Nucleotide sequence of a spinach chloroplast proline tRNA

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ABSTRACT

The nucleotide sequence of a spinach chloroplast proline tRNA (sp. chl. tRNA ^{Pro}) has been determined. This tRNA shows more overall homology to phage T4 proline tRNA (61% homology) than to eukaryotic proline tRNAs (53% homology) or mitochondrial proline tRNAs (36-49% homology). Sp. chl. tRNA ^{Pro}, like all other chloroplast tRNAs sequenced, contains a methylated GG sequence in the dihyrouridine loop and lacks unusual structural features which have been found in many mitochondrial tRNAs.

INTRODUCTION

Mitochondria and chloroplasts contain their own DNA and protein synthesizing systems. Protein synthesis in mitochondria shows significant differences from that occurring in prokaryotes and eukaryotic cytoplasm. For example specific changes in the genetic code have been observed (1-3) and a variety of unusual tRNA and tRNA genes studied (2-4). Protein synthesis in chloroplasts is not as well characterized as in mitochondria. As part of a study of protein synthesis in chloroplasts we have determined the nucleotide sequence of a spinach chloroplast proline tRNA.

MATERIALS AND METHODS

The isolation of chloroplasts from <u>Spinacia oleracea</u> L. var 424, the isolation of total spinach chloroplast tRNA and the methods used in sequence determination and hybridization have been described previously (6-8)

RESULTS

<u>Purification of tRNA</u> The pure tRNA^{PTO} used for sequence analysis and hybridization studies was obtained by a three step chromatographic procedure consisting first of a benzoylated DEAE cellulose column rum at pH 7.6. Fractions 209-231 of this column (see Fig. 1 of reference 9) were pooled and further purified on an RPC-5 column at pH 4.5. Finally, tRNA^{PTO} was purified on an RPC-5 column at pH 7.5 to homogeneity as judged by a

single band on a denaturing 20% polyacrylamide gel, stained with ethidium bromide (data not shown). This tRNA was shown by subsequent sequence analysis and aminoacylation (with <u>E. coli</u> synthetase) to be a proline tRNA.

Sequence Analysis The nucleotide sequence of sp. chl. tRNA pro was determined using formamide fragment analysis, RNA sequence gels and mobility shifts. The formamide fragment analysis method (10,11) provided the most sequence information of the three methods employed and allowed an unambiguous assignment of residues 3 to 74 with the exception of residue G₁₀ for which there was no corresponding band in the formamide generated "ladder". Missing bands, compression effects and other anomalies have occasionally been observed in the formamide fragment procedure (6,7,11) and underscore the need for confirmatory sequence procedures. Residue 10 was identified as G on the basis of an RNase Tl band on several RNA sequence gels. However, since RNA sequence gels cannot distinguish between a G and a modified G, and since this residue was not detected on the formamide fragment analysis method, we conclude that residue 10 is either a G or a modified G.

The dinucleotide GmG in the dihydrouridine loop and all other modified nucleotides obtained using formamide fragment analysis were identified as described previously (7). RNA sequence gels (6,12-14) were used to analyze both $5'-^{32}P$ and $3'-^{32}P$ labeled sp. chl. tRNA pro. These gels confirmed the assignment of residues 1-75. The 3' end of the tRNA was also determined by mobility shift procedures, performed as described (15).

Hybridization of Spinach Chloroplast tRNA^{PTO} The sp. chl. [3'-³²P] tRNA^{PTO} was hybridized to restriction fragments of sp. chl. DNA which were generated by the enzymes SalI and KpnI respectively using methods previously described (7,8,16-18). The sp. chl. [3'-³²P]tRNA^{PTO} hybridized to a 13.8 megadalton SalI fragment and to a 13.8 megadalton KpnI fragment, on a part of the sp. chl. genome where other tRNAs are known to hybridize (19,20).

DISCUSSION

The nucleotide sequence of sp. chl. tRNA^{PTO} is shown in Fig. 1. This tRNA contains 77 residues, the same size as phage T4 tRNA^{PTO} (21), both of which are larger than the eukaryotic proline tRNAs from mouse and chicken (21) (75 residues) and mitochondrial proline tRNAs from yeast (22) (75 residues) and human cells (3) (71 residues). In addition sp. chl. tRNA^{PTO} shows more overall homology to phage T4 tRNA^{PTO} (61% homology) than to eukaryotic proline tRNAs (53% homology) or mitochondrial tRNA^{PTO} (36-49% homology). Unfortunately no prokaryotic tRNA^{PTO} has yet been sequenced.

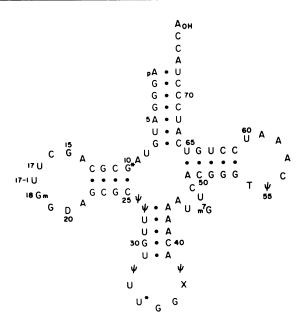


Figure 1. Nucleotide sequence of spinach chloroplast proline tRNA Residue G_{10}^{\star} may be a modified G.

Sp. chl. tRNA^{PTO} contains a methylated GG sequence in the dihydrouridine loop, a feature found in all chloroplast tRNAs sequenced to date. Sp. chl. tRNA^{PTO} contains an unknown modified uridine, denoted as U*, in the wobble position of the anticodon (residue 34). This same residue is also found in sp. chl. tRNA^{Val} and its characteristics have been discussed (23). Sp. chl. tRNA^{PTO} also contains an unknown modified residue at position 37 denoted as X. Residue X appears to be similar but not identical to t⁶A. Residue X migrates as pt⁶A in the two-dimensional system of Silberklang, et al. (15) but migrates slightly slower than pt⁶Ap in the ammonium formate system of Gupta and Randerath (11). Authentic t⁶A was obtained from bovine liver tRNA^M₁

A comparison of the sequences of the six known proline tRNAs suggests that certain residues (numbered as in Fig. 1) may be of interest as follows: A) G_3 is present in the three organelle tRNA^{PTO} while C_3 is present in the three non-organelle tRNA^{PTO}; B) G_{10} followed by a pyrimidine at residue 11 is present in all tRNA^{PTO}; C) G_{15} is present in all except mitochondrial tRNA^{PTO}; D) all tRNA^{PTO} have pyrimidines at residues 27 and 28, a U or

modified U at residue 32 and purines at residues 42 and 43; E) non-eukaryotic tRNA pro have A, while eukaryotic tRNA pro have C, These observations are based upon only six proline tRNAs and sequence analysis of additional proline tRNA will be of interest.

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