Crucial contribution of the multiple copies of the initiator tRNA genes in the fidelity of tRNA<sup>fMet</sup> selection on the ribosomal P-site in *Escherichia coli*

Suman Kapoor<sup>1</sup>, Gautam Das<sup>1</sup> and Umesh Varshney<sup>1,2,*</sup>

<sup>1</sup>Department of Microbiology and Cell Biology, Indian Institute of Science, Bangalore 560012 and <sup>2</sup>Jawaharlal Nehru Centre for Advanced Scientific Research, Bangalore 560064, Karnataka, India

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**ABSTRACT**

The accuracy of the initiator tRNA (tRNA<sup>fMet</sup>) selection in the ribosomal P-site is central to the fidelity of protein synthesis. A highly conserved occurrence of three consecutive G–C base pairs in the anticodon stem of tRNA<sup>fMet</sup> contributes to its preferential selection in the P-site. In a genetic screen, using a plasmid borne copy of an inactive tRNA<sup>fMet</sup> mutant wherein the three G–C base pairs were changed, we isolated *Escherichia coli* strains that allow efficient initiation with the tRNA<sup>fMet</sup> mutant. Here, extensive characterization of two such strains revealed novel mutations in the *metZWV* promoter severely compromising tRNA<sup>fMet</sup> levels. Low cellular abundance of the chromosomally encoded tRNA<sup>fMet</sup> allows efficient initiation with the tRNA<sup>fMet</sup> mutant and an elongator tRNA Gln, revealing that a high abundance of the cellular tRNA<sup>fMet</sup> is crucial for the fidelity of initiator tRNA selection on the ribosomal P-site in *E. coli*. We discuss possible implications of the changes in the cellular tRNA<sup>fMet</sup> abundance in proteome remodeling.

**INTRODUCTION**

*Escherichia coli* possesses four copies of the initiator tRNA genes that encode tRNA<sup>fMet</sup>. Three of these (*metZ, metW* and *metV*) are transcribed from the *metZWV* operon located at 63.5 min and the fourth one (*metY*) is transcribed as an upstream gene of the *nuxA-infB* operon at 71.5 min in the genome (1). The three genes in the *metZWV* operon encode three identical tRNAs, tRNA<sup>fMet</sup><sub>1</sub> whereas the *metY* encodes tRNA<sup>fMet</sup><sub>2</sub>. In *E. coli* K, the tRNA<sup>fMet</sup><sub>1</sub> and tRNA<sup>fMet</sup><sub>2</sub> differ from each other at a single nucleotide at position 46; whereas the tRNA<sup>fMet</sup><sub>1</sub> carries a methylated guanine (m<sup>7</sup>G), the tRNA<sup>fMet</sup><sub>2</sub> carries an adenine (A) at this position (2,3). However, in *E. coli* B, all genes encode identical tRNAs corresponding to tRNA<sup>fMet</sup><sub>1</sub> (4).

The initiator tRNAs are special in that they bind directly to the ribosomal P-site. All other tRNAs (the elongator tRNAs) first bind to the ribosomal A-site and are then, subsequent to peptide bond formation and the translocation step, located into the P-site. Selective binding of the initiator tRNAs to the P-site, crucial for the fidelity of initiation, is an outcome of the functional interaction between the unique structural features of the initiator tRNA, the elements within the ribosome, and the action of initiation factors (IFs). Among the features in tRNA<sup>fMet</sup>, a mismatch at the top of the acceptor stem (C1xA72, in *E. coli*) prevents its binding to the elongation factor Tu (EF-Tu); and together with the 2:71 and 3:70 base pairs, it constitutes a major determinant for formylation of the amino acid attached to tRNA<sup>fMet</sup> (5).

Another major structural feature of the initiator tRNAs, highly conserved in all the three domains of life, is the presence of three consecutive G–C base pairs in their anticodon stems. This feature is important for the preferential binding of the initiator tRNAs in the ribosomal P-site (6). Biochemical and genetic studies suggested that IF3 facilitates tRNA<sup>fMet</sup> binding to the 30S ribosomes via the 3GC base pairs (7). Subsequent studies showed that in the ribosome, the binding site of IF3 is located at a distance from the acceptor stem of the tRNA<sup>fMet</sup> (10). More recently, it has been observed that the presence of IF3 increases the rate of dissociation of all tRNAs from the 30S ribosomes. Because the presence of IF2 (along with IF1) strongly favors binding of fMet-tRNA<sup>fMet</sup> to the 30S subunit and docking of the 50S subunit, such a role of IF3, in essence allows for
higher accuracy of fMet-tRNA<sup>Met</sup> selection on the ribosome (12). Also, two of the universally conserved nucleotides G1338 and A1339 in the head region of the 16S rRNA are in juxtaposition to the minor groove of the 3GC base pairs (13,14). Mutational analysis of these residues (15) has supported their role in stabilizing tRNA<sup>Met</sup> binding via the ‘A-minor’ interactions with the 3GC base pairs (16). In addition, C-terminals of S9 and S13 are within the contacting distance with the anticodon stem loop of the tRNA<sup>Met</sup> (14).

To further our understanding of the biological significance of the 3GC base pairs in initiator tRNA selection on the ribosomal P-site, we developed a genetic screen (17) and isolated <i>E. coli</i> strains that allow initiation with tRNAs lacking the 3GC base pairs. Characterization of two such strains in this study shows that they possess novel mutations in the <i>metZWV</i> promoter severely compromising tRNA<sup>Met</sup> expression. Low cellular abundance of the tRNA<sup>Met</sup> allows efficient initiation not only with the mutant tRNA<sup>Met</sup> but also with tRNA<sup>Gin</sup> revealing that the high abundance of the cellular tRNA<sup>Met</sup> is crucial for the fidelity of translation in <i>E. coli</i>.

**MATERIALS AND METHODS**

**Strains, plasmids and media**

Strains and plasmids have been listed in Supplemetntary Tables S1 and S2. Bacteria were grown in Luria-Bertani (LB) liquid or LB-agar plates (18) containing 1.5% bacto-agar (Difco). Unless indicated otherwise, media were supplemented with ampicillin (Amp, 100 μg ml<sup>-1</sup>), chloramphenicol (Cm, 30 μg ml<sup>-1</sup>), kanamycin (Kan, 25 μg ml<sup>-1</sup>) or tetracycline (Tet, 7.5 μg ml<sup>-1</sup>) as required.

**Isolation and characterization of D4 and D27, and their genetic mapping**

The isolation and preliminary characterization of the suppressors of the 3GC mutations in tRNA<sup>Met</sup> have been described before (17). Methodologies used to characterize two of the suppressors, D4 and D27 and the genetic fine mappings of the mutations in these were as described (17).

**Cloning of metZWV gene promoter regions**

The promoter regions of <i>metZWV</i> genes were PCR amplified from genomic DNAs of <i>E. coli</i> KL16 (the parent strain), D4 and D27 using <i>Pfu</i> DNA polymerase in 50-μl reactions containing 20 pmols each of the forward (5’ cgctgagctgaggcaagcag 3’) and reverse (5’ cctacagccggtttatcgcagc 3’) primers. The amplicons (~363 bp) were digested with KpnI and BamHI, eluted from agarose gels and cloned into similarly digested promoterless pTKCAT (18).

**Cloning of supE**

The <i>glnVX</i> gene (<i>supE</i>) was PCR amplified from <i>E. coli</i> TG1 using a supEfp (5’ gctgagctgccggtttatcgcag 3’) and a supErp (5’ cttacagcagcctggtttatcgcag 3’) primers as 0.4-kb amplicon, digested with NcoI and EcoR1 and cloned into pACDH between the same sites to generate pACDHsupE (18).

**Isolation of tRNAs and northern blot analysis**

Total tRNA preparations from various strains were fractionated on native 15% polyacrylamide gels (6), electroblotted onto nitran membrane and analyzed by northern blotting using a <sup>32</sup>P 5’-end labeled DNA oligomer (5’ cttacagctgaggcaagcag 3’) essentially as described (19).

**Generation of <i>AmetY</i> :: <i>kan</i> strains**

The Kan<sup>R</sup> cassette from the plasmid pKD4 was amplified with <i>metYko_fp</i> (5’ ttacagctgaggcaagcagcctggtttatcgcag 3’) and <i>metYko_rp</i> (5’ ttacgctcaagcagcctggtttatcgcagcctggtttatcgcag 3’) primers, electroporated into <i>E. coli</i> DY330 and selected on Kan plates (20). Transformants were screened for replacement of <i>metY</i> with the Kan<sup>R</sup> cassette by PCR with the flanking primers, metYfp (5’ cttacagctgaggcaagcagcctggtttatcgcag 3’) and metYrp (5’ gctgagctgaggcaagcag 3’). The <i>AmetY</i> :: <i>kan</i> locus was then mobilized into <i>E. coli</i> KL16, D4 and D27 strains by P1-mediated transductions to generate their <i>AmetY</i> :: <i>kan</i> counterparts (21).

**Generation of metZWV :: <i>kan</i> strains**

The Kan<sup>R</sup> cassette from pKD4 was amplified with the primers, <i>metZWVko_fp</i> (5’ ttgtagggctggtttatcgcagcctggtttatcgcag 3’) and <i>metZWVko_rp</i> (5’ ttctactaagcagcctggtttatcgcagcctggtttatcgcag 3’) primers, electroporated into <i>E. coli</i> DY330 (20). Transformants were screened for replacement of <i>metZWV</i> with the Kan<sup>R</sup> cassette by PCR with the flanking primers, metZWVfp and metZWP (5’ ttgtagggctggtttatcgcagcctggtttatcgcag 3’ and metZWVrp 5’ gctgagctgaggcaagcag 3’) following which the <i>AmetZWV</i> :: <i>kan</i> locus was mobilized into <i>E. coli</i> KL16, D4 and D27 by P1-mediated transductions to generate their <i>AmetZWV</i> :: <i>kan</i> counterparts (21).

**Preparation of cell-free extracts and chloramphenicol acetyltransferase assays**

<i>Escherichia coli</i> cells were grown in 3-ml LB medium containing Amp to mid-log phase and processed as described (17). Total of the pixel values in the spots corresponding to 1-acetyl-, plus 3-acetyl-, chloramphenicol (Ac-Cm, P) and the left over substrate, chloramphenicol (Cm, S) were quantified using a BioImageAnalyzer (FLA5000, Fuji). The chloramphenicol acetyltransferase (CAT) assays were done for 15min using appropriate amounts of total cell extracts as defined in the figure legends. The CAT activities were calculated as picomoles of Cm converted to Ac-Cm per microgram total protein ([by multiplying the P/(S+P) ratio with the total picomoles of Cm taken in the reaction] divided by the total protein in the cell extract). Assays were carried out from three independent colonies and the averages (picomoles of Cm converted to Ac-Cm per microgram total protein per
15 min, along with the associated standard deviations) were used to generate histograms.

RESULTS

The assay for initiation in *E. coli* and characterization of D4 and D27

The plasmid, pCAT<sub>am1</sub><sub>metY<sub>_CUA</sub></sub> (Figure 1A) harbors a CAT<sub>am1</sub> reporter gene wherein the AUG initiation codon has been changed to UAG codon along with mutations in the second and the fifth codons to eliminate secondary sites of weak initiation (22). The presence of an initiator tRNA gene harboring a complementary mutation in the anticodon, CAU to CUA (Figure 1A, panel i) fosters efficient initiation from the UAG initiation codon of the reporter and confers resistance to chloramphenicol (Cm<sup>R</sup>) to *E. coli*. As the met<sub>Y</sub><sub>_CUA</sub> encoded tRNA is aminoacylated with Gln, initiation occurs with formyl-Gln (22). Expectedly, when the three consecutive G–C base pairs in the anticodon stem of the initiator tRNA were mutated to those found in the elongator tRNA<sup>Met</sup> (G29G30G31/C39C40C41 to U29C30A31/ψ39G40A41; hereafter referred to as ‘3GC’ mutations or the 3GC tRNA), in the plasmid pCAT<sub>am1</sub><sub>metY<sub>_CUA</sub>/3GC</sub> (Figure 1A, panel ii) initiation did not occur to any detectable level and *E. coli* harboring the pCAT<sub>am1</sub><sub>metY<sub>_CUA</sub>/3GC</sub> plasmid were sensitive to chloramphenicol (Cm<sup>S</sup>) even though the encoded tRNA is fully aminoacylated and formylated [(17,23); see also Figure 1B, panel ii, sectors 1 and 2].

In our earlier study, we described isolation of *E. coli* KL16 (K strain) derivatives harboring chromosomal mutations that alleviate the initiation defect of the 3GC tRNA. Characterization of two such suppressor strains, called D4 and D27, is the focus of this study. As a part of the strain validation, introduction of

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**Figure 1.** (A) Assay for initiation in *E. coli*. (i) The plasmid, pCAT<sub>am1</sub><sub>metY<sub>_CUA</sub></sub> possesses pCAT<sub>am1</sub> and met<sub>Y</sub><sub>_CUA</sub> genes. The pCAT<sub>am1</sub> gene produces CAT<sub>am1</sub> reporter mRNA wherein initiation occurs with the UAG codon. The met<sub>Y</sub><sub>_CUA</sub> gene makes tRNA<sub>fMet</sub> wherein the CAU anticodon has been mutated to CUA. The encoded tRNA is shown as tRNA<sub>fMet</sub> (CUA) and the relevant sequence detail of the anticodon stem loop (boxed with the dotted lines) are as shown. Schematic representation shows that in the presence of the wild-type sequence (3GC base pair) in the anticodon stem of the tRNA, initiation from the reporter mRNA confers chloramphenicol resistance (Cm<sup>R</sup>) to the host. (ii) The plasmid, pCAT<sub>am1</sub><sub>metY<sub>_CUA/3GC</sub></sub> is a derivative of pCAT<sub>am1</sub><sub>metY<sub>_CUA</sub></sub> wherein met<sub>Y</sub><sub>_CUA</sub> gene has been replaced with met<sub>Y</sub><sub>_CUA/3GC</sub> gene. The encoded tRNA is shown as tRNA<sub>Met</sub> (CUA/3GC) and the relevant sequence detail of the anticodon stem loop (boxed with the dotted lines) are as shown. The mutated sequence in the anticodon stem (3GC mutations) of the tRNA makes it deficient in initiation and renders the host sensitive to chloramphenicol (Cm<sup>S</sup>). (B) Growth of *E. coli* KL16 parent (KL16) and the suppressor strains (D4, D27) harboring pCAT<sub>am1</sub><sub>metY<sub>_CUA/3GC</sub></sub> (Amp<sup>R</sup>) on LB-agar plates containing, (i) Amp and, (ii) Amp plus Cm. Details of the strains (and the resident plasmids), streaked in various sectors are shown on the right. Plates were streaked with overnight cultures and incubated at 37°C for ~15 h.
pCAT_{am1}metY_{CUA/3GC} plasmid into the plasmid-free versions of D4 and D27 conferred Cm\(^R\) (Figure 1B, panel ii, sectors 3 and 4 and 5 and 6). Under the same conditions, introduction of this plasmid into the parent strain (KL16) did not confer Cm\(^R\) (sectors 1 and 2). As a control, all transformants grew on Amp plate (Figure 1B, panel i). In yet another control, we showed that initiation from CAT_{am1} in D4 and D27 was dependent on the presence of the 3GC tRNA (Supplementary Figure S1).

**Mapping of the suppressor mutations in D4 and D27**

The mutation(s) responsible for the phenotypes of D4 and D27 were mapped to a site between 63.4 and 63.6 min on the *E. coli* genome (Supplementary Figures S2 and S3). Sequence analysis of a high-probability candidate within this region, the *metZWV* operon encoding for the majority of tRNA\(^{fMet}\) in *E. coli*, revealed that D4 and D27 possessed novel mutations within the promoter region of *metZWV* (Figure 2A). We then introduced targeted mutations in the *E. coli* genome to ensure that these single mutations in the promoter regions of *metZWV* of D4 and D27 were responsible for their phenotypes (Supplementary Figure S4).

**The mutations in *metZWV* locus downregulate expression of tRNA\(^{fMet}\)**

Based on the transcriptional start site mapping of the *metZWV* transcripts (24), the mutations in D4 and D27 could impact the promoter quality. Hence, to analyze the influence of these mutations, we carried out northern blot analysis of the total tRNA preparations using a native polyacrylamide gel, which allows separation of tRNA\(^{1fMet}\) and tRNA\(^{2fMet}\) (Figure 2B). As shown in Figure 2B (panel ii), the levels of tRNA\(^{1fMet}\) in D4 and D27 were significantly decreased [compare the relative intensities of the bands corresponding to tRNA\(^{1fMet}\) and tRNA\(^{2fMet}\) in KL16 (lane 1) with those in D4 and D27 (lanes 2 and 3)]. It may be noted that the decrease in the tRNA\(^{1fMet}\) is quite severe and significantly more amounts of total tRNA were needed to detect its presence on the northern blot (Figure 2B, panel i). In fact, in the ethidium bromide stained gel, while the tRNA\(^{1fMet}\) is detectable in Lanes: 1, KL16; 2, D4; and 3, D27. (C) Promoter regions of the *metZWV* locus from KL16, D4 and D27 were cloned upstream of a promoterless CAT reporter in pTKCAT and introduced into *E. coli* KL16. Three independent colonies from each were grown to mid-log phase, and the cell-free extracts (±90 ng total protein) used to assay for CAT activities. Average activities (±SD) are shown as picomoles Ac-Cm (1-acetyl-, and 3-acetyl-, chloramphenicol) formed per microgram total cell protein in 15 min.

**Figure 2.** Mapping of the mutations in D4 and D27 and their initial characterization. (A) Comparative analysis of the promoter regions of D4 and D27 *metZWV* locus with that of the parent (wild-type, WT) reveals that there is a G to A mutation in D4 just upstream to the Pribnow box (−10) and a T to C mutation in D27 within the Pribnow box (−10). The mutations have been indicated by arrows emanating from the sequence of the WT promoter and asterisks above and below the D4 and D27 sequences. The −35 region, the spacing between the −10 and the −35 sequences and the transcription start site (+1) are also indicated. (B) Northern blot analysis of tRNA\(^{1fMet}\) and tRNA\(^{2fMet}\) in the parent (KL16), and the D4 and D27 strains. Total tRNA preparations (−4, 6 and 10 µg, respectively, for KL16, D4 and D27) were separated on the native polyacrylamide gel (15%) stained with ethidium bromide (panel i), transferred onto nytran membrane and analyzed by northern blotting (panel ii). Positions of tRNA\(^{1fMet}\) and tRNA\(^{2fMet}\), and the ratios of the tRNA\(^{1fMet}\) to tRNA\(^{2fMet}\) are indicated. Lanes: 1, KL16; 2, D4; and 3, D27. (C) Promoter regions of the *metZWV* locus from KL16, D4 and D27 were cloned upstream of a promoterless CAT reporter in pTKCAT and introduced into *E. coli* KL16. Three independent colonies from each were grown to mid-log phase, and the cell-free extracts (±90 ng total protein) used to assay for CAT activities. Average activities (±SD) are shown as picomoles Ac-Cm (1-acetyl-, and 3-acetyl-, chloramphenicol) formed per microgram total cell protein in 15 min.
the parent (KL16), the same is not seen in D4 and D27 even though the total tRNA loaded for these samples was in 1.5- and 2.5-fold excess, respectively, of the amounts used for the KL16 sample (Figure 2B, panel i). To ensure that the decreased levels of tRNA\textsubscript{1fMet} were a direct consequence of the promoter down mutation, we cloned equivalent regions of the promoters from the KL16 (wild-type), D4 and D27 strains upstream of a promoterless CAT reporter in pTKCAT vector, and assayed for the CAT activities in \textit{E. coli}. As shown in Figure 2C, the mutations in D4 and D27 indeed led to drastic decreases in the promoter strength.

**D4 and D27 confer a minor growth defect and cold sensitivity to \textit{E. coli}**

To understand the phenotypes of D4 and D27, and the mechanism of initiation by the 3GC tRNA in these strains, we generated strains of \textit{E. coli} KL16 wherein either the \textit{metZWV} or \textit{metY} were knocked out by a Kan\textsuperscript{R} marker. As shown in Figure 3A, while the analysis of the KL16 total tRNA sample on the native polyacrylamide gel (panel i) revealed the presence of both the tRNA\textsubscript{1fMet} and tRNA\textsubscript{2fMet} (lane 1), the tRNA\textsubscript{1fMet} and the tRNA\textsubscript{2fMet} bands, as expected, were missing in the \textit{AmetZWV::kan} and \textit{metY::kan} strains, respectively. The observation was further confirmed by the northern blot analysis (panel ii). It may also be noted that the deletion of \textit{metZWV} did not lead to an increase in the levels of tRNA\textsubscript{2fMet}. Likewise, deletion of \textit{metY} did not impact the levels of tRNA\textsubscript{1fMet}. These observations suggest that the levels of the initiator tRNA in \textit{E. coli} correspond to their gene copy numbers.

To analyze the phenotypes of D4 and D27, we compared their growth kinetics with the \textit{metZWV} and \textit{metY} knockout strains. During their exponential phases, saturated cultures were diluted 1000-fold in LB broth, and growth monitored at 1-h intervals in a Bioscreen C growth reader. All strains were taken in replicates of three, and the average values were plotted. (C) Growth of \textit{E. coli} KL16, its \textit{AmetZWV::kan} and \textit{metY::kan} derivatives and the suppressor strains (D4, D27) harboring pCAT\textsubscript{am\textit{metY\textit{CUA}/3GC}} or pCAT\textsubscript{am\textit{metY}} (Amp\textsuperscript{R}), as indicated, on LB-agar plates at 37\textdegree C (i) and 22\textdegree C (ii). Plates were streaked with overnight cultures and incubated at the indicated temperatures for \textasciitilde 15 h.
the growth of the D4 and D27 strains was slightly compromised when compared with the parent strain (KL16) harboring all four copies of the initiator tRNA genes but slightly better than its AmetZWV::kan derivative (Figure 3B). Further, it was earlier reported that deletion of tRNA1Met locus resulted in cold sensitivity to E. coli (25). Consistent with these observations, deletion of metZWV locus (AmetZWV::kan) which encodes for the majority of the cellular initiator tRNA in E. coli, but not of metY (metY::kan) which contributes to minor amounts of the same, in the background of KL16 (harboring pCATam1metYCUA3GC) resulted in a cold sensitive phenotype at 22°C (Figure 3C, panel ii, compare sector 4 with sectors 3 and 5). Interestingly, E. coli strains with both the D4 and D27 mutations also displayed cold sensitivity (sectors 1 and 7). However, the cold sensitivity could be rescued by the presence of a plasmid borne copy of the metY gene, which encodes for the wild-type tRNA2Met (sectors 2 and 6). As a control, all strains grew well at 37°C (Figure 3C, panel i).

Taken together with the results shown in Figure 2, these observations suggested that a major consequence of the D4 and D27 mutations is to lower the abundance of the tRNA1Met in the cell.

**Mechanism of initiation with the 3GC tRNA in D4 and D27**

Based on the observations in Figures 2 and 3, we hypothesized that the 3GC tRNA has to compete with the chromosomally encoded tRNA1Met for its binding to the ribosomal P-site, and that the lower abundance of the chromosomally encoded tRNA1Met in D4 and D27 results in an increased probability of the P-site occupancy by the 3GC tRNA leading to an increased opportunity of initiation by it. Consistent with this hypothesis, when a compatible plasmid, pACDHNmetYCUA3GC was introduced into the KL16 strain already harboring pCATam1metYCUA3GC (for an elevated level of the 3GC tRNA), it conferred Cm resistance to the transformants such that they grew on a plate with 30μg/ml Cm (Figure 4A, panel ii, sectors 3–6). As a control, introduction of the vector alone (pACDHN) did not confer such a resistance (sectors 1 and 2). Further, knockout of the metY gene (AmetY::kan), a gene which contributes to minor amounts to the cellular pools of tRNA1Met, in KL16 harboring pCATam1metYCUA3GC resulted in gain of initiation by the 3GC tRNA such that it grows at 30–50μg/ml Cm (Figure 4B, sector 3 in all panels). And, a knockout of the metZWV locus (AmetZWV::kan) conferred efficient initiation with the 3GC tRNA to allow good growth even at 100μg/ml Cm (Figure 4B, sector 2 in all panels). As a control, in the absence of these knockouts, the KL16 strain did not foster a detectable initiation with the 3GC tRNA (Figure 4B, sector 1 in all panels).

The knockouts of metY and metZWV were also introduced into D4 and D27 (Figure 4B, sectors 5 and 8, 9). However, for a better assessment of the impact of these knockouts, we carried out CAT assays using cell-free extracts prepared from the strains (Figure 4C). The results from these assays reveal a clear correlation between increase in initiation by the 3GC tRNA and a decrease in the abundance of the chromosomally encoded tRNA1Met. For example, in the KL16 background, knockout of the metZWV gene locus (which is responsible for a major decrease in the tRNA1Met levels) leads to a greater increase in initiation by the 3GC tRNA than does the knockout of the metY gene (which is responsible for a minor decrease in the tRNA1Met levels) (Figure 4B, sectors 1–3; and Figure 4C, compare bars 1–3). On the other hand, in the D4 and D27 strains wherein the metZWV is already substantially downregulated, the metY gene knockout causes a greater increase in initiation (by the 3GC tRNA) than does the metZWV knockout (Figure 4C, compare bars 4–6 and 7–9). In fact, the knockout of metY in D27 (which sustains growth on very low levels of tRNA1Met encoded by the downregulated metZWV) has the highest level of initiation by the 3GC tRNA (bar 9). However, we should add that this strain grows slowly in the liquid medium and is unstable to continued subculturing.

Thus, data in Figure 4 support the concept that the 3GC tRNA has to compete with the wild-type tRNA1Met for the occupancy of the P-site; and the lower the cellular abundance of the wild-type tRNA1Met, the better the initiation with the 3GC mutant. Intuitively, such an inference also implies that high level of tRNA1Met in E. coli (encoded together by the four copies of the initiator tRNA genes) must contribute to the maintenance of the fidelity of initiation by competing out the elongator tRNAs from the ribosomal P-site.

**Initiation with elongator tRNA (supE) and maintenance of the fidelity of initiation by the cellular levels of initiator tRNA**

To check if the elongators could initiate in E. coli, we made use of the TG1 strain which harbors the supE tRNA (tRNA1Gln with CUA anticodon). To assay for the participation of supE at the step of elongation, we used pCATam27 which harbors a CAT gene possessing a UAG codon at position 27 in the reading frame (26). In addition, for these assays, we made use of a two plasmid system. The second pAC1 origin of replication-based plasmid, pACDH (TetR) is compatible for its existence with the colE1-based plasmid in E. coli, and was used to overproduce the supE tRNA. As expected, when the pCATam1 plasmid was introduced into the strain, no detectable levels of CAT protein were produced and the strain remained CmS (Figure 5A, panel i, sector 3; and Figure 5B, bar 1). Overproduction of supE from pACDHsupE, also did not allow production of CAT to any detectable level (sector 1 and bar 3). However, as a control, when pCATam27 was introduced in TG1, it conferred CmR to the host (sector 2 and bar 2). These observations suggest that in a strain possessing full complement of the four initiator tRNA genes, the fidelity of initiation is maintained and the elongator tRNAs do not participate at the step of initiation to any detectable level.
Figure 4. (A) Elevated expression of the 3GC tRNA allows initiation. Plasmids, pACDHN (pAC1 origin of replication, TetR) or its derivative pACDHNmetyCUA/3GC were introduced into E. coli KL16 harboring pCATam1metyCUA/3GC (a colE1 ori of replication, AmpR), and the transformants grown in LB containing Amp plus Tet were streaked onto Amp plus Tet, and Amp plus Tet plus Cm (as indicated) and incubated at 37°C for ~15 h. (B) Growth of the E. coli KL16, D4 and D27 and their ΔmetZWV::kan and ΔmetY::kan derivatives harboring pCATam1metyCUA/3GC (as indicated) on LB-agar plates containing Amp and, Amp plus Cm (30–100 μg ml⁻¹). Plates were streaked with overnight cultures and incubated at 37°C for ~15 h. (C) CAT assays to determine initiation in E. coli KL16, D4 and D27 and their ΔmetZWV::kan and ΔmetY::kan derivatives harboring pCATam1metyCUA/3GC (as indicated). Three independent colonies of each were grown to mid-log phase, and the cell-free extracts (500-ng total protein) used to assay for CAT activities. Average activities (±SD) are shown as picomoles Ac-Cm (1-acetyl-, and 3-acetyl-, chloramphenicol) formed per microgram total cell protein in 15 min.

We next performed these assays in the background of decreased copy number of the initiator tRNA genes (ΔmetZWV::kan). We observed that the overexpression of supE, did confer CmR to the host containing pCATam1 (Figure 5A, panel ii, sector 8; and Figure 5B, bar 6). However, in the presence of chromosomally encoded supE alone, under the assay conditions used, we did not observe CmR phenotype (sector 6) or an increase in the CAT activity above the background (compare bars 1 and 4). Although when the plates were incubated for longer periods (24–30 h), isolated colonies appeared in the transformants of pCATam1 in TG1ΔmetZWV::kan but not in TG1 (data not shown). Importantly, when we overexpressed tRNA^{Met} from a plasmid borne copy, it purged the initiation activity of supE (Figure 5A, panel ii, sector 5), suggesting that the initiation by supE is a specific phenomenon and that the elongator tRNAs do compete with initiator tRNAs in vivo. Furthermore, this experiment showed that higher levels of tRNA^{Met} in the cell are essential to maintain fidelity of initiation in that they prevent initiation by the non-initiator tRNAs. Similar observations were also made in the strains D4 and D27 (Supplementary Figure S5).

DISCUSSION

Barring a few species of mycoplasma (17,27), the presence of the three consecutive G–C base pairs in the anticodon stems of the initiator tRNAs is a highly conserved feature throughout the three domains of life, and any substitutions in the 3GC base pairs lead to a deficient utilization of the mutant tRNA^{Met} in initiation (6). To address the long-standing question of how the 3GC base pairs contribute to initiator tRNA selection on the ribosomal P-site and to gain insight into the mechanisms that maintain the fidelity of translation initiation in E. coli, we used a genetic screen to isolate suppressor strains of
E. coli which due to mutations in their genomes, allow initiation by the 3GC mutant tRNA (17). In the present study, we have characterized two such strains, D4 and D27, and found that they both harbor single nucleotide mutations in the \( \text{C}_{0} \text{C}_{10} \) promoter regions of the \( \text{metZWV} \) operon (a G to A mutation in D4; and a T to C mutation in D27) and possess reduced levels of tRNA\(^{\text{Met}}\) which allows initiation not only by the 3GC mutant but also the elongator tRNA. It is plausible that these mutations impact various aspects of promoter recognition and decrease transcription initiation by the RNA polymerase (28). Simulating conditions of reduced RNA\(^{\text{Met}}\) levels using knockouts of the initiator genes (Figures 4 and 5) has revealed a previously unknown connection between tRNA\(^{\text{Met}}\) abundance and maintenance of the fidelity of initiation by tRNA\(^{\text{Met}}\) by rejecting non-initiator tRNA binding in the ribosomal P-site.

How does the cellular abundance of tRNA\(^{\text{Met}}\) regulate translation of CAT\(_{\text{am1}}\) mRNA with the 3GC or the \( \text{supE} \) tRNAs lacking the 3GC base pairs to the P-site is disallowed. Hence, even though the CAT\(_{\text{am1}}\) mRNA may be engaged on the 30S subunit with the tRNA\(^{\text{Met}}\) containing CAU anticodon (Figure 6A, Step 1), because the mRNA lacks AUG- or AUG-related codons in the vicinity of the Shine–Dalgarno (S. D.) sequence, the codon–anticodon pairing and the conformational changes required to give rise to a 30S pre-initiation complex (29), would not take place. The initial complex would decompose (Figure 6A, Step 1), and prevent translation initiation to proceed through to Steps 2 and 3. Consequently, the CAT protein would not be synthesized and the cells remain Cm\(^{30}\). However, when the abundance of the tRNA\(^{\text{Met}}\) is decreased (as is the case with D4, D27 and the KL16 strains deleted for \( \text{metZWV} \) or \( \text{metY} \)), it now allows for a finite probability of the P-site occupancy with the 3GC mutant or the \( \text{supE} \) tRNAs (possibly because of the delays in the occupancy of the P-site by the tRNA\(^{\text{Met}}\) due to decreased mass effect) (Figure 6B, Step 1). Even though the 3GC or \( \text{supE} \) tRNAs so bound could fall off of the 30S ribosome due to their poor affinity (Figure 6B, Step 1), the increased potential for binding of the tRNAs containing CUA anticodon (complementary to the UAG initiation codon) would lead to an increased potential for formation of productive 30S pre-initiation
complexes with the CATam1 mRNA which can then proceed through steps 2 and 3. The CAT protein so synthesized would confer CmR to cells (Figure 6B). Since initiation is one of the most regulated and rate-limiting step in protein synthesis (30), a high cellular level of tRNA^{Met} (encoded by four genes) has thus far been solely attributed to a necessity for efficient initiation. Based on the observations that a high cellular abundance of initiator tRNA makes significant contribution to preventing initiation with the 3GC mutant, as well as with the elongator tRNAs (Figures 4, 5 and 6A), we propose that a high abundance of initiator RNA is important not only in overcoming the rate-limiting step of initiation but also crucial in maintaining the fidelity of initiation by offering an effective discrimination against selection of the elongator tRNAs in the P-site (and thus initiation from codons unrelated to AUG). This direct relationship between fidelity and levels of initiator tRNA could also be an explanation behind the multiple copies of the initiator tRNA genes in various organisms.

Fluctuations in the initiator tRNA levels have been reported during diverse physiological conditions. It has been observed that metZWV is downregulated during stringent response by ppGpp (24). Likewise, metY is downregulated by the global regulator, cAMP–CAP complex and also by ArgR, a specific regulator of arginine metabolism (31). In addition, studies have shown that there could be major decreases in the charged initiator tRNA levels in E. coli during leucine starvation (32). However, the significance of such observations has remained unknown. Although in yeast, it is known that depletion of initiator tRNA levels leads to increased translation of GCN4, and GCN4-dependent transcription of target genes (33). Also, it has been shown that the levels of initiator and elongator Met-tRNA are inversely correlated in cell proliferation versus quiescence. In quiescent cells, the initiator tRNA levels go down but the elongator tRNA levels go up (34). Under the special circumstances of nutritional deprivation, etc., it may be advantageous for the cell to
allow a ‘flexibility/relaxation’ in the fidelity of protein synthesis. Such a proposal has parallels in other biological processes. For example, fidelity of replication is paramount to the maintenance of the genetic blue print of the organism. Yet, during SOS repair/replication, error-prone incorporation of nucleotides by DNA polymerases IV and V makes an important contribution to the viability of the cell (35). Likewise, a major reprogramming of gene expression by transcriptional and translational regulation is a well-established phenomenon during conditions of stress or starvation (36). However, in these cases of proteome remodeling, the issue of possible ‘relaxation in fidelity’ for initiation has never been raised. Our preliminary computational analysis shows that in E. coli (and possibly all other eubacteria), Shine–Dalgarno (S. D.) like sequences are found in the context of a variety of non-AUG codons. This observation, together with the possibility of downregulation of the initiator tRNA abundance under stress/starvation conditions, makes it feasible that relaxation in the fidelity of initiation may offer a mechanism to contribute to regulation of the proteome remodeling/plasticity in the cell.

An accurate selection of the initiator tRNA on the ribosomal P-site is a combined outcome of several coordinated events. For example, in eubacteria, formylation of initiator tRNA which facilitates its binding to IF2, and effectively decreases its off rate upon IF3 binding (12) is perhaps one of the most crucial contributors to the specificity of the initiator tRNA selection on the ribosome. However, this feature alone is neither sufficient nor essential for the initiator tRNA selection on the ribosomal P-site. For instance, despite the fact that 3GC mutant is fully formylated in E. coli (17), it fails to initiate to any significant level in the wild-type strain background. And although they suffer a mild to severe impact on the growth rates, it is known that knockouts of the formylase gene in Pseudomonas aeruginosa and E. coli are viable (37,38). Furthermore, in Salmonella mutants, efficient initiation could occur in the absence of formylation. Interestingly, the mutants compensated for the deficiency of formylation by amplification of the initiator tRNA genes (39). Among the other mechanisms that contribute to the fidelity of initiation are the rates with which the 50S and 30S subunit interact with each other (12), and possible effects of the modified nucleosides in rRNA (both in the 30S and 50S subunits) (17,40). In this scenario, the role played by the abundance of the initiator tRNA adds another level of complexity to the fidelity maintenance.

Given the numerous factors that could influence the translation initiation fidelity, a redundancy between these is an expected possibility. Although it is difficult to determine the relative contribution of various factors, it is feasible that in organisms having a single copy of functional initiator tRNA gene (such as mycobacteria) or a single initiator tRNA gene lacking the full complement of the 3GC base pairs in their anticodon stems (such as certain species of mycoplasma), other mechanisms compensate for the decreased fidelity. Alternatively, the organism’s survival strategy in the host might itself dictate a requirement of a certain ‘relaxed’ fidelity of translation. Thus, a study of translation fidelity in these organisms (mycoplasma and mycobacteria) would be rewarding not only from the point of view of understanding basic life processes but also to gain insight into their pathogenic behavior.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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