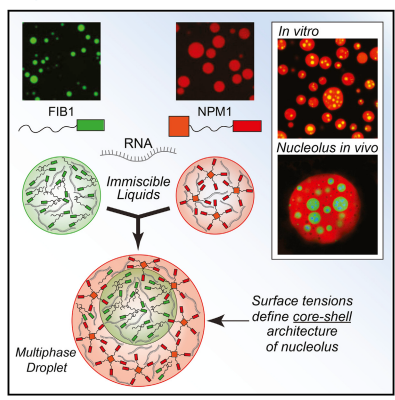


Coexisting Liquid Phases Underlie Nucleolar Subcompartments

Graphical Abstract



Authors

Marina Feric, Nilesh Vaidya, Tyler S. Harmon, ..., Richard W. Kriwacki, Rohit V. Pappu, Clifford P. Brangwynne

Correspondence

cbrangwy@princeton.edu

In Brief

Immiscibility of different liquid-like phases comprised of RNA and protein can recapitulate the subcompartmentalized organization characteristic of nucleoli.

Highlights

- Multiple subcompartments of nucleoli are immiscible liquid phases
- Mixing nucleolar proteins in vitro yields similar multiphase behavior
- Disordered domains drive phase separation; RNA binding domains confer immiscibility
- Differences in droplet surface tensions organize core-shell structure



Article

Cell

Coexisting Liquid Phases Underlie **Nucleolar Subcompartments**

Marina Feric, 1,5 Nilesh Vaidya, 1,5 Tyler S. Harmon, 2,3 Diana M. Mitrea, 4 Lian Zhu, 1 Tiffany M. Richardson, 1 Richard W. Kriwacki, A Rohit V. Pappu, and Clifford P. Brangwynne^{1,7}

- ¹Department of Chemical and Biological Engineering, Princeton University, Princeton, NJ 08544, USA
- ²Department of Physics, Washington University in St. Louis, St. Louis, MO 63130, USA
- ³Department of Biomedical Engineering and Center for Biological Systems Engineering, Washington University in St. Louis, St. Louis, MO 63130, USA
- ⁴Department of Structural Biology, St. Jude Children's Research Hospital, Memphis, TN 38103, USA
- 5Co-first author

*Correspondence: cbrangwy@princeton.edu http://dx.doi.org/10.1016/j.cell.2016.04.047

SUMMARY

The nucleolus and other ribonucleoprotein (RNP) bodies are membrane-less organelles that appear to assemble through phase separation of their molecular components. However, many such RNP bodies contain internal subcompartments, and the mechanism of their formation remains unclear. Here, we combine in vivo and in vitro studies, together with computational modeling, to show that subcompartments within the nucleolus represent distinct, coexisting liquid phases. Consistent with their in vivo immiscibility, purified nucleolar proteins phase separate into droplets containing distinct non-coalescing phases that are remarkably similar to nucleoli in vivo. This layered droplet organization is caused by differences in the biophysical properties of the phases—particularly droplet surface tension which arises from sequence-encoded features of their macromolecular components. These results suggest that phase separation can give rise to multilayered liquids that may facilitate sequential RNA processing reactions in a variety of RNP bodies.

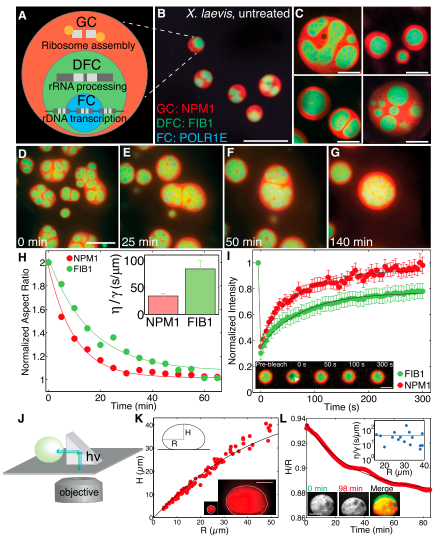
INTRODUCTION

The cellular interior is organized into organelles whose structures have evolved to facilitate their functions. The most well-known examples are the canonical membrane-bound organelles such as secretory vesicles, the Golgi apparatus, and the endoplasmic reticulum. However, many intracellular compartments are membrane-less bodies comprised of RNA and protein, often referred to as RNP bodies; these include stress granules and processing bodies in the cytoplasm and Cajal bodies and nucleoli in the nucleus, among many others. Despite their lack of a delimiting membrane, these organelles nevertheless maintain a coherent size and shape, with a well-defined boundary that compartmentalizes different types of proteins and RNA. By concentrating molecules within a micro-compartment, while allowing dynamic molecular interactions, these RNP bodies may function to control reaction efficiencies much like conventional membrane bound cytoplasmic organelles (Balagopal and Parker, 2009; Spector, 2001).

Many of these RNP bodies exhibit liquid-like biophysical properties, and growing evidence suggests they assemble via liquidliquid phase separation (Brangwynne et al., 2009; Nott et al., 2015; Li et al., 2012; Weber and Brangwynne, 2015). Intracellular phase transitions can result in switch-like changes in molecular organization and the spontaneous formation of micron-scale membrane-less organelles. Such behavior is reminiscent of well-known in vitro observations in protein crystallization, where soluble proteins are observed to condense into concentrated liquid phases or crystalline solid phases. A number of recent papers suggest that intrinsically disordered proteins or low complexity sequences (IDP/LCS) drive phase transitions underlying assembly of the nucleolus (Berry et al., 2015), stress granules (Wippich et al., 2013; Molliex et al., 2015; Patel et al., 2015), P granules and nuage bodies (Elbaum-Garfinkle et al., 2015; Wang et al., 2014), and nuclear speckles (Hennig et al., 2015).

The liquid-like nature of the nucleolus may facilitate its function in ribosome biogenesis. The nucleolus forms around regions of chromosomes containing stretches of tandem rDNA gene repeats, known as nucleolar organizer regions (NORs). In most eukaryotes (including human, Xenopus laevis, and Caenorhabditis elegans), a precursor rRNA transcript is generated from the rDNA gene and contains each of the co-transcribed 18S, 5.8S, and 28S rRNA subunits, separated by two intervening transcribed sequences (Figure 1A). The nucleolus may facilitate increased reaction rates by locally concentrating rRNA processing factors involved in transforming the precursor rRNA transcript into individual rRNA subunits. Due to its role in producing this protein translational machinery, the structure and function of the nucleolus are intimately connected with cell growth and size homeostasis (Derenzini et al., 2000; Frank and Roth, 1998).

Despite the biological importance of the nucleolus, a mechanistic biophysical understanding of its assembly and internal organization is lacking. The simplest picture of the nucleolus as a unitary liquid phase body becomes difficult to reconcile with its well-known complex and multi-component nature. Indeed, in addition to the various types of RNA in the nucleolus, the nucleolar proteome consists of hundreds of different proteins that are



segregated into at least three distinct compartments (Boisvert et al., 2007). This layered tripartite organization consists of the fibrillar center (FC), where the RNA polymerase I (POL1) machinery is active; the dense fibrillar component (DFC) that is enriched in the protein fibrillarin (FIB1); and the granular component (GC) that is enriched in the protein nucleophosmin (NPM1/B23) (Figures 1A and 1B). This multi-layered structure is not unique to the nucleolus, as stress granules and other liquid-like RNP bodies exhibit similar "core-shell" structuring (Jain et al., 2016; Hubstenberger et al., 2013).

The multi-layered structure of the nucleolus is thought to facilitate assembly line processing of rRNA. Nascent rRNA transcripts undergo sequential processing steps by enzymes that localize to the distinct compartments, ultimately exiting the nucleolus and being exported for final ribosome assembly in the cytoplasm (Figure 1A). Although recent work has shown that the entire nucleolus can exhibit active liquid-like properties (Brangwynne et al., 2011) and its assembly may represent a type of liquid-liquid phase transition (Weber and Brangwynne,

Figure 1. Liquid-like Behavior Distinct Nucleolar Subcompartments

(A) Schematic diagram of ribosome biogenesis in nucleolus.

(B) Nucleoli in an untreated X. laevis nucleus. Scale bar, 20 $\mu\text{m}.$ For all images, granular component (GC) is visualized with NPM1 (red), dense fibrillar component (DFC) with FIB1 (green), and fibrillar center (FC) with POLR1E (blue).

(C) Examples of nucleoli after coarsening in X. laevis nuclei treated with Lat-A. Scale bar, 20 μm.

(D-G) Time-course of nucleolar component fusion after actin disruption by Lat-A. Scale bar, 20 µm. (H) Normalized aspect ratio versus time for nucleolar components fusing after actin disruption. Inset shows η/γ for 59 nucleoli analyzed from 20 nuclei.

(I) FRAP recovery curves for NPM1 (red) and FIB1 (green) in X. laevis oocytes. Inset: FRAP of FIB1labeled DFC (green). Scale bar, 5 μm.

(J) Schematic illustrating XZ imaging with a right angle prism.

(K) Height, H, versus radius, R, of different sized nucleoli at steady-state (91 nucleoli, from 61 nuclei). Black line is the fit from the average surface tension for all nucleoli. Bottom inset: example of the shape of a small versus large nucleolus. Scale bar, 40 um.

(L) Example of nucleolar height to radius ratio, H/R, versus time for one nucleolus deforming under gravity. Black line is an exponential fit. Top inset: η/γ for 16 nucleoli in 16 different nuclei. Blue line indicates average. Bottom inset shows example deforming nucleolus: Scale bar, 40 µm.

See also Figure S1 and Movies S1, S2, S3, and S4.

2015), reconciliation of these findings with the multi-layered structure of the nucleolus has proven elusive. Indeed, if the nucleolus is a type of liquid, what

mechanism prevents the three components from mixing and fusing to form a single liquid phase?

Here, we uncover a physical mechanism for intranucleolar organization: differences in miscibility between proteins from different nucleolar compartments keep the compartments phase separated, giving rise to the layered, multiphase droplet nature of nucleoli. By isolating protein domains from key nucleolar proteins, we provide evidence for a molecular mechanism whereby intrinsically disordered regions (IDRs) help drive protein condensation into droplets, while associated RNA binding domains confer subcompartment specificity by making the two droplet phases immiscible with one another.

RESULTS

Nucleolar Subcompartments Behave as Liquid-like Phases In Vivo

To gain insight into the biophysical assembly principles underlying nucleolar structure, we took advantage of the numerous large nucleoli, ranging in size from one to ten microns, found in the nucleus (germinal vesicle [GV]) of X. laevis oocytes (Figure 1B). This system is also convenient because the nucleoli are extra-chromosomal, forming around amplified stretches of rDNA, allowing us to disentangle the confounding effects of somatic chromosome architecture on nucleolar structure. Nucleoli in X. laevis oocytes will fuse with one another when brought into contact (Brangwynne et al., 2011), but the frequency of such coalescence events is slowed significantly by the presence of a nuclear actin network (Figures S1A-S1D) (Feric and Brangwynne, 2013; Feric et al., 2015).

We visualized intranucleolar organization by labeling individual components of nucleoli with fluorescent fusion proteins as follows: granular component with nucleophosmin (GC, NPM1:: Cerulean), dense fibrillar component with fibrillarin (DFC, FIB1::RFP), and the fibrillar center with RNA polymerase 1E (FC, GFP::POLR1E). To test the role of nuclear actin in organizing these nucleolar substructures, we utilized the actin-disrupting compound Latrunculin-A (Lat-A). As shown previously, the entire nucleolus undergoes liquid-like coalescence events, which can be seen by fusion of the NPM1 (GC) region of one nucleolus with the NPM1 (GC) region from another nucleolus (Figures 1D-1G; Movies S1, S2, and S3). The FC regions (POLR1E) rarely came into close contact with each other, and consequently, we did not observe fusion between multiple FCs, but we did see rearrangements into more spherical FC domains. Strikingly, however, we typically observe that the FIB1 (DFC) cores from one nucleolus will fuse when in close proximity with FIB1 (DFC) cores from a different nucleolus (Figures 1D-1G and S1E-S1L). These DFC regions, which were initially irregular in shape, would round up and coalesce, exhibiting classic liquid-like behavior. After long times (~1 hr), these coalescence events ultimately resulted in DFCs located in the center of the nucleolus, surrounded by one continuous phase of GC, representing an outer-most enveloping compartment (Figure 1C).

In Vivo Subcompartments Have Different Biophysical **Properties**

To gain insight into the biophysical properties of different nucleolar subcompartments, we quantitatively analyzed fusion events. We found that homotypic fusion between the GC (NPM1) or DFC (FIB1) occurs by exponential relaxation to a single larger spherical shape; this is characteristic of coalescing liquid droplets and can be used to determine the ratio of droplet viscosity, η , to surface tension, γ , known as the inverse capillary velocity: η/γ (Brangwynne et al., 2011) (Figure 1H). We find that FIB1-labeled DFC tends to exhibit slower fusion dynamics, with a larger value $\eta/\gamma = 80 \pm 15 \text{ s/}\mu\text{m}$ (mean \pm SEM) compared to NPM1-labeled GC with $\eta/\gamma = 30 \pm 5$ s/ μ m (mean \pm SEM) (Figures 1H, inset, S1M, and S1N). This behavior suggests that these nucleolar components behave as distinct liquid-like phases within the nucleolus, with different properties that could underlie nucleolar structural organization.

Interfaces represent sharp concentration gradients and surface tension, with units of energy per unit area, is the energetic cost of increasing the interfacial area. Surface tension is a key parameter that governs how two different droplets interact with one another. We therefore hypothesized that different surface

tension values of the nucleolar sub-phases could explain their immiscibility and multi-layered organization. To measure the surface tension of the outermost GC compartment (i.e., interfacial energy of GC/nucleoplasm interface), we disrupted actin and allowed nucleoli to fuse for several hours. This resulted in a single large coalesced nucleolar droplet, which becomes measurably flattened at the bottom of the nucleus due to gravity. Because this flattening is resisted by surface tension, we could use a right-angle prism to measure the (XZ) shape of the droplet and determine the surface tension: $4\pm 1 \times 10^{-7}$ N/m (mean \pm SEM) (Figures 1J and 1K; Supplemental Experimental Procedures); this value is very low, roughly five orders of magnitude lower than water-oil surface tension values (Than et al., 1988), but is comparable to values reported for colloidal liquids (Aarts et al., 2004). Complementary measurements of the droplet flattening timescale combined with droplet fusion measurements are consistent with this low value for the surface tension of the NPM1-rich GC compartment (Figure 1L; Movie S4; Supplemental Experimental Procedures).

To confirm the liquid-like dynamics of the nucleolar subcompartments, we performed fluorescence recovery after photobleaching (FRAP) experiments of X. laevis nucleoli in vivo (Figure 11). NPM1 exhibits fast dynamics with a nearly complete recovery on a timescale of τ = 64 \pm 8 s for a bleach spot of 1 μ m, consistent with the expected response for diffusion within a liquid. However, FIB1 recovery was slightly slower ($\tau = 75 \pm 7$ s). Moreover, the recovery of FIB1 was not complete, but only reached ~80%; this suggests that the DFC subcompartment may not be a simple liquid, but instead may exhibit partially solid-like properties (i.e., viscoelasticity).

Purified FIB1 and NPM1 Can Phase Separate into Droplets Similar to Nucleoli In Vivo

To gain further insight into how nucleolar proteins could give rise to distinct liquid-like nucleolar sub-phases, we purified recombinant FIB1 and NPM1 and studied their behavior in vitro. Consistent with our previous work, we find that FIB1::GFP (hereafter referred to simply as FIB1) can phase separate in vitro under near physiological protein and salt concentrations (Berry et al., 2015), giving rise to condensed liquid-phase droplets that are \sim 50-fold more concentrated with protein than the surrounding dilute phase (Figure 2A). Indeed, in the presence of 5 μg/ml rRNA and 150 mM NaCl, FIB1 condenses into droplets at a protein concentration of $\sim\!600$ nM. FIB1 can phase separate even in the presence of non-specific poly-U50 and heparin, suggesting that electrostatic interactions contribute to droplet assembly (Figure S2A). NPM1 has also recently been demonstrated to undergo phase separation into liquid-like droplets (Figure 2B) (Mitrea et al., 2016). However, at 150 mM NaCl, NPM1 requires significantly higher concentrations of protein (2 μM) and rRNA (100 µg/ml). Moreover, phase separation of NPM1 required rRNA and cannot be induced by the addition of heparin or poly-U50.

Given the distinct biophysical properties of the nucleolar subcompartments, we hypothesized that the two different types of in vitro droplets would also exhibit different material properties. As with the in vivo subcompartments, in vitro droplets undergo homotypic fusion events when brought into close contact, but

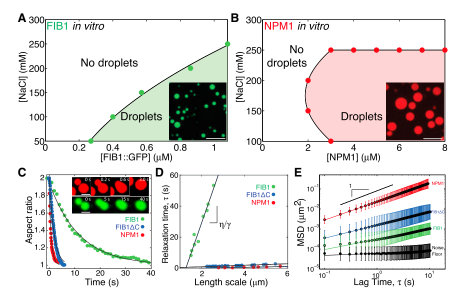


Figure 2. Purified Nucleolar Proteins Can **Phase Separate into Droplets with Different Biophysical Properties**

- (A) Phase diagram of purified FIB1 in the presence of 5 µg/ml of rRNA. Inset: FIB1 droplets. Scale bar, 10 μm.
- (B) Phase diagram of purified NPM1 in the presence of 100 $\mu g/ml$ of rRNA. Inset: NPM1 droplets. Scale bar, 10 µm.
- (C) Aspect ratio versus time for fusing droplets of FIB1 (green), NPM1 (red), and FIB1ΔC (blue). Inset: FIB1 fusing (scale bar, 2 µm) and NPM1 fusing (scale bar, 5 µm).
- (D) Relaxation time versus length scale for fusion data from multiple FIB1 (green), NPM1 (red), and FIB1ΔC droplets (blue).
- (E) MSD versus lag time of microrheological probe particles (R = 50 nm) embedded in droplets of FIB1 (green), NPM1 (red), or FIB1ΔC (blue); black data points represent the noise floor (black).

See also Figure S2.

do so with markedly different timescales: FIB1 droplets typically take roughly a hundred times longer than NPM1 droplets of comparable size to coalesce and relax into a single larger sphere (Figure 2C). Using an analysis similar to that performed in vivo, we find that FIB1 has an inverse capillary velocity of $\eta/\gamma = 40 \pm 10 \text{ s/}\mu\text{m}$ (95% confidence interval), comparable to that measured in vivo (Figure S2C). Also mirroring the in vivo data, purified NPM1 droplets have a lower value of 0.30 \pm 0.07 s/µm (Figures 2D and S2C). A series of experiments confirmed that the presence of the GFP tag does affect droplet properties, but does not qualitatively impact our findings (Figures S2B and S2C; Supplemental Experimental Procedures).

NPM1 readily formed large droplets in vitro, allowing for direct measurement of the surface tension, using a method similar to our in vivo set up with the prism (Figure 1J). We estimated a surface tension of γ_{NPM1} = 8 ± 2 × 10⁻⁷ N/m (mean ± SEM) (Figure S2D). This value is again surprisingly low and on the same order of magnitude as the value obtained for X. laevis nucleoli (Figure S2C).

Viscoelasticity and Time-Dependence of In Vitro **Droplets**

To further shed light on the different properties of the two droplet subtypes, we performed microrheology experiments using the fluctuating motion of probe particles (R = 50 nm) (Elbaum-Garfinkle et al., 2015; Zhang et al., 2015). These data reveal that NPM1 droplets exhibit a diffusive exponent of $\alpha = 0.92 \pm 0.06$, (Figure 2E, red symbols), consistent with that of a simple viscous liquid for which $\alpha = 1$. We can thus calculate a viscosity of NPM1 droplets, η = 0.74 \pm 0.06 Pa-s, which is several hundred times more viscous than water (Figure S2C). By contrast, probe particle motion in FIB1 droplets is significantly reduced (Figure 2E, green symbols), in agreement with the slowed coalescence dynamics observed with FIB1 droplets. Interestingly, FIB1 droplet microrheology reveals a sub-diffusive exponent ($\alpha = 0.5 \pm 0.1$), which implies that these are not simple viscous liquid droplets, but are instead viscoelastic.

To determine how FIB1 droplet viscoelasticity may arise, we performed FRAP experiments on phase-separated in vitro droplets (Figure 3). After 30 min of initiating phase separation, we find that NPM1 has near complete recovery (84% \pm 3%) on very short timescales, with a recovery constant of $\tau = 23 \pm 1$ s (Figures 3A) and S3C); this is consistent with the purely viscous microrheology results, as well as the nearly complete NPM1 FRAP recovery in vivo. By contrast FIB1 has low recovery (37% \pm 2%) with a timescale of $\tau = 56 \pm 5$ s (Figures 3B and S3C). Such incomplete FRAP recovery is expected for a viscoelastic material, since not all molecules exhibit dynamic, fluid-like exchange. Moreover, this in vitro FIB1 behavior agrees well with the incomplete FIB1 FRAP recovery in vivo, in both cultured mammalian cells expressing FIB::GFP and amphibian nucleoli (Figure 3F).

To test whether droplet material properties change with time. we performed FRAP experiments on in vitro droplets as a function of time. We find that NPM1 always exhibits a near complete recovery, even for droplets that have been sitting for several hours (Figures 3A and 3D). However, FIB1 FRAP dynamics are strongly impacted by the droplet age. By 2 hr, the percent recovery has dropped by a factor of \sim 4, to 8% \pm 0.5% (Figures 3B and 3D); this suggests that these droplets become increasingly solidlike with time, potentially due to the formation of fibers. Consistent with this, we see liquid-like FIB1 droplets evolve into sticky gel-like structures over a 2-hr time period (Figure S3A). Also, we find that replacing the GFP tag with the monomeric GFP (A206K) did not alter the FIB1 aging behavior (Figure S3B). As we describe further below, the N-terminal R/G-rich domain of FIB1, designated as FIB1ΔC, drives phase separation as an autonomous unit. However, it exhibits nearly complete FRAP recovery (Figures 3C and 3D). Moreover, unlike full-length FIB1, the percent recovery is stable over 4 hr (Figure 3D). Therefore, we conclude that the C-terminal methyltransferase domain of FIB1 plays a key role in promoting viscoelastic maturation of FIB1 droplets in vitro.

ATP-dependent active processes have been hypothesized to play an important role in promoting dynamics within cells, in a

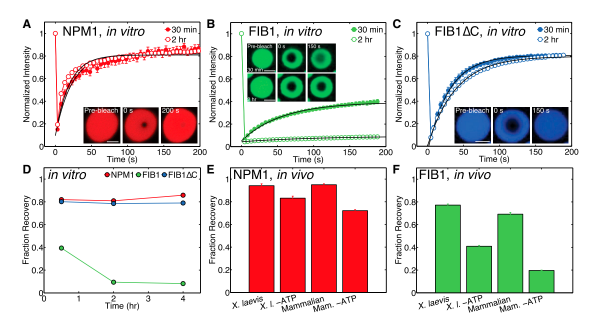


Figure 3. Nucleolar Protein Droplets Exhibit Liquid-like Dynamics, but FIB1 Shows Evidence for Aging

(A-C) FRAP recovery curves for NPM1 (red), FIB1 (green), and FIB1\(\Delta\)C (blue) droplets, 30 min (closed circles) and 2 hr (open squares) after phase separation was initiated. (A) Inset: example FRAP time course. Scale bar, 5 µm. (B) Insets: example FRAP time courses after 30 min (top) and 2 hr (bottom). Scale bar, 2 µm. (C) Inset: example FRAP time course. Scale bar, 2 μm .

(D) Fraction recovery after FRAP experiment as a function of time after phase separation for NPM1 (red), FIB1 (green), and FIB1∆C (blue) droplets. (E and F) Fraction recovery for NPM1 (E) and FIB1 (F) in X. laevis nucleoli and mammalian cell culture nucleoli in vivo, for native and ATP depletion conditions. See also Figure S3.

process known as "active diffusion" (Brangwynne et al., 2011; Parry et al., 2014; Kroschwald et al., 2015). Depleting ATP from X. laevis oocytes or mammalian cells (Figure 3E) had relatively little effect on FRAP recovery of NPM1, with nearly full FRAP recovery, comparable to that seen with in vitro NPM1 droplets (Figures 3A and 3D). By contrast, ATP-depletion resulted in significantly slowed FIB1 dynamics, with longer recovery times in X. laevis (Figure S3C) and a 2- to 3-fold decrease in the percent recovery in both systems (Figure 3F). Moreover, the low percent recovery of FIB1 in ATP-depleted cells (20%-40%) is similar to that measured for in vitro FIB1 droplets (Figures 3D and 3F). This suggests that ATP-dependent enzymatic activity is essential for actively maintaining the fluidity of the aging-prone, FIB1-rich DFC.

In Vitro FIB1 and NPM1 Coexist as Multiphase Droplets

Given that FIB1 and NPM1 individually phase separate into liquid droplets in the presence of rRNA, we next tested how these proteins behave when mixed together. At relatively low concentrations, both proteins colocalize in the same condensed droplets; depending on the relative amount of FIB1 to NPM1, these droplets are either enriched in FIB1 (FIB1-rich/NPM1-lean phase) (Figures 4B and S4B) or they are enriched in NPM1 (FIB1-lean/ NPM1-rich phase) (Figures 4C and S4C). Thus, considering the soluble "buffer" phase, in these cases the system still resides within a two-phase region of the phase diagram (Figure 4D). However, when these proteins are both mixed at relatively high concentrations, we observe a three-phase system with both FIB1-rich/NPM1-lean droplets coexisting with FIB1-lean/

NPM1-rich droplets, surrounded by the buffer phase (Figure 4A). Interestingly, the NPM1 rich phase tends to partially envelope the FIB1 rich phase (Figure S4A); in the absence of the GFP tag, this envelopment becomes even more pronounced, with FIB1 droplets fully embedded within NPM1 (Figure S4D). This droplet organization is very similar to what is observed in X. laevis and mammalian nucleoli, where the FIB1 DFC is always internalized within the NPM1 GC. A phase diagram can be constructed by determining the threshold concentrations of FIB1 and NPM1 required to phase separate into a three-phase system (Figure 4D). These results suggest that the "layered" structural organization of nucleolar proteins could be self-organized by liquidliquid phase separation alone.

To test whether qualitatively similar phase behavior may be observed upon changing the relative protein concentrations in living cells, we injected nucleolar proteins into living X. laevis nuclei (Figure 4E). Consistent with the expectation from equilibrium phase coexistence, we observed that the volume fraction of the corresponding component increased after microinjection (Figure 4H). Typically, the DFC visualized by FIB1 is $25\% \pm 2\%$ (mean \pm SEM) of the volume in the nucleolus. When more FIB1 was injected, the protein localized preferentially to the DFC causing the fibrillar cores to increase in size, occupying \sim 37% \pm 3% (mean \pm SEM) of the volume. Conversely, when NPM1 was injected, the protein localized preferentially to the GC and caused the nucleoli to swell to large sizes, causing the fibrillar cores to occupy a lower volume fraction of only 15% \pm 1% (mean ± SEM). Moreover, small extranucleolar droplets of the respective protein appeared to form de novo (Figures 4F

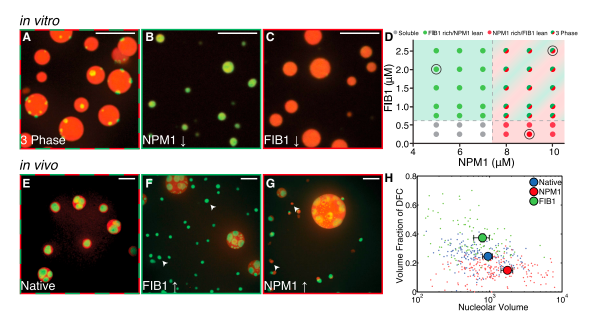


Figure 4. FIB1 and NPM1 Form Immiscible Droplets In Vitro and In Vivo

(A–C) In vitro images of mixtures of purified NPM1 and FIB1. Scale bar, 10 μm. (A) High concentrations of both proteins (FIB1: 2.5 μM, NPM1: 10 μM) give rise to FIB1-rich droplets (green) that are immiscible with and partially enveloped by NPM1-rich droplets (red). (B) For much lower concentrations of NPM1 (NPM1: 5 µM, FIB1: 2 μM) only FIB1-rich/NPM1-lean droplets are observed. (C) For much lower concentrations of FIB1 (FIB1: 0.25 μM, NPM: 9 μM) only NPM1-rich/FIB1-lean droplets are observed.

(D) Phase diagram for varying concentrations of NPM1 and FIB1 in vitro. Colors represent observed phase (gray, soluble phase; green, FIB1 rich/NPM1 lean phase: red. NPM1 rich/FIB1 lean phase: red/green, three phase). Black circles indicate concentrations shown in (A)–(C).

(E-G) Images of nucleoli in X. laevis; red, NPM1; green, FIB1. Scale bar, 10 μm. (E) Untreated nuclei. (F) Nuclei after microinjection of FIB1. (G) Nuclei after microinjection of NPM1.

(H) Volume fraction of the DFC (identified by FIB1) in each nucleolus for native nuclei (blue), after NPM1 injection (red) and after FIB1 injection (green). Large symbols represent mean ± SEM.

See also Figure S4.

and 4G). This is possible if the saturation concentration in the nucleoplasm has been reached, causing spontaneous condensation of nucleolar proteins, without requiring NORs for nucleation (Berry et al., 2015).

Protein Domains Required for Phase Separation and Immiscibility

To gain insight into the molecular-scale driving forces underlying phase separation and droplet immiscibility, we created deletion mutants of both FIB1 and NPM1 that contained individual domains. Full-length FIB1 consists of two domains: a disordered N-terminal arginine (R)/glycine (G)-rich domain with lowsequence complexity (R/G domain) and an RNA methyltransferase domain (MD) that together with small nucleolar RNA (snoRNA) can methylate substrate rRNA (Figure 5A). We find that the R/G domain (FIB1 Δ C) is sufficient to form liquid-like droplets in vitro, while the MD alone (FIB1 AN) is unable to form droplets in vitro (Figure 5A). Interestingly, FIB1ΔC can phase separate into liquid-like droplets in vitro, even in the absence of RNA (Figure S2E). By contrast, full-length FIB1 requires rRNA; however, this may be a non-specific consequence of the polyanionic nature of rRNA since heparin can also drive phase separation of full-length FIB1 (Figures S2A and S2F). Furthermore, we find that FIB1ΔC droplets undergo homotypic fusion with an inverse capillary velocity of $\eta/\gamma = 0.5 \pm 0.04$ s/µm (95% confidence interval); these dynamics are significantly faster than for full-length FIB1 ($\eta/\gamma \approx 40 \pm 10 \text{ s/}\mu\text{m}$) and comparable to NPM1 ($\eta/\gamma \approx 0.3 \pm 0.07$ s/ μ m) (Figures 2C, 2D, and S2C). Consistent with this, in microrheology experiments, FIB1ΔC droplets also exhibit faster dynamics than full-length FIB1 (Figures 2E, blue symbols, and S2C).

When these mutant proteins were injected into X. laevis nuclei, FIB1ΔN strongly partitions to the DFC, similar to the full-length FIB1 protein (Figure 5C). Similarly, we observed that FIB1ΔN does not colocalize with NPM1 droplets in vitro (Figures 5C and S5A). DFC enrichment of FIB1ΔN in vivo may not reflect immiscibility, but may instead reflect co-recruitment due to the presence of full-length FIB1 in the native system. Moreover, we observed that injected FIB1 Δ C localizes to the entire nucleolus and has nonspecific interactions with the GC and DFC. Consistent with this, we find that the FIB1\DC colocalizes with in vitro NPM1 droplets, rather than forming a third immiscible droplet phase (Figure S5A). Taken together, these data suggest that the N-terminal R/G domain of FIB1 is sufficient for droplet formation, but does not encode for a separate liquid-like DFC subcompartment; instead, the C-terminal MD of FIB1, which alone is not sufficient for droplet formation, confers immiscibility with proteins in the GC.

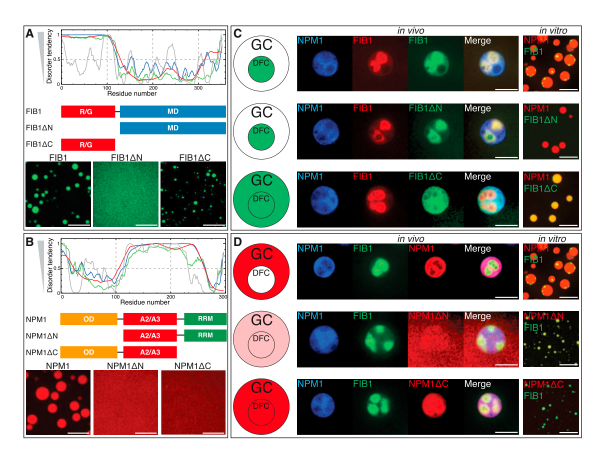


Figure 5. Molecular Mechanism of Phase Partitioning in X. laevis Oocytes

(A) Domain analysis of FIB1. Plot shows predicted disorder across full-length FIB1 using various algorithms, P-FIT (green line), VSL2B (blue line), VL3 (red line), and VLXT (gray line). Schematic diagrams show three constructs: FIB1 full-length, R/G deletion (FIB1 \(\Delta \N \)), and deletion of MD (FIB1 \(\Delta C \)), with images below testing for constructs' ability to form droplets. Scale bar, 10 um.

(B) Domain analysis of NPM1. Predicted disorder across full-length NPM1 for the four algorithms. Schematic diagrams show three constructs: NPM1 full-length, oligomerization deletion (NPM1ΔN), and RNA binding deletion (NPM1ΔC) with images below testing for constructs' ability to form droplets. Scale bar, 10 μm. (C and D) The left most panel shows schematic summary of center panels. Center panels contain images from X. laevis nucleoli in vivo. Left channel contains expression of mRNA for NPM1::Cerulean, followed by expression of mRNA for FIB1::RFP or GFP, followed by injection of various constructs (FIB1, FIB1ΔN, FIB1ΔC, NPM1, NPM1ΔN, NPM1ΔC), and final image is the overlay of all three channels. Scale bar, 10 μm. The right most panel shows in vitro images of FIB1 or mutants (green) mixed with NPM1 droplets (red) or NPM1 or mutants (red) mixed with FIB1 droplets (green). Scale bar, 10 μm. See also Figure S5.

We next probed the importance of the three domains of NPM1: an N-terminal oligomerization domain (OD) which has been shown to be necessary to form an ordered pentameric structure (Mitrea et al., 2014), a central disordered domain containing acidic tracts (A2/A3), and a C-terminal RNA binding domain (RRM) (Figure 5B). Furthermore, the OD of NPM1 can form a pentamer to generate multivalency and could potentially increase the affinity of its RRM domain for rRNA. We deleted the N-terminal oligomerization domain to create NPM1 Δ N, and we deleted the C-terminal RNA binding domain to create NPM1ΔC. We find that neither mutant is able to form droplets in vitro, consistent with phase separation of NPM1 requiring the oligomerization of NPM1 into multivalent pentamers that can bind to rRNA (Mitrea et al., 2016).

When NPM1 \Delta N is injected into X. laevis nuclei, we find that it localizes only very weakly to the nucleolus (Figure 5D). This is consistent with the finding that pentameric state of NPM1 is necessary to retain this protein in the nucleolus (Mitrea et al., 2016). When NPM1ΔN is mixed with FIB1 droplets in vitro, we see strong co-localization of NPM1 Δ N within FIB1 droplets (Figures 5D and S5A). To determine whether this strong co-localization is associated with the presence of rRNA in FIB1 droplets, we used poly-U50 to drive the phase-separation of FIB1. Interestingly, we find that the localization of NPM1ΔN within FIB1 droplets is reduced significantly with poly-U50 (Figure S5B); this suggests that the NPM1 \Delta N can localize within FIB1 droplets through its RRM interacting with rRNA. When NPM1ΔC is injected into X. laevis oocytes, we see that the protein strongly localizes to both the GC and DFC. However, NPM1 Δ C localizes very weakly to FIB1 droplets in vitro (Figures 5D and S5A); this in vitro colocalization does not appear to be affected by the types of RNA used, suggesting that the interaction between NPM1 Δ C and FIB1 in vivo is not driven by RNA, because NPM1ΔC lacks an RRM (Figure S5C), but rather by interactions between the

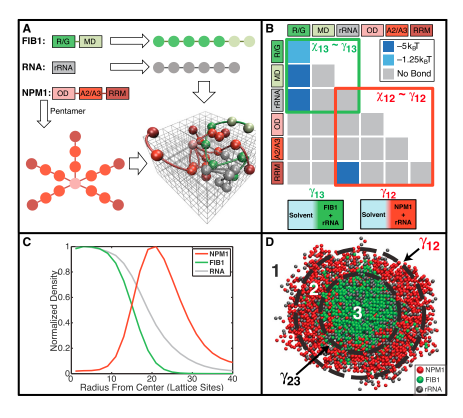


Figure 6. Preferential Interaction Model Captures the Formation of Spatially Organized Droplets for the Ternary System Comprising of FIB1, NPM1, and rRNA

(A) Mapping of the sequences of FIB1, NPM1, and rRNA to linear/branched polymers of modules on three-dimensional lattices. FIB1 is modeled as a linear polymer comprising of seven modules, five corresponding to the R/G domain and two corresponding to the MD. Similarly, the rRNA sequence is modeled as a linear polymer comprising six modules. NPM1 is modeled as a branched polymer with five arms. Here, the ODs of five NPM1 molecules occupy the base for each branch: two modules correspond to the intrinsically disordered acid-rich regions (A2/A3) and a single module captures the RNA recognition module (RRM). A representative snapshot is shown of polymers on the cubic lattice.

- (B) The matrix of module interaction strength.
- (C) The normalized mean radial density of FIB1 (green), NPM1 (red), and RNA (gray) for representative largest cluster observed throughout a simulation.
- (D) Visual depiction of a slice through representative phase separated droplet; FIB1 (green) and NPM1 (red).

See also Figure S6 and Movie S5.

R/G domain of FIB1 and the OD/A2/A3 domains of NPM1 (Mitrea et al., 2014).

In summary, our domain analysis supports three key conclusions: (1) the disordered R/G domain of FIB1 can drive phase separation, but the time-dependent viscoelastic properties of full-length FIB1 require the RNA-binding MD; (2) the disordered domains of both FIB1 and NPM1 appear capable of localizing equally to both components of nucleoli; and (3) the RNA binding domains, multivalent in the case of NPM1, play a key role in driving each protein to their respective subcompartment.

A Minimalist Computational Model for Three-Phase Behavior

Our data lead to the hypothesis that spatial organization within the nucleolus derives from the sequence-encoded interaction preferences of the different domains of nucleolar proteins. To test this hypothesis, we asked if the observed spatial organization could be reproduced in coarse-grained computer simulations. The simulation is comprised of 900 of each of the three polymers, performed on three-dimensional lattices to reduce the computational complexity. FIB1 and rRNA were modeled as linear polymers of interaction modules, while the pentameric nature of NPM1 was captured using a branched polymer with five arms, each comprising the appropriate number of interaction modules (Figure 6A).

Interactions between modules of each of the protein- and RNA-like polymers are governed by parameters of an interaction matrix. These parameters represent effective pairwise affinities in the presence of the competing effects of module-solvent and module-module interactions (Figure 6B). The interaction matrix in Figure 6B is sufficient to reproduce the totality of experimental observations. Figure 6C shows the normalized density profiles of FIB1, NPM1, and rRNA within droplets that form in the simulations, revealing a FIB1-rich core and NPM1-rich outer shell, with rRNA distributed across the two regions. Figure 6D shows a representative cutaway snapshot from the simulations (Movie S5). Numerous distinct matrix parameterizations fail to reproduce one or more aspects of the in vitro data, although there are other specific choices of matrix parameters that do reproduce all of the experimental data (Figure S6).

An exploration in the space of interaction matrix parameters suggests that the computational model must include three necessary features in order to reproduce all of the in vitro data. First, the R/G modules must have favorable homotypic interactions. Second, the OD of NPM1 should generate the requisite multivalency of RRM modules that drives the phase separation of NPM1 through interactions with rRNA. Third, the A2/A3 modules of NPM1 must be preferentially solvated, thus ensuring that they form weak or no bonds (Figures 6B and S6L).

The minimalist model supports the presence of three distinct phases: phase 1 is the solvent and includes water plus the solution ions; phase 2 is NPM1 + rRNA; and phase 3 is FIB1 + rRNA. The balance of interactions can be quantified in terms of pairwise interaction coefficients designated as χ_{12} (solvent-NPM1), χ_{13} (solvent-FIB1), and χ_{23} (NPM1-FIB1) that are derived from the Flory-Huggins theory for polymer solutions and blends (Rubinstein and Colby, 2003). The individual χ values quantify the free energy gained or lost when modules exchange homotypic

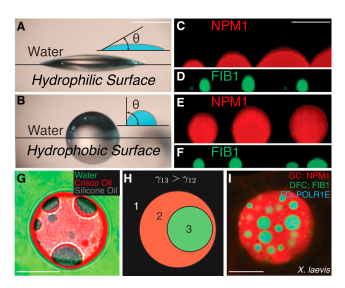


Figure 7. Surface Tension Drives Organization of Multiphase **Droplets**

(A-F) Images of droplets on hydrophilic surfaces (Pluronic-treated, A) or hydrophobic surfaces (Sigmacote-treated, B). (A) Water droplet on hydrophilic surface. (B) Water droplets on hydrophobic surface. Scale bars, 1 mm. (C) NPM1 droplets on hydrophilic surface. Scale bar, 5 μm . (D) FIB1 droplets on hydrophilic surface. (E) NPM1 droplets on hydrophobic surface. Scale bar, 5 μm. (F) FIB1 droplets on hydrophobic surface. Scale bar, 5 μm.

(G) Image of non-biological multiphase droplets: green, water; red, Crisco oil; gray, silicone oil. Scale bar, 20 µm.

(H) Schematic organization of immiscible multiphase droplets. The more hydrophobic phase (green), has a higher surface tension with water than the more hydrophilic phase (red), which has a lower surface tension with water. (I) Image of multiphase nucleoli after actin disruption in X. laevis. Scale bar, 20 μm.

See also Figure S7.

interactions for heterotypic ones. A direct consequence of the structure of the interaction matrix (Figure 6B) is that the values of each of $\chi_{12},~\chi_{13},$ and χ_{23} are positive. The three-phase behavior observed in experiments and reproduced in simulations implies that χ values must obey the relation: $\chi_{13} > \chi_{12} >$ χ_{23} > 0. Since surface tension is directly proportional to the Flory parameter, $\gamma \sim \chi$, it follows that $\gamma_{13} > \gamma_{12} > \gamma_{23}$ (i.e., the surface tension of FIB1 droplets is larger than that of NPM1 droplets). It is thus energetically more favorable to envelope the FIB1 droplet within the NPM1 droplet, as opposed to NPM1 being enveloped within a FIB1 droplet (Figure S7D).

Nucleolar Organization Arises from Differential Surface Tension of Subcompartments

The physical picture that emerges from our computational model is consistent with the very low values we obtained for the surface tension of the in vitro NPM1 droplets, as well as for the in vivo NPM1-rich GC. Unfortunately, the small size of in vitro FIB1 droplets, as well as their viscoelasticity, makes it difficult to undertake direct surface tension measurements. As an alternate route to evaluate the relative droplet surface tensions, we measured droplet wetting behavior on hydrophobic and hydrophilic coverslips. Wetting refers to the contact between liquids and surfaces and is a consequence of surface tension; for example, water droplets will spread over a Pluronic-treated hydrophilic surface (low contact angle), while water droplets will round up and avoid contact with a Sigmacote-treated hydrophobic surface (high contact angle), as shown in Figures 7A and 7B.

On hydrophobic surfaces, we find that NPM1 droplets behave as water droplets and exhibit minimal wetting, with a contact angle of $130^{\circ} \pm 10^{\circ}$ (mean \pm SD) (Figures 7E, S7A, and S7C). On hydrophilic surfaces they exhibit increased wetting, with a contact angle of $60^{\circ} \pm 10^{\circ}$ (Figures 7C, S7A, and S7C). In contrast, FIB1 droplets tended to better wet the hydrophobic coverslips, with a contact angle of $70^{\circ} \pm 10^{\circ}$ (mean \pm SD) (Figures 7F, S7A, and S7C), as compared to the hydrophilic coverslips, on which they exhibited a contact angle of $130^{\circ} \pm 10^{\circ}$ (Figures 7D, S7A, and S7C). The differential hydrophobicity of NPM1 and FIB1 droplets explains our key observation, which we describe in vivo (Figures 1A-1G), in vitro (Figure 4A), and also in silico (Figures 6C and 6D): FIB1 and NPM1 form multiphase droplets where FIB1 is at least partially encapsulated by NPM1 (Figure 7I). This organization is guite similar to how immiscible liquids are organized in non-biological multiphase systems (Neeson et al., 2012). To demonstrate this with a simple example, we prepared a system of water, Crisco vegetable oil, and silicone oil, which are immiscible liquids with known surface tensions (Than et al., 1988) (Figure 7G). Silicone oil is more hydrophobic than Crisco oil, i.e., $\gamma_{\text{silicone/water}} > \gamma_{\text{Crisco/water}},$ and as a result, the silicone oil droplets are always enveloped by the Crisco oil droplet. Similarly, since FIB1 is more hydrophobic than NPM1, $\gamma_{\text{FIB1/water}} >$ $\gamma_{NPM1/water}$, FIB1 droplets will tend to be encapsulated within NPM1 droplets (Figures 7H and S7D). We note that in both cases there is also the requirement that a third surface tension, $\gamma_{\text{silicone/Crisco}}$ or $\gamma_{\text{FIB1/NPM1}},$ must not be too high, or the two droplets would never interact (Figure S7D).

DISCUSSION

The nucleolus is the most prominent of numerous membraneless RNP bodies and was recognized over 150 years ago by early microscopists (Pederson, 2011). However, a mechanistic biophysical understanding of the principles governing the wellknown subcompartmental organization has been elusive (Thiry and Lafontaine, 2005). Our findings suggest that these distinct compartments arise as a consequence of liquid phase immiscibility, supported by: (1) the different layers can undergo coalescence upon contact, relaxing toward round droplet-like structures; (2) protein components of the different layers are highly dynamic as determined by FRAP; (3) when expressed and purified, key enriched nucleolar proteins undergo phase separation into droplets with properties comparable to those found in their corresponding nucleolar subcompartment; (4) when mixed, purified proteins exhibit phase immiscibility, resulting in subcompartmentalized in vitro droplets that are strikingly similar to in vivo nucleoli; (5) a minimalist coarse-grained model is sufficient for reproducing phase immiscibility and the observed colocalization of different truncation constructs and further shows how a network of molecular interactions determines surface tensions, which dictate the core-shell droplet architecture; and (6) biophysical characterization and droplet wetting behavior indicate that the



FIB1/DFC phase exhibits a higher surface tension than the NPM1/GC phase, which results in the former being embedded within the latter.

What are the functional implications of liquid phase immiscibility and surface-tension driven subcompartmentalization of the nucleolus? The most important nucleolar function is ribosome biogenesis (Boisvert et al., 2007), which involves the coordinated modification and assembly of rRNA through multiple processing steps (Tschochner and Hurt, 2003; Henras et al., 2008). Analogous to an assembly line or the staged processes in a modern chemical plant, the spatial separation and distinct physical and compositional features of the FIB1-rich DFC and NPM1-rich GC may tune the vectorial transport and associated processing of rRNA into mature pre-ribosomal particles. Indeed, continuous transcription within the FC causes radial flux of rRNA through the DFC into the GC and finally into the nucleoplasm. The DFC is effectively an enzymatic bath, which facilitates correct base-pairing with small nucleolar guide RNAs (snoRNAs), for example those associated with methylation by FIB1 as part of a box C/D snoRNP and pseudouridinylation by H/ACA snoRNPs, as well as cleavage reactions and other rRNA modifications (Henras et al., 2008). These modifications are critical for correct rRNA folding and stability, subsequent assembly with ribosomal proteins, and ultimately ribosome function (i.e., translational fidelity) (Gigova et al., 2014).

We propose that the viscoelastic properties of the DFC serve to lower the flux of incompletely or incorrectly processed/folded pre-ribosomal particles, ensuring that DFC-associated enzymatic processes are completed, before passage of pre-ribosomal particles into the outer GC layer, where they encounter NPM1 and early binding ribosomal proteins. Indeed, NPM1 phase separates with both rRNA and ribosomal proteins (Mitrea et al., 2016), consistent with fluorescence imaging studies suggesting that ribosomal proteins localize to the GC, but not the DFC (Krüger et al., 2007). Our findings indicate that pentameric NPM1 is integral to the fluid features of the GC, whose relatively low viscosity may allow ribosomal proteins to remain dynamically accessible to pre-ribosomal particles emerging from the DFC.

Our data show that FIB1 droplets, but not NPM1 droplets, are metastable and can age with time both in vitro and in vivo. These data are consistent with the hypothesis that disordered regions can facilitate the transition from liquid-like to solid-like structures (Weber and Brangwynne, 2012), which is supported by recent studies on a variety of RNA binding proteins (Zhang et al., 2015; Patel et al., 2015; Lin et al., 2015; Molliex et al., 2015; Xiang et al., 2015). We note that the name "Fibrillarin" was given due to its localization to fibrillar structures within the DFC (Ochs et al., 1985); these structures could reflect droplet aging/fibril formation. Consistent with this, we observe apparent aging of nucleoli in non-dividing C. elegans intestinal cells: FIB1 recovers less completely in older adult worms compared with younger larvae (Figure S7E). However, given that RNA can impact the fluidity of related phase-separated droplets (Elbaum-Garfinkle et al., 2015; Zhang et al., 2015), the rate of FIB1-rich DFC aging could be impacted by the flux of newly synthesized rRNA transiting through the DFC. Future studies will be required to elucidate the biophysical origin of nucleolar droplet maturation

and the role of RNA transcription and other ATP-dependent processes.

Our findings underscore the importance of surface tension, whose role is well established in physical systems, as is readily visualized with immiscible oils in water (Figure 7G). In living systems, effective surface tension may be important for the organized demixing of cell populations: cell types that exhibit a relatively high apparent surface tension will tend to be enveloped by cell types with a relatively low apparent surface tension (Foty et al., 1996). Our data show that this same basic principle is important for structuring the nucleolus, with possible implications for other RNP bodies. For example, histone locus bodies (HLBs) in the frog nucleus are commonly found to have B-snurposomes attached to their surface (Gall, 2000); incomplete internalization of B-snurposomes suggests that their surface tension may be similar to HLBs. Interestingly, this partial internalization is reminiscent of the altered nucleolar structure observed in actinomycin-D-treated nucleoli, wherein rRNA transcriptional inhibition results in a more lobulated nucleolus (Shav-Tal et al., 2005; Wachtler and Stahl, 1993). A similar organization is also seen with processing bodies (Kedersha et al., 2005; Buchan and Parker, 2009; Hubstenberger et al., 2013). Recently, stress granules have been shown to contain less dynamic cores, which exhibit a qualitative similarity to the FIB1/DFC cores of the nucleolus (Jain et al., 2016). Building on the biophysical groundwork we have laid here, it may be possible to alter or even invert the organization of such RNP bodies, by using surfactants to modulate droplet surface tensions; this could significantly impact sequential RNA processing steps and the overall flow of genetic information.

Organelle subcompartmentalization is well-known in membrane-bound organelles, such as mitochondria. Our data show that membrane-less liquid phase organelles can also generate significant substructure. Phase separation and the coexistence of multiple distinct liquid RNA/protein phases thus provide a simple but elegant mechanism for the cell to control the spatial localization and processing of molecules, without relying on membrane boundaries.

EXPERIMENTAL PROCEDURES

Detailed methods are available in the Supplemental Experimental Procedures.

Preparation of X. laevis, Mammalian, and C. elegans Nucleoli

Frogs were anesthetized with 0.1% MS-222 solution, and oocytes were surgically removed from female *X. laevis* frogs following an IACUC approved protocol. mRNA of endogenous proteins (FIB1, NPM1, and POLR1E) with fluorescent tags and recombinant proteins were microinjected into oocytes. Nuclei were manually dissected in mineral oil and subsequently imaged. Actin wals disrupted using Lat-A, and ATP was depleted using Apyrase. Mammalian cells expressing fluorescent fusion proteins (FIB1 and NPM1) were maintained at 37°C using standard conditions, and ATP was depleted using sodium azide and deoxyglucose. *C. elegans* expressing intestinal FIB1::GFP were maintained at 20°C under standard conditions and anesthetized with levamisole in M9 prior to imaging.

Purification and Phase Separation of In Vitro Droplets

FIB1 and NPM1 variants were expressed using a standard *E. coli* expression system, purified using either a 6x-His or GST tag, and stored in a high salt buffer. Phase separation was initiated by lowering the salt concentration of



stock protein in the presence or absence of rRNA. Non-biological multiphase droplets were obtained by vortexing water, Crisco oil, and silicone oil.

Biophysical Characterization of In Vivo and In Vitro Droplets

For fusion relaxation experiments, the aspect ratio was measured as a function of time for droplets of different sizes to obtain the inverse capillary velocity. Surface tension of non-wetting droplets with measured densities was estimated from non-spherical XZ shape profiles obtained using a right-angle prism. For microrheology experiments, time-lapse images of fluctuating R = 50 nm particles inside protein droplets were acquired and analyzed using particle-tracking MATLAB code to obtain the mean squared displacement as a function of lag time; from the Stokes-Einstein relation, the viscosity was determined. For FRAP experiments, 1 μm spots inside in vivo and in vitro droplets were photobleached, and percent fluorescent recovery and recovery times were determined using standard techniques. Wetting behavior of in vitro droplets was observed for surfaces treated with Sigmacote (hydrophobic) or Pluronic (hydrophilic), and the contact angle was measured at the interface between the glass and line tangent to the droplet.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, seven figures, and five movies and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2016.04.047.

AUTHOR CONTRIBUTIONS

M.F. performed and analyzed data in X. laevis, in vitro surface tension, microrheology experiments, a non-biological multiphase system, and contact angle experiments. N.V. performed and analyzed in vitro experiments with FIB1 and NPM1 and FRAP experiments in C. elegans. T.S.H. performed computational simulations and analysis. D.M.M. created in vitro NPM1 constructs and created NPM1 phase diagrams. L.Z. analyzed experiments in mammalian tissue culture cells. T.M.R. performed and analyzed FRAP experiments in X. laevis with help from M.F. and ATP depletion in mammalian tissue culture cells with help from L.Z. C.P.B, M.F., and N.V. designed the study and wrote the paper with help from all authors.

ACKNOWLEDGMENTS

We thank members of the C.P.B. lab for discussions and Adrienne Fung for preliminary work on surface tension measurements. We acknowledge funding from the Princeton Center for Complex Materials, a MRSEC supported by NSF grant DMR 1420541. This work was also supported by an NIH Director's New Innovator Award (1DP2GM105437-01 to C.P.B.), an NSF CAREER award (1253035 to C.P.B.), a Helen Hay Whitney Fellowship (to N.V.), an NSF grant (MCB 1121867 to R.V.P.), NIH grants (5RO1NS056114 to R.V.P. and 5R01GM115634 to R.W.K.), an NCI Cancer Center Support grant (P30CA21765 at St. Jude Children's Research Hospital to R.W.K.), and ALSAC (R.W.K.).

Received: February 1, 2016 Revised: March 11, 2016 Accepted: April 13, 2016 Published: May 19, 2016

REFERENCES

Aarts, D.G., Schmidt, M., and Lekkerkerker, H.N. (2004). Direct visual observation of thermal capillary waves. Science 304, 847-850.

Balagopal, V., and Parker, R. (2009), Polysomes, P bodies and stress granules: states and fates of eukaryotic mRNAs. Curr. Opin. Cell Biol. 21, 403-408.

Berry, J., Weber, S.C., Vaidya, N., Haataja, M., and Brangwynne, C.P. (2015). RNA transcription modulates phase transition-driven nuclear body assembly. Proc. Natl. Acad. Sci. USA 112, E5237-E5245.

Boisvert, F.-M., van Koningsbruggen, S., Navascués, J., and Lamond, A.I. (2007). The multifunctional nucleolus. Nat. Rev. Mol. Cell Biol. 8, 574-585.

Brangwynne, C.P., Eckmann, C.R., Courson, D.S., Rybarska, A., Hoege, C., Gharakhani, J., Jülicher, F., and Hyman, A.A. (2009). Germline P granules are liquid droplets that localize by controlled dissolution/condensation. Science 324, 1729-1732.

Brangwynne, C.P., Mitchison, T.J., and Hyman, A.A. (2011). Active liquid-like behavior of nucleoli determines their size and shape in Xenopus laevis oocytes. Proc. Natl. Acad. Sci. USA 108, 4334-4339.

Buchan, J.R., and Parker, R. (2009). Eukaryotic stress granules: the ins and outs of translation. Mol. Cell 36, 932-941.

Derenzini, M., Trerè, D., Pession, A., Govoni, M., Sirri, V., and Chieco, P. (2000). Nucleolar size indicates the rapidity of cell proliferation in cancer tissues. J. Pathol. 191, 181-186.

Elbaum-Garfinkle, S., Kim, Y., Szczepaniak, K., Chen, C.C.-H., Eckmann, C.R., Myong, S., and Brangwynne, C.P. (2015). The disordered P granule protein LAF-1 drives phase separation into droplets with tunable viscosity and dynamics. Proc. Natl. Acad. Sci. USA 112, 7189-7194.

Feric, M., and Brangwynne, C.P. (2013). A nuclear F-actin scaffold stabilizes ribonucleoprotein droplets against gravity in large cells. Nat. Cell Biol. 15, 1253-1259

Feric, M., Broedersz, C.P., and Brangwynne, C.P. (2015). Soft viscoelastic properties of nuclear actin age oocytes due to gravitational creep. Sci. Rep. 5. 16607.

Foty, R.A., Pfleger, C.M., Forgacs, G., and Steinberg, M.S. (1996). Surface tensions of embryonic tissues predict their mutual envelopment behavior. Development 122, 1611-1620.

Frank, D.J., and Roth, M.B. (1998). ncl-1 is required for the regulation of cell size and ribosomal RNA synthesis in Caenorhabditis elegans. J. Cell Biol. 140, 1321-1329.

Gall, J.G. (2000). Cajal bodies: the first 100 years. Annu. Rev. Cell Dev. Biol. 16, 273-300.

Gigova, A., Duggimpudi, S., Pollex, T., Schaefer, M., and Koš, M. (2014). A cluster of methylations in the domain IV of 25S rRNA is required for ribosome stability, RNA 20, 1632-1644.

Hennig, S., Kong, G., Mannen, T., Sadowska, A., Kobelke, S., Blythe, A., Knott, G.J., Iyer, K.S., Ho, D., Newcombe, E.A., et al. (2015). Prion-like domains in RNA binding proteins are essential for building subnuclear paraspeckles. J. Cell Biol. 210, 529-539.

Henras, A.K., Soudet, J., Gérus, M., Lebaron, S., Caizergues-Ferrer, M., Mougin, A., and Henry, Y. (2008). The post-transcriptional steps of eukaryotic ribosome biogenesis. Cell. Mol. Life Sci. 65, 2334–2359.

Hubstenberger, A., Noble, S.L., Cameron, C., and Evans, T.C. (2013). Translation repressors, an RNA helicase, and developmental cues control RNP phase transitions during early development. Dev. Cell 27, 161-173.

Jain, S., Wheeler, J.R., Walters, R.W., Agrawal, A., Barsic, A., and Parker, R. (2016). ATPase-modulated stress granules contain a diverse proteome and substructure. Cell 164, 487-498.

Kedersha, N., Stoecklin, G., Ayodele, M., Yacono, P., Lykke-Andersen, J., Fritzler, M.J., Scheuner, D., Kaufman, R.J., Golan, D.E., and Anderson, P. (2005). Stress granules and processing bodies are dynamically linked sites of mRNP remodeling. J. Cell Biol. 169, 871-884.

Kroschwald, S., Maharana, S., Mateju, D., Malinovska, L., Nüske, E., Poser, I., Richter, D., and Alberti, S. (2015). Promiscuous interactions and protein disaggregases determine the material state of stress-inducible RNP granules. eLife 4. e06807.

Krüger, T., Zentgraf, H., and Scheer, U. (2007). Intranucleolar sites of ribosome biogenesis defined by the localization of early binding ribosomal proteins. J. Cell Biol. 177, 573-578.

Li, P., Banjade, S., Cheng, H.-C., Kim, S., Chen, B., Guo, L., Llaguno, M., Hollingsworth, J.V., King, D.S., Banani, S.F., et al. (2012). Phase transitions in the assembly of multivalent signalling proteins. Nature 483, 336-340.



Lin, Y., Protter, D.S., Rosen, M.K., and Parker, R. (2015). Formation and maturation of phase-separated liquid droplets by RNA-binding proteins. Mol. Cell 60. 208-219.

Mitrea, D.M., Grace, C.R., Buljan, M., Yun, M.-K., Pytel, N.J., Satumba, J., Nourse, A., Park, C.-G., Madan Babu, M., White, S.W., and Kriwacki, R.W. (2014). Structural polymorphism in the N-terminal oligomerization domain of NPM1. Proc. Natl. Acad. Sci. USA 111, 4466-4471.

Mitrea, D.M., Cika, J.A., Guy, C.S., Ban, D., Banerjee, P.R., Stanley, C.B., Nourse, A., Deniz, A.A., and Kriwacki, R.W. (2016). Nucleophosmin integrates within the nucleolus via multi-modal interactions with proteins displaying R-rich linear motifs and rRNA. eLife 5, e13571.

Molliex, A., Temirov, J., Lee, J., Coughlin, M., Kanagaraj, A.P., Kim, H.J., Mittag, T., and Taylor, J.P. (2015). Phase separation by low complexity domains promotes stress granule assembly and drives pathological fibrillization. Cell 163. 123-133.

Neeson, M.J., Tabor, R.F., Grieser, F., Dagastine, R.R., and Chan, D.Y. (2012). Compound sessile drops. Soft Matter 8, 11042-11050.

Nott, T.J., Petsalaki, E., Farber, P., Jervis, D., Fussner, E., Plochowietz, A., Craggs, T.D., Bazett-Jones, D.P., Pawson, T., Forman-Kay, J.D., and Baldwin, A.J. (2015). Phase transition of a disordered nuage protein generates environmentally responsive membraneless organelles. Mol. Cell 57, 936-947.

Ochs, R.L., Lischwe, M.A., Spohn, W.H., and Busch, H. (1985). Fibrillarin: a new protein of the nucleolus identified by autoimmune sera. Biol. Cell 54, 123-133.

Parry, B.R., Surovtsev, I.V., Cabeen, M.T., O'Hern, C.S., Dufresne, E.R., and Jacobs-Wagner, C. (2014). The bacterial cytoplasm has glass-like properties and is fluidized by metabolic activity. Cell 156, 183-194.

Patel, A., Lee, H.O., Jawerth, L., Maharana, S., Jahnel, M., Hein, M.Y., Stoynov, S., Mahamid, J., Saha, S., Franzmann, T.M., et al. (2015). A liquid-to-solid phase transition of the ALS protein FUS accelerated by disease mutation. Cell 162, 1066-1077.

Pederson, T. (2011). The nucleolus. Cold Spring Harb. Perspect. Biol. 3, a000638.

Rubinstein, M., and Colby, R.H. (2003). Polymer Physics (Oxford University

Shav-Tal, Y., Blechman, J., Darzacq, X., Montagna, C., Dye, B.T., Patton, J.G., Singer, R.H., and Zipori, D. (2005). Dynamic sorting of nuclear components into distinct nucleolar caps during transcriptional inhibition. Mol. Biol. Cell 16, 2395-2413.

Spector, D.L. (2001). Nuclear domains. J. Cell Sci. 114, 2891–2893.

Than, P., Preziosi, L., Josephl, D., and Arney, M. (1988). Measurement of interfacial tension between immiscible liquids with the spinning road tensiometer. J. Colloid Interface Sci. 124, 552-559.

Thiry, M., and Lafontaine, D.L. (2005). Birth of a nucleolus: the evolution of nucleolar compartments. Trends Cell Biol. 15, 194-199.

Tschochner, H., and Hurt, E. (2003). Pre-ribosomes on the road from the nucleolus to the cytoplasm. Trends Cell Biol. 13, 255-263.

Wachtler, F., and Stahl, A. (1993). The nucleolus: a structural and functional interpretation. Micron 24, 473-505.

Wang, J.T., Smith, J., Chen, B.-C., Schmidt, H., Rasoloson, D., Paix, A., Lambrus, B.G., Calidas, D., Betzig, E., and Seydoux, G. (2014). Regulation of RNA granule dynamics by phosphorylation of serine-rich, intrinsically disordered proteins in C. elegans. eLife 3, e04591.

Weber, S.C., and Brangwynne, C.P. (2012). Getting RNA and protein in phase. Cell 149, 1188-1191.

Weber, S.C., and Brangwynne, C.P. (2015). Inverse size scaling of the nucleolus by a concentration-dependent phase transition. Curr. Biol. 25, 641-646.

Wippich, F., Bodenmiller, B., Trajkovska, M.G., Wanka, S., Aebersold, R., and Pelkmans, L. (2013). Dual specificity kinase DYRK3 couples stress granule condensation/dissolution to mTORC1 signaling. Cell 152, 791-805.

Xiang, S., Kato, M., Wu, L.C., Lin, Y., Ding, M., Zhang, Y., Yu, Y., and McKnight, S.L. (2015). The LC domain of hnRNPA2 adopts similar conformations in hydrogel polymers, liquid-like droplets, and nuclei. Cell 163, 829-839.

Zhang, H., Elbaum-Garfinkle, S., Langdon, E.M., Taylor, N., Occhipinti, P., Bridges, A.A., Brangwynne, C.P., and Gladfelter, A.S. (2015). RNA controls PolyQ protein phase transitions. Mol. Cell 60, 220-230.