Internal Modification of U2 Small Nuclear (sn)RNA Occurs in Nucleoli of Xenopus Oocytes

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Abstract. U2 small nuclear (sn)RNA contains a large number of postranscriptionally modified nucleotides, including a 5′-trimethylated guanosine cap, 13 pseudouridines, and 10 2′-O-methylated residues. Using Xenopus oocytes, we demonstrated previously that at least some of these modified nucleotides are essential for biogenesis of a functional snRNP. Here we address the subcellular site of U2 internal modification. Upon injection into the cytoplasm of oocytes, G-capped U2 that is transported to the nucleus becomes modified, whereas A-capped U2 that remains in the cytoplasm is not modified. Furthermore, by injecting U2 RNA into isolated nuclei or enucleated oocytes, we observe that U2 internal modifications occur exclusively in the nucleus. Analysis of the intranuclear localization of fluorescently labeled RNAs shows that injected wild-type U2 becomes localized to nucleoli and Cajal bodies. Both internal modification and nucleolar localization of U2 are dependent on the Sm binding site. An Sm-mutant U2 is targeted only to Cajal bodies. The Sm binding site can be replaced by a nucleolar localization signal derived from small nucleolar RNAs (the box C/D motif), resulting in rescue of internal modification as well as nucleolar localization. Analysis of additional chimeric U2 RNAs reveals a correlation between internal modification and nucleolar localization. Together, our results suggest that U2 internal modification occurs within the nucleolus.

Key words: snRNP • Sm binding site • modified nucleotide • snoRNA • localization

Introduction

The spliceosomal small nuclear (sn)RNAs (U1, U2, U4, U5, and U6), which are essential for pre-mRNA splicing, are all posttranscriptionally modified (Reddy and Busch, 1988; Massenet et al., 1998). Aside from 5′ cap trimethylation, numerous internal nucleotides are pseudouridylated or 2′-O-methylated. U2 is the most extensively modified of all the spliceosomal snRNAs. There are 10 2′-O-methylated residues and 13 pseudouridines in the Xenopus U2 snRNA (Fig. 1; Yu et al., 1998).

The biogenesis of U2 and the other spliceosomal snRNPs has been extensively studied, principally using Xenopus oocytes (Mattaj, 1988; Mattaj and Englmeier, 1998). With the exception of U6, all major spliceosomal snRNAs are transcribed by RNA polymerase (Pol) II in response to specific promoter and termination signals (Dahlberg and Lund, 1988). These Pol II transcripts are rapidly transported to the cytoplasm, where the RNAs are trimmed at their 3′ ends and a common set of proteins called the “core” Sm proteins are assembled onto the conserved Sm binding sequence present in each of these snRNAs (Mattaj and De Robertis, 1985; Luhrmann, 1988). Upon binding of the Sm proteins, the 5′-monomethylated guanosine cap becomes 2,2,7-trimethylated, an event that is sufficient for recognition by the protein snurportin 1, which facilitates the transport of these RNA-protein particles back to the nucleus (Huber et al., 1998). In addition to the common Sm proteins, each mature Sm snRNP contains specific proteins, the number of which varies between snRNPs and can be quite large (Luhrmann, 1988; Behrens et al., 1993; Gottschalk et al., 1998, 1999). Two distinct U2 snRNPs have been observed in nuclear extracts from HeLa cells (Behrens et al., 1993) and Xenopus oocytes (Yu et al., 1998), corresponding to 12S or 17S particles. The 12S particle contains all the core Sm proteins and two U2-specific proteins, whereas the 17S particle possesses at least nine additional loosely associated U2-specific proteins (Behrens et al., 1993). Seven of these nine proteins are the subunits of splicing factor 3 (SF3; Krämer, 1996). Using...
microinjection into the *Xenopus* oocyte, we recently demonstrated that the modified nucleotides within the 5’-27 nucleotides of U2 are essential for conversion of the 12S particle to the mature 17S particle (Yu et al., 1998).

Although it is clear that U2 and other Sm snRNAs undergo Sm protein binding and 5’ cap hypermethylation in the cytoplasm, it is not known where within the cell these Sm snRNAs are internally modified. It is established that the nucleolus harbors a large number of small nucleolar (sno)RNPs responsible for pseudouridylation and 2'-O-methylation of Pol I-transcribed ribosomal RNAs (Maxwell and Fournier, 1995; Smith and Steitz, 1997; Tollervey and Kiss, 1997; Yu et al., 1999). Recent studies indicate that 2'-O-methylation of U6 snRNA, a Pol III transcript, is likewise mediated by snoRNPs (Tycowski et al., 1998; Ganot et al., 1999). Intranuclear localization studies indicate that the U6 snRNA travels through the nucleolus (Lange and Gerbi, 2000; Narayanan, A., R.M. Terns, and M.P. Terns, unpublished data). The striking similarities in 2'-O-methylation between ribosomal RNAs (rRNAs) and the U6 snRNA raises the possibility that internal modification of Pol II–transcribed U2 and other Sm snRNAs also occurs in the nucleolus. In this study, we use the *Xenopus* oocyte microinjection system to show that internal modification of U2 (a product of Pol II transcription) occurs exclusively in the nucleus, and likely within the nucleolus.

**Materials and Methods**

**Preparation of U2 Mutant and Chimeric RNAs**

PCR was used to generate the Sm-mutant U2 construct, starting with a plasmid containing the wild-type U2 sequence (Yu et al., 1998). First, a 5’ and a 3’ Sm-mutated U2 fragment were produced using two pairs of DNA oligonucleotides. One pair (the 5’ primer [primer 1], 5’-TAATACGACTCACTATAGG-3’ corresponding to the T7 promoter sequence located immediately upstream of the 5’ terminus of U2, and the 3’ primer [primer 2], 5’-CTATATATAGTATACGATCATCAG-3’, complementary to nucleotides 84–112 of *Xenopus* U2 except for two mismatches [underlined] in the second and third nucleotides of the Sm binding site) was used to amplify a 5’ half U2 fragment. The other pair (the 5’ primer [primer 3], 5’-TAATATGGAATGTTGAACTGGAAG-3’, corresponding to nucleotides 92–115 of *Xenopus* U2, except for two changes in the second and third nucleotides of the Sm binding site [underlined], and the 3’ primer [primer 4], 5’-GGGAAGTGCCACCGGTCTCGAGGT-3’, containing three G’s at a 21-nucleotide sequence complementary to the 3’ end of *Xenopus* U2) was used to create a 3’ half U2 fragment. These two half DNA PCR products (containing an overlapping sequence, base pairs 92–112 with respect to the U2 sequence) were then mixed with primer 1 and primer 4 for a new round of PCR amplification, generating a mutant U2 construct with the second and the third U’s of the Sm binding site changed to G’s. The final PCR product was cloned into the Smal site of pGEM-3Z vector (Promega) and sequenced. After linearization with Smal, the plasmid was used as a template for GpppG-primed (for wild-type U2, ApppG was also used) transcription by T7 RNA polymerase.

To create the MID U2-U7 chimera, two oligodeoxynucleotides were used. The 5’ primer, 5’-CTTATACGACTCACTATAGGATTGATATATAATTTTTGACAATCGGCTTTG-3’, corresponding to the T7 promoter sequence, the 5’ 32 nucleotides of the yeast U14 snRNA (including the terminal stem and box C sequences [Samarsky et al., 1998]), and the 5’ 19 nucleotides of U2. The 3’ primer, 5’-ATATATATAGTATACGATCATCAGACCAAGTGCAC-3’, was complementary to the 3’ 27 nucleotides of yeast U14 (including the terminal stem and box D sequences) and the 3’ 20 nucleotides of *Xenopus* U2 RNA. After PCR, the DNA product was gel purified and used as a template for in vitro T7 RNA polymerase transcription.

To create the MID U2-U7 chimera, two oligodeoxynucleotides were used. The 5’ primer, 5’-CTTATACGACTCACTATAGGATTGATATATAATTTTTGACAATCGGCTTTG-3’, corresponding to the T7 promoter sequence, nucleotides 19–46 of *Xenopus* U2, and nucleotides 19–34 of *Xenopus* U7 snRNA. The 3’ primer, 5’-TGTTGGCTCTACAGATTAGAAACGGTCTCGAGACAAAATAG-3’, was complementary to the 3’ 38 nucleotides of *Xenopus* U7 snRNA. These two primers have a complementary sequence, nucleotides 19–34 with respect to the *Xenopus* U7 snRNA. After annealing, both strands were extended with Klenow DNA polymerase. The double-stranded DNA product served as a template for in vitro transcription of MID U2-U7 chimera by T7 RNA polymerase.

**Anti-Sm (Y12) Immunoprecipitation**

Immunoprecipitation with anti-Sm (Y12) antibodies was performed according to Lerner et al. (1981) and Yu et al. (1998). In brief, after injection of labeled U2 RNAs, nuclei were broken by sonication in Net-2 buffer (50
mM Tris, pH 7.5, 150 mM NaCl, 0.05% NP-40). The nuclear suspension was clarified by centrifugation at 14,000 g for 5 min. The supernatant was mixed with Y-12 antibodies at 4°C for 2 h. After centrifugation, the pellet was washed three times with Net-2 buffer and digested with proteinase K. RNAs were recovered by phenol-chloroform-isooamylalcohol extraction and ethanol precipitation.

**Xenopus Oocyte Microinjection and RNA Modification Assays**

The procedures for microinjection and modification assays were essentially as described previously (Yu et al., 1998). In brief, various U2 RNAs (differentially capped wild-type U2, Sm-mutant U2, U2-C/D motif chimeras, and the U2-U7 chimera) uniformly labeled with α[32P]UTP (10⁶ cpm/μl) were individually injected into Xenopus oocytes (32 nl for cytoplasmic injection, and 9 nl for nuclear injection [in most cases, 9 nl of U2 RNA constructs was directly injected into isolated nuclei under oil]). After 5 h at room temperature, RNAs were recovered from the nuclei or cytoplasm and assayed for modifications.

To assay pseudouridylation, RNAs were digested with nuclease P1 in 3 μl of 20 mM sodium acetate, pH 5.2, at 37°C for 1 h. The digested samples were analyzed on cellulose TLC PEI plates (Baker; Patton, 1991). To determine the site specificity of pseudouridylation, endogenous U2 was first assayed for pseudouridylation. The positions of uridylate, pseudouridylate, uncleaved U2, and 3’ fragments generated by RNase H site-specific cleavage are indicated on the side of each gel. In these experiments, >80% of the expected level of pseudouridylation and 2’-O-methylation was observed.

**Results**

**U2 Modification Occurs Exclusively in the Nucleus**

Given the fact that U2 snRNA cycles through the cytoplasm before returning to the nucleus, we first tested RNA constructs were analyzed on 6% denaturing gels (Yu et al., 1997, 1998).

**Analysis of the Intranuclear Distribution of U2 snRNA and Variants**

The procedure used to prepare Xenopus oocyte nuclear spreads and to determine the intranuclear distriution of injected fluorescent-labeled RNAs has been described (Narayanan et al., 1999). In brief, fluorescent-labeled RNAs were generated by in vitro transcription of linearized plasmid DNA in the presence of α[32P]GTP and fluorescent-12-UTP (1:1 ratio with UTP, Boehringer). The fluorescent-labeled RNAs (1 fmol each) were injected into the nuclei of stage V or stage VI oocytes. α[32P]-labeled U6 snRNA (retention control), U1 snRNA, and rRNA (export controls) were co-injected with each fluorescent-labeled test RNA to help make conclusions about the stability and nucleocytoplasmic distribution of wild-type U2 snRNA and U2 variants. The injected oocytes were incubated at 18°C for 5 h before nuclei were dissected and nuclear spreads were prepared. Images were obtained on a 63× magnification inverted fluorescence microscope (Axiovert S 100; Zeiss) equipped with differential interference contrast (DIC) optics (Thornwood) using a cooled charge-coupled device camera (Quantix-Photometrics) and IPLab Spectrum software (Signal Analytics). Localization analysis was performed using multiple independent sets of RNAs and oocytes, and at least four slides per time point. Representative data are shown (see Fig. 4). To determine the stability and nucleocytoplasmic distribution of the injected RNAs, RNAs present in the nuclear (N) and cytoplasmic (C) fractions were purified 5 h after injection (from the same set of oocytes analyzed by microscopy) and analyzed by 8% denaturing PAGE and autoradiography.

**Figure 2.** U2 internal modifications occur exclusively in the nucleus. α[32P]UTP uniformly labeled G- or A-capped U2 was injected into the cytoplasm of oocytes (A, lanes 1 and 2; B, lanes 1–6) or directly into nuclei (A, lanes 4 and 5; B, lanes 10–15). After cytoplasmic injection, only G-capped U2 entered the nucleus; after nuclear injection, both G- and A-capped U2 RNAs were retained in the nucleus. RNAs recovered from nuclei (A, lanes 1, 4, and 5; B, lanes 1–3, 10–15) or from cytoplasm (A, lane 2; B, lanes 4–6) were assayed for pseudouridylation (A) and 2’-O-methylation (B). In the 2’-O-methylation assay, two different chimeras were used to test two positions (G11 and A30). The control is un.injected U2 RNA. (C) Oocyte nuclei were separated from the cytoplasm under oil. α[32P]UTP uniformly labeled U2 was then injected into the isolated nuclei (lane 1) or enucleated oocytes (lane 2). After a 5-h incubation, U2 RNAs were recovered and assayed for pseudouridylation. The positions of uridylate, pseudouridylate, uncleaved U2, and 3’ fragments generated by RNase H site-specific cleavage are indicated on the side of each gel. In these experiments, >80% of the expected level of pseudouridylation and 2’-O-methylation was observed.
isolated nuclei, not from the enucleated oocytes, was cleated oocytes. Again, only U2 RNA recovered from A or G cap directly into the isolated nuclei or the enucleated oocytes. This suggests that the cap structure does not affect the ability to serve as a substrate for internal modification, we injected both A-capped and G-capped U2 RNA directly into nuclei and a control RNAs (U1 snRNA, U6 snRNA, and tRNA), were injected into oocyte nuclei and the intranucleus of the snRNA by the modifying machinery. Alternatively, the Sm site is required for internal modification. The Sm site is critical for U2 snRNP biogenesis (Lührmann, 1988; Mattaj, 1988) and is located in the 3′ half of the RNA, where essentially no modified nucleotides are present. We mutated this element by changing the second and the third uridines in the Sm binding site to guanosines (Fig. 1). 5 h after injection of RNA that was uniformly labeled with [32P]UTP into isolated nuclei, total RNAs were recovered and modification assays were performed. Although the wild-type U2 was pseudouridylated (Fig. 3, lane 2) and 2′-O-methylated (data not shown), no modified nucleotides were detected in the Sm mutant RNA (Fig. 3, lane 3 and data not shown), even after a prolonged (10-h) incubation (Fig. 3, lane 7), suggesting that the Sm site is required for internal modification. The Sm site is important as a site for binding of Sm proteins, which normally occurs in the cytoplasm; however, assembly of the Sm RNP can occur in the nucleus under some conditions, apparently via exchange of proteins with endogenous snRNPs (Terns and Dahlberg, 1994; Terns et al., 1995). Assembly of the injected wild-type U2 into particles in our experiments was confirmed by anti-Sm immunoprecipitation: ~50% of injected RNAs were immunoprecipitated (Fig. 3, compare lanes 4 and 5). The U2 RNA bound by Sm proteins was pseudouridylated (Fig. 3, lane 5), whereas only a trace amount of pseudouridylate was detected in the supernatant (unbound) fraction (Fig. 3, lane 4), presumably because of incomplete immunoprecipitation. We conclude that the Sm binding site, and likely the binding of Sm proteins, is required for U2 internal modification.

Wild-Type U2, but Not Sm-mutant U2, Localizes to the Nucleolus

The Sm binding site could contribute to U2 internal modification in several ways. For instance, mutation of the Sm binding site could directly or indirectly impair recognition of the snRNA by the modifying machinery. Alternatively, the nucleus contains several subcompartments (e.g., nucleoplasm, nucleoli, Cajal bodies, gems, speckles, and snurpsomes) and the binding of Sm core proteins could target the U2 snRNA to nuclear subcompartments where internal modification occurs. Alteration of the Sm binding site resulting in mislocalization would thus affect modification.

We analyzed the intranuclear distribution of fluorescently labeled RNAs to pinpoint the location of the wild-type and the Sm-mutant U2 within the nucleus. Fluorescently and radio-labeled U2 RNAs, along with radiolabeled control RNAs (U1 snRNA, U6 snRNA, and tRNA), were injected into oocyte nuclei and the intranuclei, not from the enucleated oocytes, was

whether the internal modification (pseudouridylation and 2′-O-methylation) occurs in the nucleus and/or in the cytoplasm. In these experiments, we exploited the fact that after injection into the cytoplasm of Xenopus oocytes, GpppG-capped U2 is hypermethylated and rapidly transported into the nucleus, whereas ApppG-capped U2 is not trimethylated and remains in the cytoplasm (Fischer et al., 1991). 5 h after injection of differentially capped, uniformly labeled U2 snRNA into the cytoplasm, the cytoplasm and the nucleus were manually separated. The RNA in each cellular compartment was assayed for pseudouridylidation by complete nuclease P1 digestion followed by TLC analysis (Fig. 2 A) and for 2′-O-methylation by RNase H site-specific cleavage directed by 2′-O-methyl RNA–DNA chimeras (Fig. 2 B; Yu et al., 1997, 1998). The G-capped U2 recovered from nuclei was efficiently pseudouridylated (Fig. 2 A, lane 1) and 2′-O-methylated (Fig. 2 B, lanes 1–3). In contrast, the A-capped U2 recovered from the cytoplasm was not detectably modified (Fig. 2, A and B, lanes 2 and 4–6, respectively). To ensure that the cap structure does not affect the ability to serve as a substrate for internal modification, we injected both A-capped and G-capped U2 RNA directly into nuclei and found that both were modified to a comparable extent (Fig. 2 A, compare lanes 4 and 5, and 2 B, compare lanes 10–12 and 13–15).

To confirm these results, we physically separated oocyte nuclei from the cytoplasm under oil (Lund and Paine, 1990) and then injected U2 snRNA containing either the A or G cap directly into the isolated nuclei or the enucleated oocytes. Again, only U2 RNA recovered from isolated nuclei, not from the enucleated oocytes, was pseudouridylated (Fig. 2 C, compare lanes 1 and 2) and 2′-O-methylated (data not shown). To ensure that the enucleated oocytes were functional, we conducted anti-Sm and anti-trimethyl guanosine cap immunoprecipitation experiments. Our data showed that both G- and A-capped U2 snRNAs injected into the cytoplasm were able to assemble into immunoprecipitable Sm snRNPs and that G-capped U2 snRNA became 5′ trimethylated (data not shown). We conclude that pseudouridylation and 2′-O-methylation occur exclusively in the nucleus and do not require prior 5′ cap trimethylation.

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clear distribution of U2 was analyzed by fluorescence microscopy of nuclear spreads prepared 5 h after injection (Fig. 4 A). We also extracted the RNAs from the nucleus and cytoplasm and analyzed their nucleocytoplasmic distribution and stability by gel electrophoresis (Fig. 4 B). The wild-type U2 snRNA was localized to nucleoli and Cajal bodies (Fig. 4 A, U2). The U2 nucleolar signal was less than that observed for U6 snoRNA, but it was significantly greater than the level observed for U1 snRNA and the background fluorescence of uninjected oocytes (Fig. 4 A, compare U2, U1, U65, and uninjected). In contrast, the Sm-mutant U2 RNA was observed only in Cajal bodies;
no signal was detected in nucleoli (Fig. 4 A, U2sm-CD). Control RNAs (U1, U6, and tRNA) behaved as expected (Fig. 4 B). The correlation between nucleolar localization and efficient modification suggests that U2 internal modifications take place in the nucleolus.

**Nucleolar Targeting of Sm-mutant U2 Restores Internal Modification**

The observation that a double point mutation in the Sm binding site resulted both in a loss of nucleolar localization and internal modification of U2 RNA (Sm-mutant U2) prompted us to test whether targeting of this RNA to nucleoli can restore internal modification. Therefore, we introduced a well-characterized nucleolar localization signal, the snoRNA Box C/D motif (containing boxes C and D and the terminal stem; Samarsky et al., 1998; Narayanan et al., 1999; Speckmann et al., 1999), into Sm-mutant U2 as well as into wild-type U2 RNA. Specifically, the 5'-terminal sequence of U14 snoRNA, including box C and one strand of the terminal stem, was fused to the 5' terminus of U2, while the 3'-terminal sequence of U14, including box D and the other strand of the terminal stem, was linked to the 3' terminus of U2 (Fig. 5 A). In addition, the 5' half (nucleotides 1–100) and the 3' half (101–188) of U2 were each inserted into the same box C/D motif (Fig. 5 A). These chimeric RNAs were injected into isolated nuclei, incubated for 5 h, and assayed for modification.

Remarkably, the Sm-mutant U2-C/D motif chimera not only became modified (Fig. 5 B, lane 3), but was modified to a level equivalent to that of the wild-type U2-C/D motif chimera (Fig. 5 B, lane 2). As expected, since all the internally modified nucleotides are located exclusively in the 5' half of U2, the 5' half U2-C/D motif chimera was efficiently modified (lane 4). No modification was detected in the 3' half U2-C/D motif chimera (lane 5). Modification was dependent on the C/D motif, as changes in the box elements completely abolished modification (lane 6, and data not shown). The accuracy of the restored modifications was confirmed by RNase H site-specific cleavage directed by 2'-O-methyl RNA–DNA chimeras (for 2'-methylation; Yu et al., 1997, 1998) and by CMC modification followed by reverse transcription (for pseudouridylation, see Materials and Methods; Bakin and Ofengand, 1993; data not shown).

Visual evidence of nucleolar localization conferred on these chimeric RNAs by the box C/D motif was obtained by fluorescence microscopy using nuclear spreads prepared 5 h after injection of fluorescently labeled chimeric RNAs into oocyte nuclei. Fig. 6 shows that introduction of the C/D motif indeed resulted in targeting of the Sm-mutant U2-C/D motif chimera and the 5' half U2-C/D motif chimera to nucleoli (U2sm-CD and 5' half U2-CD). The nucleolar localization of these chimeric RNAs was dependent on the presence of the C/D motif, as point mutations within the motif resulted in loss of localization to nucleoli but not Cajal bodies (Fig. 6, 5' half U2-CDΔC). Introduction of the C/D motif into wild-type U2 did not alter its nucleolar targeting (compare Fig. 6, U2-CD and Fig. 4 A, U2).

Although our observations suggest that the nucleolus is the site of U2 internal modification, U2 constructs that become efficiently modified are detected in both nucleoli and Cajal bodies. We attempted to assess any potential role of Cajal bodies in modification by fusing a 5' fragment (nucleotides 19–46) of U2 RNA to a 3' half sequence of U7 snRNA, which contains a distinct Sm binding site known to target U7 snRNAs to Cajal bodies (Wu et al., 1996). Surprisingly, this U2-U7 chimera became modified (Fig. 7 A, lane 3). As expected, the RNA localized to Cajal bodies (Fig. 7 B, MIDU2-U7). However, we also detected nucleolar signals that were weak but distinctly greater than background, suggesting that these U2-U7 chimeric constructs may transiently cycle through nucleoli for modification (see Discussion).

**Discussion**

Using the *Xenopus* oocyte microinjection system, we have demonstrated that U2 internal modification is a nuclear event. A combination of mutant analyses and fluorescence microscopy further showed that efficiently modified U2 RNA variants are localized to nucleoli (and Cajal bodies),...
but that RNAs that are not localized to nucleoli are not modified. Strikingly, a modification-deficient Sm-mutant U2 RNA can be converted to a modification proficient state by addition of an exogenous nucleolar localization signal (the box C/D motif). Our findings suggest that U2 internal modifications occur in the nucleolus.

Previous studies suggested that U2 internal modification might not occur in the nucleolus. Ganot et al. (1999) reported that a fragment of U2 was not modified when fused to an rRNA minigene and transcribed by RNA Pol I in the nucleolus. Although the studies were done using a different cell system (mouse versus Xenopus oocytes), two additional points should be considered. First, it is possible that the short U2 sequence (22 nucleotides), fused to a large rRNA sequence, formed a structure that prevented U2 from being accessed by the modifying machinery. Second, modification of U2 may require prior exposure to other nuclear environments, which might be forfeited when the RNA is synthesized in the nucleolus. In our experiments RNAs were injected into nuclei of oocytes, which may allow trafficking of the RNA in the nucleus and association with relevant nuclear factors or structures. In other work it was shown that pseudouridylation of position 44 of yeast U2 snRNA can be catalyzed by a yeast tRNA pseudouridylase, Pus1p (Massenet et al., 1999), which resides in the nucleoplasm (Simos et al., 1996). Several other known yeast pseudouridylases, including Cbf5p (NAP57), were tested and found to be negative (Massenet et al., 1999), suggesting that pseudouridylation of this site is not mediated by snoRNPs in the nucleolus in yeast. It is possible that internal modification of yeast Sm snRNAs is different from that of vertebrate Sm snRNAs and that modification of the other residues of yeast Sm snRNAs is catalyzed by snoRNPs in the nucleolus.

**Do Cajal Bodies as Well as Nucleoli Harbor Machinery for Internal Modification of U2?**

U7 snRNA is targeted to Cajal bodies by its Sm site (Wu et al., 1996), but is not detected in nucleoli (Fig. 7 C; Wu et al., 1996). We were surprised to find that when attached
to the U7 Sm binding site, a 5′ fragment of U2 was efficiently modified (Fig. 7, A and B). The U2-U7 chimera was targeted to Cajal bodies, but distinct nucleolar signals were also observed (Fig. 7 C). We favor the explanation that the U2-U7 chimeras are modified during a transient association with nucleoli. The work of several labs suggests that many types of RNAs progress through nucleoli and Cajal bodies during their maturation, even if they ultimately come to reside in other cellular subcompartments (Pederson, 1998; for review see Olson et al., 2000). For instance, box C/D snoRNAs, including U3, U8, and U14, transiently associate with Cajal bodies before localization to nucleoli (Narayanan et al., 1999; Speckmann et al., 1999). On the other hand, U6 snRNA transiently localizes to nucleoli (Narayanan et al., 1999; Speckmann et al., 1999). On the other hand, U6 snRNA transiently localizes to nucleoli. (Lange and Gerbi, 2000; Lukowiak, A. Narayanan, R.M. Terns, and M.P. Terns, manuscript submitted for publication) presumably for modification mediated by box C/D snoRNAs (Tycowski et al., 1998; Ganot et al., 1999). Fibrillarin and NAP57, which are the presumed 2′-O-methylase and pseudouridine synthase components of the box C/D and box H/ACA snoRNPs, respectively, have been reported to be present in both nucleoli and Cajal bodies (Raska et al., 1991; Meier and Blobel, 1994; Gall et al., 1999). The Lamond group has also shown that mammalian spliceosomal snRNPs accumulate in Cajal bodies, which, after treatment with a phosphatase inhibitor, coalesce with nucleoli (Lyon et al., 1997; Sleeman et al., 1998). The accumulation of spliceosomal snRNPs in Cajal bodies and nucleoli is a transient event that occurs before localization to speckles (Sleeman and Lamond, 1999), suggesting that Cajal bodies and/or nucleoli may be the site(s) for spliceosomal snRNA modification (Böhm et al., 1995a, b; Sleeman and Lamond, 1999). Our data indicate that internal modification of U2 snRNA occurs in the nucleolus, but does not exclude a role for Cajal bodies. Further analyses are needed to establish whether U2 internal modification also takes place within Cajal bodies.

Role of the Sm Binding Site in Intracellular Trafficking

We have shown that the Sm binding site is required not only for U2 internal modification but also for nucleolar localization. A double point mutation in the U2 Sm binding site results in a modification defective phenotype and disruption of nucleolar targeting, even though a significant fraction of the RNA remained in nuclei (including Cajal bodies) at the time analyzed (Fig. 4). This observation suggests a role for the Sm binding site in nucleolar targeting. Interestingly however, two other Sm snRNAs, U1 and U7, were detected in Cajal bodies but not nucleoli (Figs. 4 and 7). Why do these two Sm RNAs have different intranuclear localization profiles? If U2 is internally modified in the nucleolus, where are U1 and U7 modified? One explanation is that U1 and U7 may pass through nucleoli too quickly to generate detectable signals. The different nucleolar transit times of the RNAs could correlate with the number of modifications acquired by the various RNAs. U2 is modified at 23 internal sites (by pseudouridylation and 2′-O-methylation; Massenet et al., 1998; Reddy and Busch, 1988) and is detected in the nucleolus (Fig. 4), whereas U1 is modified at only five sites (Reddy and Busch, 1988; Massenet et al., 1998) and is not detected there (Fig 4). Similarly, U7 snRNA contains no pseudouridylation (Fig. 7 B, lane 1) (it is unknown whether U7 is 2′-O-methylated) and is not detected in the nucleolus. Interestingly, we also detect a transient nucleolar localization of U6 snRNA (Lukowiak, A. Narayanan, R.M. Terns, and M.P. Terns, manuscript submitted for publication), which contains 13 internal modifications (Reddy and Busch, 1988). Finally, addition of a 5′ fragment (nucleotides 19–46) of U2, which lacks an Sm site but contains 10 sites of modification, to U7, which is not detected in nucleoli, results in detectable levels of nucleolar localization (Fig. 7; U2-U7). Alternatively, differences in nucleolar transit time may be a result of functional differences between the Sm binding sites. Comparing sites from various species, we note a striking difference in the sequences for U1 and U7 versus U2: AUUUGUG for U1 (AUUUCUG in frog), AUUUGUC for U7, and AUUUUUUG for U2. Perhaps a difference in the fifth position of the Sm site mediates association with distinct factors that affect nucleolar trafficking rates. Interestingly, the Sm binding sites of U1 and U5 (AUUUUUUG, more similar to U2 than to U1 and U7 Sm binding sites) are not functionally exchangeable in protein binding (Jarmolowski and Mattaj, 1993). Even between U1 and U7, the Sm binding sites are not exchangeable, since replacement of the U7 Sm site with the U1 Sm site generates a nonfunctional U7 snRNP (Stefanovic et al., 1995). Further studies are required to understand the detailed differences between the functions of various Sm binding sites. Our work indicates that the Sm site targets U2 to nucleoli where it is modified. Other snRNAs may transit through nucleoli with different kinetics.

At What Stage in U2 Biogenesis Does Modification Occur?

Although our results demonstrate that U2 internal modification occurs in the nucleus rather than the cytoplasm, we still have no definitive answer to the question of whether modifications are introduced before export to the cytoplasm or after reentry into the nucleus. For three reasons, we consider the latter more likely. First, it is well established that the Sm snRNAs are rapidly transported to the cytoplasm after synthesis in the nucleus (Yang et al., 1992; Terns et al., 1993; Jarmolowski et al., 1994). By contrast, U2 internal modification is relatively slow, in our hands requiring more than 6 h to complete (Yu et al., 1998; data not shown). Second, we have shown previously that internal modifications directly contribute to the conversion of the 12S U2 particle into the functional 17S particle, which possesses at least nine extra proteins (Behrens et al., 1993). This suggests that the internal modifications are introduced at the stage between 12S and 17S snRNP assembly, i.e., after U2 transport back to the nucleus. Third, we have demonstrated here that an intact Sm site, and presumably the binding of Sm proteins, is required for U2 internal modification (Fig. 3). Since assembly with Sm proteins normally occurs in the cytoplasm (Mattaj, 1988), internal modification could occur only after reentry into the nucleus.

Is Internal Modification of Sm snRNAs Guided by snoRNAs in the Nucleolus?

Our evidence that U2 internal modification occurs in the nucleolus strongly supports the notion that the nucleolus...
harbors the modifying machinery responsible for pseudo-uridylation and 2’-O-methylation of several different types of RNAs. These include ribosomal RNAs (Pol I transcripts; Cavaille et al., 1996; Kiss-Laszlo et al., 1996, 1998; Ganot et al., 1997; Ni et al., 1997; Bortolin et al., 1999), U6 snRNA (a Pol III transcript; Tycowski et al., 1998; Ganot et al., 1999; Lange and Gerbi, 2000), and U2 and perhaps other Sm snRNAs (Pol II transcripts; this study). Modification of eukaryotic rRNAs and U6 snRNA is catalyzed by snoRNPs, where the snoRNA component serves as a guide to direct modification at a specific site (Cavaille et al., 1996; Kiss-Laszlo et al., 1996, 1998; Ganot et al., 1997, 1999; Ni et al., 1997; Tycowski et al., 1998; Bortolin et al., 1999). Are the Pol II–transcribed Sm snRNAs (U2 and others) modified by the same mechanism? Since all of these RNAs share a common modification subcompartment, the nucleolus, we predict the existence of similar snoRNPs responsible for internal modification (2’-O-methylation and pseudouridylation) of Pol II–transcribed Sm snRNAs.

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