrease of tRNA modification levels might be achieved by 2 possible scenarios. The first includes the participation of tRNA erasers that actively remove existing modifications from the tRNA. While RNA erasers are known to exist for mRNAs, only recently ALKBH1 was observed...
in vitro to demethylate 1-methyladenosine to adenosine in certain tRNAs.8 Another potential way to remove RNA modifications is degradation of the existing, modified RNA and synthesis of new, unmodified RNA. However, current methods cannot distinguish the modification profile of the existing RNA pool and the new RNA pool. Here we present a new technique, NAIL-MS (nucleic acid isotope labeling coupled mass spectrometry) which allows observation of modification dynamics in existing and newly synthesized RNAs from S. cerevisiae during growth.

Results

**tRNA modification quantities fluctuate during growth**

tRNA from yeast was purified from total RNA by size-exclusion chromatography (SEC) as described previously9 and the tRNA fraction collected. Fig. 1 shows an exemplary chromatogram of total RNA before SEC and purified tRNA after SEC. The large RNA is a mixture of rRNA and other large RNA molecules like mRNA (Fig. S1). The isolated tRNA was digested to nucleosides and subjected to liquid chromatography coupled mass spectrometry (LC-MS). An exemplary chromatogram for detection of the canonical nucleosides and 13 abundant modified nucleosides can be found in Fig. 1. The less common modifications like 2′-O-methyluridine (Um) or wobble-uridines like 5-methoxy carbonylmethyl-2-thiouridine (mc5’s2U) were not included in the current study due to low detection sensitivity of uridine derivatives. Dihydrouridine was not available as a synthetic standard and could not be included in quantification analysis. Using a biosynthetic stable isotope labeled internal standard (SILIS), produced according to10 and calibration with synthetic standards, we quantified the absolute level of modification per tRNA in log phase growing cells (calibration curves in Fig. S2). Fig. S3 shows a scheme how RNA modifications are quantified and which calculations and normalization steps are necessary to arrive at the number of a given modification per tRNA.

In total tRNA from log growing cells we find pseudouridine (Ψ) as the most abundant modified nucleoside, followed by the methylated nucleosides 5-methylcytidine (m5C, 1-methylguanosine (m1G) and 5-methylguanosine (m5G) as expected (compare Table S1 prepared from11). Here, the modification profile of 34 tRNA sequences is listed. For total tRNA the exact number of modified nucleosides depends on the ratio of the isoacceptors and isodecoders. This ratio is dynamic and used as an adaptation mechanism to overcome changes in growth conditions and external stress.12,13 This adaptation and the unknown modification status of most tRNAs make it difficult to assign a quantitative number of modifications to the total tRNA pool.

During these initial experiments to quantify tRNA modifications in S. cerevisiae BY4741 we noticed fluctuations in the quantities of many modifications in dependence of the growth phase (Fig. S4). 2′-O-methylcytidine (Cm), 1-methylguanosine (m1G) and 7-methylguanosine (m7G) levels increased in log phase (Fig. 1 and Fig. S4). In contrast, we observed a drop in 5-methylcytidine (m5C), 5-methyluridine (m5U) and pseudouridine (Ψ) quantities in log phase. For modified nucleosides 2′-O-methylguanosine (Gm), 2′-O-methyladenosine (Am), 2-methylguanosine (m2G), 2,2-dimethylguanosine (m22G), inosine (I) and 1-methyladenosine (m1A) the levels stayed constant throughout growth (Fig. S4).

While we cannot explain this phenomenon at the moment we were curious to understand how the levels of modifications are de- or increased mechanistically. This prompts the question whether a change of modification quantity is caused by hyper- or hypomodified new tRNAs while the pre-existing tRNAs stay unaltered or by dynamic adaptation of the pre-existing tRNAs.

**Developing media for nucleic acid isotope labeling**

Intrigued by these results we wanted to design an isotope based assay which allows discrimination of the original tRNA pool and newly synthesized tRNAs by mass spectrometry. For this

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**Figure 1.** Workflow to determine modification levels in tRNA during yeast growth cycle. The isolated total RNA was purified by size exclusion chromatography (SEC) (‘5.8S/5S tRNA peak, #unknown contamination). The tRNA fraction was then digested and the obtained nucleosides separated and quantified over LC-MS to receive the shown chromatogram on the upper right side. The average amounts of the respective modifications per molecule tRNA in log growing cells were calculated and plotted (error bars represent SD of n = 3). The quantification was repeated during growth (arrows indicate sample time points) and the quantities of the modifications Cm and m7G were plotted over time. Abbreviations: D = dihydrouridine, Ψ = pseudouridine, ζ = cytidine, m1A = 1-methyladenosine, U = uridine, m2G = 5-methylcytidine, m5G = 7-methylguanosine, Cm = 2′-O-methylcytidine, I = inosine, G = guanosine, m5U = 5-methyluridine, m2G = 1-methylguanosine, Gm = 2′-O-methylguanosine, m22G = N2-methylguanosine, A = adenosine, m6G = N2,N2-dimethylguanosine, Am = 2′-O-methyladenosine.
type of experiments it is necessary to have 2 growth media that result in differential labeling and therefore different masses of the nucleosides. We have tested commercial rich growth medium (Silantes, Munich, Germany) which allows efficient labeling of nucleobases with either $^{15}$N or $^{13}$C while the ribose remains unlabeled when supplemented with regular $^{12}$C-glucose (see Fig. S5). In these media it is possible to introduce additional heavy carbons in the ribose by supplementing with $^{13}$C-glucose. Furthermore it is possible to supplement with D$_3$-methionine which resulted in a 60-70% labeling efficiency of methylated nucleosides with D$_3$-labeled methyl groups (see Fig. S6). The commercial medium was chosen for preparation of the stable isotope labeled internal standard (SILIS) to receive a max. labeled internal standard. The SILIS nucleosides have a +9 mass shift for uridine and cytidine, and +12 for methylated uridine and cytidine modifications (see Table S2). The purines, guanosine and adenosine, have a +10 while their methylated derivatives have a +13 mass shift (except m$_2$G with a mass shift of +16).

For the time course experiments, the commercial media was not suitable since some modifications like 5-methylcytidine would have formed isotopomers with the same m/z as its corresponding SILIS nucleoside. Therefore, we tested the labeling efficiency in minimal medium (YNB) supplemented with isotope labeled amino acids, glucose and nucleobases. With D$_3$-methionine, labeling of methylated nucleosides was achieved quickly (increase in deuterated over unlabeled nucleosides shown in Fig. S7). As S. cerevisiae BY4741 is a uracil auxotroph strain, we supplemented with $^{15}$N-uracil which resulted in a complete labeling of pyrimidines and their modifications. For purine labeling we added the purine biosynthesis precursors $^{15}$N-amide labeled glutamin or $^{15}$N-aspartic acid. Although these amino acids are the nitrogen source of purines, the incorporation efficiency was very low (Fig. S8). However, a quantitative +1 purine base label could be achieved by growth in $^{15}$C$_6$-glucose supplemented YNB medium which results in a purine nucleoside mass increase of +6 (+1 from base and +5 from ribose, Fig. S8 and Fig. S9) and +5 for pyrimidines (+5 from ribose). From these results a combination of $^{15}$N-uracil with $^{13}$C$_6$-glucose minimal medium was chosen as the first medium for our experiments. The nucleoside and nucleobase m/z values and their mass increase compared with unlabeled nucleosides for the canonnals are found in Table 1 and Fig. S10.

For the second medium we decided to use YNB minimal medium but supplemented with regular uracil ($^{14}$N-uracil) and regular $^{13}$C$_6$-glucose which results in unlabeled canonnals (Fig. 2a). By combination of these 2 media we can now distinguish the pre-existing nucleosides (heavy $^{15}$N/$^{13}$C-label) and the newly synthesized tRNA nucleosides (unlabeled). To further allow a detailed analysis of the pre-existing tRNA modifications, we added D$_3$-methionine to the growth medium which led to the incorporation of heavy methyl groups after medium exchange (Fig. 2b for newly synthesized modifications). In combination with the differential labeling of the nucleoside structure, this step allows to detect newly methylated nucleosides in the pre-existing tRNA pool by a mass increase of +3 (Fig. 2d and Fig. S10) compared with the pre-existing methylated nucleosides. We refer to this event as post-methylation as it happens after experiment initiation.

**NAIL-MS (nucleic acid isotope labeling coupled mass spectrometry) reveals dilution of the pre-existing tRNA pool**

In a next step we combined the 2 growth media in a single assay. The principle is shown in Fig. 3a. The cells are grown overnight in $^{15}$N-uracil/$^{13}$C$_6$-glucose supplemented YNB medium. To achieve complete labeling of all nucleosides with the heavy label the cells were centrifuged, resuspended in fresh $^{15}$N/$^{13}$C YNB medium and grown overnight a second time. At this step high labeling efficiency of all nucleosides is required which was verified by high-resolution mass spectrometry (see Fig. S9 and S10). Minor signals of unlabeled nucleosides are due to impurities of the $^{15}$C$_6$-glucose and $^{15}$N-uracil starting material (99% ATOM $^{13}$C and 98% ATOM $^{15}$N purity) which causes <5% of unlabeled nucleosides at experiment start. For experiment initiation the cells are briefly centrifuged and

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**Figure 2.** Mass spectra of guanosine (G) and 7-methylguanosine (m7G) in different growth media. a unlabeled medium: [M+H]$^+$ (G) 284, [M+H]$^+$ (m7G) 298. b D$_3$-methionine containing medium: [M+H]$^+$ (G) 284, [M+H]$^+$ (m7G) 301. c $^{15}$N$_2$ uracil and $^{13}$C$_6$ glucose containing medium: [M+H]$^+$ (G) 290, [M+H]$^+$ (m7G) 304. d $^{15}$N$_2$ uracil, $^{13}$C$_6$ glucose and D$_3$ methionine: [M+H]$^+$ (G) 290, [M+H]$^+$ (m7G) 307. e SILIS: [M+H]$^+$ (G) 294, [M+H]$^+$ (m7G) 311.
suspended in D$_3$-methionine supplemented YNB medium. Total RNA isolation and digestion to nucleosides followed by mass spectrometry revealed an increase of newly synthesized canonical nucleosides (light) from the new RNA pool during growth (Fig. S11, black line) while the original RNA was diluted (red). For modified nucleosides we performed mass spectrometry in scan mode after experiment initiation. As expected we could detect the signal of the new, D$_3$-methylated nucleosides and the H$_3$-methylated original nucleosides. However, we could also detect the predicted post-methylated nucleoside species with a mass increase of +3 from the original heavy isotope labeled RNA pool (Fig. 2d and Fig. S10). This nucleoside species is the result of additional methylation of canonicals that were not methylated in the pre-existing tRNA pool at experiment initiation. In summary, we can observe 3 different isotopomers of each methylated nucleoside (Fig. 3c) which originate from different RNA pools: 1) methylated nucleosides from the new RNA pool, 2) original nucleosides already methylated at experiment initiation and 3) original nucleosides, methylated after experiment initiation.

With this assay at hand we decided to grow yeast for 24 hours and observe the fate of canonical and modified nucleosides in tRNAs by sensitive triple quadrupole mass spectrometry. The mass transitions of canonical and modified nucleosides were programmed for each observed isotopomer into a multiple reaction monitoring (MRM) method (mass transitions in Table S2 and mass spectra in Fig. S9). The total RNA was isolated from 6 biologic replicates before experiment initiation (T = 0 hours), after 1, 2, 4 and 6 hours (lag phase and log phase) and 24 hours (stationary) as the last time point. Using size-exclusion chromatography we purified total tRNA and subjected all samples to the developed NAIL-MS method after digestion to nucleosides. By injection of tRNA digests we could quantify the amount of tRNA injected and separate the signal of the pre-existing and novel tRNA pool. This quantification revealed an initial degradation of the existing tRNAs in the first 60 to 120 minutes followed by constant levels thereafter (Fig. 3b). We did not observe further tRNA degradation which is consistent with the reported long half-life of tRNA.$^{15,16}$ In similar pulse chase studies using tritium labeled uracil and subsequent gel electrophoretic quantification of the radioactively labeled yeast tRNA, no initial drop in tRNA levels was observed.$^{17}$ While this difference might be caused by technical errors during our sample collection and RNA isolation, we currently investigate the possibility of stress-induced tRNA degradation caused by the necessary pelleting of yeast cells before experiment initiation.

**NAIL-MS reveals differences in the modification profile of the original and new tRNA pool**

Fig. 3d shows all signals of 7-methylguanosine (m$^7$G) which potentially occur in a NAIL-MS experiment after medium exchange. For quantification of newly synthesized m$^7$G, the signal for the unlabeled and D$_3$-labeled isotopomers were summed and normalized to the amount of unlabeled tRNA injected (calculated from the signal of unlabeled adenosine). Thereby we

**Figure 3.** Principle of the developed NAIL-MS assay and RNA species emerging over time. a Yeast is grown in heavy isotope medium which results in heavy labeled nucleosides. The experiment is initiated by medium exchange with D$_3$-methionine labeled medium. New canonicals are unlabeled while pre-existing nucleosides remain $^{15}$N/$^{13}$C labeled. b Original tRNA quantities stay constant over time (red), while new tRNAs are synthesized (black line) as cells grow (OD 600 gray line) (error bars are SD of n = 3). c Different modified nucleosides can be distinguished according to their labeling. New modified nucleosides have an unlabeled canonical building block and a D$_3$-methyl mark, original modified nucleosides derive from labeled canonicals but carry the unlabeled methylation and original, labeled canonicals which receive a methyl mark after experiment initiation have an additional D$_3$-methyl group. d Mass spectrum of potential isotopomers of 7-methylguanosine (m$^7$G) in a NAIL-MS experiment: $^{13}$unlabeled m$^7$G as a reference (298), from new tRNA (301), from original tRNA with pre-existing methylation (304), from original tRNA but from post-methylation (307) and the signal of the m$^7$G SILIS (311).
received the number of m7G per new tRNA. Similarly we added the signal of the original m7G isotopomer and the post-methylated m7G (heavy nucleoside + heavy D3-methyl group) and normalized to the amount of pre-existing tRNA in the sample (calculated from the signal of labeled adenosine).

The plots of Fig. 4 show the number of modifications per original tRNA or new tRNA, respectively, on the y-axis. In the unlabeled experiment (Fig. 1) we observed growth phase dependent fluctuations for 7-methylguanosine (m7G). By NAIL-MS we identified the reason for the observed increase in a slow and steady increase of m7G in new tRNAs during log phase (Fig. 4a). The number of m7G in the original tRNAs appears to stay constant throughout the experiment. However, a detailed analysis of the pre-existing tRNA pool reveals a constant decrease in the number of pre-existing m7G while new m7G is incorporated into the original tRNAs.

For 2'-O-methylcytidine (Cm) we observed a growth cycle dependent fluctuation of modification level (Figs. 1 and 4b). By NAIL-MS we observed an initial increase of Cm levels in pre-existing tRNAs while the new tRNA pool forms Cm rather slowly in the first 2 hours. While one explanation for the increase of Cm in the pre-existing tRNAs is degradation of those tRNAs that are not Cm modified during lag phase, the quantification of D3-labeled pre-existing cytidines revealed a substantial incorporation of new methyl marks in the pre-existing tRNAs (box of Fig. 4b). Curiously we observe a sudden decrease of pre-existing Cm in the original tRNA pool after entering log phase. This could be explained by several scenarios. The most likely explanation is sudden degradation of pre-existing tRNAs with Cm at a certain position (e.g. tRNAs with Cm at position 34, see Fig. S13) as these tRNAs might be no longer needed while entering log phase. In parallel, Cm might become necessary at the other positions of pre-existing tRNAs (e.g., at position 4) which leads to increased post-methylation. Another scenario for the decrease in pre-existing Cm is active demethylation of Cm by an unknown RNA eraser. Quantification of specific Cm-modified tRNAs by northern blotting might provide an answer to this open question.

The unmethylated nucleoside modifications pseudouridine and inosine are rapidly introduced into the nascent tRNA as these modifications reach their maximum abundance in the first 60 minutes (Fig. 4c and d). While new inosine levels in new tRNA are immediately comparable to the original tRNA inosine levels, pseudouridine levels are lower in the newly synthesized tRNAs compared with the levels in original tRNA. Furthermore, we observed a slight decrease of pseudouridine modification in the original tRNA pool after 6 hours, which might be linked to degradation of pseudouridine containing tRNAs. As both pseudouridine and inosine are non-methylated tRNA modifications, we cannot further distinguish between pre-existing modifications and those introduced after experiment initiation.

NAIL-MS results of the other methylated nucleosides are shown in Fig. S12. Like Cm, 5-methylcytidine (m5C) is found at various positions of tRNA (positions 34, 40, 48, 49) and has a pronounced drop in its abundance in the pre-existing tRNAs. This drop can be observed in both the pre-existing and post-methylated m5C molecules. The other methylated nucleosides Gm, m1A, m22G, m2G and Am are like inosine rather unresponsive with similar quantities throughout the experiment. Analysis of the incorporation speed into the nascent tRNA pool shows an almost immediate incorporation of m7G, m22G, Gm and Am while Cm, m2G, m1A, m2G, m3U and m3C are less abundant during lag phase compared with stationary and log phase.

For all methylated nucleoside modifications we observed incorporation of the heavy D3-methyl mark into the pre-existing tRNAs. To compare the extent of this delayed post-methylation we formed the ratio of e.g., post-methylated Cm to all Cm in original tRNA and plotted the % of the post-methylated

**Figure 4.** Levels of modifications in tRNA during a time range of 24 h discriminated by the original and newly-synthesized tRNA pool. *The quantity of modification was normalized to its respective origin (e.g., new m7G was normalized to abundance of new tRNA to receive the number of m7G per new tRNA while original m7G was normalized to original tRNA). A level of 7-methylguanosine (m7G) in new and original tRNAs over time. The constant level of original m7G is caused by a constant increase in pre-existing m7G, while post-methylated m7G increases (box). (error bars reflect the SE of n = 6) b 2’-O-methylcytidine (Cm) increases fast in newly synthesized tRNAs, while the level of pre-existing Cm drops (box). This drop is masked by substantial post-methylation of original tRNAs. (error bars reflect the SE of n = 6) c Pseudouridine quantities are lower in newly-synthesized tRNAs while d the number of inosine per respective tRNA is constant throughout the experiment (both n = 3)
modifications (Fig. 5). After 6 hours the maximum of post-methylation is reached for all methylated nucleosides. Except Cm, all nucleosides plateau at 10% post-methylation. Cm has a 1.8-fold higher plateau of post-methylation compared with the other methylated nucleosides, indicating the necessity to quickly adapt Cm levels in tRNAs to overcome lag phase.

Discussion

Using stable isotope labeling mass spectrometry with a biosynthetically produced internal standard we quantified 13 modified nucleosides in total tRNA from S. cerevisiae grown in minimal medium. The detected quantities from yeast log growing cells were lower compared with the expectation from a database search (Table S1). These differences are partly explained by the low number of tRNA sequences with reported modification profile in the database (only 34 tRNA sequences available) and the general error introduced by absolute quantification approaches. In our quantification we use an “averaged” tRNA generated from the database under the assumption of equimolar contribution of each listed tRNA sequence. Therefore, absolute quantification results might slightly deviate from the true number of modification per tRNA molecule. Another explanation can be found in the dynamic nature of tRNA modifications which adapt depending on the growth phase as observed in Fig. 1 and Fig. S4 or stress.6 Especially during log phase the quantities of modified nucleosides increased for Cm, m7G and m6G, while m5C, m5U and Ψ levels decreased and Gm, Am, m7G, m52G, I and m1A stayed constant. While the biology of these changes remains unclear, we have developed a new tool to separately analyze the methylated and non-methylated tRNAs (Fig. 4 and Fig. S12).

The mechanism, by which the modification level adapts, depends on the modified nucleoside and potentially the distribution in the RNA. For 7-methylguanosine (m7G, pos. 46) in tRNA, increased incorporation into newly synthesized tRNAs was observed. We observed a decrease in pre-existing m7G while post-methylated m7G increased, resulting in an apparent constant level of m7G in the pre-existing tRNA.

We made similar observations for 1-methylguanosine (m1G), 2,2-dimethylguanosine (m22G) and 2′O-methylguanosine (Gm) (see Fig. S12). While these methylated nucleosides showed a limited adaptation during growth, 2′O-methylcytidine (Cm, pos 34 and 4 of tRNAs) and 5-methylcytidine (m5C pos 34, 40, 48 and 49) quantities fluctuated depending on the growth phase. Cm showed an interesting modification profile over time which was partly caused by reduced modification levels in original tRNAs, high modification levels in the new tRNAs and post-methylation of the existing tRNA pool. Especially the reduced modification profile in the original tRNAs after 4 hours is intriguing (compare Fig. 4b and Fig. S12 for m5C after 6 hours). One potential explanation is degradation of Cm and m5C modified tRNAs (e.g., tRNA^Gly, tRNA^Phe or tRNA^Leu) which would result in a reduced number of these modifications in the original tRNAs. It is possible that the observed changes in the tRNA epitranscriptome reflect fluctuations in the ratio of isoacceptors and isodecoders. Another potential mechanism that was recently described for 1-methyladenosine (m1A) in human cell culture is active removal of the methylation mark by an RNA eraser. While our current data cannot distinguish these mechanisms, it highlights m5C and Cm as potential candidates for RNA erasers. Future analysis of single modifications within their respective tRNA subtype will reveal the underlying mechanism.

We interpret methylation of original canonicals as an addition of new methyl marks into existing, intact RNA to adapt the modification profile to the physiologically needed modification status. However, it is possible that the observed signal is caused by salvage of existing heavily-labeled canonicals which are introduced into newly synthesized tRNAs followed by subsequent methylation. However, if salvage was the main reason for added methylation to existing canonicals, the quantities of added methylation should be similar among the different modified nucleosides and the level would constantly increase over time as more and more RNA gets degraded and original canonicals get released. Both were not observed in our studies as the % of post-methylated nucleosides in the original tRNAs plateau after 6 hours. Although outside the scope of this manuscript, oligonucleotide mass spectrometry of e.g., a partial RNase T1 digest of NAIL-MS samples would allow to assess the origin of these methylated nucleosides.

We used the NAIL-MS method to analyze 13 modified nucleosides, 2 of those are non-methylated tRNA modifications. For non-methylated nucleosides it is not possible to determine the quantity of added modification to the original RNA pool with the developed assay. However, we could monitor the modification extent in new and original RNAs and compare their quantities of modification.
In principle, it is possible to monitor all methylated nucleosides in original and new RNA and even observe the addition of the methyl mark to original RNA. Especially methylated uridine modifications like 5-methoxycarbonylmethyl-2-thiouridine (mcm5S2U) or 5-methoxycarbonylmethyluridine (mcm5U) found at position 34 of tRNA would be interesting. It can be applied to heavily modified RNAs, as shown here for tRNA, but also to any other purified RNAs species like rRNA, mRNA or small non-coding RNAs.

**Materials and methods**

**Growth medium and conditions for S. cerevisiae BY4741**

*S. cerevisiae* BY4741 was grown in 13C labeled Yeast OD 2 (+ 10 g/L 13C6 Glucose (99% ATOM) (Euriso-top, Saclay, France)) rich medium (Silantes, Munich, Germany) for SILIS preparation or in YNB minimal medium for NAIL-MS experiments. 10x YNB (Carl Roth GmbH, Karlsruhe, Germany) was prepared according to manufacturer’s manual. 1x YNB (Carl Roth GmbH, Karlsruhe, Germany) was added to a final concentration of 1 mM in the digest solution. The resulting digest/theophylline mixture is referred to as 10x SILIS which was added to a final concentration of 1x to samples and calibration solutions.

**Calibration**

Synthetic modified nucleosides for preparation of calibration solutions were purchased from: Sigma-Aldrich, Munich, Germany: Nucleoside test mix: inosine (I), pseudouridine (Ψ), 5-methyluridine (m5U), 2'-O-methylcytidine (Cm), 5-methylcytidine (m5C); solid nucleosides: adenosine (A), 7-methylguanosine (m7G), 2'-O- methyladenosine (Am), 1-methyladenosine (m1A), 1-methylguanosine (m1G), 2-methylguanosine (m2G) and N2,N2-dimethylguanosin (m22G); solid nucleosides from Berry&Associates, Dexter, MI, USA: 2'-O-methyluridine (Um) and 2'-O-methylguanosine (Gm). Each nucleoside powder was weighed into a clean tube (5–10 mg per nucleoside) and dissolved in pure water to reach a final concentration of 10 mM. The nucleosides and the test mix were combined to a final concentration of 500 pmol/μL of A, 200–500 fmol/μL of each modified nucleoside and 1x of the prepared SILIS. The calibration mix was serial diluted 1:5 and 1:10 with 1x SILIS until the lowest calibration concentration of 5 fmol/μL canonical and 5 amol/μL modified nucleoside were reached. 10 μL of each calibration solution was injected onto LC-MS/MS. After LC-MS/MS measurement the value of the integrated MS signals of the unlabeled synthetic nucleosides were set into relation to the integrated MS signals of the heavier SILIS nucleosides. The results were plotted against the nucleoside concentrations and the regression lines from the diagrams (Fig. S2) were used to calculate the respective RFN values, necessary for quantification.

**NAIL-MS experiments**

Five mL of 15N and 13C-Glucose labeled medium was inoculated with a single *S. cerevisiae* BY4741 culture from an YPD-agar plate in a 50 mL falcon tube. The cells were grown in a shaking incubator over night at 30 °C and 250 rpm. The next day 1 mL of the lysate was centrifuged, the supernatant discarded and the pellet resolved in 5 mL fresh 15N and 13C-Glucose labeled medium to achieve better labeling. After incubation over night at 30°C and 250 rpm again, the OD was determined at 600 nm using an Eppendorf Biophotometer. For experiment initiation part of the culture was centrifuged at 3000 xg for 5 minutes at room temperature. The cell pellet was resuspended in 20 mL of D2O-labeled YNB medium to achieve a starting OD of 1. Using a 100 mL glass flask, the cells were incubated for 24 hours at 30 °C and 250 rpm in a shaking incubator. At set times (T = 0, 1, 2, 4, 6 and 24 hours) the OD was measured and a 2 mL sample was taken for RNA isolation. After centrifugation at 3000 xg for 5 minutes, the pellet was dissolved in 1 mL of TRI reagent® transferred into a screw-top tube with an amount of acid-washed beads equivalent to 200 μL and vortexed for 10 minutes. Isolation of RNA was done according to user manual. The experiment was repeated with 6 biologic replicates.

*tRNA isolation via SEC* and tRNA digestion below. The tRNA digest from 100 mL OD 6 culture was diluted in 1 mL pure water which resulted in mass spec peak areas 50–100x above the limit of quantification (LOQ) of the modified nucleosides of interest in a 1/10 dilution of the digest. As an external standard 10 mM theophylline was added to a final concentration of 1 mM in the digest solution.
260 nm, autosampler, column thermostat (60 °C) and fraction collector. A size-exclusion column (Agilent Bio SEC-3, 3μm, 300A, 7.8 × 300 mm, Agilent, Waldbronn, Germany) allowed collection of RNA fractions after isotropic elution with 100 mM ammonium acetate at pH 7. The tRNA fraction was concentrated to about 150 μL in a Speed-Vac (Christ, Osterode am Harz, Germany). 5 M NH₄OAc was added to a final concentration of 0.5 M and after addition of 2x Vol. ice-cold ethanol (100%) the RNA was precipitated over night at −20°C. After centrifugation at 12,000 xg for 30 minutes at 4°C the RNA pellet was subjected to an additional ethanol (80 %) wash step to verify complete removal of ammonium acetate and then resuspended in 20 μL pure water. The quality of the thus isolated tRNA was verified by analytical SEC using the same method.

**tRNA digestion**

tRNA was digested using 0.6 U nuclease P1 (Sigma-Aldrich) in the presence of ZnCl₂, ammonium acetate pH 5.3, THU, pentostatine and BHT from²¹ at 50°C for one hour followed by overnight incubation at room temperature, the next day, 10 U benzonase (Sigma-Aldrich, Munich, Germany), 0.1 U phosphodiesterase I (VWR, Ismaning, Germany) and MgCl₂ in a final concentration of 1 mM were added and incubated at 37°C for one hour. Then 20 U alkaline phosphatase (Sigma-Aldrich, Munich, Germany) and the proper amount to reach a final concentration of 0.1 M Tris-HCl pH 8 were added. After 2 hours at 37°C the digestion mix was filtered through a 10 kDa MWCO filter (AcroPrep™ Advance, 350 μL, Omega™ 10K MWCO, Pall, Dreieich, Germany) at 3000 xg for 30 minutes. 18 μL of filtrate were mixed with 2 μL of the 10x SILIS and subjected to LC-MS/MS analysis.

**High resolution mass spectrometry**

For determination of m/z precursor and product values 5 mL cultures of each growth medium was inoculated with a colony of *S. cerevisiae* BY4741. Total RNA was isolated at OD 6 and digested as described for NAIL-MS experiments. The ribonucleosides were separated using a Dionex Ultimate 3000 HPLC system on an Interchim Uptisphere120–3HDO C18. Mobile phase A was 2 mM ammonium acetate and mobile phase B was 80% acetonitrile containing 2 mM ammonium acetate. Gradient elution started with 0% B and increased to 12% B after 10 minutes and to 80% after 12 minutes. After 4 minutes elution at 80% B and subsequently regeneration of starting conditions to 100% A after 5 minutes, the column was equilibrated at 100% A for 8 minutes. The flow rate was 0.2 mL/minute and the column temperature 30°C. High-resolution mass spectra of precursor and product ions were recorded by a ThermoFinnigan LTQ Orbitrap XL.

**LC-MS / QQQ**

Ribonucleosides were separated using a Synergy Fusion RP, 2.5 μm particle size, 100 A pore size, 100 mm length, 2 mm inner diameter from Phenomenex (Torrance, CA), on an Agilent 1290 series HPLC system equipped with a diode array detector. Mobile phase A was 5 mM ammonium acetate adjusted to pH 5.3 with glacial acetic acid and mobile phase B was pure acetonitrile. Gradient elution started with 100% A for one minute, increase to 10% B after 10 minutes, 40% after 14 minutes and regeneration of starting conditions with 100% A for 3 additional minutes. The flow rate was 0.35 mL/minute and the column temperature 35°C. For mass spectrometric measurements an Agilent 6490 Triple Quadrupole mass spectrometer set to dynamic multiple reaction monitoring (MRM) mode was used. The MS was operated in positive ion mode with the following parameters: electro-spray ionization (ESI-MS, Agilent Jetstream), Fragmentor Voltage (set in tunefile to) 250 V, Cell Accelerator Voltage 2 V, N2-Gas temperature 150°C, N2-Gas flow 15 L/min, Nebulizer 30 psi, Sheath gas (N2) temperature 275 °C, Sheath gas flow 11 l/min, Capillary 2500 V and Nozzle Voltage 500 V. The mass transition for each modified nucleoside and its isotopomers are found in Table S2.

**Data analysis**

Using Agilent’s Quantitative Data Analysis software, the mass signal areas of the different isotope labeled ribonucleosides were integrated. For methylated ribonucleosides the SILIS isotopomer, the ¹⁵N/¹³C labeled isotopomer (original), the ¹⁵N/¹³C/D₃ labeled isotopomer (original+new methyl mark) and the D₃ labeled isotopomer (new) could be identified. For the canonicals the isotopomers of the SILIS, the original and the unlabeled m/z transitions were analyzed to quantify the amount of injected tRNA. The values of the integrated signals were used for further calculations. The nucleosides were quantified by calculating the nucleoside isotope factor (NIF) and the response factors (RFN) were received by plotting the NIF over the concentration of injected synthetic standard.¹⁰

\[
\text{NIF} = \frac{\text{area}_{MS}(\text{isotopomer})}{\text{area}_{MS}(\text{SILIS})}
\]

The signal of adenosine in each sample was divided by the response factors and the amount of injected adenosine was received. The value for adenosine was divided by the number of adenosine per tRNA molecule (see also Table S1). Thereby the injected amount of original and new tRNA molecules could be determined.

The modified nucleosides were analyzed by correcting first the original+D value under consideration of the presence of signals with the same m/z introduced by the biosynthetic SILIS (only for 5-methylcytidine and 5-methyluridine). Therefore the integral of the mass signal corresponding to original+D of an independent SILIS measurement was divided by the integrated SILIS mass signal of the same measurement. The resulting correction factor was then multiplied by the SILIS value of the actual sample and subtracted from the sample’s original+D mass signal area.

Using the RFN the signal of each modified nucleoside iso- tomer to SILIS ratio was converted into the amount of injected modification. The amount of new modified nucleoside was normalized by the amount of new tRNA injected and the number of the modification in new tRNA thus revealed. For original and original/post-methylated modified nucleosides the
same procedure was applied but their amounts related to the amount of original tRNA injected.

**Disclosure of potential conflicts of interest**

No potential conflicts of interest were disclosed.

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