Signal recognition particle RNA localization within the nucleolus differs from the classical sites of ribosome synthesis

Joan C. Ritland Politz
University of Massachusetts Medical School

Laura B. Lewandowski
University of Massachusetts Medical School

Thoru Pederson
University of Massachusetts Medical School

Follow this and additional works at: http://escholarship.umassmed.edu/oapubs

Part of the Biochemistry Commons, Molecular Biology Commons, and the Pharmacology Commons

Repository Citation
Politz, Joan C. Ritland; Lewandowski, Laura B.; and Pederson, Thoru, "Signal recognition particle RNA localization within the nucleolus differs from the classical sites of ribosome synthesis" (2002). Open Access Articles. 923.
http://escholarship.umassmed.edu/oapubs/923
Signal recognition particle RNA localization within the nucleolus differs from the classical sites of ribosome synthesis

Joan C. Politz, Laura B. Lewandowski, and Thoru Pederson

Introduction

That the nucleolus is the site of ribosome biosynthesis was discovered over 35 years ago (Vincent and Miller, 1966), but recently it has emerged that additional processes are likely to be taking place in this region of the nucleus (Pederson, 1998a, 1998b; Carmo-Fonseca et al., 2000; Olson et al., 2000; Visintin and Amon, 2000). One initial clue to this idea was the discovery that signal recognition particle (SRP)* RNA injected into the nucleus of mammalian cells transiently passes through the nucleolus before its appearance in the cytoplasm (Jacobson and Pederson, 1998). The biological relevance of this observation was indicated by the finding that the nucleolar transit depends on particular sequence domains in the SRP RNA molecule (Jacobson and Pederson, 1998). These results were reinforced by the subsequent finding that endogenous SRP RNA as well as expressed GFP-labeled SRP protein components are present in the nucleolus of mammalian cells (Politz et al., 2000). Subsequent work in Saccharomyces cerevisiae has also revealed the nucleolar presence of SRP proteins (Ciufo and Brown, 2000; Grosshans et al., 2001).

The significance of SRP components in the nucleolus is not presently understood. A plausible explanation is that the SRP is at least partially assembled in the nucleolus. In addition, because mature SRP interacts with cytoplasmic ribosomes, SRP assembly may be coordinated with ribosome assembly within the nucleolus, perhaps as a mutual quality control mechanism assuring proper assembly of each particle (Pederson and Politz, 2000). Because the tripartite structural organization of the nucleolus has been so extensively studied and defined in relation to the steps of ribosome biosynthesis (Goessens, 1984; Hadjioff, 1985; Hernandez-Verdun, 1991; Spector, 1993; Shaw and Jordan, 1995; Scheer and Hock, 1999; Huang, 2002), we reasoned that an important step toward defining the role of SRP RNA in the nucleolus, and any potential interaction with ribosomal components, would be to determine the precise sites within the nucleolus at which it is localized.
Results

Previously, we detected endogenous SRP RNA in the nucleoli of rat NRK fibroblasts by in situ nucleic acid hybridization using conventional phosphodiester (PO) backbone oligodeoxynucleotide probes (Politz et al., 2000), but the signal intensity was not high enough for high resolution intranucleolar mapping studies. Oligonucleosides linked by peptide bonds (peptide nucleic acid [PNA] oligos) form more stable DNA/RNA hybrids (Nielsen, 1999); therefore, we investigated the use of PNA oligos as a way to increase the detection of SRP RNA in the nucleolus. Fig. 1 shows that we detected an increase in both nucleolar and cytoplasmic signal of SRP RNA using PNA oligos as compared with PO backbone probes. The specificity of this result was demonstrated by the lack of detectable hybridization when a PNA oligo complementary to the evolutionary diverged yeast SRP RNA was used (Fig. 1 E). Signal was also reduced to background levels if cells were treated with RNase before in situ hybridization (unpublished data), demonstrating that the PNA probes were hybridizing to RNA as expected. It should be noted that the respective efficiencies with which PNAs and PO oligos hybridize to identical RNA target sequences cannot be directly compared because PNAs are typically much shorter than PO backbone probes and often must be designed to target different sequences because of PNA solubility limitations.

The three domains of the nucleolus defined originally by electron microscopy and now as well by specifically localized proteins are (a) the fibrillar centers, in which the rDNA resides, for which upstream binding factor (UBF) is a marker; (b) the dense fibrillar component (DFC), into which nascent ribosomal RNA extends and rRNA processing and ribosome assembly commences, for which fibrillarin is a marker; and (c) the granular component, which contains partially processed rRNA and hosts the final stages of ribosome assembly, for which the protein B23 is a marker (Goessens, 1984; Hadjiolov, 1985; Hernandez-Verdun, 1991; Spector, 1993; Shaw and Jordan, 1995; Scheer and Hock, 1999).

UBF binds upstream of the rDNA promoters, and is thought to bind both actively transcribing genes as well as those open for transcription (Junera et al., 1997). As shown in Fig. 2 (A–F), SRP RNA only minimally overlapped with sites in NRK nucleoli that were immunostained for UBF. Similarly low levels of colocalization were also observed in HeLa cells expressing GFP-UBF to mark the fibrillar centers (Fig. 2, G–L). Next, three-dimensional optical stacks of immunostained, in situ–hybridized cells were processed using a constrained interactive deconvolution algorithm (Swedlow et al., 1997; Wallace et al., 2001) to obtain a higher resolution map of the intranucleolar space occupied by SRP RNA and UBF. Again, at this refined resolution, only a limited amount of overlap at the edges of a few fibrillar centers was observed (Fig. 3).

SRP RNA was detectable in only low levels in the DFC, as defined by immunostaining of fibrillarin (Fig. 4) or by the localization of a yellow fluorescent protein fusion of fibrillarin (unpublished data). Colocalization between SRP RNA signal and fibrillarin protein was limited to a minimal spatial concordance of a minor fraction of SRP RNA along the edges of the DFC.

The same techniques were next used to map SRP RNA with respect to the granular component of the nucleolus. B23 is a multifunctional protein that is involved in nucleolar rRNA processing and ribosome assembly (Savkur and Olson, 1998; Szebeni and Olson, 1999; Okuwaki et al., 2002). It is primarily localized in the granular component in many cell types (Ochs et al., 1996). In agreement with these previous observations, we observed no overlap between concentrated regions of GFP-B23 and fibrillarin immunostaining in the NRK cells used in this study (unpublished data), demonstrating that B23 is a good marker for the granular component in these cells. A portion of SRP RNA overlapped with B23 in both immunostaining experiments (Fig. 5 F, arrow, yellow) and in the nucleoli of cells expressing GFP-B23 (Fig. 5 L, yellow). As can be seen in Fig. 5 (L), these areas of colocalization were distributed throughout the nucleolus, and were not limited to the edges of two adjoining signal regions. However, although colocalization was observed, a significant portion of SRP RNA was present in nucleolar regions where B23 was not concentrated (Fig. 5, F and L, red within nucleolus).

Using constrained iterative deconvolution to increase the resolution of subnucleolar regions, the intensity distribution of SRP RNA signal (Fig. 6, red) was found to clearly differ from that of B23 (Fig. 6 F, green). Specifically, the most concentrated regions of SRP RNA (Fig. 6, G and H, red peaks in linescans) often did not overlap with the most concentrated regions of B23 (Fig. 6, G and H, green peaks in
Although the fraction of SRP RNA that colocalized with B23 varied among nucleoli in the same cell and between cells, all nucleoli contained some SRP RNA signal that was concentrated in intranucleolar regions where B23, fibrillarin, and UBF were least concentrated.

Because these results so far indicated that a considerable portion of SRP RNA resides in intranucleolar regions that do not correspond with any of the three classically defined sites of ribosome synthesis, we sought to confirm this conclusion by determining the spatial relationship of SRP RNA to 28S ribosomal RNA in the nucleolus by performing double in situ hybridization experiments. As shown in the deconvolved images in Fig. 7, although some SRP RNA signal was coincident with 28S rRNA (Fig. 7 F, yellow signal; overlapping green and red peaks in linescans), a significant portion was not (Fig. 7 C, red; red peaks in linescans). Thus, all our results taken together reveal a region extending throughout the nucleolus from which the markers characteristic of the three classical stations of ribosomal biosynthesis are absent (or present in low concentration), and in which 28S rRNA is also deficient. SRP RNA is concentrated within this previously unidentified region of the nucleolus.

**Discussion**

The discovery that there is a connection between the nucleolus and SRP RNA was first made when fluorescent SRP RNA was microinjected into the nucleus of mammalian cells and observed to transiently pass through the nucleolus (Jacobson and Pederson, 1998). It was subsequently found that endogenous SRP RNA is also present in the nucleolus of mammalian cells as are some of the SRP's protein components (Politz et al., 2000). Our goal in the present study was to determine the intranucleolar distribution of SRP RNA at higher spatial resolution than in our earlier work, and this was made possible by our use of PNA probes for in situ hybridization. This
resulted in a substantial increase in the detection of SRP RNA in the nucleolus. Indeed, the levels of SRP RNA detected in the nucleolus with PNA probes were similar to or greater than, on an average intensity per pixel basis, that observed in the most SRP RNA–rich regions of the cytoplasm. Although we do not know the efficiency with which our PNA probes detect SRP RNA, it is to be noted that SRP RNA was readily observed biochemically in previous studies with purified rat hepatoma and HeLa cell nucleoli (Reddy et al., 1981; Mitchell et al., 1999). Our results confirm these cell fractionation studies and further suggest that, at least as determined by in situ nucleic acid hybridization, the level of SRP RNA in the nucleolus is quantitatively substantial.

The major goal of the present study was to resolve the intranucleolar distribution of SRP RNA within the nucleolus, particularly in relation to the stations of ribosome synthesis as defined by the classical tripartite organization of the nucleolus. Because we have hypothesized that SRP RNA in the nucleolus is related to the regulated construction of the overall translation machinery (Pederson and Politz, 2000), we wished to determine whether SRP RNA might be uniquely present in one, and only one, of the three nucleolar compartments, as a clue to its possible role at a discrete step in ribosome biosynthesis. In the present study we have used the RNA polymerase I–specific transcription factor UBF to demarcate the fibrillar centers, the protein fibrillarin as a fiduciary landmark of the DFC of the nucleolus, and the protein B23 to identify the rRNA containing regions of the granular component. In all cases, we used both specific antibodies as well as fluorescent protein expression to identify these components of the nucleolus, combined with the highly sensitive detection of SRP RNA. Additionally, we performed dual in situ hybridization experiments to resolve the spatial relationship between 28S rRNA and SRP RNA. We found
that very little SRP RNA was present in the fibrillar centers and the DFC. Rather, a portion of the SRP RNA was localized with B23 in the granular component and, surprisingly, the remainder was concentrated in rRNA deficient regions of the nucleolus.

The intranucleolar distribution pattern of SRP RNA observed in the present investigation renders unlikely a model where the SRP RNA is intimately associated with precursor ribosomes throughout their synthesis and assembly. SRP RNA is not concentrated in either fibrillar centers or the DFC and therefore it probably does not play a role during rRNA transcription or early processing in any direct, interactive fashion, although it remains possible that very low concentrations of SRP RNA might carry out functional interactions with formative ribosomes at these sites.

Rather, our results indicate that SRP RNA might interact with nucleolar ribosomes much later during assembly, perhaps within the B23-rich portion of the granular component. However, the SRP RNA that was localized within the granular component displayed a heterogeneous distribution and was not closely correlated with the abundance of B23 protein. The most concentrated regions of SRP RNA were often not coincident with concentrated regions of B23 protein, indicating that portions of SRP RNA and B23 protein have different locations within the granular component.

Studies at the electron microscopic level have generally conveyed the granular component as having a fairly compact and generally homogenous particulate appearance (for review see Hadjiolov, 1985). A protein termed No55 identified by Ochs et al. (1996) using a rare human patient autoimmune serum, and not presently known to be involved in ribosome biosynthesis, was found to be distributed fairly homogeneously throughout the granular component. However, B23, which is involved in rRNA processing step (Savkur and Olson, 1998; Okuwaki et al., 2002), exhibits a somewhat uneven distribution within the granular component (Ochs et al., 1996; unpublished data). Our finding that a substantial portion of the SRP RNA is present in regions at which B23 is not concentrated (and yet are not

Figure 5. SRP RNA signal colocalizes with portions of the nucleolar granular component. (A–F) SRP RNA and B23 immunostaining. (A) B23. (B) SRP RNA. (C) Merged. (D–F) Nearest neighbor deconvolutions of A–C, respectively. (G–L) SRP RNA and GFP-B23. (G) GFP-B23. (H) SRP RNA. (I) Merged. (J–L) Nearest-neighbor deconvolutions of G–I, respectively. (C, F, I, and L, insets) Nucleolus at higher magnification.
DFC) may indicate that within the granular component itself, particular activities are spatially organized into different functional domains. Such activities may require different levels of SRP RNA as compared with B23. Indeed, it is not known whether all of the “granules” of the granular component that are visualized by electron microscopy actually represent nascent ribosomal subunits, or whether there are other particles present also. Our results raise the possibility that a portion of the particulate texture of the granular component might in part represent nascent SRPs, and

Figure 6. **Constrained iterative deconvolution of SRP RNA and B23 protein signal in the nucleolus.** Raw (A) and deconvolved (D) midplane of a NRK cell nucleolus containing GFP-B23. Raw (B) and deconvolved (E) midplane of same nucleolus showing SRP RNA signal. (C and F) Overlays of B23 (green) and SRP RNA (red) images showing regions of similar intensities in yellow. (G) Linescan of left vertical line in F. (H) Linescan of right vertical line in F. Each linescan shows, from left to right, the intensities (arbitrary units) of B23 (green) and SRP RNA (red) along the line indicated in F, proceeding downward from top to bottom. Linescans are displayed with the minimal linescan intensity at the origin of the y-axis.

Figure 7. **SRP RNA and 28S rRNA distribution in the nucleolus revealed by double in situ hybridization.** Deconvolved midplane of NRK cell containing 28S rRNA signal (A), SRP RNA signal (B), and these two images combined (C) where 28S rRNA is green and SRP RNA is red. Overlapping regions of similar intensities are shown in yellow. (D–F) Deconvolved images of nucleolus shown in boxed area in A. (D) 28S rRNA signal. (E) SRP RNA signal. (F) Color combined images of D and E coded as in the C above. (G) Linescan of leftmost line in F. (H) Linescan of rightmost line in F. Each linescan shows, from left to right, the intensities (arbitrary units) of 28S rRNA (green) and SRP RNA (red) along the line indicated in F, proceeding downward from top to bottom. Linescans are displayed with the minimal linescan intensity at the origin of the y-axis.
SRP RNA and the nucleolus | Politz et al. 417

Antibodies and immunofluorescence
UBF, a RNA polymerase I–specific transcription factor specifically localized in fibrillar centers of the nucleolus, was detected with an antibody (Chan et al., 1991; Roussel et al., 1993; Dousset et al., 2000) provided by Daniele Hernandez-Verdun (Institut Jacques Monod, Paris, France). Fibrillarin, a protein specific to the DFC of the nucleolus, was detected with a monoclonal antibody (Ochs et al., 1983) provided by Pui-Kwong Chan (Baylor College of Medicine, Houston, TX). Immunostaining and sequential in situ hybridization was performed using a minor modification of a protocol provided by Sui Huang (Northwestern University School of Medicine, Chicago, IL). HeLa cells were grown on coverslips and fixed for 12 min in 4% (vol/vol) formaldehyde in PBS, containing 5 mM MgCl₂. Fixation and all subsequent steps were performed at room temperature unless otherwise noted. Coverslips were washed three times in PBS containing 1% bovine serum albumin (PBSB) for 5–10 min each, incubated for 5 min in 0.5% Triton X-100 in PBSB, and then again washed three times in PBSB for 5–10 min each time. Coverslips were then incubated with the desired primary antibody (1:2,000 for anti-B23, 1:75 for anti-fibrillarin and 1:100 for anti-UBF) in PBSB for 1 h in a humidified chamber, washed three times in PBSB (10 min each), incubated with the appropriate secondary antibody (1:200 anti–mouse IgG for B23, 1:80 anti–mouse IgG for fibrillarin and 1:750 anti–human IgG for UBF; all secondary antibodies were from Sigma) in PBSB in humidified chambers for one hour. Coverslips were then washed three times in PBSB (10 min each). Cells were then refixed by immersing the coverslips in 2% (vol/vol) formaldehyde in PBS containing 5 mM MgCl₂ for 5 min, followed by three rinses in 70% (vol/vol) ethanol. Coverslips were stored in absolute ethanol at 4°C for 18–24 h before performing in situ hybridization as described above.

Fluorescent fusion proteins
Plasmids encoding green fluorescent protein fusions to human UBF and human B23 were obtained from Sui Huang (Northwestern University School of Medicine; Chen and Huang, 2001). A plasmid encoding a yellow fluorescent protein fusion to human fibrillarin was obtained from Angus Lamond (University of Dundee, Dundee, Scotland). These fluorescent protein–encoding plasmids were transfected into NRK cells with Lipofectamine 2000 (Invitrogen Life Technologies) following manufacturer’s instructions. In the case of YFP-fibrillarin, a stable cell line was constructed. HeLa cells were transfected as above and 18 h later the medium was replaced with fresh medium containing 800 µg/ml Geneticin (G-418; Gibco BRL). The medium was changed twice weekly and the cells were cultured for 6–8 wk before subcloning and further selection.

Microscopy and image processing
Results were analyzed with a Leica DMIRB microscope equipped with a 100× objective (N.A. 1.4) and appropriate filter sets, and images were captured using a Quantix 57 CCD camera (Roper Scientific Photometrics). For high resolution spatial mapping, three-dimensional optical stacks (containing 21 consecutive 0.25 micron slices) were captured using a PIFOC microscope focusing drive (Polytec PI). Images were dark current subtracted, intensity scaled, and in some cases, subjected to two-dimensional nearest-neighbor deconvolution using Metamorph software. Alternatively, image stacks were processed by constrained iterative deconvolution (Applied Precision) using an empirical point-spread function (Swedlow et al., 1997; Wallace et al., 2001).

We thank Christina N. Alavian in our laboratory for her important role in the initial application of PNA oligos for detection of SRP RNA in the nucleolus. We are very grateful to Jason Swedlow (University of Dundee) for his most generous assistance in carrying out the constrained iterative deconvolution of some of our images. We also thank Craig Brumwell in our laboratory for constructing the YFP–fibrillarin stable cell line, and Christine Powers for help with the B23 experiments. We also gratefully acknowledge the following scientists, who provided key materials: Pui-Kwong Chan, Edward Chan, Daniele Hernandez-Verdun, Sui Huang, Angus Lamond, and Eng Tan.

This work was supported by National Institutes of Health grant GM21595, which requires us to state that the content of our paper is not the official position of the U.S. government.

Submitted: 6 August 2002
Revised: 25 September 2002
Accepted: 25 September 2002