Mapping and engineering RNA-driven architecture of the multiphase nucleolus

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Biomolecular condensates are key features of intracellular compartmentalization^{1,2}. As the most prominent nuclear condensate in eukarvotes, the nucleolus is a multiphase liquid-like structure in which ribosomal RNAs (rRNAs) are transcribed and processed, undergoing multiple maturation steps to form the small (SSU) and large (LSU) ribosomal subunits³⁻⁵. However, how rRNA processing is coupled to the layered organization of the nucleolus is poorly understood owing to a lack of tools to precisely monitor and perturb nucleolar rRNA processing dynamics. Here we developed two complementary approaches to spatiotemporally map rRNA processing and engineer de novo nucleoli. Using sequencing in parallel with imaging, we found that rRNA processing steps are spatially segregated, with sequential maturation of rRNA required for its outward movement through nucleolar phases. By generating synthetic nucleoli in cells using an engineered rDNA plasmid system, we show that defects in SSU processing can alter the ordering of nucleolar phases, resulting in inside-out nucleoli and preventing rRNA outflux, while LSU precursors are necessary to build the outermost layer of the nucleolus. These findings demonstrate how rRNA is both a scaffold and substrate for the nucleolus, with rRNA acting as a programmable blueprint for the multiphase architecture that facilitates assembly of an essential molecular machine.

Biomolecular condensates have emerged as a ubiquitous feature of intracellular compartmentalization^{1,2,6}, forming through phase separation and related phase transitions of interacting multivalent biomolecules^{7,8}. Nuclear bodies like nucleoli and speckles are examples of such structures, concentrating DNA, RNA and proteins involved in multiple functions, including transcription and RNA processing. Among their protein and nucleic acid components, RNA can have a central role in the formation of various condensates, while the physicochemical environment within condensates is thought to impact their RNA-associated functions^{9,10}. However, dissecting this relationship between the structure of condensates and the biochemical reactions occurring within them has remained a key challenge.

The nucleolus provides an ideal model for studying condensate structure–function relationships. It is a multiphase condensate composed of three nested subcompartments: the innermost fibrillar centre (FC), middle dense fibrillar component (DFC) and outer granular component (GC)¹¹⁻¹³ (Fig. 1a). Precursor rRNA (pre-rRNA) is transcribed as a 13.3 kb transcript by RNA polymerase I (Pol I) at the FC–DFC boundary and fluxes radially outward as it is processed and assembled into preribosomal ribonucleoproteins (RNPs)^{3,4,14,15}. For example, pre-rRNA

is cleaved and modified by small nucleolar RNA (snoRNA)-guided RNPs (snoRNPs) and other processing enzymes as it progresses through the nucleolus. While rRNA maturation correlates with its outward flux, it remains unclear whether rRNA movement between phases is driven by rRNA processing, or is simply due to rRNA processing coinciding with outward transport. This question has been difficult to address owing to the tight coupling between nucleolar structure and function (rRNA transcription and processing): inhibition of Pol I transcription or disruption of rRNA processing induces considerable reorganization of nucleolar morphology^{12,16–22}. Despite these connections, how rRNA and its maturation directly contribute to the formation and organization of the multiphase nucleolus and, in turn, whether these phases gate rRNA outflux, remain poorly understood.

Addressing these questions has been hampered by the absence of techniques for dissecting where and when specific pre-rRNA processing steps occur in the nucleolar phases, and how these steps contribute to nucleolar organization. Traditional methods like pulse–chase radiolabelling or northern blotting lack spatial resolution^{23,24}, while RNA fluorescence in situ hybridization (FISH) reveals spatial localization but not processing dynamics^{25,26}. Moreover, approaches to understand nucleolar

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Fig. 1 | **Sequencing and imaging of nascent rRNA flux provides a spatiotemporal map of processing in the nucleolus.** FCs, DFCs and GCs in MCF10A cells were visualized through RNA Pol I subunit RPA194 (immunofluorescence (IF)), FBL (IF) and endogenously tagged mTagBFP2-NPM1, respectively. **a**, The three nucleolar phases: FC (green), DFC (red) and GC (blue). **b**, Cells were pulsed with 5eU (15 min) to label nascent RNA and chased to measure the rRNA flux (imaging) or cleavage and modification (sequencing). **c**, The radial outflux of 5eU-labelled pre-rRNA over time in nucleoli (dashed lines); the averaged signal around the FCs is shown. **d**, Min-max-normalized FC, DFC and GC intensities by distance from the FC centre from 4,274 nucleoli. The colour bar indicates FC, DFC and GC localization. **e**, Min-max-normalized 5eU intensities by distance from the FC centre across chase timepoints. *n* = 717, 459, 499, 603, 470, 451, 550 and 525 nucleoli. **f**, Pre-rRNA cleavage steps categorized as early (green), middle (red), late (blue) and mature (purple) based on **i**.

assembly are limited by the repetitive nature of rDNA genes, which makes it challenging to endogenously mutate and directly study how the rRNA sequence contributes to nucleolar morphology²⁷⁻²⁹. A number of studies

g, Schematic of pre-rRNA cleavage, modification (m), and outflux during assembly into SSU (in the FC and DFC) and LSU (in the FC, DFC and GC) based on **i**. **h**, 18S and 28S pre-rRNA 2'-O-methylation (2'-O-Me) (ScoreC) detected using 5eU–seq over time. The colour bar relates the chase time to 5eU peak localization (Extended Data Fig. 2a). The dashed lines demarcate the phase boundaries. *n* = 2 per timepoint. **i**, The fraction of pre-rRNA cleaved at early, middle and late sites over time. *n* = 2 per timepoint. **j**, Schematic of the RNA-FISH probes (top). Bottom, the averaged intensity around FCs (example images are shown in Extended Data Fig. 2e). **k**, The min–max-normalized RNA-FISH intensity by distance from the FC centre. *n* = 72 (5' ETS), 95 (3' ETS), 24 (site 01), 111 (site 1), 230 (site 2), 38 (site 3'), 105 (site 4'), 318 (ITS2-28S), 72 (18S) and 310 (28S) nucleoli. For **a**, **c** and **j**, scale bars, 1 μm. Data are mean ± s.e.m. (**d**, **e**, **h**, **i** and **k**).

have used 'bottom-up' in vitro reconstitution approaches using purified nucleolar proteins and rRNAs to form nucleoli mimics^{13,30,31}, but they are vast simplifications of the complexity of real nucleoli.

Here, to overcome these challenges, we introduce and use two complementary approaches that reveal insights into the relationship between rRNA processing and nucleolar structure. By combining a pulse-chase nucleotide analogue approach with sequencing and imaging, we precisely measure the kinetics of pre-rRNA cleavage and modification at single-nucleotide resolution. Our findings reveal that rRNA processing occurs in a spatially segregated manner within the nucleolus, with sequential maturation steps driving the outward progression of rRNA through nucleolar phases. Most small-subunit assembly steps occur within the DFC phase, while large-subunit assembly steps occur in both the DFC and GC phases. We also use an engineerable rDNA plasmid system to assemble synthetic nucleoli in living cells. By mutating rDNA, we find that LSU precursors are necessary for the formation of the GC phase. Notably, perturbing SSU pre-rRNA processing in both synthetic and endogenous nucleoli leads to a marked 'inversion' of nucleolar phases and a lack of rRNA outflux from the DFC, establishing an essential role of SSU processing in nucleolar phase organization and RNA flux through the nucleolar phases. Finally, we develop a mathematical model predicting how altered concentrations of rRNA intermediates could change interfacial tensions to shape the multiphase nucleolus. Taken together, our study provides a detailed dissection of how rRNA sequences and processing build and arrange the nucleolar phases, providing a programmable blueprint to assemble the multiphase nucleolus that contributes to proper rRNA flux and ribosome assembly.

Mapping rRNA processing in time and space

To measure rRNA processing in time and space, we developed a 5-ethynyl uridine (5eU) nucleotide analogue-based pulse-chase labelling approach to measure both the nucleolar RNA localization using super-resolution fluorescence microscopy and the processing state using RNA sequencing (RNA-seq; Fig. 1b). This approach requires modifying existing nascent RNA-seq strategies³² to remove the background signal arising from the abundance of mature rRNAs, which we found impairs the detection of nascent pre-rRNA processing (Extended Data Fig. 1a-d). In brief, we labelled nascently transcribed RNA for 15 min (pulse) with the 5eU nucleotide analogue and then chased with excess unlabelled uridine over time. As 5eU nucleotides can be conjugated with a dye using click chemistry (5eU-imaging)¹⁹, we can visualize the radial outflux of nascently transcribed rRNA from the FC-DFC interface. where RNA Poll is synthesizing rRNA, to the outer GC layer over time³³ (Fig. 1b). Alternatively, 5eU nucleotides can be conjugated with biotin, enabling the purification of these nascently transcribed RNAs with streptavidin beads followed by sequencing (5eU-seq)³² to measure both rRNA cleavage and modification (2'-O-methylation using RiboMeth-Seq³⁴) steps in one experiment, at the single-nucleotide resolution (Extended Data Fig. 1a,b,h,i). Importantly, 5eU incorporation does not substantially impair pre-rRNA synthesis, processing or modification (Extended Data Fig. 1e-g).

Using this approach, we can quantitatively measure both rRNA localization (Fig. 1c–e and Extended Data Fig. 2a,b) and processing (Fig. 1f–i) over time. We find that the timing of rRNA cleavage steps is correlated with rRNA movement through the nucleolar phases (Fig. 1i). Specifically, early cleavage steps (sites 01 and 02) begin immediately after transcription (around 0–15 min) (Fig. 1i), which coincides with SeU rRNA localization near the FC–DFC boundary at early chase time-points (Fig. 1c–e), consistent with these cleavage steps occurring co-transcriptionally or immediately after transcription^{35,36}. Next, we observed subsequent cleavage steps (sites 1 and 2) as rRNA fluxes from the DFC to GC (about 30–60 min), and late cleavages occur as RNA fluxes from the GC to nucleoplasm (around 60–120 min) (Fig. 1c–e,i). This spatiotemporal map suggests that nucleolar SSU-processing (18S rRNA precursors) steps may be largely completed before RNA enters the GC, while LSU-processing (5.8S/28S rRNA precursors) steps, which

take markedly longer, may occur throughout the nucleolus (Fig. 1f,g,i). This is consistent with research in yeast showing that the nuclear dwelling time of SSU precursors is far less than those of LSU^{37} . Moreover, the vast majority of 2'-O-methylations on rRNA occur within 30 min after transcription (defined as fast sites; Fig. 1h), consistent with findings that most 2'-O-methylations are placed co-transcriptionally in yeast³⁶. However, a handful of post-transcriptional 2'-O-methylations (for example, 28S Um4498, Gm4499) occur substantially later (around 60–90 min) (Fig. 1h and Extended Data Fig. 1h–j), corresponding to when 5eU rRNA moves into the GC phase.

To validate the 5eU-seq and imaging approach, we performed RNA SABER-FISH³⁸ (hereafter, RNA-FISH) using cleavage-spanning probes designed to no longer hybridize after cleavage (Fig. 1f, j). RNA-FISH shows early pre-rRNA species (site 01) at the FC-DFC boundary, consistent with previous work²⁵ (Fig. 1j,k and Extended Data Fig. 2c-e). Site 1 and 2 RNA-FISH signals are primarily localized within the DFC and at the DFC-GC boundary (Fig. 1j,k and Extended Data Fig. 2c-e), consistent with their cleavages occurring as 5eU rRNA moves from the DFC to GC (around 30-60 min) (Fig. 1i). Late LSU cleavage steps (such as ITS2), which occur (around 60-120 min) after 5eU RNA enters the GC (Fig. 1i), are indeed localized within the GC (Fig. 1j,k and Extended Data Fig. 2c-e), confirming that the late LSU-processing steps occur in the GC, while nucleolar SSU-processing steps are completed before RNA enters the GC (Fig. 1g,i). Using a kinetic model to deconvolve the contributions of individual rRNA intermediates, these FISH data were directly compared to our 5eU-seq and imaging data and appear comparable (Supplementary Note 1). Taken together, this map of where different rRNA processing steps occur demonstrates that the cleavage steps are spatially segregated in different phases of the nucleolus.

Processing controls flux and morphology

Our findings above show that rRNA processing is correlated with rRNA outflux through the nucleolus; however, it is unclear whether these processing steps are strictly required for rRNA movement through the nucleolus. To test this, we perturbed rRNA processing with chemical inhibitors or knockdowns (KDs) of processing factors and measured the impact on rRNA processing, flux and nucleolar morphology (Fig. 2a). First, we treated cells with flavopiridol (FVP)-a CDK9 (Pol II) inhibitor that is known to block rRNA processing but not Pol I transcription¹⁷. Using 5eU-seq to label nascently transcribed rRNA after 1 h of FVP treatment, we observed a major impairment of all but one rRNA cleavage step-only the earliest co-transcriptional 01 cleavage proceeds normally (Fig. 2b and Extended Data Fig. 3a, b, f). Notably, 2'-O-methylation levels also decreased substantially (Fig. 2d and Extended Data Fig. 3c-e), with stronger reductions on 28S than 18S (Extended Data Fig. 3g). Consistent with previous studies, FVP-treated nucleoli form a typical 'necklace' morphology¹⁷, whereby the outer GC phase appears detached from the inner phases (Fig. 2e and Extended Data Fig. 4a,c).

Given that FVP inhibits almost all of the processing steps, we next performed pulse-chase 5eU imaging analysis of cells pretreated with FVP for 1 h to determine whether unprocessed rRNA species can flux into the GC phase. While newly transcribed rRNAs move into the GC after a 60-min chase in the DMSO-control cells (Fig. 2c), unprocessed 5eU-labelled rRNAs in the FVP-treated cells accumulate around the GC surface, but do not partition into the GC (Fig. 2c and Extended Data Fig. 4d,e). RNA-FISH confirmed that only processed rRNA enters the GC: probes targeting early and middle cleavage sites appear excluded from the GC, while late probes localize inside the GC (Extended Data Fig. 4f). Supporting this, FVP-washout experiments demonstrated that restarting pre-rRNA processing led to 5eU-labelled RNA and processed rRNA species entering the GC, coincident with the reattachment of the FC and DFC with the GC phase, within about 30-60 min (Extended Data Fig. 4b,c,f-h and Supplementary Videos 1 and 2). This timing is consistent with when rRNA cleavage and flux into the GC occurs (around



Fig. 2 | **Impaired rRNA processing impacts rRNA flux and alters the nucleolar morphology. a**, Pre-rRNA processing was perturbed in MCF10A cells and analysed using 5eU-seq (processing) or 5eU imaging (flux and morphology). **b**, The normalized cleavage efficiency (treatment versus control; Methods) for 5' ETS (1) or 3' ETS (02) sites measured by 5eU-seq after treatment with 2 μ M FVP (general processing inhibitor, purple), KD of SSU-processing factors (U3 snoRNA or FBL, red) or LSU-processing factors (RPL5 or U8 snoRNA, blue). *P* values were calculated using two-tailed *t*-tests; ****P* = 0.0004, *****P* < 0.0001. *n* = 12 (FVP), 8 (U3), 15 (FBL), 8 (RPL5) and 6 (U8) measurements. ND, not determined; NS, not significant; estimation of cleavage efficiency and statistical testing was not possible for FVP due to complete cleavage inhibition (Extended Data Fig. 3f). **c**, Representative images of 5eU-labelled RNA (white) in nucleoli (mTagBFP2–NPM1) after a 60 min chase in control and treated cells (top). Bottom, 5eU–GC (NPM1) correlation from 0–120 min. *n* ≥ 50 cells per timepoint per condition. Example images are provided in Extended Data Figs. 4e and 6. **d**, The average 18S and 28S rRNA 2'-O-methylation levels (ScoreC) over O-90 min from 5eU-seq for controls (solid lines) and treatments (dashed). n = 101-208 measurements per timepoint. **e**, Representative images and schematics of GC (NPM1, blue), DFC (FBL, red) and FC (RPA194, green) (IF) for all conditions except FBL KD, where mTagBFP2-NPM1, NOP56-mCherry and RPA16-GFP are visualized. RPA194 puncta outside the nucleoli are immunostaining background, and were not observed in the fluorescence-tagged lines (Extended Data Fig. 5d, e). **f**, FC and DFC nucleolar rim enrichment in controls and perturbations (normalized to control). The violin plots show the median (solid line) and quartiles (dashed lines). Statistical analysis was performed using two-tailed Mann–Whitney *U*-tests; ****P < 0.0001. n = 1,116 (controls), 97 (FBL), 228 (RPL5), 153 (U3) and 105 (U8) cells. All scale bars, 3 µm. For **c** and **d**, data are mean ± s.e.m.

30–60 min) (Fig. 1c,i). Taken together, our results suggest that rRNA processing is required for the flux of rRNAs into the GC, and that this processing-gated movement of RNAs has a key role in shaping nucleolar morphology.

As FVP broadly affects rRNA processing and other cellular functions (Pol II inhibition)^{16,39}, we next tested more specific perturbations of rRNA processing. We used antisense oligonucleotides (ASOs) to deplete the box C/D snoRNA U3 (ref. 40), a key SSU processome component that is involved in 5' ETS cleavage^{40,41} (Extended Data Fig. 5a–c). U3 KD disrupted the two cleavages within the 5' ETS (01 and 1) in the SSU pathway (Fig. 2b and Extended Data Fig. 3h) while LSU processing remained unaffected, as expected⁴⁰ (Fig. 2b and Extended Data Fig. 3h). Given that rRNA processing rates were reduced, we next performed 5eU imaging and found that RNA flux into the GC was slowed as well (Fig. 2c). Together, these data show that 5eU–seq enables quantitative measurement of rRNA cleavage and modification rates at single-nucleotide resolution and, when coupled with 5eU imaging, reveals that pre-rRNA processing rates are tightly connected to rRNA movement through the nucleolus.

SSU defects cause 'inside-out' nucleoli

We next examined the consequences of processing defects on nucleolar morphology after U3 KD. Notably, U3 KD results in an 'inside-out' morphology whereby the FC and DFC phases move towards the nucleolar periphery, reversing the order of the nucleolar phases (Fig. 2e, f and Extended Data Fig. 5d, e). This phenotype begins within 8 h and fully develops within 24–48 h (Extended Data Fig. 5h–j); furthermore, the number of nucleoli reduces from about 2–3 per cell to 1 per cell after U3 KD (Extended Data Fig. 5g). Importantly, the inversion is distinct from the well-known nucleolar segregation phenotype^{17,22}, whereby FCs also move towards the periphery of the nucleolus, but fuse to form large caps due to transcriptional inhibition (Extended Data Fig. 5e, f). Instead, the number of FCs remains high after U3 snoRNA KD, consistent with active Pol I transcription (Extended Data Fig. 5e, f).

We next examined whether nucleolar inversion could be phenocopied by knocking down fibrillarin (FBL), another U3 snoRNP component (Extended Data Fig. 5k-l). 5eU-seq reveals that partial KD of FBL slows SSU cleavage and broadly reduces modifications (Fig. 2b,d and Extended Data Fig. 3j,k), consistent with its function as a 2'-O-methyltransferase and SSU processome component^{41,42}. Notably, the levels of 28S modifications change more substantially than 18S (Extended Data Fig. 3k), except for the late-forming Gm4499 modification, not placed by FBL⁴³, which is unaffected (Extended Data Fig. 5m). FBL KD also results in a partially inverted morphology, like that seen with U3 KD (Fig. 2e, f), underscoring the role of SSU processing in nucleolar morphology. Inversion is not a generic trait of processing defects, as depletion of RPL5 (uL18) or the box C/D snoRNA U8, which are primarily involved in LSU processing, results in distinct nucleolar morphologies and slowed rRNA flux (Fig. 2c,e,f and Extended Data Figs. 5n-p and 6). In addition to impaired LSU processing (Fig. 2b), these RPL5 and U8 KDs also impact certain sites in the SSU pathway (Extended Data Fig. 31,n), consistent with studies showing that LSU-processing defects can impair SSU maturation^{18,40}. Note that 2'-O-methylation is not broadly impacted after U3, U8 or RPL5 KD (Extended Data Fig. 3i, m-o). Together, these findings suggest that rRNA processing shapes both nucleolar morphology and rRNA flux.

Engineering synthetic nucleoli in cells

To better understand how pre-rRNA processing shapes nucleolar morphology, we engineered synthetic nucleoli in living cells using a plasmid system that was previously shown to produce mature ribosomal subunits^{44,45}. These rDNA plasmids contain unique sequences in 18S and 28S (18S* and 28S*), enabling visualization of plasmid rRNA using FISH^{44,45} (Fig. 3a and Extended Data Fig. 7a). Transient transfection led to the formation of a de novo nucleolus with characteristic FCs and ring-like DFCs embedded within a GC phase, recapitulating the size and morphology of native nucleoli (Fig. 3b,c and Extended Data Fig. 8a,b and 9b). Synthetic nucleoli were distinguished from native nucleoli using FISH for both plasmid-specific sequences (18S*/28S*) and endogenous rRNA sequences absent in the plasmid (Δ 1,2,3 segments of 5' ETS) (Fig. 3a–d and Extended Data Fig. 8c). Moreover, we observed plasmid-derived rRNAs in the cytoplasm (Fig. 3b,f,h), where they incorporate into monosomes and polysomes (Extended Data Fig. 8d), demonstrating that they are functional. Finally, despite not being tethered to chromosomes, synthetic nucleoli recruit heterochromatin, marked by DNA and H3K9me2/3, to their periphery–another hallmark of endogenous nucleoli^{46–48} (Fig. 3e and Extended Data Fig. 10a).

Notably, synthetic nucleoli sometimes fuse with endogenous nucleoli, generating hybrid nucleoli containing endogenous and plasmid-derived rRNA (Fig. 3b–d). In these hybrids, 18S* rRNA localizes in discrete territories while 28S* rRNA spreads throughout the GC (Extended Data Fig. 8e). This is consistent with our observations that LSU processing continues into the GC and takes much longer than SSU processing (Fig. 1i).

LSU precursors build the GC phase

Given the spatiotemporal differences in SSU and LSU processing observed in these synthetic nucleoli (Extended Data Fig. 8e) and 5eU-seq (Fig. 1i), we used this engineerable system to test whether SSU or LSU precursors independently shape the nucleolar phases. To do this, we generated truncated plasmids expressing only SSU or LSU pre-rRNAs (Fig. 3f and Extended Data Fig. 7b). Notably, plasmids expressing only SSU precursors form ring-like DFC structures like those in native nucleoli, but lack a surrounding GC (Fig. 3f,g). These synthetic structures recruit FC and DFC components (such as the Pol I subunit RPA194 and the SSU processome component NOP56) (Figs. 3f,g and 4f), but lack GC components, such as the well-established GC marker NPM1 and the two LSU-processing factors SURF6 and RRP1 (Fig. 3f,g and Extended Data Fig. 8f). This is consistent with previous findings showing a partial 18S rRNA sequence, which failed to recruit NPM1²⁹. Conversely, plasmids encoding only a large-subunit precursor (5.8S and 28S pre-rRNAs) generate a synthetic structure containing both DFC and GC components (Fig. 3f.g), suggesting that LSU rRNA is necessary to build the GC phase. Nonetheless, the ring-like DFC morphology within LSU-only nucleoli was less pronounced, indicating that SSU pre-rRNAs have a key role in shaping the DFC.

Notably, 18S rRNA expressed from the SSU-only plasmid is exported into the cytoplasm (Fig. 3f,h) and incorporated into monosomes and polysomes (Extended Data Fig. 8d), further suggesting that the GC may be dispensable for SSU processing (Supplementary Note 3). Conversely, a signal for 28S* rRNA from the LSU-only plasmid was not observed in the cytoplasm, even when co-transfected with an SSU-only plasmid, suggesting that LSU assembly is compromised in the absence of upstream SSU pre-rRNA elements (Fig. 3f,h, Extended Data Fig. 8g,h and Supplementary Note 4). Taken together, these findings demonstrate that LSU precursors, but not SSU precursors, actively build the GC phase, consistent with LSU precursors undergoing cleavage in the GC.

SSU processing drives nucleolar layering

Having shown that plasmid-expressed pre-rRNAs recruit assembly factors generating each nucleolar subphase, we next used this system to directly examine how rRNA processing shapes nucleolar morphology. Indeed, although altered nucleolar morphology is often linked to rRNA processing defects^{18,49,50}, it remains unclear whether this arises directly through defective processing or secondary effects such as cellular stress. This engineered plasmid system enables us to directly



Fig. 3 | **Engineered synthetic nucleoli in cells recapitulate normal multiphase architecture with the expression of the LSU precursors required for GC recruitment. a**, Schematics of endogenous rDNA and a plasmid expressing an rDNA sequence containing three deletions (Δ1, Δ2 and Δ3) in the 5' ETS region and unique insertions in 18S and 28S (18S* and 28S*). **b**, RNA-FISH detection of endogenous pre-rRNA (endo 5' ETS probes; yellow) and plasmid-derived pre-rRNA (18S* probe, white) in HEK293T cells. The dashed lines show individual nucleoli; the solid lines indicate individual nuclei. **c**, Representative images of nucleoli classified as de novo (plasmid derived), endogenous and hybrid (fusion), labelled for DFC (FBL IF) and GC (NPM1 IF). The FISH probe control is shown in Extended Data Fig. 8c. **d**, RNA-FISH intensities of 18S* and endogenous 5' ETS probes across nucleolar classes. a.u., arbitrary units. **e**, Heterochromatin (H3K9me2/3) and chromatin (DAPI) staining around endogenous and de novo nucleoli (GC: RRP1 IF, NPM1 IF). **f**, Schematics of SSU-only and LSU-only plasmids with truncated rDNA sequences (Extended Data Fig. 7a, b). Left, representative

test the effect of rRNA processing on nucleolar architecture without broad changes to cellular function.

As KD of U3 snoRNP components (that is, U3 snoRNA or FBL) results in inverted nucleoli (Fig. 2e), we predicted that designing sequence-specific changes in the plasmid rDNA to block U3-mediated

images of de novo nucleoli formed from rDNA, SSU-only and LSU-only plasmids, visualized by RNA-FISH (28S* for rDNA and LSU only, 18S* for SSU only) and nucleolar markers DFC (NOP56–mCherry), GC (mTagBFP2–NPM1). Right, the corresponding cytoplasmic FISH signals. **g**, Quantification of the mean nucleolar DFC (NOP56–mCherry) and GC (mTagBFP2–NPM1) intensity from **f**. *P = 0.0248, **P = 0.0012, ****P < 0.0001. n = 24 (wild type, WT), 14 (SSU only) and 17 (LSU only) nucleoli. **h**, Quantification of cytoplasmic RNA-FISH intensities for rDNA, SSU-only and LSU-only plasmids from **f**. **P = 0.0016, ****P < 0.0001. n = 32 (WT; 18S* rRNA), 47 (WT; 28S* rRNA), 90 (SSU-only) and 7 (LSU-only) cells. Scale bars, 3 µm (**b** and **f** (right; whole cells)) and 1 µm (others). The box plots show the median (centre lines), 25th–75th percentiles (box limits) and minimum–maximum values (whiskers). HEK293T cells were used for all of the experiments. Statistical analysis was performed using two-tailed Mann–Whitney *U*-tests.

processing should result in a similar morphological inversion. We designed plasmids containing U3 snoRNA in addition to the rDNA locus (Fig. 4a). Using the human SSU processome structure as a template⁴⁴, we mutated two sites (5' and 3' hinges) within the 5' ETS rRNA spacer that directly base-pair with U3 snoRNA⁵¹⁻⁵³ (Fig. 4a,b and Extended Data



Fig. 4 | **U3-mediated SSU processing drives the layering of the multiphase nucleolus and rRNA outflux. a**, Plasmids co-expressing U3 snoRNA and rDNA with various mutations (red) (Extended Data Fig. 7c–g). **b**, Synthetic nucleoli from normal, 3' and 3'5' hinge mutants or base-pairing-rescued plasmids, visualized by GC (mTagBFP2–NPM1), DFC (NOP56–mCherry), identified by a lack of endogenous rRNA and the presence of 18S* rRNA. Right, cytoplasmic 18S* and 28S* rRNA signals. Scale bars, 1 μ m (left), 3 μ m (right). The solid lines indicate nuclei; the arrows indicate a lack of cytoplasmic 18S* signal. See Extended Data Fig. 9a,b for morphologies reproduced with IF and all phases visualized. **c**, Quantification of DFC rim enrichment. *****P* < 0.0001, ***P* = 0.0024. *n* = 95, 12, 19, 9, 18 and 13 nucleoli. **d**, The Pearson correlation between 18S* rRNA and NPM1. **P* = 0.0221, ***P* = 0.0012. *n* = 7, 9, 6, 15 and 8 nucleoli. **e**, Quantification of the cytoplasmic 18S* signal from **b**. *****P* < 0.0001. *n* = 32, 38, 58, 79 and 102 cells. **f**, Morphology (left) and FC rim enrichment score (right) of de novo nucleoli (FC, RPA194 IF; DFC, NOP56-mCherry) from WT (n = 15) or 3'5' hinge mutant (n = 29) SSU-only plasmids (Extended Data Fig. 7b). ****P < 0.0001. **g**, Localization of proteins of interest (POI, green: FBL (IF), EXOSC10 (IF) and RPS6-Halotag) relative to DFCs (NOP56-mCherry) of WT SSU-only nucleoli (left). Right, the radial distribution of 18S* RNA and POIs around the DFC boundary (dashed line, 0 µm). n = 74 (Nop56, 18S), 87 (FBL), 21 (RPS6) and 12 (EXOSC10) nucleoli. **h**, The 18S* RNA distribution around DFCs (NOP56) in WT (n = 74) versus mutant (n = 164) SSU-only nucleoli. **i**, The cytoplasmic 18S* rRNA levels from WT (n = 90) and mutant SSU (n = 34) plasmids. Scale bar, 3 µm. ****P < 0.0001. Statistical analysis was performed using two-tailed Mann-Whitney *U*-tests. For **g** and **h**, data are mean ± s.e.m. Scale bars, 1 µm (**f**-**h**). The box plots in **c**-**f** and **i** show the median (centre lines), 25th-75th percentiles (box limits) and minimum-maximum values (whiskers). HEK293T cells were used for all experiments.



Fig. 5 | **An RNA-dependent multiphase model of nucleolar architecture. a**, The proposed model of how rRNA transcription and processing shape the multiphase nucleolus. A 13.3 kb pre-rRNA is transcribed from rDNA, processed and cleaved to assemble the SSU and LSU. Here we show that the LSU precursors are necessary for assembling the nucleolar GC phase (blue) and SSU processing drives the ordering of the DFC (red) and GC (blue) phases. Different arrangements of multiphase structures (such as normal or inversion) can arise from changes in interfacial tensions across multiple interfaces: nucleoplasm (NP)–DFC ($\gamma_{NP,DFC}$), NP–GC ($\gamma_{NP,GC}$) and GC–DFC ($\gamma_{GC,DFC}$). Under normal U3-mediated cleavage of the 5′ ETS from SSU pre-rRNA, the DFC localizes inside the GC. After impaired U3-mediated SSU processing, SSU pre-rRNAs build up in the DFC phase and no longer flux into the GC phase. This results in a change in the interfacial tensions and the nucleolar morphology inverts, whereby the GC is now enveloped by the DFC. **b**, Different nucleolar morphologies are recapitulated in a phase-field model that considers the partitioning of different rRNA precursors (for example, SSU before and after 5' ETS cleavage) into the different nucleolar phases (DFC and GC). For simplicity, the FC and DFC are modelled as one nucleolar phase. Changes in U3-mediated processing, RNA Pol Itranscription or LSU production (SSU-only) alter the concentrations of rRNA precursors in each phase, resulting in different nucleolar morphologies. **c**, Modelling of impaired SSU processing (5' ETS cleavage) over time leads to an accumulation of SSU precursors (before 5' ETS cleavage) and inversion of the nucleolar phases. **d**, Modelling of RNA Pol I transcriptional inhibition over time; a decreased concentration (normalized to the concentration at *t* = 0) of all SSU and LSU rRNA precursors results in the nucleolar cap morphology.

Fig. 7c–g). These constructs formed synthetic nucleoli that recapitulate the inversion phenotype observed after U3 KD, with FCs and DFCs localizing at the GC periphery (Figs. 2e and 4b,c and Extended Data Fig. 9a,b). Inversion is observed with multiple DFC markers (Extended Data Fig. 9a,c), including FBL and NOP56 (components of box C/D snoRNPs), NOPP140 (important for snoRNP biogenesis and function), UTP23, ESF1 and KRR1 (SSU assembly factors)^{44,54}, as well as GC marker NPM1 and LSU assembly factors RRP1 and SURF6 (ref. 45). Notably, the co-existence of a normal endogenous nucleolus in the same cell as an inverted synthetic nucleolus demonstrated that inversion is a direct result of impaired SSU processing rather than altered cellular function (Extended Data Fig. 9d).

Inspired by previous research in yeast⁵¹, we next produced U3 variants to rescue base-pairing with the mutant rRNA (Fig. 4b and Extended Data Fig. 7c–g). These compensatory mutations largely restored the order of nucleolar phases (Fig. 4b,c) and rescued cytoplasmic export of 18S rRNA (Fig. 4b,e), indicating that both the nucleolar structure and processing are largely rescued. As a control, 28S rRNA expressed from these plasmids was successfully exported to the cytoplasm in all conditions (Fig. 4b and Extended Data Fig. 9e). Notably, inverted synthetic nucleoli are no longer associated with peripheral heterochromatin (Extended Data Fig. 10a), which is also reproduced in endogenous nucleoli after U3KD (Extended Data Fig. 10b). This suggests that FC and DFC localization at the edge of the nucleolus impairs heterochromatin tethering, consistent with studies suggesting that nucleolar heterochromatin association occurs through the GC^{46-48} .

As loss of U3 snoRNA function inverted the nucleolar phases (FC– DFC–GC becomes GC–DFC–FC; Fig. 2e), we next used the SSU-only plasmid, which generates synthetic nucleoli lacking a GC (Fig. 3f), to dissect the role of rRNA processing in the reorganization of the FC and DFC subphases (Fig. 4f). While the wild-type SSU-only plasmid produced the classical arrangement of an FC embedded inside a ring-like DFC, a mutant SSU-only plasmid deficient in U3 base-pairing instead resulted in an FC on the surface of the DFC (Fig. 4f). This mirrors the peripheral FC localization observed in synthetic nucleoli produced from rDNA plasmids lacking U3 binding (Extended Data Fig. 9b). These results demonstrate that base-pairing between U3 snoRNA and the 5' ETS, which is essential for SSU-mediated processing, is required for the ordering of the FC and DFC phases, even in the absence of a GC.

Pre-rRNA processing gates rRNA outflux

Given that pre-rRNA processing is tightly coupled with its movement through the nucleolus (Fig. 2), we next used synthetic nucleoli to study whether impaired U3 processing alters the outflux of 18S pre-rRNA from the DFC. In contrast to wild-type synthetic nucleoli, in which 18S rRNA precursors are localized throughout the nucleolus, U3-binding mutant 18S precursors remain localized in the DFC, and do not move into the GC (Fig. 4b,d; see the schematics of RNA distribution in grey). Compensatory mutations in U3 snoRNA restore RNA flux into the GC (Fig. 4b,d,e), indicating that U3-mediated processing is required for 18S pre-rRNA outflux.

We next used the SSU-only plasmid system (Fig. 4f), which provides a unique opportunity to study the outflux of SSU precursors without the complicating presence of the GC. 18S rRNA precursors and factors demarcating various SSU-processing stages are radially organized around the DFC (Fig. 4g and Extended Data Fig. 10c,d). Early processing factors (NOP56, FBL, KRI1 and KRR1) localized closest within the DFC. 18S pre-rRNA and ribosomal proteins RPS4X and RPS6 extend beyond the DFC, consistent with ribosomal proteins remaining part of maturing ribosomal particles as they are exported. Further out, the RNA exosome component EXOSC10 is localized around the periphery of the DFC. This positioning of the RNA exosome, marking the degradation of 5' ETS and release of DFC assembly factors⁴⁴, further underscores that processing and removal of dozens of assembly factors could enable SSU precursors to be released from the DFC.

To test whether processing directly enables outflux from the DFC, we used the mutant SSU-only plasmids defective in U3-mediated processing and found that 18S precursors, ribosomal protein RPS6 and the late processing factor EXOSC10 are no longer localized at the DFC periphery (Fig. 4h and Extended Data Fig. 10e.f), indicating that successful SSU processing is required for outflux. Indeed, mutant SSU-only nucleoli show accumulation of 18S pre-rRNA and early processing factors (ESF1, NAT10, NOP56 and FBL) within enlarged DFCs (Extended Data Fig. 10g-i), consistent with the retention of unprocessed intermediates in the DFC phase. This lack of outflux of mutant 18S rRNA is also correlated with loss of cytoplasmic 18S rRNA export (Fig. 4i). Together, in both mutant SSU-only and mutant rDNA nucleoli, 18S rRNA precursors appear trapped within the DFC phase and do not flux outward into the GC or nucleoplasm (Fig. 4b.d.h-i). These results suggest that the DFC phase, composed of multivalent interactions between early processing factors and pre-rRNA, can serve as a checkpoint to prevent the outflux of unprocessed SSU intermediates.

Modelling RNA-dependent nucleolar form and function

Our findings indicate that irreversible steps in pre-rRNA processing like rRNA cleavage directly shape the morphology of the nucleolus. Specifically, we hypothesize that rRNA processing modulates the biophysical properties of its liquid-like phases. As a multiphase condensate, the organization of the nucleolar layers is thought to be governed by interfacial tensions (y_{ij}) between phases (i,j), which depend on local concentrations of rRNA species and associated proteins^{13,55-57} (Fig. 5a). To examine this physical picture, we adapted a phase-field model from material science using a Flory-Huggins framework^{1,7,58,59}. The model comprises three phases-DFC, GC and nucleoplasm (for simplicity, FC and DFC were modelled as a single component, DFC)-and pre-rRNA. Pre-rRNA production, processing, degradation and flux were incorporated based on the kinetic model presented earlier (Methods and Supplementary Notes 1 and 2). A three-body interaction term between rRNA and two of the nucleolar components was essential to capture the effect of rRNA on interfacial tensions⁶⁰.

This minimal model recapitulates the inversion morphology observed after inhibition of 5' ETS cleavage, which results in a significant increase in unprocessed species in the DFC and a lack of mature 18S rRNA in the GC (Fig. 5a–c and Supplementary Note 2). This altered rRNA composition in each phase changes their interfacial tensions, resulting in a relocalization and inversion of the GC and DFC phases (Fig. 5c and Supplementary Video 3). Moreover, our model can also explain two other marked nucleolar morphologies: classical nucleolar caps in response to reduced concentrations of all rRNA intermediates, mimicking Pol I inhibition, or the SSU-only phenotype lacking a GC observed after selective reduction of LSU intermediates (Fig. 5b, d and Supplementary Video 4). While this model is a major simplification of the full complexity of pre-rRNA processing, it nonetheless clearly demonstrates how changes in rRNA processing can reshape nucleolar morphology by altering interfacial tensions (Fig. 5).

Discussion

The nucleolus has been known for almost two centuries, but the precise origin of its multiphase architecture and its relationship to the critical function of ribosome biogenesis has been veiled by a lack of tools to probe this structure–function relationship. Here we have developed and used a set of tools to map and dissect nucleolar organization and function, identifying the essential components required to generate and topologically program the nucleolus. By combining spatiotemporal mapping of pre-rRNA processing, engineering synthetic nucleoli in human cells and mathematical modelling of the determinants of multiphase nucleolar architecture, we bridge the gap between molecular-scale functions of the nucleolus and its structure at the

micrometre scale. Indeed, the approaches developed here provide single-nucleotide resolution of cleavage and modification rates and allow for the parallel measurement of processing in space and time, revealing how the sequential maturation of rRNA is required for its outward flux. The ability to precisely engineer a synthetic nucleolus based on nucleotide-level changes within a recombinant rDNA locus highlights that both rRNA transcription and accurate ribosome assembly blueprint the architecture of the multiphase nucleolus.

A key finding is that, although nucleolar rRNA is generated by the transcription of a single 13.3 kb pre-rRNA transcript, its cleavage, chemical modifications and overall maturation results in SSU and LSU precursors that differentially contribute to the nucleolar structure: SSU processing controls the order of the phases, whereas the LSU is required for GC formation. Moreover, faithful pre-rRNA processing is required for the flux of pre-ribosomal particles through the successive phases. Specifically, pre-ribosomal particles that cannot undergo key maturation events remain trapped within the DFC phase, suggesting that the nucleolar phases could ensure the fidelity of ribosome assembly by preventing the release of immature precursors into the next compartment. Our study therefore underscores the role of nucleolar multiphase organization in functional staging, whereby the physical separation of nucleolar phases may serve as distinct processing or 'retention' compartments segregating the different reactions driving the sequential maturation of SSU and LSU particles.

Together, we used a set of tools to identify how pre-rRNAs build and arrange the nucleolar phases that facilitate their processing, revealing how defects in ribosome assembly can manifest in substantial changes in the topology of the nucleolus. Our work therefore not only underscores basic principles of how RNA and its processing gives rise to the multiphase architecture of the nucleolus, but also provides a foundational toolkit for dissecting how ribosome biogenesis and nucleolar morphology are dysregulated in disease.

Online content

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41586-025-09207-4.

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Methods

Statistics and reproducibility

All statistical tests were performed using GraphPad Prism 10, using either two-tailed Mann–Whitney *U*-tests or two-tailed *t*-tests, as specified in the legends. Tests for normality were performed when appropriate. Sample sizes were not predetermined. For all experiments, cells were randomized such that within a biological or technical replicate all cells were analysed equally with no sub-sampling. Microscopy imaging, western blotting and quantitative PCR with reverse transcription (RT– qPCR) were repeated independently at least three times with similar results. *P* values and the numbers of observations (*n*) are provided in the figure legends.

Cell culture and cell lines used in this study

All cells were cultured in a humidified chamber at 37 °C with 5% CO₂ and 1% streptomycin and penicillin (Gibco, 15140122). HEK293T cells (ATCC), HEK293T cells tagged with UBTF-sfGFP, NPM1-mtagRFP and FBL-Halotag (gift from the Leonetti laboratory), HCT116 cells (gift from Y. Kang) and MCF7 cells (gift from Y. Kang) were cultured in DMEM (GIBCO, 11995065) supplied with 10% FBS (R&D Systems, S11150H). MCF10A (gift from Y. Kang) cells were cultured in DMEM/ F12 medium (Thermo Fisher Scientific, 11320082) supplied with 5% horse serum (Sigma-Aldrich, H1138), 20 ng ml⁻¹ EGF (Novoprotein, C029), 10 ng ml⁻¹insulin (Sigma-Aldrich, 91077 C) and 1 µg ml⁻¹hydrocortisone (Sigma-Aldrich, H0888). All cells were checked for Mycoplasma and authenticated. For imaging, cells were treated with trypsin (trypsin-EDTA 0.05%, Thermo Fisher Scientific, 25300054) for dissociation and then seeded into 96-well glass-bottom dishes (Cellvis, P96-1.5H-N) coated with bovine fibronectin (Sigma-Aldrich, F1141) diluted 1:4 in 1× DPBS (Thermo Fisher Scientific, 14190144). All rDNA plasmid transfection experiments were performed in HEK293T cells, and all other experiments were performed in MCF10A cells unless otherwise noted.

Recombinant human rDNA plasmid designs

A plasmid containing a previously described minimized 5' ETS (mini 5' ETS) that is compatible with human SSU biogenesis was used as a starting point (pSK M323) for the design of plasmids described in this Article⁴⁴ (Extended Data Fig. 7a). The structure of the human SSU processome in state pre-A1 (Protein Data Bank: 7MO8) was used to redesign the 3' and 5' hinge RNA duplexes between U3 snoRNA and the 5' ETS (Extended Data Fig. 7c,d). Nucleotide substitutions were introduced that maintain the overall nucleotide composition of the duplexes while only allowing matching variants to base pair (Extended Data Fig. 7e, f). Variants of 3' and 5' hinges of the 5' ETS and U3 snoRNA were combined (pSK M432-pSK M435) by including variants of the complete human U3 gene upstream of the RNA polymerase I promoter, resulting in a bidirectional promoter for pre-rRNA (RNA polymerase I) and U3 snoRNA (RNA polymerase II) (Extended Data Fig. 7g). Plasmids coding for either wild-type or mutant SSU pre-rRNAs were generated by terminating transcripts after the first 48 nucleotides of the 5.8S gene and a plasmid coding for the LSU pre-rRNAs contained the first 53 nucleotides of the mini 5' ETS followed by ITS1, 5.8S, ITS2, 28S and the 3' ETS (Extended Data Fig. 7b). A probe for ITS2 was introduced from a previously published plasmid (pSK_M349) that was shown to give rise to mature human LSUs⁴⁵.

Plasmid construction

FM5-Nop56-mcherry was a gift from D. W. Sanders. FM5-mTagBFP2-NPM1, FM5-RPA16-GFP and FM5-RPS6-Halotag lentiviral DNA plasmids were generated using the FM5 lentiviral vector (gift from D. W. Sanders)⁵⁷. A DNA fragment encoding human RPS6 was amplified from original plasmids (DNASU Plasmid Repository, HsCD00043827) by PCR using the Q5 High-Fidelity 2X Master Mix (NEB) with oligonucleotides synthesized by IDT. gBlocks encoding RPA16 or RPS4X proteins were ordered from IDT. DNA fragments containing *NPM1* and *RPA16* were PCR amplified from gifted plasmids from J. A. Riback. The In-Fusion HD cloning kit (Takara) was used to insert the fragments into the desired linearized vector featuring a GS-linker–fluorescent tag fusion. All of the constructs were confirmed by Sanger sequencing (GENEWIZ).

IF analysis

Cells were fixed in 96 well glass bottom plates with 4% PFA for 10 min, washed with 1× PBS twice and then permeabilized with 1× PBST (with 0.5% Triton X-100) for 15 min. The samples were then blocked in 2% BSA in PBS for 30 min and then incubated with primary antibodies in 2% RNase-free BSA (VWR, 97061-420) for 2 h at 37 °C (a detailed antibody list is provided in Supplementary Table 1). Three 1× PBS washes were conducted for 5 min each. For non-conjugated antibodies, anti-mouse or anti-rabbit secondary antibodies with the desired fluorophores were used at a 1:1,000 dilution for 2 h at 37 °C. Three 1× PBS washes were conducted with 5 min each before imaging.

RNA-FISH

SABER FISH was performed as previously described^{38,61}. Probes were designed across the ribosomal RNA sequence and additional hairpin sequences are appended at the 3' end of the probe for primer exchange reaction concatemerization with hairpin_28 (paired with 488 fluorescent oligos for imaging) or hairpin 25 (paired with 647 fluorescent oligos for imaging). A list of all of the probe sequences is provided in Supplementary Table 1. Note that, because early cleavage sites are cleaved and degraded rapidly, we used FISH probes tiling 5' ETS (upstream of 01 site) and 3' ETS (downstream of 02 site) to improve detection. For probing ribosomal RNAs transcribed from synthetic plasmids, 18S* and 28S* probes were designed (Supplementary Table 1) to probe the unique (around 16-20 nucleotides) insertion within the 18S and 28S sequence from plasmids⁴⁴. Moreover, an antisense 18S* probe was designed to ensure the RNA-FISH signal specifically comes from RNA instead of DNA (Extended Data Fig. 8c and Supplementary Table 1). To specifically probe endogenous ribosomal RNA, we used probes hybridized to parts of the 5' ETS region that are excluded from the synthetic rDNA plasmids $(\Delta 1, 2, 3)^{44}$. Moreover, as a unique sequence is inserted within 28S (28S*), we designed a FISH probe flanking the insertion, located upstream and downstream of 28S*, for selective hybridization to endogenous 28S rRNA (endogenous 28S in Supplementary Table 1). For analysing whether plasmid-derived RNAs are exported to the cytoplasm, we used the endogenous 28S probe for segmenting the nuclei and cytoplasm (details are provided in the the 'Quantitative image analysis' section). IF analysis was conducted after completing all of the FISH steps, starting from the blocking step as described in the 'IF analysis' section. Murine RNase inhibitors (NEB, M0314L) were used at a 1:200 dilution in all of the steps of IF to preserve the RNA-FISH signals.

Microscopy

A Nikon CSU-W1 SoRa spinning-disc confocal microscope equipped with a Yokogawa SoRa pixel reassignment-based super-resolution device was used for rapid super-resolution imaging. The system was built around the Nikon Ti2-E fully motorized microscope and is equipped with dual Hamamatsu Fusion BT sCMOS cameras. The W1 Sora system is equipped with 405, 488, 514, 532, 561, 594 and 640 nm laser lines. For this work, a Nikon CFI Plan Apo Lambda D ×60 oil (MRD71670) lens was used with a ×2.8 SoRa magnification and 405, 488, 561 and 640 nm lasers. A Mad City Labs piezo z stage was used for z-stack acquisition. All acquisitions were performed using the Fusion BT in Ultra Quiet readout using correlated double sampling. In some cases, Denoise.ai (Nikon software) was used for images shown and analysed in this study.

Lentiviral packaging and transduction

Lentiviruses were made using HEK293T cells seeded in a six-well plate at 70–80% confluence. The desired plasmids were transfected together with helper plasmids VSVG and PSP using Lipofectamine 3000 (Invitrogen, L3000008) according to previously described protocols^{57,62}. Viruses were collected 48 h after transfection and filtered through syringe filters with a 0.45-µm pore size (VWR). Lentiviral transduction was conducted in 96-well plates at 30% cell confluency for 2 days and cells were then expanded to make stable expression lines and sorted using fluorescence-activated cell sorting for polyclonal lines, with a tight window for each fluorescent protein intensity.

Transient transfection

rDNA plasmids (including engineered mutant plasmids) were transfected into HEK293T cells (for one well of a 24-well plate, 600 ng plasmids were transfected into 150,000 HEK293T cells) using lipofectamine 3000 (Invitrogen). Unless otherwise stated, for all plasmid transfection experiments, cells were seeded into 96-well glass-bottom plates 24 h after transfection and fixed for SABER FISH 48 h after transfection. Specifically, all cytoplasmic export analyses were performed at 48 h after transfection (Figs. 3f-h and 4e,i and Extended Data Fig. 9e). This ensured that all plasmids were compared at the same time after transfection. For the wild-type and mutant SSU-only plasmid transfection experiments comparing nucleolar morphology and radial outflux in Fig. 4f-h and Extended Data Fig. 10c-h, we optimized the timing of transfection to be 24 h to achieve the optimal number of de novo SSU nucleoli.

$\label{eq:starses} Endogenous\,N\mbox{-}terminal\,tagging\,of\,NPM1\,with\,mTagBFP2\,using\,CRISPR\mbox{-}Cas9$

Endogenous N-terminal tagging of NPM1 in MCF10A cells was performed as previously described^{63,64}. An oligonucleotide pair encoding an NPM1-targeting gRNA (TGTCCATCGAATCTTCCAT) was cloned into a modified lentiCRISPRv2-puro plasmid (from A. Lin) through the BsmBI restriction site. MCF10A cells were transfected using the FuGENE HD transfection reagent (Promega, E2311) with plasmids containing the cloned gRNA and a donor plasmid. The donor plasmid was constructed by cloning the tag with a flexible linker flanked by 300 bp homology arms complementary to the N terminus of the *NPM1* gene into the pUC19 vector (Thermo Fisher Scientific, SD0061). Then, 3 days after transfection, mTagBFP2-positive cells were single-cell sorted into 96-well plates. These single-cell clones were then cultured and expanded for tagging validation through western blotting and junction PCR of the specific genomic locus (Extended Data Fig. 2f,g) and imaging to confirm correct subcellular localization.

All ribosomal RNA processing perturbations used in this study

CX-5461 Pol I inhibition. Cells were treated with 10μ M final CX-5461 (MedChem Express, HY-13323) for 90 min before fixation and imaging. DMSO was used for the control group.

FVP broad rRNA processing inhibition. Cells were treated with FVP at a final concentration of 2 μ M (MedChem Express; HY-10005). As a control, cells were treated with DMSO. For the 5eU imaging and 5eU-seq, cells were pretreated for 1 h to broadly inhibit processing prior to a 15 min pulse and subsequent 0–90 min chase. All pulse–chase medium contained 2 μ M FVP to ensure that processing inhibition was maintained throughout the 5eU pulse–chase labelling. For imaging the reformation of the multiphase nucleolus after FVP washout, cells were treated with 2 μ M FVP for 2 h and then washed twice quickly with 1× DPBS before imaging in regular medium. Videos were taken at 37 °C with 5% CO₂, every 2 min after FVP removal for 90 min. For 5eU imaging after FVP washout, cells were treated with 2 μ M FVP for 2 h and then washed twice quickly with 1× DPBS before imaging in regular medium. Videos were taken at 37 °C with 5% CO₂, every 2 min after FVP removal for 90 min. For 5eU imaging after FVP washout, cells were treated with 2 μ M FVP for 2 h and then washed twice quickly with 1× DPBS before a 15 min 5eU pulse followed by a 0–90 min chase.

RPL5 shRNA. For shRNA vector cloning into a lentiviral plasmid for expressing shRNAs with puromycin selection. RPL5 shRNA (GATGA TAGTTCGTGTGACAAA) sequences and a negative control (GCTCT TAACTAACGTCACCTA) sequence were separately cloned into pLKO.1 TRC vector after digestion with Agel and EcoRI. Lentiviruses were produced as described above and added to cells for 5eU-seq and imaging experiments with MCF10A cells at 30% confluency. After 1 day, virus-containing medium was removed and replaced with fresh medium including 5 µg ml⁻¹ of puromycin. After selection for another 3-4 days, cells in the negative control well that were not treated with virus were dying as expected, indicating that selection was effective. Selection was terminated by replacing the puromycin medium with normal MCF10A medium. Cells were then split for 5eU sequencing, imaging or RNA extraction followed by RT-aPCR (a list of the primer sequences used is provided in Supplementary Table 1) 4-5 days after adding shRNA viruses.

U3/U8 snoRNA ASO. U3 and U8 snoRNA ASO treatment was performed as previously described⁴⁰ with several adaptations. For 5eU-seq experiments, in each well of a 6-well plate, 1.5 µl of 40 µM (stock) ASO diluted in 125 µl Opti-MEM (Thermo Fisher Scientific, 31985062) was combined with 7.5 µl Lipofectamine RNAiMAX (Thermo Fisher Scientific, 13778075) diluted in 125 µl Opti-MEM. After a 30 min incubation at room temperature, 1.75 ml of a suspension containing 250,000 cells in antibiotic-free medium (see the 'Cell culture and cell lines used in this study' section) was added to each well. Cells were incubated for 1.5 days before analysis using 5eU-seq (Fig. 2b,d and Extended Data Fig. 3h,i,n,o). For imaging experiments, in each well of a 96-well plate coated with bovine fibronectin (Sigma-Aldrich, F1141) diluted 1:4 in 1× DPBS (Thermo Fisher Scientific, 14190144), 0.05 μ l of 40 μ M ASO diluted in 4.165 µl Opti-MEM was combined with 4.165 µl Lipofectamine RNAiMAX diluted in 0.25 µl Opti-MEM. After a 30 min incubation at room temperature, 100 µl of a suspension containing 8,000 cells in antibiotic-free medium was added to each well. For 5eU-imaging experiments, cells were incubated for 2.5 days before fixation (Fig. 2c and Extended Data Fig. 6e). For experiments monitoring the inversion morphology, cells were treated again with ASO after 2.5 days according to the protocol described above, then incubated for an additional 2 days before fixation (Fig. 2e, f). We also tested the morphology change at shorter times (at 8, 12, 24 and 48 h), and observed that FCs and DFCs start moving towards the edge of the nucleolus within 8-12 h, with the phenotype fully developing within 24-48 h (Extended Data Fig. 5h-j). All incubations were performed at 37 °C and 5% CO₂. ASO sequences were published in a previous paper⁴⁰ and are listed in Supplementary Table 1.

FBL siRNA. *FBL* siRNA treatment was performed as described in the U3 and U8 snoRNA ASO treatment protocol with the following modifications. For 5eU-seq experiments and western blotting, 2 µl of 20 µM *FBL* siRNA (Supplementary Table 1) or control siRNA (Thermo Fisher Scientific, 4390843) diluted in 250 µl Opti-MEM was combined with 6 µl Lipofectamine RNAiMAX diluted in 250 µl Opti-MEM. For imaging experiments, 0.066 µl of 20 µM *FBL* siRNA or control siRNA diluted in 8.33 µl Opti-MEM was combined with 8.33 µl Lipofectamine RNAiMAX diluted in 0.2 µl Opti-MEM. For all of the experiments, cells were treated again after 24 h with *FBL* siRNA or control siRNA. Cells were incubated for an additional 3 days before performing 5eU-seq, collection for western blotting or fixation for 5eU imaging.

Total RNA isolation

For each well of a six-well plate, total RNA was collected in $200 \ \mu l 1 \times$ Buffer RLT (QIAGEN, 79216) and isolated using the QIAGEN RNeasy Mini Kit (74104), followed by 1 h of DNase treatment using TURBO DNase (Thermo Fisher Scientific, AM2238). DNase-digested RNA was then further purified using the Zymo RNA Clean and Concentrator-25 kit (R1017).

RNA electrophoresis

The RNA integrity after isolation was analysed and the ratios of 28S to 18S rRNA in U3 and U8 snoRNA ASO- and SCR ASO-treated samples were assayed using the Agilent RNA High Sensitivity Assay (Agilent, 5067-5579) on the 4150 TapeStation system (Agilent Technologies) according to the manufacturer's instructions.

RT-qPCR for validation of perturbation

U3 snoRNA, U8 snoRNA and *RPL5* mRNA KD efficiency were assayed using RT–qPCR using the Luna Universal One-Step RT-qPCR Kit (NEB, E3005) according to the manufacturer's instructions, except that each reaction was scaled to $60 \,\mu$ l to allow for four technical replicates (12.5 μ l) per sample. U6 snoRNA served as the loading control. RT–qPCR was performed on the Applied Biosystems QuantStudio 3 Real-Time PCR System instrument (A28567). Primer sequences used are listed in Supplementary Table 1. All primers were synthesized by Integrated DNA Technologies (IDT). U3, U8 and U6 snoRNA absolute amounts were determined using standard curves prepared for each primer set. U3 and U8 KD efficiency was assayed by comparing the absolute amounts of U3 or U8 snoRNA normalized to the absolute amounts of U6 snoRNA, in U3 or U8 snoRNA ASO- versus SCR ASO-treated samples.

Polysome fractionation and analysis

HEK293T cells after 48 h of transfection with the wild-type rDNA plasmid, SSU-only rDNA plasmid or no transfection control were treated with 100 µg ml⁻¹ cycloheximide (CHX) for 10 min at 37 °C with 5% CO₂ and polysome fractionation was performed as described previously with minor modifications⁶⁵. Cells were then lysed in 400 µl polysome lysis buffer (25 mM HEPES, pH 7.3, 150 mM NaCl, 15 mM MgCl₂, 1% Triton X-100, 8% glycerol, 0.5% sodium deoxycholate, 100 µg ml⁻¹CHX, 1 mM DTT, RNase inhibitors (NEB, M0314L, 1:60 dilution) and DNase (Thermo Fisher Scientific, AM2239, 1:400)) by incubating on ice for 15 min, followed by two consecutive centrifugations at 800g for 5 min, one centrifugation at 8,000g for 5 min and one centrifugation at 20,000g for 5 min (all at 4 °C) to remove nuclei and mitochondria. A 50 µl portion of the resulting supernatant was set aside as the input, while the remainder was loaded onto a 10-50% (w/v) sucrose gradient prepared in polysome gradient buffer (25 mM HEPES, pH 7.3, 150 mM NaCl, 15 mM MgCl₂, 100 µg ml⁻¹CHX and 1 mM DTT). Ultracentrifugation was performed at 40.000 rpm for 2.5 h at 4 °C in a SW41 Ti rotor (Beckman Coulter). After ultracentrifugation, gradients were fractionated using a piston gradient fractionator (Biocomp) with continuous monitoring of absorbance at 254 nm to visualize ribosomal profiles. The fractions corresponding to 'monosomes (80S)' and 'polysomes (>2 ribosomes)' were collected and pooled separately. RNA was extracted from each fraction using ethanol precipitation followed by proteinase K (NEB, P8107S) digestion for 1 h at 50 °C, and cleaned using the Zymo RNA Clean and Concentrator-5 kit (R1016). Then, 25 ng of isolated RNA from input, monosome or polysome fractions was put into the RT-qPCR reaction. RT-qPCR was performed as described in the previous section using primers for 18S* (plasmid rRNA) and 18S (endogenous rRNA) (Supplementary Table 1). For each fraction (monosome or polysome), we calculated the incorporation of plasmid-derived rRNA across all ribosomes by normalizing the plasmid rRNA (18S*) abundance to the total rRNA (18S) abundance. Standard curves were generated to normalize for primer efficiencies.

5eU imaging

The SeU imaging protocol was modified from previous studies³³. In brief, cells were seeded 1 day before at 40% confluency in 96-well glass-bottom plates. The volumes of all reagents were kept at 100 μ l for each well of the 96-well plate. SeU (Thermo Fisher Scientific, E10345) solution was prepared at 0.5 mM in cell culture medium and added to cells with medium removed from the well. Cells were then kept in an incubator (37 °C with 5% CO₂) for 15 min (pulse). Next, SeU

containing medium was removed and quickly washed twice with 1× DPBS containing 10 mM (excess) uridine (Sigma-Aldrich, U6381-5G) to outcompete the leftover 5eU. The culture medium containing 10 mM uridine was then added to the well before incubating for different chase timepoints (0 to 120 min) at 37 °C with 5% CO₂. Note that all solutions mentioned above were kept at 37 °C using a heat block to minimize temperature-induced effects on RNA transcription and processing. For wells with different chase times on the same 96-well plate, the starting time of the 5eU pulse was staggered so that all the chase timepoints ended at the same time. For fixation, 4% paraformaldehyde (PFA) diluted from 16% PFA (Thermo Fisher Scientific, PI28906) with 1× PBS was used for 15 min at room temperature. Fixed cells were then washed twice with 1× PBS and permeabilized with 1× PBST (1× PBS and 0.5% Triton X-100) for 15 min. For click chemistry, the Click-iT Plus Alexa Fluor 647 Picolyl Azide Toolkit (Thermo Fisher Scientific, C10643) was used according to the manufacturer's instructions, with the exception of using AZDye 647 Picolyl Azide (Click Chemistry Tools, CCT-1300-1) instead of the azide from the kit. After 30 min of applying the click chemistry reaction cocktail, cells were washed once with 1× PBS before imaging. For combining IF with 5eU imaging, IF steps were performed as described above before click chemistry.

5eU-seq

All 5eU pulse-chase labelling experiments combined with sequencing were performed in MCF10A cells except for method validation described in Extended Data Fig. 1c-g, which was performed on HEK293T cells. Cells were seeded for 5eU pulse-chase labelling such that their confluency was about 50-60% at the time of collection. Cells were pulse labelled with 5eU (Jena Biosciences, CLK-N002-10) for 15 min at 37 °C with 5% CO₂. Cells were then removed from the incubator and quickly washed twice with 1× DPBS to remove 5eU from the cells. Cells were then chased with medium containing 10 mM uridine (Sigma-Aldrich, U6381) to outcompete the leftover 5eU in cells over different chase timepoints. All solutions were kept at 37 °C using a heat block to minimize temperature-induced changes to the cells, which could impact RNA transcription and processing. After the given chase time, cells were collected in 1× buffer RLT (Qiagen, 79216) and frozen at -80 °C for RNA isolation. RNA isolation was performed using the Qiagen RNeasy Mini kit followed by DNase digestion (TURBO DNase, Invitrogen, AM2238) to remove genomic DNA. RNA concentrations were measured using the Oubit RNA BR (Thermo Fisher Scientific, O10211) system and the RNA integrity was analysed on the Agilent RNA High Sensitivity (HS) Tapestation.

Total RNA (10-15 µg) isolated after 5eU pulse-chase was click-reacted with biotin picolyl azide (Click Chemistry Tools, 1167-25) as described $previously^{32} with the following modifications. Capture of biotinylated\\$ RNA was performed using 20 µl Dynabeads MyOne Streptavidin C1 beads (Invitrogen, 65002) after click chemistry. Captures and washes were performed as previously described except for a modification to the three 5 min washes of captured material, which was changed to 75 °C in no-salt urea buffer (4 M urea, 10 mM HEPES, pH 7.5, 10 mM EDTA, 0.5% Triton X-100, 0.2% SDS, 0.1% Na-DOC). We found that three rounds of sequential captures (as described previously³²) as well an optimized protocol introduced here performing washes at high temperature and in a buffer lacking salt were essential to reduce the background of highly abundant mature rRNA (Extended Data Fig. 1a-d). RNA-seq library preparation was performed as previously described⁶⁶. 5eU-seq libraries were then sequenced on the NovaSeq 6000 (Illumina) system with paired-end reads (either 150 × 150 or 100 × 200). The samples that failed quality control, such as failed click reactions, streptavidin captures or library amplifications, were excluded from downstream analyses.

High-performance computing

The analyses presented in this Article were performed on computational resources managed and supported by Princeton Research

Computing, a consortium of groups including the Princeton Institute for Computational Science and Engineering (PICSciE).

Computational analysis of 5eU–seq cleavage and 2'-O-methylation

Sequencing read analysis pipeline. A custom snakemake pipeline was used to perform alignments, 2'-O-methylation analysis and cleavage analysis (https://github.com/SoftLivingMatter/5eU-seq-pipelines). Sequencing reads were trimmed with Trimmomatic (v.0.39)⁶⁷ to remove adaptor sequences and bases containing low quality scores. Trimmed reads were then aligned to the rDNA genome (GenBank: U13369.1) using STAR (v.2.7.11a)⁶⁸. Reads were sorted and indexed using Samtools $(v.1.9-4)^{69}$ and only uniquely mapped reads were kept for further analysis. The fraction of reads that are cleaved at a given site was calculated as the number of non-spanning reads divided by the total number of reads (spanning and non-spanning) at a given site (Extended Data Fig. 1b) using featureCounts (v.1.6.4)⁷⁰ (Subread package). rRNA cleavage sites were annotated based on the positions described previously¹⁴. All rRNA cleavage sites used in this study were manually inspected to define the sites of cleavage based on where the 5' or 3' end of reads piled up, demarcating a precise cleavage site. Specifically, well-defined cleavage sites correspond to those that undergo endonuclease cleavage and can be mapped at a base-pair resolution using 5eU-seq (examples shown in Extended Data Fig. 1b). In cases in which endonucleolytic rRNA cleavage events are followed by gradual degradation, a broader window (several nucleotides) downstream of an annotated site was used (Supplementary Table 1).

Cleavage efficiency measurement. Cleavage efficiency, defined as 1 over the time to reach 50% cleavage (t_{50}), was estimated by fitting the observed 5eU–seq fraction cleaved over time (Extended Data Fig. 3) to a sigmoidal 4PL curve using Prism 10 software. We then computed the normalized cleavage efficiency (Fig. 2b) by dividing each by its respective control sample. Two-tailed *t*-tests were performed to estimate the significance of each treatment cleavage efficiency relative to its respective control. We note that, because FVP treatment completely blocks almost all rRNA cleavage steps, it is not possible to accurately estimate the time to reach 50% cleavage, so statistics were not determined.

2-O-methylation analysis. To determine 18S and 28S ribosomal RNA 2'-O-methylation levels over time, we applied RiboMethSeq computational analysis methods^{34,71} to 5eU-seq sequencing reads. In brief, 2'-O-methylated nucleotides are substantially more resistant to alkaline hydrolysis compared to non 2'-O-methylated nucleotides, resulting in fewer RNA fragmentation events at these sites. To map the nucleotides where fragmentation occurred, sequencing adaptors are ligated to the 3' ends of fragmented RNA and 3' ends of cDNA. This enables the identification of 2'-O-methylation sites at the single-nucleotide resolution by performing end mapping of reads. Specifically, the 5' ends of read 1 sequences, which correspond to the 3' end of the fragmented RNA, are mapped to the reference rRNA sequence. Next, modification levels at a given position were estimated by calculating the number of 5'-end read counts at a given 2'-O-methylated site compared with that of their neighbouring nucleotides, defined as ScoreC (see below). Nucleotides that are 2'-O-methylated, due to their resistance to hydrolysis, have substantially fewer 5' end read counts, and therefore appear as 'dips' or areas of low coverage in the end-mapping profile (Extended Data Fig.1h,i).

We calculated ScoreC at known 2'-O-methylation sites⁴² using a weighted average of the 5' end read counts in a ± 2 nucleotide window, recommended previously⁷², around each site. Nucleotides in the ± 1 and ± 2 neighbouring positions were assigned weight contributions of 0.9 and 1, respectively. If a separate 2'-O-methylation site was found within the ± 2 window around a site, the 5' end read counts at the former were skipped and those at the immediately preceding nucleotide were

used alternatively. For the calculation of ScoreC, nucleotide positions of 2'-O-methylation sites on 18S rRNA were converted to those on 47S rRNA (U13369.1 human ribosomal DNA) by adding +3,655 to the 18S 2'-O-methylation positions (except for Am1678, Cm1703 and Um1804, to which +3,657 was added). For sites on 28S rRNA, nucleotides were converted to 28S 2'-O-methylation positions on 47S rRNA by adding +7,924, +7,920, +7,912, +7,911 and +7,903 for Am398 to Am400, Gm1316 to Gm2876, Cm3701 to Gm3944, Gm4042 to Cm4054, and Gm4196 to Gm4637, respectively. For data visualization, negative ScoreC values were clipped to zero. The heat map in Extended Data Fig. 3d was generated in Prism using the heat map function.

Defining fast and slow 2'-O-methylation sites. We measured the rates at which 2'-O-methylation occurs across all known 18S and 28S sites in our 5eU-seq data performed on MCF10A cells (n = 2 replicates) to determine which occur at a fast versus slow rate. 18S and 28S rRNA 2'-O-methylation sites that do not attain near steady state 2'-O-methylation levels within 30 min from transcription are defined as slow. All other 2'-O-methylation sites that meet this criterion and are defined as fast. Sites with high variation between replicates that made it challenging to confidently infer 2'-O-methylation rates were manually excluded. A table of high-confidence fast and slow 18S and 28S rRNA 2'-O-methylation sites is provided in Supplementary Table 1. We have provided examples of 5eU-seq data for individual high-confidence fast (Gm4494) and slow sites (18S-Cm1272, 28S-Cm3808, 28S-Um4498) in Extended Data Fig. 1i, j and Fig. 1h. The average of all high-confidence fast sites for each of 18S and 28S is shown in the SSU and LSU panels of Fig. 1h, respectively.

RiboMethSeq

RiboMethSeq for data in Extended Data Fig. 1g was conducted exactly as described previously³⁴. For analysis, adapter sequences were trimmed from raw reads using Trimmomatic v0.39 with the following parameters: LEADING:30, TRAILING:30, SLIDINGWINDOW:4:15, AVG-QUAL:30 and MINLEN:17. Quality control of the raw and trimmed reads was assessed using FastQC (v.0.11.9). Alignment to the reference rRNA sequence (18S: NR_003286.4; 5.8S: NR_003285.2; 28S: NR_003287.4) was done using Bowtie2 (v.2.3.5.1) with the default parameters. Sorting, indexing and extraction of mapped reads was done using Samtools (v.1.15.1), with the option -F 4 to exclude the unmapped reads. Subsequent analysis was conducted using R: the quantification of ribosomal RNA (rRNA) 2'-O-methylated residues was performed using the RNAmodR.RiboMethSeq package (v.1.18.0), the final processed data were exported using the writexl (v.1.5.0) package and display output was generated using Prism (v.10.3.0).

Pre-rRNA processing analysis by northern blotting

Total RNA was extracted from HEK293 cells labelled with or without 5eU with TRI reagent solution (AM9738, Thermo Fisher Scientific), according to the manufacturer's instructions. Then, 5 µg total RNA was resolved on a 1.2% agarose/6% PFA denaturing gel, transferred overnight by capillarity onto a nylon Hybond N+ membrane (RPN203B, Cytiva), and hybridized with radioactively labelled probes (Supplementary Table 1). The probes were designed to detect all major pre-rRNA intermediates. The signal was acquired with a phosphorimager (FLA-7000, Fujifilm) and quantified using native multi-gauge software (v.3.1, Fujifilm).

Quantitative image analysis

All quantitative imaging measurements were performed using CellProfiler⁷³ v.4.2.6. Cell segmentation was performed using cellpose⁷⁴ v.2.3.2 and the RunCellpose plugin for cellprofiler. Full computational methods, pipelines and notebooks describing the calculation of the derived methods can be found at the GitHub repository (https://github.com/ SoftLivingMatter/image-analysis-quinodoz-jiang-2024). To deploy cellprofiler plugins on HPC systems, a snakemake workflow⁷⁵ wrapper was developed. It controls the resources and behaviour of each invocation and can be found at GitHub (https://github.com/softLivingMatter/ snakemake-cellprofiler/).

Morphology. To quantitatively examine changes in cell morphology as a function of nucleolar perturbations, the following steps were performed in CellProfiler. First, the multichannel images were split into separate channels, such as GC, FC, DFC, FISH/SeU/other markers. The GC channel was used to define initial GC objects using minimum cross-entropy thresholding with a diameter range of 30-400 pixels, where each pixel corresponds to $0.0387 \,\mu$ m. Clumped objects were not separated as it produced too many falsely separated GC objects. As the images do not have a nuclear or cytoplasmic stain and some metrics are best measured across an entire cell, we computationally assigned nucleoli to the same cell based on if they were within 100 px to each other, producing a merged object mask. The mask was dilated by 50 px to measure metrics within the prospective nucleoplasm and find nucleolar features that may be outside the GC, such as FCs after FVP treatment, which causes detachment of the phases.

To find FC objects, the dilated GC objects were used to mask the FC image before performing 'enhancing speckles' with a feature size of 20 px using the Fast setting in the EnhanceOrSuppressFeatures module. The FC objects were found in the enhanced FC image using an adaptive, three-class Otsu threshold with the middle intensity class assigned to the background. This was selected to ensure that the identified FC objects were primarily in plane and in focus. The adaptive window was 100 px, clumped objects were separated by intensity and shape, and FC objects were selected with diameters of 7–50 px. As these FCs included objects within and outside of the GC, each FC was assigned to either a nucleolar or extra-nucleolar class using an overlap threshold of 50%. For example, an FC that overlaps an initial GC object by more than 50% was considered a nucleolar FC.

DFC objects were defined by masking the DFC image with the dilated GC objects. As the DFC phase organization is highly variable between treatments, the built-in IdentifyPrimaryObjects module was not able to perform adequately in all cases. Instead, DFCs were found by first thresholding with an adaptive, three-class otsu cut-off with a 150 px window. The middle class was assigned to background to ignore the diffuse DFC phase that develops after certain perturbations. The threshold image was converted to objects and then prospective DFCs were filtered to ensure they had an area of at least 20 px.

The GC, DFC and FC objects were combined to a single mask which was used as the support for measuring Pearson's correlation and overlap of the image channels. The size and shape of each object set was also measured and used for scaling some metrics, discussed below. The intensity of the FC channel was measured in FC objects, and GC and probe (for example, FISH probe or 5eU) were measured in the GC and DFC phases. The distribution of FC and DFC intensity was measured over the initial GC objects using 20 scaled bins. A binary image of the initial GC objects was also measured to facilitate combining bins during post-processing. Each object was related to its corresponding dilated GC object before exporting to a csv for further analysis in Python.

The raw data produced by cellprofiler was further processed using jupyter notebooks and custom analysis scripts. Each object csv was read into a pandas dataframe and merged into a final result by the image and dilated GC unique identifier. Values such as total area or FC count were summed to provide a per-cell measurement. Average intensities per cell were calculated by first multiplying the mean intensity from cellprofiler with the object area, summing the result, then dividing by the total area per cell. Rim enrichment was calculated by summing the radial distribution fraction for the bins of interest and dividing by the fraction of the GC object over the same range of bins. Investigation of a range of rim widths found that the outer 20% of the GC rim provided the best discrimination between U3 KD and control cells. Circularity was estimated as $4 \times \pi \times area/perimeter^2$.

RDF estimation. To facilitate broader utilization of Radial distribution function (RDF) measurements, we developed a cellprofiler plugin, MeasureRDF, for calculating the radial distributions of object sets, such as the intensities of FISH or 5eU signals within a 1 µm radius from the FC centre, on a given input image. A separate plugin allows for further flexibility in performing upstream object finding and filtering and simplifies integration into other analysis tasks. In contrast to the MeasureObjectIntensityDistribution method available in cellprofiler, the MeasureRDF plugin uses a fixed distance in pixels and attempts to resolve overlapping objects as described below.

In the point-based measurement mode, the MeasureRDF plugin operates on a set of objects as the point sources along with a containing object. Here we used the GC boundary as the masking objects and FCs as the point sources. For each GC, the set of FCs are considered together. The image intensities are mean-centred with unit variance and pixel distances from each FC are determined. The RDF distribution is estimated by minimizing the difference between the scaled image intensity and a superposition of each FC point source with the same RDF. This allows for deconvolution of neighbouring point sources while providing an averaged estimate of the RDF. The estimated intensities are rescaled to the original intensity units before reporting to the user. In all of the plots, the intensity is either min–max scaled between the minimum and maximum intensities in the RDF or max scaled by dividing by the maximum intensities in the RDF to highlight intensity dynamics.

To visualize the location of each phase from the FC centre, the RDF distributions of FC, DFC and GC can be plotted on a colour bar, where the intensity of a phase is mapped to a given colour (FC, green; DFC, red; GC, blue). To identify the boundaries between the distinct phases, the intensities of each phase were subtracted from the intensity of their adjacent phase before plotting. Specifically, the distance at which the intensity of pairs of phases is equal is used to estimate the boundary between phases. Finally, the 5eU RDF distributions were used to map the distance the 5eU RNA has travelled from the FC centre to its corresponding chase time. For each chase time, the location of maximum EU intensity (5eU peak) was determined and used in a piecewise-linear interpolation between time and distance.

The MeasureRDF plugin can also operate in a boundary-measurement mode, which was used for the SSU-only and nucleolar periphery analyses performed in Fig. 4 and Extended Data Fig. 10a–f. In this setting, the source object boundaries are considered at r = 0, defined as the radial position of 50% DFC or GC normalized intensity, and the image intensity is measured as a function of distance from the boundary. The distance of each pixel to the object boundary is determined and used to estimate the RDF curve. In cases in which neighbouring point sources overlap, pixel intensities are assigned to the closest object.

Engineered plasmid measurements. For the images of de novo nucleoli, we manually classified individual nucleoli as endogenous, de novo or hybrid on the basis of their intensities for plasmid-expressed 18S*/28S* RNA and endogenous 5' ETS rRNA. Specifically, endogenous nucleoli were those with high endogenous 5' ETS RNA-FISH and no detectable plasmid-expressed 18S*/28S* RNA-FISH signal. Conversely, de novo nucleoli were those with high 18S*/28S* plasmid RNA-FISH signal, and no detectable endogenous 5' ETS signal. Hybrid nucleoli contained FISH signals for both channels. Figure 3d demonstrates the range of nucleolar intensities of 5' ETS and plasmid-expressed FISH intensities observed before further manual classification, where only high-confidence de novo versus endogenous nucleoli were analysed and any ambiguous nucleoli were excluded.

As the SSU-only rDNA plasmid produced nucleoli without detectable NPM1 intensity, a separate workflow was required to evaluate a subset of the morphology measurements described above. First, each channel was background corrected by subtracting the bottom 5 percentile value of each image. Next, the DFC (NOP56 or FBL) channel was blurred with a 5-px sigma Gaussian filter, which was found to produce better

segmentation of SSU-only nucleoli. The blurred DFC image was used to detect nucleoli with a two-class global Otsu threshold and diameters between 10 and 150 px without declumping. The initial nucleoli objects were dilated by 10 px to act as support for the rim-based RDF measurement. Each channel's intensity was measured in the nucleoli and the 10 -px rim as well as the object size and shape. For quality control, we manually checked the segmented SSU-only nucleoli objects and excluded those that were out of focus or incorrectly segmented.

To measure cytoplasmic and nuclear intensities of the 18S* or 28S* RNA expressed from engineered plasmids, nuclei were segmented from the endogenous 28S FISH using cellpose with the nuclei model (inverted mask) and an expected object diameter of 300 px. The entire cell extents were then segmented from the endogenous 28S FISH stain (non-inverted mask) using the cyto2 cellpose model and an object diameter of 500 px. The whole-cell and nucleus objects were masked to ensure that each cell had a nucleus and vice versa. Then, the cell objects were masked with the nucleus to define the cytoplasm. The background subtracted endogenous 28S and engineered 18S* or 28S* RNA intensities were measured in the nuclei and cytoplasm.

Assumptions used for nucleolar phase-field modelling

As multiphase condensates, the organization of the nucleolar phases should be strongly impacted by the relative interfacial tensions ($\gamma_{i,j}$) between each of the phases (i,j), which is a measure of the energy per unit area associated with interfaces^{13,55–57,76}. The relative interfacial tensions are expected to depend on the local concentrations of rRNA species and associated proteins (Fig. 5a), flux and biochemical nature of the underlying molecular species⁷⁷.

Specifically, we modelled the affinities and localization of different species to be consistent with our experimental observations: nascently transcribed pre-rRNA in the FC/DFC boundary, 18S rRNA precursors before 5' ETS cleavage in the DFC, and 18S rRNA and 28S rRNA precursors in the GC (Fig. 5a). Importantly, these rRNAs are not alone and form hundreds of interactions with proteins. Indeed, owing to its interactions with many assembly proteins and RNAs, the SSU processome represents a high-valency particle, of which the maturation, including the cleavage and degradation of 5' ETS, releases 50 assembly factors to generate a pre-40S particle^{44,78-80}.

A standard pairwise interaction term in the Flory–Huggins model encodes the affinity of rRNA to each nucleolar component and therefore the rRNA partitioning. Specifically, given our findings that U3-processing-deficient 18S precursors appear trapped within the DFC (Fig. 4b,d,h), our model assumes that these species have a high affinity for DFC, and a low affinity for the GC. Conversely, because 18S pre-rRNAs with normal processing flux into GC or nucleoplasm (Fig. 4b,d,h,i), we assume that 18S pre-rRNAs after 5' ETS cleavage have a low affinity for the DFC and a higher affinity for the GC. Additional details are provided in Supplementary Note 2.

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

All data supporting the findings of this study are available in the Article and its Supplementary Information. All genomics data are available at the Gene Expression Omnibus (GEO) under accession numbers GSE296080 and GSE296162. Source data are provided with this paper.

Code availability

All codes and pipelines used for this study are provided at GitHub and Zenodo: https://github.com/SoftLivingMatter/image-analysis-quinodoz-jiang-2024 (https://zenodo.org/records/14908468)⁸¹;

https://github.com/SoftLivingMatter/snakemake-cellprofiler (https:// zenodo.org/records/14908470)⁸²; and https://github.com/SoftLiving Matter/SeU-seq-pipelines (https://zenodo.org/records/14908466)⁸³. Example images and analysis notebooks are provided at Zenodo (https://doi.org/10.5281/zenodo.14910509)⁸⁴.

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Additional information

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Extended Data Fig. 1| 5eU-seq method description and validation. a, Schematics of the 5eU-sequencing protocol with 3 rounds of sequential captures. b, The "fraction cleaved" metric is calculated by measuring the number of "cleaved" reads (ending at a cut site) divided by the total number of reads (uncleaved (spanning a cut site) + cleaved). Bottom, examples of 5eU-seq data demonstrating single nucleotide resolution mapping of rRNA cleavage at specific sites (1 and 02) over time in MCF10A cells. c, Optimization of wash temperature and buffer conditions to reduce background from mature rRNA. d, The fraction of reads cleaved at 02 and 1 sites comparing Protocols 1, 2, and 3 on 5eU pulse-labelled material (15 min pulse, 0 min chase) from HEK293T cells. Protocol 3 is the optimized protocol used for all datasets in this paper. Total RNA is a reference for mature rRNA. n = 1 (Protocol 2 and 3), 2 (Protocol 1, total RNA) replicates. e, Northern blot analysis of 5eU pulse-labelled material (30 min) and unlabelled material as a control (no pulse) from HEK293T cells. Two replicates were performed. **f**, Northern blot probes used in e. **g**, RiboMethScore of 2'-O-Methylation for 5eU 30 min pulse labelled material (+5eU, 60 or 90 min chase) and unlabelled material as a control (-5eU). Modification levels are not significantly (n.s.) different across n = 108 2-O-Me sites between conditions (two-tailed t-test). **h**, Schematic (adapted from⁸⁵) explaining the detection of 2'-O-methylation levels by mapping 5' ends of reads. Drops in read end counts are observed at modified sites; see methods. **i**, Example 5' end read coverage at fast (Gm4494, 28S) and slow (Um4498, Gm4499, 28S) 2'-O-methylation sites over 0-240 min chase timepoints. **j**, Quantification of 2'-O-methylation levels (ScoreC) over 0-240 min chase timepoints at 28S Gm4494, Um4498, Gm4499 2'-O-methylation sites shown in i. n = 2 per timepoint. Error bars are s.e.m.



Extended Data Fig. 2 | **RNA FISH and 5eU-imaging of rRNA flux as well as characterization of endogenously tagged mTagBFP2-NPM1 cells. a**, Peak of 5eU signal (distance from the centre of FCs) over chase time quantified in Fig. 1e. **b**, Max normalized 5eU intensity over distance from the centre of FCs over time, quantified in Fig. 1e. **c**, Max normalized FISH intensity over distance from FC centre, quantified from images in e. **d**, Pearson correlation coefficient between all FISH probes and GC from images in e. Number of nucleoli (n) = 72 (5' ETS), 95 (3' ETS), 24 (Site 01), 111 (Site 1), 230 (Site 2), 38 (Site 3'), 105 (Site 4'),

318 (ITS2-28S), 72 (18S), 310 (28S). Violin plots are centred by median and quartiles are shown. **e**, Example images of RNA FISH probes in Fig. 1j, with FC (RPA194 IF), DFC (FBL IF), and GC (mTagBFP2-NPM1) shown. Scale bar = 3 μ m. Bottom: averaged FISH images around individual FCs. Scale bar = 1 μ m. **f**, Junction PCR of a 400 bp genomic region spanning the inserted mTagBFP2 in MCF10A -/- (parental) and +/- (one copy of NPM1 tagged) cells. For gel source data, see Supplementary Fig. 1b. **g**, Western blot for NPM1 in MCF10A -/- and +/- cells with β -actin as loading control. For gel source data, see Supplementary Fig. 1a.



Extended Data Fig. 3 | Example 5eU-seq data and altered pre-rRNA cleavage and modification measured by 5eU-seq upon all perturbations. a, 5eU-seq reads over 47S pre-rRNA for 15 min pulse labelled material over 0-90 min chase timepoints in DMSO-treated (black) and FVP-treated (red) MCF10A cells. FVPtreated cells were pretreated with 2 µM FVP for 1 hr. prior to 5eU pulse-chase and throughout the time course. Arrows indicate where the sequencing reads are changing over time upon cleavage. b, Zoom-in examples of 5eU-seq reads at 01,1,2/NA, 3'/ITS2-28S and 02/3' ETS regions in a. Dashed lines demarcate the cleavage sites and arrows indicate where the sequencing reads are changing over time upon cleavage. c, Zoom-in on 5' end read counts in DMSO-control or FVP-treated conditions over 0-90 min chase timepoints in 28S rRNA at Gm1522 and Am1524 2'-O-methylation sites. FVP causes impaired 2'-O-methylation, observed by loss of characteristic drops at 2'-O-Me sites. d, Heatmap of 2'-O-Me levels (ScoreC) at all 18S and 28S rRNA sites in control (DMSO) and FVP-treated

conditions over 0-90 min chase timepoints. **e**, Quantification of 2'-O-Me levels (ScoreC) over 0-90 min chase timepoints at the 28S Gm1522 and 28S Am1524 sites (plotted in c) upon DMSO and FVP treatment. n (replicates) = 2 per time point. **f-n**, Quantification of the fraction of RNA cleaved at each site displayed in b for MCF10A cells upon perturbations to rRNA processing (red): FVP-treatment or knockdown of U3 snoRNA (U3 ASO), Fibrillarin (FBL siRNA), RPL5 (RPL5 shRNA), and U8 snoRNA (U8 ASO) compared to their respective controls (black; DMSO for FVP, or scramble/negative control for ASO, siRNA, and shRNA treatments). n = 1 (U8, RPL5) or 2 (FVP, U3, Fib) per time point. **g-o**, Average 2'-O-Me levels (ScoreC) on 18S (left) and 28S (right) rRNA in perturbations (red) and control conditions (black) over 0-90 min chase timepoints. n = 1 (U8, RPL5) or 2 (FVP, U3, Fib) per time point. n = 1 (U8, RPL5) or 2 (FVP, U3, Fib) per time point. n = 1 (U8, RPL5) or 2 (FVP, U3, Fib) per time point. n = 1 (U8, RPL5) or 2 (FVP, U3, Fib) per time point. n = 1 (U8, RPL5) or 2 (FVP, U3, Fib) per time point. n = 1 (U8, RPL5) or 2 (FVP, U3, Fib) per time point. n = 1 (U8, RPL5) or 2 (FVP, U3, Fib) per time point. All error bars are s.e.m. All data collected from MCF10A cells.



Extended Data Fig. 4 | Altered localization of RNA species and nucleolar morphology upon FVP treatment. a, Nucleolar morphology (IF for RPA194, FBL, and NPM1 for FC, DFC and GC) in DMSO (-FVP) and after 30-90 min of 2 µM FVP treatment. b, Nucleolar morphology–FC (RPA16-GFP), DFC (NOP56-mCherry), and GC (mTagBFP2-NPM1)–after 0-90 min of FVP washout. Dashed lines demarcate nuclei; arrows indicate examples of GC reattachment to FC/DFC. c, Pearson correlation between DFC and GC signals from a and b. Left: Schematics illustrate GC detachment. Detachment: n = 303, 383, 327, 87 cells (0-120 min); Reattachment: n = 464, 291, 490, 182, 230, 182 cells (0-90 min). d-e, Schematic of 5eU pulse-chase scheme and example images of 5eU labelled RNA upon FVP perturbations. d, Representative images of 5eU-labelled RNA (30 min pulse, 60 min chase) and GC (mTagBFP2-NPM1) upon DMSO or 2 µM FVP treatment. Arrows demarcate 5eU labelled rRNA at the periphery of GC. e, Example images of 5eU RNA (15 min pulse, 0-90 min chase) and GC (mTagBFP2-NPM1) for correlation analysis in Fig. 2c (FVP-treated samples). **f**, RNA FISH upon 2 μ M FVP treatment for 0 (-FVP), 60 (+FVP), and 120 min (+FVP), or 60 min FVP washout; GC (mTagBFP2-NPM1). Right: Pearson correlation between GC and FISH probes (5' ETS: n = 91, 209, 320, 366; Site 1: n = 183, 58, 247, 291; Site 2: n = 127, 69, 272, 263; Site 4': n = 139, 156, 86, 265; 28S rRNA: n = 138, 285, 196, 851 cells). **g**, 5eU pulse-chase scheme (15 min pulse, 0-90 min chase) and flux upon continuous FVP treatment or FVP washout. Bottom: FC (RPA16-GFP), DFC (NOP56-mCherry), and GC (mTagBFP2-NPM1), and 5eU RNA. **h**, Pearson correlation between 5eU and GC from g (Continuous: n = 221, 344, 358, 230, 273, 392 cells (0-90 min); Washout: n = 464, 291, 490, 182, 230, 182 cells (0-90 min)). Error bars are s.e.m. All scale bars = 3 μ m. Violin plots are centred by median and quartiles are shown. MCF10A cells used except for d (HEK293T).





Extended Data Fig. 5 | Validation of all knockdowns performed in this study and comparison of nucleolar morphology between U3 snoRNA KD and Pol I inhibition. a, Quantification of U3 snoRNA FISH intensity in nucleoli from scramble (SCR, n = 479) and U3 ASO-treated (n = 329) cells. **** p-value < 0.0001. b, RT-qPCR analysis for U3 snoRNA levels 72 hrs. post-treatment with SCR or U3 ASO (n = 3 biological replicates per condition). *** p-value = 0.0006. c, 18S/28S rRNA ratio (RNA electrophoresis) in total RNA isolated 72 hrs. after SCR, U3, or U8 ASO treatment (n = 3 biological replicates per condition). * p-value = 0.0241, **** p-value < 0.0001. d, Nucleolar morphology in HCT116, HEK293T, and MCF7 cells following SCR or U3 ASO treatment. e, Nucleolar morphology following U3 ASO or CX-5461 (Pol I inhibition) treatment. Markers: GC (mTagBFP2-NPM1), DFC (NOP56-mCherry), and FC (RPA16-GFP) for d and e. f, Number of FCs per nucleolus in U3 ASO (n = 30), SCR ASO (n = 30), CX-5461 (n = 35), and control (n = 31) nucleoli. *** p-value = 0.0002, **** p-value < 0.0001. g, Number of GCs per cell in SCR (n = 384) and U3 ASO (n = 177) conditions. **** p-value < 0.0001. h, Time $course of nucleolar reorganization following 8-48 \,h \, of \, U3 \, ASO \, treatment \, in$ MCF10A (immunofluorescence for FC: RPA194; DFC: FBL; GC: RRP1) and HEK293T (endogenously tagged UBTF-sfGFP (FC), FBL-Halotag (DFC), and NPM1-mtagRFP (GC)) cells. i, Quantification of U3 snoRNA FISH from h.

**** p-value: <0.0001 (two-tailed t-test), HEK293T: n = 160, 110, 46, 284, 27 nucleoli; MCF10A: n = 204, 147, 177, 97, 135 nucleoli. j, DFC rim score following U3 ASO treatment from h. ** p-value = 0.0069 (HEK293T), 0.0088 (MCF10A), **** p-value = <0.0001, HEK293T: n = 160, 14, 27, 167, 17 cells; MCF10A: n = 1050, 94, 138, 83, 68 cells. k, Mean nucleolar FBL intensity by IF upon FBL (n = 278) or control siRNA (n = 541) treatment. I, Western blot of FBL protein levels; β -actin serves as a loading control; For gel source data, see Supplementary Fig. 1c. m, 2'-O-Me levels (ScoreC) at 28S Gm4499 (right), a site modified independently from FBL, and averaged across all other 18S and 28S sites (left) in control and FBL KD treatment conditions measured by 5eU-seq over 15 min pulse, 0-120 min chase timepoints. n = 1-2 per time point. n, Fold change in U8 snoRNA levels (RT-qPCR; n = 3 biological replicates per condition). o, Quantification of U8 snoRNA FISH intensity in nucleoli upon SCR (n = 174) or U8 (n = 40) ASO treatment. p, Fold change in RPL5 mRNA expression by RT-qPCR (n = 3 biological replicates per condition). **** p-value < 0.0001, ** p-value = 0.0011. All scale bars = 3 µm. Box plots show medians (lines), boxes (25th-75th percentiles), whiskers (min-max). Violin plots: centred by median and quartiles shown. All error bars are s.e.m. Statistical tests are two-tailed Mann Whitney tests unless otherwise noted. MCF10A cells used unless otherwise stated.



Extended Data Fig. 6 | **5eU-imaging examples for processing perturbations. a**, Schematic of experimental workflow: cells were treated with various pre-rRNA processing perturbations followed by a 15 min 5eU pulse and chase time course (0-120 min). **b**, Representative images of 5eU-labelled RNA (white) and nucleoli (GC: mTagBFP2-NPM1, blue) in MCF10A cells over indicated chase timepoints under control (SCR) and perturbation conditions. Bottom,

averaged 5eU intensity relative to individual FCs for each condition over time. Scale bars = 3 μ m (cells) and 1 μ m (averaged images). **c-f**, Quantification of 5eU peak distance from the centre of FCs over time for all perturbation conditions shown in b compared to corresponding controls. Number of nucleoli >100 for each time point per condition (see source data for specific n numbers). All error bars are s.e.m.



Extended Data Fig. 7 | Engineered rDNA plasmid designs used in this study. a, Schematics of endogenous (top) 47S rDNA and synthetic (bottom, pSK_ M323) rDNA plasmid with minimized (mini) 5' ETS. Sequences of 3' and 5' hinge regions of 5' ETS-U3 snoRNA base pairing are shown. b, Schematics of wildtype (WT) synthetic SSU-only, 5' ETS 3' hinge mutant SSU-only, and wildtype LSU-only rDNA plasmids. c, Structure of SSU processome in state pre-A1 (PDB: 7mq8). d, Zoom-in on 5' and 3' hinge RNA duplexes between the U3 snoRNA and 5' ETS in c. e-f, Sequences of 3' (e) and 5' (f) hinges of 5' ETS-U3 snoRNA base pairing in wildtype (WT), mutant 5' ETS with mutant U3, and mutant 5' ETS with WT U3 conditions. Sequence substitutions for mutants are marked by double-sided arrows. g, Schematics of complete U3 snoRNA gene combined with rDNA plasmids with 5' ETS and U3 snoRNA 5' hinge and 3' hinge mutations.



Extended Data Fig. 8 | Characterizing the synthetic nucleoli and mature rRNA produced from the engineered rDNA system. a, Size comparison of endogenous and synthetic nucleoli (IF for FBL and NPM1 for DFC and GC, respectively) from endogenous, full-length plasmid, and $\Delta 1,2,3$ plasmid rDNA. Scale bar = 1 µm. b, Quantification of nucleolar area from a. Nucleolar size is not significantly different between any two conditions (two-tailed t-test, endogenous (n = 829), full length rDNA (n = 13), and $\Delta 1,2,3$ rDNA (n = 16)). c, Strand-specific RNA FISH (sense and antisense probes) demonstrate that FISH signal is specific to RNA (not DNA) in cells transfected with $\Delta 1,2,3$ rDNA plasmid (GC, mTagBFP2-NPM1). Scale bar = 10 µm. d, Left, polysome profiling of cells transfected with SSU-only or rDNA plasmids, compared to untransfected controls. Right, RT-qPCR quantification of plasmid-derived rRNA (18S*) and total 18S rRNA in input, monosome, and polysome fractions. e, Representative image of a "Hybrid" nucleolus containing endogenous (endo.) 5' ETS pre-rRNA (yellow) and plasmid-derived rRNAs: 18S* (white) and 28S* (magenta). Scale bar

= 3 μm. **f**, Localization of two GC markers, RRP1 and SURF6 (both detected by IF, blue), in endogenous and SSU-only nucleoli (DFC, NOP56-mCherry). Right, quantification of RRP1 (endogenous: n = 37; SSU-only: n = 10) and SURF6 (endogenous: n = 38; SSU-only: n = 13) intensity. Scale bars = 3 μm (left), = 1 μm (right). Box plots show medians (lines), boxes (25th-75th percentiles), whiskers (min-max). *** P-value < 0.0001 (two-tailed Mann-Whitney test). **g-h**, g, Cells transfected with LSU-only plasmid (28S* FISH, magenta) form a "hybrid" with endogenous nucleoli (endo. 5′ ETS, yellow). Note the absence of cytoplasmic 28S* signal despite the colocalization between plasmid-expressed 28S rRNA in endogenous nucleoli. h, Co-transfection of SSU-only (18S* FISH, white) and LSU-only (28S* FISH, magenta) plasmids to test whether co-localization between plasmid-expressed 18S* and 28S* rRNA is sufficient to rescue cytoplasmic 28S* export. Dotted lines outline nucleoli; solid lines outline nuclei. Arrows indicate colocalization events. DFC: NOP56-mCherry; GC: mTagBFP2-NPM1. Scale bar = 3 μm. HEK293T cells used in all panels.





Extended Data Fig. 9 | Nucleolar morphology changes for engineered rDNA plasmids with impaired U3 snoRNA base pairing. a, De novo nucleolar morphology in HEK293T cells transfected with mutations with rDNA plasmids containing mutations in the 3' or 5' hinge U3 snoRNA binding sites within the 5' ETS, along with compensatory U3 snoRNA mutations. Nucleoli were labelled by IF for DFC (FBL, red), GC (NPM1, blue), and RNA FISH for plasmid-derived 18S* rRNA (white) and endogenous 5' ETS (yellow). b, De novo nucleoli labelled with markers for three nucleolar layers (FC, RPA194 IF; DFC, NOP56-mCherry; GC, mTagBFP2-NPM1) in HEK293T cells transfected with the same plasmids as in a. c, Validation of inverted nucleolar morphology using immunofluorescence staining for additional markers of the DFC (KRR1, ESF1, NOPP140 and UTP23) and GC (NPM1, RRP1, SURF6) in HEK293T cells. **d**, Top, HEK293T cell transfected with 3' hinge mutant rDNA plasmid. Endogenous and de novo nucleoli can be distinguished using RNA FISH for endogenous 28S rRNA (endo 28S, yellow) and plasmid-derived 18S* rRNA (white). Bottom, zoom in on the endogenous and de novo nucleoli labelled with DFC (NOP56-mCherry, red) and GC (mTagBFP2-NPM1, blue) markers. **e**, Quantification of cytoplasmic 28S* plasmid rRNA signal from cells transfected with the indicated plasmids (corresponding to Fig. 4b). n = 36, 74, 84, 103, 92 cells. ** p-value = 0.0026; **** p-value < 0.0001 (two-tailed Mann-Whitney test). Box plots show medians (lines), boxes (25th-75th percentiles), whiskers (min-max). Scale bars = 1 µm (a-c), 3 µm (d).



Extended Data Fig. 10 | **Characterization of perinucleolar chromatin upon inversion and rRNA outflux defects in mutant SSU-only nucleoli. a**, Staining of heterochromatin (H3K9me2/3; GC marked by RRP1IF) and chromatin (DAPI; GC marked by NPM1IF) surrounding de novo nucleoli from wild-type (WT) or 3' hinge mutant rDNA-transfected cells. Right, radial distribution of (hetero) chromatin signal around the GC boundary (dashed line = 0 µm) in endogenous nucleoli, WT de novo nucleoli, or 3' hinge mutant de novo nucleoli. DAPI: n = 1779 (Endogenous), 15 (rDNA plasmid), 15 (3' hinge Mutant); H3K9me2/3 n = 37 (Endogenous), 33 (rDNA plasmid), 30 (3' hinge Mutant). **b**, Radial distribution of chromatin (Hoechst) or heterochromatin (H3K9me2/3) around endogenous nucleoli in scramble (SCR) or U3 ASO treated cells. Hoechst: n = 1958 (SCR), 901 (U3 ASO), H3K9me2/3: n = 2556 (SCR), 562 (U3 ASO). **c-f**, Visualization of SSU processing factors and ribosomal proteins (proteins of interest (POIs), green) in WT and mutant SSU-only nucleoli demarcated by NOP56-mCherry. c-d, Radial distribution of KRII (IF, n = 17), KRR1 (IF, n = 26) and RPS4X-Halotag (n = 15) around the DFC boundary of WT SSU-only nucleoli. See Fig. 4g for images of the other quantified POIs. e-f, Radial distribution of EXOSC10 and RPS6 in WT and mutant SSU-only nucleoli. EXOSC10: n = 12 (WT), 38 (Mutant), RPS6: n = 21 (WT), 47 (Mutant). g, Mean nucleolar 185* rRNA intensity in WT (n = 87) and mutant (n = 57) SSU-only nucleoli. **** P-value < 0.0001. h, Mean nucleolar intensity of early SSU processing factors ESF1, NAT10, and FBL (IF) and NOP56-mCherry in WT and Mutant SSU-only nucleoli. WT: n = 38, 49, 74, 87; Mutant: n = 33, 24, 164, 57 for ESF1, Nat10, Nop56, and Fib. **** P-value = 0.0005; **** P-value < 0.0001. i, DFC area (NOP56-mCherry) in WT (n = 74) and mutant (n = 164) SSU-only nucleoli. **** P-value < 0.0001. All scale bars = 1 μ m. Violin plots are centred by median. Box plots show medians (lines), boxes (25th-75th percentiles), whiskers (min-max). Statistical tests are two-tailed Mann-Whitney tests. All error bars are s.e.m. HEK293T cells are used except for b (MCF10A).

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	\boxtimes	Estimates of effect sizes (e.g. Cohen's d, Pearson's r), indicating how they were calculated
	I	Our web collection on <u>statistics for biologists</u> contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Images were acquired with the NIS-Elements AR (Nikon, 5.42.02) with Nikon CSU-W1 SoRa spinning disk confocal system. Denoising (when Data collection applicable) was performed using the built-in NIS-Elements Denoise.ai module. All statistical analyses and graphing were performed using Graphpad Prism 10 and Microsoft Excel. For creating example images, imageJ Data analysis (1.53f) was used. All codes and pipelines used for this study are provided in the following repositories: https://github.com/SoftLivingMatter/image-analysis-quinodoz-jiang-2024 (https://zenodo.org/records/14908468) https://github.com/SoftLivingMatter/snakemake-cellprofiler (https://zenodo.org/records/14908470) https://github.com/SoftLivingMatter/5eU-seq-pipelines (https://zenodo.org/records/14908466) Example images & analysis notebooks: https://doi.org/10.5281/zenodo.14910509

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All data supporting these findings are available in the manuscript and associated source data. All genomics data are available at the Gene Expression Omnibus (GEO) with accessions: GSE296080 and GSE296162.

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Recruitment	n.a.
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Sample size	Sample sizes were not predetermined. For imaging experiments, sample size were chosen to make sure a representative population were captured. For sequencing experiments, number of replicates, time points, and cells were determined based on technical requirements that ensure reproducibility and maintain high data quality.
Data exclusions	For imaging, cells that are out of focus are excluded. For 5eU sequencing, samples that failed quality control (such as failed click reactions, streptavidin captures, or library amplifications) were excluded from downstream analysis.
Replication	Microscopy imaging, western blotting, and RT-qPCR were repeated independently for at least three times with similar results. 5eU-sequencing were conducted with 1 or 2 replicates per time point. P-values and number of observations (n) are provided in figure legends. All attempts at replication were successful.
Randomization	For all experiments, cells were randomized such that within a biological or technical replicate all cells were analyzed equally with no sub- sampling.
Blinding	Blinding not performed due to standardized data collection procedures. Imaging and sequencing analyses were performed using automated pipelines whenever possible to minimize observer biases.

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	Eukaryotic cell lines
\boxtimes	Palaeontology and archaeology
\boxtimes	Animals and other organisms
\boxtimes	Clinical data
\boxtimes	Dual use research of concern
\boxtimes	Plants

Methods

n/a Involved in the study

- ChIP-seq
- Flow cytometry
- MRI-based neuroimaging

Antibodies

Antibodies used	1 Antihodies for Immunofluresence
Antiboules used	Artikody Name runplier name catalog number, working dilution
	Antibody Ivanie supplier hante datalog humbel - working diddon
	DDA104 499 Sonto Cruz os 49395 1:200
	NPA134-400 Salid Clu2 SC40505 1.200
	NP/MI-486 INE/INO FISHEI MIAS-23200-A488 1:400
	Fibrillatin Abdalli absoli 1:300
	Pipiliarin-568 Abcam abz02540 1:300
	KNP1B SIGMA HPA017893 1:100
	SURF6 Thermo Fisher PA5-54841 1:100
	Nat10 Thermotisher 13365-1-AP 1:100
	ESF1 proteinTech 23496-1-AP 1:100
	EXOSC10 Sigma-Aldrich HPA028484 1:100
	Nucleolin ThermoFisher 39-6400 1:100
	DDX21 Sigma-Aldrich HPA036593 1:100
	KRI1 proteintech 16243-1-AP 1:100
	KRR1 Assay Genie CAB4487-20 1:100
	H3K9me2/3 Cell Signaling Technology 5327T 1:100
	UTP23 proteinTech 15950-1-AP 1:100
	NOPP140 Santa Cruz Biotechnology sc-374033 1:200
	Goat anti-Mouse IgG (H+L) Cross-Adsorbed Secondary Antibody, Alexa Fluor™ 568 Thermofisher A-11004 1:1000
	Goat anti-Rabbit IgG (H+L) Cross-Adsorbed Secondary Antibody. Alexa Fluor™ 488 Thermofisher A-11008 1:1000
	Goat anti-Rabbit IgG (H+L) Cross-Adsorbed Secondary Antibody, Alexa Fluor™ 405 Thermofisher A-31556 1:1000
	2. Antibodies for Western blotting
	Antibody Name -Source -Identifier-Working dilution
	Fibrillarin Abcam ab5821 1:1000
	8-Actin Antibody #4967 Cell Signalling 49675 1:1000
	NPM1 Monoclonal Antibody (4TOLI-182) Thermofisher MA3-086 1:1000
	Perovidase difinitive Goat Anti-Rabbit InG (H+1) larkson ImmunoResearch 111-035-144 1:10,000
	Peroxidase AffiniPure™ Goat Anti-Mouse IgG (H+L) Jackson ImmunoResearch 115-035-062 1:10, 000
Validation	All antibodies used for immunofluorescence (IE) are validated by confirming that the intracellular localization matches the reported
Validation	cellular localization. Western blotting (WB) antibodies are validated by the molecular weight of the band detected and the reduced
	intensity upon knockdown for fibrillarin In addition antibodies were validated by the manufacturer e_{σ} ESE1 (proteinTech 23496-1)
	A) has been tested in mouse heart tissue for WB and MCF-7 cells for IF and it has tested reactivity against mouse and human
	Fibrillarin (Abram ab5821 and ab202540) has been tested for WB and IE in human mouse and Prosphila melanogaster RP1B

Eukaryotic cell lines

Policy information about cell lines and Sex and Gender in Research

Cell line source(s)	The following cell lines are used in this study: MCF10A (gift from Yibin Kang, ATCC, female), HEK293T (ATCC, female), HCT116 (gift from Yibin Kang, ATCC, male) and MCF7 (gift from Yibin Kang, ATCC, female). All cell lines were derived from human participants
Authentication	All the gifted cell lines were validated by STR profiling with > 90% match. Vendors of each of the commercially-available cell lines (ATCC) provide further information about the authentication of cells on their website.
Mycoplasma contamination	All cell lines used in this study tested negative for mycoplasma contaminiation.
Commonly misidentified lines (See <u>ICLAC</u> register)	No cell lines used in this study were among the commonly misidentified lines

Cruz sc-48385) have been tested in IF with localization to nucleoli.

(Sigma HPA017893), SURF6 (Thermo Fisher PA5-54841), Nat10 (Thermofisher 13365-1-AP), DDX21 (Sigma-Aldrich HPA036593), NPM1-488 (Thermo Fisher MA3-25200-A488), Nucleolin (ThermoFisher 39-6400), RRP1B (Sigma HPA017893), and RPA194 (Santa

Plants

Seed stocks	n.a.	ļ		
Novel plant genotypes	n.a.			
Authentication	n.a.			

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