Domains of initiator tRNA and initiation codon crucial for initiator tRNA selection by *Escherichia coli* IF3

Dieter Hartz, Jonathan Binkley, Tamy Hollingsworth, and Larry Gold

Department of Molecular, Cellular, and Developmental Biology, University of Colorado, Boulder, Colorado 80309 USA

Initiation factors are used by *Escherichia coli* to select the initiator tRNA over elongator tRNAs during translation initiation. IF3 appears to 'inspect' the anticodon end of the tRNA, probably along with the initiation codon. The anticodon stem and loop of the initiator tRNA, together with part of the initiation codon of the mRNA, can be thought of as a unit. Changes made in the anticodon stem, the anticodon loop, or the anticodon of an initiator tRNA fragment result in a loss of selection by IF3 in an in vitro assay for translation initiation. IF3 allows the selection of an initiator tRNA anticodon stem and loop fragment on GUG and UUG codons but does not select that tRNA fragment in response to AUU.

[Key Words: Translation; initiation; IF3; tRNA]

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Results

To analyze 30S complex formation on various translation initiation sites, we used the extension inhibition technique, also called toeprinting. cDNA synthesis on a template mRNA is terminated when the reverse transcriptase encounters a 30S ribosomal subunit plus tRNA bound on the mRNA. The short cDNA is visualized as a toeprint band on a sequencing gel. The toeprint usually appears 15 bases downstream of the A of the initiation codon if initiator tRNA is bound in the complex or at +15 from the first base of the cognate codon if elongator tRNA is bound. When compared to the band of the full-length cDNA, the toeprint band is a quantitative measure of ternary complex formation [Hartz et al. 1989].

Variations of A22Met

Using the toeprinting assay, we have shown previously that the synthetic initiator tRNA fragment A22Met [Fig. 1] behaves very similarly to the whole tRNAMet in 30S initiation complex formation; A22Met is able to bind to the ribosomal P site in ternary complexes with 30S subunits and T4 gene 32 mRNA [the pr5170 transcript] at the true translation initiation site. A22Met is also selected over the elongators tRNA_Cys and tRNA_Phe by IF3 [Hartz et al. 1989]. A22Met binds to 30S complexes with a lower affinity than the whole initiator tRNA.

To elucidate further the parts of the initiator tRNA that contribute to ternary complex formation and IF3 selection, we synthesized variants of A22Met [Fig. 1]. The fragments DA39Met and AT42Met contain an additional, unmodified D and TΨC loop, respectively, whereas...
Figure 1. Synthetic tRNA fragments. Note the triphosphates at the 5' ends and the extra random nucleotide (N) on the 3' ends. Some fragments (with N in parentheses) were also isolated without the 3' random nucleotide.

A13\textsuperscript{Met}, as the shortest fragment, contains only the three GC base pairs of the anticodon stem. We further synthesized "flip-stem," with the base pairs of the anticodon stem switched, and "flop-loop," with the two 5' bases of the anticodon loop switched with the two 3' bases of the loop. Finally, we synthesized anticodon variants with complementarity to UGU/UUC ("sphinx-cys"), UUU/UUC ("sphinx-phe"), and AUU ("sphinx-ile") codons. The fragments were tested for their ability to form ternary complexes with 30S subunits and the pRS170 [gene 32] in vitro transcript in the toeprinting assay. All fragments, except flop-loop, yield strong toeprints 15 bases downstream from their respective codons (Fig. 2). Sphinx-cys and sphinx-phe recognize the UGU and UUU codons, respectively, as indicated by a corresponding shift of the toeprinting position. Sphinx-ile has no cognate codon on this mRNA (see below).

Binding affinities

The fragments of Figure 1 were titrated for their ability to form ternary complexes on the pRS170 transcript. At a constant 30S subunit concentration, increasing amounts of tRNA fragments were added to toeprinting reactions (data not shown). The relative toeprints were determined by radioscanning of the gels [Fig. 3]. The relative toeprints do not reach 100% at apparently saturating concentrations of tRNA fragments because the reactions have not gone to completion during the 10-min preincubation period. Reaction completion, at saturating concentrations of tRNA\textsuperscript{Met} and A13\textsuperscript{Met}, takes ~100 min; at this time, relative toeprints of 100% are obtained with both tRNA\textsuperscript{Met} and A13\textsuperscript{Met} [S. Ringquist, pers. comm.]. Thus, the equilibrium is strongly toward ternary complex formation, and the number of complexes displaced by reverse transcriptase is negligible. The tRNA-binding step cannot be rate limiting because the reaction saturates as a function of tRNA concentration, even after 10 min. The same is true for the mRNA-binding step (data not shown). The rate-limiting step is most likely the rearrangement of a preternary complex, in which both mRNA and tRNA are bound at the ribosome, but within which the tRNA anticodon is not base paired with the codon. The rearrangement creates a ternary complex in which the anticodon of the tRNA is base paired with a codon of the mRNA [Gualerzi and Pon 1981]. Preternary complexes might equilibrate with tRNA, mRNA, and

Figure 2. Detection of ternary complexes on the pRS170 transcript with several tRNA fragments. Reactions contained 0.2 \mu M 30S subunits and 0.5 \mu M of the fragments indicated above the lanes. Toeprint bands are marked with arrows. [Lane 1] A sequencing reaction containing ddTTP [A-lane]. [Bottom] The location of the toeprints within the initiation domain of the pRS170 transcript, 15 nucleotides 3' of the first base of the codons AUG, UGU, and UUU. The Shine and Dalgarno sequence (SD) is underlined.
30S subunits quickly, whereas the speed of the rearrangement step limits the formation of the ternary complex (Gualerzi et al. 1977; Ellis and Conway 1984). The speed of ternary complex formation with tRNA\textsuperscript{Met} and A13\textsuperscript{Met} at saturating tRNA concentrations is practically identical and approximately linear over the 10-min preincubation period (data not shown). The apparently better binding of tRNA\textsuperscript{Met} is thus not due to different speeds of the rearrangement step with these tRNAs but probably reflects a higher binding affinity of tRNA\textsuperscript{Met} in the preternary complex. The binding of DA39\textsuperscript{Met} and AT42\textsuperscript{Met} appears to be slightly better than that of the fragments A13\textsuperscript{Met}, A22\textsuperscript{Met}, sphinx-cys, sphinx-phe, flip-stem, and flop-loop, which all show very similar binding (Fig. 3). The binding of flop-loop was much poorer. Even the best binding fragment DA39\textsuperscript{Met} still binds much worse than tRNA\textsuperscript{Met}.

**Selection of initiator tRNA fragments by IF3**

To test the ability of IF3 to select the synthetic initiator tRNA fragments, we devised competition experiments in which one AUG-reading fragment species was added together with the UGU- and UUU-reading fragments (sphinx-cys and sphinx-phe) in toeprinting reactions. The incubations contained 30S subunits and the pRS170 transcript plus or minus IF3. The concentrations of the AUG-reading fragments were chosen so that they would yield a reasonable toeprint in competition with sphinx-cys and sphinx-phe (Fig. 4, lanes 1, 4, 7, 10, 13, and 16). Addition of IF3 leads to a clear selection of the toeprint band generated by the AUG-reading fragment only if it is a “true” initiator tRNA fragment (Fig. 4). Even the smallest fragment A13\textsuperscript{Met} is selected. The fragments flip-stem and flop-loop, with base changes in stem or loop, are not selected by IF3 over the fragments sphinx-cys and sphinx-phe (Fig. 4, lanes 14, 15, 17, and 18). In those latter incubations, IF3 destabilizes all ternary complexes. Bases in the anticodon stem, loop, and the anticodon itself must be important for the selection of the initiator tRNA by IF3.

**Loop conformation of the initiator tRNA fragments**

The anticodon loop conformation of several initiator tRNAs is different from that of the elongator tRNAs when the accessibility of loop phosphodiester bonds is probed with S1 nuclease (Wrede et al. 1979). The special loop conformation of the initiator tRNA (and its function as initiator tRNA) is dependent on the three GC base pairs of the anticodon stem (Seong and RajBhandary 1987). Here, we used the same technique to probe the anticodon loop conformation of several 3’ \textsuperscript{32}P-labeled tRNA fragments. The positions of the S1 nuclease cleavages were determined on a gel by comparison with the alkaline hydrolysis ladder of the fragments, which itself was aligned with the sequence-specific cleavage pattern obtained with RNase T\textsubscript{1} and RNase U\textsubscript{2}.

Flip-stem, flop-loop, sphinx-cys, and sphinx-phe all show S1 cuts in the loop region (Fig. 5). Cleavage positions are indicated in Figure 6. The cuts in the loop must occur more rapidly than cuts in the 3’ single-stranded region of the fragments to appear as labeled bands on the gel. The loops of the different fragments vary considerably in their sensitivity toward S1 nuclease, and so do the positions of the S1 cuts. Especially prone to S1 cleavage is the loop of the flop-loop fragment. The loop sequence of A13\textsuperscript{Met}, however, is highly resistant to S1 cleavage. Only one potential cleavage site was detected (Fig. 6). Strong resistance to S1 nuclease was also shown by the loop region of 5’ \textsuperscript{32}P-labeled A22\textsuperscript{Met}. No loop cleavages by nuclease S1 were observed under conditions in which the loops of 5’ \textsuperscript{32}P-labeled flip-stem and sphinx-cys showed extensive S1 cuts (data not shown). It thus appears that true initiator tRNA fragments have a
different loop conformation than the tRNA fragments with base changes in anticodon stem, loop, or the anticodon.

Selection of yeast and wheat germ initiator tRNA by IF3

The cytoplasmic initiator tRNAs of yeast and wheat germ (when charged with formylmethionine) are able to initiate translation properly in an E. coli in vitro system (Takeishi and Ukita 1968; Ghosh et al. 1974). Interestingly, their sequences differ extensively from that of E. coli initiator tRNA (Sprinzl et al. 1987); however, the three GC base pairs of the anticodon stem are conserved. The anticodon loops show differences mainly in nucleotide modifications [Fig. 7]. We asked whether either eukaryotic initiator tRNA could be recognized by E. coli IF3 in our in vitro system. An equimolar mixture of each initiator tRNA with E. coli tRNA\textsuperscript{Cy} and tRNA\textsuperscript{Phe} produces toeprints at +15 from the cognate codons on the pRS170 in vitro transcript [Fig. 8]. If IF3 is added in an equimolar amount to the 30S subunits, the initiator tRNA toeprints prevail, whereas the elongator tRNA toeprints disappear. Clearly, yeast and wheat germ initiator tRNAs are selected by E. coli IF3. Most likely, the similar anticodon units of the initiator tRNAs are responsible for the selection [as suggested by the data in Fig. 4].

Importance of the initiation codon

In the experiments reported so far, we have identified parts of the initiator tRNA crucial for its selection by IF3. The base pairing of the mRNA codon with the tRNA anticodon and the strength of that interaction might also play a role in initiator tRNA selection by IF3 (Berkhout et al. 1986). We tested the initiator tRNA fragment A13\textsuperscript{Met} on mRNAs with varied initiation codons [UUG, GUG, AUU] in competition with sphinx-cys and sphinx-phe on the codons UGC and UUC, respectively. The synthetic mRNA SD8AUG (the T7 transcript of pT7SD8AUG) contains a UGC codon in the +1 reading frame [Fig. 9], which should allow for a CG base pair between the cytosine in the third position of the codon and the guanine in the sphinx-cys anticodon. An AUG codon is also present in the SD8UUG and SD8GUG mRNAs [Fig. 9]. A13\textsuperscript{Met} was competed against sphinx-cys in 30S ternary complexes on those mRNAs. Two major toeprint stops at +15 from the AUG and UGC codons, respectively, were obtained [Fig. 9, lanes 1 and 7; Fig. 10, lane 4]. Part of the lower reverse transcriptase stop is also contributed by A13\textsuperscript{Met}-containing complexes which, by themselves, yield a double toeprint with a strong +15 and a weaker +16 band [Fig. 10, lane 3]. This makes the selection of A13\textsuperscript{Met} against sphinx-cys less obvious in 30S complexes on SD8AUG, SD8UUG, and SD8GUG plus IF3 [Fig. 9, lanes 2, 3, 8, and 9; Fig. 10, lanes 5 and 6]. Nevertheless, the control experiment with flip-stem and sphinx-cys plus IF3 clearly shows that neither sphinx-cys nor flip-stem are selected by IF3 because both toeprint bands disappear [Fig. 9, lanes 5, 6, 11, and 12]. Thus, sphinx-cys is in no case selected by IF3, even when codon-anticodon base-pairing with two GC base pairs can occur. On the other hand, A13\textsuperscript{Met} is selected by IF3 on the initiation codons UUG and GUG. Weak toeprint bands appear above the
Figure 5. Probing of the loop conformation of \( ^{32}\pmb{p} \)Cp 3'-end-labeled tRNA fragments with S1 nuclease. For each fragment, an alkaline hydrolysis ladder is shown at left and the S1 nuclease digest is shown at right (two lanes with different S1 nuclease concentrations for A13\(^{Met}\)-Met). The bracket on the right indicates the loop region of the fragments [except A13\(^{Met}\)-Met] where S1 nuclease cuts were found; the bracket on the left indicates the same loop region for A13\(^{Met}\)-Met.

main stops, mainly at +11 and +12 from the AUG codon. They are not +15 from any cognate codon and are not understood.

We then investigated the consequences of a base change in the third position of the initiation codon for selection of initiator tRNA by IF3. The SD8AUU synthetic mRNA [with an AUU initiation codon] contains a UUC codon in the +1 reading frame, complementary to the anticodon of sphinx-phe (Fig. 9). The AUU initiation codon is of particular interest, because so far it has only been found at the translational start site of infC, which encodes IF3 itself (Sacerdot et al. 1982). Models for autogenous regulation of IF3 translation have been proposed based on that unique initiation codon (Gold et al. 1984; Berkhout et al. 1986). A13\(^{Met}\) was added either alone or together with sphinx-phe in toeprinting reactions containing the SD8AUU mRNA. A13\(^{Met}\) at high concentration yields a weak toeprint band at +15 and +12 from the AUU codon (Fig. 11, lane 3). Both bands disappear when IF3 is added in equimolar amount to the 30S subunits (Fig. 11, lane 4). IF3 excludes A13\(^{Met}\) from the AUU codon. The same result was obtained when

Figure 7. Comparison of the anticodon stem and loops of E. coli, yeast, and wheat germ initiator tRNAs. E. coli tRNA\(^{Met}\) has Cm (2'-O-methylcytidine) instead of cytidine at position 32. Yeast and wheat germ initiator tRNAs, both have t6A (N-(9-β-D-ribofuranosylpurine-6-yl)carbamoyl-threonine) instead of adenosine in position 37. Furthermore, wheat germ initiator tRNA has a cytidine at position 33 instead of the “universal” uridine. The sequences are from Sprinzl et al. (1987).
Initiator tRNA, initiation codon, and IF3

anticodon stem and a proper anticodon loop are sufficient for IF3 selection. Larger RNAs are not selected better by IF3 and show only modestly enhanced affinity for ribosomes during ternary complex formation. The flop-loop fragment shows much poorer binding than the other fragments. Weaker binding might be due to the two adenosines 5’ to the anticodon (Uhlenbeck et al. 1982). A13Met, with only three GC base pairs of the anticodon stem maintained [the minimum requirement for ternary complex formation is 2 bp (Dube et al. 1969;]

Figure 8. Selection of E. coli, yeast, and wheat germ initiator tRNAs by IF3 on the pRS170 transcript. The initiator tRNAs were competed against E. coli tRNA^{cys} and tRNA^{phe}. Reactions contained the components indicated above the lanes at the concentrations (mM) indicated in parentheses. Toeprints corresponding to ternary complexes with the three tRNAs are indicated by arrows.

A13Met was competed with sphinx-phe. Toeprints originating from the respective complexes vanish in incubations containing IF3 [Fig. 11, lanes 6 and 7]. The third base of the initiation codon must play a crucial role in the IF3 selection of initiator tRNA.

The AUU-reading fragment sphinx-ile [Fig. 1], which restores the base pair with the uridine in the third position of the AUU codon, was also tested in toeprinting reactions with SD8AUU mRNA, either alone or in competition with sphinx-phe. Signals from ternary complexes were obtained with both fragments at +15 from the cognate codons [Fig. 11, lane 9], but they disappeared in incubations including IF3 [Fig. 11, lanes 10 and 11].

Discussion

IF3 selection of tRNA fragments

E. coli initiation factor IF3 is able to select initiator tRNA against elongator tRNAs in 30S ternary complexes [Hartz et al. 1989]. A most striking new finding of this study is that only synthetic RNA fragments with the anticodon stem and loop sequences of initiator tRNA are selected by IF3. Three GC base pairs of the

Figure 9. Selection of A13Met by IF3 on an AUG and UUG initiation codon against sphinx-cys on a UGC codon. The left six lanes contained reactions with SD8AUG mRNA; the right six lanes contained reactions with SD8UUG mRNA. Besides A13Met [lanes 1–3 and 7–9], flip-stem [lanes 4–6 and 10–12] was competed against sphinx-cys as a control experiment. Reactions contained all components indicated above the lanes at the concentrations (mM) indicated in parentheses. Toeprints resulting from ternary complexes at +15 from the AUG and UGC codons are marked with arrows. Note that the upper toeprint band and part of the lower toeprint band are contributed by ternary complexes with A13Met and flip-stem. [Bottom] The initiation domain of SD8AUG mRNA is shown 3’ to the Shine and Dalgarno (SD) sequence with the changes in the initiation codon that yield SD8UUG, SD8GUG, and SD8AUU mRNA.

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The requirement for a base in a specific position does not imply the recognition of that base by IF3; rather, the entire loop conformation could be a unit that is sensitive to base substitutions.

Other initiator tRNAs

Yeast and wheat germ initiator tRNAs are selected by IF3. Because the important features of the tRNA for IF3 selection are probably all located in the anticodon stem and loop, the minor differences in the anticodon loops of the initiator tRNAs (Fig. 7) do not interfere with IF3 selection. The anticodon loop conformations of E. coli, yeast, and mammalian initiator tRNA (which has the three GC base pairs of the anticodon stem and the same loop as wheat germ initiator tRNA) appear to be iden-

Rose et al. 1983), is selected by IF3. The three GC base pairs of A13Met are conserved in all cytoplasmic initiator tRNAs (Sprinzl et al. 1987). Changing these base pairs in E. coli initiator tRNA hampers initiation complex formation and translation in vitro (Seong and RajBhandary 1987). In the fragment flip-stem, which IF3 fails to select, these GC base pairs are changed to CG base pairs. Thus, the GC base pairs of the anticodon stem are crucial for IF3 recognition of the initiator tRNA.

The three GC base pairs help to create a special conformation of the anticodon loop of the initiator tRNA, as measured by S1 nuclease cleavage (Wrede et al. 1979; Seong and RajBhandary 1987). This special loop conformation probably is one important feature for initiator tRNA selection by IF3. Strikingly, we find high resistance to S1 nuclease in the anticodon loop of A13Met and A22Met but lower resistance in flip-stem. The same effect was observed previously by Seong and RajBhandary (1987), when they changed the three GC base pairs in the anticodon stem of initiator tRNA. Fragments not selected by IF3 (flip-stem, flop-loop, sphinx-cys, sphinx-phe) all share a higher accessibility in their loops to S1 cleavage and, thus, probably have different loop confor-

Figure 10. Selection of A13Met by IF3 on a GUG initiation codon against sphinx-cys on a UGC codon. In addition to SD8GUG mRNA, reactions 3-6 contained the components indicated above the lanes at the concentrations (μM) indicated in parentheses. Toeprints are indicated by arrows. Note that the upper band and part of the lower band are contributed by ternary complexes with A13Met. (Lane 1) A sequencing reaction including ddCTP [G-lane].

Figure 11. A13Met and sphinx-ile both fail to be selected by IF3 on the AUU initiation codon of SD8AUU mRNA. A13Met [lanes 5-7] and sphinx-ile [lanes 9-11] were each competed against sphinx-phe, which forms ternary complexes on the UUC codon in the +1 frame. Lanes 3-11 contained the components above the lanes at the concentrations (μM) indicated in parentheses. Toeprints at +15 from the cognate codons are indicated by arrows. (Lanes 1 and 2) Sequencing lanes containing ddTTP [A-lane] and ddATP [U-lane], respectively.
tical [Wrede et al. 1979]. We speculate that those positions with varied nucleotides are not directly inspected by IF3 action but still allow the special anticodon loop conformation of initiator tRNAs.

What biological/biochemical function is served by the yeast initiator tRNA structure that responds to E. coli IF3? We see this result as implying the capacity in yeast to select the initiator tRNA as the sole occupant of the ribosomal P site during initiation, exactly as occurs in E. coli. Two wonderful papers from Donahue’s lab [Cigan et al. 1988; Donahue et al. 1988] ask what requirements exist for initiation codon selection. The initiator tRNA anticodon was changed to 3'-UCC-5', and that tRNA initiates at AGG codons following all the rules of translation initiation in yeast. Thus, yeast contains some mechanism for allowing an initiator to use codon/anticodon pairing to select the initiation codon, and that selection continues to operate with single nucleotide substitutions at the middle positions of the codon and anticodon [nucleotide substitutions that we have not tried yet for IF3 selectivity]. As importantly, amino acid substitutions in the eIF-2β protein create the dominant phenotype in yeast of allowing yeast initiator tRNA to start translation at an AUU codon (as occurs without difficulty in E. coli). We see Donahue’s work as suggesting that a yeast initiation factor (perhaps even eIF-2B) functions in yeast of allowing yeast initiator tRNA to start translation in vivo (Gold 1988). CUG also serves as an initiation codon in E. coli, and free 30S particles are probably saturated with IF3 (Howe and Hershey 1981).

The binding of IF3 to the 16S tRNA of 30S subunits near the 3’ end of 16S tRNA (Wickstrom 1983), which must be in proximity to the ribosomal P site, makes it plausible that IF3 itself “inspects” the anticodon stem and loop of the tRNA bound in the ribosomal P site. This inspection must at least include the third position of the codon/anticodon interaction (Fig. 12). IF3 promotes the exchange of elongator tRNAs bound in ternary complexes in the ribosomal P site; elongator tRNAs are bound very stably in the absence of IF3 (Pon and Gualerzi 1974). Ternary complex formation with initiator tRNA is also catalyzed by IF3; initiator tRNA, however, is stably bound in the complex when IF3 is present [Risuelo et al. 1976]. If IF3 lowers the activation energy for the codon/anticodon base-pairing step (Gualerzi et al. 1974), it might bind to some part of the anticodon stem and loop of initiator tRNA to keep the initiator tRNA from being exchanged after ternary complex formation. Most likely, the third position of the initiation codon is part of such a binding site. Whereas only several bases need to be contacted directly, the other bases of the anticodon stem and loop are crucial to maintain certain bases in the conformation that can be bound by IF3.

Methods

tRNA

Total E. coli tRNA and uncharged E. coli tRNAMet and tRNAPro were purchased from Boehringer Mannheim. E. coli tRNAPro was purchased from Subhrden RNA. Yeast and wheat germ tRNAMet were a gift of P. Sigler and R. Basavappa. The synthetic tRNA fragments shown in Figure 1 were generated by T7 tran-

Figure 12. Model for IF3 selection of the initiator tRNA. IF3 binds to the 3’ end of 16S tRNA and contacts some part of the anticodon stem and loop and the initiation codon.
cription from the oligodeoxynucleotide pts as top strand and oligodeoxynucleotides as bottom strand, which contain the complementary sequence to the generated fragments [Table 1]. T7 transcription was performed according to Milligan et al. (1987). The sphinx-ile fragment was synthesized by S. Beckmann. The tRNA fragments were purified on a 20% polyacrylamide gel. Some fragments were purified with and without an extra 3' nucleotide [indicated by N in parentheses in Fig. 1]. A13Met was only purified as a mixture of the two fragments. The 3' nucleotide of purified tRNA fragments was determined by 3'-end labeling with 32pCp according to Beckett and Uhlenbeck [1984], followed by RNase digestion and two-dimensional thin-layer chromatography [Nishimura 1972]. Fragments without the extra 3' nucleotide exhibited the expected 3' nucleotide, whereas fragments with an extra nucleotide were random [but with a preference for cytidine]. Furthermore, 3'-end-labeled fragments were partially sequenced with RNase T1 and RNase U2, according to Donis-Keller et al. (1977). The expected partial sequences were obtained.

Enzymes

AMV reverse transcriptase and MMLV reverse transcriptase were obtained from Life Sciences, Inc. and Bethesda Research Laboratories, respectively. T7 RNA polymerase was provided by O. Uhlenbeck. T4 polynucleotide kinase and T4 RNA ligase were obtained from Boehringer Mannheim. RNase T1 and RNase U2 were obtained from Calbiochem.

Initiation factor IF3 and Ribosomes

Purified E. coli initiation factor IF3 was a gift from C. Gualerzi. 30S subunits, prepared according to Kenney et al. [1979], were a gift from R. Traut.

mRNAs

In vitro RNAs from the PvuII-cut plasmids pRS170, pT7SD8AUG, pT7SD8UG, and pT7SD8AUU were synthesized with T7 polymerase according to Lowary et al. [1986]. The RNAs were purified on a 6% polyacrylamide gel.

Table 1. Description and use of oligodeoxynucleotides

| Sequence | Use |
|----------|-----|
| taatagcactcacttag | pts top strand, T7 transcription |
| ctcgccggtatagcgccag cgctctctagcgacttagctgctgta | T7 transcription DA39Met |
| ggctagctggcagctagctata | T7 transcription AT42Met |
| ctgcggcagctagctata | T7 transcription A22Met |
| ggttctactagctata | T7 transcription A13Met |
| ctcggaagcagctagctata | T7 transcription flip-stem |
| ctggcagctagctata | T7 transcription flop-loop |
| gttggagctagctata | T7 transcription sphinx-cys |
| gttggagctagctata | T7 transcription sphinx-phe |
| gttggagctagctata | T7 transcription sphinx-ile |
| gttggagctagctata | LP134 primer, compl. to lacZ gene from 57 to 73 |
| gttggagctagctata | cloning of pT7SD8AUU |
| gttggagctagctata | cloning of pT7SD8UG |
| gttggagctagctata | cloning of pT7SD8AUU |

Plasmids

The plasmid pRS170 contains the sequence from –92 to 107 of bacteriophage T4 gene 32 [Krisch and Allet 1982] and has been described in Hartz et al. [1989]. To construct plasmids pT7SD8AUG, pT7SD8UG, and pT7SD8AUU, the oligodeoxynucleotides shown in Table 1 were notch-cloned by D. Barrick into the large fragment of BglII and PstI-digested pBC39 vector according to Childs et al. [1985]. pBC39 was constructed by replacing the Pra2 promoter between EcoRI and HindIII of pBC29 [Childs et al. 1985] with the tacII promoter. The tacII promoter of the pBC39 clones was subsequently substituted by the T7 φ10 promoter to yield the plasmids pT7SD8AUG, pT7SD8UG, and pT7SD8AUU. Plasmid pT7SD8AUU was constructed by cloning the two complementary oligodeoxynucleotides shown in Table 1 as double-stranded DNA into the large fragment of HindIII- and BamHI-digested plasmid pTS27. pTS27 is a derivative of pTS24 with the F2 origin between Sall–EcoRI cut out and replaced by a Sall–SmaI–EcoRI linker. pTS24 was constructed by replacing the EcoRI–HindIII Pra2 promoter fragment of pRS210 [McPheeters et al. 1988] with the phase T7 φ10 promoter (T. Schneider, unpubl.). Cloning protocols were adapted from Maniatis et al. [1982].

Oligodeoxynucleotides

Oligodeoxynucleotides were synthesized on an Applied Biosystems Model 380A DNA synthesizer and purified by preparative polyacrylamide gel electrophoresis. Oligodeoxynucleotides used as bottom strands to synthesize the tRNA fragments of Figure 1 were sequenced with AMV reverse transcriptase by using a protocol for RNA sequencing [McPheeters et al. 1986]. The pts top strand oligodeoxynucleotide was used as primer. The sequences and use of the oligodeoxynucleotides are shown in Table 1.

Extension inhibition

Extension inhibition (or toeprinting) was performed essentially as described in Hartz et al. [1988]. Ten-microliter toeprinting reactions contained 6.7 nm of the in vitro transcripts, annealed to primer LP134 (Table 1), along with ribosomes, tRNAs, and IF3, as specified in the Results. Reactions with all ingredients were prepared on ice and preincubated for 10 min at 37°C, fol-
allowed by primer extension with MMLV reverse transcriptase for 15 min at 37°C.

Labeling of tRNA fragments

tRNA fragments were 3'-end-labeled with 32P-cytidine according to Beckett and Uhlenbeck [1984]. With the exception of A13Met, only fragments without the extra random nucleotide on the 3' end were 3'-end-labeled. To 5' 32P-labeled A22Met, flip-stem, and spinhex-cys, 10 pmoles of each fragment was dephosphorylated with 1 unit calf intestinal alkaline phosphatase in a 50-μl reaction containing 50 mM Tris-Cl (pH 8.0) and 0.1 mM EDTA. The reaction was incubated for 30 min at 45°C. The samples were phenol extracted four times, and the fragments were precipitated with three volumes of ethanol. 5' 32P-labeling was performed with [γ-32P]ATP, using a standard protocol for oligodeoxynucleotide labeling (Maniatis et al. 1982). Only fragments with an extra random nucleotide on the 3' end were 5'-end-labeled.

Partial alkaline hydrolysis of tRNA fragments

Four-microliter reactions contained 1.6 pmoles of 3'- or 5'-end-labeled fragments in 50 mM sodium carbonate buffer (pH 9.0), 1 mM EDTA. They were incubated for 8 min at 95°C and cooled, and 10 μl of loading buffer was added. Five-microliter samples were analyzed on a 20% polyacrylamide gel containing 8 M urea.

Probing of anticodon loop conformation with nuclease S1

Nuclease S1 digestions were performed essentially as described by Wrede et al. [1979]. Five-microliter reactions contained 0.8 pmoles of 3'- or 5'-end-labeled fragments and 1 μg of carrier tRNA in 25 mM sodium acetate (pH 4.5), 5 mM MgCl2, 50 mM KCl, and 1 mM ZnCl2. Nuclease S1 was added at the concentrations indicated in the figure legends, and the reactions were incubated for various time intervals at 37°C. Only reactions from 10-min incubations, which showed the most extensive loop cleavage, are shown in Figure 5. Reactions were stopped by adding 10 μl of loading dye [Hartz et al. 1988] containing 4 mM ATP and 0.2 μg/μl carrier tRNA. Five-microliter samples were analyzed on a 20% polyacrylamide gel containing 8 M urea. Cleaved RNAs that carry the 3'-end label and are created by S1 digestion [with 5'-phosphate ends] should migrate faster than those created by alkaline hydrolysis [with 5' hydroxyl ends] [Wrede et al. 1979]. However, in our experiments, S1-cleaved fragments appear to run slightly slower than alkaline-cleaved fragments; the dinucleotides cleaved off in the 3' single-stranded region appear to migrate slightly slower than the di-nucleotides created by alkaline hydrolysis, and even the full-length fragments appear to run slower in the lanes with S1 incubation [Fig. 5]. This effect of S1 is not observed with 5'-end-labeled fragments [data not shown]. If the S1 preparation contained a 3' phosphatase as a contaminant, only the electrophoresis of 3'-end-labeled fragments would be affected. We assigned the S1 cleavage sites based on these observations.

Relative toeprints

To quantify the toeprint stop and the 5'-end stops, the gels were scanned on an AMBIS Systems radiodensitometric imaging system and the bands were electronically integrated. Relative toeprints (%) were calculated: [toeprint/toeprint + 5' ends] × 100. To quantify the 5' ends, all bands 5' of the toeprint band were integrated.

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