In vitro trans-translation of *Thermus thermophilus*: Ribosomal protein S1 is not required for the early stage of trans-translation

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ABSTRACT

Transfer-messenger RNA (tmRNA) plays a dual role as a tRNA and an mRNA in trans-translation, during which the ribosome replaces mRNA with tmRNA encoding the tag-peptide. These processes have been suggested to involve several tmRNA-binding proteins, including SmpB and ribosomal protein S1. To investigate the molecular mechanism of trans-translation, we developed in vitro systems using purified ribosome, elongation factors, tmRNA and SmpB from *Thermus thermophilus*. A stalled ribosome in complex with polyphenylalanyl-tRNAPhe was prepared as a target of tmRNA. A peptidyl transfer reaction from polyphenylalanyl-tRNAPhe to alanyl-tmRNA was observed in an SmpB-dependent manner. The next peptidyl transfer to aminoacyl-tRNA occurred specifically to the putative resume codon for the tag-peptide, which was confirmed by introducing a mutation in the codon. Thus, the in vitro systems developed in this study are useful to investigate the early steps of trans-translation. Using these in vitro systems, we investigated the function of ribosomal protein S1, which has been believed to play a role in trans-translation. Although *T. thermophilus* S1 tightly bound to tmRNA, as in the case of *Escherichia coli* S1, it had little or no effect on the early steps of trans-translation.

Keywords: tmRNA; trans-translation; S1; *Thermus thermophilus*

INTRODUCTION

Transfer-messenger RNA (tmRNA, also known as SsrA RNA or 10Sa RNA) is a bifunctional RNA widely distributed among eubacteria. tmRNA has a tRNA-like domain (TLD) and a peptide-coding region (mRNA domain). Both terminal regions form a tRNA-like structure (Komine et al. 1994; Ushida et al. 1994), while the middle part typically forms four pseudoknots (PK) and some helices (Felden et al. 1996; Williams and Bartel 1996). The 3’ end of tmRNA, like tRNA, is aminoacylated by alanyl-tRNA synthetase (AlaRS) (Komine et al. 1994; Ushida et al. 1994). tmRNA was later shown to act also as mRNA, to facilitate a non-canonical translation, trans-translation (Keiler et al. 1996; Himeno et al. 1997). It produces a chimera polypeptide from two different RNAs, a problematic mRNA and tmRNA with a message for degradation. Thus, tmRNA plays a role in rescuing the stalled ribosome and tagging a signal for degradation to the incomplete protein from a terminator-less mRNA (Muto et al. 2000; Withey and Friedman 2003).

tmRNA forms a complex with several protein factors, such as SmpB, elongation factor Tu (EF-Tu), and S1, in vivo and in vitro (Karzai et al. 1999; Wower et al. 2000, 2004; Barends et al. 2001; Karzai and Sauer 2001; Okada et al. 2004). SmpB binds TLD (Hanawa-Suetsugu et al. 2002; Gutmann et al. 2003; Nameki et al. 2005) to perform functions in the processes of trans-translation, recruitment of tmRNA to the stalled ribosome (Karzai et al. 1999;

S1 is a component of the small subunit of the ribosome (Subramanian 1983; Shiryaev et al. 2002). Due to the weak association with the ribosome, some of S1 seems to be apart from ribosomes in the cytoplasm. S1 specifically binds to the uridine-rich region upstream of the SD sequence (Boni et al. 1991; Zhang and Deutscher 1992) or the 5’-untranslated region of an mRNA without an SD sequence (Tzareva et al. 1994) and possesses a nucleic acid unwinding ability (Thomas and Szer 1982). It ensures the translation of mRNAs bearing no or weak SD sequences (Roberts and Rabinowitiz 1989; Tzareva et al. 1994) of mRNAs that bear long SD sequences (Komarova et al. 2002) or of most natural mRNAs in vivo (Sorensen et al. 1998). The low-G+C group of Gram-positive bacteria lacks S1 but has a strong SD sequence in each mRNA instead (Isoson and Isoson 1976; Farwell et al. 1992). Thus, S1 is thought to help the association between the ribosome and the ribosomes besides the SD and anti-SD interaction, which is significant for efficient translation of mRNAs, especially those with a weak SD sequence (Boni et al. 1991; Zhang and Deutscher 1992; Tzareva et al. 1994; Tedin et al. 1997; Sengupta et al. 2001; Komarova et al. 2002).

Since S1 cross-links to the upstream region of the tag-encoding sequence of tmRNA that is significant for trans-translation, the interaction between S1 and tmRNA is presumed to be critical for initiation of tag-translation (Williams et al. 1999; Wower et al. 2000; Lee et al. 2001). Wower et al. (2000) have shown that Escherichia coli S1 enhances the binding of tmRNA to the ribosome in the absence of SmpB. E. coli S1 induces a conformational change in tmRNA, making the nucleotides around the residue codon accessible to a solvent (Bordeau and Felden 2002). A conformational change in tmRNA by S1 has also been suggested by a cryo-electron microscopic study on a Thermus thermophilus Ala-tmRNA-SmpB-EF-Tu-GDP-kirromycin-ribosome complex (Valle et al. 2003). Collectively, these findings raise the possibility that S1 binds tmRNA, induces it to the ribosome, anchors the upstream region of the residue codon on the ribosome, and destabilizes the surrounding structured region to precisely set the residue codon at the A site. On the other hand, some studies have argued against the significance of S1 for trans-translation. S1 is absent in the low-G+C group of Gram-positive bacteria, and tmRNA from Bacillus subtilis that lacks S1 can facilitate trans-translation in E. coli that has S1 (Ito et al. 2002). Overproduction of the N-terminal fragment inhibits general translation but not trans-translation in E. coli, also implying the insignificance of S1 for trans-translation (McGinness and Sauer 2004). S1 binds to the junction of the head, platform, and main body of the 30S subunit according to a cryo-electron microscopic study (Sengupta et al. 2001), which is reasonable for its function in binding and unwinding the upstream region of the initiation point of structured mRNA to initiate translation. However, upon tmRNA binding to the ribosome, the S1 binding region on tmRNA is apparently far from the position of S1 on the 30S subunit, leading to an idea that even if S1 has a role in trans-translation, that role may be independent of its role in canonical translation.

In the present study, we developed in vitro trans-translation systems composed of factors from T. thermophilus to study the role of S1 in trans-translation. Many biochemical and genetic studies on biological events in bacteria have been performed using E. coli, although structural studies have often been carried out using factors from thermophiles. T. thermophilus is a thermophile in which the structures of factors required for trans-translation, such as the ribosome (Yusupova et al. 1991; Carter et al. 2001) and SmpB (Someya et al. 2003), have been extensively studied. However, there have been only a few biochemical and no genetic studies (Yusupov et al. 1991; Yusupova et al. 1991; Uzawa et al. 2002; Stepanov and Nyborg 2003). In contrast, several kinds of cell-free trans-translation systems have been developed in E. coli (Himeno et al. 1997; Shimizu and Ueda 2002; Asano et al. 2005), although with few structural studies. Thus, T. thermophilus cell-free trans-translation systems might be expected to link structural studies to biochemical and genetic studies on trans-translation. The in vitro systems developed here can assess the early steps of trans-translation, which revealed that T. thermophilus S1 has little or no effect on these steps.

RESULTS AND DISCUSSION

In vitro poly (U)-dependent poly(Phe) synthesis

The ribosome translates poly (U) into poly (Phe) and probably stalls at the very 3′ end of the poly (U). This type of stalled ribosome has already been reported to be a target of tmRNA for trans-translation in E. coli (Himeno et al. 1997; Shimizu and Ueda 2002). A highly purified S1-free ribosome from T. thermophilus was incubated with poly (U), [14C]Phe-tRNAPhe, EF-Tu, EF-Ts, EF-G, and GTP. As expected, poly (U)-dependent [14C]phenylalanine incorporation was observed. Poly (U), GTP, and EF-G were all essential for the reaction (Fig. 1, -poly (U), -GTP, and -G). The efficiency decreased to 28% and 55% by exclusion of EF-Tu and EF-Ts, respectively, from the reaction mixture (Fig. 1, -Tu and -Ts). Considering that trans-translation can occur, but inefficiently, on the ribosomes stalled in the middle of mRNA (Ivanova et al. 2004; Asano et al. 2005), only a limited amount of ribosomes was added in the
acylation of tmRNA by AlaRS, as in the case of 4 in the absence of SmpB, while it was 50% in the presence of SmpB. The binding of SmpB to tmRNA was confirmed by gel mobility shift assay (Nameki et al. 2005). Furthermore, SmpB enhanced the aminoacylation of tmRNA from T. thermophilus, as shown in Fig. 2A. This result indicates that the transfer actually occurred as expected in consideration of the indispensability of SmpB in E. coli (Asano et al. 2005).

After the transfer reaction and presumably translocation of tmRNA from the A site to the P site, the first codon of the mRNA domain should be set at the A site. The resume codon has been predicted to be G91C92C93, an alanine codon, for T. thermophilus tmRNA from a sequence comparison (Gueneau de Novoa and Williams 2004). In order to monitor the decoding of the resume codon, the stalled ribosome was incubated with unlabeled Ala-tmRNA and [3H]Ala-tRNAAla together. This reaction was expected to result in the production of poly (Phe)-Ala-[3H]Ala. To ascertain the validity of this system, a mutant tmRNA (tmRNA92U), in which the putative resume codon was altered from GCC to GUC encoding a valine, was constructed, and it was used for the resume codon decoding assay in combination with [3H]Ala-tRNAAla or [3H]Val-tRNAVal.

First, [3H]Ala-tRNAAla together with Ala-tmRNAwt or Ala-tmRNA92U was added to the stalled ribosomes (Fig. 2B, left panel). [3H]Ala was incorporated into poly (Phe)-Ala-[3H]Ala when wild-type tmRNA was used, whereas no incorporation of [3H]Ala was observed when tmRNA92U was used. Next, [3H]Val-tRNAVal was added instead of [3H]Ala-tRNAAla (Fig. 2B, right panel). In this case, [3H]Val was incorporated into poly (Phe)-Ala-[3H]Val only when tmRNA92U was used. Taken together, the results show that the resume codon decoding occurred only when the predicted resume codon matched the anticodon of the aminoacyl-tRNA used. These results confirmed that our in vitro system is valid for monitoring the initial steps of trans-translation, including the two steps (1) binding of Ala-tmRNA to the ribosome to undergo a peptide transfer reaction, and (2) setting of the resume codon to the A site to be translated. These results also indicate that the resume codon of T. thermophilus tmRNA is G91C92C93 (an alanine codon), and the amino acid sequence of the tag-peptide of T. thermophilus is therefore (A)ANTNYALAA, as predicted from a phylogenetic study (Gueneau de Novoa and Williams 2004).

Effect of S1 on translation and trans-translation

Almost all populations of the ribosomes used in the present study were free of S1, which was checked by SDS polyacrylamide gel electrophoresis. No evident band corresponding to S1 was detected in the ribosome fraction (data not shown). It has been reported that poly (U)-dependent poly (Phe) synthesis occurred in E. coli even in the absence of S1 and that the addition of S1 to the reaction promoted the
incorporation of phenylalanine (Van Dieijen et al. 1975; Yokota et al. 1977). Poly (Phe) synthesis without S1 was also observed in the present *T. thermophilus* system (Figs. 1, 3A). Exogenous addition of S1 to the reaction remarkably promoted both the initial rate and the plateau level of polyphenylalanine synthesis (Fig. 3A), and 1.2 mM S1 enhanced the rate by fourfold. This result indicates that S1 is functional in the in vitro translation system.

*E. coli* S1 has been reported to bind tightly to tmRNA (Wower et al. 2000; Hanawa-Suetsugu et al. 2001). In this study, we examined the binding property of S1 from *T. thermophilus* to tmRNA by gel mobility shift assay (data not shown). Shifted bands were observed with increasing concentration of S1. The Kd value was estimated to be 5–10 nM from the free tmRNA bands. Thus, S1 from *T. thermophilus* binds tmRNA as tightly as that from *E. coli*, implying a significance of S1 for trans-translation. We reproducibly detected plural shifted bands with increase in the concentration of S1 as in the case of *E. coli* S1, which may be reflected by binding of multiple molecules of S1 to a single tmRNA (Bordeau and Felden 2002).

The effect of S1 on the trans-transfer step was examined by adding exogenous S1 to the Ala-tmRNA-SmpB complex prior to the addition of the complex to the stalled ribosomes free of S1. Since the concentration of S1 used was far higher than the apparent Kd value estimated from the gel mobility shift assay, Ala-tmRNA was expected to predominantly form a complex with S1. The reaction was started by adding a mixture containing [3H]Ala-tmRNA, SmpB, EF-Tu, GTP, and S1 to S1-free stalled ribosome. As already shown in Figure 2A, alanine incorporation was observed even in the absence of S1 (Fig. 3B, closed circles). Addition of S1 to the reaction showed no enhancement of trans-transfer, although a slight inhibition was observed at high concentration (Fig. 3B). The effect of the lower concentration of S1 was also examined, but no effect was observed (data not shown).

The effect of S1 on the setting of the resume codon at the A site to be decoded after trans-transfer was examined. It should be noted that because Ala-tmRNA and Ala-tRNAAla were added together to the stalled ribosome, the observed effect of S1 includes the effects not only on the resume codon decoding but also on trans-transfer. A mixture containing [3H]Ala-tRNAAla and a complex of unlabeled Ala-tmRNA, SmpB, and various concentrations of S1 was added to the stalled ribosome. Incorporation of [3H]Ala into a peptide was observed in the absence of S1 (Figs. 2B, 3B). S1 slightly inhibited rather than enhanced this reaction, which can be attributed to the effect of S1 on the trans-transfer step. No effect of S1 at its lower concentration was observed. These results strongly suggest that S1 is not involved in the step of the resume codon decoding.
The above results from the cell-free trans-translation systems demonstrate that *T. thermophilus* S1, even if it causes a structural change around the resume codon, does not contribute to the early steps of trans-translation involving tmRNA recognition by the ribosome, peptidyl-transfer from peptidyl-tRNA to Ala-tmRNA, setting of the resume codon to the A site, and peptidyl-transfer from peptidyl-tmRNA to Ala-tRNA<sub>Ala</sub>. It is possible that S1 functions in later steps of trans-translation, such as in the exit of TLD from the E site. Alternatively, it may function only in the protection of tmRNA from degradation in the cell.

In the present study, we demonstrated that *T. thermophilus* S1 was not essential for the early steps of trans-translation, although it tightly bound to tmRNA. This suggests that S1 has no role in the functional interaction between tmRNA and the stalled ribosome or in a structural link between the tRNA domain and the mRNA domain for functional accommodation of tmRNA in the stalled ribosome. SmpB, another tmRNA-binding protein, may be deeply involved in this mechanism. Our *T. thermophilus* cell-free system should contribute to further clarification of the molecular mechanism of trans-translation in concert with recent and future progress of structural studies.

**MATERIALS AND METHODS**

**Cloning and purification of tmRNA**

The *ssrA* gene was amplified from genomic DNA from *T. thermophilus* HB8 by polymerase chain reaction with two primers, 5′-TTACTCTACacgGgGTTGGAACGGCTGACGCGGCT-3′ and 5′-GGCTGCTGGTACCgagggagactcGCTGAGCTGAGGGAAGTC-3′, in which the sense or anti-sense sequence of either end of the gene for tmRNA from *T. thermophilus* (GenBank, Y15063) is underlined. For proper processing of the primary transcript into a mature tmRNA in *E. coli*, the flanking sequences of the *E. coli ssrA* gene (GenBank, D12501) shown as lowercase letters, were added. A restriction enzyme cleavage sequence, shown in italics, was also added to each primer. The amplified DNA fragment was ligated into a plasmid vector, pGEM-EX2 (Promega), after digestion with XbaI and SphI. The resulting plasmid was termed pTtmRNA<sup>wt</sup>. The sequence of the recombinant tmRNA was revealed to be version 2 of the *T. thermophilus* tmRNA sequence in the tmRNA Web site that has G<sub>310</sub> instead of A<sub>310</sub> (Guinou and Williams 2004). The plasmid for preparation of a mutant tmRNA<sub>92U</sub> was constructed by PCR using the primers 5′-GGCTGCTGGTACCgagggagactcGCTGAGCTGAGGGAAGTC-3′ and 5′-GGCTGCTGGTACCgagggagactcGCTGAGCTGAGGGAAGTC-3′. The amplified DNA and pTtmRNA<sup>wt</sup> were ligated after digestion with HsSIII and XbaI, resulting in pTtmRNA<sup>92U</sup>. The plasmids were transformed into *E. coli* strain BL21 (DE3) (*ssrA<sup>kan</sup>*) constructed by P1 transduction from W3110 Δ*ssrA* (Komine et al. 1994). *T. thermophilus* tmRNA<sup>wt</sup> and tmRNA<sup>92U</sup> were overexpressed and purified as described previously (Hanawa-Suetsugu et al. 2001).
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Preparation of EF-Tu, EF-Ts, EF-G, S1, AlaRS, Smb, and ribosome from *T. thermophilus*

Expression vectors of EF-Tu, EF-Ts, and EF-G were kind gifts from Dr. Sprinzl. Recombinant proteins were overexpressed in the *E. coli* strain BL21 (DE3) and purified as previously reported (Blank et al. 1995). The genes for SmbP, AlaRS, His-tagged EF-Tu (EF-Tu-His), and ribosomal protein S1 were cloned from the *T. thermophilus* HB8 genome by the RIKEN Structural Genomics Initiative (Yokoyama et al. 2000).

Ribosomal protein S1 cloned into pET11b (Novagen) was expressed in *E. coli* strain BL21 (DE3) codonplus RIL (Stratagene). Recombinant S1 was precipitated by the addition of 50% saturated ammonium sulfate to the supernatant, which was obtained after heat treatment of the cell lysate for 20 min at 70°C. The supernatant was precipitated in 50 mM Tris-Cl (pH 7.6), 10 mM MgCl₂, 4.6 mM 2-mercaptoethanol, and 20 μM GDP containing 100 mM NH₄Cl. The supernatant of the cell extract that had been subjected to heat treatment at 65°C was subjected to anion-exchange chromatography (HiTrap Q HP), EF-Tu-His fractions enriched by a 50–500 mM linear gradient were eluted with 20 mM HEPES-NaOH (pH 7.0) containing 150 mM NaCl and 4 mM 2-mercaptoethanol. The protein yield was 1 mg per 1 g of frozen cells.

EF-Tu-His cloned into pET11b was overexpressed in *E. coli* strain BL21 (DE3). Cells were disrupted with a French Press in buffer A (20 mM Tris-HCl [pH 7.6], 10 mM MgCl₂, 4.6 mM 2-mercaptoethanol, and 20 μM GDP) containing 100 mM NH₄Cl. The supernatant of the cell extract that had been subjected to heat treatment at 65°C was used for anion-exchange chromatography (HiTrap Q HP), EF-Tu-His fractions enriched by a 50–500 mM linear gradient elution of KCl were loaded onto Ni-NTA agarose (QIA-GEN). Pooled fractions were dialyzed against buffer A containing 50 mM KCl. The protein yield was 15 mg per 1 g frozen cells.

AlaRS cloned into pET11b was overexpressed in *E. coli* strain BL21 (DE3). Cells were disrupted with a French Press in buffer A with 20 mM Tris-HCl (pH 8.0) buffer containing 100 mM NH₄Cl, 0.5 mM EDTA, 4.6 mM 2-mercaptoethanol, and 5% glycerol. The supernatant of the cell extract that had been subjected to heat treatment at 65°C was loaded onto Ni-NTA agarose. After the resin had been washed with 20 mM Tris-HCl (pH 7.6) buffer containing 10 mM imidazole and 1 M NH₄Cl, absorbed proteins were eluted with 200 mM imidazole, pooled, and dialyzed against buffer B (HEPES-NaOH [pH 7.0] and 100 mM KCl). Further purification was performed with anion-exchange chromatography (Resource Q) and size-exclusion chromatography (Superdex-200). The protein yield was 0.5 mg per 1 g frozen cells.

Ribosomes from *T. thermophilus* were prepared from frozen cells harvested in the middle-log phase as previously reported (Gogiya et al. 1986) with slight modification. 70S ribosome yield was 1 mg per 1 g of frozen cells.

For the preparation of AlaRS, Val-tRNA synthetase (ValRS), and phenylalanyl-tRNA synthetase (PheRS) were prepared from *E. coli* cells using DEAE-Toyopearl 650M (Tohos) (Tamura et al. 1991, 1994). AlaRS was further purified by hydroxyapatite column chromatography, Gigapite (Seikagaku Corporation).

Preparation of aminoacyl-RNA

AlaRS, valyl-tRNA synthetase (ValRS), and phenylalanyl-tRNA synthetase (PheRS) were prepared from *E. coli* cells using DEAE-Toyopearl 650M (Tohos) (Tamura et al. 1991, 1994). AlaRS was further purified by hydroxyapatite column chromatography, Gigapite (Seikagaku Corporation).

Poly (Phe) synthesis was performed by incubating 0.5 pmol of S1-free ribosomes, 1.2 μg of poly (U) (Sigma), 5 pmol of [3-3H]phenylalanine (18.4 GBq/mmol, PerkinElmer) or 10 μM L-[3-3H]phenylalanine (1.96 TBq/mmol, Amersham Biosciences) in aminoacylation buffer. Ninety-three percent and 86% of total tRNAPhe were aminoacylated by AlaRS and ValRS from *E. coli* by incubating 1 μM tRNA and AlaRS from *E. coli* with 8 μM L-[3-3H]alanine (15 TBq/mmol, PerkinElmer) or 30 μM unlabeled alanine in aminoacylation buffer for 10 min at 37°C. Twenty-eight percent of total tRNA was aminoacylated with [3H]alanine. Ala-tRNA was further purified by an immobilized EF-Tu column (Ribeiro et al. 1995) with slight modifications as follows: The immobilized EF-Tu column was prepared by incubating 2 ml of Ni-NTA (Qiagen) and 8 mg of His-tagged EF-Tu from *T. thermophilus* in a buffer (50 mM Tris-HCl [pH 7.5], 50 mM NH₄Cl, 50 mM KCl, 10 mM MgCl₂, 1 mM GTP, 5 mM phosphoenolpyruvic acid, 15 units of pyruvate kinase [Sigma], and 10 mM 2-mercaptoethanol) for 10 min at 37°C. The Ni-NTA/EF-Tu/GTP was mixed with 1.6 nmol of aminoacylated RNA and incubated for 2 min on ice. After purifying the mixture into an empty column (Poly-Prep Chromatography Columns, Bio-Rad Laboratories), deacylated RNA was eluted with 4 ml of buffer B (50 mM Tris-HCl [pH 7.5], 50 mM NH₄Cl, 50 mM KCl, 10 mM MgCl₂, 1 mM GTP, and 5 mM 2-mercaptoethanol) and then with 4 ml of buffer C (50 mM HEPES-Na [pH 7.5], 50 mM NH₄Cl, 150 mM NaCl, 10 mM MgCl₂, 50 μM GTP, and 5 mM 2-mercaptoethanol). Then, aminoacylated RNA was eluted with 4 ml of buffer D (50 mM Tris-HCl [pH 7.5], 1 M NaCl, 10 mM MgCl₂, 1 mM GDP, and 5 mM 2-mercaptoethanol) and precipitated with ethanol. We confirmed that nonaminoacylated tmRNA did not bind to the immobilized EF-Tu column.

In vitro translation and trans-translation

Poly (Phe) synthesis was performed by incubating 0.5 pmol of S1-free ribosomes, 1.2 μg of poly (U) (Sigma), 5 pmol of [3H]Phe-tRNAVal, 5 pmol of EF-Tu, and 2 pmol of EF-G in 40 μL of TMNDS buffer containing 20 mM Tris-HCl (pH 7.5), 4 mM MgCl₂, 50 mM NH₄Cl, 1 mM dithiothreitol, 0.2 mM spermine, and 0.5 mM GTP at 65°C. Aliquots were withdrawn, diluted in 4 ml of 5% trichloro acetic acid, and incubated for 10 min at 90°C. The precipitated peptide was recovered by filtration with a mixed cellulose membrane (Advantec). The filter was dried, and the
radioactivity on the filter was counted by a liquid scintillation counter (Alola).

The stalled ribosome was prepared from the reaction mixture of poly (U)-dependent poly (Phe) synthesis as described above. After reaction for 10 min, the mixture was filtered through a Microcon YM-100 (Millipore) and washed with 100 μl of TMNDS buffer four times. The ribosome fraction that remained on the filter was recovered, and the volume was adjusted to half of the poly (Phe) synthesis reaction volume.

The trans-transfer reaction was initiated by adding a 10 μl aliquot of a mixture containing 0.2 μM [3H]Ala-tmRNA, 2 μM SmpB, and 0.5 μM EF-Tu with or without S1 in TMNDS buffer to 40 μl of stalled ribosomes. The reaction mixture was incubated at 65°C and aliquots were withdrawn. The product was detected by the same procedure as that used for the detection of poly (Phe) described above.

The resume codon decoding was performed by mixing and incubating the stalled ribosome (40 μl) and a mixture (10 μl) containing 0.5 μM unlabeled Ala-tmRNA, 0.2 μM [3H]Ala-tmRNA or [3H]Val-tmRNA, 2 μM SmpB, 1 μM EF-Tu, and 0.2 μM EF-G in TMNDS buffer. The product was detected by trichloroacetic precipitation and filtration as described above.

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