Analysis of genomic tRNA revealed presence of novel genomic features in cyanobacterial tRNA

Tapan Kumar Mohanta a,1,⇑, Dhananjay Yadav b,1, Abdullatif Khan a, Abeer Hashem c,d, Elsayed Fathi Abd_Allahe, Ahmed Al-Harrasi a

a Natural and Medical Sciences Research Center, University of Nizwa, Nizwa 616, Oman
b Dept. of Medical Biotechnology, Yeungnam University, Gyeongsan 38541, Republic of Korea
c Botany and Microbiology Department, College of Science, King Saud University, P.O. Box 2460, Riyadh 11451, Saudi Arabia
d Mycology and Plant Disease Survey Department, Plant Pathology Research Institute, Agriculture Research Centre, Giza, Egypt
e Plant Production Department, College of Food and Agriculture Science, King Saud University, P.O. Box 2460, Riyadh 11451, Saudi Arabia

Abstract
Transfer RNAs (tRNA) are important molecules that involved in protein translation machinery and acts as a bridge between the ribosome and codon of the mRNA. The study of tRNA is evolving considerably in the fields of bacteria, plants, and animals. However, detailed genomic study of the cyanobacterial tRNA is lacking. Therefore, we conducted a study of cyanobacterial tRNA from 61 species. Analysis revealed that; cyanobacteria contain thirty-six to seventy-eight tRNA gens per genome that encodes for 20 tRNA isotypes. The number of iso-acceptors (anti-codons) ranged from thirty-two to forty-three per genome. tRNAIle with anti-codon AAU, GAU, and UAU was reported to be absent from the genome of Gleocapsa PCC 73,106 and Xenococcus sp. PCC 7305. Instead, they were contained anti-codon CAU that is common to tRNAMet and tRNAIle as well. The iso-acceptors ACA (tRNACys), ACC (tRNAGly), AGA, ACU (tRNASer), AAA (tRNAPhe), AGG (tRNAPro), AAC (tRNAVal), GCG (tRNAArg), AUG (tRNAHis), and AUC (tRNAAsp) were absent from the genome of cyanobacterial lineages studied so far. A few of the cyanobacterial species encode suppressor tRNAs, whereas none of the species were found to encode a selenocysteine iso-acceptor. Cyanobacterial species encode a few putative novel tRNAs whose functions are yet to be elucidated.

1. Introduction

The study of tRNA biology has brought an unprecedented level of understanding as well as various novel discoveries in the fields of genetics and molecular biology. The principal function of tRNA is to transfer the amino acids to the ribosome machinery during the process of protein synthesis (Hopper and Phizicky, 2003; Kirchner and Ignatova, 2014). Essentially, tRNA functions as an adaptor molecule and delivers amino acids to the ribosome by reading the genetic information encoded within the mRNA (Mohanta and Bae, 2017; Rihas de Pouplana and Dedon, 2014; Xiaolong and Enduo, 2013). The tRNAs are universally found in every cell type and are the most abundant small non-coding cellular molecules, constituting approximately 4–10% of all cellular RNA. The importance and evolution of tRNA are based on its comprehensive relationship to the origin and development of the genetic code; the discovery of the universal nature of the genetic code in prokaryotes and eukaryotes has been incredibly important to the field of molecular biology (Koonin and Novozhilov, 2009). In 1966, Robert W. Holley first proposed the clover leaf model of tRNA containing four arms and three loops. These components are the acceptor arm, the D-arm, the D-loop, the anti-codon arm, the variable loop, the Ψ-arm, and the Ψ-loop (Clark, 2006; Kirchner and Ignatova, 2014). Nuclear encoded eukaryotic and prokaryotic tRNAs have a variable length of 73–90 nucleotides (nts) (Kirchner and Ignatova, 2014; Sharp et al., 1985). The acceptor arm contains 7 bp, the D-arm 3–4 bp, the anti-codon (AC) arm 5 bp, the variable

Abbreviations: tRNA, transfer RNA; U, uridine; Ψ, pseudouridine; A, adenine; C, cytosine; G, guanine.

⇑ Corresponding author.
E-mail address: tapan.mohanta@unizwa.edu.om (T.K. Mohanta).
1 Contributed equally.
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a Natural and Medical Sciences Research Center, University of Nizwa, Nizwa 616, Oman
b Dept. of Medical Biotechnology, Yeungnam University, Gyeongsan 38541, Republic of Korea
c Botany and Microbiology Department, College of Science, King Saud University, P.O. Box 2460, Riyadh 11451, Saudi Arabia
d Mycology and Plant Disease Survey Department, Plant Pathology Research Institute, Agriculture Research Centre, Giza, Egypt
e Plant Production Department, College of Food and Agriculture Science, King Saud University, P.O. Box 2460, Riyadh 11451, Saudi Arabia
loop 4–23 bp, the Ψ-arm 5 bp and the Ψ-loop 7 bp. In some species, tRNA has variable numbers of nucleotide residues in the D-loop and the variable arms (Kirchner and Ignatova, 2014). Regardless, the variable arm starts after nucleotide 44, and the anti-codon is always numbered as nucleotides 34, 35, and 36, while the C-C-A tail is always numbered 74, 75, and 76 (Kirchner and Ignatova, 2014). The acceptor arm provides the point of attachment for the amino acid through the C-C-A tail. The D-arm is named for the presence of an unusual pyrimidine nucleotide called dihydrouracil, and it contains a synthetase site that recognizes the amino acid activating enzyme. The anti-codon loop reads the nucleotide codon of mRNA. The Ψ-arm is named for the presence of the uracil-pseudouridine nucleotide and plays a crucial role in the recognition of the ribosome. The genetic code includes sixty-four codons, of which sixty-one are sense codons, and three are nonsense codons. The sense codons are read by sixty-one tRNA iso-acceptors. To date, all of this information has been obtained from studies confined to the bacterial, plant and animal kingdoms, and only very limited genomic data are available on cyanobacterial tRNAs. The availability of the complete genome sequence of several cyanobacterial species prompted us to examine the cyanobacterial tRNA sequences. To this end, we have identified and analysed the tRNAs from sixty-one cyanobacterial species, and here, we reported the novel genomics details of cyanobacterial tRNAs.

2. Materials and methods

2.1. Retrieval of genomic tRNA sequence data

The annotated genomic tRNA sequences of the cyanobacterial species were downloaded from the Joint Genome Institute (JGI) Genome Portal (Nordberg et al., 2014). Based on the availability of the genome sequence data at the start of this study, we downloaded the tRNA sequences from sixty-one cyanobacterial species (Supplementary data 1). Further details of the species used in this study are shown in Supplementary Table 1.

2.2. tRNA analysis

The genomic tRNA sequences of all the species were submitted to the tRNAscan-SE server to determine whether all the downloaded sequences encoded tRNA or not. The analysis revealed the presence of a clover leaf-like structure tRNA in the studied sequences, thus confirming that the sequences encode tRNA. The statistical parameters used for the analysis of tRNAs in tRNAscan-SE were as follows: sequence source: bacterial/general tRNA; search mode: default; query sequence: formatted (Fasta); genetic code for tRNA isotype prediction: universal (Lowe and Eddy, 1997). The tRNAscan-SE server was used for the analysis because tRNAscan-SE produces only one false positive report per 15 gigabases (Lowe and Eddy, 1997). tRNAscan-SE is one of the most successful programs for identifying tRNAs in genome sequences. tRNAscan-SE utilizes EufindtRNA to search for conserved non-coding tRNA sequences and evaluates the co-variance in the tRNA. The algorithm of tRNAscan-SE can detect and identifying tRNAs at 99–100% accuracy with a very low error rate.

2.3. Prediction of tRNA clover leaf-like structure and novel tRNA

The clover leaf-like structures of cyanobacterial tRNAs were predicted using the tRNAscan-SE server, and the resulting novel tRNA structures were retained for further analysis. The numbers of nucleotides in the acceptor arm, the D-arm, the D-loop, the anti-codon arm, the variable loop, the Ψ-arm and the Ψ-loop were recorded by manual counting for individual tRNAs by observing the tRNA structures that resulted from tRNAscan-SE server.

3. Results and discussion

3.1. Cyanobacterial tRNA gene families are diverse among species

Analysis revealed, the cyanobacterial tRNA gene family ranged from thirty-six to seventy-eight tRNAs per genome (Supplementary Table 1). Xenococcus sp. PCC 7305 encoded the lowest number (36), while Nostoc sp. PCC 7107 encoded the highest number (78) of tRNAs (Mohanta et al., 2017). Cluster analysis of the number of tRNAs per genome showed four clusters (Supplementary Fig. 1). Most of the species were clustered within 40–50 tRNAs per genome, and some were clustered within 60–70 tRNAs, whereas only a few were clustered within 70–80 tRNAs per genome (Supplementary Fig. 1). The abundance of tRNAs in the cyanobacterial genome ranged from 0.59% to 1.62% with Fischereilla sp. PCC 9605 having the lowest (0.59%) abundance and Synechococcus elongatus PCC 7942 having the highest (1.62%) (Supplementary Table 1). Although the genome sizes of Calothrix sp. PCC 7103 and Calothrix desertica were 11.584 Mb and 11.416 Mb, respectively, the number of tRNAs accounted for them was only 0.63% and 0.62% of their genomes (Supplementary Table 1). Cluster analysis revealed that the distribution of tRNA percentage per genome is very dynamic (Supplementary Fig. 2). The number of iso-acceptors in the cyanobacterial genome ranged from thirty-two to forty-three (Supplementary Table 1). The lowest number (32) of iso-acceptors was found in Xenococcus sp. PCC 7305, while the highest number (43) of iso-acceptors was found in Oscillatoria nigro-viridis PCC7112. These numbers show that cyanobacteria do not encode all of the iso-acceptors in their genome and that none of the species were found to contain all of sixty-one iso-acceptors. At least twelve species were found to contain pseudo-tRNAs in their genomes (Supplementary Table 1). The Nostoc sp. PCC 7524 and Syctonema hofmannii UTEX 2349 suppressor tRNAs contain CUA anti-codon, whereas the Oscillatoria nigro-viridis PCC 7112 suppressor tRNA contains UUA. Cluster analysis revealed that the number and distribution of tRNA iso-acceptor fall within the range of 32 to 43 per genome, showing a greater correlation of iso-acceptor number than of the genome size (Supplementary Fig. 3).

The number of individual tRNA family members (isotypes) in the genomes of different species ranged from zero to 10 (Supplementary Table 2). The tRNAiso gene with AAU, GAU, and UAU anti-codons were found to be absent from the genome of Gleocapsa sp. PCC 73,106 and Xenococcus sp. PCC 7305 (Supplementary Table 2). Analyses revealed that, cyanobacterial tRNAiso also encode CAU anti-codon that codes for tRNAMet as well. In total, 53 tRNA genes were found to encode CAU anti-codon (Table 1). The absence of tRNA genes from certain genomes might be due to the independent deletion. The sequence constraints of tRNA function suggest that tRNAs are by large intolerant to mutation. Mutation in tRNA can occur at any stem or loop of the tRNA, and it is reported that more than 230 mutations have been detected in human tRNA and that all of them were found to be associated with various diseases (Guy et al., 2014; Yarham et al., 2010). Therefore, there is a possibility of mutation in the anti-codon loop, and hence absence of AAU, GAU, and UAU anti-codons in tRNAiso in the genome of cyanobacteria. The absence of AAU, GAU, and UAU anti-codons in tRNAiso forced the genome to encode the isoleucine codon by CAU anti-codon. In some organisms, the UCU tRNA anticodon translate to AGA and AGG codons. Hence, the UCU tRNA improves the efficiency of the translation machinery for the AGG codon. A few anti-codons were also found very rarely in the cyanobacterial
tRNA. For example, the anti-codons AGU (threonine) AAU (isoleucine) and AGC (alanine) were found only once in the sixty-four studied cyanobacterial species, whereas AAU (tyrosine) and UAU (isoleucine) were found only twice, and CUC (glutamate) was found only thrice (Table 1). Each of the AAU (asparagine), AAG (leucine), and UCG (arginine) iso-acceptors were found only four times. Thus, the iso-acceptors AGU, AAU, AAU, CUC, UAU, AAG, and UCG are the rarest forms of iso-acceptor in the cyanobacterial earlier Mark and Grosjean (2002) reported regarding the presence of CAU anti-codons in tRNAiso which is in accordance with our finding (Mark and Grosjean, 2002). Nucleotide C34 at wobble position of the anti-codon modified to lysidine and lysidine modified tRNA CAU gets charged in-vitro by isoleucine. When modified lysidine is replaced by unmodified C nucleotide, the resulting CAU anti-codon aminoacylated by methionine instead of isoleucine (Mark and Grosjean, 2002). Nucleotide at 44th position (1st position of variable arm) of tRNA CAU possibly play important role for tRNAfMet, fMet (N-formylmethionine) is a derivative of methionine where a formyl group is attached to the amino group. tRNAfMet is used for initiation of protein synthesis in prokaryotic organisms which are subsequently removed during post-translational modifications.

3.2. The nucleotide length of tRNAser was greater than that of other tRNAs

The nucleotide composition of the different tRNAs gene varied greatly. We found that the nucleotide length of cyanobacterial tRNAs ranged from fifty-five to ninety-six nucleotides. tRNA^Thr^ (gene id: 2508648704) of Xenococcus sp. PCC 7305 was found to be the smallest tRNA gene, containing only fifty-two nucleotides whereas tRNAasp found from Cylindrospermum stagnale (gene id: 2509768958) contained 96 nucleotides. At least 33 tRNAser genes were found to contain 92 nucleotides whereas 12 genes were found to contain 90 nucleotides (Supplementary Table 3). The nucleotide lengths of cyanobacterial tRNAiso were higher than those of other tRNAs (Supplementary Table 3). The nucleotide length for tRNAleu was the longest tRNA found in cyanobacteria.}

Table 1

| tRNA Isootypes | Iso-acceptors |
|----------------|--------------|
| Polar          |             |
| Asparagine     | AUU (4)      |
| Cysteine       | GCA (78)     |
| Glutamine      | CUC (11)     |
| Glycine        | ACC (0)      |
| Serine         | GGA (63)     |
| Threonine      | AGU (1)      |
| Tyrosine       | AUA (2)      |
| Non-polar      |             |
| Alanine        | GGC (67)     |
| Isoleucine     | GAU (120)    |
| Leucine        | GAG (68)     |
| Methionine     | CAU (146)    |
| Phenylalanine  | ATT (92)     |
| Proline        | GCC (62)     |
| Tryptophan     | CCA (83)     |
| Valine         | AAC (0)      |
| Positively charged |   |
| Arginine       | GGC (50)     |
| Histidine      | AUU (2)      |
| Lysine         | UAU (53)     |
| Negatively charged |   |
| Aspartic acid  | UUC (84)     |
| Glutamic acid  | UUC (84)     |
| Suppressor     | CUA (2)      |
| Selenocysteine | UCA (0)      |

The genomes of Oscillatoria formosa PCC 6407 and Oscillatoria nigro-viridis PCC7112 possess ten tRNAiso genes, which was the highest number of individual tRNA (isotypes) genes found in the cyanobacterial genome (Supplementary Table 2). The cyanobacterial species encoded only one or two tRNAiso genes in their genomes, and the numbers of tRNAiso, tRNAiso, and tRNAiso genes varied from one to two per genome (Supplementary Table 2). Only the genomes of Microchaeo sp. PCC 7126 and Nostoc sp. PCC 7107 were found to contain three tRNAiso genes. The abundance of tRNAiso and tRNAiso genes was lower than that of tRNAiso, tRNAiso, and tRNAiso genes (Supplementary Table 2). The abundance of tRNAiso genes was found to be much higher than that of tRNAiso, tRNAiso, and tRNAiso genes (Supplementary Table 2). From a total of sixty-four codons, the cyanobacterial kingdom encoded only fifty-four anti-codons (Table 1). The anti-codons ACA (cysteine), ACC (glycine), AGA and ACU (serine), AAA (phenylalanine), AGG (proline), AAC (valine), GCG (arginine), AUA, (histidine) and AUC (aspartate) were found to be absent. The UCA anti-codon for selenocysteine was also found to be absent in cyanobacterial tRNAs. However, Oscillatoria nigro-viridis PCC 7112 encoded UUA suppressor tRNA, whereas Nostoc sp. PCC 7524 and Scytonema hofmanni UTEX_2349 encoded UUA suppressor tRNA (Table 1). From a total of 146 tRNAiso, 125 were found to encode for tRNAMet. fMet (N-formylmethionine) is a derivative of methionine where a formyl group is attached to the amino group. tRNAMet is used for initiation of protein synthesis in prokaryotic organisms which are subsequently removed during post-translational modifications.

The nucleotide composition of the different tRNAs gene varied greatly. We found that the nucleotide length of cyanobacterial tRNAs ranged from fifty-five to ninety-six nucleotides. tRNA^Thr^ (gene id: 2508648704) of Xenococcus sp. PCC 7305 was found to be the smallest tRNA gene, containing only fifty-two nucleotides whereas tRNA^asp^ found from Cylindrospermum stagnale (gene id: 2509768958) contained 96 nucleotides. At least 33 tRNAser genes were found to contain 92 nucleotides whereas 12 genes were found to contain 90 nucleotides (Supplementary Table 3). The nucleotide lengths of cyanobacterial tRNAiso were higher than those of other tRNAs (Supplementary Table 3). The nucleotide length for tRNAleu was the longest tRNA found in cyanobacteria.
3.3. tRNAs contain a variable number of nucleotides in the arms and loops

Overall, the nucleotide length of the acceptor arm ranged from four to eight nucleotides (Supplementary Table 5). Only two of the 3092 studied tRNAs were found to contain five nucleotides in the acceptor arm, whereas fifteen were found to contain six nucleotides, and two were found to contain eight nucleotides (Supplementary Table 5). In contrast, 99.38% of the tRNAs were found to contain seven nucleotides in the acceptor arm. tRNA^{Cys} from Fischerella sp. PCC 9431 (gene id: 2512980262) was found to contain four nucleotides each in the acceptor arm, whereas tRNA^{Asp}, tRNA^{Asn}, tRNA^{Glu}, and tRNA^{Met} were found to contain six nucleotides in the acceptor arm (Supplementary Table 5). Interestingly, except for Cylindrospermum stagnale PCC 7417, no other tRNAs from any cyanobacterial species were found to contain eight nucleotides in the acceptor arm (Supplementary Table 5).

Study revealed, cyanobacterial tRNAs were found to contain 3'-CCA tail sequence in the tRNA genes. From 3092 tRNA genes, 916 (29.62%) genes were found to contain 3'-CCA sequence. Previous study led by Ardell and Hou (2016) also reported regarding the presence of 3'-CCA sequence in bacteria (Ardell and Hou, 2016). From the studied 146 tRNA^{Met} genes, 116 (79.45%) genes were found to contain 3'-CCA tail sequence and from 125 tRNA^{Asp} (initiator tRNA) genes, 109 (74.65%) genes were found to contain 3'-CCA sequence. Ardell and Hou (2016) reported 74.1% of prokaryotic initiation tRNA contain 3'-CCA sequence and the result from cyanobacterial initiator tRNA genes (74.65%) closely match with the report of Adrell and Hou (Ardell and Hou, 2016).

The D-arm of cyanobacterial tRNAs were found to contain two to four nucleotides (Supplementary Table 5). Of 3092 studied tRNAs, only three were found to contain two nucleotides in the D-arm. They were tRNA^{Cys} from Synechococcus elongatus PCC 7942 (gene id: 640711018) and tRNA^{Ser} from Cyanotohe sp. BHE3E ATCC 51472 (gene id: 2507502103) and from Pleurocapsa sp. PCC7319 (gene id: 2509700995). Overall, 29.04% of cyanobacterial tRNAs were found to contain three nucleotides, whereas 70.86% of tRNAs was found to contain four nucleotides in the D-arm. Although the majority of tRNA^{His} tRNA^{Ile} tRNA^{Leu} tRNA^{Lys} tRNA^{Met} tRNA^{Phe} tRNA^{Pro} and tRNA^{Thr} genes were found to contain four nucleotides, a few were found to contain three nucleotides in the D-arm. This result suggests that these tRNA families inherently possess four nucleotides in the D-arm, and a deletion of one nucleotide most likely caused the others to have only three nucleotides.

The nucleotide length of the D-loop in cyanobacterial tRNAs varied from seven to thirteen (Supplementary Table 5). Of 3092 studied tRNAs, only one tRNA^{Met} from Calothrix sp. PCC 7507 (gene id: 2505798780) was found to contain seven nucleotides in the D-loop. For tRNA^{Ala}, the D-loop was found to contain seven to eleven nucleotides (Supplementary Table 5).

The length of the anti-codon arm ranged from two to seven nucleotides. The majority of the tRNA genes contained five nucleotides in the anti-codon arm (Supplementary Table 5). However, some variation was observed in the length of the anti-codon arm in different cyanobacterial tRNAs. All of the cyanobacterial tRNA^{The} and tRNA^{Met} genes were found to contain five nucleotides in the anti-codon arm without any variation. Eight of the tRNA^{His} genes were found to contain four nucleotides in the anti-codon arm, while the remaining of the genes had only five nucleotides. In the case of tRNA^{Glu}, one gene was found to contain two nucleotides; ninety-two (72.44%) were found to contain four nucleotides, and thirty-four (26.77%) were found to contain five nucleotides (Supplementary Table 5). tRNA^{Met} genes were found to be the most variable with regard to the nucleotide composition in the anti-codon arm. They contained three to five nucleotides in the anti-codon arm. Among 204 tRNA^{Met} genes, twenty (9.80%) were found to contain three nucleotides, forty-four (21.56%) were found to contain four nucleotides, and the remaining 140 (68.62%) were found to contain five nucleotides. The anti-codon loop was found to contain five to nine nucleotides (Supplementary Table 5). However, more than 95% of the tRNA molecules examined were found to contain seven nucleotides in the anti-codon loop. tRNA^{Cys} from Microchae sp. PCC 7126 (gene id: 2509780286) had five nucleotides in the anti-codon loop. tRNA^{Glu} from Calothrix sp. PCC 7507 (gene id: 2505798780) and tRNA^{His} from Chroococcidiopsis thermals PCC 7203 (gene id: 2503610603) and Nostoc sp. PCC 7107 (gene id: 2503744091) were also found to contain five nucleotides in the anti-codon loop. Among 3092 tRNAs, sixty-nine (2.23%) tRNAs were found to contain nine nucleotides in the anti-codon loop. tRNA^{Cys} from Dactylococcopsis salina PCC 8305 (gene id: 2509553070) and tRNA^{His} from Nostoc sp. PCC7524 (gene id: 2509613151), and tRNA^{His} from Fischerella sp. PCC9431 (gene id: 2512981565) were found to contain at least one tRNA gene with nine nucleotides in the anti-codon loop. tRNA^{Cys} from Microcolus sp. PCC 7113 (gene id: 2509433893), Microcoleus vaginatus fGP-2 (gene id: 2506348930), Oscillatoria anacutemel PCC 6304 (gene id: 2509421382), and Pseudanabaena sp. PCC 6802 (gene id: 2507089029) were found to contain nine nucleotides in the anti-codon loop. tRNA^{Cys} genes from Leptolyngbya sp. PCC 7375 (gene id: 2509846846), Leptolyngbya sp. PCC 7375 (gene id: 2509846848), and Pseudanabaena sp. PCC 6802 (gene id: 2507089039) were also found to contain nine nucleotides in the anti-codon loop. Of a total of 204 tRNA^{Met} genes, sixty-one tRNAs were found to contain nine nucleotides in the anti-codon loop. Surprisingly, not a single tRNA was found to contain an even number (six, eight or ten) of nucleotides in the anti-codon loop.

The length of the variable loop of the cyanobacterial tRNAs ranged from three to twenty-three nucleotides (Supplementary Table 5). The variable loop of tRNA^{Cys} of Synechococcus elongatus PCC 7942 (gene id: 640711018) was found to contain only three nucleotides and, interestingly, that was found to contain only two nucleotides in the D-arm. For tRNA^{Ala}, all genes were found to contain five nucleotides in the variable loop with a minor variation. In the case of tRNA^{His}, most of the sequences were found to contain five nucleotides, while twenty of them were found to contain six nucleotides. tRNA^{Ala} of Microcolus sp. PCC 7113 genes (gene id: 2509433873) was found to contain only four nucleotides. The length of the variable loop in tRNA^{Cys} genes ranged from three to seven nucleotides (Supplementary Table 5). tRNA^{Gin} genes were found to contain either four or five nucleotides, and most of them were found to contain four nucleotides. The variable loop of the tRNA^{Gin} was found to contain four to seven nucleotides. Among 193 tRNA^{Gin} genes, 113 (58.54%) were found to contain five nucleotides, while the remaining eighty (41.45%) were found to contain four nucleotides. Seven of the tRNA^{Ala} genes were found to contain six nucleotides, while the remainder of the tRNA^{Ala} genes were found to contain five nucleotides in the variable loop. Among tRNA^{Ala} genes, eighteen (14.51%) were found to contain five nucleotides, and 106 (85.48%) were found to contain six nucleotides. The variable loops of tRNA^{Ala} genes varied from five to eighteen nucleotides; the number of nucleotides in the tRNA^{Ala} variable loop could be five, six, seven, nine, eleven, twelve, thirteen, fourteen, fifteen, sixteen and seventeen. These data showed that tRNA^{Ala} genes had eleven different nucleotide variants in the variable loop. tRNA^{Ala} genes contained five (27.55%) or six (71.65%) nucleotide while tRNA^{Met} genes were found to contain five to seventeen nucleotides in the variable loop. Among ninety-one tRNA^{Met} genes, except for...
two, all others were found to contain five nucleotides in the variable loop. The variable loops of tRNA<sup>Pro</sup> genes were found to contain five to six nucleotides. The variable loops of tRNA<sup>Thr</sup> genes were very diverse and contained four to twenty-three nucleotides (Supplementary Table 5). The variable loops of tRNA<sup>Val</sup> genes were found to contain four to six nucleotides. The majority of tRNA<sup>Thr</sup> genes were found to contain five nucleotides, while only seven of them were found to contain four nucleotides, and only six were found to contain six nucleotides (Supplementary Table 5). tRNA<sup>Thr</sup> genes contained four to six nucleotides in the variable loops. Among eighty-three tRNA<sup>Thr</sup> genes, fourteen were found to contain four nucleotides; one was found to contain six nucleotides, while the remaining sixty-eight (81.92%) were found to contain five nucleotides. tRNA<sup>Val</sup> genes were found to contain five to eighteen nucleotides in the variable loops. For tRNA<sup>Tyr</sup>, all genes were found to contain five nucleotides in the variable loop. Importantly, it was found that only tRNA<sup>Leu</sup> and tRNA<sup>Thr</sup> and tRNA<sup>Val</sup> genes contained ten or more nucleotides in their variable loops, whereas the remainder of the tRNA genes contained less than ten nucleotides in their variable loops. Previous reports suggest that the variable loop is less critical for the biological function of tRNA (Brennan and Sundaralingam, 1976; Kjems et al., 1989). However, the length of the variable arm is important for the recognition of tRNA by aminoacyl tRNA synthetase, and it also helps to maintain the stability of the tRNA. However, the long variable loop consisting of a helical stem loop emerges from the deep groove side of the dihydrouridine helix (Brennan and Sundaralingam, 1976). The long variable loop most possibly resulted from retaining a splicing-deficient intron (Kjems et al., 1989).

The length of the Ψ-arm ranged from two to six nucleotides. tRNA<sup>Leu</sup> of Calothrix sp. PCC 7507 (gene id: 2505802880) had only two nucleotides in the Ψ-arm. tRNA<sup>Thr</sup> from Chroococcidiopsis thermalis PCC 7203 (gene id: 2503610603) was found to contain only four nucleotides in the Ψ-arm. Apart from these tRNAs, all other tRNA genes were found to contain five nucleotides in the Ψ-arm. The fact that the Ψ-arm is the ribosome recognition site and, apart from a few tRNAs, all Ψ-arms were found to contain five nucleotides suggests that it is a conserved structural feature of the tRNA molecule.

The length of the Ψ-loop ranged from two to thirty nucleotides in the cyanobacterial tRNA. Approximately 3087 tRNA genes (99.83%) were found to contain seven nucleotides in the Ψ-loop. The tRNA<sup>Thr</sup> gene from Oscillatoria sp. PCC 10,802 (gene id: 2509509666) was found to contain nine nucleotides in the Ψ-loop (Fig. 1). Among tRNA<sup>Leu</sup> genes, the Ψ-loops of Anabaena cylindrica PCC 7122 (gene id: 2504131825) and Calothrix sp. PCC 7507 (gene id: 2505802880) were found to contain thirteen nucleotides (Fig. 1). The Ψ-loop, along with the Ψ-arm, recognizes the ribosome during protein synthesis. The presence of five nucleotides in the Ψ-arm and seven nucleotides in the Ψ-loop is a characteristic feature of tRNAs. However, the presence of a lower or a higher number of nucleotides in the Ψ-arm and the Ψ-loop is very intriguing. The existence of non-standard Ψ-arms and Ψ-loops suggests that further novel translational mechanisms that remain to be elucidated might be occurring in these species.

### 3.4. Based on the D-loop and the variable loop, cyanobacterial tRNA can be classified into two classes

tRNA research has received considerable attention since the inception of the clover leaf-like model by Robert W. Holley in 1965. Initially, tRNAs were classified based on the amino acid they transferred to the translation machinery. Further classification was then performed based on the number and position of introns present in the tRNA (Tocchini-Valentini et al., 2009). The acceptor arm, the D-arm, the D-loop, the anti-codon arm, the anti-codon loop, the variable loop, the Ψ-arm and the Ψ-loop all contain a specified number of nucleotides. The presence of specific numbers of nucleotides in different parts of the tRNA has its own significance and functional importance. Hence, it was very important for us to classify the tRNAs according to the distribution nucleotides in different parts of the tRNA molecule. As a result, we have classified the cyanobacterial tRNAs based on the nucleotide composition in the D-arm and the variable loop. Based on the number of nucleotides in the D-arm and the variable loop, we were able to classify these tRNAs into two classes, referred to as class I and class II tRNAs (Table 2). The D-arm of class I tRNAs has 2–4 nucleotides, whereas the D-arm of class II tRNAs has only four nucleotides. Similarly, the variable loops in class I tRNA have 0–7 nucleotides, whereas the tRNAs of class II have 8–23 nucleotides. Based on the D-arm nucleotide numbers, the tRNAs in class I were tRNA<sup>Asp</sup>, tRNA<sup>Glu</sup>, tRNA<sup>Ile</sup>, tRNA<sup>Leu</sup>, tRNA<sup>Met</sup>, tRNA<sup>Phe</sup>, tRNA<sup>Pro</sup>, tRNA<sup>Thr</sup>, tRNA<sup>Tyr</sup>, tRNA<sup>Val</sup>, and tRNA<sup>Val</sup>, whereas those in class II were tRNA<sup>Ala</sup>, tRNA<sup>Asp</sup>, tRNA<sup>Glu</sup>, tRNA<sup>His</sup>, tRNA<sup>Pro</sup>, tRNA<sup>Thr</sup>, tRNA<sup>Tyr</sup>, and tRNA<sup>Val</sup>. However, based on the variable loop nucleotide numbers, the tRNAs in class I were tRNA<sup>Ala</sup>, tRNA<sup>Asp</sup>, tRNA<sup>Ile</sup>, tRNA<sup>Leu</sup>, tRNA<sup>Met</sup>, tRNA<sup>Phe</sup>, tRNA<sup>Pro</sup>, tRNA<sup>Thr</sup>, tRNA<sup>Tyr</sup>, and tRNA<sup>Val</sup> (Table 2). Class I tRNAs have a short variable loop with no helical stem and hairpin structure, whereas class II tRNAs might possess an additional stem and hairpin loop.

### 3.5. Cyanobacteria encode putative novel tRNAs

The acceptor arm usually contains seven nucleotides. However, in our study, we found a few tRNAs that contained zero (Pseudanabaena sp. PCC 7367, gene id: 2504679288), four (Fischerella sp. PCC 9431, gene id: 2512980262); six (Rivularia sp. PCC 7116, gene id: 2,510,091,557 and others) or eight (Cylindrospermum stagnale PCC 7417, gene id: 2,509,767,605 and 2509767604) nucleotides (Supplementary Table 5, Fig. 2). At least fifteen tRNAs were found to contain six nucleotides in the acceptor arm (Supplementary Table 5). The amino acids bind to the 3’ end of the acceptor arm during the translation process, and the presence of a smaller number nucleotide in the acceptor arm might impact the binding efficiency of the tRNA to the cognate amino acid. Usually, the D-arm contains three to four nucleotides. However, in our study, we found a few tRNAs that contained only two nucleotides in the D-arm (Synechococcus elongatus PCC 7942, gene id: 640711018; Cyanotheca sp. BH63E ATCC 51472, gene id: 2507592103) (Fig. 3). Bovine and human mitochondrial tRNA are reported to lack the D-arm (Arcari and Brownlee, 1980; de Bruijn et al., 1980; Salinas-Giegé et al., 2015). In general, the anti-codon arm contains five nucleotides. However, we observed that cyanobacterial tRNA contained three (Chroococcidiopsis thermalis PCC 7203, gene id: 2503612106; Cyanobacterium aponinum PCC 10605, gene id: 2503745044, etc.), four or six (Leptolyngbya sp. PCC 7376, gene id: 2503887456; Nostoc sp. PCC 7524, gene id: 2509813153) nucleotides instead of the usual five nucleotides in the anti-codon arm (Fig. 4). Several of the tRNAs were found to have three or six nucleotides in the anti-codon arm. The tRNAs found to contain four nucleotides in the anti-codon arm were tRNA<sup>Asp</sup>, tRNA<sup>Glu</sup>, tRNA<sup>His</sup>, tRNA<sup>Pro</sup>, tRNA<sup>Thr</sup>, tRNA<sup>Tyr</sup>, and tRNA<sup>Val</sup> (Supplementary Table 5). However, except for tRNA<sup>Ala</sup>, no tRNAs were found to possess three nucleotides in the anti-codon arm. The anti-codon loop of tRNAs contains seven nucleotides. However, dramatic variation in the nucleotide composition was observed in our study. We discovered several tRNAs that contained more or fewer than seven nucleotides in the anti-codon loop. Some of the tRNAs were found to contain nine nucleotides in the anti-codon loop (Microcoleus sp. PCC 7113, gene id: 2509433893;
Calothrix desertica PCC 7102, gene id: 2510026762; Calothrix sp. PCC 7507, gene id: 2505802880; Microcoleus vaginatus FGP-2, gene id: 2503648930; Pseudanabaena sp.PCC6802, gene id: 2507089029; Chroococcidiopsis thermalis PCC 7203, gene id: 2503741210; Nostoc sp. PCC 7524, gene id: 2509813151 (Fig. 5).

The tRNAs those were found to contain nine nucleotides in the anti-codon loop had a variable number of nucleotides in the anti-codon arm (Fig. 5). The tRNAs those were found to contain only five nucleotides in the anti-codon loop were from the species Microchaete sp. PCC 7126 (gene id: 2509780286), Calothrix sp. PCC 7507 (gene id: 2505798780), Chroococcidiopsis thermalis PCC 7203 (gene id: 2503610603), and Nostoc sp. PCC 7107 (gene id: 2503744091) (Supplementary Figure 4). Interestingly, none of the tRNAs were found to contain either six or eight nucleotides in the anti-codon loop. Also, of interest, the anti-codon loop was found to be absent in Leptolyngbya sp. PCC 7376 (gene id: 2503887456), and Nostoc sp. PCC 7107 (gene id: 2503742551). The presence of such variation in the nucleotide sequences in the anti-codon loop suggests that there might be some novel protein translation machinery in cyanobacteria that remains to be elucidated.

The region of the tRNA between the anti-codon loop and the Ψ-arm, which has a bulge-like structure, is commonly known as the variable loop. In our study, we found that unlike the D-arm, the anti-codon arm and the Ψ-arm, which each contained a loop, the variable region also contained an arm and a loop structure. We referred to this arm as the variable arm and to the loop as

**Table 2**

Classification of cyanobacterial tRNAs based on the number of nucleotides in the D-arm and variable loop regions. Based on nucleotide variation in the D-arm, the tRNAs that contained two to four nucleotides in the D-arm were classified as class I, whereas those contained four nucleotides in the D-arm were classified as class II tRNAs. Similarly, based on the nucleotide variation in the variable region, the tRNAs that contained zero to seven nucleotides were classified as class I, whereas those that contained eight to twenty-three nucleotides were classified as class II tRNAs.

| tRNA class | No. of nucleotides in D-arm | tRNA class | No. of nucleotides in variable region |
|------------|-----------------------------|------------|-------------------------------------|
| Class I    | 2–4                         | Class I    | 0–7                                 |
| Class II   | 4                           | Class II   | 8–23                                |

The tRNAs those were found to contain nine nucleotides in the anti-codon loop had a variable number of nucleotides in the anti-codon arm (Fig. 5). The tRNAs those were found to contain only five nucleotides in the anti-codon loop were from the species Microchaete sp. PCC 7126 (gene id: 2509780286), Calothrix sp. PCC 7507 (gene id: 2505798780), Chroococcidiopsis thermalis PCC 7203 (gene id: 2503610603), and Nostoc sp. PCC 7107 (gene id: 2503744091) (Supplementary Figure 4). Interestingly, none of the tRNAs were found to contain either six or eight nucleotides in the anti-codon loop. Also, of interest, the anti-codon loop was found to be absent in Leptolyngbya sp. PCC 7376 (gene id: 2503887456), and Nostoc sp. PCC 7107 (gene id: 2503742551). The presence of such variation in the nucleotide sequences in the anti-codon loop suggests that there might be some novel protein translation machinery in cyanobacteria that remains to be elucidated.

The region of the tRNA between the anti-codon loop and the Ψ-arm, which has a bulge-like structure, is commonly known as the variable loop. In our study, we found that unlike the D-arm, the anti-codon arm and the Ψ-arm, which each contained a loop, the variable region also contained an arm and a loop structure. We referred to this arm as the variable arm and to the loop as
the variable loop (Supplementary Fig. 5). Unlike the D-arm, the anti-codon arm and the W-arm, the variable arm has base pairing, and the number of nucleotides in the variable arm ranged from four to seven, whereas the number of nucleotides in the variable loop ranged from five to nine, in contrast to the anti-codon loop. Some of the tRNAs that were found to contain the variable loop were Chroococcidiopsis thermals PCC7203 (gene id: 2503615137), Pseudanabaena sp.PCC6802 (gene id: 2507087115), Calothrix deser-
Fig. 4. The anti-codon loop of cyanobacterial tRNAs with five nucleotides. Several of the cyanobacterial tRNAs were found to possess only five nucleotides instead of the usual seven nucleotides in the anti-codon loop. The cyanobacterial tRNAs that contained five nucleotides in the anti-codon loop possessed either four or five nucleotides in the anti-codon arm.

tica PCC7102 (gene id: 2510025188), Cylindrospermum stagnale PCC7417 (gene id: 2509769172), Calothrix sp. PCC6303 (gene id:2504098647), Calothrix sp.PCC7507 (gene id: 2505801912), Calothrix sp. PCC7103 (gene id: 2507478119), Cyanobacterium stanieri PCC7202 (gene id: 2,503,365,868 and 2503366170), Cylindrospermum stagnale PCC7417 (gene id: 2509768958) was found to contain twenty-nine nucleotides in the tRNA-Ala-W-arm, whereas the tRNA-Glu-W-arm and nine nucleotides in the tRNA-Glu-W-loop (Fig. 1).

The canonical tRNAAla isodecoders associate with mRNA and regulate canonical tRNAAsp isodecoders. Interestingly, the tRNAAla gene from Synechococcus elongatus PCC 7942 (gene id: 640711018) was found to contain four nucleotides in the Ψ-arm. Alternatively, the tRNAAla gene from Cylindrospermum stagnale PCC 7417 (gene id: 2509768958) was found to contain twenty-nine nucleotides in the Ψ-loop (Fig. 1). These dynamic variations in the nucleotide length in different parts of the tRNA molecule & novel structure clearly reflect the presence of possible alternate translation machinery in the cyanobacteria. They might also be playing competitive role and performing extra-translational function as well by bypassing ribosomal machinery. Additionally, these could also bring strong tRNA structure and can interfere with the translation process. Francklyn and Minajigi (2010) reported that tRNA is a multi-functional molecule that participates in several processes of the cellular metabolism (Francklyn and Minajigi, 2010). Therefore, the presence of several putative novel forms of cyanobacterial tRNA indicates that cyanobacterial tRNA might be involved in diverse cellular metabolism in addition to the protein translation. Nucleotide sequence change in tRNA can bring changes in the efficiency of aminoacylation and modification of ribosome’s (Dale and Uhlenbeck, 2005; Ledoux et al., 2009; Siegfried et al., 2010). Suppressing the stop codon of isodecoders sequence from the iso-acceptor family of tRNA shows diverse ranges of suppression efficiencies in HeLa cell line (Geslain and Pan, 2010). The isodecoders sequence that do not occur naturally resulted best suppression efficiency, suggesting not all of the naturally occurring iso-decoder are ideal for best translational efficiency (Geslain and Pan, 2010). Charged tRNA can be used as a carrier of activated amino acids whereas uncharged tRNAs that activates protein kinase GCN2 that regulates translation during cellular nutrient availability (Dong et al., 2000). tRNA prevents premature apoptosis by binding to cytochrome C and it has potential to cleave to produce small fragments that can be used as small interfering RNA to regulate translation factors (Ivanov et al., 2011; Mei et al., 2010; Yamasaki et al., 2009). Non-canonical tRNAAla isodecoders associate with mRNA and regulate alternative 3' end formation, tRNAs are also interacts with diverse arrays of cellular protein involved in cellular metabolism where tRNA interacts with histone H3K9 or farnesyl-transferase mediated protein modification (Parisien et al., 2013b; 2013a). To exactly decipher the function of these putative novel tRNAs, first of all it needs to identify their cognate interacting partners that may be protein or RNA.
4. Conclusion

An analysis of cyanobacterial tRNA sequences has revealed the presence of a diverse and dynamic tRNA structure and genomic architecture of cyanobacterial tRNA. The analyses revealed the presence of putative novel tRNA structures in cyanobacteria. The presence of putative novel tRNAs in cyanobacteria is very interesting, and laboratory-based experimental analysis to confirm their functional role can be very promising. However, it is difficult to accurately determine the function of individual tRNA. Although we have gained considerable success in RNA sequencing approach, the presence of post-transcriptional modifications in tRNA makes it difficult to accurately sequence them. The presence of putative novel tRNA in cyanobacteria raises speculation about the presence of diverse evolutionary function that might not be directly linked to translation. Further, the presence of such putative novel tRNA gives a brief idea about their potential to change themselves due to evolutionary pressure.

5. Data availability

All the data used during this study was taken from publicly available genomic database and details are mentioned in the materials and methods section.

Declaration of Competing Interest

Authors don't have any competing of interest to declare

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Author’s contribution

TKM: Conceived the idea, performed the experiment and analysis, drafted and revised the manuscript, DY: revised the manuscript, AK: analysed & revised the manuscript, AH: drafted and revised the manuscript, EFA: drafted and revised the manuscript, AAH: Revised the manuscript.

Appendix A. Supplementary material

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