

# Guide RNA acrobatics: positioning consecutive uridines for pseudouridylation by H/ACA pseudouridylation loops with dual guide capacity

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Site-specific pseudouridylation of human ribosomal and spliceosomal RNAs is directed by H/ACA guide RNAs composed of two hairpins carrying internal pseudouridylation guide loops. The distal “antisense” sequences of the pseudouridylation loop base-pair with the target RNA to position two unpaired target nucleotides 5'-UN-3', including the 5' substrate U, under the base of the distal stem topping the guide loop. Therefore, each pseudouridylation loop is expected to direct synthesis of a single pseudouridine ( $\Psi$ ) in the target sequence. However, in this study, genetic depletion and restoration and RNA mutational analyses demonstrate that at least four human H/ACA RNAs (SNORA53, SNORA57, SCARNA8, and SCARNA1) carry pseudouridylation loops supporting efficient and specific synthesis of two consecutive pseudouridines ( $\Psi\Psi$  or  $\Psi N\Psi$ ) in the 28S ( $\Psi 3747/\Psi 3749$ ), 18S ( $\Psi 1045/\Psi 1046$ ), and U2 ( $\Psi 43/\Psi 44$  and  $\Psi 89/\Psi 91$ ) RNAs, respectively. In order to position two substrate Us for pseudouridylation, the dual guide loops form alternative base-pairing interactions with their target RNAs. This remarkable structural flexibility of dual pseudouridylation loops provides an unexpected versatility for RNA-directed pseudouridylation without compromising its efficiency and accuracy. Besides supporting synthesis of at least 6% of human ribosomal and spliceosomal  $\Psi$ s, evidence indicates that dual pseudouridylation loops also participate in pseudouridylation of yeast and archaeal rRNAs.

[*Keywords:* pseudouridine; guide RNA acrobatics; box H/ACA RNAs; RNA-guided RNA modification; pseudouridylation; RNA–RNA interaction]

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Mature cellular RNAs carry >160 different types of post-transcriptionally modified nucleotides (Boccaletto et al. 2018). The most prevalent modified nucleotide is pseudouridine ( $\Psi$ ), the 5-ribosyl isomer of U, which is present in transfer RNAs (tRNAs), ribosomal RNAs (rRNAs), small nuclear RNAs (snRNAs), and also in polyadenylated mRNAs (Reddy and Busch 1988; Bakin and Ofengand 1998; Carlile et al. 2014; Schwartz et al. 2014; Spenkuch et al. 2014; Li et al. 2015; Taoka et al. 2018). U and  $\Psi$  share identical Watson-Crick faces, but the endocyclic N1 group of  $\Psi$  can function as an extra proton donor supporting water-coordinated hydrogen bonding that increases the thermodynamic stability of base stacking and the rigidity of the phosphodiester backbone (Arnez and Steitz 1994; Davis 1995; Newby and Greenbaum 2002a,b). Cor-

rect pseudouridylation of spliceosomal snRNAs, rRNAs, and tRNAs is important for efficient and accurate pre-mRNA splicing and protein synthesis (Ge and Yu 2013; Spenkuch et al. 2014; De Zoysa and Yu 2017; Sloan et al. 2017; Bohnsack and Sloan 2018; Adachi et al. 2019). The functional significance of mRNA pseudouridylation is still unclear, but artificially introduced  $\Psi$ s can alter stop codon reading, increase mRNA stability and translation efficiency, and suppress RNA immunogenicity (Karikó et al. 2008; Karijovich and Yu 2011; Fernández et al. 2013; Hoernes et al. 2016).

RNA pseudouridylation is achieved by stand-alone and guide RNA-associated  $\Psi$  synthases. The stand-alone  $\Psi$  synthases recognize local RNA structures and frequently show “region and/or multisite specificity” (Spenkuch et al. 2014). Site-specific synthesis of eukaryotic

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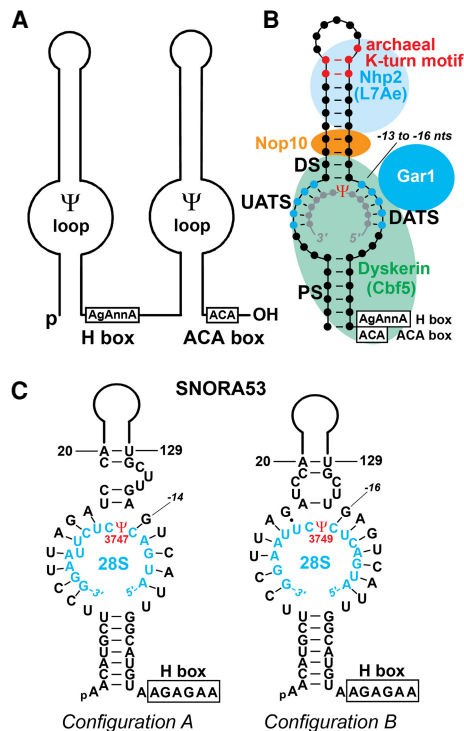
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spliceosomal and eukaryotic and archaeal ribosomal  $\Psi$ s located in diverse sequence and structural environments is mediated by H/ACA guide RNA-associated  $\Psi$  synthases (Ganot et al. 1997a; Ni et al. 1997; Darzacq et al. 2002). Through base-pairing with the target RNA, the guide RNAs provide high sequence specificity for the associated  $\Psi$  synthase, making the pseudouridylation reaction independent from local two-dimensional RNA structures. While archaeal H/ACA RNAs contain one to three hairpin structures, eukaryotic H/ACA RNAs are composed of two hairpins that are connected and tailed with short single-stranded sequences carrying the conserved H (AgAnnA) and ACA (less frequently, AUA) boxes, respectively (Fig. 1A; Balakin et al. 1996; Kiss et al. 1996; Ganot et al. 1997b). The molecular principles of RNA-guided pseudouridylation were established more than two decades

ago (for reviews, see Kiss et al. 2010; Watkins and Bohnsack 2012; Lui and Lowe 2013; Yu and Meier 2014; Czekay and Kothe 2021). The H/ACA hairpins contain internal pseudouridylation guide loops carrying bipartite target recognition (antisense) elements preceding and following the distal stem (DS) that closes the pseudouridylation loop (Ganot et al. 1997a). Base-pairing of the bipartite antisense sequences with the substrate RNA forms an RNA three-way helical junction positioning two unpaired substrate nucleotides (5'-UN-3') into the "pseudouridylation pocket" at the base of the DS (Fig. 1B). Invariantly, the unpaired 5' U that is separated by 13–16 nt from the H or ACA box of the guide RNA is converted into  $\Psi$ .

Each H/ACA hairpin associates with four evolutionarily conserved proteins: Nop10, Gar1, Nhp2 (L7Ae in archaea), and the  $\Psi$  synthase dyskerin (also called NAP57 in mice and Cbf5 in yeast and archaea) (Fig. 1B). First, the structure of archaeal single-hairpin ACA RNP has been solved (Li and Ye 2006; Liang et al. 2007, 2009; Duan et al. 2009). The ACA RNA hairpin is anchored to the core trimer of Cbf5, Nop10, and L7Ae through bindings of Cbf5 to the ACA box and the proximal stem (PS) and L7Ae to the kink (K)-turn or K-loop in the apical stem-loop. These interactions position the selected target U at the catalytic center of Cbf5. Gar1 associates with the catalytic domain of Cbf5 and controls its activity through promoting accurate substrate placing and releasing (Duan et al. 2009; Wang et al. 2015). Eukaryotic double-hairpin H/ACA RNPs have similar structural organization (Egan and Collins 2010; Li et al. 2011). In contrast to archaeal L7Ae, eukaryotic Nhp2 binds to the distal stem-loop independently from a K-turn motif. While Nhp2 alone has little H/ACA RNA binding affinity, upon incorporation into the dyskerin–Nop10–Nhp2 core complex, it binds to the distal stem-loop with high affinity and specificity (Wang and Meier 2004; Baker et al. 2005; Li et al. 2011). Eukaryotic H/ACA RNPs directing rRNA pseudouridylation accumulate in the nucleolus and are called small nucleolar RNPs (snoRNPs). Guide RNPs mediating snRNA pseudouridylation concentrate in the nucleoplasmic Cajal body (CB) and are termed small CB-specific RNPs (scaRNPs) (Darzacq et al. 2002).

Due to their stringent target selection strategy, the H/ACA pseudouridylation guide loops are believed to position only one substrate U for pseudouridylation in their target sequences (Fig. 1B). However, arguing against this notion, the 3'-terminal pseudouridylation loops of *Xenopus* pugU2-43, yeast snR82, and archaeal sR-h45 H/ACA RNAs have been implicated in synthesis of two consecutive spliceosomal and ribosomal  $\Psi$ s ( $\Psi\Psi$  and  $\Psi N\Psi$ ) through unclear mechanisms (Schattner et al. 2004; Deryusheva and Gall 2013, 2017; Majumder et al. 2020). Here, functional characterization of human H/ACA RNAs reveals that at least four human H/ACA RNAs possess pseudouridylation loops with dual guide capacity. We demonstrate that the 5'-terminal pseudouridylation loop of SNORA53 and the 3'-terminal pseudouridylation loops of SNORA57, SCARNA8, and SCARNA1 direct formation of  $\Psi$ 3747/ $\Psi$ 3949 in the 28S rRNA,  $\Psi$ 1045/ $\Psi$ 1046 in the 18S rRNA, and  $\Psi$ 43/ $\Psi$ 44



**Figure 1.** Selection of substrate uridines for pseudouridylation by box H/ACA guide RNAs. (A) Schematic structure of eukaryotic bipartite box H/ACA pseudouridylation guide RNAs. The consensus sequences of the conserved H and ACA box motifs and the internal pseudouridylation guide loops ( $\Psi$ -loop) are shown. (B) Schematic consensus structure of the 5' and 3' hairpins of H/ACA pseudouridylation guide RNPs. The antisense sequences (blue dots) and the selected target sequences (gray dots) with the target U ( $\Psi$ ) positioned for pseudouridylation are shown. The proximal and distal stems (PS and DS) together with the upstream and downstream antisense target stems (UATS and DATS) are indicated. The four H/ACA core proteins (dyskerin, Nop10, Nhp2, and Gar1) are shown. Archaeal ACA RNAs contain K-turns or K-loops. (C) Potential base-pairing interactions of the proposed 5'-terminal pseudouridylation guide loop of human SNORA53 with 28S rRNA positioning U3747 or U3749 for pseudouridylation.

and  $\Psi$ 89/  $\Psi$ 91 in the U2 spliceosomal snRNA, respectively. A detailed mutational analysis of SNORA53, SNORA57, and SCARNA1 confirmed that dynamic alternative guide RNA and substrate RNA base-pairing interactions, termed guide RNA acrobatics, support the sequence specificity and efficacy of the dual pseudouridylation guide loops of human H/ACA RNAs. Since the previously reported dual pseudouridylation guide loops of *Xenopus* pugU2-43, yeast snR82, and archaeal sR-h45 H/ACA RNAs can also form alternative guide-target interactions that can correctly position their target Us for pseudouridylation, we propose that H/ACA guide RNA acrobatics is an evolutionarily conserved strategy of pseudouridylation guide RNAs for accurate and efficient synthesis of two proximate  $\Psi$ s.

## Results

### *Human SNORA53 is required for synthesis of both $\Psi$ 3747 and $\Psi$ 3749 in the 28S rRNA*

Human SNORA53, originally named ACA53 (Kiss et al. 2004), is a box H/ACA “orphan” snoRNA, as it has not been assigned for RNA modification or for any other cellular function. Closer inspection of SNORA53 revealed that its 5'-terminal hairpin can be folded into two alternative structures accommodating slightly different putative pseudouridylation loops that, in principle, could direct synthesis of  $\Psi$ 3747 and  $\Psi$ 3749 in the 28S rRNA (Fig. 1C). Thus far, no H/ACA snoRNAs have been proposed to govern  $\Psi$ 3747 and  $\Psi$ 3749 synthesis (Jorjani et al. 2016). However, in the newly suggested SNORA53–28S rRNA interactions, the alleged pseudouridylation loops, instead of being closed by perfect distal stems, are topped by unusually unstable structures encompassing several bulged and unpaired nucleotides. Moreover, in configuration A assigned for synthesis of  $\Psi$ 3747, the upstream anti-sense target stem (UATS) is interrupted with the bulged U3742 residue in the 28S rRNA sequence. To test SNORA53 participation in  $\Psi$ 3747 and/or  $\Psi$ 3749 synthesis, expression of SNORA53 was knocked out in human HAP1 cells. Mammalian box H/ACA guide RNAs are synthesized within introns of protein-coding genes and are processed from the removed and debranched host introns. The human SNORA53 gene is located within the fifth intron of the *SLC25A3* gene in sense orientation, indicating that SNORA53 is a canonical intronic snoRNA (Fig. 2A). A fragment of the fifth intron of *SLC25A3* encompassing the SNORA53 gene was excised by CRISPR/Cas9 genome editing. The correctness of SNORA53 deletion was verified by PCR amplification and gel electrophoretic analysis of the truncated *SLC25A3* gene fragment (Fig. 2B). RT-PCR detected comparable levels of correctly spliced mature *SLC25A* mRNA in the SNORA53 knockout (KO) and control HAP1 cells, demonstrating that the removal of the SNORA53 gene had no detectable impact on the expression of the *SLC25A3* host gene (Fig. 2C). Finally, RNase A/T1 mappings performed with an internally labeled sequence-specific RNA probe failed to detect SNORA53 expression in SNORA53-KO cells, although accumula-

tion of the U85 control intronic scaRNA remained unaltered (Fig. 2D).

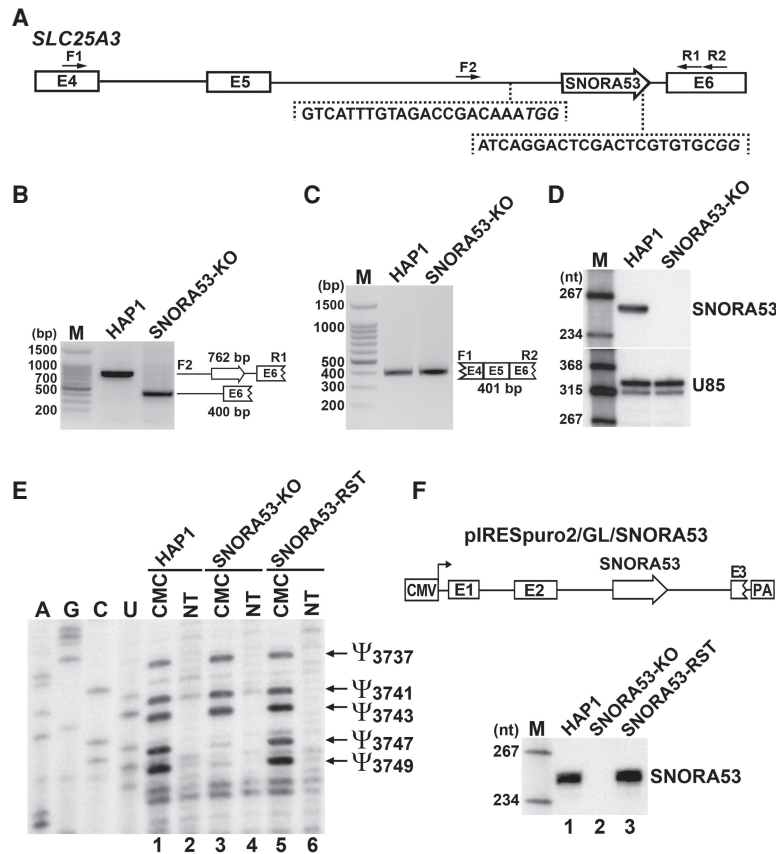
To compare the pseudouridylation states of 28S rRNA at U3747 and U3749 in the presence and absence of SNORA53, cellular RNAs extracted from HAP1 and SNORA53-KO cells were incubated with CMCT [N-cyclohexyl-N'(2-morpholinoethyl)carbodiimide metho-p-toluenesulfonate], which efficiently and irreversibly reacts with  $\Psi$  (Bakin and Ofengand 1998). The covalently bound bulky CMCT molecule blocks reverse transcriptase elongation 1 nt before the modified  $\Psi$  (Fig. 2E). Primer extension analysis of CMCT-treated control HAP1 RNAs with a terminally labeled 28S rRNA-specific oligonucleotide primer detected several known  $\Psi$ s, including  $\Psi$ 3737,  $\Psi$ 3741,  $\Psi$ 3742,  $\Psi$ 3747, and  $\Psi$ 3749 (Fig. 2E, lane 1). In contrast, primer extension mapping of CMCT-modified 28S rRNA from SNORA53-KO cells failed to detect pseudouridylation of U3747 and U3749, although synthesis of  $\Psi$ 3737,  $\Psi$ 3741, and  $\Psi$ 3742 remained unaffected (Fig. 2E, lane 3).

To exclude the formal possibility that disruption of U3747 and U3749 pseudouridylation was a consequence of undesired CRISPR/Cas9 off-target activity, we restored SNORA53 accumulation in the SNORA53-KO cell line by using the pIRESpuro/GL expression plasmid (Fig. 2F; Vitali and Kiss 2019). After selection of a puromycin-resistant monoclonal cell line, accumulation of SNORA53 was confirmed by RNase A/T1 mapping (Fig. 2F, lane 3). Primer extension mapping of 28S rRNA pseudouridylation showed that restoration of SNORA53 accumulation re-established pseudouridylation of U3747 and U3749, demonstrating that SNORA53 is responsible for the synthesis of both  $\Psi$ 3747 and  $\Psi$ 3749 in the human 28S rRNA (Fig. 2E, lane 5).

### *The 5'-terminal pseudouridylation loop of SNORA53 directs synthesis of both $\Psi$ 3747 and $\Psi$ 3749 in the 28S rRNA*

The 3' hairpin of SNORA53 lacks evolutionarily conserved sequences complementary to rRNAs or other stable cellular RNAs (Supplemental Fig. S1A), suggesting that the elements supporting pseudouridylation of 28S-U3747 and 28S-U3749 are located in the 5'-terminal hairpin of SNORA53. To validate this assumption and to learn about the molecular mechanism supporting the predicted double-guide activity of the 5'-terminal pseudouridylation loop of SNORA53, we performed mutational analysis of SNORA53.

The 5' hairpins of SNORA53 RNAs derived from phylogenetically distant species fold into similar two-dimensional structures composed of a long major hairpin and a short internal stem-loop branched out from the main hairpin (Fig. 3A; Supplemental Fig. S1A). However, upon expression of truncated SNORA53 RNAs lacking either the short internal stem-loop (SNORA53d1) or the distal part of the major 5' hairpin (SNORA53d2) in SNORA53-KO cells (Fig. 3A,B), synthesis of both  $\Psi$ 3747 and  $\Psi$ 3749 was restored (Fig. 3C, lanes 5,7), indicating that all guide elements supporting U3747 and U3749 pseudouridylation



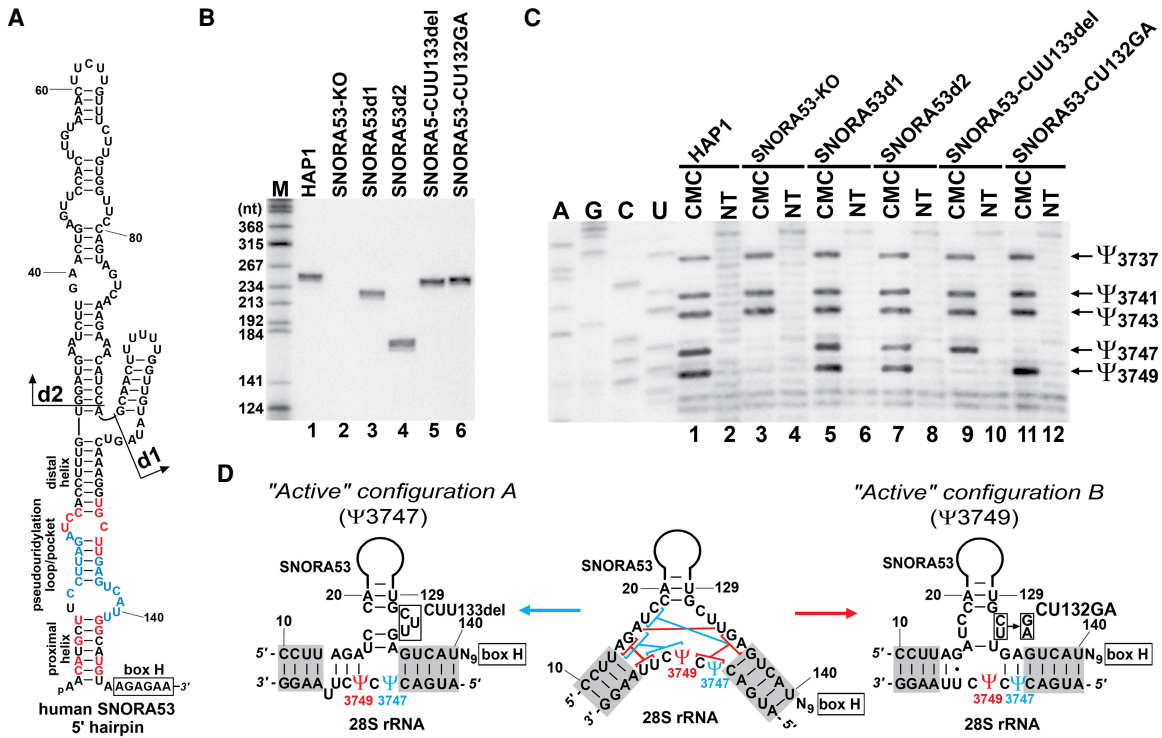
**Figure 2.** Human SNORA53 is required for synthesis of both Ψ3747 and Ψ3749 in the 28S rRNA. (A) A strategy for CRISPR/Cas9-mediated deletion of human *SNORA53* gene in HAP1 cells. Schematic structure of a fragment of the human *SLC25A3* gene encompassing exons 4–6 (E4–E6) and the intronic *SNORA53* gene (open arrow) is shown. Positions of the forward (F1 and F2) and reverse (R1 and R2) primers used for genomic and RT-PCR analysis are indicated. The genomic sequences targeted by sgRNAs are presented in sense orientation. The PAM sequences are in italics. (B) PCR analysis of genomic DNAs extracted from *SNORA53* knockout (SNORA53-KO) and parental HAP1 cells. The amplified DNA fragments were analyzed on 1% agarose gel. (Lane M) DNA size markers in base pairs (bp). Schematic structures and expected lengths of the amplified fragments are shown at the right. (C) RT-PCR analysis of the accumulation of spliced *SLC25A3* mRNA in SNORA53-KO and HAP1 cells. The structure of the amplified cDNA fragment is shown. (D) SNORA53 accumulation. RNAs isolated from HAP1 and SNORA53-KO cells were annealed with internally labeled SNORA53- or U85-specific antisense RNA probes and digested with a mixture of RNase A and T1. The protected RNA fragments were analyzed on a 6% sequencing gel. (Lane M) Single-stranded DNA size markers in nucleotides (nt). (E) Mapping of 28S rRNA pseudouridylation in HAP1 and SNORA53-KO cells before and after restoration of SNORA53 expression (SNORA53-RST). Ψs covalently modified by CMCT were detected by primer extension analysis using a terminally labeled 28S-specific primer and

AMV reverse transcriptase. (Lanes A, G, C, U) Dideoxy sequencing reactions performed on 28S ribosomal DNA with the same 28S-specific primer. (Lanes NT) Primer extension reactions performed on nontreated control RNAs. Ψs corresponding to the detected RT stop signals are indicated at the right. (F) Restoration of SNORA53 accumulation in SNORA53-KO cells. Schematic structure of the pIRESpuro2/GL/SNORA53 expression construct is shown. The coding region of the human *SNORA53* gene (open arrow) was inserted into the second intron of the truncated β-globin gene (E1–E3) placed under the control of the cytomegalovirus (CMV) promoter and the globin polyadenylation signal (PA). The resulting CMV-globin-SNORA53 expression unit was inserted into the pIRESpuro2 expression plasmid and transfected into SNORA53-KO cells. Puromycin-resistant cell colonies were isolated, and expression of SNORA53 in SNORA53-RST cells was verified by RNase A/T1 mapping.

are confined to the proximal part of the SNORA53 5' hairpin. Consistent with its functional importance, the proximal part of the SNORA53 5' hairpin contains many phylogenetically conserved nucleotides (Fig. 3A; Supplemental Fig. S1). Significantly, all nucleotides predicted to participate in alternative intra- and intermolecular base-pairing interactions positioning either U3747 or U3749 for pseudouridylation are conserved in the known SNORA53 sequences (Fig. 1C; Supplemental Fig. S1).

To unambiguously demonstrate that the 5'-terminal pseudouridylation loop region of SNORA53 is able to direct synthesis of both 28S-Ψ3747 and 28S-Ψ3749, we attempted to separate the two pseudouridylation guide activities of SNORA53 through construction of mutant SNORA53 RNAs directing either Ψ3747 or Ψ3749 formation. In the proposed SNORA53–28S interactions, in both configuration A and B, the C10-U13 upstream and the G136-U140 downstream antisense sequences are base-paired with the A3753-G3756 and A3742-C3746 regions of 28S rRNA (Fig. 3D, shaded helices). In the center of the guide RNA–substrate RNA three-way helical

junction, alternative base-pairing interactions are predicted to govern the structural rearrangements positioning either U3747 or U3749 for pseudouridylation (Fig. 3D, middle panel). Deletion of the 131-CUU-133 bulged nucleotides (CUU133del) was expected to lock the SNORA53–28S rRNA interaction in configuration A and to prevent formation of functional configuration B. Replacement of the 131-CU-132 nucleotides for GA was predicted to stabilize configuration B and to interfere with folding into configuration A. When expressed in SNORA53-KO cells, the SNORA53-CUU133del snoRNP efficiently synthesized Ψ3747, but failed to convert U3749 into Ψ (Fig. 3C, lane 9). On the contrary, the SNORA53-CU132GA snoRNP supported Ψ3749 formation, but failed to synthesize Ψ3747 (Fig. 3C, lane 11). Thus, demonstration that sequence alterations can restrict the pseudouridylation guide ability of SNORA53 to the synthesis of either Ψ3747 or Ψ3749 in a predictable way validated the proposed dual guide capacity of the 5'-terminal pseudouridylation loop of SNORA53 and strongly supported the correctness and functional relevance of the



**Figure 3.** The 5'-terminal pseudouridylation loop of SNORA53 directs pseudouridylation of U3747 and U3749 in the 28S rRNA. (A) A computer-predicted secondary structure of the 5' hairpin of human SNORA53 based on minimum free energy calculation (Zuker 2003). The evolutionarily invariant nucleotides predicted to base-pair with 28S rRNA are in blue. Other conserved nucleotides are in red. Arrows indicate internal deletions d1 and d2. (B) Expression of mutant SNORA53 RNAs in SNORA53-KO cells monitored by RNase A/T1 protection with sequence-specific antisense RNA probes. As controls, accumulation of endogenous SNORA53 was also tested in HAP1 and SNORA53-KO cells. (Lane M) Single-stranded DNA size markers. (C) Primer extension mapping of 28S rRNA pseudouridylation in HAP1 and SNORA53-KO cells either lacking or expressing mutant SNORA53 RNAs as indicated above the lanes. For details, see the legend for Figure 2E. (D) Proposed structural rearrangements of the three-way helical junctions formed by the 5'-terminal pseudouridylation loop of human SNORA53 and 28S rRNA positioning either U3747 or U3749 for pseudouridylation. The proposed alternative base-pairing interactions leading to "active" configuration A or B are indicated by blue and red lines, respectively. The common base-pairing interactions present in both configurations are shaded. The deleted or altered nucleotides in mutant SNORA53-CUU133del and SNORA53-CU132GA RNAs are boxed.

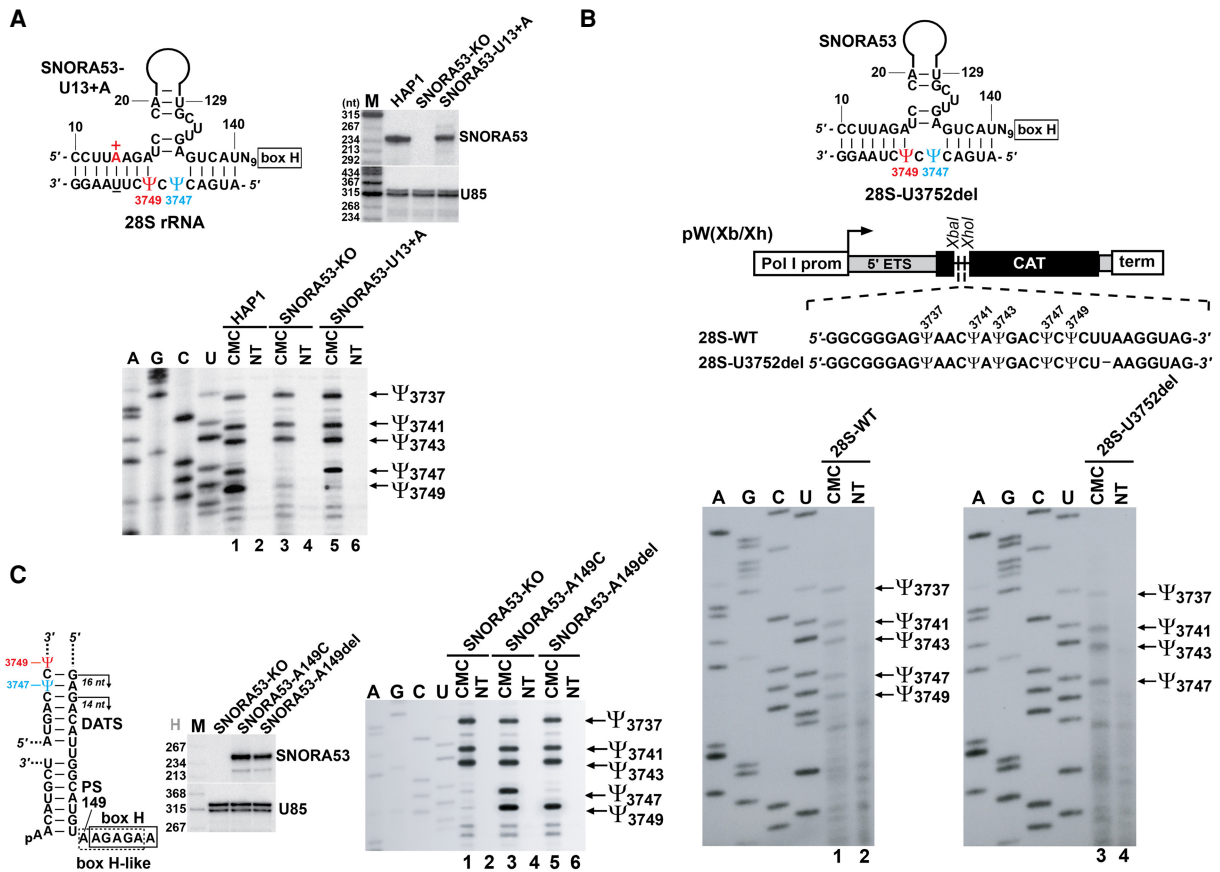
proposed alternative base-pairing interactions of SNORA53 with 28S rRNA positioning U3747 or U3749 for pseudouridylation.

*The bulged ribosomal U3752 residue supports the dual guide activity of SNORA53*

Normally, the antisense elements of pseudouridylation guide RNAs form perfect double helices with their target sequences (Fig. 1B; Ganot et al. 1997a). However, in configuration A positioning U3747 for pseudouridylation, the upstream antisense target stem (UATS) formed by the upstream antisense element of SNORA53 (10-CCUUAGA-17) and its 28S rRNA target sequence (3749-UCUUAAGG-3756) is interrupted by the bulged U3752 ribosomal residue (Figs. 1C, 3D). Given that the SNORA53 antisense and the 28S target sequences are conserved, we speculated that bulging of 28S-U3752 may contribute to the dual guide capacity of the 5'-terminal pseudouridylation loop of SNORA53. To test this hypothesis, an extra A residue was inserted into SNORA53

between U13 and A14 in order to base-pair with the bulged ribosomal U3452 residue (Fig. 4A). Expression of the mutant SNORA53-U13+A RNA in SNORA53-KO cells restored pseudouridylation of U3747, but failed to support formation of Ψ3749 (Fig. 4A, lane 5).

Next, we tested the *in vivo* pseudouridylation of 28S rRNA sequences lacking U3752. Small fragments of the human 28S rRNA from position G3729 to G3759 either containing (28S-WT) or lacking (28S-U3752del) U3752 were transiently expressed in mouse cells by using the pW(Xb/Xh) mouse ribosomal minigene expression plasmid (Fig. 4B; Hadjiolova et al. 1994). Since the ribosomal minigene is transcribed by RNA polymerase I in the nucleolus, rRNA sequences imbedded within the nascent minigene transcript can undergo snoRNP-mediated modifications (Ganot et al. 1997a). Primer extension analysis after CMCT treatment clearly demonstrated that synthesis of Ψ3749 was largely inhibited or fully abolished in the 28S-U3752del transcript (Fig. 4B, lane 3), but it was unaffected in the control 28S-WT RNA (Fig. 4B, lane 1). Importantly, both 28S-U3752del and 28S-WT RNAs were



**Figure 4.** Structural requirements of the dual pseudouridylation guide activity of the 5' hairpin of SNORA53. (A) In vivo pseudouridylation of 28S rRNA with a mutant SNORA53 RNA. (Top left panel) A predicted interaction of endogenous 28S rRNA with an ectopically expressed mutant SNORA53 RNA carrying an extra A (A+, in red) inserted between U13 and A14. The 28S-U3752 residue is underlined. Please note that 28S-U3751 could also bulge out in configuration A, but the upstream antisense element of SNORA53-U13+A is able to base-pair with both U3751 and U3752. Accumulation of SNORA53-U13+A in SNORA53-KO cells was verified by RNase mapping. As controls, accumulation of endogenous SNORA53 and U85 RNAs was also tested. (Bottom panel) Pseudouridylation of 28S rRNA was monitored by primer extension mapping in HAP1 and SNORA53-KO cells lacking or expressing SNORA53-U13+A. For other details, see the legend for Figure 2E. (B) In vivo pseudouridylation of transiently expressed mutant 28S rRNA (28S-U3752del) sequences lacking U3752 in mouse 3T3 cells. (Top panel) A predicted interaction of endogenous mouse SNORA53 with 28S-U3752del rRNA and schematic structure of the mouse pW(Xb/Xh) expression construct are shown. The mouse RNA polymerase I promoter (Pol I prom) and terminator (term), the 5'- and 3'-terminal regions of the 5' and 3' external transcribed sequences (ETS; gray boxes), and a fragment of the chloramphenicol acetyltransferase (CAT) gene are indicated (Hadjiolova et al. 1994). The nucleotide sequences of wild-type (28S-WT) and mutant (28S-U3752del) 28S rRNA fragments with the known ribosomal Ψs are shown. (Bottom panel) Pseudouridylation of the expressed 28S-WT and 28S-U3752del ribosomal minigene transcripts was monitored by primer extension analysis. (C) Correct positioning of the H box relative to the selected substrate Us is essential for the dual guide activity of the 5' hairpin of SNORA53. (Left panel) Interaction of the descending strand of the 5' hairpin of SNORA53 with 28S rRNA positioning U3747 and U3749 for pseudouridylation. A putative H-box-like motif is highlighted in the dashed box. The A149 residue was deleted in SNORA53-A149del and replaced for C in SNORA53-A149C. (Middle panel) Accumulation of ectopically expressed SNORA53-A149C and SNORA53-A149del RNAs and endogenous U85 scaRNA in SNORA53-KO cells was measured by RNase mapping. (Right panel) Primer extension mapping of 28S rRNA pseudouridylation in SNORA53-KO cells lacking or expressing SNORA53-A149C or SNORA53-A149del RNAs.

faithfully pseudouridylated at Ψ3747 and other known pseudouridylation sites: Ψ3737, Ψ3741, and Ψ3743. Thus, stabilization of the upstream antisense target stem (UATS) of SNORA53 in configuration A, either by removal of the bulged U3752 ribosomal residue or by insertion of a complementary A residue into the upstream antisense sequence of SNORA53, promoted synthesis of Ψ3747 and prevented formation of Ψ3749, suggesting that accurate function of the dual guide loops necessitates comparable

thermodynamic stabilities of the alternative antisense target base-pairing interactions.

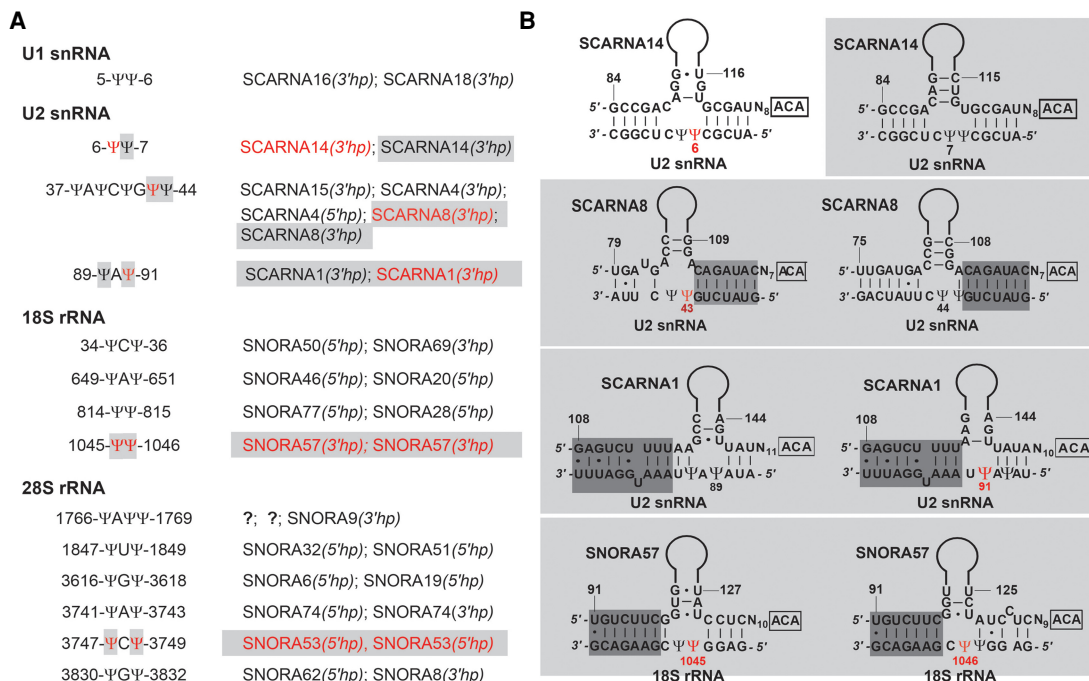
In most H/ACA guide RNA–substrate RNA interactions, the selected target Us are separated from the H or ACA boxes by 14- to 15-nt-long guide RNA sequences (Supplemental Fig. S2). In the proposed alternative SNORA53–28S rRNA interactions, U3747 is positioned at the optimal 14-nt distance from the H box, but U3749 is separated by 16 nt, which is a less common

configuration (Fig. 4C, left panel). Besides the canonical H box (150-AGAGAA-155), SNORA53 contains another overlapping H-box-like sequence (149-AAGAGA-154) that is separated only by 15 nt from U3749. However, disruption of the potential dyskerin binding capacity of the H-like motif through conversion of A149 into C had no effect on the synthesis of 28S-Ψ3747 and 28S-Ψ3749, suggesting that dyskerin bound to the canonical H box of SNORA53 can convert both U3747 and U3749 into Ψ (Fig. 4C, lane 3). More unexpectedly, deletion of A149 that reduced the distance between the H box of SNORA53-A149del and the U3747 target residue from 14 to 13 nt fully abolished pseudouridylation of U3747 (Fig. 4C, lane 5), contrary to the fact that 13-nt-long spacers separating target Us from the H or ACA boxes are relatively frequent (Supplemental Fig. S2). In addition to the H and ACA motifs, Cbf5/dyskerin also interacts with the minor groove of the proximal stem (PS) and the downstream antisense target stem (DATS) through its PUA (pseudouridine synthase and archaeosine transglycosylase) and catalytic domains, respectively (Fig. 1B). The PS-DATS joint helical stack seems to serve as a molecular ruler measuring the distance between the target Us and H/ACA boxes. In the SNORA53–28S rRNA interaction, the PS and DATS helices are separated by only one unpaired

loop nucleotide (U141) (Fig. 4C, left panel). This may limit the structural plasticity of the PS-DATS helical stack of SNORD53-A149del required for correct positioning of U3747 for pseudouridylation.

*Human H/ACA pseudouridylation loops with potential dual guide capacities*

To identify additional human H/ACA pseudouridylation loops with potential dual guide capacities, we collected the human spliceosomal and ribosomal Ψs that are located next to each other (ΨΨ) or are separated by 1 nt (ΨNΨ) and listed them together with the corresponding H/ACA hairpins assigned for their site-specific selection (Fig. 5A; Lestrade and Weber 2006; Jorjani et al. 2016; Taoka et al. 2018). This compilation confirmed that most Ψs, even in densely pseudouridylated regions, are selected individually by dedicated pseudouridylation loops. Nevertheless, inspection of the listed H/ACA hairpins revealed that the 3'-terminal pseudouridylation loops of SCARNA14, SCARNA8, and SCARNA1, which had been earlier assigned for pseudouridylation of the U2 snRNA at Ψ7, Ψ44, and Ψ89, respectively (Darzacq et al. 2002; Kiss et al. 2004; Schattner et al. 2006), might also direct synthesis of the neighboring “orphan” Ψ6, Ψ43, and



**Figure 5.** Human pseudouridylation guide loops with possible dual guide capacities. (A) Closely located human spliceosomal and ribosomal Ψs are listed together with the corresponding pseudouridylation guide RNA hairpins (5'hp and 3'hp) connected to their synthesis. Hairpins newly implicated in dual pseudouridylation reactions and their predicted target Ψs are highlighted in red. Participation of the shaded H/ACA hairpins in the synthesis of their predicted target Ψs has been experimentally validated in this study. (B) Potential alternative base-pairing interactions of H/ACA pseudouridylation guide loops with their target RNAs positioning consecutive ribosomal and spliceosomal Us for pseudouridylation. Direction of the synthesis of Ψ7, Ψ44, and Ψ89 in the U2 snRNA by human SCARNA14, SCARNA8, and SCARNA1, respectively, has been proposed earlier (Darzacq et al. 2002; Kiss et al. 2004; Schattner et al. 2006). The newly predicted target Ψs are highlighted in red. The functional interactions shaded in light gray have been experimentally validated in this work. Base-pairing interactions common in the alternative functional configurations are shaded in dark gray. For the interaction of SNORA53 and 28S rRNA, see Figure 3D.

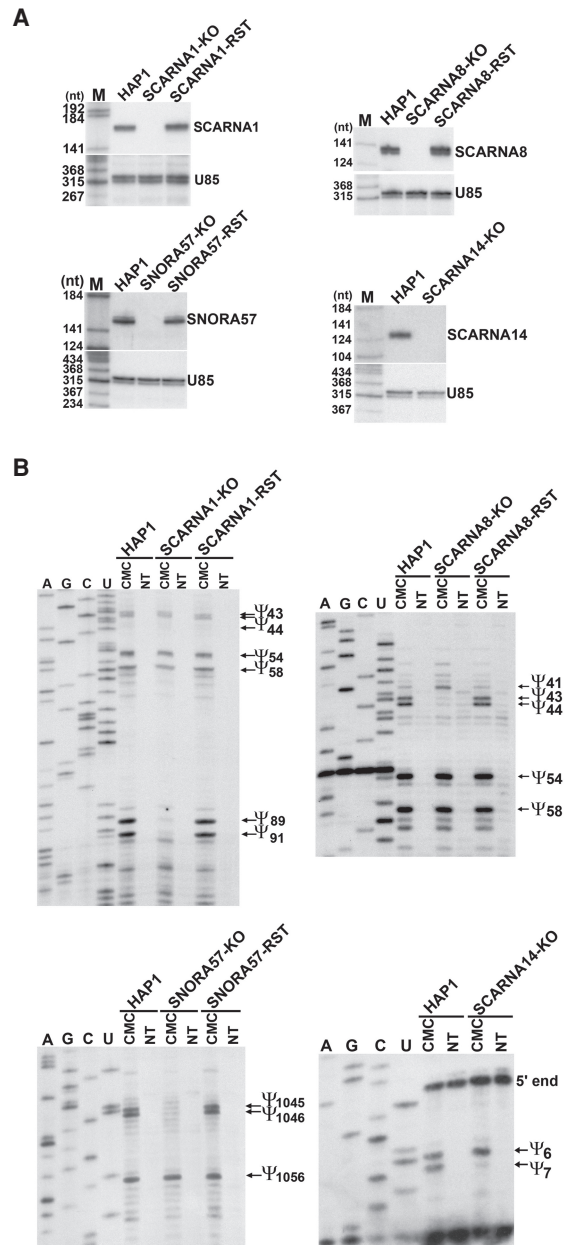
$\Psi$ 91 residues through formation of alternative base-pairing interactions (Fig. 5B). In fact, the 3' hairpin of the *Xenopus* equivalent of human SCARNA8, pugU2-34/44, has already been reported to direct pseudouridylation of the U2 snRNA at both U43 and U44 (Deryusheva and Gall 2013). Finally, re-examination of human orphan H/ACA RNAs revealed that the 3' hairpin of SNORA57 carries a putative pseudouridylation loop that, in principle, could direct synthesis of both  $\Psi$ 1045 and  $\Psi$ 1046 in the 18S rRNA. Thus, our search identified four novel putative dual pseudouridylation guide loops amenable for experimental testing.

*The 3' hairpins of SCARNA1, SCARNA8, and SNORA57 direct synthesis of two adjacent  $\Psi$ s in the U2 and 18S RNAs*

To examine the *in vivo* pseudouridylation guide competences of SCARNA1, SCARNA8, SCARNA14, and SNORA57, the single-copy genes of these intronic RNAs were deleted with CRISPR-Cas9 genome editing in human HAP1 cells (Supplemental Fig. S3). Disruption of the accumulation of each target H/ACA RNA in the CRISPR-treated KO cells was confirmed by RNase A/T1 mapping (Fig. 6A), and pseudouridylation of the predicted target Us in U2 and 18S RNAs was monitored by primer extension analysis (Fig. 6B). Abrogation of SCARNA1 and SCARNA8 accumulation fully obliterated pseudouridylation of the U2 snRNA at the U89/U91 and U43/U44 residues, respectively. Lacking U2- $\Psi$ 43 formation in the absence of SCARNA8 was somewhat unexpected, because formation of *Xenopus* U2- $\Psi$ 43 had been reported to be a redundant reaction implicating both SCARNA8 and a stand-alone  $\Psi$  synthase, Pus1 (Deryusheva and Gall 2017). Disruption of SNORA57 accumulation abolished  $\Psi$ 1045 and  $\Psi$ 1046 synthesis in the 18S rRNA. Importantly, restoration of SCARNA1, SCARNA8, and SNORA57 accumulations in the corresponding KO cells re-established synthesis of U2- $\Psi$ 89/ $\Psi$ 91, U2- $\Psi$ 43/ $\Psi$ 44, and 18S- $\Psi$ 1045/ $\Psi$ 1046, corroborating the notion that the 3'-terminal pseudouridylation loops of SCARNA1, SCARNA8, and SNORA57 possess dual guide activities. Finally, disruption of SCARNA14 expression demolished formation of  $\Psi$ 7 in the human U2 snRNA, but it had no impact on the synthesis of  $\Psi$ 6, excluding the proposed dual guide capacity of the 3' hairpin of SCARNA14 and confirming the idea that it directs synthesis of U2- $\Psi$ 7 (Schattner et al. 2006).

*Alternative base-pairings with the H/ACA antisense sequences are the major determinants of dual guide activities*

We have already demonstrated above that the dual pseudouridylation guide activity of the 5' hairpin of SNORA53 can be confined to the synthesis of one  $\Psi$  through structural manipulation of its DS region (Fig. 3). These results seem to support the functional importance of the DS regions of dual pseudouridylation loops. The tested CUU133del and CU132GA mutations of SNORA53 (Fig.



**Figure 6.** The 3'-terminal pseudouridylation loops of human SCARNA1, SNORA57, and SCARNA8 possess dual guide capacities. (A) Expression of SCARNA1, SNORA57, SCARNA8, and SCARNA14 measured by RNase mappings. After deletion of the *SCARNA1*, *SNORA57*, *SCARNA8*, and *SCARNA14* genes in HAP1 cells (Supplemental Fig. S3), each KO cell line was transformed by pIRESpuro/GL plasmids expressing SCARNA1, SCARNA8, or SNORA57 (RST cells). (B) Primer extension mapping of U2 snRNA and 18S rRNA pseudouridylation in the absence and presence of SCARNA1, SCARNA8, SNORA57, and SCARNA14 as indicated above the lanes. For other details, see the legend for Figure 2E.

3D), while promoting formation of one of the two alternative active configurations of SNORA53, prevented formation of the other one. To further understand the



importance of the DS region in the activities of H/ACA dual pseudouridylation guide loops, a series of mutant SNORA57 and SCARNA1 RNAs were expressed in SNORA57-KO or SCARNA1-KO cells, and their pseudouridylation guide capacities were determined (Fig. 7).

Through increasing the stability of the DS regions, the 124-UUC-126 deletion and the U125-to-A conversion in SNORA57 were predicted to promote formation of active configuration A or B and synthesis of 18S- $\Psi$ 1045 or 18S- $\Psi$ 1046, respectively (Fig. 7A). However, while the 124-UUC-126 deletion was predicted to interfere with formation of active configuration B, the U125A mutation of SNORA57 did not exclude the formal possibility of folding into active configuration A. Upon expression in SNORA57-KO cells (Fig. 7B), the SNORA57-UUC126del RNA proved to be efficient in 18S- $\Psi$ 1045 synthesis, but failed to govern 18S- $\Psi$ 1046 formation (Fig. 7B, lane 5). In contrast, the SNORA57-U125A RNA retained its dual guide ability to direct synthesis of both  $\Psi$ 1045 and  $\Psi$ 1046 (Fig. 7B, lane 7).

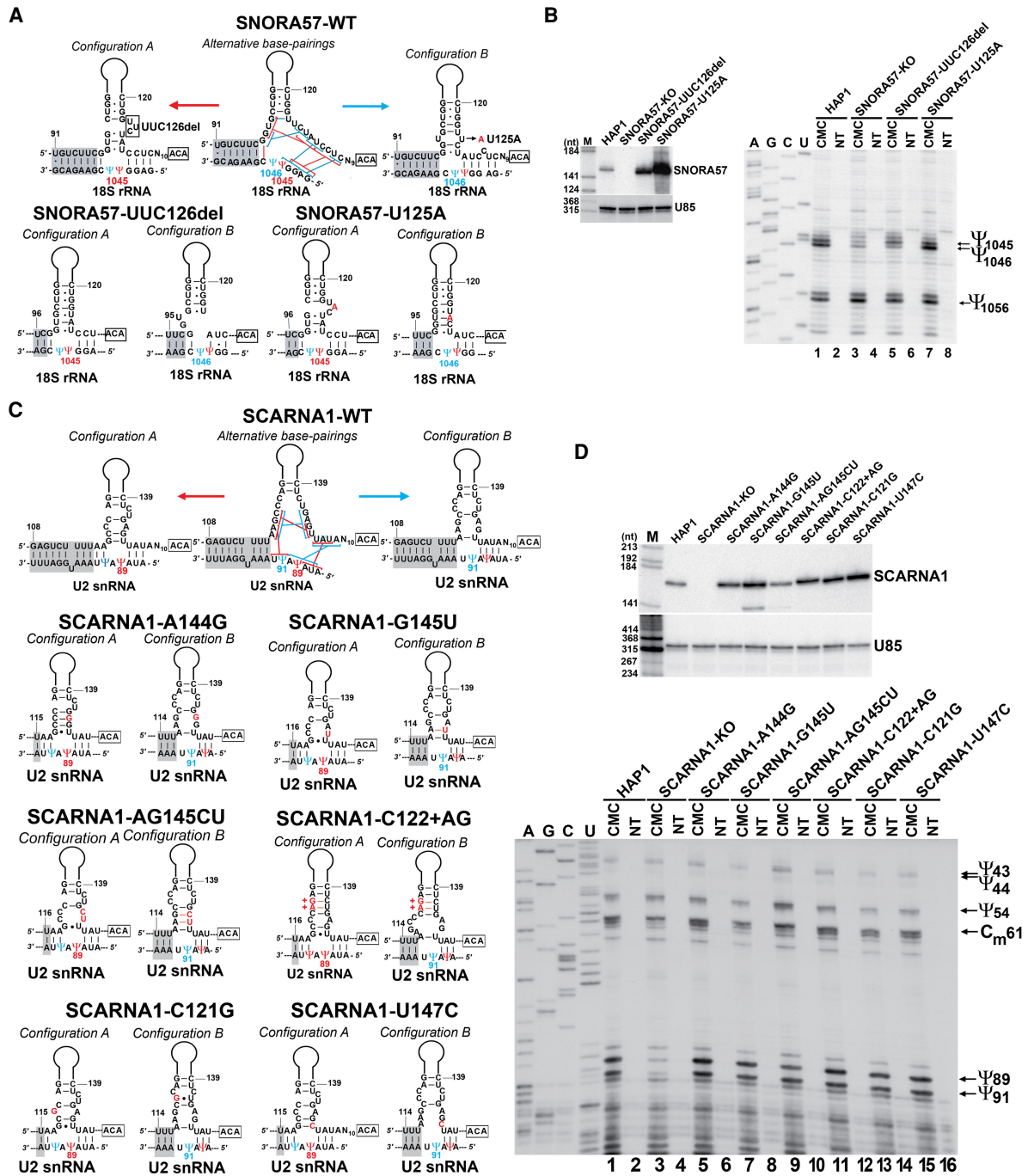
Next, we further assessed the contribution of the distal stem (DS) region to the dual pseudouridylation guide activity of the 3' hairpin of SCARNA1. A series of mutant SCARNA1 RNAs carrying nucleotide alterations slightly increasing or reducing the stability of the DS region were expressed in SCARNA1-KO cells (Fig. 7C). Importantly, none of the tested mutations were expected to fully exclude folding into any of the alternative active configurations of the 3' hairpin of SCARNA1. The A144-to-G transition in SCARNA1-A144G was expected to promote DS stability in configuration A directing U2- $\Psi$ 89 synthesis and to slightly destabilize the DS region in configuration B governing U2- $\Psi$ 91 synthesis. In contrast, the G145-to-U (G145U), the 144-AG-145-to-CU (AG145CU), and the C121-to-G (C121G) base conversions in the DS region of SCARNA1 were predicted to support formation of configuration B and to disfavor folding into configuration A. Insertion of AG dinucleotides between the C122 and A123 residues of SCARNA1 (C122+AG) was predicted to increase the stem stability in both configurations A and B, but leaving six unpaired nucleotides highly destabilizing the proximal part of the DS region in configuration B. Finally, replacement of U147 for a C residue was predicted to disrupt the A117:U146 base-pairing interaction closing the pseudouridylation loop in configuration B. However, when expressed in SCARNA1-KO cells (Fig. 7D, top panel), the mutant SCARNA1 RNAs retained dual guide activity directing efficient synthesis of both  $\Psi$ 89 and  $\Psi$ 91 in the U2 snRNA (Fig. 7D, bottom panel), confirming that structural alterations in the distal stem region, if they do not exclude formation of one functional configuration, have no significant impact on the dual pseudouridylation guide capacity of the 3'-terminal guide loop of SCARNA1. These results, together with the observations that minor structural alterations in the upstream antisense target stem (UATS) could completely abolish the dual guide activity of the 5' hairpin of SNORA53 (Fig. 4), suggest that the alternative base-pairing interactions formed between the antisense sequences and the

target RNA are the major determinants of the activity of dual pseudouridylation guide loops.

## Discussion

Post-transcriptional pseudouridylation of eukaryotic spliceosomal snRNAs and rRNAs is essential for efficient and correct pre-mRNA splicing and protein synthesis. Site-specific synthesis of  $\sim$ 130  $\Psi$ s in human snRNAs and rRNAs is achieved mostly by H/ACA pseudouridylation guide RNPs. In contrast to stand-alone  $\Psi$  synthases that recognize local RNA structures and frequently synthesize several closely placed  $\Psi$ s (Spenkuch et al. 2014), the H/ACA RNPs have been believed to direct synthesis of a single  $\Psi$  in their target sequences. However, here we have demonstrated that the pseudouridylation guide loops in the 5'-terminal hairpin of human SNORA53 and in the 3' hairpins of human SNORA57, SCARNA1, and SCARNA8 H/ACA RNAs are able to support synthesis of two consecutive  $\Psi$ s ( $\Psi\Psi$  or  $\Psi N\Psi$ ) through formation of alternative base-pairing interactions with their target RNAs, breaking with the notion that pseudouridylation guide RNPs lack site promiscuity. Apparently, synthesis of two  $\Psi$ s is not a consequence of defective or inaccurate functions of the dual guide loops of human SNORA53, SNORA57, SCARNA1, and SCARNA8, because their substrate Us are pseudouridylated with practically 100% efficiency (Reddy and Busch 1988; Taoka et al. 2018). Even more telling, all guide RNA nucleotides implicated in inter- and intramolecular base-pairings promoting alternative target RNA interactions are perfectly conserved during evolution (Supplemental Figs. S1, S4). These strong evolutionary constraints clearly demonstrate the functional importance of these nucleotides in correct and efficient dual pseudouridylation reactions.

In principle, synthesis of two  $\Psi$ s by dual pseudouridylation guide loops could be accomplished through one or two independent target binding events, depending on whether the guide RNP dissociates from or remains associated with its target RNA after the first pseudouridylation reaction. Although we cannot unambiguously distinguish between these two possibilities, we prefer the "one binding–two pseudouridylation" model. Continuous guide RNA–target RNA association could increase the efficiency of the two subsequent pseudouridylation reactions directed by the same guide loop. In this context, it is noteworthy that the proposed alternative guide substrate configurations always share common interacting core domains that could continuously anchor the guide RNP to the target RNA without interfering with the structural rearrangements supporting the two pseudouridylation reactions (Figs. 3D, 5B, shaded helices). In the "two-binding model," after dissociation of the guide RNP following the first pseudouridylation reaction, the dual pseudouridylation loop region of the released guide RNP is expected to undergo structural rearrangements to conform to the second pseudouridylation reaction. Such conformational changes in the guide RNA could



**Figure 7.** Mutational analyses of SNORA57 and SCARNA1. (A) Predicted base-pairing interactions of wild-type (WT) and mutant SNORA57 RNAs with 28S rRNA in configurations A and B directing synthesis of  $\Psi_{1045}$  and  $\Psi_{1046}$ , respectively. (B) Accumulation of mutant SNORA57 RNAs in SNORA57-KO cells and pseudouridylation of endogenous 28S rRNA. (Left panel) Accumulation of SNORA57-UUC126del and SNORA57-U125A RNAs was detected by RNase protection with sequence-specific probes. (Right panel) The pseudouridylation state of 18S rRNA in wild-type HAP1 and SNORA57-KO cells lacking or expressing the indicated mutant SNORA57 RNAs was measured by primer extension. (C) Predicted interactions of wild-type and mutant SCARNA1 RNAs with U2 snRNA directing pseudouridylation of U89 (configuration A) or U91 (configurations B). The nucleotide alterations in the mutant SCARNA1 RNAs are highlighted in red. (D) In vivo pseudouridylation activity of mutant SCARNA1 RNAs. (Top panel) Expression of mutant SCARNA1 RNAs in SCARNA1-KO cells was monitored by RNase protection. The lower band in the SCARNA1-G145U lane likely represents a degradation product of SCARNA1-G145U. (Bottom panel) Primer extension mapping of U2 snRNA pseudouridylation at U89 and U91 in the absence (SCARNA1-KO) or presence of wild-type (HAP1) and mutant SCARNA1 RNAs as indicated above the lanes. For other details, see the legend for Figure 2E.

be governed by the distal stem region through slightly altering and repositioning the single-stranded antisense sequences. However, the dual guide loops of SNORA53, SNORA57, and SCARNA1 are topped by loosely structured and thermodynamically unstable distal stem regions, arguing against a role in shaping their pseudouridylation loops (Figs. 3D, 5B). In line with this conclusion, nucleotide alterations in the distal stem region had no significant impact on the dual pseudouridylation capacity of the 3'-terminal guide loops of SCARNA1 and SNORA57 (Fig. 7). Instead, our data supported the idea that correct remodeling of the target RNA–guide RNA three-way helical junctions is dictated predominantly by formation of alternative antisense target interactions (Fig. 7).

Dissection of the molecular mechanisms driving the appropriate guide RNA–target RNA structural rearrangements supporting the correct and efficient function of dual pseudouridylation guide RNPs remains a challenge for the future. In the secondary structures of H/ACA RNAs defined by minimal free energy calculation, the pseudouridylation loops frequently fold into partially closed structures (see Fig. 3A; Supplemental Fig. S1; data not shown), suggesting that the associated H/ACA proteins contribute to the correct folding of functionally competent pseudouridylation loops. Indeed, the catalytic surface of Cbf5 loosely and nonspecifically associates with the pseudouridylation loop regions of H/ACA RNAs (Li and Ye 2006). In the substrate-bound H/ACA RNP, the conserved arginine-rich thumb region of Cbf5 interacts with the substrate RNA locking the target U at the active site of the RNP (Duan et al. 2009). Gar1 interacts with the catalytic domain of archaeal Cbf5 and controls transitions between the closed and open states of the Cbf5 thumb loop (Duan et al. 2009; Wang et al. 2015). Although it seems that archaeal Gar1 makes no direct RNA contacts, the N- and C-terminal parts of eukaryotic Gar1 contains glycine–arginine-rich RGG domains (Girard et al. 1992). RGG domains are frequent in RNA-binding proteins and they play accessory roles in RNA binding (Thandapani et al. 2013). The RGG domains of yeast Gar1 have been found to facilitate pseudouridylation of the bound substrate RNA and to inhibit its release, raising the possibility that eukaryotic Gar1 may directly interact with target RNAs though its RGG domains (Trucks et al. 2021).

It seems that dual pseudouridylation guide loops are not confined to vertebrate H/ACA RNAs. Genetic depletion of yeast snR82 abolished pseudouridylation of the 27S rRNA at U2350 and also at the nearby U2348 residue (Schattner et al. 2004). While synthesis of  $\Psi$ 2350 was predicted to be guided by the 3'-terminal pseudouridylation loop of snR82, snR82-dependent synthesis of  $\Psi$ 2348 could not be reasoned, as U2348 was imbedded in the downstream antisense target stem (DATS) of snR82 directing  $\Psi$ 2350 formation (Supplemental Fig. S5A, configuration A). However, the 3'-terminal pseudouridylation loop of snR82 can form an alternative interaction with the 27S rRNA to position U2348 for pseudouridylation (configuration B). More recently, the 3' hairpin of the archaeal

*Haloferax volcanii* sR-h45 H/ACA RNA has been shown to govern pseudouridylation of the 23S rRNA at both  $\Psi$ 1940 and  $\Psi$ 1942 (Majumder et al. 2020). The investigators proposed that the dual guide activity of the 3' hairpin of sR-h45 was due to its unusual interaction with 23S sequences, placing four unpaired ribosomal nucleotides, 1940-UAUG-1943, including the U1940 and U1942 substrate Us, into the pseudouridylation pocket of the 3'-terminal pseudouridylation loop (Supplemental Fig. S5B, configuration A). In fact, pseudouridylation pockets can accommodate more than two unpaired nucleotides at the base of the distal stem (De Zoysa et al. 2018). The investigators suggested that in this context, Cbf5 bound to the ACA box of sR-h45 was able to convert both the canonically positioned U1940 and the downstream unpaired U1942 into  $\Psi$ . Instead, we propose that sR-h45 forms an alternative interaction with 23S rRNA in order to specifically select U1942 for pseudouridylation (configuration B). Importantly, the results of the mutational analysis presented by the investigators are consistent with the existence and functional significance of configuration B supporting  $\Psi$ 1942 synthesis. For instance, the presence of U1940 in the 23S rRNA was required for the synthesis of  $\Psi$ 1942, supporting the importance of U1940 in formation of DATS in configuration B. In contrast, replacement of the U1942 residue failed to affect U1940 pseudouridylation, as formation of functional configuration A is independent from U1942. Based on these observations, we propose that the 3' hairpins of yeast snR82 and *Haloferax* sR-h45 RNAs carry dual pseudouridylation guide loops directing synthesis of two consecutive  $\Psi$ s in the yeast 25S and archaeal 23S rRNAs through formation of alternative base-pairing interactions. Thus, H/ACA guide RNA acrobatics seems to be an evolutionarily widespread strategy for synthesis of two consecutive  $\Psi$ s in vertebrate, yeast, and archaeal spliceosomal and ribosomal RNAs.

How many human H/ACA RNAs contain dual pseudouridylation guide loops? Apparently, synthesis of isolated  $\Psi$ s dispersed in ribosomal and spliceosomal sequences is directed by specific pseudouridylation guide loops. Although it is unlikely that our search has overlooked many H/ACA RNAs with dual pseudouridylation guide loops (Fig. 5), it is possible that some pseudouridylation loops assigned for synthesis of  $\Psi$ s in heavily pseudouridylated sequence environments possess dual guide capacity, as efficient synthesis of some  $\Psi$ s might be accomplished by cooperative actions of single and dual pseudouridylation guide loops. Nevertheless, demonstration that four H/ACA RNAs carry dual pseudouridylation loops already shows that synthesis of at least 6% of the human ribosomal and spliceosomal  $\Psi$ s is directed by dual guide loops, indicating that formation of closely located  $\Psi$ s by dual pseudouridylation loops is a relatively frequent strategy of H/ACA RNPs.

In summary, guide RNA acrobatics supporting accurate and efficient conversion of nearby substrate Us (UU or UNU) into  $\Psi$  is a novel, unexpected, relatively frequent, and, most probably, evolutionarily widespread strategy of box H/ACA pseudouridylation guide RNPs.

## Materials and methods

### General procedures

Unless stated otherwise, all DNA and oligonucleotide manipulations were performed according to standard laboratory procedures. Oligodeoxynucleotides were purchased from Eurofins MWG Operon. Human HAP1 cells were grown in Iscove's modified Dulbecco's medium (IMDM) supplemented with 10% (v/v) fetal bovine serum, 1 mM sodium pyruvate (Gibco), and 100 µg/mL penicillin and streptomycin (Gibco). Cells were transfected by electroporation with Gene Pulser (Bio-Rad) using exponential decay at settings 270 V and 950 µF.

### Expression plasmids

Human HAP1 genomic fragments encompassing the *SNORA53*, *SNORA57*, *SCARNA1*, and *SCARNA57* intronic H/ACA RNA genes were PCR-amplified by using sequence-specific primers encompassing the *Cl*I and *X*hoI restriction sites (Supplemental Table S1A). The resulting DNA fragments were inserted into the *Cl*I and *X*hoI sites of the pBST vector and the pCMV/globin expression vector (Darzacq et al. 2002). The H/ACA RNA gene together with its host globin gene was PCR-amplified and inserted into the *E*coRI and *N*otI sites of pIRESpuro2 expression plasmid (Addgene). Mutant guide RNA genes were generated by the megaprimer PCR amplification approach using appropriately designed mutagenic oligonucleotide primers and wild-type *sno*/*sca*RNA gene-containing recombinant pBST plasmids as templates. For nucleolar expression of mouse ribosomal minigene transcripts, appropriate oligonucleotides encoding the wild-type and mutant fragments of the human 28S rRNA from G3729 to G3759 were annealed and inserted into the *X*baI and *X*hoI sites of the pW(Xb/Xh) expression plasmid (Hadjiolova et al. 1994). The resulting pW-28SWT and pW-28S-U3752del plasmids were transfected into 3T3 mouse cells with JetPrime transfection reagent (Polyplus Transfection SA). Two days after transfection, cellular RNA was extracted. The identities of all expression constructs were verified by sequence analysis.

### Excision of HAP1 intronic H/ACA RNA genes by CRISPR/Cas9

To generate synthetic DNA encoding appropriately designed sgRNAs, appropriately designed oligodeoxynucleotides (Supplemental Table S1B) were annealed and inserted into the *B*bsI site of pSpCas9(BB)-2A-Puro (PX459) V2.0 by overnight incubation with T4 DNA ligase (NEB) at room temperature. The resulting pX459/sgRNA expression vectors were transfected into HAP1 cells. Twenty-four hours after transfection, cells were transferred into puromycin-containing (1.5 µg/mL) selective medium and grown for 48 h before establishment of monoclonal stable cell lines by limited dilution under nonselective conditions.

### Genomic PCR and RT-PCR

Cellular DNA was purified with GenElute mammalian genomic DNA miniprep kit (Sigma), and the targeted genomic fragments were PCR-amplified with appropriately designed oligonucleotide primers (Supplemental Table S1C) and Phusion DNA polymerase (Thermo Scientific). cDNAs were generated with AMV reverse transcriptase (Promega) using total cellular RNAs as templates and sequence-specific oligonucleotide primers. The PCR-amplified genomic and cDNA fragments were analyzed by electrophoresis on 1% agarose gels.

### RNA purification and primer extension mapping of pseudouridylation

The guanidine thiocyanate-phenol-chloroform purification procedure was used for RNA extraction from human HAP1 and HAP1-derived cells as described before (Kiss and Jády 2004). Cellular RNAs were reacted with N-cyclohexyl-N'-(2-morpholinoethyl)carbodiimide metho-p-toluenesulfonate (CMCT) and subjected to primer extension analyses (Bakin and Ofengand 1998; Kiss and Jády 2004). The oligonucleotide primers used for pseudouridylation mapping of 18S and 28S rRNAs, U2 snRNA, and mouse ribosomal minigene transcripts are listed in Supplemental Table S1D.

### RNase A/T1 protection analysis

RNase A and T1 protection analysis was performed according to Goodall et al. 1990. For mapping of endogenous and ectopically expressed wild-type and mutant *SNORA53*, *SNORA57*, *SCARNA1*, *SCARNA8*, and *SCARNA14* RNAs, complementary RNA probes were synthesized in vitro by T3 RNA polymerase (Promega), using 30 µCi [ $\alpha$ -<sup>32</sup>P]CTP and *Cl*I-linearized pBST-*SNORA53*, pBST-*SNORA57*, pBST-*SCARNA1*, pBST-*SCARNA8*, pBST-*SCARNA14*, and pBST-U85 plasmids as templates. Before hybridization with test RNAs, each probe was purified on a 6% denaturing gel. The protected fragments of the probe RNAs were analyzed on 6% sequencing gels.

## Competing interest statement

The authors declare no competing interests.

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*Author contributions:* B.E.J., A.K., and D.M. carried out experiments. B.E.J. and T.K. conceived and designed experiments and analyzed results. T.K. wrote the manuscript.

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