Human intron-encoded Alu RNAs are processed and packaged into Wdr79-associated nucleoplasmic box H/ACA RNPs

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Alu repetitive sequences are the most abundant short interspersed DNA elements in the human genome. Full-length Alu elements are composed of two tandem sequence monomers, the left and right Alu arms, both derived from the 7SL signal recognition particle RNA. Since Alu elements are common in protein-coding genes, they are frequently transcribed into pre-mRNAs. Here, we demonstrate that the right arms of nascent Alu transcripts synthesized within pre-mRNA introns are processed into metabolically stable small RNAs. The intron-encoded Alu RNAs, termed AluACA RNAs, are structurally highly reminiscent of box H/ACA small Cajal body (CB) RNAs (scaRNAs). They are composed of two hairpin units followed by the essential H (AnAnnA) and ACA box motifs. The mature AluACA RNAs associate with the four H/ACA core proteins: dyskerin, Nop10, Nhp2, and Gar1. Moreover, the 3'-hairpin of AluACA RNAs carries two closely spaced CB localization motifs, CAB boxes (UGAG), which bind Wdr79 in a cumulative fashion. In contrast to canonical H/ACA scaRNPs, which concentrate in CBs, the AluACA RNPs accumulate in the nucleoplasm. Identification of 348 human AluACA RNAs demonstrates that intron-encoded AluACA RNAs represent a novel, large subgroup of H/ACA RNAs, which are apparently confined to human or primate cells.

[Keywords: Alu elements; AluACA RNAs; box H/ACA RNAs; intron-encoded RNAs; RNA processing]

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Mammalian cells express thousands of non-protein-coding RNAs [ncRNAs], which, in the form of ribonucleoproteins [nRNP], function in all steps of gene expression [Mattick and Makunin 2006; Wilusz et al. 2009; Taft et al. 2010]. Functional ncRNPs are produced through diverse and complex biosynthetic processes, which usually include the synthesis and processing of precursor ncRNAs, the assembly of mature ncRNAs with RNP proteins, and finally, the targeting of nascent ncRNPs to the appropriate subcellular compartments where they function.

Box H/ACA RNAs represent an abundant and evolutionarily conserved group of ncRNAs [for reviews, see Meier 2005; Terns and Terns 2006; Matera et al. 2007; Kiss et al. 2010; Watkins and Bohnsack 2012]. Most box H/ACA RNAs function as guide RNAs in site-specific pseudouridylation of tRNAs and spliceosomal snRNAs [Ganot et al. 1997a; Ni et al. 1997; Darzacq et al. 2002] but also participate in pre-rRNA processing, telomere synthesis, and probably other processes [Morrissey and Tollervey 1993; Mitchell et al. 1999; Chen et al. 2000; Atzorn et al. 2004; Fayet-Lebaron et al. 2009]. The majority of human H/ACA RNAs are processed from pre-mRNA introns, and the mature RNAs are typically composed of two 55- to 60-nucleotide (nt)-long hairpins connected and followed by short single-stranded hinge and tail sequences [Fig. 1A; Kiss et al. 1996; Ganot et al. 1997b]. While the hinge region contains the conserved H box [consensus AgAnnA], the 3'-terminal tail carries the ACA or, less frequently, the AUA box motif located always 3 nt before the RNA end [Balakin et al. 1996; Ganot et al. 1997b]. The bipartite guide sequences occupy the rising and descending strands of the internal pseudouridylation loop in the 5’ and/or 3’ hairpin [Ganot et al. 1997a]. Each hairpin unit associates with four H/ACA core proteins: Nhp2, Nop10, Gar1, and the pseudouridine synthase dyskerin [for reviews, see Reichow et al. 2007; Ye 2007; Li 2008; Kiss et al. 2010; Watkins and Bohnsack 2012]. In the H/ACA RNP, base-pairing of the H/ACA guide sequences
with the complementary target RNA places the unpaired substrate uridine at the active site of dyskerin (Li and Ye 2006; Duan et al. 2009; Liang et al. 2009). Optimally, the so-called “pseudouridylation pocket” of the guide RNA is located 14 nt upstream of the H or ACA box that is a fundamental coordinator of correct dyskerin binding. Box H/ACA RNPs mediating rRNA pseudouridylation accumulate in the nucleolus and are termed the small nucleolar RNPs (snoRNPs), while those H/ACA RNPs that

direct snRNA pseudouridylation reside in the nucleoplasmic Cajal bodies (CBs) and are called the small CB RNPs (scaRNPs) (Darzacq et al. 2002; Kiss et al. 2006). The terminal loop of the 5’ and/or 3’ hairpin of scaRNAs carries the CAB box (consensus, uAAG) that binds Wdr79 [also known as Wrap53] and targets scaRNPs into CBs (Richard et al. 2003; Tycowski et al. 2009; Venteicher et al. 2009).

Alu elements are abundant primate-specific repetitive sequences that are present in >1 million similar but not identical copies in the human genome (Lander et al. 2001). Full-length Alu elements are ~300 nt long and are composed of two tandem monomer units, called the left and right arms, both followed by short A-rich sequences (Fig. 1B; for reviews, see Batzer and Deininger 2002; Dewannieux et al. 2003; Hasler et al. 2007; Kriegs et al. 2007). Due to their preferential interspersion within gene-rich regions, Alu elements are coincidentally transcribed by Pol II into pre-mRNAs (Lander et al. 2001; Chen et al. 2002; Versteeg et al. 2003). While Alu sequences imbedded in the untranslated regions of mRNAs may control mRNA stability and translation, the intronic Alu RNAs are believed to be rapidly degraded after pre-mRNA splicing (Cordaux and Batzer 2009; Ponicsan et al. 2010; Berger and Strub 2011).

Using the internal A and B Pol III-binding motifs inherited from the 7SL gene, some Alu elements are independently transcribed, albeit with low efficiency, into 300- to 400-nt-long Alu RNAs that terminate in at least four consecutive U residues encoded by downstream DNAs (Fig. 1B). The Pol III-specific primary Alu transcripts can be further processed into more stable small cytoplasmic Alu (scAlu) RNAs encompassing only the left monomer (Chang et al. 1996; Sarrowa et al. 1997). Both full-length and processed Alu RNAs interact with the heterodimer of the Spr9 and Srp14 7SL SRP proteins (Bovia et al. 1995; Hsu et al. 1995). Alu RNAs accumulate at very low levels (100–1000 copies per cell), but upon stress induction, cellular Alu concentrations show 10-fold to 20-fold increases, indicating that Alu RNAs play a role in stress response (Hasler et al. 2007; Ponicsan et al. 2010; Berger and Strub 2011). Nuclear full-length Alu RNAs have been reported to repress the transcription of protein-coding genes during heat shock because synthetic Alu RNAs were found to bind Pol II to prevent its interaction with promoter DNA in in vitro assembled preinitiation complexes (Mariner et al. 2008; Yakovchuk et al. 2009). Although both Alu arms seem to bind Pol II, the transcriptional inhibitory activity is limited to the right arm. More recently, aberrant overaccumulation of Alu RNAs in retinal pigment epithelium [RPE] cells has been reported to induce cytotoxicity, leading to the development of geographic atrophy, a severe form of age-related macular degeneration (Kaneko et al. 2011).
that the right arms of nascent intronic Alu RNAs, instead of being degraded after pre-mRNA splicing, can be processed and packaged into metabolically stable box H/ACA RNP. We also demonstrate that the newly discovered Alu-derived H/ACA RNPs, contrary to their association with the CB localization protein Wdr79, accumulate in the nucleoplasm of human cells. Since AluACA RNAs are processed from the primate-specific right Alu monomers, they represent a large group of ncRNAs that are expressed exclusively in human or primate cells.

Results

**Human Wdr79 copurifies with novel Alu-derived H/ACA RNAs**

To identify novel Wdr79-associated RNAs, a Flag-tagged version of human Wdr79 (FL-Wdr79) was transiently expressed in human HeLa cells and purified by immunoprecipitation (IP) with an anti-Flag antibody. RNAs coprecipitated with FL-Wdr79 were converted into cDNAs, and after PCR amplification, their 3′-terminal nucleotide sequences were determined by strand-specific high-throughput sequencing with an Illumina Genome Analyzer [Fasteris AG]. The majority (60%) of the 5.75 million total reads that aligned to the human genome sequence represented contaminating RNAs, mainly spliceosomal snRNAs, box C/D and H/ACA snorRNAs, and rRNAs. The abundant snRNAs and snoRNAs transiently accumulate in CBs and might associate, most probably nonspecifically, with the Wdr79 CB protein. Nevertheless, 36% of the positive reads corresponded to known or predicted H/ACA and C/D scRNA [data not shown]. To our surprise, the remaining 235,000 reads, ~4% of the total reads, represented Alu repetitive sequences. A detailed analysis of the obtained Alu sequences resulted in the identification of 348 novel putative Alu RNAs (Supplemental Table 1). To unambiguously distinguish between closely related Alu sequences, we considered only those Alu reads that were at least 75 nt long and perfectly aligned with a unique region of the reported human genome sequence (University of California at Santa Cruz [UCSC] Genome Browser). Likewise, Alu RNAs represented by <20 reads were also excluded from the list of Alu RNAs. The 3′-terminal sequences of the new Alu RNAs were present within the right arms of dimeric or, in a few cases, monomeric Alu elements, which fell into 19 different subgroups of the major Alu subfamilies (AluS, AluJ, and AluY), indicating that expression and association of Alu RNAs with Wdr79 is not an Alu family-specific affair (Fig. 2A, Supplemental Table 2).

We noticed that the newly discovered Wdr79-associated Alu RNAs carried an ACA or, in two instances, an AUA motif located invariantly 3 nt before the RNA 3′ end (Fig. 2A; Supplemental Table 1). Inspection of the genomic coding sequences of the new Alu RNAs revealed that they always contained a potential H box motif (consensus, AGAgaA) situated 67–76 nt upstream of the 3′ end of the RNA. Computer-mediated structure predictions revealed that the putative H and ACA motifs of all Alu RNAs were folded together by 50- to 60-nt-long hairpins that are highly reminiscent of the 3′ hairpins of canonical box H/ACA RNAs. The predicted 3′-terminal structures of 19 Alu RNAs representing different Alu subgroups are shown in Figure 2B. Finally, we also noticed that the 3′-terminal hairpin of the great majority of Alu RNAs carried two closely spaced potential CB localization signals: the highly conserved proximal CAB box (pCAB) located in the raising strand of the hairpin and the less conserved distal CAB box (dCAB) in the terminal loop (Fig. 2A,B, Supplemental Table 1). These observations together strongly suggested that the newly detected Wdr79-associated Alu RNAs, hereafter called AluACA RNAs, represent a novel, large subclass of human H/ACA RNAs.

**AluACA7 is processed from the second intron of the β-globin pre-mRNA reporter in an H and ACA box-dependent manner**

Apart from a few exceptions (see the Discussion), the genomic Alu DNAs encoding the new AluACA RNAs were found within introns of established or predicted protein-coding genes (Fig. 2A; Supplemental Table 3). Importantly, the expressed RNA sequences lay invariably on the sense intronic DNA strand, suggesting that they were synthesized within and processed from their host pre-mRNAs. To test this notion, the full-length human ALUACA7 element that encodes the AluACA7 RNA and resides within the first intron of the PARP1 [poly-ADP-ribose polymerase family, member 1] gene was PCR-amplified and inserted into the second intron of the human β-globin gene in the pGL expression vector [Fig. 3A]. The resulting pGL/ALU7 and the empty pGL expression plasmids were transfected into HeLa cells, and accumulation of the processed AluACA7 and β-globin RNAs was monitored by RNase A/T1 protection using internally labeled sequence-specific antisense RNA probes [Fig. 3A]. Mapping of the RNAs derived from pGL-transfected cells with a β-globin-specific probe (probe 2) confirmed the accumulation of the three processed β-globin mRNA exons: E1, E2, and E3 [Fig. 3B, lane 2]. In pGL/ALU7-transfected cells, besides the spliced globin exons, a novel, ~105-nt-long RNA was detected with the GL/ALU7-specific probe 1 [Fig. 3B, lane 3]. Mapping of the same RNA with probes complementary to the full-length ALUACA7 DNA [probe 3] or to the 3′-terminal right arm of ALUACA7 [probe 4] demonstrated that the accumulating new RNA was processed from the right arm of the expressed intronic ALUACA7 RNA [Fig. 3B, lanes 4,5]. This conclusion was further corroborated by the observation that upon transfection of HeLa cells with the pGL/ALU7′ expression plasmid that carried only the right arm of ALUACA7, the 105-nt-long AluACA7 RNA efficiently accumulated [Fig. 3B, lanes 8,9]. This also demonstrated that the accumulating AluACA7 RNA, instead of being synthesized from putative internal Pol III promoter elements located in the left arm of ALUACA7 DNA, was processed from the globin pre-mRNA synthesized by Pol II from the CMV promoter.
To test whether the predicted H, ACA, pCAB, and dCAB boxes are essential for the in vivo processing of AluACA7 from the transiently expressed globin pre-mRNA, these sequence motifs were altered in the pGL/ALU7 expression construct (Fig. 3A). The resulting mutant pGL/ALU7H, pGL/ALU7ACA, pGL/ALU7pCAB, and pGL/ALU7dCAB expression plasmids were transfected into HeLa cells. RNase mappings with specific probes demonstrated that disruption of the H and ACA motifs fully abolished accumulation of the mutant AluACA7 RNAs but had no
Pol III-transcribed Alu RNAs were separated on a sequencing gel, protected by AluACA7 or partially protected by other AluACA and ALU elements. The endogenous AluACA7 RNA was mapped with a high-specific-activity RNA probe complementary to the ALUACA7 DNA. HeLa cellular RNA (30 μg) was mapped with sequence-specific RNA probes. (Fig. 3C, lanes 1–4) Size markers in nucleotides. (Fig. 3B [lanes 1,6], C [lane 1]). However, when the amount of HeLa RNA used for mapping, the specific activity of the ALUACA7-specific probe (probe 3), and the time of autoradiography were increased ~10-fold each, the endogenous HeLa AluACA7 RNA was readily detected and, similarly to the transiently expressed RNA, protected an ~105-nt-long fragment of the RNA probe (Fig. 3D, lane 2). Thus, we concluded that the AluACA7 RNA was faithfully processed from the second intron of the transiently expressed globin pre-mRNA.

The 5′-terminal hairpin of AluACA RNAs shows strong structural variation

The correct 5′ and 3′ termini of transiently overexpressed AluACA7 RNA were determined by primer extension and 3′ end race analysis, respectively (Fig. 4A,B). These experiments confirmed that the human AluACA7 RNA is composed of 105 nt and therefore represents the shortest human H/ACA RNA reported thus far. Notably, AluACA7 carries a 29-nt-long 5′-terminal hairpin that is unusually short for H/ACA RNAs (Fig. 4C). To test whether a short 5′ hairpin is a common hallmark of AluACA RNAs, four other RNAs (AluACA15, AluACA17, AluACA21, and AluACA177), representing different Alu subfamilies (Fig. 2B), were transiently overexpressed in HeLa cells using the pGL expression vector (Fig. 4D). RNase mappings demonstrated that all Alu RNAs were processed from the transiently expressed globin pre-mRNA, albeit with different efficiency (Fig. 4D, lanes 3,4,7,8,11,12,15,16). The accumulating AluACA RNAs showed strong size variations. While the apparent length of AluACA15 was similar to that of AluACA7, the other three RNAs (AluACA17, AluACA21, and AluACA177) migrated above the 152-nt-long first exon (E1) of the globin mRNA, albeit with different efficiency (Fig. 4D, lanes 3,4,7,8,11,12,15,16). The accumulating AluACA RNAs showed strong size variations. While the apparent length of AluACA15 was similar to that of AluACA7, the other three RNAs (AluACA17, AluACA21, and AluACA177) migrated above the 152-nt-long first exon (E1) of the globin mRNA. Importantly, mapping of endogenous HeLa AluACA15, AluACA17, AluACA21, and AluACA177 using increased amounts of cellular RNAs and high specific activity probes confirmed that the transiently expressed AluACA15, AluACA17, AluACA21, and AluACA177 RNAs were faithfully processed from the globin pre-mRNA [Fig. 5A; data not shown]. Computer folding of the 5′-terminal genomic sequences predicted by the measured lengths of the mature RNAs revealed that the human AluACA15, AluACA17, AluACA21, and AluACA177 RNAs carry 27-, 85-, 79-, and 82-nt-long 5′-terminal hairpins, respectively (Fig. 4C). While the short AluACA7 (105-nt) and AluACA15 (107-nt) RNAs were processed exclusively from the right arms of intronic Alu dimers, the 5′-terminal nucleotides of the AluACA RNAs were processed from the second intron of the transiently expressed globin pre-mRNA.
long AluACA17 (162-nt) and AluACA21 (155-nt) RNAs derived from the terminal regions of the upstream left Alu arms. Interestingly, the 5’-terminal sequences of the 158-nt-long AluACA17 RNA that is encoded by a right Alu monomer originated from upstream non-Alu-flanking sequences. Thus, we propose that in contrast to the highly conserved 3’-terminal hairpin unit, the 5’-terminal hairpins of human intron-encoded AluACA RNAs show strong sequence and structural variations.

AluACA RNAs associate with box H/ACA core proteins and Wdr79

We next assayed whether the newly discovered human AluACA RNAs are associated with the four box H/ACA RNP proteins: dyskerin, Nhp2, Gar1, and Nop10. Flag-tagged dyskerin (FL-Dys) and green fluorescent protein (GFP)-tagged Nhp2-GFP, Gar1-GFP, and Nop10-GFP proteins were transiently expressed in HeLa cells and, after extract preparation, immunoprecipitated with anti-Flag and anti-GFP antibodies [Fig. 5A]. IP of the Wdr79 scaRNP protein from a HeLa cell extract with a specific antibody recovered U85, AluACA7, and AluACA15, confirming the specificity of the observed interaction of transiently expressed Flag-tagged Wdr79 with HeLa AluACA RNAs (Fig. 5A, lane 20). As expected, none of the tested H/ACA scaRNP proteins interacted with HeLa 7SL RNA (Fig. 5A, lanes 4, 8, 12, 16, 20). On the other hand, IP of HeLa Srp14, a component of the Srp9/Srp14 heterodimer that binds to Pol III-specific Alu RNAs and the “Alu domain” of 7SL RNA [Bovia et al. 1995; Hsu et al. 1995], efficiently recovered the 7SL RNA but failed to pull down U85, AluACA7, and AluACA15 [Fig. 5A, lane 24].

Finally, we wanted to confirm that similar to the endogenous HeLa AluACA RNAs, transiently overexpressed AluACA RNAs processed from the globin pre-mRNA are accurately packaged into H/ACA scaRNPs.

Figure 4. Mapping of transiently overexpressed AluACA RNAs. (A) Determination of the 5’ terminus of transiently overexpressed AluACA7 with primer extension analysis. The 5’-terminal nucleotides of AluACA7 are indicated. (B) Mapping of the 3’ terminus of AluACA7 with RNA 3’ end race. The 3’-terminal AluACA7 and the 5’-terminal oligonucleotide tag sequences are shown. (C) Proposed secondary structures of the 5’ hairpins of AluACA RNAs. The 5’-terminal sequences of AluACA15, AluACA17, AluACA21, and AluACA117 were inferred from the lengths of transiently expressed RNAs measured by RNase mappings. The predicted H boxes are indicated. Nucleotides derived from left Alu arms or from upstream non-Alu-flanking sequences are shown. (D) Transient expression of AluACA RNAs in HeLa cells. RNAs from HeLa cells nontransfected (NT) or transfected with the indicated expression plasmids were analyzed by RNase mappings with sequence-specific probes indicated above the lanes. For other details, see the legend for Figure 3B.
The AluACA7 and AluACA15 RNAs, together with FL-Dys and FL-Wdr79, were transiently expressed in HeLa cells [Fig. 5B]. Upon IP of the accumulating FL-Dys and FL-Wdr79 proteins, RNase mapping demonstrated that similar to the control U85 scaRNA, both AluACA7 and AluACA15 were present in the pellets of the IP reactions, which, on the other hand, lacked the 7SL RNA [Fig. 5B, lanes 3,5,10,12]. As predicted, IP of the Srp14 SRP protein, which recovered the 7SL RNA, failed to precipitate the AluACA7, AluACA15, and U85 RNAs [Fig. 5B, lanes 7,14]. These observations demonstrated that human AluACA RNAs associate with the four H/ACA core proteins dyskerin, Nhp2, Nop10, and Gar1 as well as with the proposed CB localization protein Wdr79.

The CAB boxes of AluACA7 are essential for both Wdr79 binding and efficient RNA accumulation

Binding of the Wdr79 scaRNP protein is directed by a short sequence motif, the CB localization signal (CAB box), located in the terminal loop of the 5' and/or 3' hairpin of box H/ACA scaRNAs. Surprisingly, the terminal stem–loop region of the 3' hairpin of AluACA RNAs carries two closely spaced potential Wdr79-binding motifs: the distal and proximal CAB boxes [Fig. 2]. To test the functional significance of the two potential CAB boxes of AluACA RNAs, mutant AluACA7 RNAs with altered proximal [AluACA7pCAB] or distal [AluACA7dCAB] CAB motifs were transiently expressed in HeLa cells [Fig. 6A]. Association of the accumulating mutant AluACA7 RNAs with the four H/ACA core proteins dyskerin, Nhp2, Nop10, and Gar1 was monitored by Western blot analysis. RNA was prepared from cell extracts (Ext) and pellets of IP reactions were analyzed by RNase mapping with sequence-specific RNA probes indicated on the right. (Lane M) DNA and protein size markers. [B] Association of transiently overexpressed AluACA RNAs with scaRNP proteins. HeLa cells transfected with pGL/ALU7 or pGL/ALU15 were cotransfected with the pFL-DYS or pFL-WDR expression plasmids as indicated. Extracts prepared from nontransfected (NT) or transfected cells were subjected to IP with the indicated antibodies. RNAs prepared from cell extracts and pellets of IP reactions were analyzed by RNase protection.
RNAs with HeLa Wdr79 was measured by co-IP experiments using an anti-Wdr79 antibody. Compared with the control wild-type AluACA7 [Fig. 6A, lane 2], the mutant AluACA7pCAB and AluACA7dCAB RNAs showed reduced Wdr79-binding capacities [Fig. 6A, lanes 4, 6]. Simultaneous disruption of both CAB boxes fully abolished the Wdr79-binding ability of the double mutant AluACA7pdCAB RNA [Fig. 6A, lane 8]. Moreover, as compared with the wild-type RNA, the double mutant AluACA7pdCAB RNA lacking functional CAB boxes accumulated at a highly reduced level, contrary to the fact that accumulation of the host globin mRNA showed no alteration [Fig. 6A, cf. lanes 1 and 7]. This suggested that besides binding of Wdr79, the CAB boxes are also critical for efficient accumulation of AluACA RNAs.

To further assay the role of the two CAB boxes of AluACA RNAs in RNA accumulation, we constructed and transiently expressed a hybrid AluACA RNA encompassing the 3′ hairpin unit of AluACA7 and the 5′-terminal hairpin region of the U64 box H/A CA snoRNA [Fig. 6B]. The U64-AluACA7 composite RNA accumulated with 15–20 times higher efficiency than the wild-type AluACA7, indicating that the inherently weak expression of AluACA7 is at least partially due to its unusually short 5′ hairpin [Fig. 6B, lane 2]. Upon IP of HeLa Wdr79, the U64-AluACA7 composite RNA coprecipitated with Wdr79, demonstrating that the 5′ hairpin unit of AluACA7 is dispensable for Wdr79 binding [Fig. 6B, lane 3]. Alteration of both the distal and proximal CAB boxes of U64-AluACA7, besides fully disrupting Wdr79 binding, largely reduced the accumulation of the double mutant U64-AluACA7pdCAB RNA [Fig. 6B, lanes 4, 5]. Based on these observations, we concluded that the two CAB box motifs of AluACA7, besides recruiting Wdr79 in a cumulative fashion, contribute to the efficient accumulation of AluACA7.

Transiently expressed tagged AluACA7 RNAs accumulate in the nucleoplasm

While all box H/ACA RNAs tested so far accumulated within the nucleus, Alu RNAs have been detected in both the nucleus and cytoplasm (for reviews, see Kiss et al. 2006; Berger and Strub 2011). To learn about the subcellular localization of human AluACA RNAs, HeLa cells were fractionated into nuclear and cytoplasmic fractions, and the distribution of the endogenous AluACA7 and AluACA15 RNAs was monitored by RNase mapping [Fig. 7A]. Together with the control U85 scaRNA that accumulates in the nucleoplasmic CBs (Darzacq et al. 2002), both AluACA7 and AluACA15 RNAs were detected in the nuclear fraction and were missing from the cytoplasm, where the majority of the 5.8S rRNA accumulated [Fig. 7A, lanes 3, 4].

We used fluorescent in situ hybridization microscopy to determine the precise subnuclear localization of AluACA RNAs. To avoid undesired cross-hybridization of AluACA-specific oligonucleotide probes with Pol III-synthesized full-length or 3′-terminally processed Alu RNAs or with pre-mRNAs containing Alu sequences, modified AluACA7 RNAs carrying an MS2 coat protein-binding motif in either the hinge (AluACA7-MSh) or the 5′-terminal hairpin [AluACA7-MS5'] were transiently expressed in HeLa cells using the pGL expression vector [Fig. 7B]. RNase mappings confirmed that both MS2-tagged AluACA RNAs were correctly and efficiently processed from the β-globin host pre-mRNA [Fig. 7B, lane 2]. Coexpression and IP of FL-Dys and FL-Wdr79
proteins revealed that the accumulating AluACA7-MS\(h\) and AluACA7-MS\(5'\) RNAs were packaged into Wdr79-associated H/ACA RNPs [Fig. 7B, lanes 4,5]. Moreover, cell fractionation experiments showed that the transiently expressed AluACA7-MS\(h\) RNPs, similar to the endogenous AluACA7 and AluACA15 RNPs, accumulated within the nucleus of transfected cells [data not shown].

Probing of cells expressing AluACA7-MS\(h\) or AluACA7-MS\(5'\) RNAs with a fluorescently labeled MS2 coat protein-binding motif (in red) are shown. For the structure of U64-AluACA7, see Figure 6B. Extracts (Ext) prepared from HeLa cells nontransfected (NT) or transfected (TR) with the indicated combination of the pGL/ALU7-MS\(h\), pGL/ALU7-MS\(5'\), pGL/ACA26, and pFLWDR79 expression plasmids were reacted with an anti-Flag antibody. RNAs prepared from cell extracts or the pellets of IP reactions were analyzed by RNase mapping. Lane Cont shows mapping with E. coli tRNA. (C) Fluorescent in situ hybridization. HeLa cells transiently expressing the indicated tagged or chimeric version of AluACA7 RNA were stained with sequence-specific fluorescent oligonucleotide probes. Nucleoli were visualized by expression of fibrillarin-GFP or Gar1-GFP. CBs were immunostained with antibodies against Wdr79 or coilin. DAPI staining of nuclear DNA was omitted at U2 localization. Bar, 10 μm.

Figure 7. Subcellular localization of AluACA RNAs. (A) HeLa AluACA7 and AluACA15 RNAs accumulate in the nucleus. RNA samples isolated from either HeLa cells (T) or the nuclear (Nu) and cytoplasmic (Cy) fractions of HeLa cells were mapped by RNase A/T1 protection with antisense RNA probes as indicated on the right. (B) Expression of internally tagged AluACA7 RNAs in HeLa cells. Schematic structures of transiently expressed AluACA7-MS\(h\) and AluACA7-MS\(5'\) RNAs with the inserted MS2 coat protein-binding motifs (in red) are shown. For the structure of U64-AluACA7, see Figure 6B. Extracts (Ext) prepared from HeLa cells nontransfected (NT) or transfected (TR) with the indicated combination of the pGL/ALU7-MS\(h\), pGL/ALU7-MS\(5'\), pGL/ACA26, and pFLWDR79 expression plasmids were reacted with an anti-Flag antibody. RNAs prepared from cell extracts or the pellets of IP reactions were analyzed by RNase mapping. Lane Cont shows mapping with E. coli tRNA. (C) Fluorescent in situ hybridization. HeLa cells transiently expressing the indicated tagged or chimeric version of AluACA7 RNA were stained with sequence-specific fluorescent oligonucleotide probes. Nucleoli were visualized by expression of fibrillarin-GFP or Gar1-GFP. CBs were immunostained with antibodies against Wdr79 or coilin. DAPI staining of nuclear DNA was omitted at U2 localization. Bar, 10 μm.
a predominant nucleoplasmic distribution. A weak accumulation of U64-AluACA7 within the nucleolar territories was probably due to its high level of expression. Nevertheless, these results confirm that the cis-acting elements supporting the nucleoplasmic accumulation of AluACA7 are located in the 3′-terminal hairpin of the RNA. Simultaneous disruption of the distal and proximal CAB box motifs failed to alter the nucleoplasmic localization of the mutant U64-AluACA7-pdCAB and the AluACA7-MSS′-pdCAB RNAs, indicating that Wdr79 binding is not required for the nucleoplasmic localization of AluACA RNAs.

Instead of being evenly dispersed in the nucleoplasm, the MS2-tagged AluACA7 RNAs showed a speckled localization pattern highly reminiscent of the nucleoplasmic distribution of spliceosomal snRNAs (Carmo-Fonseca et al. 1991a,b). Upon costaining of HeLa cells with fluorescent oligonucleotide probes specific for the transiently expressed AluACA7-MS RNAs, the AluACA7 and U2 signals failed to perfectly overlap in the nucleoplasm. Moreover, in contrast to U2 that concentrated in the nucleoplasmic speckles, the AluACA7-MSh RNA showed no significant enrichment in speckles and paraspeckles (data not shown). In conclusion, in situ localization experiments confirmed that the newly discovered AluACA RNPs represent a novel subgroup of human box H/ACA RNPs that accumulate in the nucleoplasm of human cells.

Discussion

Despite the fact that Alu repetitive elements constitute >10% of the human genome, expression of Alu RNAs is remarkably low (Berger and Strub 2011). Genomic Alu elements can be transcribed by either Pol III using the weak internal promoters of Alu DNAs or Pol II as parts of long pre-mRNA transcripts. In this study, we demonstrated that intron-embedded nascent Alu RNAs can be processed into metabolically stable RNAs. The novel intron-encoded Alu RNAs, termed AluACA RNAs, feature the characteristic hallmarks of box H/ACA scaRNAs: They fold into the common hairpin–hinge–hairpin–tail structure and carry the conserved H, ACA, and CAB boxes of H/ACA scaRNAs. Consequently, the mature AluACA RNAs associate with the five H/ACA scaRNPs proteins dyskerin, Nop10, Nhp2, Gar1, and Wdr79.

Our in vivo processing studies confirmed that the intronic AluACA RNAs, instead of being transcribed independently by Pol III, are post-transcriptionally processed from their host pre-mRNAs in a box H- and ACA-dependent fashion [Fig. 3]. Consistent with this biosynthetic pathway, the genomic copies of 337 new AluACA RNAs, out of 348, were found within introns of known or predicted protein-coding genes invariably in the sense orientation [Supplemental Table 3]. Although the remaining 11 new RNAs [AluACA11, AluACA63, AluACA82, AluACA83, AluACA100, AluACA111, AluACA134, AluACA324, AluACA328, AluACA342, and AluACA348] were mapped to intergenic regions lacking established Pol II transcription units, it is unlikely that these RNAs are synthesized independently by Pol III because primary H/ACA RNA transcripts generated by Pol III are not recognized as substrates for in vivo H/ACA RNA processing [Richard et al. 2006]. Therefore, we favor the idea that these intergenic Alu elements are transcribed within yet-undetected Pol II transcripts or are misplaced in the human genome sequence.

We identified 348 novel AluACA RNAs [Supplemental Table 1], but the final number of human AluACA RNAs is apparently much higher. During the processing of our sequence data, we followed a stringent filtering strategy to definitively distinguish between highly similar genomic Alu elements. We ignored almost 200 potential AluACA RNA sequences that were either <75 nt or absent from the human genome reference sequence [UCSC Genome Browser]. It is important to mention that dozens of our Alu reads, although missing from the UCSC genome sequence, were detected in other databases, indicating that they likely represent bona fide AluACA RNAs. We can also predict that human cells in different tissues and at different stages of development express many additional intronic AluACA RNAs. Thus, we propose that human cells express several hundreds or maybe thousands of AluACA RNAs. However, contrary to their high number, AluACA RNAs show a low level of global accumulation in HeLa cells. As predicted by the distribution of AluACA and scaRNA reads in our deep sequencing data set, AluACA RNAs accumulate at a 10 times lower level than do canonical scaRNAs.

Human Alu elements fall into three major subfamilies of different evolutionary ages: the young AluY and the more ancient AluS and AluJ subfamilies [Batzer and Deininger 2002]. Pol III preferentially transcribes AluY elements, which constitute the smallest Alu subfamily [Shaikh et al. 1997]. In contrast, the subfamily distribution of the 348 new intronic AluACA RNAs—specifically, 37 AluY, 78 AluJ, and 233 AluS RNAs—shows a strong correlation with the genomic abundance of AluY, AluJ, and AluS elements [Supplemental Table 2]. In primates, fusion of the more ancient free left Alu monomer with the primate-specific free right Alu monomer led to the prevailing dimeric Alu organization [Quentin 1992; Kriegs et al. 2007]. The new AluACA RNAs, with no exception, derived from right Alu monomers. As both originated from the 7SL RNA, the left and right Alu monomers are highly similar, except that the right monomer contains a 31-nucleotide insertion missing from the left monomer [Supplemental Fig. 2]. In fact, this inserted extra sequence accounts for the AluACA-coding capacity of the right Alu monomer because it constitutes the terminal stem–loop region and carries the proximal and distal CAB boxes of the 3′ hairpins of AluACA RNAs. Given that right Alu arm sequences are confined to primate genomes, the AluACA RNAs represent a human-specific or maybe primate-specific subgroup of box H/ACA RNAs [Quentin 1992]. Therefore, it seems that gaining AluACA RNA-coding capacity by Alu DNAs is a recent event of primate genome evolution, although we cannot exclude the possibility that primordial SRP RNA originated from ancient H/ACA RNAs.
Besides their apparent similarity to canonical H/ACA scaRNPs, the AluACA RNPs also possess distinctive structural features. The AluACA RNAs may carry unusually short 5′ hairpins that are not anticipated to support stable H/ACA RNP assembly [Fig. 4]. The archaeal homolog of Nhp2, called L7Ae, tethers Nop10 and the catalytic domain of dyskerin to the upper stem of H/ACA RNAs by forming a specific interaction with the conserved Kink turn motif [Rozhdestvensky et al. 2003; Baker et al. 2005; Charpentier et al. 2005; Liang et al. 2007]. Eukaryotic H/ACA RNAs lack Kink turns, suggesting that Nhp2, contrary to its structural similarity to L7Ae [Koo et al. 2011], forms a more flexible interaction with the structurally variable upper stems of H/ACA RNAs [Li et al. 2012]. Accumulation of AluACA7 RNA requires the H box motif [Fig. 3], indicating that the short 5′ hairpin of AluACA7 still binds dyskerin that recognizes the H box and the preceding basal stem of H/ACA RNAs [Li and Ye 2006]. Eukaryotic dyskerin can form a stable complex with Nop10, Nhp2, and Gar1 even in the absence of H/ACA RNAs [Henras et al. 2004; Wang and Meier 2004; Li et al. 2012]. This suggests that the short 5′ hairpins of AluACA RNAs can bind the complete set of H/ACA core proteins, contrary to the apparent absence of a docking surface for Nhp2. Nevertheless, it seems that suboptimal structural organization of the 5′ hairpin units is the major reason of the remarkable weak cellular accumulation of AluACA RNAs. Supporting this idea, replacement of the short and long 5′ hairpin units of AluACA7 and AluACA17, respectively, for the 5′ hairpin of the U64 box H/ACA snRNA dramatically increased the accumulation levels of both RNAs [Fig. 6B; data not shown].

The 3′-terminal or 5′-terminal hairpin of canonical H/ACA scaRNAs contains one CAB box motif that is essential for Wdr79 binding and CB localization but dispensable for RNA accumulation [Richard et al. 2003; Jady et al. 2004]. In contrast, the 3′ hairpin of AluACA RNAs carries two closely spaced CAB motifs that bind Wdr79 in a cumulative fashion and are crucial for efficient RNA expression [Fig. 6]. It is unclear whether the two CAB boxes promote RNA accumulation directly through binding Wdr79 or indirectly through recruiting putative trans-acting RNA processing and/or RNP assembly factors [Theimer et al. 2007]. Wdr79 has been proposed to function as a localization signal protein that targets box H/ACA and C/D scaRNPs into CBs [Tycowski et al. 2009; Venteicher et al. 2009]. However, the demonstration that Wdr79-associated AluACA RNPs accumulate in the nucleoplasm suggests that the molecular mechanism supporting the subnuclear localization of human box H/ACA RNPs is more complex than anticipated before [Fig. 7].

The function of AluACA RNPs remains elusive. The great majority of box H/ACA RNAs direct site-specific RNA pseudouridylation [see above]. Intriguingly, the 3′-terminal hairpins of AluACA RNAs contain perfect pseudouridylation guide loops that, in principle, could position the putative target uridine 14 nt upstream of the ACA box, which is an optimal configuration for efficient pseudouridylation [Fig. 2B]. Compared with the bracketing basal and upper stems, the putative pseudouridylation loops of AluACA RNAs show increased sequence variability. Thus, AluACA RNAs could direct pseudouridylation of many different target RNAs. We are currently testing whether in vitro and in vivo assembled AluACA RNPs possess pseudouridine synthase activity. The AluACA RNAs might also function as transcriptional regulators, as the right arms of human Alu RNAs have been reported to inhibit Pol II transcription through binding to core Pol II at promoters of transcriptionally repressed genes [Mariner et al. 2008; Yakovchuk et al. 2009]. However, mature AluACA RNAs partially lack the internal L region and frequently miss the entire A-rich motif, which are essential for Alu-mediated Pol II repression [Fig. 1]. Understanding the nucleoplasmic function of human AluACA snRNPs remains an exciting task for the future.

Playing important roles in all steps of gene expression, ncRNAs contribute to the biological complexity of eukaryotic organisms. In this study, besides uncovering a novel, unpredicted role for human Alu repetitive elements, we identified and partially characterized a novel, large subgroup of box H/ACA RNP that are apparently confined to human or primate cells. Although the nucleoplasmic role of the newly discovered AluACA RNPs remains unknown, they likely contribute to the genetic and biochemical complexity of human cells.

Materials and methods

Unless stated otherwise, all manipulations were performed according to standard laboratory protocols [Sambrook et al. 1989]. Modified and unmodified oligodeoxynucleotides were purchased from Eurofins MWG Operon.

Expression constructs

Full-length human Alu DNA elements encoding the AluACA7, AluACA15, AluACA17, AluACA21, and AluACA177 RNAs, together with their 69–45/58–80/65–74/71–75/159–124-base-pair [bp] 5′- and 3′-flanking sequences, respectively, were PCR-amplified using HeLa nuclear DNA as a template. To obtain the pBST/ALU7, pGL/ALU7, pBST/ALU15, pGL/ALU15, pBST/ALU17, pGL/ALU17, pBST/ALU21, pGL/ALU21, pBST/ALU177, and pGL/ALU117, the amplified Alu fragments were inserted into the ClaI and Xhol sites of the pBluescript plasmids [pBST] and pGL plasmids [Darzacq et al. 2002]. Construction of the pGL/ALU7H, pGL/ALU7pCAB, pGL/ALU7dCAB, pGL/ALU7pdCAB, pGL/ALU7ACA, pGL/ALU73′, pGL/ALU7MSb, pGL/ALU7MSs, and pGL/ALU7MSs’pdCAB expression vectors carrying mutants versions of the ALUACA7 gene was performed by PCR amplification using the appropriate mutagenic primers. The MS2 coat protein-binding sequence 5′-AAACAU GAGGAGUCACCCAUGUC-3′ was inserted between the A9 and U20 [AluACA-MSs’] or U38 and A40 [AluACA-MSb] hairpins of AluACA7. To generate pGL/U64-ALUACA7, the 5′ hairpin and the adjacent H box of the human U64 snoRNA gene [1–72 nt] were PCR-amplified and fused to the coding DNA of the second hairpin for use in the Clal and Xhol sites of the pBluescript plasmids. To obtain pGL/ACA26-MSs′, pGL/U64-ALUACA7pdCAB expression plasmid was constructed by PCR amplification using an appropriate mutagenic oligonucleotide primer and pGL/U64-ALUACA7 as a template. To obtain pGL/ACA26-MSs′.
the MS2 coat protein-binding motif was inserted between the A31 and A32 residues of the ACA26 gene in the pGL/ACA26 expression plasmid (Kiss et al. 2004). The pFL-WDR79 expression vector was purchased from GenoCopeia (EX-V0379-M12). To obtain FL-
DYS, the human dyskerin cDNA was PCR-amplified with an upstream oligonucleotide that contained a Flag motif and was inserted into the KpnI and BamHI sites of pcDNA3. The pNHP2-
GFP, pGAR1-GFP, pNOP10-GFP, and pFB-GFP expression plasmids have been reported (Dundr et al. 2000; Darzacq et al. 2006). The identity of all of the constructs was verified by sequence analysis.

RNA analyses

RNAs from HeLa cells and from the nuclear and cytoplasmic fractions of HeLa cells were extracted by the guanidine thocyanate/phenol-chloroform extraction method (Goodall et al. 1990). Primer extension analysis was performed using a 5’ end-labeled oligonucleotide primer complementary to AluACA7 from G59 to C79. RNA 3’-terminal sequence race and RNase A/T1 protection analysis have been described (Goodall et al. 1990; Kiss and Filipowicz 1993). To generate low and high specific activity (sp. act.) antisense RNA probes, the pGL/ALU and pBST/ALU recombinant plasmids were digested with HindIII and ClaI and used as a template for in vitro transcription with SP6 and T7 RNA polymerases, respectively, in the presence of [α-32P]CTP (sp. act. 220 or 2700 Ci/mmol). All probes were purified on denaturing polyacrylamide gels. About 2 μg of human HeLa RNA coimmunoprecipitated with transiently expressed Flag-tagged Wdr79 was used as a template for generation of a cDNA library that was analyzed by high-throughput sequencing using Solexa technology (Fastersi SA).

Protein analyses

Preparation of HeLa cell extracts, cell fractionation, IP of snRNP complexes, and Western blot analysis of proteins have been described (Tyc and Steitz 1989). For Western blot analysis, we used the following antibodies: mouse monoclonal anti-Flag M2 (1:10,000 dilution; Sigma), anti-GFP (1:1000 dilution; Roche), rabbit polyclonal anti-WDR (1:200 dilution; Abcam ab99376), and anti-GFP, pGAR1-GFP, pNOP10-GFP, and pFB-GFP expression plasmids. We are indebted to M. Carmo-Fonseca (Universidade de Lisboa, Portugal) for providing us with anti-coilin antibody. We are grateful to H. Setz for his help in initial data analysis. Our work was supported by la Fondation pour la Recherche Medicale grant to T.K.

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Human intron-encoded Alu RNAs are processed and packaged into Wdr79-associated nucleoplasmic box H/ACA RNPs

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