



## Arabidopsis tRNA-derived fragments as potential modulators of translation

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1   **Arabidopsis tRNA-derived fragments as potential modulators of**  
2   **translation**

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13  
14   **ABSTRACT**

15   Transfer RNA-derived fragments (tRFs) exist in all branches of life. They are involved  
16   in RNA degradation, regulation of gene expression, ribosome biogenesis. In  
17   archaeabacteria, kinetoplastid, yeast and human cells, they were also shown to regulate  
18   translation. In Arabidopsis, the tRFs population fluctuates under developmental or  
19   environmental conditions but their functions are yet poorly understood. Here, we show  
20   that populations of long (30-35 nt) or short (19-25 nt) tRFs produced from Arabidopsis  
21   tRNAs can inhibit *in vitro* translation of a reporter gene. Analyzing a series of  
22   oligoribonucleotides mimicking natural tRFs, we demonstrate that only a limited set of  
23   tRFs possess the ability to affect protein synthesis. Out of a dozen of tRFs, only two  
24   deriving from tRNA<sup>Ala</sup>(AGC) and tRNA<sup>Asn</sup>(GUU) strongly attenuate translation *in vitro*.  
25   Contrary to human tRF(Ala), the 4 Gs present at the 5' extremity of Arabidopsis  
26   tRF(Ala) are not implicated in this inhibition while the G<sub>18</sub> and G<sub>19</sub> residues are  
27   essential. Protein synthesis inhibition by tRFs does not require complementarity with  
28   the translated mRNA but, having the capability to be associated with polyribosomes,  
29   tRFs likely act as general modulation factors of the translation process in plants.

30  
31   **KEYWORDS:** protein synthesis; polyribosomes; small non-coding RNAs; tRNAs; plant

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33

1     **Introduction**

2         Beyond their primordial role in protein synthesis, transfer RNAs (tRNAs) appear  
3         to have many other multiple functions [1] [2]. Among them, the production of regulatory  
4         small non-coding RNAs (sncRNAs) called tRFs (tRNA-derived Fragments) has been  
5         put forward in the last few years. These tRFs have various origins within tRNA  
6         transcripts. They can originate from tRNA precursor molecules, but most of them derive  
7         from mature tRNAs (for reviews see e.g. [3] [4]). Mature tRNAs generate two main  
8         classes of tRFs depending on the cleavage site: long tRFs (l-tRFs, 30-35 nt in size)  
9         when cleavage occurs in the anticodon region and short ones (s-tRFs, 19-25 nt) if the  
10        cleavage is located in the D or T regions of tRNA molecules. Several nomenclatures  
11        exist and we will use here the nomenclature we have proposed in [3] where l-tRFs  
12        mainly correspond to tRF-5A or tRF-3A and s-tRFs to tRF-5D or tRF-3T whether the  
13        RNA fragments derive from the 5' or 3' extremities of mature tRNAs.

14         The cleavage of tRNAs likely represents an essential process to recycle  
15        molecules such as phosphate or nitrogen upon nutrient starvation. Nevertheless, in all  
16        evolutionary divergent organisms from bacteria to human cells, a growing list of other  
17        important functions have been assigned to tRFs (for a review, see [2]). For instance,  
18        numerous l-tRFs were often shown to be upregulated upon stress ([5] [6] [7] and  
19        references therein). Others are involved in tumorigenesis [8] and hematopoiesis [9].  
20         The implication of tRFs to promote RNA degradation [10], to prime viral reverse  
21        transcription [11] or to regulate gene expression via the RNA silencing pathway (e.g.  
22        [12] [13]) has also been described. Recent work also showed that tRFs can regulate  
23        epigenetic inheritance [14], the expression of retro-elements [15], or plant nodulation  
24        [16].

25         Regulation of translation appears to be the most documented process where  
26        tRFs are involved. Upon abiotic stress, l-tRFs produced in human cells promote stress  
27        granule assembly and translation repression [17]. Reduction of translation was also  
28        observed when some human stress-induced l-tRFs, including tRF-5A (Ala) and tRF-  
29        5A (Cys), can displace the eIF4F factor of the initiation elongation complex [18]. Also  
30        in human, Sobala and Hutvagner [19] showed that some human s-tRFs (e.g. a tRF-5D  
31        (Gln) of 19 nt), can inhibit protein synthesis. More recently, a proteomic approach  
32        allowed us to demonstrate that this tRF can interact with the multi-synthetase complex  
33        (MSC), a complex interacting with ribosomes [20]. Furthermore, in the yeast  
34        *Saccharomyces cerevisiae* [21] and the archaea, *Haloferax volcanii* [22], stress-

1 induced tRFs can bind ribosomes and inhibit translation *in vitro*. By contrast, it was  
2 recently demonstrated that in the kinetoplastid *Trypanosoma brucei*, a tRF-3A (Thr)  
3 produced during nutrient deprivation can associate with ribosomes and subsequently  
4 stimulate translation [23].

5 The existence of tRFs has also been proven in various plants (e.g. [7] [13] [24]  
6 [25] [26] [27]). In *Arabidopsis thaliana*, the tRFs population varies upon stress and plant  
7 development and some are specifically enriched in Argonaute (AGO)  
8 immunoprecipitates [28] [29]. This suggests that tRFs likely also play key roles in  
9 photosynthetic cells. Currently, except for the study reporting that *Arabidopsis* AGO1-  
10 associated tRFs can target and cleave transposable elements, little is known about  
11 their functions [30].

12 Some work on pumpkin provided evidence that a phloem small non-coding RNA  
13 (sncRNA) population triggers translation inhibition *in vitro* [24]. This sncRNA population  
14 comprises many tRFs, and here we have addressed the question of whether plant tRFs  
15 could act as a regulator of protein synthesis. We observed that both tRFs purified from  
16 *A. thaliana* leaves and specific synthetic tRFs deriving from the 5' extremity of  
17 *Arabidopsis* tRNA<sup>Ala</sup> and tRNA<sup>Asn</sup> efficiently inhibit protein synthesis *in vitro*. This  
18 inhibition is cap-independent and does not require a sequence complementarity with  
19 the mRNA to be translated. Furthermore, analysis of *Arabidopsis* ribosomal fractions  
20 shows that tRFs can bind and be associated with heavy polysomes, suggesting their  
21 involvement in global regulation of the translation process in the land plant.

22

## 23 **Results**

### 24 ***Arabidopsis short and long tRFs inhibit protein synthesis in vitro***

25 We previously showed that *Arabidopsis* tissues (roots, stems, leaves, flowers, siliques)  
26 contain complex and variable pools of tRFs [28]. Furthermore, several RNases T2,  
27 called RNS1, 2 and 3, expressed in these tissues are involved in the biogenesis of  
28 both I-tRFs and s-tRFs [31]. Indeed, incubation of *in vitro* synthesized cytosolic tRNA<sup>Ala</sup>  
29 transcript or *Arabidopsis* total tRNAs with a crude enzymatic extract from *Arabidopsis*  
30 leaves produces I-tRF (Ala) and s-tRF (Ala), (Figures 1A and 1B). As a first step to  
31 knowing whether plant tRFs can modulate translation, I- and s-tRFs pools were  
32 produced by incubating total *Arabidopsis* tRNAs in the presence of a crude enzymatic  
33 extract from *Arabidopsis* leaves (Figure 1C). According to our previous results, this  
34 enzymatic extract contains the RNS involved in tRFs production in *Arabidopsis* [31],

1 and the tRFs population must correspond to the previously identified one when  
2 analyzing sncRNA libraries [28]. Then, an *in vitro* coupled transcription/translation  
3 wheat germ system was used to test the effect of the tRFs pools on the translation of  
4 a mRNA coding for the pGFP (*i.e.* GFP fused at its N-terminal extremity to a  
5 mitochondrial targeting sequence) in the presence of  $^{35}\text{S}$  methionine. No difference in  
6 protein synthesis efficiency, as compared to the mock sample, was observed when  
7 purified Arabidopsis tRNA fractions were added to the system (Figure 1D). By contrast,  
8 both types of tRFs efficiently inhibit the production of radiolabeled pGFP when added  
9 to the wheat germ system. Only two concentrations of tRFs (4 and 8  $\mu\text{M}$ ) were used,  
10 but the data obtained suggest that the inhibition displays dose-dependency. At a  
11 concentration of 8  $\mu\text{M}$ , l- or s-tRFs can inhibit up to 95% of pGFP expression (Figure  
12 1E). Globally, each set of Arabidopsis tRFs (long or short) inhibits translation. However,  
13 both the Arabidopsis l- and s-tRFs populations are generated from a vast collection of  
14 Arabidopsis tRNAs [32], and the question of whether all of them can modulate protein-  
15 coding gene expression needed to be addressed.

16

17 ***Only a limited set of Arabidopsis tRFs affects translation in vitro***

18 The Arabidopsis tRFs are highly variable in origin, sequence, and length, with a  
19 majority deriving from the 5' or 3' extremities of mature tRNAs [28]. On the one hand,  
20 we previously identified a specific population of tRFs associated with AGO1, thus  
21 suggesting their implication at least in the regulation of gene expression *via* the RNA  
22 silencing pathway. On the other hand, abundant tRFs that were not found in AGO  
23 precipitates could be involved in other molecular processes, including regulation of  
24 translation as shown in other organisms. As a first attempt to know whether translation  
25 inhibition is a general function of tRFs, we synthesized ten sequences (Figure 2A;  
26 Supplemental Table S1) representative of the Arabidopsis tRFs population that we  
27 previously analyzed in [28]. These sequences correspond to three tRFs that were  
28 shown to be among the most abundant ones associated with AGO1. The two tRFs-3T  
29 reflect two different abundances: Thr-AGU is rather abundant whereas Gly-GCC is  
30 present in low amount in sncRNA libraries. Finally, among the five selected tRF-5, Ala-  
31 AGC, Gly-GCC, and Glu-CUC belong to the most abundant tRFs and by contrast, Asn-  
32 GUU and Cys-GCA are present in a low amount. We tested them at two concentrations  
33 (0.2 and 1  $\mu\text{M}$ ) for their ability to inhibit the translation of pGFP using the *in vitro* coupled  
34 transcription/translation wheat germ system (Figures 2B and 2C). As control

1 experiments, two random oligoribonucleotides of 15 and 21 nt corresponding to an  
2 internal sequence of GFP have no significant effect on expression (Supplementary  
3 Figure 1). The three tRFs previously found associated with AGO1 in leaves have no  
4 effect on protein expression (rather a slight positive effect of pGFP synthesis was  
5 noted). For five others [e.g. tRF-5D (Gly-GCC), tRF-5A (Cys-GCA) or tRF-3T (Thr-  
6 AGU)], inhibition of expression is also not observed. Only two tRFs of 20 nt, namely  
7 tRF-5D (Ala-AGC) and tRF-5D (Asn-GUU) strongly inhibit GFP expression already at  
8 a concentration of 0.2  $\mu$ M, inhibition reaching 75-90% at 1  $\mu$ M. It is important to note  
9 that several isomers exist for each of these tRFs *in vivo*. Indeed, from the 33 tRNA<sup>Ala</sup>  
10 and the 15 tRNA<sup>Asn</sup> Arabidopsis nuclear genes, various tRFs of 20 nt can be generated  
11 (Supplementary Figure 2) and are found in sncRNA libraries [28]. For simplification,  
12 these two tRFs are now called Ala20 and Asn20 and correspond to one specific isomer  
13 (see Supplementary Figure 2). To have an idea of the amount of Ala20 versus tRNA<sup>Ala</sup>  
14 *in vivo*, we quantified them by northern blot in an Arabidopsis total tRNA extract and  
15 we observed that roughly there is  $10^4$  more tRNAs than tRFs in the extract  
16 (Supplementary Figure 3). It is to note that tRFs-5D (Ala) are the most abundant tRFs  
17 found in sncRNA libraries, whereas tRF-5D (Asn) is 15 times less present than Ala20  
18 [28]. In the wheat germ extract, the tRNA concentration is about 20  $\mu$ M (Supplementary  
19 Figure 3), thus to keep physiological conditions, we should have a 2 nM tRF  
20 concentration. However, in our assays, we used 0.2 to 1  $\mu$ M. There is an excess of  
21 tRFs but we also cannot exclude that, *in vivo*, there is a higher localized concentration  
22 within the cell. Though this analysis has been performed at a small scale, our  
23 observations indicate that Arabidopsis tRFs behave differently (from null to strong  
24 effect) in their capacity to modulate expression. Interestingly, none of the major tRFs  
25 associated with AGO1 affects mRNA expression and more, the tRF capacity to  
26 modulate expression is not correlated with their abundance in the total Arabidopsis  
27 tRFs population. Thus, likely only a limited set of tRFs are efficient inhibitors, among  
28 them, Asn20 and Ala20.

29

30 **tRF Ala20 inhibits translation by a cap- and sequence-independent mechanism**

31 Interestingly, a stress-induced tRF of 29 nt deriving from human tRNA<sup>Ala</sup> and called 5'-  
32 tiRNA<sup>Ala</sup>, was shown to strongly inhibit translation [17, 18]. We thus wondered whether  
33 inhibition by tRF (Ala) is an evolutionary conserved molecular process and focused our  
34 work on Ala20. In Arabidopsis, under normal growth conditions, tRF-5 of 16, 20 and

1 33 nt deriving from tRNA<sup>Ala</sup> are abundant [28]. Thus, in addition to Ala20, we further  
2 tested two other synthetic oligoribonucleotides corresponding to tRF-5D of 16 nt  
3 (Ala16) and 33 nt (Ala33) deriving from tRNA<sup>Ala</sup> (Figures 3B, C and D). In contrast to  
4 Ala20, no significant inhibition of pGFP translation is observed when Ala16 is added in  
5 the transcription/translation system at a concentration of 1  $\mu$ M, and a 40% decrease of  
6 expression is observed at a 4  $\mu$ M concentration. When Ala33 is added in the  
7 transcription/translation system, an intermediate inhibitory effect is observed with a  
8 40% decrease of expression at 1  $\mu$ M. Altogether, this suggests that the size and/or the  
9 sequence are important for determining the efficiency of inhibition. Yet, no tRF was  
10 shown to be implicated in the inhibition of both transcription and translation. The  
11 coupled transcription/translation system does not allow us to discriminate at which step  
12 the tRF is implicated. We thus performed *in vitro* translation experiments with a wheat  
13 germ extract translation kit where no transcription is possible. In this case, in the  
14 presence of *in vitro* synthesized capped or uncapped pGFP mRNA reporter, a similar  
15 important decrease in pGFP synthesis is observed when Ala20 or Asn20 is present  
16 (Figure 3E). This further demonstrates that the decrease in protein synthesis is not due  
17 to inhibition of transcription but rather that the translation process is indeed affected by  
18 the addition of the tRF. Finally, to exclude the possibility that the Ala20 is inhibiting  
19 translation by directly interfering with the sequence of the pGFP mRNA (although no  
20 sequence similarity has been bioinformatically detected), an *in vitro*  
21 transcription/translation assay was performed in the presence of another recombinant  
22 plasmid coding for pDHFR (the mammalian DiHydroFolateReductase fused to a  
23 mitochondrial targeting sequence), [33]. As for pGFP translation, pDHFR synthesis is  
24 not affected by GFP15, GFP21 (Supplementary Figure 1) and Ala16 but is efficiently  
25 repressed by the addition of Ala20 (Figure 3F). This shows that plant tRF-5D Ala20  
26 inhibits translation by a transcript sequence-independent mechanism.

27

28 ***Conserved G<sub>18</sub> and G<sub>19</sub>, and A<sub>16</sub> are important for tRF-mediated translation  
29 inhibition in Arabidopsis***

30 Several residues of mammalian tRFs (also called tiRNA) were identified as important  
31 for translation inhibition. On the one hand, the four G residues present at the 5'-end of  
32 5'-tiRNA<sup>Ala</sup> and 5'-tiRNA<sup>Cys</sup> can form a G-quadruplex that binds the YB1 protein. This  
33 complex then displaces eIF4F complex from the cap and inhibits translation initiation  
34 [18]. Arabidopsis Ala20 also starts with four Gs (Figure 3A). To determine if these

1 residues are also involved in translation inhibition in plants, we replaced them by Us  
2 (Ala20-5U, Figure 4A). As shown in Figures 4B and 4C, the inhibition of translation is  
3 not affected when a synthetic Ala20-5U is added to the wheat germ system, thus  
4 demonstrating that in Arabidopsis, contrary to human tRF-5 (Ala), the G1 to G4  
5 residues are not essential for tRF-mediated translation inhibition. This result is in  
6 agreement with the work of Nowacka and collaborators [34] showing that human tRF-  
7 5 Ala does not inhibit translation in a wheat germ extract. On the other hand, Sobala  
8 and Hutvagner showed that in several human tRFs-5D, the conserved two Gs present  
9 in the D loop at positions 18 and 19 of tRNAs are important for translation inhibition  
10 [19]. Therefore, we also replaced G<sub>18</sub> and G<sub>19</sub> by Us in Ala20 (Ala20-3U). As compared  
11 to the wild type sequence, the synthetic tRFAla20-3U has lost its capacity to promote  
12 translation inhibition (Figures 4B and 4C), suggesting that these two residues are  
13 important for tRF(Ala)-mediated translation inhibition. To confirm these data, when the  
14 two Gs located at the same position in Asn20 oligoribonucleotide (Figure 4A) are  
15 replaced by Us, the same loss in translation inhibition is observed (Figure 4D).  
16 Furthermore, sequence comparison between Ala20 and Asn20 (the two tRFs that  
17 efficiently inhibit translation) and the sequence of Arg19 (unable to reduce translation),  
18 reveals a high degree of conservation (Figure 4E). The major difference is that G<sub>18</sub> and  
19 G<sub>19</sub> are at positions 17 and 18 in Arg19 due to the absence of an additional A at position  
20 16 in Ala20 and Asn20. To know whether A<sub>16</sub> residue is also important for translation  
21 inhibition, other oligoribonucleotides were designed. When the A<sub>16</sub> of Ala20 is deleted  
22 (Ala20-16A) or replaced by a U (Ala20-A16U), translation inhibition is strongly affected.  
23 Conversely, when an A is introduced at position 16 in Arg19 sequence (Arg19+A16),  
24 this tRF acquires the ability to inhibit translation as efficiently as Ala20 or Asn20  
25 (Figures 4E and 4F). The introduction of a U rather than an A (Arg19+U16), although  
26 less efficient, still leads to a 50% inhibition. To sum up, among tRF sequences, G<sub>18</sub>  
27 and G<sub>19</sub> appear to be crucial to get the capacity of inhibiting translation; nucleotides  
28 such as A<sub>16</sub> may have a synergistic effect, and the size of the tRF and/or the position  
29 of the nucleotides within the sequence may also be crucial.

30

### 31 **tRF-5D (Ala) associates with polyribosomes**

32 A subset of Arabidopsis s-tRFs appears to efficiently repress translation, raising the  
33 question at which level this regulation is achieved. In organisms such as Archaea [35]  
34 or human cells [19], it has been shown that regulation of translation by short tRFs can

1 be due to their association with ribosomal subunits and we hypothesize a similar  
2 mechanism in plants.

3 To determine if Arabidopsis tRF Ala20 can act similarly, *in vitro* binding studies were  
4 performed. For that purpose, crude lysates from Arabidopsis seedlings, pre-incubated  
5 in the presence of the synthetic oligoribonucleotide Ala20, were fractionated on  
6 sucrose gradient to separate free RNAs, ribosomal subunits and polyribosomes. The  
7 quality of the fractionation is attested by the ribosome profile (Figure 5A) and northern  
8 blots (Figure 5B). An enrichment of 5S rRNA was observed in the polyribosomes,  
9 ribosomes and free ribosomal subunits fractions, while few 5S rRNA is present in the  
10 free RNA fractions. As expected, most of tRNAs (as attested by tRNA<sup>Ala</sup>) is found in  
11 the low-density fractions and in the polyribosomes where active translation occurs.  
12 When polyribosomes are disrupted in the presence of puromycin, an antibiotic causing  
13 premature chain termination of translation, the distribution of 5S rRNA and tRNA<sup>Ala</sup> is  
14 shifted towards the light fractions of the sucrose gradient (right panels of Figure 5A).  
15 The profile of the binding of tRF Ala20 follows the profile of tRNA<sup>Ala</sup> (Figure 5C). As  
16 expected, more than 98% of Ala20 is recovered in the low-density fractions. It  
17 corresponds to the excess of tRF Ala20 added to the Arabidopsis extract before  
18 fractionation on the sucrose gradient. The remaining 2% of Ala20 is mainly found in  
19 the heavy polyribosomal fractions compared to the 80S and ribosomal subunit fractions  
20 (Figures 5B and 5C). By contrast, when a puromycin treatment is performed, Ala20 is  
21 not detected anymore in the polyribosomal fractions. Altogether this indicates that  
22 Arabidopsis tRF Ala20 does not efficiently associate with 80S ribosomes or with free  
23 ribosomal subunits but rather binds active polyribosomes. Similar binding experiments  
24 were performed with Ala16 and Ala33, which were previously shown to not efficiently  
25 inhibit translation *in vitro*. Both tRFs can associate with polyribosomes as Ala20 does  
26 (Supplemental Figure 4). In the same way, the Arg19 that does not affect mRNA  
27 expression *in vitro* has the same binding profile than Ala20 (Supplemental Figure 5).  
28 Thus, the capacity of tRFs to bind polyribosomes *in vitro* cannot explain their differential  
29 translation inhibition observed in wheat germ extract.

30 In Arabidopsis seedlings and leaves, under normal condition of growth, the amount of  
31 tRF Ala20 is very low as compared to the amount of the corresponding tRNA.  
32 Nevertheless, we raised the question of whether endogenous tRFs can be detected in  
33 polyribosomal fractions in a crude lysate from Arabidopsis seedlings under unstressed  
34 conditions. As shown in Figure 5D, although Ala33 is visible in the light fractions, it is

1 not detected in the polyribosomal ones, thus suggesting that *in vivo* this I-tRF is weakly  
2 or not associated with active ribosomes. Two weak signals (called a and b; Figure 5D)  
3 of around 18-19 and 24-26 nt in size, abundant in the free fractions but also observed  
4 all along the gradient, are slightly enriched in the heavy fractions (Figures 5D and 5E).  
5 They represent less than 0.5% of the total tRF Ala of the same size found in the free  
6 fractions. As different tRF-5D (Ala) in the size range of 19-26 nt were previously  
7 identified in sncRNA libraries from Arabidopsis leaves [28], it is presently impossible to  
8 precisely identify which ones are detected in polyribosomes. *In vitro*, tRF-5D Ala20  
9 strongly inhibits translation and can bind polyribosomes; however, *in vivo*, the amount  
10 of s-tRF-5D (Ala) associated with polyribosomes is very low. Thus, taking into account  
11 these two facets of Ala20, we propose that tRFs may rather act as modulators rather  
12 than strong inhibitors of translation.

13

## 14 **Discussion**

15 Among the functions attributed to tRFs, regulation of translation has been observed in  
16 various organisms such as Archaea, protozoan, yeast, and human. In plants, previous  
17 work on sncRNAs found in the phloem sap of pumpkin and on Arabidopsis tRNA  
18 degradants suggested that plant tRFs could also have an inhibitory effect on protein  
19 biosynthesis [24] [34]. Here, our work brings further evidence that indeed, in  
20 *Arabidopsis*, some specific tRFs possess the ability to modulate protein synthesis.  
21 Out of a dozen of tested tRFs, only two tRF-5D, derived from tRNA<sup>Ala</sup> and tRNA<sup>Asn</sup>,  
22 were shown to strongly inhibit translation. These two tRFs are 20 nt long, but we also  
23 showed that a total fraction of Arabidopsis I-tRFs can efficiently inhibit translation *in*  
24 *vitro*. Thus, the population of Arabidopsis tRFs able to negatively affect translation is  
25 likely more widespread. However, considering the high complexity of the Arabidopsis  
26 tRFs population [28], no rule can be established yet to predict which tRFs will affect  
27 the translation. Starting from total s-tRFs or I-tRFs populations of Arabidopsis, a  
28 SELEX approach may be a valuable tool to identify them. Besides, the *in vitro*  
29 synthesized tRFs lack modifications that can influence the affinity of nucleotide-  
30 binding, thus the use of natural post-transcriptionally modified tRFs as those generated  
31 from the Arabidopsis tRNA fraction is likely useful.

32 Intriguingly, while the function of tRFs as a translation inhibitor appears as a rather  
33 widespread process, the modes of action seem to differ depending on the organism  
34 and the type of tRF. Yet, only a few data on the mode of actions have been reported,

they are summarized in Figure 6. In the human alanine I-tRF, the four G residues of the 5' extremity enable the formation of a G-quadruplex structure essential for translation repression by interacting with the translational silencer YB-1 and displacing the eukaryotic initiation factor eIF4G/A from mRNAs, thus inhibiting the binding of the small ribosomal subunit [18] [36]. By contrast, we show here that the four Gs also present in Arabidopsis tRF (Ala) are not essential to affect protein synthesis. Interestingly, the G18 and 19 residues of tRF-5D (Ala) and tRF-5D (Asn) appear necessary. However, they belong to the conserved nucleotides of the D-loop present on mostly all tRNA molecules and are thus not sufficient to explain the specificity of inhibition. Nevertheless, similar observations were obtained in human cells with a global protein synthesis inhibition by s-tRFs, in particular, tRF-Gln19 [19]. Both in Arabidopsis and human cells, these tRFs are not or are weakly associated with AGO proteins and repression of translation is independent of the mRNA sequence, thus suggesting that their mode of action is independent of the silencing pathway. Rather, such tRFs can associate with actively elongating polyribosomes, but not efficiently with ribosomal subunits. This may suggest the existence of a still unknown but conserved process of global translation regulation by tRFs between animals and plants, and future work is required to verify this hypothesis. This is different to what has been described in the archaea *Haloferax volcanii* where s-tRFs inhibit translation by binding the small ribosomal subunit and competing with mRNA association [22] or in *Saccharomyces cerevisiae* where I-tRFs negatively affect protein synthesis by interacting with aminoacyl-tRNA synthetases and the small ribosomal subunit [37]. Finally, in *Trypanosoma brucei*, a I-tRF deriving from a tRNA<sup>Thr</sup> induced under nutrient starvation has an opposite effect, which is to stimulate translation by binding to ribosomes and polysomes [23].

How Arabidopsis tRFs associate with active polyribosomes is presently unknown. Interestingly, by providing the first tRF protein interactome, Keam et al demonstrated recently that the human tRF (Gln)19 which globally represses translation can associate with the Multisynthetase Complex MSC, a complex known to be associated with ribosomes, and paradoxically increase the translation of ribosomal and RNA-binding proteins [20]. Such a MSC complex that was shown to interact with polyribosomes in mammals or archaea also exists in protozoans and yeast [38] and is likely present in Arabidopsis. Deciphering the Arabidopsis tRF interactome will be required to see whether a similar process may be relevant in plants or whether other partners are

1 identified. In human, archaea and yeast, many tRFs are induced upon various stress  
2 conditions, from oxidative stress to nutrient deprivation or pH change. In these  
3 organisms, the regulation of translation by tRFs seems to be linked to the stress  
4 response. Here, in Arabidopsis, a small proportion of tRF (Ala) was detected in active  
5 polyribosomal fractions from young leaves of plants grown under normal conditions.  
6 This low amount of tRFs may represent a way to slightly and regularly modulate the  
7 global translation process under normal conditions and to play an important biological  
8 role by allowing a rapid change in translation efficiency upon demand, i.e. during plant  
9 development or under adverse growth conditions. Indeed, as already shown, the  
10 population of plant tRFs dynamically fluctuates upon various stresses or in different  
11 tissues or organs. For instance tRFs deriving from Arabidopsis Alanine tRNAs are  
12 among the most abundant in leaves [28] and their population may even be much higher  
13 in senescing tissues such as old leaves or senescing seeds where the RNases T2  
14 responsible for tRFs production are upregulated [31]. Unfortunately, polyribosomes  
15 fractionation from such tissues has not been successful yet to allow tRFs analysis.

16

## 17 **Materials and methods**

### 18 ***Plant material and growth conditions***

19 *A. thaliana* is of Columbia 0 (Col-0) ecotype. Eight-week old leaves were harvested  
20 from plants grown on soil at 22°C under a 16h light photoperiod. For *in vitro* grown  
21 seedlings, surface sterilized seeds stratified 2 days at 4°C in the dark were sown on  
22 agar plates containing MSP01 (Caisson Labs UT, USA) supplemented with 3% (wt/vol)  
23 sucrose and grown 8 days at 22°C under a 16h light photoperiod.

24

### 25 ***Crude leaf enzymatic preparation***

26 The protocol used is from [31]. Briefly, 200 mg of leaves powder were resuspended in  
27 1 ml of enzymatic extraction buffer (50 mM Tris-HCl pH 7.5, 10 mM MgCl<sub>2</sub>, 10%  
28 glycerol, 1 mM EDTA, 1 mM β-mercaptoethanol, 50 mM Phenylmethanesulfonyl  
29 fluoride, 1X proteases inhibitor cocktail Complete® (Roche)) and shaked at 4°C for 20  
30 min. After centrifugation at 10 000 x g for 15 min, the supernatant was used for *in vitro*  
31 cleavage assays.

32

33

34

1    **In vitro synthesized tRNA and GFP transcript**

2    Plasmid containing *A. thaliana* cytosolic tRNA<sup>Ala</sup>(UGC) gene sequence was obtained  
3    previously [39]. The tRNA gene was fused to the T7 RNA polymerase promoter at its  
4    5' terminus and included a BstNI site at its 3' terminus. The BstNI linearized constructs  
5    was used as substrate to synthesize an *in vitro* transcript with T7 RNA polymerase  
6    using Ribomax™ transcription kit (Promega, Madison WI) under conditions previously  
7    described in [40]. Capped pGFP transcript was obtained using the same kit in the  
8    presence of a pSu9GFP recombinant plasmid [33]. Reaction was performed in a final  
9    volume of 10 µl following manufacturer's protocol, with slight differences: rGTP was  
10   added to a 0.9 mM final concentration, the Cap analog used was from New England  
11   Biolabs, and 50 µM BSA and 50 µM RNase OUT™ (Invitrogen) were added. Reaction  
12   was run 25 min at 37°C, then 0.75 µl of rGTP (100 mM) was added, and reaction was  
13   pursued during 2h30. To synthesize uncapped GFP transcript, no cap analog was  
14   added and rGTP was supplied to a final concentration of 7.5 mM. Plasmid DNA was  
15   eliminated by a 40 min incubation at 37°C after addition of 80 µl of water, 10 µl of  
16   DNase buffer (400 mM Tris-HCl pH 8.0, 100 mM MgSO<sub>4</sub>, 10 mM CaCl<sub>2</sub>) and 1 µl of  
17   DNase RQ1 (1U/µl, Promega). RNAs were then extracted with phenol/chloroform (1/1)  
18   and ethanol precipitated.

19

20    **In vitro tRNA cleavage assay**

21    Assays [31] were performed at 20°C in a 175 µl reaction volume containing 1 µg of  
22    synthesized tRNA<sup>Ala</sup> transcript, 12.5 µl of leaf enzymatic extract and completed with  
23    water. At time points 0, 5, 10, 15 and 30 min, 30 µl aliquots were rapidly transferred  
24    into a microtube containing 200 µl of water saturated phenol and 170 µl of tRNA  
25    extraction solution (10 mM Tris-HCl pH 7.5, 10 mM MgCl<sub>2</sub>, 1% SDS) and vortexed until  
26    all samples were collected. After centrifugation (2300 x g, 10 min), the aqueous phase  
27    was ethanol precipitated, and RNAs were analyzed by northern blot as described in  
28    [28] using as a probe an oligonucleotide (Supplemental Table S1) complementary to  
29    the 5' extremity of most *Arabidopsis* tRNAs<sup>Ala</sup> (Supplementary Figure 2). As a negative  
30    control, tRNA<sup>Ala</sup> transcript was incubated in the absence of leaf enzymatic extract.

31

32    **Arabidopsis tRNAs and tRFs purification**

33    Total tRNA was prepared from *A. thaliana* Col-0 leaves as described in [41]. This  
34    protocol includes a LiCl precipitation step that allows enrichment, in the supernatant,

1 of RNAs of a size smaller than 150 nt (mainly 5S rRNA, tRNAs and sncRNAs). The  
2 tRNA fraction was further purified on 15% polyacrylamide gel by cutting out the piece  
3 of gel containing tRNAs after staining by methylene blue. tRNAs were eluted from the  
4 gel by a 16-hour incubation at 20°C in the presence of extraction solution (500 mM  
5 ammonium acetate, 10 mM magnesium acetate, 100 mM EDTA, 0.1% SDS). Eluted  
6 tRNAs were submitted to a phenol extraction, followed by ethanol precipitation. The  
7 tRNA pellet was dissolved in water. To produce I-tRFs and s-tRFs, 5 µg of purified  
8 tRNAs were incubated 30 min at 20°C in the presence of 15 µl of leaf enzymatic extract  
9 in a total volume of 90 µl, essentially as described above for *in vitro* tRNA cleavage  
10 assay. After phenol extraction and ethanol purification, the RNA sample was  
11 fractionated on 15% polyacrylamide gel. Purification of I-tRFs and s-tRFs was  
12 performed as described above for total tRNA sample preparation. As a negative  
13 control, tRNAs incubated in the absence of Arabidopsis leaf enzymatic extract were  
14 purified following the same protocol.

15

#### 16 ***In vitro protein synthesis assays***

17 *In vitro* radiolabeled protein synthesis assays were performed using the TnT-T7  
18 Coupled Wheat Germ Extract System kit (Promega) following manufacturer  
19 instructions. Recombinant plasmids expressing pSu9GFP (pGFP) or pSu9DHFR  
20 (pDHFR) constructs under the control of T7 promoter [33] were used and synthetic  
21 oligoribonucleotides were added at different concentrations. After fractionation of the  
22 protein samples on 15% polyacrylamide gels, gels were further stained with  
23 Coomassie blue, dried on a Whatmann paper and analyzed by autoradiography and  
24 Phosphor imaging. For translation assays only, the TnT Wheat Germ Extract kit  
25 (Promega) was used following manufacturer instructions, in the presence of 1.5 µg of  
26 capped or uncapped GFP transcripts and synthetic oligoribonucleotides.

27

#### 28 ***Polyribosomes extraction and RNA analysis***

29 Polyribosomes were extracted from *A. thaliana* seedlings and analyzed as described  
30 previously [42], with few modifications. Crude polysomal extracts were obtained from  
31 200 mg of seedlings powder (in 600 µl), and 300 µl were resolved on 15-60% sucrose  
32 gradient (9 ml) centrifuged for 3h at 178 000 x g (rotor SW41-Ti, Beckman Coulter).  
33 Analysis of polysome profiles was performed with an absorbance detector at 254 nm  
34 and sucrose gradients collected with a BioLogic Duoflow fractions collector (Biorad)

1 into 13 to 16 fractions of 600 µl each. When mentioned, 1.2 µl of a synthetic  
2 oligoribonucleotide (100 µM) was added to lysis buffer. For puromycin treatment, 200  
3 mg of seedlings powder was homogenized in the presence of 600 µl of lysis buffer  
4 (200 mM Tris-HCl pH 9.0, 200 mM KCl, 35 mM MgCl<sub>2</sub>, 1% Brij-35, 1% Triton X-100,  
5 1% NP40, 1% Tween 20). After 10 min of incubation on ice and 10 min centrifugation  
6 at 16 000 x g at 4°C, supernatant (340 µl) was supplied with 85 µl of puromycin solution  
7 (3 mg/ml puromycin, 1.5 M KCl) and 75 µl of 2 M KCl. After 30 min of incubation at  
8 37°C, 27.5 µl of 10% sodium deoxycholate and 22.5 µl of water were added. After  
9 centrifugation (5 min, 16 000 x g, 4°C), 485 µl of supernatant were loaded on a 15-  
10 60% sucrose gradient.

11 RNAs were extracted from sucrose gradient fractions or from crude polysomal extract  
12 by water-saturated phenol/chloroform (1/1), chloroform extracted, and ethanol  
13 precipitated. Then they were analyzed by northern blot as previously described [28].  
14

### 15 **Miscellaneous**

16 The oligonucleotide sequences used in this study are listed in Supplementary Table  
17 S1. The tRFs were chemically synthesized and HPLC purified by the company  
18 Integrated DNA Technologies IDT.  
19

### 20 **Supplementary data**

21 Supplementary Data are available at RNA Biology *online*.  
22

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27

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33 for technical support and Anne-Marie Duchêne for advices.  
34

1   **Disclosure statement**

2   No potential conflict of interest was reported by the authors.

3

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12

13 **Figure legends**

14 **Figure 1. *A. thaliana* l- or s-tRFs can inhibit translation *in vitro*.**

15 (A) Kinetics of cleavage of Arabidopsis *in vitro* synthesized tRNA<sup>Ala</sup>(UGC) transcript  
16 upon incubation in the presence (+) or absence (-) of a crude enzymatic extract from  
17 Arabidopsis leaves (Enz. Ex.). After incubation, RNAs were phenol extracted and  
18 fractionated on 15% polyacrylamide gel, followed by northern blot analysis with a  
19 radiolabeled oligonucleotide specific for the 5' extremity of cytosolic Arabidopsis  
20 tRNA<sup>Ala</sup>. (B) Cleavage of Arabidopsis total tRNA (Tot RNA) upon incubation in the  
21 presence (+) or absence (-) of a crude enzymatic extract from Arabidopsis leaves (Enz.  
22 Ex.). After incubation, RNAs were phenol extracted and fractionated on a 15%  
23 polyacrylamide gel, followed by northern blot analysis with a radiolabeled  
24 oligonucleotide specific for the 5' extremity of cytosolic Arabidopsis tRNA<sup>Ala</sup>. (C)  
25 Production of long (l-tRF) and short (s-tRF) tRFs by *in vitro* cleavage of total  
26 Arabidopsis tRNAs in the presence (+) of a crude enzymatic extract from Arabidopsis  
27 leaves (Enz. Ex.). After incubation, RNAs were phenol extracted, fractionated on 15%  
28 polyacrylamide gel and all l-tRFs and s-tRFs were purified separately from the gel. (D)  
29 Effect of the addition of total Arabidopsis tRNAs or tRFs purified in (B) on the synthesis  
30 of GFP in a wheat germ coupled transcription/translation system in the presence of <sup>35</sup>S  
31 methionine. Radiolabeled GFP was detected by autoradiography or phosphor imaging  
32 after fractionation on 15% acrylamide gel. Two concentrations of tRNAs or tRFs were  
33 used: 4 µM (tRNA-4 and tRF-4) and 8 µM (tRNA-8 and tRF-8). Mock = control

1 experiment without added tRNAs or tRFs. Coomassie blue staining of the gel is shown  
2 as a loading control. (E) Relative quantification of *in vitro* synthesized pGFP in  
3 experiments similar to that described in (D). Error bars show standard deviations  
4 (mean of three biological replicates). One-way ANOVA tests were used to calculate p-  
5 values. Asterisks indicate statistically significant differences between Mock and each  
6 treatment (\*\*P<0.001; \*\*P<0.01; \*P<0.1). A value of 1 has been given to the Mock  
7 sample.

8

9 **Figure 2. A subset of Arabidopsis tRFs can inhibit efficiently translation *in vitro*.**

10 (A) Features of the ten tRFs analyzed in this work. Their sequences are presented in  
11 Supplemental Table 1 and the data derive from [28]. Besides, tRF-5A (Asp) was also  
12 found in AGO1 immunoprecipitates (L. Drouard, personal communication). (B) The  
13 synthetic tRFs presented in (A) were tested for their ability to inhibit the *in vitro* protein  
14 synthesis of pGFP. Experiments were performed essentially as described in Figure  
15 1C. Two concentrations of oligoribonucleotides (0.2 and 1  $\mu$ M) were used.  
16 Quantifications of the results are presented in (C). An arbitrary value of 1 (dotted  
17 vertical line) has been given to *in vitro* synthesized pGFP in the absence of added tRF  
18 (Mock experiment: M). Error bars represent standard errors of the mean of  
19 independent biological experiments (n= 3 to 13). One-way ANOVA tests were used to  
20 calculate p-values. Asterisks indicate statistically significant differences between Mock  
21 and each treatment (\*\*P<0.01; \*P<0.1). Dark grey: 0.2  $\mu$ M, pale grey: 1  $\mu$ M.

22

23 **Figure 3. The tRF-5D (Ala) of 20 nt efficiently inhibits translation *in vitro*.**

24 (A) Sequences of the three tRF-5D of 16, 20, and 33 nt (Ala16, Ala20, and Ala33)  
25 deriving from Arabidopsis tRNA<sup>Ala</sup>(AGC). (B) Effect of the addition of Ala16, Ala20, and  
26 Ala33 on the *in vitro* synthesis of pGFP protein. The experiment was performed as  
27 described in Figure 1C. Increasing concentrations of synthetic tRFs were used: 0.2,  
28 0.4, 1, 2 and 4  $\mu$ M. M = Mock, control experiment without tRF. Coomassie blue staining  
29 of the gel is shown as a loading control. (C) Relative quantification of *in vitro*  
30 synthesized GFP in the experiment presented in (B). Ala16: white triangles, Ala20:  
31 black squares, Ala33: Grey circles. (D) Relative quantification of *in vitro* synthesized  
32 pGFP in experiments similar to that described in (A) and for a tRF concentration of 1  
33  $\mu$ M. Error bars show standard deviations (mean of three biological replicates). A value  
34 of 1 has been given to M. (E) *In vitro* protein synthesis in a wheat germ translation

1 system of synthesized pGFP transcript in the presence of  $^{35}\text{S}$  methionine. Capped or  
2 uncapped transcript was used in the absence or presence of Ala20 or Asn20  
3 oligoribonucleotide at 1  $\mu\text{M}$  concentration. M = Mock, control experiment without tRF.  
4 Coomassie blue staining of the gel is shown as a loading control. (F) Effect of the  
5 addition of Ala16 and Ala20 oligoribonucleotides on the *in vitro* synthesis of DHFR  
6 protein. The experiment was performed as described in Figure 2B with the same  
7 concentration range.

8

9 **Figure 4. Importance of A16, G18 and G19 of tRF-5D (Ala) for the inhibition of**  
10 **translation.**

11 (A) Sequence alignment of Ala20, Ala16, Asn20 with the mutated versions Ala20-5U,  
12 Ala20-3U and Asn20-3U. Us replacing Gs are in bold and underlined (B) Effect of the  
13 addition of the Ala20-5U and Ala20-3U presented in (A) on the *in vitro* synthesis of  
14 pGFP. The experiment was performed as described in Figure 3B with the same  
15 concentration range. (C) Quantification of *in vitro* synthesized pGFP in experiments  
16 similar to that described in (B). Error bars represent standard errors of the mean of  
17 independent biological experiments ( $n= 3$  to 13). One-way ANOVA tests were used to  
18 calculate p-values. Asterisks indicate statistically significant differences between Mock  
19 and each treatment ( $****P<0.0001$ ). A value of 1 has been given to M (Mock, control  
20 experiment without tRF). (D) Effect of the addition of Ala20-3U or Asn20-3U on the *in*  
21 *vitro* synthesis of pGFP. Experiment was performed as in (B). Two concentrations of  
22 oligoribonucleotides (0.2 and 1  $\mu\text{M}$ ) were used. (E) Sequence alignment of Ala20 and  
23 Arg19 with the mutated versions Ala20-A16, Ala20A16U, Arg19+A16, Arg19+U16.  
24 Mutated or added nucleotides are in bold and underlined. The A residue at position 16  
25 in Asn20 and Ala20 is under gray background. *In vitro* synthesis of pGFP in the  
26 presence of these oligoribonucleotides and quantifications of the results are presented  
27 in (F) and (G) respectively. The experiments were performed as in Figure 3B, except  
28 that only two concentrations of oligoribonucleotides (0.2 and 1  $\mu\text{M}$ ) were used. Error  
29 bars represent standard errors of the mean of independent biological experiments ( $n=$   
30 3 to 13). A value of 1 has been given to M (Mock, control experiment without tRF).  
31 One-way ANOVA tests were used to calculate p-values. Asterisks indicate statistically  
32 significant differences between Mock and each treatment ( $****P<0.0001$ ;  $**P<0.01$ ).

33

34 **Figure 5. tRF-5D (Ala) associates with polyribosomes *in vitro* and *in vivo***

1 (A) Polyribosomes profile of *Arabidopsis* seedlings pre-incubated *in vitro* with Ala20  
2 was determined using sucrose gradient sedimentation and OD<sub>254nm</sub> measurement  
3 (blue line) and conductivity (red line). Samples were treated or not with Puromycin.  
4 The positions of the heavy and light polyribosomes, 80S and ribosomal subunits and  
5 free RNAs are shown on the graphs. (B) RNAs extracted from each fraction shown in  
6 (A) were fractionated on 15% polyacrylamide gels and analyzed by northern blots  
7 using probes specific for 5S rRNA, tRNA<sup>Ala</sup> and tRF-5D (Ala). Ethidium bromide (EtBr)  
8 profiles are also shown. (C) Profiles obtained after quantification of the signals from  
9 the northern blots (5S rRNA in blue, tRNA in grey and tRF Ala20 in orange) performed  
10 with samples not treated with puromycin. Due to saturation of the free RNA fractions  
11 with the alanine probe, quantification was done only for fractions 1 to 11. An arbitrary  
12 value of 10 was given to the highest value obtained for each curve. Note that the  
13 intensity of the signals is not comparable from one curve to the other. In (D) a similar  
14 experiment was performed without adding any tRF to determine *in vivo* association of  
15 tRF-5D (Ala) with polyribosomes. The RNA fragments corresponding to the signals a  
16 and b are indicated by arrows. In (E) the profiles obtained after northern blots  
17 quantification with 5S rRNA probes (in blue) are compared to those obtained for a or b  
18 (in orange).

19

20 **Figure 6. Representative examples of modulation of translation by tRFs**

21 (a) Inhibition of translation by human I-tRF (Ala), via a G-quadruplex structure (Gq), by  
22 displacement of eIF4G/A and interaction with YB-1. (b) In human cells and this work  
23 in *Arabidopsis* shows that s-tRFs can associate with active polyribosomes. The two  
24 residues G18 and G19 (GG) are essential. Also, an interaction between the  
25 MultiSynthetases Complex (MSC) and tRFs has been characterized in humans. Note  
26 that in human cells, while global repression of translation has been observed [19], the  
27 protein synthesis of a specific set of mRNAs (coding for ribosomal and RNA-binding  
28 proteins) is, by contrast, stimulated [19]. (c) In the archaea *H. volcanii*, s-tRFs bind to  
29 the small ribosomal subunit and compete with mRNA for ribosome binding. (d) In the  
30 yeast *S. cerevisiae*, I-tRFs directly interact with the small ribosomal subunit and with  
31 aminoacyl-tRNA synthetases associated with ribosomes, thus impairing tRNA  
32 aminoacylation. (e) The tRNA<sup>Thr</sup> 3' half (depicted in green) stimulates protein synthesis  
33 in the protozoan *T. brucei*, by interacting with ribosomes.

**RNA Biology – KRNB-2019-0501R1****Lalande et al, Arabidopsis tRNA-derived fragments as potential modulators of translation****Response to reviewer 2**

The authors have addressed most of my comments and suggestions. The revised manuscript with additional data is now improved. But I do still have unresolved concerns listed below.

**Major points**

1. In Supp. Fig. 3, the oligonucleotide Ala16 mentioned in the legend is missing. Amounts of each RNA used in the blotting should be provided in the figure legend. In the blot for total RNA, the signal for tRF20 appears to be much weaker than that of tRF20. The quantification data below the blot however do not seem to translate the signals in the blot.

The reviewer is right. In the Supp Fig.3, no Ala16 was loaded on the blot. "Ala 16" has been deleted from the legend. The amounts of each RNA used in the experiment are now provided in the figure legend. I imagine that the reviewer is comparing the signals obtained for Ala33 and Ala20 in the total RNA sample. We agree with the reviewer that the Ala20 signal is weaker than the signal observed for Ala33. However, when quantification has been performed in the linear part, the contrary was obtained. This is likely because the probe is better hybridizing to Ala33 than to Ala20 (e.g. when comparing hybridization signals obtained with 50 fmol of oligonucleotides, the signal is much stronger for Ala33 than for Ala20). In our opinion, this experiment only gives a rough estimation of tRFs in a total RNA sample.

**Minor points**

1. In response to the minor point #3, the authors mentioned that "This is now mentioned in the --- , page 13". This critical description is missing in the MM section.

The information is now added in the paragraph "Miscellaneous" in the MM section, page 14

2. Consistency in writing concentration and volume terms etc. needs to be maintained (e.g., page 13 and many places in the MM section, 2M vs. 200 mM, ul vs uL, rpm vs. x g).

Writing of concentrations, volumes and centrifugation speeds have been checked for homogeneity.

3. Style in the reference is still not consistent. Too many errors!

Style errors have been corrected.

4. Consistency in P values in all figure legends: e.g. \*\*\* P<0.001 vs. \*\*\*\* P<0.001

Consistency in P values has been given.

Figure 1

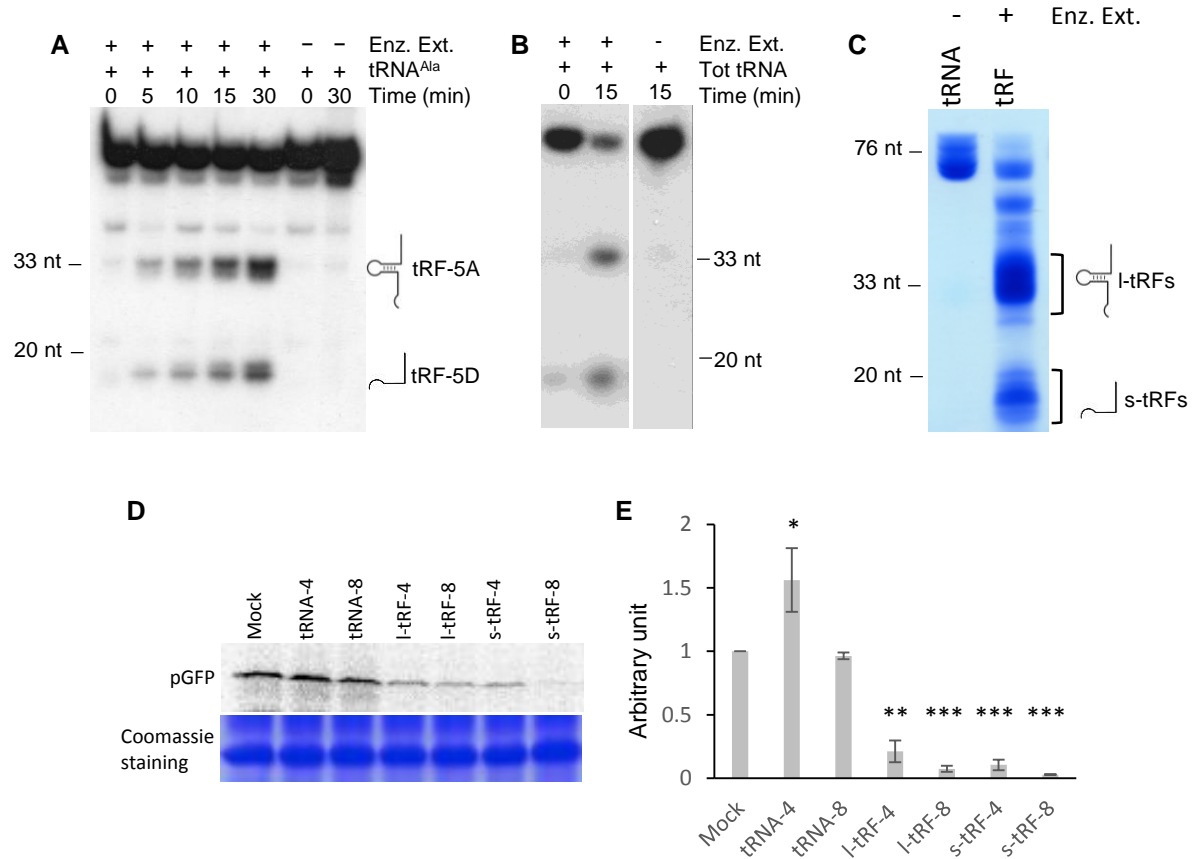


Figure 2

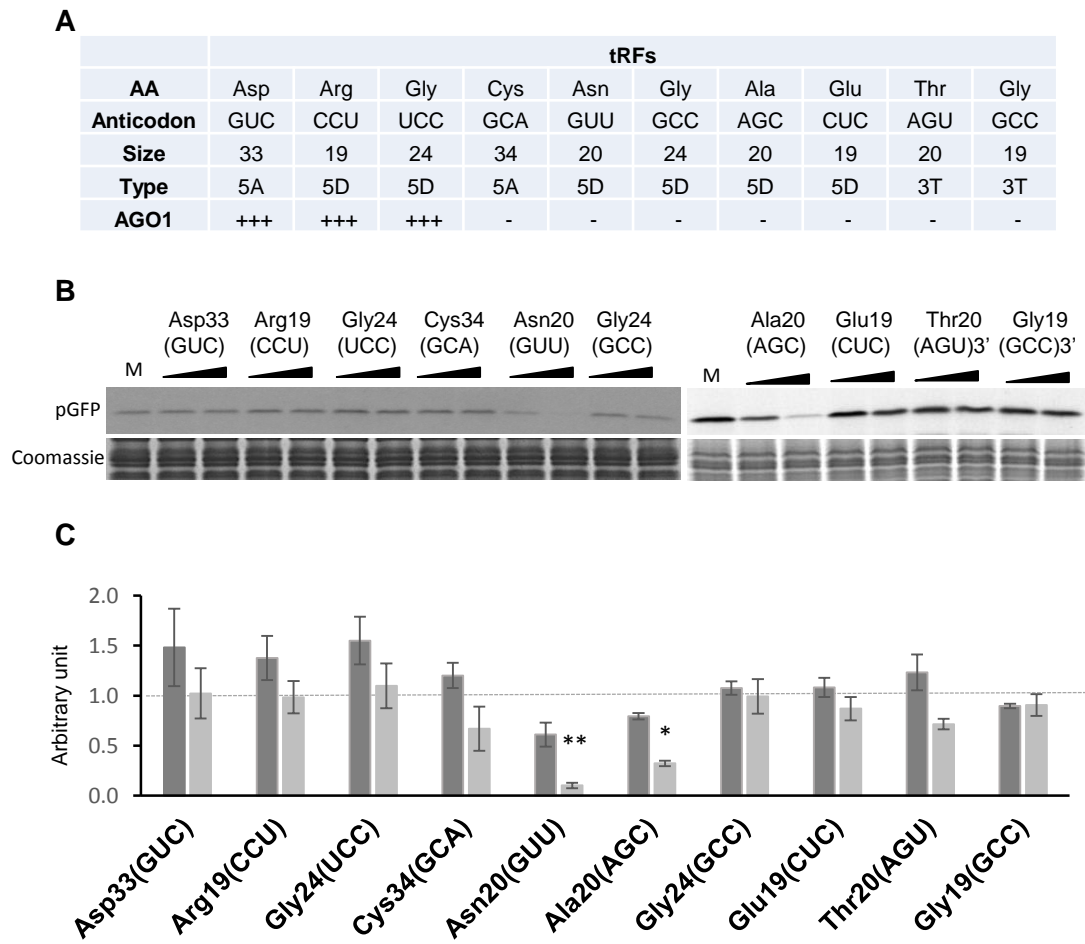


Figure 3

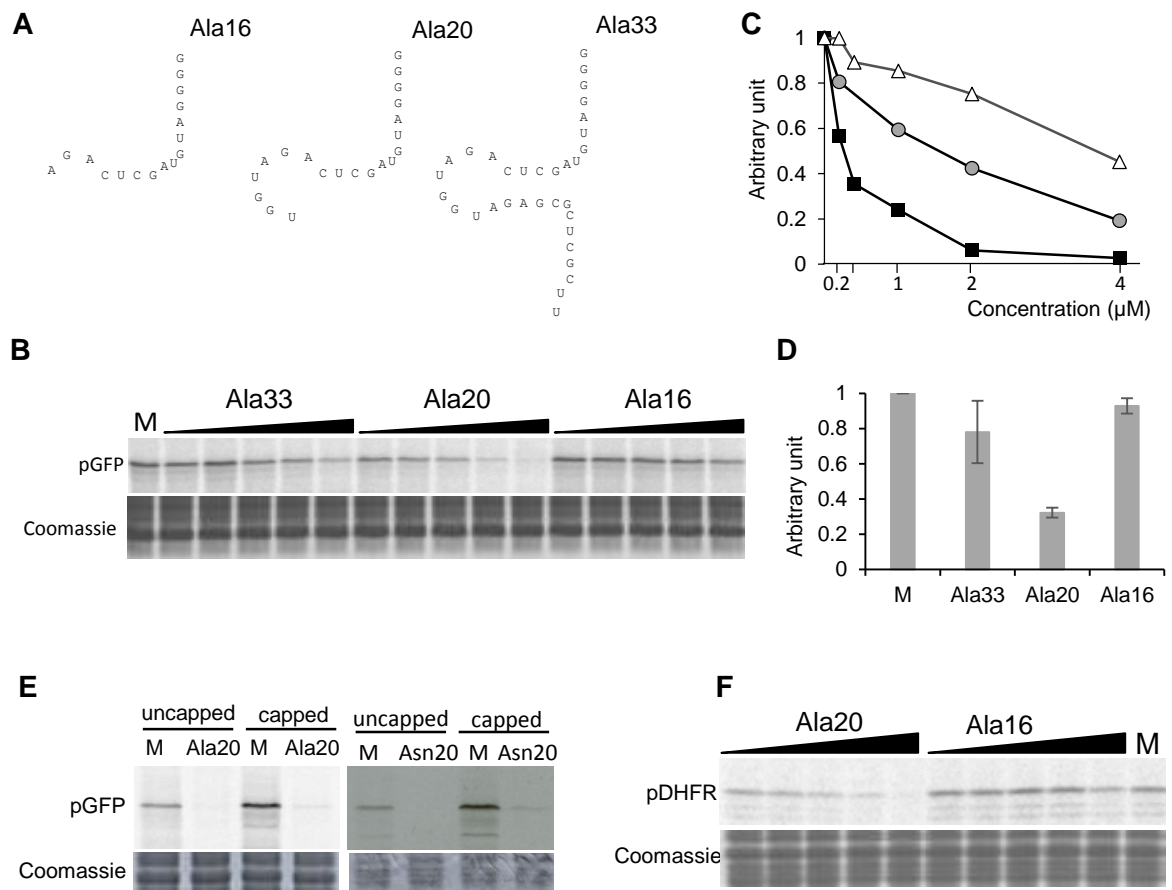


Figure 4

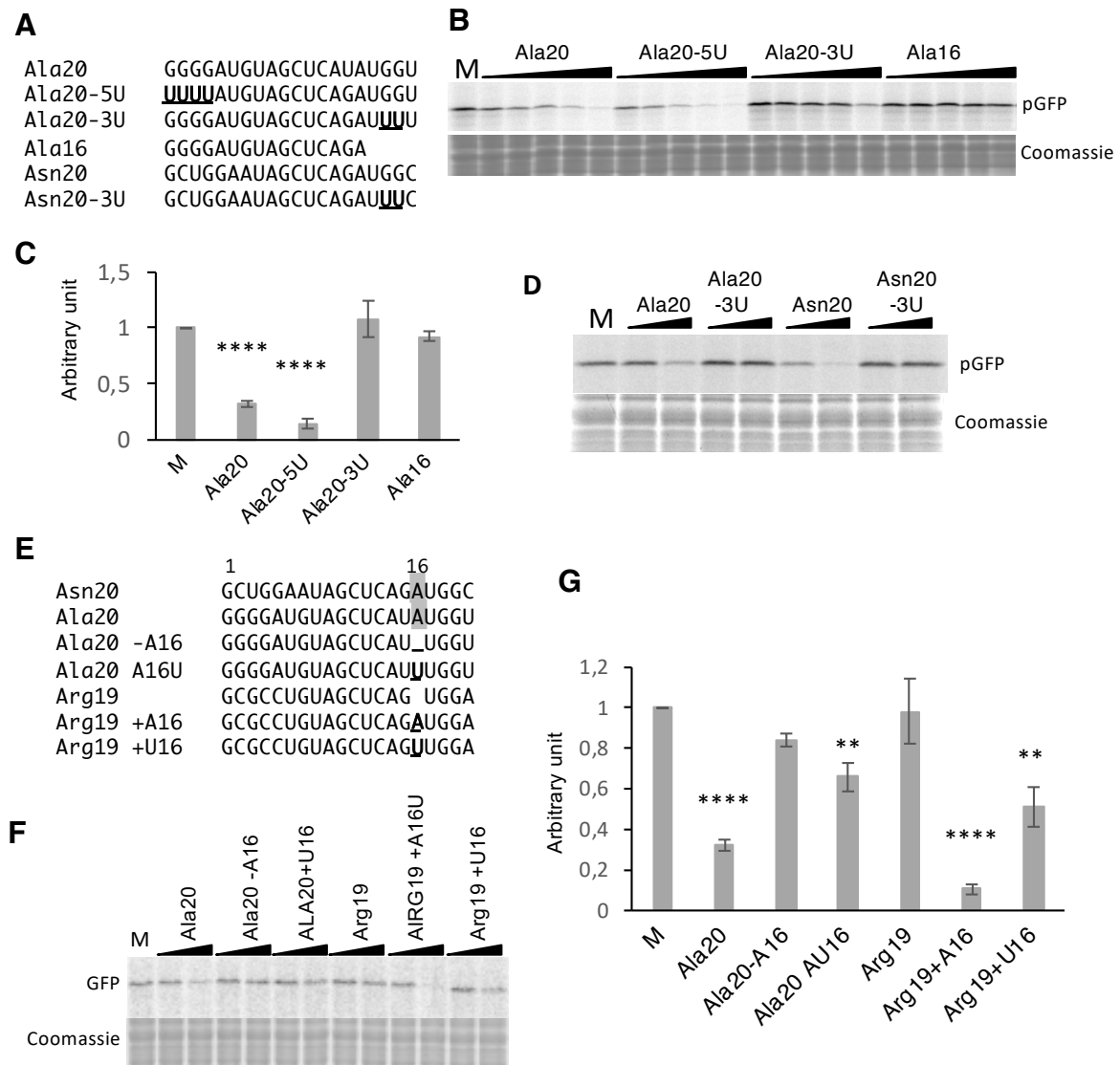


Figure 5

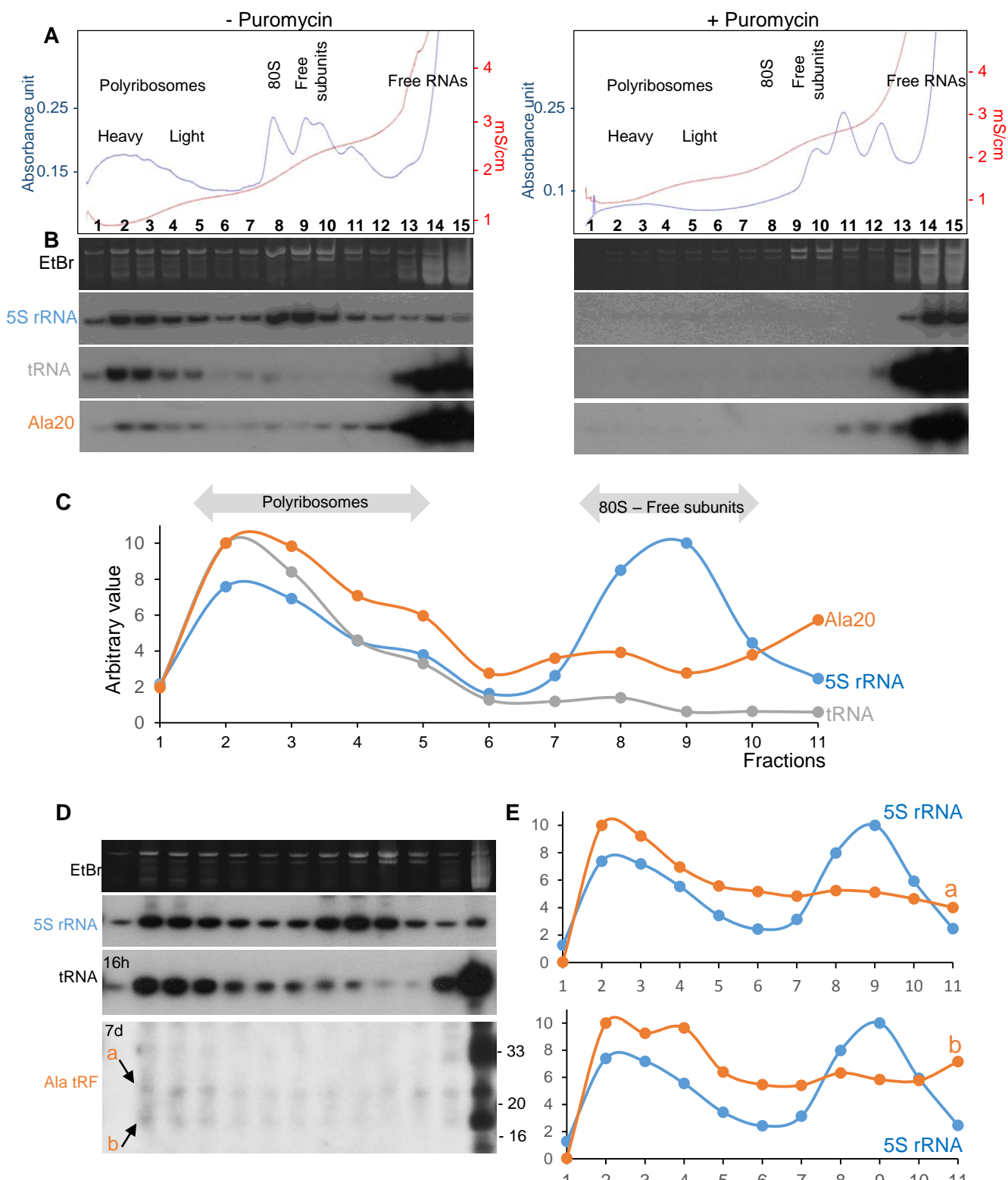
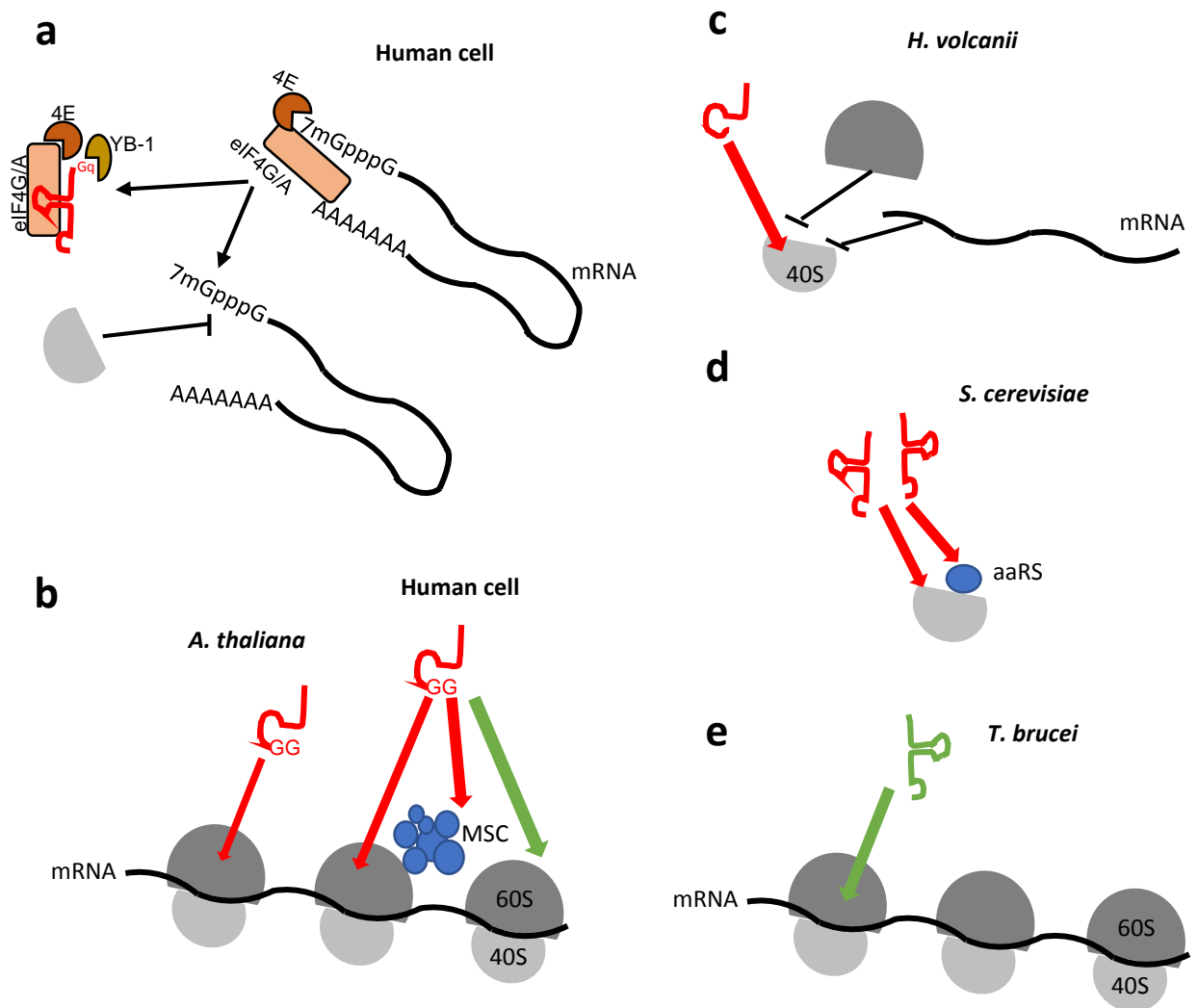
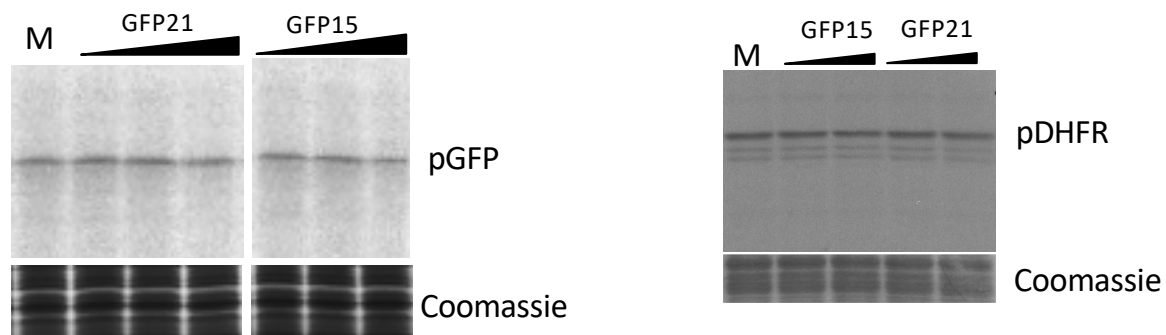


Figure 6



## Supplemental Figure 1



**Supplemental Figure 1:** Addition of GFP15 or GFP21 oligoribonucleotides have no significant effect on the *in vitro* synthesis of pGFP or pDHFR proteins. Experiments were performed as described in Figure 2B. Concentration range: 0.2, 0.4 and 1  $\mu$ M for pGFP; 0.2 and 1  $\mu$ M for pDHFR. Coomassie blue stainings of the gels are shown as loading controls. M: Mock experiment.

## Supplemental Figure 2

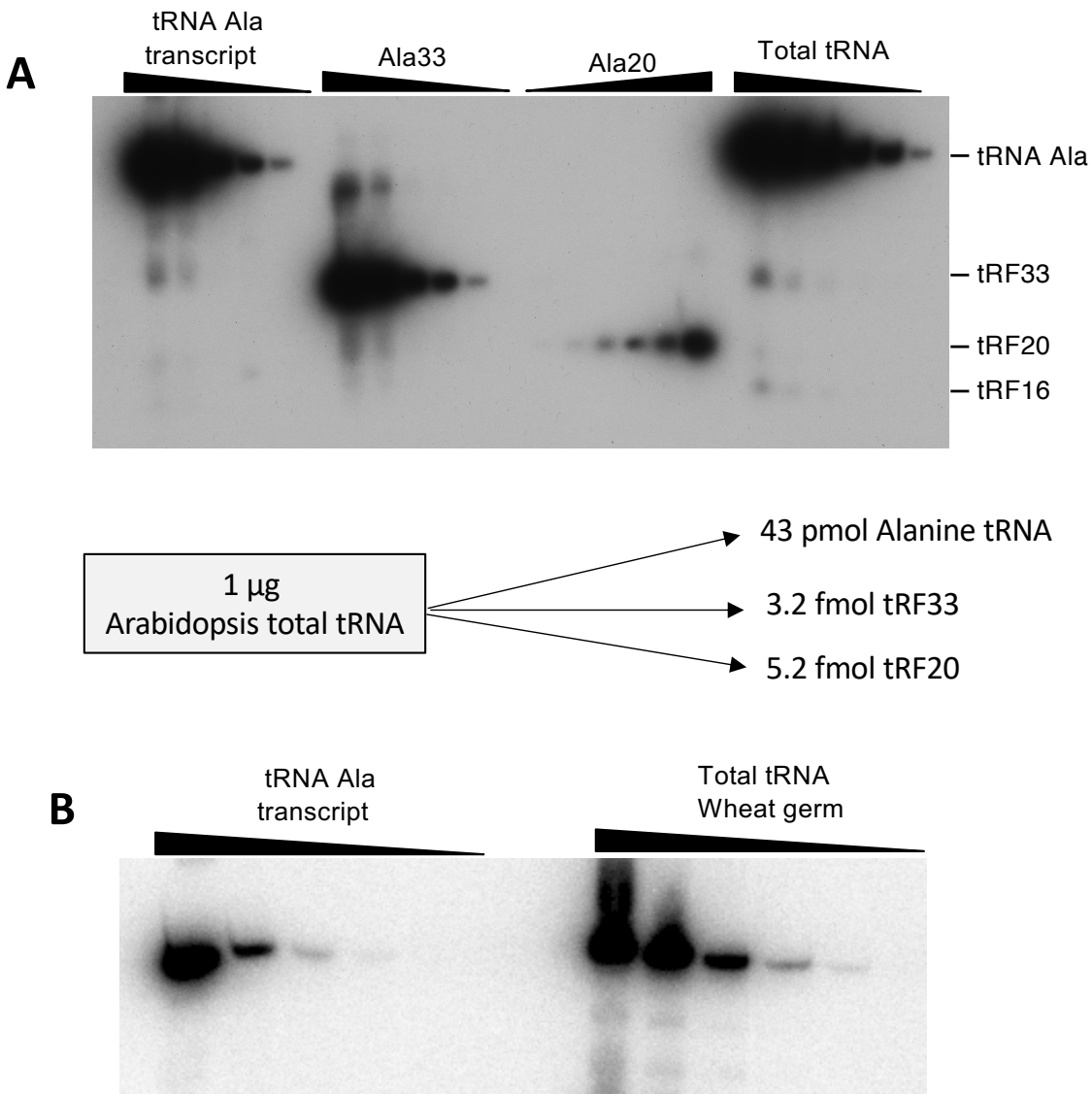
A	tRFs20	tRNAAla sequences	copies number
	tRF-5D-Ala1	<u>GGGGATGTAGCTCAGATGGT</u> AGAGCGCTCGCT <b>A</b> GCATGCGAGAGGCACGGGATCGATAACCCGCATCTCCACCA	12
	tRF-5D-Ala1	<u>GGGGATGTAGCTCAGATGGT</u> AGAGCGCTCGCT <b>A</b> GCATGCGAGAGG <b>T</b> ACGGGATCGATAACCCGCATCTCCACCA	3
	tRF-5D-Ala2	<u>GGGGATGTAGCTCA</u> <b>A</b> ATGGTAGAGCGCTCGCT <b>A</b> GCATGCGAGAGG <b>T</b> ACGGGATCGATAACCCGCATCTCCACCA	1
	tRF-5D-Ala2	<u>GGGGATGTAGCTCA</u> <b>A</b> ATGGTAGAGCGCTCGCT <b>T</b> <b>G</b> CATGCGAGAGGCACGGG <b>T</b> TCGATCCCCGCATCTCCACCA	9
	tRF-5D-Ala3	<u>GGGGATGTAGCTCA</u> <b>T</b> ATGGTAGAGCGCTCGCT <b>T</b> <b>G</b> CATGCGAGAGG <b>C</b> ACAGGG <b>T</b> TCGAT <b>T</b> CCCTGCATCTCCACCA	1
	tRF-5D-Ala3	<u>GGGGATGTAGCTCA</u> <b>T</b> ATGGTAGAGCGCTCGCT <b>C</b> <b>G</b> CATGCGAGAGG <b>C</b> ACAGGG <b>T</b> TCGAT <b>T</b> CCCGCATCTCCACCA	1
	tRF-5D-Ala4	<u>GGGGGTGTAGCTCA</u> <b>T</b> ATGGTAGAGCGCTCGCT <b>C</b> <b>G</b> CATGCGAGAGG <b>C</b> ACAGGG <b>T</b> TCGAT <b>T</b> CCCGCACCTCCACCA	5
	tRF-5D-Ala4	<u>GGGGGTGTAGCTCA</u> <b>T</b> ATGGTAGAGCGCTCG <b>T</b> <b>T</b> <b>C</b> <b>G</b> CATGCGAGAGG <b>C</b> ACAGGG <b>T</b> TCGAT <b>T</b> CCCGCACCTCCACCA	1

B	tRFs20	tRNAAsn sequences	copies number
	tRF-5D-Asn1	<u>GCTGGAATAGCTCAG</u> <b>T</b> GG <b>T</b> .AGAGCGTGTGG <b>C</b> <b>T</b> <b>T</b> TAACCACAAGGT <b>C</b> GGAGGTT <b>C</b> GAACCCCT <b>C</b> <b>T</b> CTAGCGCCA	6
	tRF-5D-Asn1	<u>GCTGGAATAGCTCAG</u> <b>T</b> GG <b>T</b> .AGAGCGTGTGG <b>C</b> <b>T</b> <b>T</b> TAACCACAAGGT <b>C</b> GGAGGTT <b>C</b> GA <b>T</b> CCCT <b>C</b> <b>T</b> CTAGCGCCA	1
	tRF-5D-Asn2	<u>GCTGGAATAGCTCAG</u> <b>C</b> GG <b>T</b> .AGAGCGTGTGG <b>C</b> <b>T</b> <b>T</b> TAACCACAAGGT <b>C</b> GGAGGTT <b>C</b> GAACCCCT <b>C</b> <b>T</b> CTAGCGCCA	1
	tRF-5D-Asn3	<u>GCTGGAG</u> TAGCTCAG <b>T</b> GG <b>T</b> .AGAGCGTGTGG <b>C</b> <b>T</b> <b>T</b> TAACCACAAGGT <b>C</b> AGAGGTT <b>C</b> GAACCCCT <b>C</b> <b>T</b> CTAGCGCCA	4
	tRF-5D-Asn3	<u>GCTGGAG</u> TAGCTCAG <b>T</b> GG <b>T</b> .AGAGCGTGTGG <b>C</b> <b>T</b> <b>T</b> TAACCACAAGGT <b>C</b> AGAGGTT <b>C</b> GAACCC <b>T</b> T <b>C</b> TCTAGCGCCA	1
	tRF-5D-Asn4	<u>GCTGGAATAGCTCAG</u> <b>T</b> AG <b>G</b> <b>T</b> .AGAGCGTGTGG <b>C</b> <b>T</b> <b>T</b> TAACC <b>T</b> CAAGGT <b>C</b> GGAGGTT <b>C</b> GAACCCCT <b>C</b> <b>T</b> TCAGCGCCA	1
	tRF-5D-Asn5	<u>GCTGGAATAGCTCAG</u> <b>A</b> GG <b>C</b> <b>T</b> .AGAGCGTGTGG <b>C</b> <b>T</b> <b>T</b> TAACCACAAGGT <b>C</b> GGAGGTT <b>C</b> GAACCCCT <b>C</b> <b>T</b> CTAGCGCCA	1

**Supplemental Figure 2:** Alignment of Arabidopsis tRNAAla and tRNAAsn unique sequences. The number of copies of each sequence is indicated. The tRFs of 20 nt (tRFs20) generated from these tRNAs and found in the small non coding RNA libraries are underlined. Anticodons are in bold. Nucleotide differences compared to the sequence of reference (i.e. the sequence in blue of the Ala20 and Asn20, we synthesized) are in red.

## Supplemental Figure 3

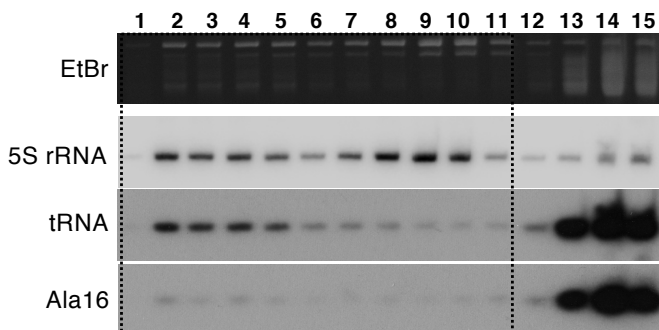


**Supplemental Figure 3:** (A) Exemple of quantification of tRFAla 20 and 33 and Alanine tRNA in an Arabidopsis total tRNA extract. Known amount of synthetic oligoribonucleotides Ala20 (10, 25, 50, 75, 100 and 250 fmol) and Ala33 (10, 50, 100, 500 and 1000 fmol) and of alanine tRNA transcript (1, 5, 10, 50 and 100 pmol) were used to obtain standard ranges and to quantify tRFs Ala and tRNA alanine in an arabidopsis total RNA fraction (10, 50, 100, 500 and 1000 ng). One example of the blot is presented here but several times of exposure were required to use non-saturated signals for the same blot. The scheme summarizes the quantification. (B) Example of northern blot experiment allowing the quantification of the Alanine tRNA in the TnT wheat germ extract using alanine tRNA transcript as a standard.

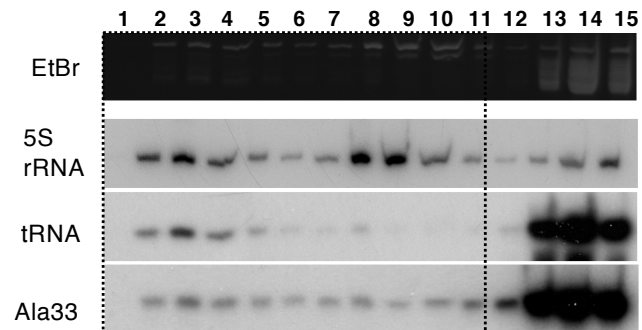
## Supplemental Figure 4

**A**

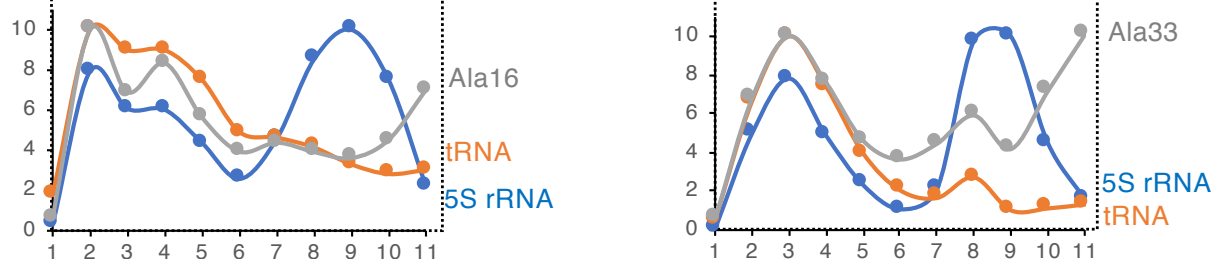
**+ Ala16**



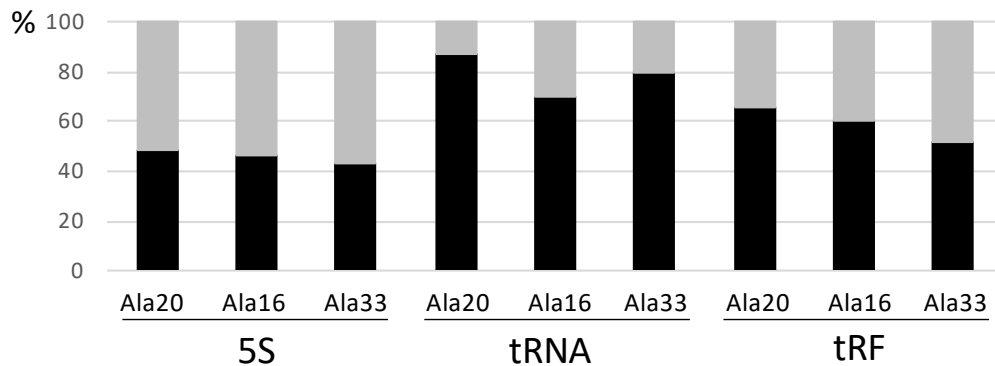
**+ Ala33**



**B**

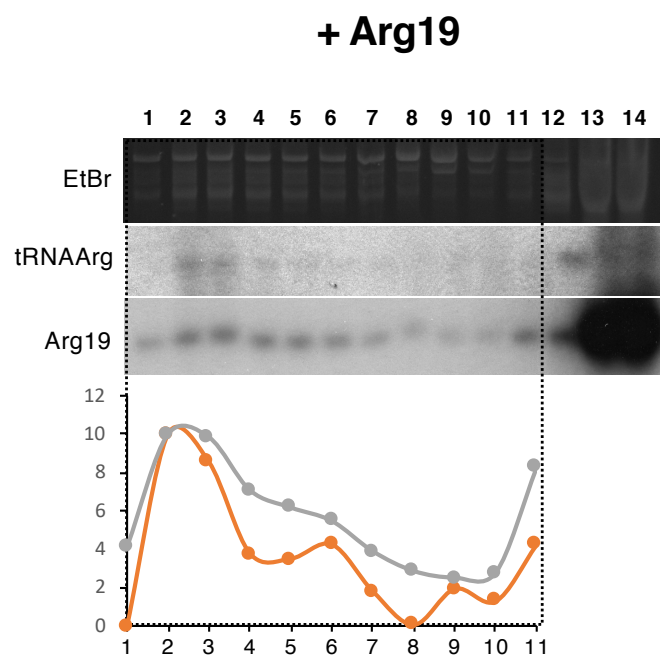


**C**



**Supplemental Figure 4:** Binding experiments performed with Ala16 and Ala33 on *Arabidopsis* seedlings, as described for Ala20 in Figure 5A-C. **A)** After polysomal fractionation on a sucrose gradient, RNAs were extracted from each fraction and analyzed by northern blots using probes specific for 5S rRNA, tRNA<sup>Ala</sup> and tRF-5D (Ala). Ethidium profiles (EtBr) are also shown. **B)** Curves obtained after quantification of the signals from the northern blots (5S rRNA in blue, tRNA in orange and tRF Ala16 or Ala33 in grey). Due to saturation of the free RNA fractions with the alanine probe, quantification was done only for fractions 1 to 11. An arbitrary value of 10 was given to the highest value obtained for each curve. Note that the intensity of the signals is not comparable from one curve to the other. **C)** Global quantification of the signals obtained in polyribosomal fractions (fractions 2 to 5; black) and in 80S-free ribosomal subunit fractions (fractions 8 to 10; grey) from the experiments shown in Figure 5B (Ala20) and in Supplemental Figure 3A (Ala16 and Ala33).

## Supplemental Figure 5



**Supplemental Figure 5:** Binding experiments performed with Arg19 on Arabidopsis seedlings, as described for Ala20 in Figure 5A-C. After polysomal fractionation on a sucrose gradient, RNAs were extracted from each fraction and analyzed by northern blots using a probe specific for, tRNA<sup>Arg(CCU)</sup> and tRF-5D (Arg). Ethidium profile (EtBr) profile is also shown. The curves obtained after quantification of the signals from the northern blots are presented (tRNA in grey and tRF in orange).

**Table S1: Oligodeoxynucleotide and oligoribonucleotide sequences used in this work**

**Oligodeoxynucleotides used for Northern blots**

<b>Names</b>	<b>Sequences (5' to 3')</b>
Ala	ACCATCTGAGCTACATCCCC
5S rRNA	GCATTAGTGCTGGTATGATCGCATCC
Arg	ACCACTGAGCTACAGGCGC

**Oligoribonucleotides used for translation inhibition assays**

<b>Names</b>	<b>Sequences (5' to 3')</b>	<b>Description</b>	<b>Size</b>
Ala16	GGGGAUGUAGCUCAGA	tRF-5D Ala(AGC)	16 nt
Ala20	GGGGAUGUAGCUCAGAUUGGU	tRF-5D Ala(AGC)	20 nt
Ala33	GGGGAUGUAGCUCAGAUUGGUAGAGCGCUCGUU	tRF-5D Ala(AGC)	33 nt
Ala20-5U	UUUAUGUAGCUCAGAUUGGU	G to U mutations at the 5' extremity of tRF-5D Ala(AGC)	20 nt
Ala20-3U	GGGGAUGUAGCUCAGAUUUU	G to U mutations at the 3' extremity of tRF-5D Ala(AGC)	20 nt
Ala20 A16U	GGGGAUGUAGCUCAUUUGGU	A to U switch at position 16 of tRF-5D Ala(AGC)	20 nt
Ala20 -A16	GGGGAUGUAGCUAUUGGU	Deletion of A16 of tRF-5D Ala(AGC)	19 nt
Arg19	GCGCCUGUAGCUCAGUGGA	tRF-5D Arg(CCU)	19 nt
Arg19 +A16	GCGCCUGUAGCUCAGAUGGA	Addition of A at position 16 of tRF-5D Arg(CCU)	20 nt
Arg19 +U16	GCGCCUGUAGCUCAGUUGGA	Addition of U at position 16 of tRF-5D Arg(CCU)	20 nt
Asn20	GCUGGAAUAGCUCAGAUGGC	tRF-5D Asn(GUU)	20 nt
Asn20-3U	GCUGGAAUAGCUCAGAUUUC	G to U mutations at the 3' extremity of tRF-5D Asn(GUU)	20 nt
Gly(UCC)24	GCGUCUGUAGCUAACGGGUAGGA	tRF-5D Gly(UCC)	24 nt
Gly(GCC)24	GCACCAUGGUCUAGUGGUAGAAU	tRF-5D Gly(GCC)	24 nt
Glu(CUC)19	UCCGUCGUAGCUAGCUGG	tRF-5D Glu(CUC)	19 nt
Asp(GUC)33	GUCGUUGUAGUAUAGUGGUAGUAUUCGGCCU	tRF-5D Asp(GUC)	33 nt
Cys(GCA)34	GGGCCAUAGCUCAGUGGUAGAGCAUUCGACUGC	tRF-5D Cys(GCA)	34 nt
Gly(GCC)19-3'	AUUCCCGGCUGGUGCACCA	tRF-3T Gly(GCC)	19 nt
Thr(AGU)20-3'	AACUCUCAACGAAAGCACCA	tRF-3T Trh(AGU)	20 nt
GFP21	GUAAACGGCCACAAGUUCAGC	random GFP sequence	21 nt
GFP15	GUAAACGGCCACAAG	random GFP sequence	15 nt