

Distinct subcellular localization of tRNA-derived fragments in the infective metacyclic forms of *Trypanosoma cruzi*

Larissa Reifur^{1,3}, Maria Rosa Garcia-Silva², Saloê Bispo Poubel¹, Lysangela Ronalte Alves¹, Paulo Arauco¹, Diane Kelly Buiar³, Samuel Goldenberg¹, Alfonso Cayota², Bruno Dallagiovanna^{1/+}

¹Instituto Carlos Chagas-Fiocruz, Curitiba, PR, Brasil ²Functional Genomics Unit, Institut Pasteur de Montevideo, Mataojo, MV, Uruguay

³Departamento de Patologia Básica, Setor de Ciências Biológicas, Universidade Federal do Paraná, Curitiba, PR, Brasil

Small non-coding RNAs derived from transfer RNAs have been identified as a broadly conserved prokaryotic and eukaryotic response to stress. Their presence coincides with changes in developmental state associated with gene expression regulation. In the epimastigote form of Trypanosoma cruzi, tRNA fragments localize to posterior cytoplasmic granules. In the infective metacyclic form of the parasite, we found tRNA-derived fragments to be abundant and evenly distributed within the cytoplasm. The fragments were not associated with polysomes, suggesting that the tRNA-derived fragments may not be directly involved in translation control in metacyclines.

Key words: *Trypanosoma cruzi* - tRNA-derived fragments - subcellular localisation

The non-infective epimastigote form of *Trypanosoma cruzi* undergoes metabolic and morphological adaptations to differentiate into the pathogenic metacyclic trypomastigote form, which causes Chagas disease in mammals (Figueiredo et al. 2000). Morphological changes are associated with and preceded by a shift in metabolism, which is correlated with the differential expression of several genes (Manning et al. 2009). Although the metabolism of the replicative epimastigote is fully active, with constitutive polycistronic transcription, starvation reduces the transcriptional rates of the RNA polymerases in metacyclines (Ferreira et al. 2008). Despite the reduced level of transcription, translation is detected in these non-replicative forms, indicating that gene expression control in metacyclines occurs primarily at the post-transcriptional level (reviewed in Haile & Papadopoulou 2007).

Considering the almost exclusive post-transcriptional control of gene expression in *T. cruzi*, it would be surprising if this organism did not have an alternative pathway to compensate for the absence of an RNA interference system. A myriad of small non-coding RNAs have been reported in trypanosomatids (Dumas et al. 2006, Garcia-Silva et al. 2010, Michaeli et al. 2012), canonical microRNAs, and siRNAs have not been detected in *T. cruzi* (Franzén et al. 2011). Short (20-35 nt) RNAs derived from tRNAs were first observed in cytoplasmic granules in the epimastigote form of *T. cruzi* after an ini-

tial fingerprint sequencing of 348 clones (Garcia-Silva et al. 2010). Although 26% of the sequenced clones represented tRNA-derived fragments, a more thorough sequencing of the small RNAs from epimastigotes showed that 65.3% of more than 282.000 clones represented tRNA-derived fragments (Franzén et al. 2011). The composition of the small RNA population in *T. cruzi* was strikingly different from that observed in *Trypanosoma brucei*, reflecting clear differences in the molecular biology of the two parasites (Michaeli et al. 2012). Nonetheless, the relative abundance of tRNA-derived fragments could be related to gene expression control under various types of cellular stress, as has been suggested for prokaryotes, yeast, mammalian cells, the protozoans *Giardia lamblia* and *Tetrahymena thermophila* (Lee & Collins 2005, Li et al. 2008, Pederson 2010).

In this study, we conducted further analyses of the relative abundance and subcellular localization of *T. cruzi* tRNA-derived fragments during the infective stage of the parasite. We used *T. cruzi* metacyclic trypomastigotes derived from Dm28c epimastigotes cultured in vitro, as described by Contreras et al. (1985). Total RNA was extracted using TRIzol (Invitrogen) and size fractionated on a denaturing 15% polyacrylamide gel electrophoresis (PAGE) gel. Subsequently, 18-40 nt RNAs were excised from the gel, purified and cloned as described by Garcia-Silva et al. (2010). To recover the small RNAs, specific oligonucleotide adaptors containing Ban 1 restriction sites were ligated to the 5' and 3' ends. The RNAs were then reverse transcribed, amplified by polymerase chain reaction for 10 cycles, Ban 1-digested, concatemerized, cloned into the pGEM T-easy vector (Promega Corp) and sequenced. Analyses were performed using the public GenBank (ncbi.nlm.nih.gov/genbank), GeneDB (genedb.org) and TriTrypDB (trityrpd.org/trityrpd) databases.

From a total of 844 clones analysed, 509 sequences aligned with the *T. cruzi* genome, whereas 2.16% showed no matches, indicating that these sequences may correspond to regions of the genome that have not been se-

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+ Corresponding author: brunod@tecpar.br

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quenced, exogenous contaminating DNA or differences between the reference strain and the strain used in this study. No sequences matched small nuclear RNA sequences, but 0.98% mapped to small nucleolar RNAs, 2.95% mapped to intergenic regions, 6.09% mapped to mRNAs and 24.56% mapped to rRNAs. Most of the cloned sequences (63.26%) were fragments derived from tRNAs that appeared to be the result of a specific cleavage at or around the anticodon loop (Supplementary data). The tRNA fragments averaged 33 nt in length and were mostly derived from the 3' end (86.96%) of a restricted group of isoacceptors. In contrast with the RNAs obtained from an epimastigote population sequenced by Garcia-Silva et al. (2010), the tRNA-derived fragments obtained in this study from metacyclines exhibited differences with respect to abundance (tRNA fragments composed 63.26% of the small RNAs in metacyclines versus 26% in epimastigotes), orientation (most of the tRNA fragments were derived from the 3' side of the tRNAs in metacyclines vs. the 5' side in epimastigotes) and origin (in metacyclines, fragments were derived mostly from tRNA^{Glu}, tRNA^{Thr} and tRNA^{Val}; in epimastigotes, they were generally derived from tRNA^{Asp} and tRNA^{Glu}). The relative abundances of most fragments did not correspond to either tRNA gene copy number or codon usage (Horn 2008, Padilla-Mejia et al. 2009), consistent with previous reports (Franzén et al. 2011). As reported by Franzén et al. (2011), we observed that the majority of the fragments were derived from the 3' side of the tRNAs and a significant proportion of these fragments (24.53%) carried the 3' CCA sequence, indicating that both mature and pre-tRNAs undergo the cleavage process. The biological importance of these tRNA fragments is unknown, but we expected that a higher percentage of these fragments would be observed in metacyclines because stressed epimastigotes showed only a slight increase in tRNA-derived fragments (Garcia-Silva et al. 2010). This phenomenon is likely observed because tRNA cleavage is a conserved process in cells under various types of stress and metacyclic trypomastigotes are the product of epimastigote differentiation triggered by nutritional stress (Contreras et al. 1985). A comparison of our results with those of Garcia-Silva et al. (2010) demonstrates that the percentage of tRNA fragments is clearly higher in metacyclines. In contrast, this difference is not observed by Franzén et al. (2011). Moreover, the epimastigote tRNA-derived fragments sequenced by Franzén et al. (2011) are mostly derived from the 3' arm of tRNA^{His}. These discrepancies could be due to differences in the strains analysed in the two studies and to the cloning and sequencing methods used. Franzén et al. (2011) used a different strain (CL Brener) and analysed a much larger number of clones using RNAseq, which provided a higher coverage of the parasite genome.

The cloned RNA sequences were aligned using the LocARNA server (Will et al. 2007) and the secondary structures identified using the RNAalifold server (Bernhart et al. 2008) from the Vienna RNA package and were adjusted manually according to the secondary structural domains of canonical tRNAs. The predicted secondary structures adopted by the most abundant 3' tRNA frag-

ments revealed that, upon cleavage around the anticodon nucleotides, the tRNA fragment maintained the TψC loop conformation and was extended a few extra base pairs into the double-stranded stem, resulting in an energetically favourable structure (Supplementary data). Although this result is based only on computational calculations and does not confirm the true molecular structure *in vivo*, the fact that the most abundant tRNA-derived fragments can assume a similar, relatively stable configuration suggests a structural (or functional) significance.

The subcellular localization of the tRNA-derived fragments in the metacyclic forms was evaluated through fluorescence *in situ* hybridisation (FISH) using probes complementary to the 5' and 3' ends of tRNA^{Glu-UUC} (the most abundantly cloned tRNA fragment). FISH assays showed that these fragments are dispersed throughout the cytoplasm in metacyclines, whereas stressed epimastigotes exhibited the same posterior granular distribution as non-stressed epimastigotes (Fig. 1). To further analyze the cellular localization of the tRNA-derived fragments, metacyclic cells were gently lysed in lysis buffer (300 mM KCl, 10 mM MgCl₂, 10 mM Tris-HCl 7.4 pH and 0.5% NP40) for 5 min. Two fractions were collected: the supernatant, containing only the soluble cytosol contents (S), and the insoluble fraction, or pellet (P), containing insoluble organelles, vesicles and cellular membranes. Both fractions were loaded on a sodium dodecyl sulphate-PAGE (10%) and transferred onto Hybond-C membranes (Amersham). Western blots were performed using antibodies to TcPUF6 (1/250), a cytosolic RNA binding protein (Dallagiovanna et al. 2005), and dynamin (1/200), a membrane protein present in endocytic vesicles (Pucadyil & Schmid 2009) (Fig. 2F, G). Northern blots were conducted using total RNA extracted from the two cell fractions (S and P) and a ra-

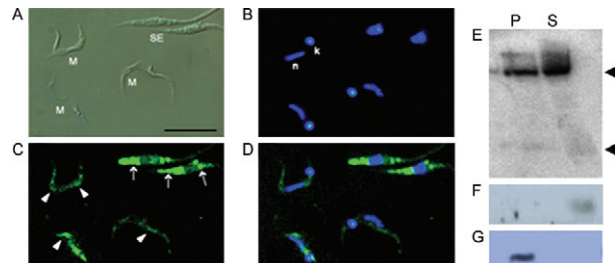


Fig. 1: subcellular localization of tRNA-derived RNAs in metacyclic trypomastigote forms. A: differential imaging contrast of metacyclic (M) and stressed epimastigote forms (SE); B: 4'-6-diamidino-2-phenylindole staining was used as a reference to identify nuclei (n) and kinetoplast (k); C: fluorescein isothiocyanate (FITC)-labeled probes recognizing the 3' fragment tRNA^{Glu-UUC} (5'TTCCGATACCGGAATCGA3') in SE (arrows) and M forms (arrowheads); D: merged images (Bar = 10 µm). Cells were observed under an Olympus IX 81 microscope coupled to a Hamamatsu Orca-ER camera (Diagnostic Instruments); E: total RNA from soluble (S) and insoluble (P) metacyclic cell fractions were analyzed with the radioactively labelled probe 3'tRNA^{Glu-UUC}, as in Garcia-Silva et al. (2010). Western blot assays with F (anti TcPUF6) and G (anti-Dynamin) sera were performed to assess the identity of both fractions.

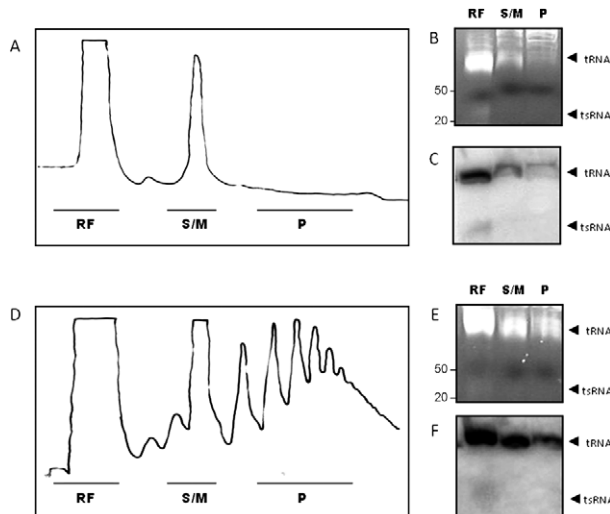


Fig. 2: tRNA-derived fragments are not associated with polysomes. Polysome profiles of (A) metacyclic trypomastigote and (D) epimastigote forms. Fractions were collected and pooled (RF: ribosome free; S/M: subunits and monomers; P: polysomes). RNA was purified and separated into 15% acrylamide gels (B, E). Northern blots of metacyclic (C) and epimastigote RNA (F) analyzed with the radioactively labelled probe tRNA^{Glu-UUC}, as in Fig. 1.

diolabeled probe specific to the tRNA^{Glu-UUC} fragment. The tRNA-derived fragments were detected in both fractions, supporting the dispersed pattern observed in the immunofluorescence assays (Fig. 2E). The signal observed for the insoluble cellular fraction could indicate the presence of these RNAs in smaller intracellular vesicles. Nonetheless, their dispersed pattern in the cytoplasm in metacyclines contrasts with the granular and posterior localization in epimastigotes, in which a partial co-localization with reservosomes has been inferred. The dismantling of the tRNA fragments in metacyclines could be correlated with the absence of reservosomes in this life stage (Figueiredo et al. 2000) and is consistent with the described cytoplasmic localization of the type II tRNA-derived fragments. Type II tRNA-derived fragments are most likely generated in the cytosol by RNaseZ cleavage and by RNA polymerase III termination and these fragments have been found to coimmunoprecipitate with Argonaute proteins (Elbarbary et al. 2009, Haussecker et al. 2010). This dramatic change in subcellular localization is puzzling and suggests distinct roles for these molecules during the different stages of the parasite life cycle.

To determine whether the dispersed tRNA-derived fragments were associated with translating polysomes, metacyclic and epimastigote forms of *T. cruzi* were treated with cycloheximide and the polysomes were purified and separated on sucrose gradients, as previously described (Nardelli et al. 2007) (Fig. 2A, D). Parasite extracts were also treated with puromycin as a negative control (not shown). The polysomes, monosomes and ribosome-free fractions were pooled and the total RNA was extracted. After acrylamide gel separation and staining, a high concentration of small RNAs was

observed in the ribosome-free pool for the metacyclic and epimastigote forms and a faint small RNA signal was observed in the ribosome-containing fractions (Fig. 2B, E). Northern blot analysis confirmed that tRNA-derived fragments from metacyclines and epimastigotes were concentrated in the ribosome-free fractions (Fig. 2C, F). Therefore, tRNA-derived fragments may not be related to the repression of the translational machinery. The knockdown of the tRNA-derived fragments and structural analysis should be used to help uncover the biological importance of these RNAs.

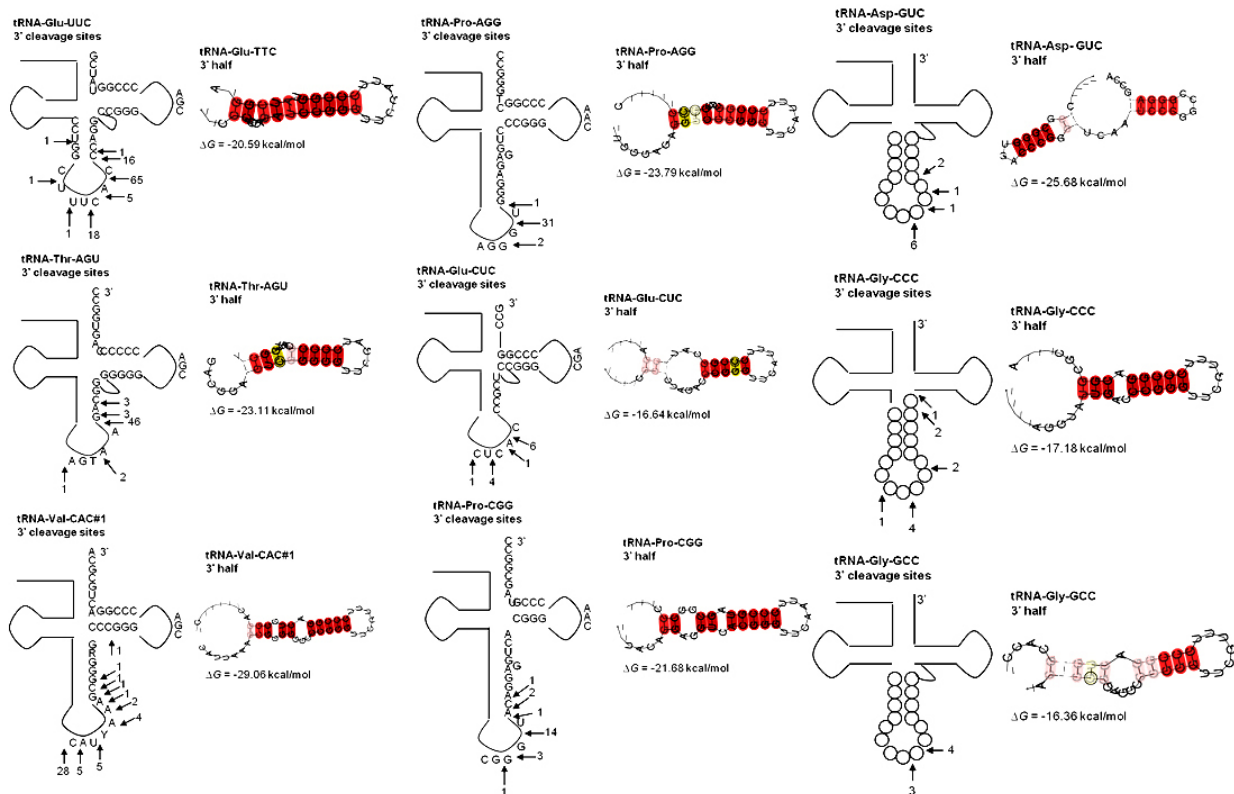
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Predicted secondary structures adopted by the most abundant 3' tRNA fragments.

TABLE

List of cloned tRNA-derived fragments, their identity obtained by Basic Local Alignment Search Tool against database sequences from *Trypanosoma cruzi* and if the sequence matches either the 5', mid portion or 3' portion of the tRNA

Clone	tRNA	Cloned sequence
332	tRNA-Glu:anticodon TTC, 3'	ACCAGGAGATCCGGGTTCGATTCCCGGTATCGGA
634	tRNA-Glu:anticodon TTC, 3'	ACCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA
70	tRNA-Glu:anticodon TTC, 3'	ACCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA AC
713	tRNA-Glu:anticodon TTC, 3'	ACCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA AC
831	tRNA-Glu:anticodon TTC, 3'	ACCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA CA
247	tRNA-Glu:anticodon TTC, 3'	ACCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA CA
172	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTTC
102	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTTC
803	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTCGATTCCCGGTATCG
685	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTCGATTCCCGGTATCGA
133	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA
150	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA
513	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA
334	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA
614	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA AC
637	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA AC
686	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA CA
5	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA CA
703	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTTGATTCCCGGTATCGGA
387	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTTGATTCCCGGTATCGGA
444	tRNA-Glu:anticodon TTC, 3'	CAGGAGATCCGGGTTCGATTCCCGGTATCGGA
124	tRNA-Glu:anticodon TTC, 3'	CCAGGAAATCCGGGTTCGATTCCCGGTATCGGA AC
487	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTCGATTCCCGGTATCCCGG
397	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTCGATTCCCGGTATCG
352	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTCGATTCCCGGTATCGG
456	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTCGATTCCCGGTATCGG
481	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTCGATTCCCGGTATCGG
516	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTCGATTCCCGGTATCGG
319	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTCGATTCCCGGTATCGGA
380	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTCGATTCCCGGTATCGGA
399	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTCGATTCCCGGTATCGGA
791	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTCGATTCCCGGTATCGGA
792	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTCGATTCCCGGTATCGGA
833	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTCGATTCCCGGTATCGGA CA
549	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTCGATTCCCGGTATCGGA CA
311	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTCGATTCCCGGTATCGGAT
722	tRNA-Glu:anticodon TTC, 3'	CCAGGAGATCCGGGTTTGATTCCCGGTATCGGA CA
702	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGGATAGGA
472	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGGATCGG
427	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGTATCG
87	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGTATCGG
248	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGTATCGG
331	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGTATCGG
390	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGTATCGG
429	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGTATCGG
506	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGTATCGG
602	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGTATCGG
779	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGTATCGG
793	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGTATCGG
542	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGTATCGG
117	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA
151	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA



Clone	tRNA	Cloned sequence
164	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
252	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
354	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
402	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
403	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
407	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
408	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
416	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
419	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
424	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
439	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
452	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
466	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
480	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
483	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
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492	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
494	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
505	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
507	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
511	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
515	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
519	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
701	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
727	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
734	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
749	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
751	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
776	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
790	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
834	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
840	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
141	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
25	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
357	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
7	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA C
652	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA C
710	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA CC
626	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA CC
656	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA CC
342	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA CC
647	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA CC
9	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA CC
479	tRNA-Glu:anticodon TTC, 3'	CCCAGGAGATCCGGGTTTGATTCCC GG TATCGG
484	tRNA-Glu:anticodon TTC, 3'	CCCGCTAGACCCGGATTCA TTCCC GG CATCGGA
774	tRNA-Glu:anticodon TTC, 3'	CCCTCGAGATCCGGGTTCGATTCCC GG TCTCGGA CTCAGGAGATCCGGGTTCGATTCCC CAA
375	tRNA-Glu:anticodon TTC, 3'	TCCAGGAGATCCGGGTTCGATTCCC GG TATC
283	tRNA-Glu:anticodon TTC, 3'	TCCGATATGGTATAACGGTTAGAACACCTGGC
760	tRNA-Glu:anticodon TTC, 3'	TCCGATATGGTATAACGGTTAGAACACCTGGC
708	tRNA-Glu:anticodon TTC, 3'	TCCGATATGGTATAACGGTTAGAACACCTGGC
442	tRNA-Glu:anticodon TTC, 3'	TCCGATATGGTATAACGGTTGGAGCCAACAGGC
348	tRNA-Glu:anticodon TTC, 3'	CACCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA
184	tRNA-Glu:anticodon TTC, 3'	TTCACCCAGGAGATCCGGGTTCGATTCCC GG TATCGGA C
314	tRNA-Glu:anticodon TTC, 3'	



Clone	tRNA	Cloned sequence
339	tRNA-Glu:anticodon TTC, 5'	CACCCAGGAGATCCGGGTTCGATTCCCGGTATCGGA ACC
723	tRNA-Glu:anticodon TTC, 5'	CCGATATGGTATAACGGTTAGAACACCCGGC
860	tRNA-Glu:anticodon TTC, 5'	CCGATATGGTATAACGGTTAGAACACCTGG
778	tRNA-Glu:anticodon TTC, 5'	CCGATATGGTATAACGGTTAGAACACCTGGC
282	tRNA-Glu:anticodon TTC, 5'	CCGATATGGTATAACGGTTAGAACACCTGGC
520	tRNA-Glu:anticodon TTC, 5'	CCGATATGGTATAACGGTTAGAACACCTGGC
580	tRNA-Glu:anticodon TTC, 5'	CGATATGGTATAACGGTTAGAACACCTGG
471	tRNA-Glu:anticodon TTC, 5'	TCCGATATGGTATAACGGTT
623	tRNA-Glu:anticodon TTC, 5'	TCCGATATGGTATAACGGTTAGAACACCTGGC
358	tRNA-Glu:anticodon TTC, 5'	TCCGATATGGTATAACGGTTAGAACACCTGGC
425	tRNA-Glu:anticodon TTC, 5'	TCCGATATGGTATAACGGTTAGAACACCTGGC
495	tRNA-Glu:anticodon TTC, 5'	TCCGATATGGTATAACGGTTAGAACACCTGGC
568	tRNA-Glu:anticodon TTC, 5'	TCCGATATGGTATAACGGTTAGAACACCTGGC
225	tRNA-Thr:anticodon AGT, 3'	AAAGGAGGTGGGGGTTCGATCCCCCCCAGTGGCC
601	tRNA-Thr:anticodon AGT, 3'	AAGACGGAGGTGGGGGTTCGATCCCCCCCAGTGGC
817	tRNA-Thr:anticodon AGT, 3'	AAGACGGAGGTGGGGGTTCGATCCCCCCCAGTGGC CCA
355	tRNA-Thr:anticodon AGT, 3'	ACGGAGGTGGGGGTTCGATCCCCCCCAGTGGCC
512	tRNA-Thr:anticodon AGT, 3'	ACGGAGGTGGGGGTTCGATCCCCCCCAGTGGCC
251	tRNA-Thr:anticodon AGT, 3'	CGGAGGTGGGGGTTCGATCCCCCCCAGTGGCC
398	tRNA-Thr:anticodon AGT, 3'	CGGAGGTGGGGGTTCGATCCCCCCCAGTGGCC
720	tRNA-Thr:anticodon AGT, 3'	CGGAGGTGGGGGTTCGATCCCCCCCAGTGGCC
795	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTCGATCCCCCCCAGTGGC
781	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGCGGC
567	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGCGGC
595	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
78	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
160	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
185	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
443	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
474	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
490	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
498	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
526	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
546	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
575	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
635	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
719	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
728	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
761	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
846	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
851	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
13	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
56	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
63	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
85	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
551	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
61	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
2	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
96	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
120	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
318	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
423	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
455	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC
504	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTGGGGGTTAGATCCCCCCCAGTGGC



Clone	tRNA	Cloned sequence
578	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTCTGGGGGTTCGATCCCCCCAGTGGCCT
721	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTCTGGGGGTTCGATCCCCCCAGTGGCCT
745	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTCTGGGGGTTCGATCCCCCCAGTGGCCT
780	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTCTGGGGGTTCGATCCCCCCAGTGGCCT
176	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTCTGGGGGTTCGATCCCCCCAGTGGCCT
345	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTCTGGGGGTTCGATCCCCCCAGTGGCCTC
295	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTCTGGGGGTTCGATCCCCCCAGTGGCCTCA
530	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTCTGGGGGTTCGATCCCCCCAGTGGCCTCA
785	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTCTGGGGGTTCGATCCCCCCAGTGGCCTCA
593	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTCTGGGGGTTCGATCCCCCCAGTGGCCTGA
799	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTCTGGGGGTTCGATCCCCCCAGTGGCCT
818	tRNA-Thr:anticodon AGT, 3'	GACGGAGGTCTGGGGGTTCGATCCCCCCAGTGGCCT
271	tRNA-Val:anticodon CAC#1, 3'	AAAGCGGGGGGCCGGGTTCTGTTCCGGACTGCGCAC
784	tRNA-Val:anticodon CAC#1, 3'	AAAGCGGGGGGCCGGGTTCTGTTCCGGACTGCGCAC
737	tRNA-Val:anticodon CAC#1, 3'	AAAGCGGGGGGCCGGGTTCTGTTCCGGACTGCGCACCC
321	tRNA-Val:anticodon CAC#1, 3'	AAAGCGGGGGGCCGGGTTCTGTTCCGGACTGCGCACCA
577	tRNA-Val:anticodon CAC#1, 3'	AAGCGGGGGGCCGGGTTCTGTTCCGGACTGCGCAC
532	tRNA-Val:anticodon CAC#1, 3'	ACGCGGGAGGCCGGGTTCTGTTCCGGACTGCGCAC
246	tRNA-Val:anticodon CAC#1, 3'	CAAAGCGGGGGGCCGGGTTCTGATTCCGGACTGCGCAC
197	tRNA-Val:anticodon CAC#1, 3'	CAAAGCGGGGGGCCGGGTTCTGTTCCGGACTGCGCAC
581	tRNA-Val:anticodon CAC#1, 3'	CAAAGCGGGGGGCCGGGTTCTGTTCCGGACTGCGCAC
137	tRNA-Val:anticodon CAC#1, 3'	CAAAGCGGGGGGCCGGGTTCTGTTCCGGACTGCGCAC
261	tRNA-Val:anticodon CAC#1, 3'	CAAAGCGGGGGGCCGGGTTCTGTTCCGGACTGCGCAC
564	tRNA-Val:anticodon CAC#1, 3'	CATCAAAGCGGGGGGCCGGGTTCCGGACTGCGCACCA
356	tRNA-Val:anticodon CAC#1, 3'	CGCGGGAGGCCGGGTTCTGTTCCGGACTGCGCA
679	tRNA-Val:anticodon CAC#1, 3'	CGGGGGGCCGGGTTCTGTTCCGGACTGCGCAC
540	tRNA-Val:anticodon CAC#1, 3'	CGGGGGGGGCCGGGTTCTGATTCCGGACTGCGCAC
289	tRNA-Val:anticodon CAC#1, 3'	GGGCCGGGTTCTGTTCCGGACTGCGCACCAT
717	tRNA-Val:anticodon CAC#1, 3'	GGGGGGGCCGGGTTCTGTTCCGGACTGCGCACCC
450	tRNA-Pro:anticodon AGG, 3'	GTGGGAGAGGTCCCGGGTTTCCCAGGCTGGGCC
861	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTTCCCAGGCTGGGCC
393	tRNA-Pro:anticodon AGG, 3'	ACAGGAGGTACGGGTTCATTCGGCTGGGCC
335	tRNA-Pro:anticodon AGG, 3'	GGGAGAGGTCCCGGGTTCATTCGGCTGGGCC
410	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
92	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
180	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
527	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
250	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
67	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
69	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
337	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
361	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
370	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
379	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
384	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
400	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
517	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
651	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
665	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
762	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
775	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
191	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
349	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC
422	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCCGGGTTCAATTCCGGCTGGGCC



Clone	tRNA	Cloned sequence
431	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCC GGTTCA TTCCGGCTGGGCC C
458	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCC GGTTCA TTCCGGCTGGGCC C
664	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCC GGTTCA TTCCGGCTGGGCC C
189	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCC GGTTCA TTCCGGCTGGGCC C
272	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCC GGTTCA TTCCGGCTGGGCC CC
814	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCC GGTTCA TTCCGGCTGGGCC CC
824	tRNA-Pro:anticodon AGG, 3'	TGGGAGAGGTCCC GGTTCA TTCCGGCTGGGCC CC
396	tRNA-Pro:anticodon AGG, 3'	TGGGGAGGTCCC GGTTCA TTCCGGCTGGGCC C
496	tRNA-Glu:anticodon CTC, 3'	ACCCGCTAGACCCGGGTT C. ATTCCCGGCATCGGA
159	tRNA-Glu:anticodon CTC, 3'	CACCCAGGAGATCCGGGTT CATTCCCGGTATCGG
128	tRNA-Glu:anticodon CTC, 3'	CACCCGCTAGACCCGGGTT CATTCCCGGCATCGGAA
669	tRNA-Glu:anticodon CTC, 3'	CACCCGCTAGACCCGGGTT CATTCCCGGCATCGGAA CCA
765	tRNA-Glu:anticodon CTC, 3'	CACCCGGTAGACCCGGGTT CATTCCCGTCATCGGAA CCA
338	tRNA-Glu:anticodon CTC, 3'	CCCAGGAGATCCGGGTT CATTCCCGGTATCGGA
378	tRNA-Glu:anticodon CTC, 3'	CCCAGGAGATCCGGGTT CATTCCCGGTATCGGA
453	tRNA-Glu:anticodon CTC, 3'	CCCAGCTAGACCCGGGTT CATTCCCGGCATCGG
486	tRNA-Glu:anticodon CTC, 3'	CCCAGCTAGACCCGGGTT CATTCCCGGCATCGGA
571	tRNA-Glu:anticodon CTC, 3'	CCCAGCTAGACCCGGGTT CATTCTCGGCATC
820	tRNA-Glu:anticodon CTC, 3'	CCCAGCTAGACCCGGGTT CATTCTCGGCATC
536	tRNA-Glu:anticodon CTC, 5'	CCGGTGTGGTATA GTGGTTAGAACAAAGCGGC
692	tRNA-Glu:anticodon CTC, 5'	CCGGTGTGGTATA GTGGTTAGAACAAAGCGG
598	tRNA-Glu:anticodon CTC, 5'	CCGGTGTGGTATA GTGGTTAGAACAAAGCGGC
340	tRNA-Glu:anticodon CTC, 5'	TCCGGTGTGGTATA GTGGTTAGAACAAAGCGGC
543	tRNA-Glu:anticodon CTC, 5'	TCCGGTGTGGTATA GTGGCTAGAACAAAGCGGC
259	tRNA-Glu:anticodon CTC, 5'	TCCGGTGTGGTATA GTGGCTAGAACAAAGCGGC
426	tRNA-Glu:anticodon CTC, 5'	TCCGGTGTGGTATA GTGGCTAGAACAAAGCGGC
454	tRNA-Glu:anticodon CTC, 5'	TCCGGTGTGGTATA GTGGCTAGAACAAAGCGGC
508	tRNA-Glu:anticodon CTC, 5'	TCCGGTGTGGTATA GTGGCTAGAACAAAGCGGC
343	tRNA-Glu:anticodon CTC, 5'	TCCGGTGTGGTATA GTGGCTAGAACAAAGCGGC
437	tRNA-Glu:anticodon CTC, 5'	TCCGGTGTGGTATA GTGGCTAGAACAAAGCGGC
459	tRNA-Glu:anticodon CTC, mid	TAGTGGTTAGAACAAAGCGGC
666	tRNA-Asp:anticodon GTC, 3'	ACGGGGTGACCCGGGTT CA ATTCCCGGCCGGGAAG CCA
239	tRNA-Asp:anticodon GTC, 3'	CACGCGGGAGACCCGGGTT CCTT
612	tRNA-Asp:anticodon GTC, 3'	CACGCGGGTGACCCGGGTT CA ATTCCCGGCCGGGA
752	tRNA-Asp:anticodon GTC, 3'	CACGCGGGTGACCCGGGTT CA ATTCCCGGCCGGGAAG CCA
77	tRNA-Asp:anticodon GTC, 3'	CACGCGGGTGACCCGGGTT CA ATTCCCGGCCGGGAAG CCA
131	tRNA-Asp:anticodon GTC, 3'	CGCGGGTGACCCGGGTT CA ATTCCCGGCCGGGAAG CC
430	tRNA-Asp:anticodon GTC, 3'	CGGGGTGACCCGGGTT CA ATTCCCGGCCGGGA
447	tRNA-Asp:anticodon GTC, 3'	CGGGGTGACCCGGGTT CA ATTCCCGGCCGGGA
155	tRNA-Asp:anticodon GTC, 3'	CACGCGGGTGACCCGGGTT CA ATTCCCGGCCGGGAAG CCA
156	tRNA-Asp:anticodon GTC, 3'	CACGCGGGTGACCCGGGTT CA ATTCCCGGCCGGGAAG CCA
620	tRNA-Asp:anticodon GTC, 5'	CTCGGTAGTATA ATGGTAAGTATACCGCC
4	tRNA-Asp:anticodon GTC, 5'	CTCGGTAGTATA ATGGTAAGTATACCGCC
57	tRNA-Asp:anticodon GTC, 5'	CTCGGTAGTATA ATGGTAAGTATACCGCC
500	tRNA-Asp:anticodon GTC, 5'	CTCGGTAGTATA ATGGTAAGTATACCGCC
46	tRNA-Asp:anticodon GTC, 5'	CTCGGTAGTATA ATGGTAAGTATACCGCC
254	tRNA-Asp:anticodon GTC, 5'	TCGGTAGTATA GTGGTAAGTATACCGCC
320	tRNA-Asp:anticodon GTC, 5'	TCGGTAGTATA GTGGTAAGTATACCGCC
88	tRNA-Asp:anticodon GTC, 5'	TCTCGGTAGTATA GTGGTAAGTATACCGCC
678	tRNA-Asp:anticodon GTC, 5'	TCTCGGTAGTATA GTGGTAAGTATACCGCC
363	tRNA-Asp:anticodon GTC, 5'	TCTCGGTAGTATA GTGGTAAGTATACCGCCT
741	tRNA-Pro:anticodon CGG, 3'	AGGAGGTACGGGTT CA ATTCCCGTAGCGGCC
368	tRNA-Pro:anticodon CGG, 3'	CAGGAGGTACGGGTT CA ATTCCCGTAGCGGCC
470	tRNA-Pro:anticodon CGG, 3'	CAGGAGGTACGGGTT CA ATTCCCGTAGCGGCC



Clone	tRNA	Cloned sequence
697	tRNA-Pro:anticodon CGG, 3'	TACAGGAGGTACGGGTTCATTCCTCGTAGCGGC
140	tRNA-Pro:anticodon CGG, 3'	TACAGGAGGTACGGGTTCAATTCCCGTAGCGGCC
71	tRNA-Pro:anticodon CGG, 3'	TACAGGAGGTACGGGTTCAATTCCCGTAGCGGCC
351	tRNA-Pro:anticodon CGG, 3'	TACAGGAGGTACGGGTTCAATTCCCGTAGCGGCC
636	tRNA-Pro:anticodon CGG, 3'	TACAGGAGGTACGGGTTCAATTCCCGTAGCGGCC
350	tRNA-Pro:anticodon CGG, 3'	TACAGGAGGTACGGGTTCATTCCTCGTAGCGGCC
420	tRNA-Pro:anticodon CGG, 3'	TACAGGAGGTACGGGTTCATTCCTCGTAGCGGCC
457	tRNA-Pro:anticodon CGG, 3'	TACAGGAGGTACGGGTTCATTCCTCGTAGCGGCC
465	tRNA-Pro:anticodon CGG, 3'	GTACAGGAGGTACGGGTTCAATTCCCGTAGCGGCC
521	tRNA-Pro:anticodon CGG, 3'	GTACAGGAGGTACGGGTTCAATTCCCGTAGCGGCC
84	tRNA-Pro:anticodon CGG, 3'	TACAGGAGGTACGGGTTCAATTCCCGTAGCGGCC
256	tRNA-Pro:anticodon CGG, 3'	TACAGGAGGTACGGGTTCATTCCTCGTAGCGGCC
385	tRNA-Pro:anticodon CGG, 3'	ACAGGAGGTACGGGTTCATTCCTCGTAGCGGCC
523	tRNA-Pro:anticodon CGG, 3'	GGTACAGGAGGTACGGGTTCAATTCCCGTAGCGGCC
497	tRNA-Pro:anticodon CGG, 3'	GTACAGGAGGTACGGGTTCAATTCCCGTAGCGGCC
369	tRNA-Pro:anticodon CGG, 3'	TACAGGAGGTACGGGTTCAATTCCCGTAGCGGCC
802	tRNA-Pro:anticodon CGG, 3'	TACAGGAGGTACGGGTTCAATTCCCGTAGCGGCC
449	tRNA-Pro:anticodon CGG, 3'	TACAGGAGGTACGGGTTCAATTCCCGTAGCGGCC
460	tRNA-Pro:anticodon CGG, 3'	TACAGGAGGTACGGGTTCATTCCTCGTAGCGGCC
341	tRNA-Pro:anticodon TGG, 3'	TACAGGAGGTACGGGTTCAATTCCCGTAGCGGCC
138	tRNA-Gly:anticodon CCC, 3'	AGGTATTGACCCGGGCTCGTTCCGGACGGCGCACCA
152	tRNA-Gly:anticodon CCC, 3'	AGGTATTGACCCGGGTTCGTTCCGGACGGCGCA
16	tRNA-Gly:anticodon CCC, 3'	CAAGGTATTGACCCGGGTTCGATTCCCGAACGGCGCA
34	tRNA-Gly:anticodon CCC, 3'	CAAGGTATTGACCCGGGTTCGATTCCCGAACGGCGCA
590	tRNA-Gly:anticodon CCC, 3'	CAAGGTATTGACCCGGGTTCGTTCCGGACGGCGCACCA
114	tRNA-Gly:anticodon CCC, 3'	CCAAGGTATTGACCCGGGTTCGTTCCGGACGGCGCA
223	tRNA-Gly:anticodon CCC, 3'	TGACCCGGGTTCGTTCCGGACGGCGC
473	tRNA-Gly:anticodon CCC, 3'	TTGACCCGGGTTCGTTCCGGACGGCGC
40	tRNA-Gly:anticodon CCC, 3'	TTGACCCGGGTTCGTTCCGGACGGCGC
325	tRNA-Gly:anticodon CCC, 3'	ACCCAGGAGATCCGGGTTCGATTCCCGACTGCGCACCA
296	tRNA-Gly:anticodon GCC, 3'	ACCGGGAGGCCCCGGGTTCGTTCCCGACTGCGCACCA
233	tRNA-Gly:anticodon GCC, 3'	ACCGGGAGGCCCCGGGTTCGTTCCCGACTGCGCACCA
51	tRNA-Gly:anticodon GCC, 3'	ACCGGGAGGCCCCGGGTTCGTTCCCGACTGCGCACCA
857	tRNA-Gly:anticodon GCC, 3'	CACGCGGGAGGCCCCGGGTTCGTTCCCGACTGCGCACCA
236	tRNA-Gly:anticodon GCC, 3'	CACGCGGGAGGCCCCGGGTTCGTTCCCGACTGCGCACCA
74	tRNA-Gly:anticodon GCC, 3'	CACGCGGGAGGCCCCGGGTTCGTTCCCGACTGCGCACCA
53	tRNA-Gly:anticodon GCC, mid	ACGAGGGAGGCCGGGTT
798	tRNA-His:anticodon GTG, 3'	CTCTGAATACCCGGGTTCGATTCCCGTCTCCCTCCA
418	tRNA-His:anticodon GTG, 3'	CTGAATACCCGGATTGATTCGATTCCCGTCTCCCTC
336	tRNA-His:anticodon GTG, 3'	GTTCGATTCCCGGTCTCCCT
260	tRNA-Lys:anticodon TTT, 3'	AACCGTGTGGTCGTGGGTTCGATCCCCACGGATGGCAC
801	tRNA-Ala:anticodon AGC, 3'	TGCGGGAGGTATTGGGATCGTTACCCAACCTCTCC
10	tRNA-Ala:anticodon CGC, 3'	TACGGAAGGCCTAGGGTTCGATCCCCTACTCGTCCA
301	tRNA-Ala:anticodon CGC, 3'	TACGGAAGGCCTAGGGTTCGATCCCCTACTCGTCCA
208	tRNA-Val:anticodon CAC#2, 5'	GCGATGGTCGTCTAGGGGTTAGGACATTCTC
446	tRNA-Val:anticodon CAC#2, 5'	GCGATGGTCGTCTAGGGTTAGGACATTGCC
856	tRNA-Val:anticodon CAC#2, 5'	GCGATGGTCGTCTAGGGTTAGGACATTCGC
395	tRNA-Arg:anticodon TCG, 3'	CAGAGGGTTGCAGGTTCGGATCCTGTACGGAT
381	tRNA-Gly:anticodon TCC, 5'	GCAATCGTGGTCAACGGTTAAGATCCCCGCC
428	tRNA-Gly:anticodon TCC, 5'	GCAATCGTGGTCAACGGTTAAGATCCCCGCC
573	tRNA-Val:anticodon AAC 5'	GCGCTGATGGCTAGGTGGTTATGACGTCGTTTG
421	tRNA-Val:anticodon AAC, 3'	GACACGGCGAAGGTCTCGGGTTCGGGTCCGATCGGC
518	tRNA-Trp:anticodon CCA, 3'	TCCAGGGGTCGCAGGTTCAATCCCTGAGTCCTC
475	tRNA-Tyr:anticodon GTA, 5'	CCTCTGTAGCTAATTGGTAGAGCATGTGACTGTAG

the non-encoded nucleotides are in red.