Environmental cues induce a long noncoding RNA–dependent remodeling of the nucleolus

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ABSTRACT The nucleolus is a plurifunctional organelle in which structure and function are intimately linked. Its structural plasticity has long been appreciated, particularly in response to transcriptional inhibition and other cellular stresses, although the mechanism and physiological relevance of these phenomena are unclear. Using MCF-7 and other mammalian cell lines, we describe a structural and functional adaptation of the nucleolus, triggered by heat shock or physiological acidosis, that depends on the expression of ribosomal intergenic spacer long noncoding RNA (IGS lncRNA). At the heart of this process is the de novo formation of a large subnuclear structure, termed the detention center (DC). The DC is a spatially and dynamically distinct region, characterized by an 8-anilino-1-naphthalenesulfonate–positive hydrophobic signature. Its formation is accompanied by redistribution of nucleolar factors and arrest in ribosomal biogenesis. Silencing of regulatory IGS lncRNA prevents the creation of this structure and allows the nucleolus to retain its tripartite organization and transcriptional activity. Signal termination causes a decrease in IGS transcript levels and a return to the active nucleolar conformation. We propose that the induction of IGS lncRNA by environmental signals operates as a molecular switch that regulates the structure and function of the nucleolus.

INTRODUCTION Cellular organelles are composed of distinct subdomains that assemble into functional architectures. The individual proteins that take part in this organization are highly mobile molecules that continuously associate and dissociate in an effort to maintain dynamic frameworks of functional interactions (Vikstrom et al., 1992; Phair and Misteli, 2000; Misteli, 2001a,b; Shav-Tal et al., 2004). A primary example of molecular dynamics occurs within the nucleolus. This subnuclear organelle is composed of three distinct compartments that assemble around ~400 tandem repeats of ribosomal DNA (rDNA), thereby consolidating the different factors required for ribosomal biogenesis. RNA polymerase (pol) I transcription is believed to take place at the interface between the fibrillar centers (FCs) and dense fibrillar components (DFCs), with further posttranscriptional maturation of newly synthesized rRNA occurring within the DFC. Final processing of rRNA and assembly with ribosomal proteins takes place in the granular component (GC). Through its tremendous output of rRNA, the nucleolus plays a central role in providing the cell with the protein synthesis capabilities necessary to sustain growth and proliferation (Melese and Xue, 1995; Derenzini et al., 1998; Scheer and Hock, 1999; Lempiainen and Shore, 2009).

The remarkable plasticity of the nucleolus is demonstrated during each mammalian cell cycle, with the structure disassembling at the onset of mitosis and reassembling at mitotic exit. It is also evident under conditions of cellular stress. Actinomycin D (ActD)–mediated inhibition of transcription leads to irreversible redistribution of both nucleolar and nucleoplasmic proteins into cap structures at the periphery of the nucleolus (Journey and Goldstein, 1961; Reynolds et al., 1964; Shav-Tal et al., 2005) and a correspondingly dramatic alteration in the nucleolar proteome (Andersen et al., 2005). Changes in the nucleolar proteome are also observed in response to DNA damage and viral infection (Hiscox, 2007; Moore et al., 2011). Reversible disorganization of nucleolar structure can be induced by...
treatment with the casein kinase 2 (CK2) inhibitor 5,6-dichloro-β-D-ribofuranosylbenzimidazole (Scheer et al., 1984; Panse et al., 1999). Under these conditions, RNA pol I remains active, while nucleoli disassociate into substructures and disperse throughout the nucleoplasm as a necklace of transcribing “beads.” Nucleolar reformation upon removal of the drug is CK2 driven and ATP/GTP dependent (Louvet et al., 2006). Although the functional significance of some of these responses is unclear, they demonstrate that the structure of the nucleolus is highly responsive to a variety of stimuli (Olson and Dundr, 2005). Furthermore, the nucleolus harbors many proteins that are unrelated to ribosomal biogenesis, supporting the notion that it is a plurifunctional organelle (Pederson, 1998; Boisvert et al., 2007) involved in cell cycle regulation, senescence, and stress responses (Guarente, 1997; Visintin and Amon, 2000; Olson, 2004; Boulon et al., 2010). In fact, the nucleolus has been reported to associate with additional structural elements, including perinuclear compartments in cancer cells (Matera et al., 1995; Pollock and Huang, 2009; Pollock et al., 2011) and intranucleolar bodies (Hutten et al., 2011).

A striking example of functional nucleolar plasticity involves the capture of proteins by long noncoding RNA (lncRNA) in response to changing environmental conditions (Audas et al., 2012a; Jacob et al., 2012). Increases in temperature (heat shock) trigger the induction of ribosomal intergenic spacer (IGS) RNAs located 16 kb (IGS16RNA) and 22 kb (IGS22RNA) downstream of the rDNA transcription start site (Audas et al., 2012a). Similarly, a decrease in extracellular pH (acidosis) causes the accumulation of a transcript located 28 kb (IGS28RNA) downstream of the cassette start site (Audas et al., 2012a). These RNA pol I transcripts, which are processed into 300–400 nucleotide products, immobilize select protein in the nucleolus, away from their downstream effectors. A specific example is the nucleolar sequestration of the von Hippel– Lindau (VHL) protein (Mekhail et al., 2004a), which allows hypoxia-inducible factors to evade proteosomal degradation (Maxwell et al., 1999) and transcriptionally (Semenza, 2000) and translationally (Uniacke et al., 2012) activate their target genes, thus mediating cellular adaptation to hypoxia (Semenza, 2001; Mekhail et al., 2004b). Of interest, VHL is only one of many targets of IGS IncRNA–induced nucleolar sequestration. Other factors include the heat shock protein 70 kDa (Hsp70), ring finger protein 8 (RFN8), DNA polymerase catalytic subunit δ (POLD1), and DNA methyltransferase 1 (DNMT1; Audas et al., 2012b).

Sequestered proteins are generally characterized by the presence of a nucleolar detention sequence (NoDS), a discrete amino acid sequence composed of an arginine motif (R-R-V-L) and at least two hydrophobic triplets (L-X-V-L) (Mekhail et al., 2007). The nucleolar detention pathway places the nucleolus at the heart of a systemic re-modeling of molecular networks, which appears central to the cell’s adaptation to environmental stresses (Audas et al., 2012b).

The fundamental question, however, of how the nucleolar architecture adapts to fulfill this additional role is unanswered. Here, we report that IGS IncRNAs functionally reorganize the nucleolus by regulating the formation of the nucleolar detention center (DC), a previously uncharacterized compartment that is spatially, dynamically, and biochemically distinct from the classic nucleolar domains. Formation of the DC reversibly alters the distribution of nucleolar factors and induces arrest in the synthesis of rRNA. These data highlight a role for IncRNA in the reorganization of the nucleolar space.

RESULTS

The nucleolus responds structurally to environmental stimuli

Heat shock and acidosis induce the capture and immobilization of an array of proteins in the nucleolus (Audas et al., 2012a). To investigate how the nucleolar architecture adapts to accommodate this large influx of proteins, we looked for gross morphological changes by differential interference contrast (DIC) microscopy. Of interest, we observed that ~90% of the nucleoli of heat-shocked or acidic MCF-7 cells possessed a more irregular shape, perforated with voids, compared with untreated cells (Figure 1, A and B, and

FIGURE 1: The nucleolus responds structurally to environmental stimuli. (A) DIC images of live untreated, heat-shocked, acidic, or recovered (heat shock followed with 6-h recovery) MCF-7 cells. Scale bars, 10 μm. Right, indicated nucleoli are enlarged. (B) Quantification of untreated, heat-shocked, acidic, or recovered MCF-7 cells showing altered nucleolar morphology. Columns, mean (n = 3); error bars, SEM. (C) Transmission electron micrograph of untreated, heat-shocked (30 min), acidic (60 min), or recovered MCF-7 cells. Subnucleolar compartments (GC, DFC, FC) are indicated. Black scale bars, 2 μm; white scale bars, 0.5 μm.
the consequences of this process for the functional organization of the nucleolus, we compared the localization of VHL, a well-characterized marker of nucleolar sequestration (Mekhail et al., 2004a, 2005, 2007; Audas et al., 2012a), to several nucleolar proteins. B23 and 60S ribosomal protein L27 (RPL27) were used as GC markers, fibrillarin (FBL) and GAR1 as markers of the DFC, and RNA polymerase I polypeptide A subunits 43 and 194 (RPA43 and RPA194) as markers of the FC (Figure 2A). Under standard growth conditions, VHL exhibited its normal nucleocytoplasmic distribution, whereas B23, RPL27, FBL, GAR1, RPA43, and RPA194 localized primarily to their respective nucleolar compartments (Figure 2B and Supplemental Figure S2A). In response to heat shock, VHL relocalized to the core of the nucleolus, forming a large and irregular structure (Figure 2C and Supplemental Figure S2B). A similar intranucleolar localization was observed for sequestered proteins Hsp70 (Figure 2C) and RNF8 (Supplemental Figure S2B). Surprisingly, the structure containing VHL did not colocalize with any of the known nucleolar compartments, as the resident proteins appeared to relocalize to the nucleolar periphery (Figure 2C and Supplemental Figure S2B). On stimulus termination, VHL was quickly returned to its native conformation (Figure 1, A and B, and Supplemental Figure S1A). Similar observations were made in U-87 MG, PC-3, and NIH/3T3 cells (Supplemental Figure S1, C–E). Cell viability was not affected by these treatments (Supplemental Figure S1B).

In light of these results, we decided to use transmission electron microscopy to further monitor changes of the nucleolus at the ultrastructural level. Untreated cells possessed the characteristic tripartite nucleolar organization, with FCs and DFCs distributed throughout the GC (Figure 1C). These distinctive features were lost in response to heat shock and acidosis. Unlike the classic organization described earlier, treated cells displayed nucleoli with a noticeably transformed morphology, composed primarily of electron-dense anastomosed sheets in a reticular structure (Figure 1C). On return to normal conditions, the nucleolus regained its original tripartite organization (Figure 1C). These results suggest that the nucleolus undergoes a significant, yet reversible remodeling in response to changing environmental conditions.

The nucleolus is reorganized around the detention center

The transformation of the nucleolus described earlier correlates with an influx of proteins from other regions of the cell. To understand the consequences of this process for the functional organization of the nucleolus, we compared the localization of VHL, a well-characterized marker of nucleolar sequestration (Mekhail et al., 2004a, 2005, 2007; Audas et al., 2012a), to several nucleolar proteins. B23 and 60S ribosomal protein L27 (RPL27) were used as GC markers, fibrillarin (FBL) and GAR1 as markers of the DFC, and RNA polymerase I polypeptide A subunits 43 and 194 (RPA43 and RPA194) as markers of the FC (Figure 2A). Under standard growth conditions, VHL exhibited its normal nucleocytoplasmic distribution, whereas B23, RPL27, FBL, GAR1, RPA43, and RPA194 localized primarily to their respective nucleolar compartments (Figure 2B and Supplemental Figure S2A). In response to heat shock, VHL relocalized to the core of the nucleolus, forming a large and irregular structure (Figure 2C and Supplemental Figure S2B). A similar intranucleolar localization was observed for sequestered proteins Hsp70 (Figure 2C) and RNF8 (Supplemental Figure S2B). Surprisingly, the structure containing VHL did not colocalize with any of the known nucleolar compartments, as the resident proteins appeared to relocalize to the nucleolar periphery (Figure 2C and Supplemental Figure S2B). This segregation was also visualized in three dimensions (Supplemental Figure S2C) and with the endogenous proteins Hsp70 and FBL in both MCF-7 (Supplemental Figure S2D) and U-87MG cells (Supplemental Figure S2E). On stimulus termination, VHL was...
necessary for DC integrity. Therefore we hypothesized that high-

heat-shocked and acidotic cells migrated as monomers by SDS–

chemical properties. Of interest, VHL and Hsp70 extracted from

organization of this compartment relies on a distinctive set of bio-

VHL underwent immobilization by the nucleolus (Figure 2, E–G).

ments remained mobile under heat shock and acidosis, whereas

Figure S4, C–E and G–I, and Table 1). The ability of certain nucleolar

components, as most nuclear proteins are known to be highly

the dynamic profile of these proteins changed from a state of high

their punctuate distribution in the FCs (Figure 4, A and B). Moreover,

ANS staining (Figure 4, A and B) and sequestered VHL (Supplemen-

the detention center as the detention center.

The detention center is characterized by a hydrophobic signature. (A) Live untreated, heat-shocked, acidic, and recovered (heat shock followed with 6-h recovery) MCF-7 cells stained with ANS. (B) Quantification of untreated, heat-shocked, acidic, and recovered (1, 6 h) cells showing ANS-positive nucleoli. Columns, mean (n = 3); error bars, SEM. (C) Top, localization of mCherry-VHL, Hsp70-GFP, EGFP-FBL, EGFP-GAR1, EGFP-RPA194, and mCherry-RPA43 in live heat-shocked cells stained with ANS. Bottom, localization of mCherry-VHL and EGFP-FBL in live untreated cells. Scale bars, 10 μm.

release from the nucleolus and FBL regained its normal distribu-

Next we compared the dynamic properties of the nucleolar

The detention center is characterized by a hydrophobic signature.

The immobility of proteins within the DC suggests that the structural

affinity hydrophobic interactions could be involved in maintaining the DC architecture in a static state, a possibility consistent with the requirement for hydrophobic triplets in the nucleolar detention sequence of se-

using the membrane-permeable fluorescent dye 8-anilino-

Using the membrane-permeable fluorescent dye 8-anilino-

The remodeled nucleolus is transcriptionally inactive

Remodeling of the nucleolus involves the immobilization of a wide

variety of cellular factors, including E3 ubiquitin ligases, chaperones,

and cell cycle regulators (Audas et al., 2012b). Surprisingly, we found

that several components of the ribosomal biogenesis machinery

were also targets of this pathway. RPA16 and RPA40, two essential

subunits of RNA polymerase I and III, were captured in the DC in

both heat shock and acidosis, as indicated by colocalization with

ANS staining (Figure 4, A and B) and sequestered VHL (Supplemen-

tal Figure S4, A and B). On return to normal condition, they regained

previous reports, as the endoplasmic reticulum, Golgi, and lysosomes are rich in mis-

folded proteins and ANS-binding sites (Hadley et al., 2011). In contrast, large ANS-

positive structures formed within the nucleo-

loli of MCF-7 (Figure 3C, A and B) and U-87 MG cells (Supplemental Figure S3B) during heat shock and acidosis treatments. On re-

turn to standard conditions, nucleolar ANS signal was lost (Figure 3, A and B, Supple-

cmental Figure S3C). These data indicate that the DC is character-

ized by a hydrophobic signature that is dis-


tinct from other nucleolar compartments,
measure the levels of 45S pre-rRNA transcripts in untreated and treated cells. Results revealed ~85% decrease in both heat-shocked and acidic cells relative to untreated cells (Figure 4E). The 45S pre-rRNA levels returned to their original values upon extracellular signal termination (Figure 4E). In addition to qPCR, we labeled de novo RNA transcription sites with 5-fluorouridine (5-FU). We observed an absence of nucleolar incorporation in ~95% of heat-shocked cells (Figure 4F), with a return to normal values after signal termination (Figure 4F). Similar results were observed in U-87 MG cells (Supplemental Figure S4K). Incorporation of 5-FU is impractical under acidic conditions due to the pH of the media. These results suggest that remodeling of the nucleolus is associated with a severe, yet reversible inhibition of ribosomal biogenesis.

Environment-induced remodeling of the nucleolus is not a byproduct of transcriptional inhibition

Transcriptional inhibition induces several alterations to the nucleolar architecture, including the formation of nucleolar caps (Reynolds et al., 1964; Shav-Tal et al., 2005) and the flux of a large number of different factors (Andersen et al., 2005). Given the correlation between structure and transcriptional activity under heat shock and acidity, we wanted to know whether transcriptional inhibition was sufficient to induce some of the traits described thus far. We found that ActD treatment (5 μg/ml) failed to induce DC formation, as evidenced by the lack ANS-positive nucleoli and the diffuse nucleocytoplasmic distribution of VHL (Figure 5A). Furthermore, using polypyrimidine tract–binding protein–associated splicing factor (PSF), a common marker of nucleolar caps, we observed that only ActD-mediated transcriptional arrest triggered the formation of these cap structures (Figure 5A). This suggests that this form of reorganization is distinct from that observed in response to environmental stresses, such as heat shock. In addition, whereas RPA16 and RPA40 were immobilized in the DC under heat shock conditions (Figure 5A), both proteins migrated to the nucleolar periphery and retained their mobility during transcriptional arrest (Figure 5, A and B). Lower levels of ActD that specifically inhibit rRNA synthesis (0.04 μg/ml) also failed to induce DC formation (Supplemental Figure S5). These results demonstrate that the remodeling observed in response to environmental cues such as heat shock is not simply a byproduct of transcriptional inhibition, as ActD appears to induce a different reorganization program. Our data further suggest that the impairment in rRNA synthesis (Figure 4, E and F) is likely a consequence, not a cause, of the nucleolar detention pathway.

evade capture by the DC further highlights the specificity of this intranucleolar reorganization process (Supplemental Figure S4, F and J, and Table 1).

Ribosomal biogenesis relies on a complex and compartmentalized assortment of dynamic factors in which structure and function are intimately linked (Hernandez-Verdun, 2006). Given the drastic remodeling of the nucleolar architecture and the immobilization of several key nucleolar proteins, we asked whether rRNA synthesis could be sustained under conditions of nucleolar sequestration. We used real-time reverse transcription (RT) PCR (qPCR) analysis to

FIGURE 4: Remodeled nucleoli are transcriptionally inactive. (A, B) Localization of EGFP-RPA16 (A) and EGFP-RPA40 (B) in live untreated, heat-shocked, acidic, and recovered (heat shock followed by 6-h recovery) MCF-7 cells stained with ANS. (C, D) Nucleolar FRAP analysis of the dynamic profile of RPA16 (C) and RPA40 (D) in untreated, heat-shocked, acidic, and recovered cells. (E) Relative levels of 45S pre-rRNA in untreated, heat-shocked, acidic, and recovered cells measured by real-time RT-PCR. Actinomycin D (2 h, 50 nM) was used as a control for inhibition of RNA polymerase I–dependent transcription. (F) Left, quantification of untreated, heat-shocked, acidic, and recovered cells showing nucleolar incorporation of 5-FU. Right, de novo rRNA synthesis visualized by incorporation of 5-FU in heat-shocked and recovered cells. Scale bar, 10 μm (A, B, F). Columns in E and F, mean (n = 3); error bars, SEM.
### IGS IncRNA is required to remodel the nucleolus

Sequestered proteins are immobilized on the rDNA cassette by several stimulus-specific IGS RNAs, including IGS16RNA and IGS22RNA during heat shock (Audas et al., 2012a; Figure 6A). These transcripts rapidly accumulate in response to environmental cues (Audas et al., 2012a; Figure 6B) and return to basal levels upon signal termination (Figure 6C). Induction of IGS IncRNA readily correlates with remodeling of the nucleolus, rapid sequestration of endogenous Hsp70, formation of ANS-positive DCs, and interruption of nucleolar transcription (Figure 6D). On signal termination, IGS IncRNA expression returns to basal levels (Figure 6C) and the nucleoli regain their original morphology, as indicated by Hsp70 release, loss in nuclear ANS signal, and resumption in nucleolar 5-FU incorporation (Figure 6D). As expected, RNA FISH experiments confirmed the presence of IGS IncRNA transcripts within the DC, as evidenced by colocalization with sequestered VHL (Supplemental Figure S5; Audas et al., 2012a). These IGS IncRNA-containing regions also notably excluded FBL (Supplemental Figure S6). Together these results demonstrate that the DC correlates spatially and temporally with the presence of IGS IncRNA.

Given the presence of IGS IncRNA in the DC, we asked whether these transcripts were required to mediate the nucleolar reorganization process described here. Despite the intricacy of the IGS as a transcriptional unit, we found that cells stably expressing short hairpin RNA (shRNA) targeting IGS22RNA (Figure 7A) effectively delayed the sequestration of proteins during heat shock treatment (Audas et al., 2012a), with near-complete inhibition at the 30-min time point. Stable knockdown of IGS22RNA did not affect the proper localization of the nucleolar factors B23, FBL, RPA16, and RPA194 (Figure 7B), the tripartite ultrastructure of the nucleolus (Supplemental Figure S7), or the expression level of rRNA under standard conditions (Figure 7C), but it impaired the sequestration of Hsp70 during heat shock treatment (Audas et al., 2012a; Figure 7D). We asked whether the formation of the DC would also be compromised by inhibiting IGS22RNA expression. ShIGS22 cells failed to produce any of the previously characterized hallmarks of DC formation.

During heat shock treatment these cells did not acquire ANS-positive subnuclear structures (Figure 7D), and most of the nucleoli retained their original appearance by DIC imaging (Figure 7, E and F). These observations were supported by transmission electron micrographs that show that the majority of shIGS22 cells retained their native nucleolar morphology, with clearly visible FCs, DFCs, and GC (Figure 7G).

Finally, we tested whether an absence of reorganization would allow the nucleolus to retain its transcriptional activity during extra-cellular stimulation. Using qPCR analysis of the 45S pre-rRNA transcript, we found a 65% decrease in the levels of pre-rRNA in control cells within 30 min of heat shock treatment but no changes in shIGS22 cells (Figure 7H). Consistently, 5-FU incorporation revealed that shIGS22 cells were unable to fully repress nucleolar transcription during heat shock treatment, as 88% of shIGS22-expressing cells retained nucleolar incorporation upon treatment versus 20% of the control cells (Figure 7, I and J). These data demonstrate the requirement for IGS RNA in both structural and functional remodeling of the nucleolus.

### DISCUSSION

We propose that the nucleolus readily alternates between two distinct and environment-dependent morphologies: a tripartite, transcriptionally active conformation under normal conditions, and a remodeled, transcriptionally inert conformation under specific environmental settings (Figure 8). The latter is characterized by the presence of the detention center, a region that is distinct from the well-documented and characterized nucleolar compartments (FC, DFC, and GC). These two architectures differ functionally as well as structurally. The former provides the cell with the output of rRNA required to sustain protein synthesis under growth conditions. The latter operates as a molecular prison, detaining cellular proteins away from their associated pathways. Through the inactivation of molecular networks and the interruption of rRNA synthesis, the remodeled nucleolus likely contributes to cell viability under conditions of stress (Olson, 2004; Mayer and Grummt, 2005; Boulon et al.,

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### TABLE 1: Spatio-dynamic profile of nucleolar proteins.

| Symbol     | Full name                                                                 | Intronucleolar localization | Untreated | Heat shock | Acidosis |
|------------|---------------------------------------------------------------------------|----------------------------|-----------|------------|----------|
| RPA194     | Polymerase (RNA) I polypeptide A, 194 kDa                                  | FC (mobile)                | FC (mobile)| FC (mobile)|          |
| RPA40      | Polymerase (RNA) I polypeptide C, 40 kDa                                  | FC (mobile)                | DC (immobile)| DC (immobile)|          |
| RPA16      | Polymerase (RNA) I polypeptide D, 16 kDa                                  | FC (mobile)                | DC (immobile)| DC (immobile)|          |
| RPA43      | Polymerase (RNA) I polypeptide subunit 43 kDa                             | FC (mobile)                | FC (mobile) | FC (mobile) |          |
| FBL        | Fibrillarin                                                               | DFC (mobile)               | DFC (mobile)| DFC (mobile)|          |
| GAR1       | GAR1 ribonucleoprotein                                                     | DFC (mobile)               | DFC (mobile)| DFC (mobile)|          |
| Nop1140    | Nucleolar and coiled body phosphoprotein 1                                 | DFC (mobile)               | DFC (mobile)| DFC (mobile)|          |
| B23        | Nucleophosmin B23                                                         | GC (mobile)                | GC (mobile) | GC (mobile) |          |
| NOL1       | NOP2 nucleolar protein homologue                                           | GC (mobile)                | DC (immobile)| DC (immobile)|          |
| NOM1       | Nucleolar protein with MIF4G domain 1                                     | GC (mobile)                | DC (immobile)| DC (immobile)|          |
| NOP52      | rRNA processing protein 1 homologue A                                    | GC (mobile)                | DC (immobile)| DC (immobile)|          |
| PES1       | Pescadillo                                                                | GC (mobile)                | DC (immobile)| DC (immobile)|          |
| RPL27      | 60S ribosomal protein L27                                                  | GC (mobile)                | GC (mobile) | GC (mobile) |          |
| RRP1B      | rRNA processing protein 1 homologue B                                    | GC (mobile)                | DC (immobile)| DC (immobile)|          |
| SENP3      | SUMO1/sentrin/SMT3 specific peptidase 3                                   | GC (mobile)                | DC (immobile)| DC (immobile)|          |

**TABLE 1**: Spatio-dynamic profile of nucleolar proteins.
IGS RNA to account for part of the phenotype in heat shock. Previous studies showed that IGS RNAs directly interact with NoDS-containing proteins (Audas et al., 2012a), suggesting that they may induce a nucleation event, further stabilized by high-affinity hydrophobic interactions. We suggest that IGS transcripts operate as a macromolecular switch that is of central importance to nucleolar function and organization, although further investigation will be required to fully understand the mechanisms at work in this process.

Of interest, parallel studies have shown that another IGS RNA, derived from the promoter region of rDNA, is involved in the recruitment of the chromatin-remodeling complex NoRC and in the physiological rationale for the inherent plasticity of the nucleolar architecture.

An interesting aspect of this transformation is the reliance on a central transcriptional event that produces IncRNA from discrete regions of the IGS. Under standard conditions, IGS transcripts are kept at basal levels, allowing the nucleolus to assume its classic organization and function. In contrast, the cell harnesses the inducible nature of IGS RNA upon signal activation to orchestrate a rapid and drastic remodeling of the nucleolus. This response relies on the simultaneous induction of multiple IGS transcripts, and although silencing IGS22RNA largely inhibits nucleolar remodeling, we expect IGS22RNA to account for part of the phenotype in heat shock. Previous studies showed that IGS RNAs directly interact with NoDS-containing proteins (Audas et al., 2012a), suggesting that they may induce a nucleation event, further stabilized by high-affinity hydrophobic interactions. We suggest that IGS transcripts operate as a macromolecular switch that is of central importance to nucleolar function and organization, although further investigation will be required to fully understand the mechanisms at work in this process. Of interest, parallel studies have shown that another IGS RNA, derived from the promoter region of rDNA, is involved in the recruitment of the chromatin-remodeling complex NoRC and in the
regulation. Synthesis of RNA in the nucleolus is predominantly controlled through transcription factors and chromatin-remodeling complexes. Additional regulatory mechanisms have been reported at the initiation, elongation, and termination steps (Leary and Huang, 2001; Santoro and Grummt, 2001; Lempiainen and Shore, 2009). The pathway described here differs not only in its mode of action, but also in its generality. The capture of proteins in the DC is a control of rRNA synthesis (Mayer et al., 2006, 2008). Homologies may exist between this pathway and the process described here.

The DC described here is emerging as a discrete nuclear component, with a distinct protein composition, spatial location, dynamic profile, hydrophobic signature, and biological function. The intranucleolar relocalization of many nucleolar proteins into the DC represents an unexpected mechanism of ribosomal biogenesis regulation. Synthesis of RNA in the nucleolus is predominantly controlled through transcription factors and chromatin-remodeling complexes. Additional regulatory mechanisms have been reported at the initiation, elongation, and termination steps (Leary and Huang, 2001; Santoro and Grummt, 2001; Lempiainen and Shore, 2009). The pathway described here differs not only in its mode of action, but also in its generality. The capture of proteins in the DC is a
systemic phenomenon that affects a breadth of unrelated factors, including DNA-processing enzymes, ubiquitin ligases, cell cycle regulators, and chaperones (Audas et al., 2012b). The requirement for RPA16 and RPA40 in RNA pol III suggests that IGS RNAs may also control pol III–mediated transcription. The tools introduced here, such as the ANS marker, could be used to rapidly assess whether a protein is specifically inactivated within the DC or otherwise located within the nucleolus. More generally, nucleolar morphology may be indicative of a systemic posttranslational remodeling of molecular networks through the nucleolar detention pathway (Audas et al., 2012b).

Ongoing ribosomal biogenesis is essential to sustaining the growth and proliferation of tumors (Lempiainen and Shore, 2009). In fact, processes that promote the up-regulation of rRNA synthesis are described as oncogenic, whereas those that repress it are considered tumor suppressing (Montanaro et al., 2012). For this reason, enlarged nucleolar morphology is indicative of malignancy (Derenzini et al., 1998, 2000; Maggi and Weber, 2005), and the RNA pol I machinery is emerging as a therapeutic target for the treatment of cancer (Drygin et al., 2010). Of interest, the nucleolar stresses described in this study are relevant to a variety of pathological conditions, including cancer. Extracellular acidosis, the outcome of anaerobic metabolism, is encountered in the core of tumors (Tannock and Rotin, 1989; Engin et al., 1995), during development, and in ischemic tissues, where it has protective effects (Giffard et al., 1990; Currin et al., 1991). Similarly, hyperthermia has been reported in some metabolically hyperactive tumors (Jayasundar and Singh, 2002) and, perhaps not coincidently, is also correlated with neuroprotection of ischemic brain tissues (Laptook et al., 1994; Thoresen et al., 1995). In light of these reports, it is tempting to speculate that the tumor microenvironment may inhibit cellular proliferation by inducing the nucleolar detention pathway. Future research may link IGS IncRNA–dependent remodeling of the nucleolus to various physiological and pathological conditions.

**MATERIALS AND METHODS**

**Cell culture, transfection, treatment, and imaging**

MCF-7, U-87 MG, PC-3, and NIH/3T3 cells were grown in monolayer culture in modified DMEM (high glucose) supplemented with 5% (vol/vol) fetal bovine serum and 1% (vol/vol) penicillin–streptomycin. All cultures were maintained at 37°C and 5% CO₂ in a humidified atmosphere, and passaged every 2–3 d. Cells were plated 24 h before transfection and allowed to grow to 50–60% confluency before transfection with Effectene (Qiagen, Valencia, CA) as per manufacturer’s instructions. shControl and shIGS22 cells were previously described (Audas et al., 2012a). Heat shock was performed by transferring cells to 42°C for 3 h, unless specified otherwise. Hypoxic acidosis was obtained by placing cells in acidic (pH 6.0) DMEM at 1% O₂ for 4 h, unless specified otherwise. For transcriptional inhibition, 5 μg/ml actinomycin D was added to cells for 2 h, unless stated otherwise.

**Plasmids**

Plasmids expressing VHL-GFP, RNF8-GFP, and EGFP-B23 were previously described (Audas et al., 2012a). mCherry-VHL was cloned from plasmid cDNA into an mCherry-c1 empty vector. EGFP-NOM1 (Kathleen Conklin, University of Minnesota, Minneapolis, MN), EGFP-FBL, EGFP-NOPP140, EGFP-PSF, EGFP-RPA40, mCherry-RPA43, EGFP-RPA194, EGFP-NOL1, EFY-PES1, EFY-RPL27, and EGFP-RRP1B (Angus Lamond, University of Dundee, Dundee, United Kingdom) were kindly provided. The following plasmids were obtained from Addgene (Cambridge, MA): EGFP-RPA16 (Tom Misteli, National Institutes of Health, Bethesda, MD; Dundr et al., 2002; Addgene plasmid 17657) and EGFP-SENP3 (Mary Dasso, National Institutes of Health, Bethesda, MD; Yun et al., 2008; Addgene plasmid 34554).

**Microscopy and image processing**

Images were collected by confocal microscopy (Zeiss 200 LSM510 META; Carl Zeiss, Jena, Germany) using a 63× Plan-Apochromat/1.4 numerical aperture (NA) objective. The system was controlled with Zeiss Zen software. DIC images were collected on a Zeiss AxioObserver D1 microscope using a 63× Plan-Apochromat/1.4 NA objective. All postacquisition image processing was performed using Zeiss Zen software. No gamma or other nonlinear adjustments were made to any image.

**Fluorescence recovery after photobleaching**

Live cells expressing constructs of interest were grown on 35-mm glass-bottom culture dishes and visualized by confocal microscopy (Zeiss 200 LSM510 META) in a 37°C and 5% CO₂ environmental chamber (Zeiss Incubator XL S1) using a 63× Plan-Apochromat/1.4 NA objective. The system was controlled by Zeiss Zen software for bleaching and image acquisition. Bleached areas (nucleolus) were subjected to 40 iterations at 100% argon laser strength at 488 nm, and imaging used 5% laser strength. Three prebleach measurements were taken before the photobleaching and 60- to 120-s recovery of the region of interest, as indicated. For nucleolar proteins, a full nucleolus was photobleached. For nucleocytoplasmic proteins, a region of the nucleoplasm was bleached instead. Region intensity measurements were performed by Zen software and normalized as previously described (Misteli, 2001b). All FRAP experiment data were the average of at least 10 cells.

**ANS fluorescence imaging**

ANS was purchased from Sigma-Aldrich (St. Louis, MO; A1028) and dissolved in PBS, pH 7.4, to a 10 mM stock solution. Culture medium was supplemented with ANS at a concentration of 250 μM for at least 1 h. Media were changed before imaging. ANS was excited at 405 nm and detected using a band pass filter.

**Immunofluorescence**

Cells were seeded onto 20-mm glass coverslips and fixed with CSK (4% paraformaldehyde) for 15 min. After permeabilization in 0.5% Triton X-100 for 10 min, cells were incubated with primary
antibody (1:200), washed three times in PBS, and incubated for 1 h in Alexa Fluor (Invitrogen, Carlsbad, CA) secondary antibody (1:400).

Fluorescence in situ hybridization
Cells were fixed in methanol for 5 min. Samples were blocked in digoxigenin (DIG) hybridization (Roche, Indianapolis, IN) solution and hybridized with 5′ and 3′ DIG-labeled oligonucleotide IGS22 RNA antisense (5′-DIG-TACTCGATTTCGTAAAGCTTCCAAAAGGCGCAAAGCCCTCAGTCAAGG-DIG), IGS29 RNA antisense (5′-DIG-CCGGCCCTAACGGTTATGGAATGCGAGGA-GACTTATCGGGGAATAGGAGAAGTAGC-DIG), or IGS sense (5′-DIG-CGATACTTCTCCTTTCCCCGATAAGTCCTCAGCTTCATAAATGTTAACGGCG-DIG) probes in a 50% formamide/DIG hybridization (Roche) solution at 37°C. After hybridization, cells were washed in 0.2% SSC, and probes were detected with an anti-DIG (Roche) antibody.

5-Fluorouridine labeling and detection of nascent rRNA
Live cells were incubated in 1 mM 5-FU (FSI130; Sigma-Aldrich) for 15 min before fixation in CSK (4% paraformaldehyde). Cells were then permeabilized in 0.5% Triton X-100 and immunostained using an anti-bromodeoxyuridine mouse antibody (B2531; Sigma-Aldrich) at 1:500, followed by incubation in Alexa Fluor (Invitrogen, Carlsbad, CA) secondary antibody (1:500).

Western blotting
Western blot was performed following standard protocol. Monoclonal antibodies were used to detect FLAG (Sigma-Aldrich), Hsp70 (Santa Cruz Biotechnology, Santa Cruz, CA), and glyceraldehyde-3-phosphate dehydrogenase (GAPDH; GeneTex, Irvine, CA). Bands were detected by enhanced chemiluminescence (Pierce, Rockford, IL).

Transmission electron microscopy
Cells were treated and fixed with 2% glutaraldehyde in PBS, pH 7.4. Samples were subsequently washed and postfixed with 1% osmium tetroxide before dehydration in a series of ascending alcohols. Specimens were then embedded in Spurr’s epoxy resin onto BEEM cups and sectioned on a Leica EM UC6 ultramicrotome (Leica, Wetzlar, Germany), and the resulting sections were stained with uranyl acetate and lead citrate. Samples were imaged with a JEOL 1230 transmission electron microscope (JEOL, Peabody, MA) equipped with AMT software (Advanced Microscopy Techniques, Woburn, MA).

RNA isolation, reverse transcription PCR, and real-time PCR
Total RNA was collected using TriPure reagent (Roche) according to manufacturer’s specifications. RT-PCR was performed using the two-step High Capacity cDNA Reverse Transcription Kit (Applied Biosystems, Foster City, CA), followed by standard PCR conditions. Real-time PCRs were performed using iQ SYBR Green SuperMix (Bio-Rad Laboratories, Carlsbad, CA) and read with Stratagene MX3005P (Stratagene, Santa Clara, CA). Transcripts levels were normalized to actin mRNA. Relative fold change in expression is calculated using the ΔΔCt method. The 45S pre-rRNA and actin mRNA primers were used as previously described (Feng et al., 2010).

Statistical analysis
The p values associated with all comparisons were based on paired two-tailed Student’s t tests. Results are mean (n = 3) ± SEM.

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