Contribution of two conserved histidines to the dual activity of archaeal RNA guide-dependent and -independent pseudouridine synthase Cbf5

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ABSTRACT

In all organisms, several distinct stand-alone pseudouridine synthase (PUS) family enzymes are expressed to isomerize uridine into pseudouridine (Ψ) by specific recognition of RNAs. In addition, Ψs are generated in Archaea and Eukaryotes by PUS enzymes which are organized as ribonucleoprotein particles (RNP)—the box H/ACA s/snoRNPs. For this modification system, a unique TruB-like catalytic PUS subunit is associated with various RNA guides which specifically target and secure substrate RNAs by base-pairing. The archaeal Cbf5 PUS displays the special feature of exhibiting both RNA guide-dependent and -independent activities. Structures of substrate-bound TruB and H/ACA sRNP revealed the importance of histidines in positioning the target uridine in the active site. To analyze the respective role of H60 and H77, we have generated variants carrying alanine substitutions at these positions. The impact of the mutations was analyzed for unguided modifications U55 in tRNA and U2603 in 23S rRNA, and for activity of the box H/ACA Pab91 sRNP enzyme. H77 (H43 in TruB), but not H60, appeared to be crucial for the RNA guide-independent activity. In contrast to earlier suggestions, H60 was found to be noncritical for the activity of the H/ACA sRNP, but contributes together with H77 to the full activity of H/ACA sRNPs. The data suggest that a similar catalytic process was conserved in the two divergent pseudouridylation systems.

Keywords: Archaea; Cbf5; H/ACA sRNP; RNA modification; pseudouridine

INTRODUCTION

Pseudouridine (Ψ) generated by isomerization of uridine is the most abundant modification found within RNAs. Until recently, Ψs have only been identified in stable RNAs as transfer, ribosomal, small nuclear and nucleolar RNAs, for which many Ψs are highly conserved and present in functionally important regions (Yu et al. 1998; Decatur and Fournier 2002; Newby and Greenbaum 2002; Badis et al. 2003; King et al. 2003; Yang et al. 2005; Ejby et al. 2007; Piekna-Przybylska et al. 2008; Karijolich and Yu 2010). Development of the high-throughput method Ψ-seq allowed transcriptome-wide quantitative mapping of Ψs in eukaryotic RNAs, including noncoding and messenger RNAs (Carlile et al. 2014; Lovejoy et al. 2014; Schwartz et al. 2014).

Site-specific pseudouridylation is catalyzed by RNA:Ψ-synthase (PUS) family enzymes, which are classified based on their homology with the bacterial RluA, RsuA, TruA, TruB, TruD proteins (for review, see Hamma and Ferré-D’Amaré 2006), and with the archaeal Pus10 protein (Roovers et al. 2006; McCleverty et al. 2007). Alternatively, pseudouridylation is based on the action of box H/ACA ribonucleoprotein (RNP) enzymes—the box H/ACA small RNPs (sRNPs) in archaea and the box H/ACA small nucleolar and small Cajal Body-specific RNPs (snoRNAs and scaRNPs, respectively) in eukaryotes (for review, see Reichow et al. 2007; Kiss et al. 2010; Watkins and Bohnsack 2012).

Escherichia coli TruB and its homolog Pus4p in Saccharomyces cerevisiae carry out site-specific recognition and
A modification at position U55 in the loop of the TΨC arm of all cellular tRNAs excluding translation initiator tRNA (Fig. 1A; Nurse et al. 1995; Becker et al. 1997). Several cocystal structures of TruB associated with a tRNA fragment mimicking the TΨC stem–loop identified the interactions established between TruB and its substrate RNA (Hoang and Ferré-D’Amaré 2001; Pan et al. 2003; Phannachet and Huang 2004). Extrusion of tRNA U55 in the active cleft of TruB is achieved by base flipping (Hoang and Ferré-D’Amaré 2001). This process relies on histidine at amino acid position 43 (H43) in TruB, which stacks against the conserved U54: A58 reverse Hoogsteen base pair (Fig. 1A2, C2; Hoang and Ferré-D’Amaré 2001; Pan et al. 2003) and which is essential for exposing U55 (Pan et al. 2003).

Archaeal protein Cbf5 (alias aCBF5) has related features with the E. coli TruB enzyme as it is able to generate in vitro the universally conserved U35 in elongator tRNAs (Fig. 1A; Roovers et al. 2006; Gurha et al. 2007; Muller et al. 2007; Kamalampeta and Kothe 2012). A pseudouridine can also be introduced at this position by the archaeal Pus10 enzyme in vitro and in vivo (Roovers et al. 2006; Gurha and Gupta 2008; Blaby et al. 2013; Chatterjee et al. 2012; Joardar et al. 2013; Kamalampeta et al. 2013). In addition, Cbf5 in association with two interacting protein partners Nop10 and Gar1 (alias aNOP10 and aGAR1) pseudouridylates in vitro a fragment of the 23S rRNA mimicking the TΨC loop of the tRNA (Muller et al. 2008). Moreover, Cbf5 is also the catalytic subunit of H/ACA sRNP comprising the ribosomal protein L7Ae in Archaea or Nhp2 in Eukaryotes, and proteins Nop10 and Gar1 (Fig. 1B1). In the context of these RNP catalysts, the specificity for substrate recognition is carried out by the RNA component of the RNP, which defines the U to be modified by the catalytic subunit Cbf5 by base-pairing with the target sequence (for review, see Hamma and Ferré-D’Amaré 2010; Kiss et al. 2010; Watkins and Bohnsack 2012).

In agreement with the feature of the bacterial TruB enzyme, the histidine equivalent to H43 at position 77 (H77) in archaeal Cbf5 is also essential for U55 tRNA modification in vitro, but does not strongly impact the activity of H/ACA sRNP (Muller et al. 2007). Based on the crystallographic structure of the substrate-bound state of the H/ACA sRNP, Duan et al. (2009) proposed that histidine H60, which is conserved in Cbf5 orthologs but not in TruB orthologs, would substitute H77 and have a functional role in U extrusion into the active site of Cbf5 (Fig. 1B2). This proposal is in agreement with another comparison of the free and substrate-bound H/ACA sRNP structures, which suggests that H60 would play an important role in coordinating substrate docking in the active site (Hamma and Ferré-D’Amaré 2001). The 3D structures were superposed with Swiss protein viewer and representation prepared with PyMol. (C1) Superposition of two H/ACA sRNPs by using the PDB access number 3HAX (Duan et al. 2009) and 3HJW (Liang et al. 2009). The histidines are in purple and the nucleobases of the substrate are in brown and of the RNA guide in sand. (C2) Superposition of the guide RNA-dependent activity of Cbf5 (B2) Scheme of an archaeal box H/ACA sRNP showing the different proteins which are represented by circles, associated with the guide RNA folded as single hairpin (in black). The substrate RNA in red is shown associated with the protein L7Ae in Archaea or Nhp2 in Eukaryotes, and proteins Nop10 and Gar1 (alias aNOP10 and aGAR1) pseudouridylates in vitro a fragment of the 23S rRNA mimicking the TΨC loop of the tRNA (Muller et al. 2008). Moreover, Cbf5 is also the catalytic subunit of H/ACA sRNP comprising the ribosomal protein L7Ae in Archaea or Nhp2 in Eukaryotes, and proteins Nop10 and Gar1 (Fig. 1B1). In the context of these RNP catalysts, the specificity for substrate recognition is carried out by the RNA component of the RNP, which defines the U to be modified by the catalytic subunit Cbf5 by base-pairing with the target sequence (for review, see Hamma and Ferré-D’Amaré 2010; Kiss et al. 2010; Watkins and Bohnsack 2012).

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Histidine H60 but not H77 was found to stack over nucleobase A or C, corresponding to nucleotide N′-1, in the crystal structures of two distinct H/ACA sRNPs bound to a substrate RNA (Duan et al. 2009; Liang et al. 2009). (C3) Structural alignments showing respective position of the conserved histidines in substrate-bound H/ACA sRNPs enzymes. The numbering of histidines is from Pyrococcus furiosus (H63 and H80 correspond to, respectively, H60 and H77 in P. abyssi). The 3D structures were superposed with Swiss protein viewer and representation prepared with PyMol. (C3) Superposition of two H/ACA sRNPs by using the PDB access number 3HAX (Duan et al. 2009) and 3HJW (Liang et al. 2009). The histidines are in purple and the nucleobases of the substrate are in brown and of the RNA guide in sand. 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D’Amaré 2010). In addition, H60 is proposed to contribute to the recognition by Cbf5 of tRNA lacking the reverse Hoogsteen pair (Gurha et al. 2007). All these observations suggest that the RNA guide-dependent and -independent activities of Cbf5 likely rely on distinct molecular determinants. Here, by mutational analysis we investigated the respective role of the two conserved histidines H60 and H77 for the dual activity of Cbf5 from the hyperthermophilic archaeon *P. abyssi*. The impact of alanine substitutions of the histidines was studied for U$_{55}$ tRNA and U$_{2603}$ rRNA modifications, and for H/ACA sRNP assembly, substrate binding and pseudouridylation activity.

**RESULTS AND DISCUSSION**

**Histidine H60 is dispensable for the RNA guide-independent activity of Cbf5**

In order to compare the functional contribution of the conserved histidine H60 with that of H77, we created variant proteins carrying a single (variant H60A and variant H77A) or a double alanine substitution (variant H77A/H60A). We used a tritium release assay to measure time course formation of Ψ$_{55}$ in tRNA$^{\text{Amp}}$, which was synthesized and tritium-labeled at C5 by incorporation of [5-$^3$H]UTP during in vitro transcription. Isomerization of uridine occurs by the breakage of the C1′-C1 glycosidic bond, followed by a rotation of the uracil ring and creation of a new C1′-C5 bond. The reaction was thus monitored by detecting liberation of tritium from C5 upon formation of the new C′-C glycosidic bond. As Cbf5 is stabilized upon binding with Nop10 and Gar1, we repeatedly measured activity of the heterotrimer Cbf5–Nop10–Gar1. In single-turnover reaction, the wild-type and the variant H60A efficiently modify tRNA with a similar apparent rate $k_{\text{app}}$ of 0.75 ± 0.31 min$^{-1}$ and 0.77 ± 0.13 min$^{-1}$, respectively (Fig. 2A). Activity was significantly reduced for variant H77A ($k_{\text{app}}$ of 0.13 ± 0.02 min$^{-1}$) and for variant H60A/H77A which combines the two mutations ($k_{\text{app}}$ of 0.16 ± 0.02 min$^{-1}$). To monitor possible small effects of the H60A substitution in an even more sensitive system, we repeated the trna modification assay with the heterodimeric Cbf5–Nop10 complex. Again, the wild-type and H60A variant show similar apparent rates of 0.58 ± 0.07 min$^{-1}$ and 0.61 ± 0.06 min$^{-1}$, respectively (Fig. 2A), whereas the apparent rates variant H77A and H60A/H77A displayed a rate reduction of more than ∼13-fold ($k_{\text{app}}$ of 0.04 ± 0.01 min$^{-1}$ and 0.04 ± 0.01 min$^{-1}$, respectively). Equivalent data were also obtained when Cbf5 was used alone (data not shown). In the presence of approximately fivefold excess of unlabeled tRNA, the multiple-turnover activity of variant H60A was comparable to that of the wild-type enzyme (Fig. 2B).

We confirmed that H60 does not contribute to the rRNA:Ψ$_{2603}$-synthase activity of Cbf5. The single-turnover reactions were performed in the presence of the auxiliary proteins Nop10 and Gar1 in the conditions used in a previous work (Muller et al. 2008). Mutation H77A reduced the rate of modification approximately fivefold compared with the wild-type enzyme (Fig. 2C). Hence, both RNA guide-independent activities of Cbf5 necessitate the presence of residue H77, but not of residue H60.

These data strongly suggest that the mechanism based on the contribution of a histidine, e.g., H43(H77), for accessibility of the substrate to TruB is conserved for Cbf5. In addition, the present results indicate that the two RNA guide-independent activities of Cbf5 are based on the same determinants,

**FIGURE 2.** Both tRNA:Ψ$^{55}$- and rRNA:Ψ$^{2603}$-synthase activities of Cbf5 require H77 but not H60. (A) Time course analysis of Ψ formation by Cbf5 in the substrate *P. abyssi* tRNA$^{\text{Amp}}$ under single-turnover conditions. The extent of Ψ formation was quantified using the tritium release assay. [3H]tRNA$^{\text{Amp}}$ (0.25 µM) was incubated with Cbf5 (wt, H60A, H77A, and H60A/H77A) and Nop10 (continuous lines) or Nop10 and Gar1 (dashed lines) at 65°C. The concentration of the proteins was 0.5 µM. (B) Ψ formation by Cbf5 in tRNA$^{\text{Amp}}$ under multiple-turnover conditions as measured by the tritium release assay. 0.25 µM [3H]tRNA$^{\text{Amp}}$ was incubated with 0.05 µM Cbf5 (wt, H60A and Nop10). (C) Analysis of Ψ formation in the RNA substrate rRNA$_{2603}$. Reactions were performed under single-turnover conditions in the presence of 30 nM uniformly [α-32P]CTP radiolabeled rRNA$_{2603}$ and 5 µM of the heterotrimer composed of the wild type and variant Cbf5 enzyme (in black) and proteins Nop10 and Gar1. The reaction was performed at 65°C for 90 min. Ψs were detected by the nearest neighbor method and the quantities were expressed as for substrate RNA 22-U in Figure 3C. The bar graph represents the relative percentage of Ψ formation compared with the wild type (panel C$_1$). The panel C$_2$ shows an example of 2D-TLC plates with the amount of Ψ formation indicated.
and likely are based on a common mechanism. Conservation of H43 in all members of the TruB-like PUS family suggests that Cbf5 produces tRNA \( \Psi_{55} \) and rRNA \( \Psi_{2603} \) in vivo. As Pus10 also modifies position U55 (Roovers et al. 2006; Gurha and Gupta 2008; Blaby et al. 2011; Kamalampeta et al. 2013), it remains to be determined whether both enzymes have redundant activities in vivo to generate tRNA \( \Psi_{55} \) and 23S rRNA \( \Psi_{2603} \) modifications. A genetic disruption strategy in archaeon \textit{Haloferax volcanii} was not able to confirm the tRNA: \( \Psi_{55} \)-synthase activity of Cbf5 in vivo due to the fact that disruption of the Pus10 encoding gene is not viable (Blaby et al. 2011). Hence, demonstration of the potential redundancy between activities of Pus10 and Cbf5 and the possibility to perform structure–function analysis in vivo necessitates archaeans for which gene disruption can be introduced by genetic tools and in which Pus10 and Cbf5 encoding genes are not essential for viability.

The Pab91 RNP enzyme lacking either histidines H60 or H77 conserves the capacity to fully modify a substrate rRNA

We next tested the potential contribution of H60 in the RNA guide-dependent activity of Cbf5. As shown in Figure 1C1, equivalent residue H63 in \textit{P. furiosus} stacks on the guide RNA nucleobase, which is engaged in the Watson–Crick interaction with the nucleobase of the substrate RNA that precedes the target U (base pair N:N\(^\prime\)-1 in Fig. 1B2). In the two crystal structures of the substrate-bound H/ACA sRNP, H63 covers the A of a A:U base pair (Duan et al. 2009) and the C of a C:G base pair (Fig. 1C1; Liang et al. 2009). In both structures, residue H80 (H77 in \textit{P. abyssi}) is positioned further away from the A:U and C:G base pairs (Fig. 1C1). From these structures, residue H60 was proposed to be a key functional determinant for substrate placement in the active site of the H/ACA RNP enzyme (Hamma and Ferré-D’Amaré 2010).

To analyze the RNA guide-dependent activity of Cbf5, we used the H/ACA sRNP enzyme Pab91 which specifically modifies U2685 in 23S rRNA from \textit{P. abyssi} (Muller et al. 2008). We first tested by EMSA the impact of the mutations on the assembly of the H/ACA sRNP (Fig. 3A). Incubation of Cbf5 (C) with the radiolabeled Pab91 guide H/ACA sRNA led to very low amounts of a stable [RNA guide:Cbf5] complex as already observed in previous works (Charpentier et al. 2005; Fourmann et al. 2013). The variant proteins bound Pab91 sRNA with the same low efficiency (compare lanes 2–5). In the presence of the core proteins L7Ae (L) and Nop10 (N) (lane 7), a major complex was formed upon binding of L7Ae on the K-loop apical structure of Pab91 but only low amounts of complex [Pab91:LCN], which corresponds to an active H/ACA sRNP enzyme, were observed (Charpentier et al. 2005). As for the [Pab91:C] complex, the mutations had no impact on the low amounts of the [Pab91:LCN] RNP enzyme (lanes 7–10). A single complex CII is formed upon addition of the unlabeled RNA substrate (Charpentier et al. 2005) (lane 11), and only the double mutation affected CII formation (lane 14), while any of the single mutation did not (lanes 12–13). To confirm these results, we tested formation of the CII’ complex obtained by association of the labeled RNA substrate 22-U with the RNP enzyme. The complex was detected after 10 min of incubation and...
disappeared during the course of the reaction (Fig. 3B, lanes 3 and 4). Variants H60A or H77A formed, respectively, slightly higher (~1.7-fold, lane 5) and lower (~0.8-fold, lane 7) complex amounts as the wild-type H/A/ACA sRNP enzyme (lane 3), but the variant H60A/H77A conducted to approximately threefold reduction in the amounts of CII’ (compare lanes 3 and 9).

Activity of the LCNG RNP enzyme containing the full quartet of core proteins L7Ae, Cbf5, Nop10, and Gar1, was next assessed in single-turnover reactions by measurement of $\Psi$ formation by the nearest neighbor technique (Fig. 3C). Both variant RNP enzymes, H60A and H77A, completely modified RNA substrate 22-U. However, compared with the wild-type enzyme ($k_{\text{app}}$ of 0.21 ± 0.014 min$^{-1}$), the rate of modification for H60A and H77A variants was lower with $k_{\text{app}}$ of 0.059 ± 0.004 min$^{-1}$ and 0.022 ± 0.005 min$^{-1}$, respectively. This corresponds, respectively, to a three to fourfold and approximately ninefold decrease to the rate of the wild-type enzyme. These data revealed that although the substitution of H60 or H77 led to a slower enzyme activity of variant H77A appeared even slower than H60. Activity of variant H77A appeared even slower than in our previous measurements (Muller et al. 2007). The H60A/H77A variant sRNP was inactive, which was in agreement with the low amounts of CII and CII’ complexes. From these data we conclude that contrary to the deduction made by analysis of the crystal structures, residue H60 appears to not be a key functional determinant of the RNA-guide-dependent activity unless H77 is absent.

**FIGURE 3.** Contribution of both histidines H60 and H77 for the guide RNA-dependent activity of Cbf5 within the Pab91 RNP. (A) Analysis by electrophoretic mobility shift assay (EMSAs) of various box H/A/ACA sub-RNPs and the complex formed between the box H/A/ACA sRNP and a substrate RNA. The guide RNA Pab91 which was radiolabeled at the 5' end with $^{32}$P was incubated with the wild type (wt) or the variant forms of Cbf5 (variants H60A, H77A, and H60A/H77A) as indicated above each lane, alone (lanes 2–5) or in the presence of L7Ae and Nop10 (lanes 7–14) and in the absence (lanes 7–10) or the presence of the substrate RNA 22-U (lanes 11–14). After 10 min incubation at 65°C, the different complexes were fractionated by nondenaturing polyacrylamide gel electrophoresis. The identity of the sub-RNPs is indicated: C and LCN correspond, respectively, to the Pab91-Cbf5 and the Pab91-L7Ae-Cbf5–Nop10 complexes; the CII complex is formed upon association of the Pab91 LCN RNP and the substrate RNA (Charpentier et al. 2005). (B) Analysis by EMSAs of the complex formed between the RNP enzyme and the substrate RNA (complex CII’). The substrate RNA 22-U which was radiolabeled at the 5' end with $^{32}$P was incubated with the wild-type and variant Pab91 LCN RNP assembled with the unlabeled Pab91 sRNA and the L7Ae, Cbf5, and Nop10 proteins as described in Materials and Methods. After 10 and 60 min at 65°C, the different complexes were fractionated by nondenaturing polyacrylamide gel electrophoresis. The identity of the sub-RNPs is indicated: C and LCN correspond, respectively, to the Pab91-Cbf5 and the Pab91-L7Ae-Cbf5–Nop10 complexes; the CII complex is formed upon association of the Pab91 LCN RNP and the substrate RNA (Charpentier et al. 2005). (C) Time course analysis of $\Psi$ formation by the Pab91 LCNG RNP enzyme in the 22-U substrate RNA. The uniformly [α-$^{32}$P] CTP radiolabeled substrate RNA 22-U was incubated with the guide RNA Pab91 and proteins LCNG wild-type (in black), H60A (in dark gray), H77A (in gray), and H60A/H77A (in light gray) and analyzed by 1D-TLC as described in Materials and Methods. The radioactivity was quantified by PhosphorImager analysis. The quantities of $\Psi$ nucleotides formed are expressed in moles per mole of 22-U.

If the mechanism leading to U extrusion upon substrate base-pairing with the guide RNA necessitates the positioning of a histidine on the N nucleobase, the present results indicate that H77 might fulfill this role in the absence of H60. The most favorable conformation of the active site occupied by the substrate RNA is probably the one trapped in the two crystal structures (Fig. 1C). The partial supplementation of H60 by H77 would imply that the catalytic site of H/A/ACA sRNP possesses a degree of plasticity that could allow a repositioning of H77 over the N nucleobase. In this scenario, the alternative conformation of the H60A variant enzyme would bind substrate with a similar efficiency as the wild-type enzyme (Fig. 3B), but would not be optimal for catalysis as slower activity was measured for the H60A sRNP variant (Fig. 3C). The structure of the active site is likely also modified in the H/A/ACA sRNP enzyme lacking H77 leading to a lower ability to stably bind the substrate RNA (Fig. 3B). Hence, the optimal function of H60 relies on the presence of H77.

**CONCLUSION**

Two major conclusions can be drawn from the results of the present study. First, the universally conserved histidine H77 —corresponding to H43 in the TruB enzyme—strongly contributes to the nonguided activity of the PUS Cbf5, but not H60 which is conserved in only the Cbf5-like PUS. Second, optimal activity of the H/A/ACA sRNP enzyme necessitates presence of both histidines H60 and H77. Among the two histidines, only H77 represents a common determinant for the RNA guide-dependent and -independent activities. These data suggest that during evolution, molecular determinants were conserved for the two divergent modes of PUS activity. Knowledge on the 3D structure adopted by the substrate-bound variant H60A and H77A H/A/ACA sRNP enzymes would help to refine our appreciation on the organization and plasticity of active site of the PUS enzymes.

**MATERIALS AND METHODS**

**Protein expression and purification**

The recombinant proteins were produced in *Escherichia coli* BL21 (DE3) (Stratagene) as GST-fusion proteins from recombinant pGEX-6P1 plasmids and purified as described previously (Charpentier et al. 2005; Muller et al. 2007). Site-directed mutagenesis based on PCR was used to prepare the plasmid variants encoding proteins Cbf5 H60A, H77A, and H60A/H77A. For the H77A substitution, the PCR reaction was performed with the plasmid pGEX-6P1-Cbf5 as the template and with two primers carrying the substitution (forward: 5’-GGTGAGGTTCCACACGACCCGCTTTC TCCAAGTT-3’ and reverse: 5’-AAGTGGAGAAACGCCTGCGGCTCTGGATCAAACCTCAGC-3’). For the H60A and H60A/H77A substitutions, the PCR reactions were realized with the plasmids pGEX-6P1-Ch5 and Ch5 H77A, respectively, and two primers carrying the substitutions (forward: 5’-GCCCTACTAGGCCTGAGG TCTCAGTT-3’ and reverse: 5’-AGTGGAGAAACGCCTGCGGCTCTGGATCAAACCTCAGC-3’).
TCGTTGC-3' and reverse: 5'-GCAACGACCTAGCGCTAGTAGGGCCG-3'). The nucleotides corresponding to the substituted codon are indicated in bold.

**In vitro transcription of the tRNA^Asp, the guide RNA Pab91 and its target RNA 22-U**

The DNA template used for in vitro transcription of the various RNAs, i.e., the *P. abyssi* Pab91 guide RNA and tRNA^Asp^, were obtained by PCR amplification with the forward primer generating the sequence of the T7 promoter as previously described (Charpentier et al. 2005, 2007; Muller et al. 2007, 2008; Kamalampera et al. 2013). The template for synthesis of the substrate RNA 22-U and the 23S rRNA fragment rRNA2603 were obtained by hybridization of two complementary oligodeoxyribonucleotides, i.e., 5'-TACGACTCTACTATAAGGTGTATAGCGAGAGA-3' and 5'-TCTCTCAGACGGTCTAAACCCCTATAGGTGCTATATTA-3' for 22-U, and 5'-TAAACGACTCTACTATAAGGTGTATAGCGAGAGA-3' and 5'-ATGGGAAGGCGCAGGCCCTATAGGTGCTATATTA-3' for rRNA2603, with the T corresponding to the target U in the coding strand indicated in bold and the sequence of the T7 promoter in italic. RNAs 22-U and rRNA2603 correspond to two fragments of the 23S rRNA encompassing, respectively, nucleotides U2685 and U2603. Conditions for the reactions of in vitro transcription and purification were performed as detailed previously (Charpentier et al. 2007; Kamalampera and Kothe 2012). The substrate RNAs 22-U and tRNA^Asp^, used for activity measurements were radiolabeled during transcription in the presence of [α-32P]CTP.

The in vitro transcription of radiolabeled tRNA^Asp^ was performed in the presence of 3 mM (ATP, CTP, and GTP) and 0.1 mM [5-3H]UTP.

**H/ACA sRNP enzyme reconstitution**

As previously described (Charpentier et al. 2007), the box H/ACA sRNPs were reconstituted either by incubation of 2 pmol of unlabeld or 50 fmol of labeled Pab91 RNA with 0.2–1 μM of recombinant proteins in buffer D (20 mM HEPES KOH pH 7.9, 150 mM KCl, 1.5 mM MgCl₂, and 0.2 mM EDTA), and in the presence of competitor tRNA (250 ng) at 65°C.

**Analysis of H/ACA sRNP enzyme:substrate complex formation, and activity measurement**

Two approaches were used. The RNPs assembled with the radiolabeled guide RNA Pab91 were incubated with unlabeled 22-U (2 pmol) leading to CHI complex formation (Charpentier et al. 2005). In the second procedure, complex CHI was formed by association of the radiolabeled substrate 22-U RNA (~0.03 μM) with the H/ACA sRNP (~0.45 μM) reconstituted with unlabeled guide RNA Pab91 (Charpentier et al. 2005, 2007; Fourmann et al. 2013). The complexes were fractionated by nondenaturing polyacrylamide gel electrophoresis and visualized with a PhosphorImager (Typhoon 9410, Amersham Biosciences). The radioactivity present in complexes CHI and CHI’ or in the unbound fraction was quantified using the ImageQuant software Version 5.2 (Molecular Dynamics). The percentage of RNA in the slower migrating complex was calculated from the radioactivity in each band relative to the total radioactivity in the lane.

RNA:Ψ-synthase activity of the reconstituted sRNP Pab91 was measured by using the nearest neighbor method and the conditions for time course analysis of single-turnover reactions were as previously described (Charpentier et al. 2005; Fourmann et al. 2013). Briefly, unlabeled guide RNA (~0.5 μM) and 150 fmol (~0.03 μM) of [α-32P]CTP labeled substrate RNA were mixed at 65°C with protein combinations LCN (L7Ae–Chb5–Nop10) or LCNG (L7Ae–Chb5–Nop10–Gar1) (1–3 μM). Aliquots were collected at several time points and the reaction was stopped by phenol–chloroform extraction followed by ethanol precipitation. The recovered RNA substrates were digested with 0.4 units of RNase T2 to generate 3’ phospho-mononucleotide. The resulting mononucleotides were fractionated by mono- or bidimensional thin layer chromatography, and the radioactivity in the resulting bands was quantified with a PhosphorImager using the ImageQuant software. The quantities of Ψ residues formed were determined as described before (Charpentier et al. 2007). The obtained values in moles of Ψ per mole of substrate RNA (mol/mol⁻¹) were analyzed with the Prism software Version 5.04 (GraphPad). Initial rates of Ψ formation and extents of the reaction were determined from at least three independent experiments. The single-turnover experiments were analyzed by fitting the data with a single-exponential equation and initial rates k_app correspond to the slope of the curve for the 0–5-min interval.

**Tritium release assay**

The pseudouridylation activity radiolabeled reactions were performed in buffer D at 65°C as described previously (Kamalampera and Kothe 2012). 250 nM of [³H]tRNA^Asp^ were incubated in the presence of 500 nM or 50 nM of proteins Chb5–Nop10 for single- and multi-turnover, respectively. Samples were collected from the reaction at different times, and proteins and tRNA were removed by two extractions with 5% (w/v) activated charcoal (Norit A, EMD, CX0655-1) in 0.1 M HCl. The supernatant was further cleared by filtration through glass wool. After adding EcoLite scintillation cocktail, the quantity of [³H] in the final solution was estimated by scintillation counting. Each experiment was repeated at least twice and results were fit with an exponential equation as it was done for the RNA guide-dependent activity.

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