In vivo tmRNA protection by SmpB and pre-ribosome binding conformation in solution

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ABSTRACT

TmRNA is an abundant RNA in bacteria with tRNA and mRNA features. It is specialized in trans-translation, a translation rescuing system. We demonstrate that its partner protein SmpB binds the tRNA-like region (TLD) in vivo and chaperones the fold of the TLD-H2 region. We use an original approach combining the observation of tmRNA degradation pathways in a heterologous system, the analysis of the tmRNA digests by MS and NMR, and co-overproduction assays of tmRNA and SmpB. We study the conformation in solution of tmRNA alone or in complex with one SmpB before ribosome binding using SAXS. Our data show that Mg2+ drives compaction of the RNA structure and that, in the absence of Mg2+, SmpB has a similar effect albeit to a lesser extent. Our results show that tmRNA is intrinsically structured in solution with identical topology to that observed on complexes on ribosomes which should facilitate its subsequent recruitment by the 70S ribosome, free or preloaded with one SmpB molecule.

Keywords: tmRNA; SmpB; trans-translation; in vivo protection; SAXS; NMR

INTRODUCTION

With its protein partner SmpB, tmRNA is the major actor of bacterial trans-translation, a quality control mechanism of translation and a ribosome rescuing system. As inferred from its name, tmRNA acts both as a tRNA and an mRNA. Its length is ~350 nt, depending on bacteria. Computational studies show that its tertiary structure is a necklace of several structural blocks (1 MLD and 4 PKs) attached by a linker helix (H2) to a tRNA-like domain (TLD) formed by the pairing of the 5’ and the 3’ ends (Fig. 1) and that the occurrence of each block is extremely well conserved (Burks et al. 2005).

Trans-translation is a multistep process involving tmRNA and SmpB binding to the ribosome. In the early life of tmRNA in the cell, the TLD is charged by the alanyl-tRNA-synthetase (Komine et al. 1994). This mandatory aminoaacylation reaction is enhanced by the presence of SmpB (Hanawa-Suetsugu et al. 2002; Shimizu and Ueda 2002), which suggests that SmpB modulates the TLD structure to enhance recognition by the synthetase. Apart from the prebinding TLD-alanylation step, efforts have been focused on ribosome recognition and on the subsequent steps such as ribosome decoding center hijacking (Nonin-Lecomte et al. 2009; Neubauer et al. 2012). Cryo-EM has provided beautiful and extremely informative pictures of how the [tmRNA·SmpB] complex performs its task, visualizing various stages from pre-accommodation to post-translocation (Valle et al. 2003; Kaur et al. 2006; Cheng et al. 2010; Fu et al. 2010; Weis et al. 2010). Comparatively few papers address the prebinding 3D structure of tmRNA. NMR and X-ray structures of the TLDs of Aquifex aeolicus and Thermus thermophilus confirm the RNA L-shape structure of the TLD (Gaudin et al. 2003; Gutmann et al. 2003; Bessho et al. 2007; Ramrath et al. 2012) and the formation of a very stable pseudoknot formed

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by the PK1 sequence (Nonin–Lecomte et al. 2006). Three-di-
dimensional models were built by Burks et al. (2005). Cryo-EM
structures of partial or full-length tmRNAs bound to the ribo-
some confirm the suspected flexibility of the global confor-
mation, with a 3D structure articulated around the different
structural blocks to go through the various stages of its task.
However, no experimental information on the three-dimen-
sional structure of the full-length molecule before ribosome
binding is available yet.

In this paper, we address the question of whether the 3D
structure of tmRNA is preorganized for ribosome binding
and to what extent the binding of the first SmpB molecule
will modulate this structure. Taken together, biochemistry
and structural studies have evidenced three potential SmpB
binding sites on tmRNA. Wower et al. (2002) showed that
up to three SmpB molecules can crosslink with a large portion
of the TLD-H2 block of E. coli tmRNA. The cryo-EM struc-
ture of Kaur et al. (2006) led the authors to conclude that
two molecules of SmpB can bind the TLD. According to
chemical and enzymatic probing experiments, SmpB can
also establish contacts with the tmRNA large loop in vitro,
in particular with PK1 and PK4 (Metzinger et al. 2005).
This third binding site would be the site of highest affinity
with a $K_d$ of $\sim 3$ nM vs. 10 for the TLD (Metzinger et al.
2008). The present study is focused on the binding of the
first molecule of SmpB. The localization of the binding
site is probed in vivo by original co-production assays of heter-
ologous tmRNAs and proteins. Three tmRNAs were over-
produced in Escherichia coli (Fig. 1), the endogenous E. coli
tmRNA (ec-tmRNA), the heterologous A. aeolicus (aa-
tmRNA), and one chimeric derivative (ec-tmRNA-aaMLD).
TmRNA and SmpB were both produced in E. coli and purified
in native conditions. The tmRNA conformation in solution
before ribosome binding was probed by small angle X-ray
scattering (SAXS). From our experimental data, we could
derive models for the conformation of full-length tmRNA.
We also demonstrate that the binding of a single SmpB mol-
ecule can drive a compaction of the whole RNA structure, an
effect similar to that of high magnesium concentration.

RESULTS

Overproduction of homologous and heterologous
tmRNA in E. coli

The production of full-length ec-tmRNA in E. coli cells was
reported in the literature (Felden et al. 1997). We have cloned
the genes of the 363-nt ec-tmRNA (ec-ssrA) and of the heter-
ologous A. aeolicus 347-nt aa-tmRNA (aa-ssrA) in pBSTNAV
plasmids (Meinnel et al. 1998). Each plasmid was used to
transform JM101tr cells. This plasmid-strain association al-
lows high levels of RNAs to be transcribed under the control
of the constitutive lpp promoter and yields high amounts of
processed full-length ec-tmRNA (Fig. 2A, lane 5). A. aeolicus
is a hyperthermophile bacterium whose optimal growth tem-
perature is $\sim 90^\circ$C (Deckert et al. 1998). Its tmRNA is thus ex-
pected to be highly resistant at 37$^\circ$C. However, despite the
existence of several highly structured elements known to ex-
hibit high-resistance to RNase degradation (TLD, PK1-4),
the gel of the total RNA extracts reveals that aa-tmRNA un-
dergoes discrete and systematic degradation in E. coli (Fig.
2A, lane 3). It exhibits one faint band (F) corresponding to
the expected full-length aa-tmRNA and two strong bands
(D1 and D2) at lower than expected molecular weights (the
in vitro transcript of the aa-tmRNA was loaded on the same
gel as a migration marker). The RNA digests evidenced by

FIGURE 1. Secondary structure models of (A) ec-tmRNA and (B) aa-tmRNA. (Inset) Modified MLD of the armored aa-tmRNA with the MS2 encap-
sidation stem–loop shaded in gray.
196-nt RNA recapitulating aa-tmRNA residues encompassing the PK4-H5 region. We have overproduced a aa-tmRNA-D2 is of which repel RNA strands. According to the gels, the size of strands 8M-urea but are prevented by SDS, the negative charges strands responsible for this effect are strong enough to with-
gregates. The intermolecular interactions between RNA transcribed either in vivo or in vitro forms dimers and/or ag-
gels of total RNA extracts shows that full-length aa-tmRNAparison of Bis Tris (Fig. 2A) and 8M acryl-urea (Fig. 2C)
comparisons of aa-tmRNA-D1 and aa-tmRNA-D2, respectively (D standing for digest). The com-
bands D1 and D2 are hereafter named aa-tmRNA-D1 and aa-
tmRNA-D2, respectively (D standing for digest). The com-
parison of Bis Tris (Fig. 2A) and 8M acryl-urea (Fig. 2C) gels of total RNA extracts shows that full-length aa-tmRNA transcribed either in vivo or in vitro forms dimers and/or ag-
grates. The intermolecular interactions between RNA strands responsible for this effect are strong enough to with-
stand 8M-urea but are prevented by SDS, the negative charges strands of which repel RNA strands. According to the gels, the size of aa-tmRNA-D2 is ~200 nt, which is consistent with a digest encompassing the PK4-H5 region. We have overproduced a 196-nt RNA recapitulating aa-tmRNA residues 97–292 (Fig. 3A, below), i.e., the PK4-H5 block framed by a few flanking nucleotides. The gel of the RNA extract shows evidence of degradation similar to that observed with full-length aa-
tmRNA, with a single species migrating exactly at the same position as aa-tmRNA-D2. This shows that the necklace of pseudoknots and the neighboring helix H5 act as degradation stoppers. The gels also exhibit a faint band at a lower molecular mass, which could arise from further digestion of the flanking nucleotides and H5 helix.

Recombinant ec-tmRNAs and aa-tmRNA-D1 were puri-
ﬁed using the three-step protocol described in the Materials and Methods section. Their respective molecular mass (MM) was determined in native conditions by supra-molec-
ular TOF/mass spectrometry. Ec-tmRNA MM is 117 kDa as expected (Supplemental Fig. 1). Aa-tmRNA-D1 actually corre-
3A, below), i.e., the PK4-H5 block framed by a few flanking nucleotides. The gel of the RNA extract shows evidence of degradation similar to that observed with full-length aa-tmRNA digestion stops at the PK4-H5 block. Together with the mass spectroscopy results, it suggests that in vivo deletion occurs in the MLD. In order to assess the presence approximately corresponds to 25 nt. The two species are not resolved on the gels of RNA extracts, and for simpliﬁcation they will be hereafter considered as one single species (D1). The denaturing ionization of purified species D1 reveals the presence of a single strand with a MM of 80,650 ± 20 Da corresponding to ~250 nt. Taken together, these results show that, despite its migration as a single band on gels, aa-tmRNA-D1 is a mixture of double-stranded species.

**Band shifts experiments**

The experiment illustrated in Figure 3A shows that aa-
tmRNA digestion stops at the PK4-H5 block. Together with the mass spectroscopy results, it suggests that in vivo deletion occurs in the MLD. In order to assess the presence

![FIGURE 2.](image)

(A,B) 4%–12% Bis Tris PAGE. (A) Lanes 1,2: Markers—(1) purified aa-tmRNA (in vitro transcript), (2) purified recombinant ec-tmRNA. Lanes 3–5: Total RNA extracts for overproduction of recombinant tmRNAs—(3,4) total RNA extracts from overproduction of aa-tmRNA, respectively, in the absence and in the presence of SmpB, (5) overproduction of ec-tmRNA. (B) Total RNA extracts from ec-tmRNA-aa-MLD at two different concentrations (lanes 1,2), from ec-tmRNA (lane 3), and aa-tmRNA (lane 4). (C) 8 M urea 12% PAGE. Purified aa-tmRNA transcribed in vitro migrating as a multimer, probably a dimer (lane 1). Lanes 3–6: Total RNA extracts from JM101tr cells transformed with empty pBSTNAV (lane 3), aa-ssrA-pBSTNAV (lanes 4,5), and ec-ssrA-pBSTNAV (lane 6). Bands Dx in lanes 4 and 5 corre-
spond to an oligomeric species and have no equivalent in lane 6.

![FIGURE 3.](image)

(A) (Left) Secondary structure model of the 196-nt RNA encompassing the aa-H5-PK4 region. The corresponding gene (196-
RNA) was inserted in the pBSTNAV plasmid. (Right) Bis Tris 4%–12% PAGE (NuPAGE Novex). Total RNA extracts from JM101tr cells transformed with 196-RNA-pBSTNAV (lane 1), ec-ssrA-pBSTNAV (lane 2), and aa-ssrA-pBSTNAV (lane 3). Lane 1 exhibits a strong band similar to band D2 of lane 3. (B,C) Hybridization experiments on full-RNA extracts from aa-tmRNA overproduction in JM101tr cells (B) and on a partially purified mixture of full-length aa-tmRNA and aa-
tmRNA-D1 (C). Lane 1: extracts loaded at room temperature; lanes 2–4: extracts loaded after a heat–cool cycle, alone (lane 2) and in the presence of antiH2PK1 and antiTLD (respectively, lanes 3 and 4); lane 5: RNA marker of 70 nt; lanes 6–9: same conditions, respectively, as lanes 1–4 but on the purified mixture.
of the TLD and of the H2-PK1 regions, we conducted hybridization experiments that we monitored by band shift observations on 12% SDS PAGE gels (Fig. 3B,C). Two DNA oligonucleotides, anti-H2PK1 (15 nt, designed to hybridize nucleotides 39–54 of Fig. 1B) and anti-TLD (35 nt, designed to hybridize nucleotides 329–347 of Fig. 1B), were used as probes. The experiments were conducted either on RNA extracts (Fig. 3B) or on a partially purified mixture of full-length aa-tmRNA and aa-tmRNA-D1 (Fig. 3C). Upon heating, the intensity of band D1 disappears, while that of bands D1′ and D1″ increases (lanes 2 and 7). Migration of band D1 matches the 250-nt species observed at ∼80 kDa in MS spectroscopy. Band D1″ is consistent with the 75-nt species expected after MLD digestion and corresponding to the 5′ side of the TLD-H2-PK1 region of aa-tmRNA. Hybridization with anti-TLD shifts bands F, D1, and D1′ (lanes 4 and 9) but not D2 and D1″ as expected. This shows that the TLD is retained in aa-tmRNA-D1. Hybridization with anti-H2PK1 shifts bands F, D1, and D1″ (lanes 3 and 8) but not D1′ and D2. This shows that the D1 and its shorter component are cleaved in the MLD region, after at least half of the PK1 sequence. Altogether, this also shows that the deletion of 25 nt occurs between PK1 and H5.

**NMR analysis of aa-tmRNA-D1**

The TLD block results from the association of the 3′ and 5′ ends (Fig. 1) and its folding topology has much in common with tRNAs (12–15). The previous hybridization experiment shows that aa-tmRNA-D1 retains the 3′-side of the acceptor arm of the TLD. The TLD undergoes a single post-transcriptional methylation in the TΨC loop, on the 3′-side of the molecule (Felden et al. 1998). We checked for its presence by NMR. The presence of a methyl group transcribes into methylation in the TΨC base pairs attesting to the formation of a stable 3D structure. The region of aromatic proton resonances (between 6.5 and 8.5 ppm) displays the strong and single cross-peak at 7.47 and 1.45 ppm, as similarly observed in the aa-TLD (Gaudin et al. 2003) and the ec-tmRNA (Ranaei-Siadat et al. 2013) NOESY spectra. The presence of this methyl resonance demonstrates the presence of the TLD T-arm and T-loop in aa-tmRNA-D1, and thus of the 3′ end nucleotides of full-length aa-tmRNA. Considering the high resistance of tRNAs to RNases and our ability to overproduce the aa-TLD alone using the same plasmid and *E. coli* strain (Gaudin et al. 2003), we can conclude that aa-tmRNA-D1 contains the full TLD part. The observed mass default thus originates from an endonucleolytic digestion and is consistent with the deletion of the MLD region, and in turn, the longer strand of aa-tmRNA-D1 species would correspond to the 3′ end part of aa-tmRNA (H5-3′ end region).

**aa-MLD ec-tmRNA chimera**

To probe the weakness of the MLD, the region coding for the ec-MLD (residues 79–108) was replaced in ec-ssrA by the corresponding aa-MLD sequences (residues 70–101) with the view to recreate a substrate for degradation. JM101tr bacteria were transformed with the ec-ssrAaaMLD-pBSTNAV plasmid. The pattern of the corresponding total RNA extracts (Fig. 2B, lanes 1,2) reveals that the MLD substitution entirely destabilizes the ec-tmRNA. The faint band at 363 nt observed in the overloaded lane most probably corresponds to the endogenous production of wild-type ec-tmRNA. Modifications of the MLD were used to engineer functional trans-translation tests (Tanner et al. 2006) and were shown to be compatible with the integrity of tmRNA. In light of this result, the complete degradation of the aa-MLD ec-tmRNA chimera is totally unexpected. However, it brings evidence that the aa-MLD is a target for degradation in *E. coli* as inferred from the aforementioned MS and NMR analyses of aa-tmRNA-D1.
NMR analysis of the MLD

Aa-tmRNA-D1 is a by-product of the endonucleolytic degradation of aa-tmRNA. Because tRNA-like structures and pseudoknots are highly resistant, we suspected the aa-tmRNA initial degradation to take place within its least folded part, around the MLD. In 2D models based on chemical and enzymatic probing experiments, the MLD region appears as a stem–loop framed by unstructured single strands. The *E. coli* MLD (ec-MLD) contains a stem–loop of 4 bp (Felden et al. 1997) and that of *A. aeolicus* (aa-MLD) is a little more structured, with 6 bp (Metzinger et al. 2005). We probed by NMR the structures in solution of 22-mer and 29-mer oligonucleotides recapitulating aa- and ec-MLDs (respectively, 5′-rAGCUCCCGAAGCUGAACUCGCU and 5′-rAAAAAA UAGUCCGAAACGCGAAGACUAC).

The 1D and 2D-NOESY 1H spectra recorded in water are consistent with MFOLD structure predictions (Zuker 2003) and confirm the poor structuring of the MLD from the two species, both of which adopt small stem–loop structures (Supplemental Figs. 2, 3). In particular, the aa-MLD contains three G·C and two A·U Watson-Crick base pairs. It is less structured than modeled from the probing experiments.

Co-overproduction of tmRNA with either SmpB or His6-MS2 coat protein

In a homologous system (ec-SmpB and ec-tmRNA in *E. coli* cells), SmpB enhances tmRNA lifetime and stability in the cells and allows recovery of higher tmRNA amounts than when the RNA is produced alone (Hanawa-Suetsugu et al. 2002; Hong et al. 2005). With the idea that SmpB may protect aa-tmRNA from degradation, we decided to co-overproduce the *A. aeolicus* SmpB (aa-SmpB) and aa-tmRNA in LB medium. Both genes were cloned in our homemade plasmid pProRNA (Ponchon et al. 2013). RNA transcription is constitutively controlled by the *ppp* promoter, while that of SmpB is controlled by the inducible *lac* promoter. The cells were grown in LB growth medium. Digest D2 disappears from the total RNA extracts (Fig. 2A, lane 4). When production is carried out in rich growth medium (2XTY), a faint band reappears at position D2, probably because the relative proportion of the aa-tmRNA transcript is increased compared to that of SmpB (data not shown). This shows that SmpB blocks the degradation pathway(s) leading to aa-tmRNA-D2 and that the disappearance of band D2 is not a consequence of the inactivation by SmpB of the RNase(s) involved in the degradation pathway. The presence of band D1 shows that aa-SmpB does not fully protect aa-tmRNA from endonucleolytic degradation.

We recently reported the co-overproduction of the armored version of aa-tmRNA (Ar-aa-mRNA) with the MS2-His6 coat protein (138 aa) using the homemade p44K plasmid (Ponchon et al. 2013). The Ar-aa-MLD was modified compared to wild type to incorporate, instead of the short stem–loop of the aa-MLD, the 23-nt-long encapsidation signal stem–loop of the MS2 coat protein (Fig. 1B, inset). This replacement does not protect the MLD from degradation (Fig. 5). The His6-MS2 coat protein binds the RNA as a dimer and, contrary to aa-SmpB, totally blocks the degradation pathway D1 of Ar-aa-tmRNA (Fig. 5, lanes 4–6). As expected, the His6-MS2 coat protein does not block the degradation pathway D1 of aa-tmRNA, which lacks the encapsidation signal (Fig. 5, lane 3). If SmpB were bound to the large loop close to the MLD, we would expect a gel pattern close to that observed with the MS2 coat protein. The comparison of the two coproduction experiments thus led us to the conclusion that, when co-overproduced in vivo with aa-tmRNA, aa-SmpB preferentially binds to the TLD.

Structural investigations of ec-tmRNA and ec-tmRNA complexes using SEC-SAXS

We aimed at obtaining structural data on tmRNA in solution before ribosome binding in the presence or absence of SmpB. The SmpB sequence is extremely conserved across species (Dong et al. 2002; Someya et al. 2003), and this translates into the fact that a heterologous SmpB can efficiently be active and complement the deletion of the gene of the endogen protein (Thibonnier et al. 2010). In vitro, the heterologous system can perform *trans*-translation, albeit less efficiently (Ito et al. 2002). Wower et al. (2002) have shown that *A. aeolicus* SmpB can bind in vitro full-length and truncated ec-tmRNA. As explained in Materials and Methods, our choice finally turned to the heterologous system composed of ec-tmRNA and aa-SmpB because atomic coordinates were available for both molecules (Dong et al. 2002; Ramrath et al. 2012). Data were recorded on tmRNA samples alone and in the presence of one molar equivalent of SmpB protein, both in the presence of 5 mM Mg2+ and in its absence. All samples were loaded onto the on-line SE-HPLC column, and SAXS patterns were recorded along the elution profile. All profiles exhibited a

*FIGURE 5.* His6-MS2 coat protein protects MLD from degradation in vivo. Total RNA extracts loaded onto a 10% SDS PAGE (UV shadowing); All RNAs were produced in p44K. (−) No protein coproduction, (+) His6-MS2 coat protein production. Lane 1: aa-tmRNA; lane 2: Ar-aa-tmRNA; lane 3: as a control, aa-tmRNA coproduced with the His6-MS2 coat protein; lanes 4–6: Ar-aa-tmRNA coproduced with the His6-MS2 coat protein; three clones tested. The degradation pathway leading to D1 was suppressed by the coproduction of the protein; lane 7: partially purified ec-tmRNA as marker.
minor fraction of oligomer and a major monomer peak. Scattering patterns of tmRNA alone or in complex with SmpB are shown in Figure 6, in panel A1 without Mg\textsuperscript{2+} and in panel A2 with Mg\textsuperscript{2+}. Structural parameters derived from each curve are shown in Table 1. Guinier plots are shown in insets to Figure 6, panels A1 and A2, while \( p(r) \) functions, which represent histograms of intra-particle distances, are presented in Figure 6, panels A3 and A4.

The comparison of data and parameters directly shows that Mg\textsuperscript{2+} ions cause a significant global conformational change of the molecule to a more compact conformation (the radius of gyration \( R_g \) decreases by \( \sim 15\% \) and the maximum diameter \( D_{\text{max}} \) by \( \sim 25\% \)). This is also true for the complexes of tmRNA with SmpB. Similar conformational changes are observed upon SmpB addition in the absence of Mg\textsuperscript{2+} ions, though of reduced amplitude, with \( \sim 7\% \) reduction in \( R_g \) and almost 20\% in \( D_{\text{max}} \). As expected for solutions eluting from a SE column, molecular mass estimates derived from the intensity at the origin are close to the values calculated from tmRNA and SmpB sequences. This is further confirmed by the values of the hydrated molecular volume derived from the Porod expression, all of which are compatible with a monomer of tmRNA (respectively, complex with SmpB).

**Rigid-body modeling**

TmRNA is a highly structured molecule comprising eight modules. We, therefore, decided to use a rigid-body modeling approach as implemented in the SASREF program (Petoukhov and Svergun 2005) using the components of the pseudo-atomic model proposed by Ramrath et al. (2012) as defined in Table 2 and explained in detail in the molecular modeling section of Materials and Methods. In order to avoid any bias in the computation from their starting spatial localization, all eight modules were superimposed at the origin.

For each sample, 300 models were obtained that fitted the experimental data within experimental uncertainties. The
most striking observation is the unique arrangement of modules, with the pseudoknots, the MLD and H5 arranged in a large loop, while H2 acts like a hinge with the TLD pointing away from the loop as shown in Figure 6B, panel B1 that presents the direct result of one rigid body refinement in the case of tmRNA alone in the presence of Mg2+ ions. The same model is displayed in panel B2 after restoration of backbone connectivity, while the corresponding scattering patterns are superimposed over experimental data in panel B3. Reduced residual plots are also presented and attest to the quality of the model. This is very reminiscent of the model extracted from the ribosome-tmRNA-SmpB complex and is remarkable in view of the starting point for refinement in which all domains are superimposed at the origin. Under all conditions, models are found in which the H2-TLD “arm” is found on either side of the loop, and models with the correct orientation were selected before further analysis (see Materials and Methods for details).

Statistical analysis of tmRNA structural parameters

From a mere visual inspection of models, the loop domains do not exhibit major rearrangements in the presence of Mg2+ that could explain the more compact overall shape suggested by the structural parameters $R_g$ and $D_{max}$ (Table 1). To try and detect some conformational differences that could be correlated with the observed differences in $R_g$ and $D_{max}$ values, we undertook a few systematic calculations on all models. To characterize the shape and size of the loop, we calculated the distribution of distances between the center of mass of the loop and all atoms of the loop domains. All distributions corresponding to the same sample were then averaged, and the averages are plotted in Supplemental Figure 4A, panel A1. While the curves derived from models in the absence of Mg2+ are asymmetric, corresponding to an anisometric shape of the loop, those in the presence of Mg2+ appear to be distinctly more symmetric. Furthermore, the root mean square distance undergoes a clear reduction from 50.4 Å to 46.1 Å in the presence of Mg2+. In the absence of Mg2+, SmpB addition shifts the distribution toward smaller values by $\sim$1 Å without changing its shape. This explains, at least in part, the observed reduction in the radius of gyration of the molecule upon SmpB and Mg2+ addition. We also calculated the distribution of the distances of all atoms of the loop to the “mid-plane” (defined by the first two inertia axes of the loop) averaged over all models for a given condition (Supplemental Fig. 4A, panel A2). In the presence of Mg2+ ions, distributions are narrower around zero than in the absence of Mg2+, showing that the loop exhibits excursions of reduced amplitude normal to the mid-plane. In conclusion, the loop appears to be more plane and isometric in the presence of Mg2+.

We then turned to investigate the orientation of the TLD with respect to the loop using two parameters determined with respect to the mid-plane of the loop. First, $\alpha$ is the angle between the vector linking the center of mass of the loop O and the projection on the mid-plane M of the hinge point of the H2 domain with the loop (phosphorus atom 304) and the vector MP where P is the projection on the mid-plane of phosphorus atom 363 at the far end of the TLD (see diagram in Supplemental Fig. 4A, panel A3). Second, Z is the Z-coordinate of phosphorus 363 that reports about the elevation of the TLD. The mean values of each parameter with associated standard deviations are plotted in Supplemental

| TABLE 1. Parameter values obtained from the analysis of SAXS scattering curves |
|---------------------------------|----------------|----------------|----------------|
| $R_g$ Guinier (Å) | MW seq (kDa) | MW l(0)/c (kDa) | $R_g$, p(r) (Å) | $D_{max}$ (Å) |
| No Mg2+ | tmRNA | 64.8 ± 0.5 | 117 | 120 ± 10 | 63.5 ± 0.5 | 220 |
| 1-1 complex | 60.8 ± 0.5 | 132 | 138 ± 10 | 58.9 ± 0.5 | 180 |
| 5 mM Mg2+ | tmRNA | 55.4 ± 0.5 | 117 | 130 ± 10 | 54.6 ± 0.5 | 160 |
| 1-1 complex | 55.5 ± 0.5 | 132 | 149 ± 10 | 55.8 ± 0.5 | 160 |

| TABLE 2. Structural rigid modules used in tmRNA 3D structure reconstruction in SASREF |
|---------------------------------|----------------|----------------|----------------|
| Eight rigid modules | Boundaries | Three rigid modules | Boundaries | Four rigid modules | Boundaries |
| 1 | TLD + H2a (+SmpB)* | 1–42 311–363 | 1 | TLD + H2a (+SmpB)* | 1–42 311–363 | 1 | TLD + H2a b (+SmpB)* | 1–48 301–363 |
| 2 | H2b | 43–48 301–310 | 2 | H2b | 43–48 301–310 | 2 | PK1 | 49–300 |
| 3 | PK1 | 49–83 | 3 | PK1 | 49–300 | 3 | MLD | 84–104 |
| 4 | MLD | 84–104 | MLD | 84–104 | H5 | 105–137 |
| 5 | H5 | 105–137 | H5 | 105–137 | PK2 | 105–300 |
| 6 | PK2 | 138–198 | PK2 | 138–198 | PK3 | 138–198 |
| 7 | PK3 | 199–246 | PK3 | 199–246 | PK4 | 199–246 |
| 8 | PK4 | 247–300 | PK4 | 247–300 | |

*SmpB from 1P6V has been positioned in interaction with the TLD as observed in the crystal structure (PDB: 1P6V) (Gutmann et al. 2003).
Figure 4, panels A4 and A5, α exhibits a marked decrease from ~110° down to ~60° upon Mg²⁺ addition, while, in contrast, the distance of the end point of TLD to the mid-plane increases from an average value of 80 Å to an average distance of 100 Å. Therefore, the TLD appears to be positioned more directly above the center of the loop (smaller α) and with its extremity higher above the mid-plane (higher Z-value).

The existence of correlations between α and Z values was investigated in a coarse-grained manner using the heat maps shown in Supplemental Figure 4B. In the absence of Mg²⁺, Z and α exhibit essentially monomodal distributions peaking around 70 Å and 130°, respectively, with more than half of all models comprised within the gray box [40 Å, 100 Å]₀ and [100°–160°]₀ (Supplemental Fig. 4B, upper panel). In contrast, in the presence of Mg²⁺, a rectangular corner-shaped distribution is observed around a square center [100 Å, 140 Å]₀ and [40°–80°]₀ with a horizontal extension [120 Å, 140 Å]₀ extending down to 0° for α and a vertical one [60°–80°]₀ down to 40 Å for Z (Supplemental Fig. 4B, lower panel). The gray box contains two thirds of all models.

Clustering offers another way to organize the ensemble of solutions; they were grouped in similarity clusters based on RMSD calculations using the routine Damclust (Petoukhov et al. 2012). In the absence of Mg²⁺, 10 clusters are identified, five of which account for 75% of all solutions (the location of each centroid is represented as a star in Supplemental Fig. 4B, upper panel). In the presence of Mg²⁺, four main clusters out of nine account for 78% of all models (Supplemental Fig. 4B, lower panel). The distributions of centroids show a good overlap with both heat maps.

The conclusion of this coarse-grained analysis of our models is that the addition of Mg²⁺ to the solution of tmRNA causes a small but clear compaction of the loop, which becomes flatter and more isometric together with a movement of the TLD centered above the loop and mainly away from the mid-plane of the loop.

Comparison with cryo-EM-derived models

In a complementary approach to the previous bias-free, rigid-body analysis, we tried to evaluate existing models derived from cryo-EM studies of tmRNA and SmpB complexed with ribosomes and other factors representing different steps of the tmRNA rescue task: pre-accommodation (Kaur et al. 2006), accommodation (Weis et al. 2010), and post-translocation (Fig. 6C, panel C1; Ramrath et al. 2012). Scattering patterns were calculated and are shown in Figure 6C, panel C2. No model provides even a decent fit to our experimental data on tmRNA in 5 mM Mg²⁺ containing buffer (the cryo-EM samples contained Mg²⁺ ions) with χ values comprised between 17.5 and 21. Thus, although the topology and overall shape of tmRNA is basically unchanged between the solution state and the complex with ribosome, tmRNA undergoes large amplitude conformational changes upon ribosome binding and through the various stages of its task.

Cryo-EM data suggest that major tmRNA conformational rearrangements during trans-translation steps on the ribosome involve changes in the relative orientation of the TLD-H2 with respect to the big loop. A second trial was thus conducted with only three rigid modules, the TLD, H2, and the loop, which was “frozen” in the conformation of the post-translocation step (PDB 3J18; 12). The fits between the computed and the experimental data recorded with Mg²⁺ were improved compared to the initial models but fell short of an acceptable fit, with χ values on the order of 5.

According to NMR, the MLD solution structure composed of a small stem–loop framed by two single strands is much less constrained that those of the pseudoknots. In a third step, we, therefore, released the MLD module from both PK1 and H5 and used four rigid modules (TLD-H2, PK1, MLD, H5-PK4). The fit between the model calculated curves and the experimental data recorded in Mg²⁺ keeps improving with the increase in the number of degrees of freedom (χ values between 1.6 and 2). Interestingly, the (α, Z) distribution analysis shows that most solutions are grouped into the low α – low Z quarter of the map, in marked contrast to that observed on the models obtained starting with all modules at the origin (Supplemental Fig. 4). Keeping all four PKs in the position found in the model proposed by Ramrath et al. (2012) constrains the H2-TLD module to adopt a different set of positions as characterized by the (α, Z) distribution that, besides, lead to fits of lower quality. This suggests that the PKs are arranged differently in solution, which is not unexpected as the conformation of the loop in the cryo-EM model results from its numerous interactions with various parts of the bound ribosome, some of which are actually located inside the loop.

DISCUSSION

Origin of the systematic aa-tmRNA degradation in E. coli cells

How and why does aa-tmRNA, composed of very stable structural blocks that should resist to temperatures up to 95°C, undergo systematic and reproducible degradation in E. coli? The precise identification of the set of RNases responsible for our patterns of degradation would require testing the overproduction of aa-tmRNA in several E. coli strains deleted for several RNase genes. This ongoing work is beyond the scope of this study.

In the first place, we can imagine that degradation leading to aa-tmRNA-D2 can arise from multiple pathways, including degradation from ill-folded conformers, degradation from aa-tmRNA-D1, or degradation from abortive transcripts followed by digestion of the sequence upstream of H5 and downstream from PK4. Coproduction assays demonstrate that SmpB prevents formation of aa-tmRNA-D2, so the last hypothesis is valid if SmpB protects the neosynthetized single strand during transcription. For reasons that are exposed...
hereafter, we suspect that aa-tmRNA-D2 mostly originates from full transcripts ill-folded in the TLD region, which would destabilize and expose parts of the RNA that are naturally protected from the action of RNAses in well-folded molecules. Despite its heterologous origin, the A. aeolicus TLD (aa-TLD) can be overproduced in high quantity using the same plasmid and E. coli strain (Gaudin et al. 2003), showing that it is mostly well-folded and not degraded. In contrast, we observed that no production of a longer aa-tmRNA derivative mimicking both the aa-TLD and the connecting stem H2 could be obtained. The H2 sequence itself and the proper association of its two strands would thus be critical for proper TLD folding and vice versa. The GC content of the aa-ssrA gene is higher than that of E. coli, in particular in the TLD arms and in helix H2. It has evolved to favor active 3D folds at high temperature. It is thus not surprising that, at 37°C, a number of competing alternate structures can be produced in high amounts and be readily degraded by the host RNases. Such structures are not recognized as tRNAs and may readily undergo degradation by the host RNases. Degradation of aa-tmRNA conformers, either full-length or ΔMLD, with misfolded TLD, can thus explain band D2. This observation is also reminiscent of the degradation of the unprotected 3′ end of the coding strand in Caulobacter crescentus two-pieces tmRNA by RNase R (Hong et al. 2005). Such band is not observed in the case of E. coli either because there is no ill-folded ec-tmRNA molecule in E. coli, as suggested by Hong et al., or because ill-folded conformers exist, but E. coli pseudoknots are not strong enough to resist degradation by endogenous RNases. Recently, Lian and Deutscher (2012) showed that SmpB regulates the RNase R turnover and that the efficiency of the regulation is enhanced when SmpB is complexed to tmRNA. We are currently investigating the role of RNase R in aa-tmRNA degradation.

The origin of band D1 was, at first glance, more intriguing. PAGE and MS analyses provide evidence for a degradation product resulting from endonucleolytic cleavages and lacking the MLD region. The presence of the 3′-end region of aa-tmRNA is revealed by the methyl resonance on the NOESY spectrum. In E. coli, most RNAs are digested in a stepwise fashion, starting with an internal endonucleolytic cleavage followed by exonucleolytic digestion. The observation of aa-tmRNA-D1 is consistent with such a degradation pathway. RNase levels in the cell can be up-regulated as a result of the cellular stress caused by overproduction of a heterologous tmRNA. The endotoxin MazF that targets single-stranded ACA triplets (Zhang et al. 2003) is a potential candidate for the initial endonucleolytic cut of the MLD region. It was shown to degrade successfully mutant ec-tmRNA bearing ACA triplets (Baik et al. 2009). Our gels show that a slight proportion of full-length tmRNA is recovered, probably because, in the cell, oligomeric association protects the MLD from degradation. As a conclusion, the unexpected observation of tmRNA digests D1 and D2 was made possible because of the exceptional stability of the blocks composing the aa-tmRNA. We suspect that similar degradation pathways might exist for every heterologous tmRNA and also probably for the ec-tmRNA and that these pathways are important for the quality control of tmRNA structure in vivo and maybe for regulation of the number of functional tmRNA molecules in the cell. Heterologous production of tmRNA in E. coli thus provides a new tool to study these degradation pathways.

SmpB binding site of highest affinity from in vivo protection

The observation and characterization of aa-tmRNA-D1 and aa-tmRNA-D2 show that RNases stop when encountering strong structural blocks. These digests can be used as original tools to probe in vivo the localization of the first SmpB binding event onto tmRNA. Endogenous SmpB and tmRNA are produced in vivo in similar amounts (Sundermeier and Karzai 2007). This suggests that in vivo, outside the ribosome, most free tmRNA molecules will associate with a single SmpB. UV shadowing is quantitative and discloses a slight increase of the relative proportion of full-length aa-tmRNA with respect to digest D1 (Fig. 2). This suggests that SmpB converts ill-folded aa-tmRNAs into good tRNA-like structures. SmpB binding does not protect aa-tmRNA from the endonucleolytic degradation leading to D1 digest as the His6-MS2 coat protein dimer does (Fig. 5). SmpB is slightly larger than one monomer of the His6-MS2 coat protein dimer (157 aa vs. 2 × 138 aa). Steric hindrance resulting from SmpB binding in the PK1-MLD region would thus be expected to prevent MLD from endonuclease attacks. This is not the case. The RNA extract pattern of the coproduction assay with SmpB is thus fully understood if we suppose that TLD binds as observed in the crystallographic structures (Gutmann et al. 2003; Bessho et al. 2007; Ramrath et al. 2012). The absence of degradation from alternative wrong folds confirms the assumption that SmpB exerts a chaperoning action on TLD folding and provides a structural explanation to the observed enhanced aminocaylation in the presence of SmpB. Former probing experiments established the existence of at least two SmpB binding sites, one on the TLD and the other one within the necklace formed by the PKs and the MLD (Wower et al. 2002; Konno et al. 2007). The respective dissociation constants measured on tmRNA fragments recapitulating the two binding sites are ~40 nM and 2 nM (Metzinger et al. 2008). This apparent inconsistency with our in vivo protection results and with a recent study reporting a Kd of 0.4 nM in 100 mM Na⁺ and 2.6 nM in 10 mM Mg⁺⁺ on a synthetic two-stranded TLD (Fahrat and Rueda 2012) could arise from the nature of the RNA samples under investigation. The former full-length aa-tmRNA was transcribed in vitro, a method conveniently avoiding RNases attacks, and purified in denaturing conditions. The resulting tmRNA thus lacks the two post-transcriptional modifications (T and Ψ) that stabilize the TLD structure. The affinity of SmpB for the TLD of full-length tmRNA in vivo is probably better than in the in vitro
context. The second drawback of the in vitro RNA production is intrinsically linked to the length of the RNA. The neosynthesized RNA folds as transcription proceeds. In vitro, the aforementioned quality control of the structure does not exist, and ill-folded tmRNAs are not degraded. The in vitro tmRNA sample is, thus, certainly much more conformationally heterogeneous than its in vivo counterpart purified in native conditions. Our gel filtrations indicated the existence of a mixture of aggregates, dimers, and a slight proportion of monomers, even after thermal treatment similar to that used by Metzinger et al. (2008). The $K_d$ values reported were most probably measured on a heterogeneous population of tmRNA conformers and oligomers. Konno et al. (2007) have shown that the C-ter tail-truncated SmpB protects U85 of ec-tmRNA from chemical modification, i.e., in a region consistent with the binding site of highest affinity observed in vitro (Metzinger et al. 2008). However, the ec-tmRNA sample was subjected to isopropyl precipitation, and in such conditions, we were not able to get monodispersed sample. According to our experimental models, the SmpB molecule bound to the TLD is too far away from U85 to interact directly. If this molecule effectively conferred protection, then it must be indirect. However, we cannot rule out that a small proportion of SmpB binds in vivo to the tmRNA loop, which would be hidden by the major fraction binding TLD.

It has been proposed that SmpB and protein S1 would act in concert as a scaffold for tmRNA (Gillet et al. 2007; Felden and Gillet 2011). A preferential binding of SmpB on TLD leaves open the access of the tmRNA loop for S1 binding (Wower et al. 2000). Also, one SmpB molecule can bind the empty decoding site of free 70S ribosome in the absence of tmRNA in vivo (Gillet et al. 2007). The authors proposed that, depending on environmental conditions, several pathways of tmRNA recruitment are likely to initiate trans-translation (Felden and Gillet 2011). The most recent cryo-EM structure of the pre-accommodation state shows a TLD complexed to two SmpB molecules binding the GTPase-associated center of the 50S and the decoding center of the 30S (Kaur et al. 2006), respectively. Our finding that SmpB binds preferentially the TLD in vivo is consistent both with the most recent crystal structure of the pre-accommodation state which shows the TLD complexed to a single SmpB molecule that also binds the decoding center of the 30S (Neubauer et al. 2012) and with the existence of an alternative recruitment pathway of tmRNA by a 70S ribosome preloaded with one SmpB in the decoding site, as first suggested by Hallier et al. (2004).

**Structural insights into the [tmRNA-SmpB] complex**

Despite the existence of alternative recruitment pathways (Felden and Gillet 2011), several data suggest that tmRNA is most likely recruited by ribosome as a complex. Our goal was to shed some light on the conformation of both tmRNA and [tmRNA-SmpB] (1:1) complex in solution before ribosome binding. It is not possible to discriminate between the two SmpB binding sites on the basis of the SAXS data alone because of the too small contribution of the protein to the total scattered intensity. For our SAXS modeling, SmpB was positioned in interaction with the TLD as found in the crystal structure of the complex (Bessho et al. 2007) and in accordance with our in vivo probing. The structural parameters measured on [tmRNA-SmpB] complexes by SEC-SAXS show that a significant compaction of the molecule occurs upon SmpB binding (Table 1). We computed rigid-body models of tmRNA and [tmRNA-SmpB] complex using the eight modules (Table 2) used by Ramrath et al. (2012) in the presence or in the absence of Mg$^{2+}$. All resulting models that yield an excellent match with the experimental data exhibit the loop made of the pseudoknots and the MLD, connected to the TLD-H2 region which points outward. The presence of Mg$^{2+}$ ions is associated with a more circular and flatter loop conformation together with a TLD arm rising higher above the loop.

In contrast, the scattering curves computed from the cryo-EM structures of the tmRNA at various steps of trans-translation display major discrepancies with the experimental data (Fig. 6C, panel C2). This shows that, although the topology appears to be preserved, the tmRNA conformation before ribosome binding is different from that in all studied ribosome-bound states. In particular, our models suggest that the large loop adopts a more compact conformation in the free state than in complex with the ribosome. Although the tightly folded conformation proposed by Ivanov et al. (2002) is not supported by our data, the basic idea of a more compact conformation in isolation, also suggested by Burks et al. (2005), holds true but in a more subtle way, the various conformations explored by the tmRNA in solution and later in the different stages of its activity appearing more like variations on a common topological theme.

**CONCLUSION**

Our results provide experimental information on the conformation of [tmRNA-SmpB] complex in solution before ribosome binding. TmRNA appears to be self-structured, even as a free molecule, with a characteristic topology that makes it mechanically easy to bind ribosome and that can be modulated through its interactions with various parts of the ribosome and associated factors. We showed by original in vivo protection assays that the first SmpB preferentially binds to the TLD and that the binding causes a significant compaction of the entire tmRNA in the absence of Mg$^{2+}$ ions. The study of the degradation pattern of heterologous RNAs that evolved to withstand high temperature thus constitutes an original and valuable tool to monitor the relative stability of the different parts of the molecules in vivo and to get the footprints of protein partners in vivo.

Our overproduction assays show that the self-structuration of tmRNA explains its high resistance to RNases and thereby its exceptional longevity in a functional state within the cell.
By overproducing the heterologous aa-tmRNA in *E. coli*, we have, in a way, mimicked drastic modifications of the selection pressure. The degradation of the heterologous aa-tmRNA also demonstrates that tmRNA sequence was finely tuned by evolution to ensure its stability, even in the absence of any protein partner. However, this does not mean that SmpB has no stabilizing or chaperoning effect on tmRNA structure. Indeed, it was formerly observed that the overproduction of tmRNA with its protein partner SmpB allows the recovery of higher amounts of full-length aa-tmRNA (Hallier et al. 2004). SmpB is known to enhance TLD aminoacylation (Barends et al. 2001; Shimizu and Ueda 2002). Hanawa-Suetsugu et al. (2002) suggested that it proceeds by modulating TLD conformation to enhance recognition by AlaRS. The authors also showed that the protein protects nonaminoacylatable mutant TLD from degradation in vivo. The disappearance of the aa-tmRNA-D2 species resulting from our co-overproduction assays brings an original piece of evidence that SmpB chaperones the TLD structure.

**MATERIALS AND METHODS**

**Expression tests and RNA recovery**

Either an XL1 or JM101tr *E. coli* strain was used in all expression tests. Expression tests for total RNA extracts recovery were performed in 20 ml LB medium. Cells were pelleted and resuspended in 400 μL of 10 mM magnesium acetate, 10 mM Tris- HCl, pH 7.4. Protein-free total RNA extracts were recovered after phenol/chloroform extraction and ethanol precipitation as described before (Ponchon et al. 2009). They were analyzed by SDS-PAGE electrophoresis. Bands were visualized by UV shadowing.

**RNA production and purification**

Cells transformed with pBSTNAV, pProRNA, or p44K plasmids were grown in either LB or 2XTY growth medium and harvested as described before (Ponchon et al. 2013). The RNAs were recovered following protocols described by Ponchon et al. (2009). The aqueous phase was purified in four steps by FPLC using nondenaturing buffers. tRNA samples were first separated from the bulk of tRNAs and SS RNA on a source 15Q (GE Healthcare) pre-equilibrated in H2O, potassium phosphate pH 6.5, 20 mM, and EDTA 1 mM using a 50 mM–1 M NaCl elution gradient. Purity was further refined on a more resolutive mono Q column using the same loading and elution buffers and a 200–300 mM NaCl gradient. Last contaminants were eliminated using a hydrophobic column. A final round of gel filtration (Superdex 200, Amersham) was used to remove aggregates prior to NMR and SAXS experiments. Analytical gel filtrations were performed at 17°C using an analytical Superdex 200 column and 150 mM NaCl, 20 mM Tris HCl, pH 6.5 elution buffer with (1 mM) or without MgCl2 added. RNAs extracts and purification fractions were analyzed by either SDS or 8 M-urea PAGE (10% or 12%). RNA bands were visualized by UV shadowing (Ponchon et al. 2009). SDS-PAGE provides good resolution in the range of our RNAs of interest (60–400 nt), we sometimes used 4%–12% gradient Bis Tris gels (NuPAGE Novex) with NuPAGE MES SDS running buffer and NuPAGE LDS sample buffer.

**Hybridization experiments**

Aa-tmRNA fragments were heated for 2 min at 98°C in water with 2 μL of a 100-μM DNA oligonucleotides anti-H2PK1 (15 nt, 5′-ACCCGGCCTGCTCCT) or anti-TLD (35 nt, 5′-AAATAACCTTC TGCAATGGTGAGCGGCGGAAT) water solution for final volumes of 15 μL and cooled down to 4°C in 30 sec. Only the last 19 nt of the anti-TLD are complementary to nucleotides 329 to 347 of aa-tmRNA. Band shifts were assayed on 12% SDS PAGE. Samples were loaded on the gels at room temperature.

**NMR**

The purified aa-tmRNA-D1 was deacylated in Tris buffer, pH 8 at 37°C for 1 h and then dialyzed several times against 1 mM EDTA water solutions and finally extensively against water. Synthetic aa-tRNA and ec-tRNA oligonucleotides were purchased from Dharmacon Inc. The unprotection buffer was exchanged against water on G25 desalting column (GE Healthcare). All samples were freeze-dried before resuspension in 95% H2O–5% D2O and 0.01 mM EDTA. The pH was adjusted to 6.5 by addition of small aliquots of 0.1N NaOH under stirring. The purified aa-tmRNA-D1 strand concentration was 0.1 mM. Synthetic oligonucleotide concentrations range from 0.2 to 1 mM. All NMR spectra were acquired in 5 mm Shigemi tubes at 600 MHz on a Bruker Avance DRX600 spectrometer equipped with a cryoprobe and Z-axis gradient. Water signal in 1D and 2D NOESY experiments was either nonexcited using the Jump and Return (jr) procedure (Guéron et al. 1991) or suppressed using the W5 Watergate (wg) pulse sequence (Liu et al. 1998). The resonances were assigned using the usual set of 2D homonuclear NMR methods in D2O and in 95% H2O/5% D2O. 2D NOESY with 60-, 90-, 150-, and 300-msec mixing times, 2D TOCSY, 2D 1H-15N-jr-HSQC.

**MS experiments**

DNA and protein sample buffers were exchanged against 150 mM ammonium acetate buffer, pH 6.5 using microcentrifuge gel-filtration columns (Zeba 0.5 mL, Thermo Scientific). Concentrations were determined spectrophotometrically using absorbance at 260 and 280 nm, respectively. RNA mass measurements under denaturing conditions were performed on a quadrupole/time-of-flight instrument (µQ-TOF, Bruker) operating in the negative ion mode. Samples were diluted down to 2 μM in H2O/ACN/TEA (50/50/1). Experiments under nondenaturing conditions were performed on a time-of-flight instrument (LCT, Waters) equipped with an automated chip-based nanoESI source (Triversa Nanomate, Advion Biosciences) operating in the positive ion mode. Data analysis was performed with MassLynx 3.5 (Waters).

**Sample preparation for SAXS**

SmpB is extremely well conserved. It is able to successfully complement heterologous strains deleted for their endogenous protein...
protein in the mixture is negligible. Prepared in 20 mM Tris HCl buffer, pH 7.5, 150 mM NaCl, 0.1
ports, and purified in native conditions before mixing. The literature repression of RNA ranging from 0.25 to
irradiation time of <1 sec. About 30 successive frames of 1 sec the automatic sample changer (Agilent) at a flow-rate ensuring an
continuously through the 1.5-mm-diameter quartz capillary using
SEC-SAXS measurements were complemented with standard mea-
from the column at a flow rate of 150 µL/min. For each sample, subtracted from the 250 frames recorded while the sample eluted
the void volume and used as a background scattering pattern to be
control the stoichiometry of each partner, ec-tmRNA and aa-Δ16-
(GE Healthcare) preequilibrated with the SAXS buffer. In order to
purification of the protein sample polished on a Superdex 75 column
(GE Healthcare). After concentration on Amicon Ultra-15 3K
sepharose and a monobed anion exchange Resource S column
(Ion) and the radius of gyration (R_g) were evaluated using the
Guinier approximation (Guinier 1939) ln[I(q)] = ln[I(0)] - R^2/3
q^2. Identical frames were averaged and the resulting curve spliced
with batch data at larger angles for further analysis. The distance distribution function p(r) was determined using the Indirect Fourier Transform approach as implemented in the program Gnom (Sver-
gun 1992). p(r) is the histogram of distances between any pair of
elementary scattering volumes within the particle. Finally, an estimate of the hydrated molecular volume is obtained from the Porod expression:

\[ V = \frac{2\pi I(0)}{\int_0^\infty q^2(I(q) - B) dq} \]

where B is a constant subtracted to account for the contribution of inhomogeneities within the particle.

Molecular modeling of tmRNA and complex

A cryo-electronmicroscopy study recently led to a low-resolution model of [tmRNA-SmpB] complex bound to stalled ribosomes (Ramrath et al. 2012). Using the available atomic structure of the TLD-SmpB module (Gutmann et al. 2003) together with models of major tmRNA regions based on known homologous RNA-motifs, the authors proposed a pseudoatomic model of the whole tmRNA complexed to SmpB. This structure was divided into eight modules as summarized in Table 2. Atomic coordinates of the MLD, which are lacking in the 3J18 pdb file, were reconstructed using ModelRena (Rother et al. 2011). All modules were superimposed with their centers of mass at the origin so as to avoid any bias toward an initial starting model. Their relative positions and orientations were then randomly modified in a protocol that aimed at bringing the scattering pattern of the ensemble of all modules to fit the target experimental pattern. This rigid-body approach using a simulated annealing protocol is implemented in the program SASREF (Petoukhov and Svergun 2005). Distance restraints were imposed between 5' and 3' extremities of adjacent modules so as to make possible the RNA connectivity restoration after rigid body refinement. To that end, a fixed core was defined within each module leaving on each side a poten-
tially flexible linker (from 1 to 5 nt in length). The distance restraint between modules i and i+1 was 3.5N, Å, where N is the sum of nucleotides in the 3' linker of module i and in the 5' linker of module i+1. The phosphate–sugar backbone was resealed between adja-
cent modules using ModeRena and its extended fragment library. Although connectivity restoration somewhat alters the scattering pattern of the model, its effect is limited and does not question the validity of the rigid body refinement step as shown in Figure 6B.

In all four cases (with/without SmpB, with/without Mg^{2+}), models are found in which the H2-TLD “arm” is found on either side of the loop. This is no surprise, as the X-ray scattering pattern is spheri-
cally averaged through thermal motion so that only distance, but no orientation information, is accessible. At low resolution, the two models look like mirror images of one another that have the same p(r) function, hence the same scattering pattern. The choice of the model “hand” is thus based on external information—here, the electron microscopy model. For each scattering pattern, we selected models with the correct “hand,” ending with a number of models ranging between 100 and 200, enough to provide a basis for further statistical analysis.

The program Alpraxin from the ATSA suite (Konarev et al. 2006) orients the molecule in such a way that its principal inertia

SAXS experiments

X-ray scattering data were recorded at Synchrotron SOLEIL (Gif-
sur-Yvette, France) on the SWING beamline. The wavelength of the X-rays was λ = 1.033 Å. Data were recorded using a CCD-based
detector (AVIEX) with a sample-detector distance of 3.13 m over a
useful q-range [0.004 Å^-1, 0.25 Å^-1], where q = 4π sinθ/λ is the
momentum transfer and θ is the scattering angle. All samples were
prepared in 20 mM Tris HCl buffer, pH 7.5, 150 mM NaCl, 0.1
mM EDTA, 2% glycerol, 10 mM DTT, and 5 mM MgCl2 for
Mg^{2+}-containing samples. All measurements were performed at
10°C. Data were recorded on ec-tmRNA with and without Mg^{2+}
ions, alone and complexed with SmpB in a 1:1 ratio.

Samples were loaded on the Size Exclusion-High Performance Liquid Chromatography (SE-HPLC) column (Biosec 3 from
Agilent Inc.) on-line with the SAXS measuring cell (David and
Pérez 2009). Typically, 15 µL of an ~3 g/L RNA solution were loaded on the column. Fifty successive frames of 1 sec were recorded before the void volume and used as a background scattering pattern to be
subtracted from the 250 frames recorded while the sample eluted from the column at a flow rate of 150 µL/min. For each sample, SEC-SAXS measurements were complemented with standard mea-

SAXS data analysis

Normalization to the intensity of the transmitted beam and buffer subtraction was performed on-line for all scattering frames using the locally developed software FOXTROT. Individual patterns across the monomer peak were then examined using the program package PRIMUS (Petoukhov et al. 2012). The forward scattering I(0) and the radius of gyration (R_g) were evaluated using the

(Thibonnier et al. 2010). Therefore, ec-tmRNA and aa-SmpB can be
used together for structural investigations. Δ mutants in the C-ter tail are functionally inactive but retain high affinity for tmRNA (Jacob et al. 2005; Sundermeier et al. 2005; Hallier et al. 2006) and are much easier to purify. The protein was purified by FPLC on phenyl sepharose and a monobed anion exchange Resource S column
(GE Healthcare). After concentration on Amicon Ultra-15 3K
(Millipore), the buffer was exchanged for the SAXS buffer and the purification of the protein sample polished on a Superdex 75 column
(GE Healthcare) preequilibrated with the SAXS buffer. In order to

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vectors are aligned with the coordinate axes. We applied it to the loop domains of all models (i.e., excluding domain H2 and the TLD) so that, afterward, the Z coordinate of any point is a direct measure of its distance to the “mid-plane” of the loop defined by the first two inertia axes of the loop.

To determine the orientation of the TLD arm with respect to the loop, we considered two parameters determined with respect to the mid-plane of the loop. First, a is the angle between the vector linking the center of mass of the loop O and the projection on the mid-plane M of the hinge point of H2 domain with the loop (phosphorus atom 304) and the vector MP, where P is the projection on the mid-plane of phosphorus atom 363 at the far end of the TLD (see diagram in Supplemental Fig. 4, panel A3). Second, Z is the Z-coordinate of phosphorus atom 363 that reports about the elevation of the TLD.

The analysis of the distribution of parameter values makes the implicit assumption that our sampling of the solution space is exhaustive. The distributions derived from all 300 models are practically identical to those obtained using the selected models with the correct “hand.” This strongly suggests that our sampling is satisfactory, thereby validating our conclusions.

To create a model for the ec-tmRNA in the accommodation stage, 3tyq (T. thermophilus) structure was used in which each module of the E. coli model was superimposed onto the corresponding T. thermophilus module before substitution (Weis et al. 2010).

Finally, a sketch of the tt-tmRNA conformation in the pre-accommodation stage has been proposed by Kaur et al. (2006). It comprises only phosphorus atoms of RNA moiety together with SmpB Ca atoms (2ob7). We proceeded in a similar way to obtain a model for the ec-tmRNA in the pre-accommodation stage. Regarding the complex with SmpB, the protein was added to module TLD+H2 in agreement with the crystallographic structure of the [TLD·SmpB] complex (Bessho et al. 2007).

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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