Discharging tRNAs: a tug of war between translation and detoxification in *Escherichia coli*

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ABSTRACT

Translation is a central cellular process and is optimized for speed and fidelity. The speed of translation of a single codon depends on the concentration of aminoacyl-tRNAs. Here, we used microarray-based approaches to analyze the charging levels of tRNAs in *Escherichia coli* growing at different growth rates. Strikingly, we observed a non-uniform aminoacylation of tRNAs in complex media. In contrast, in minimal medium, the level of aminoacyl-tRNAs is more uniform and rises to approximately 60%. Particularly, the charging level of tRNA⁹¹Ser, tRNA⁹¹Cys, tRNA⁹¹Thr and tRNA⁹¹His is below 50% in complex medium and their aminoacylation levels mirror the degree that amino acids inhibit growth when individually added to minimal medium. Serine is among the most toxic amino acids for bacteria and tRNAs⁹¹Ser exhibit the lowest charging levels, below 10%, at high growth rate although intracellular serine concentration is plentiful. As a result some serine codons are among the most slowly translated codons. A large fraction of the serine is most likely degraded by L-serine-deaminase, which competes with the seryl-tRNA-synthetase that charges the tRNAs⁹¹Ser. These results indicate that the level of aminoacylation in complex media might be a competition between charging for translation and degradation of amino acids that inhibit growth.

INTRODUCTION

Transfer RNAs (tRNAs) are fundamental components of the translation machinery and link two chemically distinct entities, the nucleotide mRNA sequence and the growing polypeptide (1). Different tRNAs decode all 61 sense codons, yet wobbling at the first position of some anticodons reduces the total number of tRNAs required for decoding. For example, *E. coli* employs 46 tRNAs (2). In prokaryotes and single-cell eukaryotes the abundance of each tRNA correlates with its genomic copy number (3) and with the codon usage of highly expressed genes (4–6). Hence, tRNA isoacceptors (that are tRNA species charged with the same amino acids but with different anticodon sequences) pairing to frequent codons have higher concentrations than tRNAs reading rare codons. This asymmetric tRNA abundance leads to variations in the translation rate of each codon (7) and this non-uniform speed of elongation along mRNA provides an important benefit to facilitate protein folding and biogenesis (8).

The total concentration of individual tRNAs in *E. coli* varies with growth rate, i.e. growth in various media (2,3). Importantly, the abundance of each tRNA species changes non-uniformly with growth rate. For example, the levels of abundant tRNAs increase while minor tRNAs remain relatively unchanged when growth rate increases (2), implying that the adaptability of tRNA pools provides a plasticity to maintain translation balance under different growth conditions. In support of this view, imbalance of the naturally selected tRNA levels alters cell growth and decreases cell fitness (9,10).

tRNAs enter a translation cycle only when charged with their cognate amino acid at its 3′ end. tRNAs are aminoacylated by 20 different aminoacyl-tRNA-synthetases (AARS), each of which is specific for the 20 different amino acids (11). Experimental determination of the charging level of several individual tRNAs in exponentially growing *E. coli* has suggested that they are aminoacylated to 80–90% with their cognate amino acid (12–14). This result has been widely extrapolated to all tRNAs, thus assuming a nearly uniform charging level for all tRNAs (15). However, the aminoacylation level of the tRNA isoacceptors sensitively reacts to changes in environmental conditions (16,17). Thereby various tRNA isoacceptors within one tRNA isoacceptor family (i.e. all tRNA isoacceptors carrying the same amino acid (1)) respond differ-
ently to environmental changes, suggesting regulatory functions of aminoacylation in shaping the growth response of the cell. For example, amino acid limitations trigger preferential charging of minor tRNAs within one isoacceptor family to facilitate specific translation of amino acid biosynthetic genes enriched in codons pairing to those rare tRNAs (16). Furthermore, tandems of rare codons (18) or long consecutive repeats of a codon with even a plentiful cognate tRNA (19) results in transient ribosomal pausing and frameshifting, implying that a paucity of charged tRNA might be a causal factor. In addition, many tRNAs participate in a plethora of processes beyond translation, including N-terminal tagging of proteins for degradation by phenylalanylt-RNA_{Phe} and leucyl–tRNA_{Leu} (1), hence the turnover of aminoacyl–tRNA levels may differ from the turnover of tRNAs involved only in elongation. Together, these observations question the general assumption that tRNAs are uniformly charged. Using the power of microarray technology to determine the charging levels of all tRNA isoacceptor families (17), we set to revisit this arbitrarily established assumption which is based on only a few experimental examples (12–14). We found that the levels of aminoacyl-tRNAs are non-uniform and the low suboptimal charging of some tRNAs is determined by the trade-off between amino acid sequestration into aminoacyl–tRNAs for biosynthesis and degradation for detoxification.

**MATERIALS AND METHODS**

**Media and growth conditions**

*Escherichia coli* MC4100 was grown aerobically at 37°C in LB medium (10 g/l NaCl, 10 g/l tryptone and 5 g/l yeast extract), M9 minimal medium (MM, 12.8 g/l Na_2HPO_4·7H_2O, 3 g/l KH_2PO_4, 1 g/l NH_4Cl, 0.24 g/l MgSO_4, 0.011 g/l CaCl_2, 4 g/l glucose) or defined rich medium (M9 minimal medium supplemented with each amino acid to the following concentrations as reported in (20): Cys, Trp – 0.1 mM; His, Met, Tyr – 0.2 mM; Arg, Asn, Asp, Ile, Lys, Phe, Pro, Thr, Trp – 0.4 mM; Glu, Gln, Val – 0.6 mM; Ala, Gly, Leu – 0.8 mM; Ser 10 mM). Starter cultures for microarray analysis cultures were grown in shake flasks. Growth curves in MM supplemented with one amino acid were performed in 96-well plates sealed with gas permeable foil at 37°C and 2 mm shaking amplitude. OD_{600} data were collected with TECAN Infinite M200 plate reader and zeroed with a blank of the medium. The intracellular concentration of free amino acids was determined as described (21). Calculation of the concentration from mol/dry cell weight to mM was performed as described for calculating various intracellular metabolites using 9 \times 10^{-12}g and 3 \times 10^{-13}g for *E. coli* volume in complex media (LB and MM+AA) and MM, respectively (22). The acetate concentration in the growth medium was measured at OD 0.5 with Acetate Colorimetric Assay kit (Sigma).

**Microarray analysis**

RNA was isolated under mild acidic conditions as described (13) with slight modifications at later steps. After phenol extraction, samples were precipitated with isopropanol for at least 30 min at −80°C and the centrifugation steps were extended to 30 min at 21 000 g at 4°C. tRNAs were oxidized with NaIO_4 as described (3), e.g. 0.1 μg/μl total RNA in 100 mM NaOAc/AcOH (pH 4.5) was treated with 50 mM NaIO_4 at 22°C for 30 min. In parallel, a control non-oxidized sample was treated the same way with NaCl instead of NaIO_4. Oxidized sample was quenched by adding 100 mM of glucose and incubated at 22°C for 10 min. Both samples were spiked in with *in vitro* synthesized transcripts of yeast tRNA_{Phe}, human tRNA_{His} and human tRNA_{Gln}, each of which with a final concentration of 0.08 μM. To remove NaIO_4 from oxidized sample, a G25 spin column was used and both oxidized and control RNA samples were precipitated in ethanol. Thereafter, both RNA samples were decayed into 0.1 M Tris (pH 9) at 37°C for 45 min and neutralized with the same volume of 100 mM NaOAc/AcOH (pH 4.8), 100 mM NaCl. After ethanol precipitation, RNA samples were dissolved into H_2O. For labeling, 0.25 μg/μl RNA was mixed with 4.5 pM tagging fluorescent oligonucleotide (3) in reaction containing 20 U/μl T4 DNA Ligase (New England Biolabs), 1x T4 DNA Ligase buffer (New England Biolabs) and 15% (v/v) DMSO and incubated at 16°C for 16 h. The oxidized sample (i.e. aminoacyl–tRNAs) was labeled with Atto647-labeled oligonucleotide and the control samples representing the total tRNA was labeled with Cy3-labeled oligonucleotide.

After phenol/chloroform extraction, RNA was precipitated with ethanol. The ligation efficiency was estimated on denaturing 10% PAGE by comparing the fluorescent signal of the labeled tRNAs (Fujifilm LAS-4000) to the intensity of the total tRNA visualized with SYBR gold (Invitrogen). Two micrograms of each total labeled tRNA sample was loaded on a microarray containing full-length tDNA probes recognizing 40 *E. coli* tRNA isoacceptors (3) and the three standard spike-ins. On each array, the Atto647-labeled aminoacyl-tRNA and Cy3-labeled total tRNA were loaded and the hybridization was carried out at 60°C for 16 h on Hyb4 microarray hybridization system (Digilab). Each array contained 24 replicates of each tDNA probe. The arrays were normalized to spike-in standards and the signal of Atto647-labeled aminoacyl-tRNA was calculated as a percentage of the cognate Cy3-labeled total tRNA. The quantification was performed with in-house R scripts.

**Northern blot analysis.** Aminoacylation levels of single tRNAs were determined by acidic denaturing polyacrylamide gel as described (13) with some modifications. Untreated and deacetylated RNA samples were resolved on acidic denaturing gel (6.5% polyacrylamide, 8 M urea, 0.1 M NaOAc/AcOH pH 5.0) and electrobotted onto a nylon membrane (GE Healthcare, Amersham Hybond™-N). Hybridization was performed at 60°C for 16 h with a 5'-32P labeled full length tDNA.

**Quantitative RT-PCR.** Total RNA from three biological replicates of exponentially growing cells (OD_{600} = 0.5) was isolated with TRI reagent (Sigma, Aldrich) according to the manufacturer’s instructions with an additional washing step with 70% ethanol. Precipitated RNA was dissolved in RNase-free water, its amount ad-
justed to 1 µg and subjected to DNA depletion with DNase I (Thermo Scientific) followed by cDNA synthesis using RevertAid RT Kit (Thermo Scientific) with random hexamer primers. The mRNA expression of *sdaB* and *sdaC* were quantified using QuantiFast SYBR Green PCR Kit (Qiagen) and normalized against the housekeeping gene gapA, which encodes glyceraldehyde-3-phosphate dehydrogenase (23). Primer sequences were as follows: forward, 5'-ATTTATCTGTGTCGGTACTG-3' and reverse, 5'-AGCGGATCGGGTAGATAGC-3' for *sdaB*; forward, 5'-ACTCCTGTGTAAGTGCTGG-3' and reverse, 5'-CTCTGAGCACGGCTGATAC-3' for *sdaC*; forward, 5'-CGTCCATTTCCAGGTTAG-3' for gapA.

**Analysis of effective codon usage and ribosomal occupancy from ribosome profiling and RNA-Seq**

To determine the codon identities in the ribosomal A site over the whole transcriptome, we used our recently generated ribosome profiling and RNA-Seq data sets from exponentially growing *E. coli* MC4100 grown either in LB or MM (24); the data were deposited in Gene Expression Omnibus (GEO), GSE68762. Ribosome-protected fragments (RPFs) were binned in groups of equal read length and each group was aligned to the stop codons. We used 3′ calibration of the reads as the MNase cleavage was more variable on the 5′ side of the ribosome-protected fragment, which is consistent with prior studies (25). The calibration was performed over the stop codon that allows precise assignment of the ribosomal A site and was used to determine the distance between the middle nucleotide in the A site and 3′ end of the read for each binned read length. We found that 23–26 nt-long reads showed the most evident 3-nt periodicity in frame. The aggregated A site codons were normalized to the library depth and the effective codon usage that represents the codon usage calculated from all expressed mRNAs from the RNA-Seq experiment.

The gene expression examples were calculated by total mapped reads to the corresponding coding sequence and normalized to the gene length and library depth (reads per kilobase per million, rpkM) (26). The threshold of expression level was determined from the biological replicates and it was set for both ribosome profiling and RNA-seq data at >60 reads (24,27).

**RESULTS**

**tRNA isoacceptors are non-uniformly charged in complex LB medium**

To determine the fraction of each aminoacyl-tRNA in *E. coli* (strain MC4100) at various growth phases and in different media, we used tRNA-specific microarrays (17). The microarrays can be run in two modes: (i) to determine differences in tRNAs between two different organisms or cell types (28) or (ii) to quantify the fraction of aminoacyl-tRNA isoacceptor by comparing its amount to that of the total amount of each isoacceptor (17). Here, we used the latter application that takes advantage of the aminoacyl group acting as a protecting moiety of the 3′CCA ends against oxidation. Total tRNA was isolated at acidic pH, which preserves the aminoacyl moiety, and split into two even aliquots. One aliquot was deacylated and represents the total tRNAs. This was labeled with a Cy3-labeled hairpin oligonucleotide pairing to intact CCA-ends. The second aliquot was treated with periodate; only non-aminoacylated tRNAs with free 3′ CCA ends were vulnerable to periodate oxidation while the aminoacyl–tRNAs remained intact (17). Next, the intact aminoacyl-tRNAs were deacylated and Atto647-labeled oligonucleotide was ligated to unoxidized CCA-ends. Both aliquots were hybridized to the same array and the signal of Atto647-labeled aminoacyl-tRNA was calculated as a percentage of the cognate Cy3-labeled total tRNA (Figure 1A). Isoacceptors that differ by at least 8 nt can be unambiguously detected by the microarrays, thus some tRNA probes bind to more than one isoacceptor (17). In total, we were able to differentiate 40 out of 46 tRNAs and all 20 tRNA isoacceptor families of *E. coli* (Supplementary Table S1).

The microarray analysis of exponentially growing *E. coli* MC4100 (OD600 = 0.5) revealed non-uniform charging of tRNAs. Among these, 9 isoacceptors charged between 60–75% (e.g. tRNA isoacceptors carrying Asn, Asp, Gly, Phe, Pro) and 20 nearly completely (>75%) aminoacylated (Figure 1A). Unexpectedly, we detected quite low charging levels (under 50%) for tRNA isoacceptors carrying Cys, His, Ser and Thr (Figure 1A). This finding was also corroborated by Northern blot analysis (Supplementary Figure S1A). Notably, all tRNA isoacceptors within one tRNA–isoacceptor family were charged to similar levels (Figure 1A). Similar suboptimal charging of Cys, Ser and Thr isoacceptor families was also observed in *E. coli* BL21(DE3) grown in LB medium, suggesting that the effect is not strain-specific (Supplementary Figure S1B).

Aminoacyl–tRNAs reach the ribosomes by diffusion and their concentration determines the translation rate of each codon (7,29). Because of the extremely low concentration of seryl–tRNAsSer, we hypothesized that Ser codons will be translated with much slower velocity than any other codon whose tRNAs are completely charged. To test this idea, we used ribosome profiling (30) to determine the average speed of translation of each codon in MC4100 cells. The frequency a codon is detected in the ribosomal A site correlates with the ribosome dwell time for that particular codon and is a measure of average codon translational speed (15,31). The ribosome profiling showed a good correlation between biological replicates (24,27). We binned the RPF by their length and calibrated them separately to the A-site codon using the 3′-assignment method that gives more precise position-specific signal for prokaryotes as discussed recently (25). Frequently used codons will statistically occupy the ribosomal A site more often, thus in parallel we performed RNA-Seq (24,27) and normalized it to the codon usage in the transcriptome (i.e. the codon usage in the genome weighted by the mRNA expression level determined by RNA-Seq). Aggregating the calibrated reads for all sequencing read lengths relative to the start and stop codon revealed a clear 3-nt phasing for the RPFs which is absent in the mRNA reads (Supplementary Figure S2A). Indeed, the occupancy of three out of six Ser codons in the ribosomal A-site was markedly high and in the 90th percentile of all codons. We attributed the increase in ribosome
Figure 1. Some tRNAs have suboptimal aminoacylation level in cells grown in LB medium. (A) Microarray of aminoacyl-tRNAs of *E. coli* MC4100 at OD<sub>600</sub> = 0.5 grown in LB at 37°C. tRNA probes are shown with their anticodon and the corresponding amino acid and classified according to the properties of the amino acid. Data are means ± SEM of three biological replicates. The horizontal dashed line denotes the level at which all tRNAs are fully charged. MetCAU<sub>f</sub> and MetCAU<sub>m</sub> denote initiator and elongator tRNAs, respectively (see also Supplementary Table S1). (B) Average ribosome dwelling occupancy, i.e. the frequency of each codon in the ribosomal A site, as determined by ribosome profiling and plotted in correspondence to their frequency in the translatome (i.e. effective codon usage calculated from all mRNAs expressed in MC4100 cells using RNA-Seq). The codons with low levels of aminoacyl-tRNAs (from panel A) are color-coded: Ser codons in red, Cys codons in orange, His codons in blue and Thr codons in light blue. The dashed line denotes the upper bound (90% confidence interval) of the ribosome dwelling occupancies. (C) Microarray of seryl-tRNAs<sub>Ser</sub> in LB medium at OD<sub>600</sub> = 1.5 (black bars), OD<sub>600</sub> = 2.5 (light gray bars) and OD<sub>600</sub> = 3.5 (dark gray bars). Data are means ± S.D. For the charging pattern of the whole tRNAome see Supplementary Figure S2C.
dwellings at these Ser codons to a decrease in translation rate, i.e. these Ser codons are among the most slowly translated codons in exponentially growing MC4100 in LB medium (Figure 1B), which correlates with the paucity of their cognate aminoacyl-tRNAs (Figure 1A), despite the fact that non-aminoacylated tRNAs\textsuperscript{Ser} are plentiful (2). The ribosomes dwell longer at only three (AGU, UCA and UCG) out of six Ser codons, although the four cognate seryl-tRNAs\textsuperscript{Ser} are low but evenly charged (Figure 1A). In searching for reason for the particular shorthand of only these three codons, we realized that the transcriptome usage of AGU, UCA and UCG is an order of magnitude lower than that of the other three Ser codons (AGC, TCC and TCT); the codon usage at the whole transcriptome mirrors the abundance of their cognate tRNAs. Predictive modeling, reassuringly, identifies a potential bottleneck and effect on fitness only of rare Ser codons with rare tRNAs (10).

One of the Gly codons, the GGA codon, also exhibited higher ribosomal occupancy (Figure 1B) suggesting it is slowly translated. Its cognate tRNA\textsuperscript{Gly UCC} is plentiful (2), but has the lowest charging level among the tRNAs\textsuperscript{Gly} (65%, Figure 1A) that may lead to a delay the ribosomes awaiting the glycyl-tRNA\textsuperscript{Gly UCC}. However, tRNA\textsuperscript{Gly GCC} and tRNA\textsuperscript{Gly CCC} are charged to similar levels as tRNA\textsuperscript{Gly UCC} (Figure 1A) for which we did not detect increased ribosome occupancy (Figure 1B). Effects beyond the concentration of the cognate tRNA may cause this: a transient pausing at Gly codons has also been detected in the ribosome profiling in \textit{E. coli} BW25113 grown in LB and interpreted as a result of slower decoding or tRNA translocation from A- to P-site at Gly codons (32).

Strikingly, from all lowly-charged tRNA families (tRNA\textsuperscript{Gly}, tRNA\textsuperscript{His}, tRNA\textsuperscript{Ser} and tRNA\textsuperscript{Thr}) in both \textit{E. coli} MC4100 and \textit{E. coli} BL21(DE3) the tRNAs\textsuperscript{Ser} isoacceptor family exhibited the lowest aminoacylation level, at 5–10% (Figure 1A and Supplementary Figure S1B). Similarly to our observations, high ribosome occupancy at Ser codons has been reported in ribosome profiling of \textit{E. coli} MG1655 in LB (33,34), however, interpreted as a result of nutrient (C-source) limitation and catabolizing Ser for energy production. Indeed, it is suggested that Ser is the first amino acid to be catabolized following shortage of a carbon source (e.g. glucose) in the medium (35). Thus, we reasoned that if nutrient limitations (e.g. limitations of the uptake of the two organic ingredients in LB, tryptone and yeast extract), are a rate-limiting step and create nutrient shortage, the addition of a favorable C-source (e.g. glucose) would be preferably used and should alleviate charging of tRNAs\textsuperscript{Ser}. However, we found that supplementation of the LB medium with glucose did not offset the low charging of the tRNAs\textsuperscript{Ser} isoacceptors (Supplementary Figure S1C).

Also, supplementing LB with an excess of Ser did not enhance the charging level of tRNAs\textsuperscript{Ser} (Supplementary Figure S1C), arguing against the suggestion that the low charging level of tRNAs\textsuperscript{Ser} is a consequence of Ser or nutrient limitations.

Next, we asked whether the tRNA aminoacylation pattern changes with growth phase as protein synthesis declines with bacterial age (36). We analyzed the fraction of aminoacylated tRNA isoacceptors in late exponential phase (OD\textsubscript{600} = 1.5) and early (OD\textsubscript{600} = 2.5) and late (OD\textsubscript{600} = 3.5) stationary phase. The tRNAs\textsuperscript{Ser} charging levels gradually increased at late growth phase compared to the exponential phase (Figure 1C and Supplementary Figure S2B), implying that a low charging level of tRNAs\textsuperscript{Ser} is not an intrinsic property of the tRNAs\textsuperscript{Ser} family and depends on the physiological activities of the cell. Similarly, the charging level for the most tRNA isoacceptors increased at higher OD (Supplementary Figure S2C). A likely explanation is that at higher ODs the cell ceases growth and consequently protein biosynthesis and the demand for charged tRNAs decreases. It is also possible that extracellular amino acids are completely depleted from the medium during exponential growth (35,37), thus the supply in the stationary phase is entirely maintained by the intracellular amino acid synthetic activities of the cell, a constellation which is reminiscent of growth in minimal media (see below).

tRNA aminoacylation level is more uniform by intrinsic amino acid synthesis

Growth in various media activates different physiological programs in bacteria and alters metabolic enzymes (38) to adjust the flow of carbon that is coupled to energy and intracellular nucleoside concentrations (39). tRNA and tRNA expression are physiologically connected to the metabolic state and growth rate of the cell (2,40). Thus, we next asked whether the tRNA charging levels will change when the growth rate decreases, at which time the total concentration of the tRNA isoacceptors also decreases (2). We analyzed the aminoacylation level of exponentially growing \textit{E. coli} MC4100 (OD\textsubscript{600} = 0.5) in minimal medium (MM) with glucose as a sole carbon source; the growth rate in MM is 0.016 ± 0.0023 min\textsuperscript{-1} and is about two times slower than in LB (the growth rate in LB is 0.026 ± 0.0019 min\textsuperscript{-1}) (Supplementary Figure S3A). In MM all amino acids needed to serve translation activities are internally produced through the own biosynthetic activities of the cell. Notably, in MM the aminoacylation level of all tRNA\textsuperscript{Ser} isoacceptors increased up to 50% (Figure 2A and Supplementary Figure S3B). In general, the aminoacylation level in MM is lower for all tRNA isoacceptors (Figure 2A) but far more uniform than in LB (Figure 1A). Similar aminoacylation levels were observed for another \textit{E. coli} strain, CP78, when cultured in MOPS minimal medium (17).

In MM, serine codons were not among most slowly translated and instead exhibited an average speed of elongation (Figure 2B). Two Gly codons were among the slowest elongated codons in MM although their cognate tRNAs were charged to a similar extent as the majority of the tRNAs in MM (Figure 2A). The detection of enhanced ribosome occupancy at Gly codons in MM suggests that the intimate process of decoding Gly codons, e.g. translocation, is rate-limiting (32).

tRNA aminoacylation levels mirror the degree of amino acid toxicity

What is the reason for the low charging level of some tRNAs in LB medium? LB is an undefined medium in which tryptone and yeast extract are the sources for amino acids
Figure 2. Medium composition dictates the aminoacylation level of tRNA isoacceptors. (A) Microarray of aminoacyl-tRNAs of *E. coli* MC4100 at OD<sub>600</sub> = 0.5 grown in MM with 0.4% glucose at 37°C. For more details see the legend to Figure 1A. (B) Average ribosome dwelling occupancy as determined by ribosome profiling and plotted in correspondence to codon frequency in the translatome (for details see the legend to Figure 1B). The following codons are color-coded: Ser codons in red, Cys codons in orange, His codons in blue and Thr codons in light blue. The dashed line denotes the upper bound (90% confidence interval) of the ribosome dwelling occupancies. (C) Microarray of seryl-tRNAs<sup>Ser</sup> in LB (black), MM+AA (light gray) and MM (dark gray) at OD<sub>600</sub> = 0.5. Data are means ± S.D. of two to three biological replicates. The microarray with all aminoacyl-tRNAs in MM+AA is included in Supplementary Figure S3C.
in de novo translation. Since the composition of these complex substrates often varies and their amino acid composition may lead to a shortage of some amino acids, we next compared the tRNA aminoacylation levels in LB to those in a defined complex medium (MM+AA) containing glucose as a C-source and optimal concentrations of all 20 proteinogenic amino acids (20). The growth rate in MM+AA at 37°C of 0.023 ± 0.0015 min⁻¹ is similar to that in LB (Supplementary Figure S3A). The charging level of the tRNA isoacceptors was slightly higher in MM+AA but more similar to that in LB than in MM (Supplementary Figure S3B). Notably, the charging level of tRNAs\textsubscript{Ser} rose to ~15–20% but remained lower than that in MM (Figure 2C). We measured an internal concentration of free Ser in MM+AA of 4.0 ± 0.3 mM (Supplementary Table S2), which is one of the highest among all amino acids and argues that Ser uptake and free Ser concentration in the cell is not a bottleneck. For comparison, the intracellular free Ser concentration in \textit{E. coli} MC4100 grown in MM was two orders of magnitude lower 0.057 ± 0.005 mM (Supplementary Table S2) and of similar magnitude as measured for \textit{E. coli} NCIM3722 – 0.068 mM (22). Note that we were not able to quantify the amount of the intracellular free amino acids in LB because of the high background from medium remains. The similar growth rate and charging level between LB and MM+AA suggests that the intracellular free amino acids concentration will be within the same order of magnitude.

Despite the lower intracellular Ser concentration in MM as compared to MM+AA, the charging level of tRNAs\textsubscript{Ser} isoacceptors is much higher in MM than in MM+AA (Figure 2C and Supplementary Figure S3C). Enhancing the charging of tRNAs\textsubscript{Ser} by supplementation of Ser in MM was not possible, since Ser readily inhibited growth even at the lowest external concentration (Figure 3). Similarly, Cys also inhibited growth at very low concentrations when present in MM, while Thr and His showed an altered phenotype in the onset of exponential growth, but no notable effect on the growth rate (Figure 3 and Supplementary Table S3). Amino acids whose tRNA isoacceptors were charged >60% inhibited cell growth only at relatively high concentrations, (e.g. Phe, Ala, Leu) or even stimulated growth (e.g. Asn and Gly) (Figure 3). In sum, for the amino acids for which we observed suboptimal charging in LB medium (e.g. Ser, Cys, Thr and His), the degree of growth inhibition mirrored the charging level of different tRNA isoacceptors, with tRNAs\textsubscript{Ser} and tRNAs\textsubscript{Cys} isoacceptor families exhibiting the lowest charging level (Figure 1A) and Ser and Cys being the most toxic to \textit{E. coli} MC4100 (Figure 3).

The inhibitory effect of free Ser in the growth medium has been noticed earlier (41). Ser is readily converted to pyruvate by L-Ser deaminases (42). Three genes in the \textit{E. coli} genome encode L-Ser deaminases: \textit{sdaA} and \textit{tdcG} are expressed in anaerobic and \textit{sdaB} in aerobic conditions (43,44). \textit{sdaB} is within the same operon with \textit{sdaC} encoding a Ser/H⁺ symporter. To explore the basis of the low seryl tRNAs\textsubscript{Ser} despite the high intercellular Ser concentration in complex medium (Supplementary Table S2), we next examined the expression of \textit{sdaB} and \textit{sdaC} in different media using both quantitative RT-PCR and RNA-Seq data. The RNA-Seq data sets are available only for LB and MM and the sequencing reads were normalized to rpKM values which denote total reads detected per open-reading frame normalized to the depth of each library (26) (see also Materials and Methods section). Both \textit{sdaB} and \textit{sdaC} were highly expressed in LB medium but were barely expressed in MM (Figure 4A). Notably, in LB \textit{sdaB} scored among the 50 most highly expressed genes in \textit{E. coli} in the RNA-Seq data set.

Next, we analyzed whether \textit{sdaB} and \textit{sdaC} expression exhibits a concentration-dependent response to the amount of Ser added in the medium. Thus, to MM+AA depleted of Ser we added different concentrations of Ser (0, 5 and 10 mM; the last is the concentration in MM+AA). Both \textit{sdaB} and \textit{sdaC} were expressed in all conditions, even in MM+AA without Ser (Figure 4A). The expression of the Ser/H⁺ symporter (\textit{sdaC}) in MM+AA lacking Ser is not surprising as the expression of Ser transporters is coupled to the expression of transporters for Gly and Ala (45), both amino acids were present in this growth condition. Since \textit{sdaC} and \textit{sdaB} are encoded within one operon and their expression is coupled, the induction of \textit{sdaC} inevitably triggers the expression of \textit{sdaB} encoding the L-Ser deaminase. In all of the conditions with high expression of \textit{sdaB}, the charging of tRNAs\textsubscript{Ser} family remained lower than in MM where the \textit{sdaB} expression is nearly absent (Figure 4B), suggesting a link between low charging level of tRNAs\textsubscript{Ser} and the expression of L-Ser deaminase which most likely degrades some portion of the free intracellular Ser.

Metabolic flux analysis suggests that at high intracellular concentrations Ser is redirected through L-Ser-deaminase toward pyruvate in the glycolysis pathway. However, the majority is not efficiently metabolized for energy but is instead wastefully metabolized and expelled as acetate (37). We reasoned that if excess Ser in complex media would be detoxified through L-Ser-deaminase, it would result in higher acetate production than in MM since a minimal acetate production is a common by-product of aerobic growth on glucose (46). Thus, any amount higher than MM would correlate with the metabolism of other constituents of the medium. The acetate production was higher in both complex media (LB and MM+AA) than in MM (Table 1). Our result in complex media agrees with the acetate concentration measured previously for another \textit{E. coli} strain (37). Furthermore, we observed a clear increase in acetate production when adding Ser to the MM+AA (Table 1). The effect was not concentration-dependent and resembled the expression pattern of \textit{sdaB} and \textit{sdaC} (Figure 4A). Together, our data provide evidence that in media with low charging levels of tRNAs\textsubscript{Ser} (LB and MM+AA), the expression of L-Ser-deaminase is high and acetate is excreted indicating that serine is, at least in part, degraded wastefully. In contrast, L-Ser deaminase is nearly absent in MM so that biosynthetically produced Ser is only used for tRNAs\textsubscript{Ser} charging.

**DISCUSSION**

In this study, we determined the aminoacylation level of tRNAs by leveraging the near single-isoacceptor resolution of tRNA-based microarray technology. Strikingly, we found that in LB and MM+AA tRNA isoacceptors are not uniformly charged with their cognate amino acids, a phenomenon which is anticodon-unspecific and relates to all
Figure 3. Some amino acids inhibit growth of *E. coli*. Growth curves at 37°C in MM containing different concentrations of one single amino acid. Rainbow colors of solid lines from purple to dark green denote amino acid concentration of 0, 0.5, 1, 5, 10, 15, 30 and 50 mM, respectively. For some amino acids, additional concentrations were measured (dashed lines) to cover their concentration in MM+AA: Cys, purple dashed line—0.05 mM and dark blue dashed line—0.1 mM; His, purple dashed line—0.1 mM and dark blue dashed line—0.2 mM; Thr, Asn and Phe, purple dashed line—0.2 mM and dark blue dashed line—0.4 mM. The charging level of tRNAs charged with Ser, Cys, Thr, His, Asn and Phe was low. For comparison, the growth in the presence of Ala, Leu and Gly is included, for which cognate tRNAs were nearly completely charged.

Table 1. Acetate concentration in the growth medium at OD$_{600}$ = 0.5

| Condition                        | Acetate concentration, ng/µl |
|----------------------------------|------------------------------|
| LB                               | 403.9 ± 5.1                  |
| MM+AA, 10 mM Ser                 | 420.1 ± 8.1                  |
| MM+AA, 5 mM Ser                  | 439.7 ± 1.2                  |
| MM+AA without Ser                | 342.9 ± 11.0                 |
| MM                               | 298.8 ± 6.0                  |

tRNAs within one isoacceptor family. In MM, the charging of tRNA isoacceptors is more uniform, yet it also deviates from the assumed optimal charging level and oscillates around 50–60%. Thus, our data revisit a common assumption that *E. coli* tRNA isoacceptors are uniformly charged to 80–90% (15), a result that has been extrapolated from few single tRNAs (12–14). tRNAs$^{\text{Ser}}$ exhibit the lowest charging level in complex MM+AA and LB media followed by tRNAs$^{\text{Cys}}$, tRNA$^{\text{Thr}}$ and tRNA$^{\text{His}}$. Thereby the aminoacylation level mirrors the toxicity of these amino acids when individually added to MM: the more toxic one amino acid is, the lower its aminoacylation level. Serine and cysteine are the most toxic amino acids for *E. coli*, as even the smallest supplementation to MM ceases cell growth completely. Cys is a potent inhibitor of growth, however, bacteria can overcome this effect (47), while the inhibition by Ser is long-lasting (41). The precise mechanism of inhibition by Ser is unclear and could branch over many different pathways as usually observed for perturbation of energy-coupled factors (42,44). Despite the high toxicity of Ser, Ser transporters are very efficient ($K_m = 1–10$ µM (45)) and maintain high Ser uptake when this amino acid is present in the nutrient medium. The steady-state intracellular concentration of Ser is the highest among
all amino acids in complex medium and is two orders of magnitude higher than that in MM (Supplementary Table S2). Thus, the free intracellular Ser should not be a limiting factor in complex (MM+AA und LB) media. However, as suggested by recent metabolic flux analysis in E. coli only 6.5% (∼0.2 mM) of the intracellular concentration of Ser is used directly for tRNA<sub>Ser</sub> aminoacylation, while 51% of the Ser flux is directed toward pyruvate and acetate production and 36% for glycine synthesis (37). Our measurements of the high expression of sdaB encoding L-Ser-deaminase under aerobic growth in LB and MM+AA, but not in MM and increased acetate production in complex media support the notion that intracellular Ser, at least in part, is degraded by L-Ser-deaminase, which competes with the seryl-tRNA-synthetase (Ser-RS) that charges the tRNA<sub>Ser</sub>. Interestingly, <i>k<sub>cat</sub>/K<sub>m</sub></i> values of L-Ser-deaminase of 1.6 x 10<sup>5</sup> (48) and Ser-RS of 1.5 x 10<sup>4</sup> (49) differ by an order of magnitude for their common substrate Ser. Ser-RS has much higher affinity for serine (e.g. <i>K<sub>m</sub></i> of 53 μM (49) versus 2.67 mM (48) of the L-Ser-deaminase), whereas L-Ser-deaminase exhibits much higher turnover (e.g. <i>k<sub>cat</sub></i> of 436 s<sup>-1</sup> (48) versus 0.8 s<sup>-1</sup> of Ser-RS (49)). In LB, the concentration of intracellular Ser (4.00 ± 0.30 mM, Supplementary Table S2) is higher than the <i>K<sub>m</sub></i> values of both L-Ser-deaminase and Ser-RS, hence serine should undergo much higher utilization by the L-Ser-deaminase due to its high turnover number (<i>k<sub>cat</sub></i>) than that of the Ser-RS. In complex medium, seryl–tRNA is also depleted in translation with turnover of 8.1 s<sup>-1</sup> (50). Thus, two parallel pathways contribute to the low charging level of tRNA<sub>Ser</sub>: its utilization in translation and also depletion of Ser by L-Ser-deaminase, while the latter factor plays a dominant role at high Ser concentration (e.g. LB or MM+AA). Our acetate measurements and the metabolic flux analysis (15) corroborate this large role of L-Ser-deaminase in depleting free Ser in complex media.

Notwithstanding the measured low charging level of tRNA<sub>Ser</sub>, they are not correlated with fitness; the growth rate in LB and MM+AA is higher than MM (Supplementary Figure S3A). The six Ser codons are not rare on a genome scale, i.e. 5.85% of all 61 sense codons. However, Ser codons are rarer in the transcriptome, i.e. 3.9% (Figures 1B and 2B), suggesting that fast growth might be sustained even with a low-charged level of tRNA<sub>Ser</sub>. High intracellular Ser concentration at high growth rate may trigger mischarging of tRNA<sub>Thr</sub> and tRNA<sub>Cys</sub> (51,52) and consequently misincorporating Ser into proteins in lieu of Thr and Ala codons. Although it may take place, the lack of effect on bacterial fitness argues that amino acid mischarging and misincorporation is not a prevalent process.

The specificity of recognition between some tRNAs and their cognate AARS is facilitated by specific posttranscriptional modification in tRNA (53). While for some modifications (e.g. mmms<sup>5</sup>s<sup>2</sup>U34) the aminoacylation efficiency is invariant, for others (e.g. s<sup>4</sup>U34) the yield strongly depends on the growth rate (54). For example, only partial s<sup>4</sup>U modification of tRNA<sub>Leu</sub>, tRNA<sub>Gly</sub>, tRNA<sub>Pro</sub> and tRNA<sub>Glu</sub> is found at high growth (54). In turn, partial s<sup>4</sup>U modifications reduce tRNA charging (55). We observed reduced charging levels for tRNA<sub>Gly</sub> and tRNA<sub>Pro</sub>, whereas the charging of tRNA<sub>Leu</sub> and tRNA<sub>Glu</sub> was nearly complete at high growth rates in MM+AA and LB. Variations in the modification pattern, particularly at high growth rate (56), may contribute to the suboptimal charging of some tRNAs, yet alone they cannot explain the very low charging of tRNA<sub>Ser</sub>, tRNA<sub>Thr</sub> and tRNA<sub>Cys</sub>.

Three of the amino acids (Ser, Thr and Cys) with suboptimal charging of the cognate tRNAs are metabolically related. For example, Ser is used to produce Cys and the byproduct of this conversion, homoserine, is the precursor for Thr biosynthesis. Thus, it cannot be ruled out that Ser indirectly influences the tRNA charging and detoxification of Thr and Cys as the metabolism of these amino acids is subordinated to Ser.

In sum, our results establish a new paradigm for biological regulatory mechanisms. To fine-tune the internal concentration of some growth-inhibiting amino acids, bacteria might trade off a high level of tRNA charging for efficient degradation and detoxification.
SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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