Distinct RNA polymerase transcripts direct the assembly of phase-separated DBC1 nuclear bodies in different cell lines

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ABSTRACT The mammalian cell nucleus is a highly organized organelle that contains membrane-less structures referred to as nuclear bodies (NBs). Some NBs carry specific RNA types that play architectural roles in their formation. Here, we show two types of RNase-sensitive DBC1-containing NBs, DBC1 nuclear body (DNB) in HCT116 cells and Sam68 nuclear body (SNB) in HEK cells, that exhibit phase-separated features and are constructed using RNA polymerase I or II transcripts in a cell type–specific manner. We identified additional protein components present in DNB by immunoprecipitation–mass spectrometry, some of which (DBC1 and heterogeneous nuclear ribonucleoprotein L [HNRNPL]) are required for DNB formation. The rescue experiment using the truncated HNRNPL mutants revealed that two RNA-binding domains and intrinsically disordered regions of HNRNPL play significant roles in DNB formation. All these domains of HNRNPL promote in vitro droplet formation, suggesting the need for multivalent interactions between HNRNPL and RNA as well as proteins in DNB formation.

INTRODUCTION Nuclei of higher eukaryotic cells are highly structured and possess multiple types of nuclear compartments called nuclear bodies (NBs). NBs contain various types of proteins and RNAs, and most of them function as the sites of synthesis, storage, and sequestration of specific RNAs, proteins, and ribonucleoprotein (RNP) complexes. In general, NBs likely serve to concentrate various regulatory factors that enhance specific biochemical reactions taking place in the NBs or suppress their activity out of the NBs (Staněk and Fox, 2017). For example, the nucleolus serves as the platform for both RNA polymerase I (RNAPI) transcription and ribosome biogenesis (Iarovaia et al., 2019). Paraspeckles regulate gene expression through sequestration of the proteins, which further results in suppression of the activity of transcriptional regulation proteins (Prasanth et al., 2005; Hirose et al., 2014; Imamura et al., 2014). The nuclear stress bodies (nSBs) regulate pre-mRNA splicing through an efficient phosphorylation of the sequestered serine- and arginine-rich pre-mRNA splicing factors (SRSFs) by CLK1, which is recruited during thermal stress recovery (Ninomiya et al., 2020). Recent studies indicated that the intracellular liquid demixing (i.e., liquid–liquid phase separation), induced by multivalent interactions between intrinsically disordered regions (IDRs) in various RNA-binding proteins (RBPs), promotes the assembly of liquid droplet-like membrane-less organelles (Banani et al., 2017; Uversky, 2017; Fox et al., 2018). In some cases, RNA serves as a scaffold of these organelles in a way to interact with multiple RBPs that further promote multivalent interactions between the IDRs (Lin et al., 2015; Shin and Brangwynne, 2017). It has been proposed that a class of long noncoding RNAs...
(lncRNAs) functions as the scaffold of specific NBs, thereby termed as architectural RNAs (arcRNAs) (Clemson et al., 2009; Sasaki et al., 2009; Sunwoo et al., 2009; Chujo et al., 2016; Yamazaki et al., 2019). For example, NEAT1 arcRNA plays an essential role in paraspeckle assembly by the interaction with IDR-containing RBPs such as NONO and SFPQ (Sasaki et al., 2009; Nagaruma et al., 2012; Henning et al., 2015; Yamazaki et al., 2018). The nSBs are produced on H5ATIII arcRNAs that recruit distinct sets of RBPs such as SRSF splicing regulators (Ninomiya et al., 2020; Ninomiya and Hirose, 2020).

In our previous study, by screening of RNase-sensitive NBs, we have searched for novel NBs that are built on unidentified arcRNAs using the fluorescently tagged human cDNA library (Mannen et al., 2016). Among the identified RNase-sensitive NBs, we found that Sam68 nuclear bodies (SNBs), which were detected at perinucleolar sites requiring RNA polymerase II (RNAPII) transcripts for their formation, are composed of two distinct RNase-sensitive substructures, Sam68 and DBC1, which are connected by the adaptor protein heterogeneous nuclear ribonucleoprotein L (HNRNPL) in HeLa cells. On the other hand, HCT116 cells form different RNase-sensitive NBs, which include DBC1 but not Sam68; therefore, we termed them DBC1 nuclear bodies (DNBs) (Mannen et al., 2016). Except for DBC1, other RNA and protein components that constitute DNBs remain still unidentified.

In this study, we aimed to show that the DNB is an RNase-sensitive and liquid droplet-like NB that requires RNAPI transcripts. As the first step, we planned to identify the remaining protein components of DNB by immunoprecipitation–mass spectrometry (IP-MS) and to characterize each protein component, bearing in mind that DBC1 and HNRNPL are essential for DNB formation. A detailed domain dissection was predicted in order to examine whether the RNA-binding domains of HNRNPL, as well as the intrinsically disordered proline-rich domain (PR), can play significant roles in DNB formation and in vitro droplet formation.

RESULTS

Formation of DBC1 nuclear bodies requires both RNA and phase-separated structures

In HeLa cells, the SNBs are composed of both the Sam68 and DBC1 substructures, which are combined with a HNRNPL adaptor protein, whereas the DBC1 focal signals were detected as distinct foci lacking the Sam68 signals in HCT116 and NIH3T3 cells (Supplemental Figure S1A). The DNBs were unexceptionally detected as a single focus in HCT116 and NIH3T3 cells, whereas SNBs were detected as two foci in HeLa cells (Supplemental Figure S1B). We confirmed that the expression levels of the DBC1 and the Sam68 proteins were almost equivalent in these two cell lines (Supplemental Figure S1C), suggesting the presence of cell type–specific factor(s) in the formation of the SNB and the DNB. The DNBs found in HCT116 and NIH3T3 cells were sensitive to the RNase treatment, which was also observed for the DBC1 substructure in HeLa cells (Supplemental Figure S1D). In addition, we showed that both the DBC1 and the Sam68 substructures in HeLa cells were rapidly dispersed upon inhibition of RNAPII with 5,6-dichloro-1-β-D-ribofuranosylbenzimidazole (DRB) or a high-dose (0.3 μM) of actinomycin D (Act D) (Figure 1A and Supplemental Figure S1E), indicating the necessity for RNAPII transcript(s) for the formation of both substructures (Chen et al., 1999; Mannen et al., 2016). We also found that these two substructures were poorly sensitive to the treatment with CX5461 or a low-dose (0.03 μM) of Act D as RNAPII inhibitors in HeLa cells (Figure 1A and Supplemental Figure S1E). Unexpectedly, in HCT116 and NIH3T3 cells the DNBs were found to be highly sensitive to the treatment with CX5461 or a low-dose of Act D, but not to DRB (Figure 1A), suggesting the requirement for RNAPI transcripts for the DNB formation. RNAPI transcribes the ribosomal DNA unit, which produces a large precursor (47S) consisting of the 18S, 5.8S, and 28S rRNAs and intergenic spacer (IGS) regions (McStay and Grummt, 2008) (Supplemental Figure S2A). In response to various environmental stimuli, IGS regions produce lncRNAs called IGS RNAs, which sequester specific sets of proteins to assemble distinct phase-separated NBs called a nucleolar detention center (Audas et al., 2012; Jacob et al., 2013). We attempted to detect various IGS RNAs in DNBs; however, they were detectable only at perinucleolar sites, distinct from DNBs (Supplemental Figure S2B), suggesting that the IGS RNAs are neither localized in DNBs nor involved in DNB formation. These findings suggest that the DNBs in both HCT116 and NIH3T3 cells and the DBC1 substructures in SNBs of HeLa cells are likely different structures and may require distinct transcripts synthesized by other types of RNA polymerases.

Recent studies demonstrated that some NBs were in fact phase-separated ribonucleoprotein condensates formed by a multivalent interaction network of the NB components (Boeynaems et al., 2018). To investigate whether DNBs and SNBs exhibit phase-separated features, HCT116 and HeLa cells were treated with 1,6-hexanediol (1,6-HD), which reportedly disintegrates the subsets of phase-separated subcellular structures in vivo by disrupting their multivalent hydrophobic interactions (Lin et al., 2016; Cho et al., 2018; Chong et al., 2018; Sabari et al., 2018; Yamazaki et al., 2018). As shown in Figure 1B, the treatment with 1,6-HD in concentrations of 5% or 7.5% disrupted DNBs or SNBs, respectively. We also confirmed that Cajal bodies, which are phase-separated cellular structures detected with colin (COIL) (Lin et al., 2016), were also readily disintegrated in the same conditions (Figure 1B). On the other hand, 2,5-hexanediol (2,5-HD), which is known to have lower activity on the phase-separated structures, hardly affected the integrity of DNBs, SNBs, and Cajal bodies (Figure 1B). These findings suggest that DNBs in HCT116 cells and SNBs in HeLa cells exhibit characteristics similar to those of the other known phase-separated NBs.

HNRNPL and HNRNPK are novel DNB components

To gain insight into the assembly and maintenance of the DNB, we attempted to isolate the native complexes of DBC1 by IP. First, we established HCT116 cell lines stably expressing doxycycline-inducible DBC1-3xFLAG (HCT116/TR_DBC1-3xFLAG cells; Supplemental Figure S3A) and confirmed that the expressed DBC1-3xFLAG protein is promptly localized in the endogenous DNBs (Supplemental Figure S3B). To identify the proteins that interact with DBC1, an IP assay with anti-FLAG antibody was carried out using the DBC1-3xFLAG cell extracts either with or without RNase A treatment. This was followed by the analysis of the precipitated proteins with MS (Figure 2, A and B). The MS data showed that DBC1 coprecipitated with the mitochondrial chaperon HSPD1, HNRNPs HNRNPL, HNRNPK, PTBP1, and HNRNPA2B1, as well as with cytoplasmic poly(A)-binding protein PABPC1. Among these proteins, the coprecipitation of HNRNPL, HNRNPK, PTBP1, HNRNPA2B1, and PABPC1 was significantly decreased by the RNase A treatment (Figure 2B). We also confirmed both coprecipitation and RNase sensitivity of these proteins by Western blotting (Figure 2C and Supplemental Figure S4A). Among the coprecipitated proteins, we previously reported that PTBP1, which is known to be a marker of a distinct NB called the perinucleolar compartment, did not colocalize with DNBs in HCT116 cells (Mannen et al., 2016). To confirm the localization of the other coprecipitated proteins in DNBs, we performed an immunofluorescence (IF) assay using the antibodies against the stated proteins (Figure 2D and Supplemental Figure S4B). Both HNRNPL...
and HNRNPK, but not HSPD1 and HNRNPA2B1, were clearly detected to be localized in DNB. The focal signals of all three DNB proteins (DBC1, HNRNPK, and HNRNPL) synchronously disappeared upon treatment with CX5461 in both HCT116 and NIH3T3 cells (Supplemental Figure S5, A and B). These data strongly support HNRNPL and HNRNPK being the additional DNB components in HCT116 and NIH3T3 cells.

DBC1 and HNRNPL are required for DNB formation

To investigate the mechanism of DNB assembly with the identified components, we performed a reciprocal depletion of each of the three DNB components (DBC1, HNRNPL, and HNRNPK) in HCT116 and NIH3T3 cells by an RNA interference (RNAi) approach (Figure 3, A and C). IF of the DNB components in the siRNA-treated cells revealed that the depletion of either DBC1 or HNRNPL resulted in DNBs’ disappearance (see siDBC1 and siHNRNPL in Figure 3, B and D), while the depletion of HNRNPK hardly affected the DNBs’ integrity (see siHNRNPK in Figure 3, B and D). These results indicate that DBC1 and HNRNPL are both required for the formation and maintenance of the DNB structure in HCT116 and NIH3T3 cells, although we cannot rule out the possibility that additional unidentified core factors are present in DNBs.

RRMs and PR of HNRNPL are both required for DNB formation

We previously showed that the S1L domain in the N-terminal region of DBC1 is required for its localization and RNA binding to the DNB in HCT116 cells (Mannen et al., 2016). Here, we attempted to identify the functional domains of HNRNPL for DNB formation. We constructed a series of deletion mutants lacking the annotated domains in FLAG-tagged HNRNPL (Figure 4A), and the localization of each mutant was monitored by the anti-FLAG IF. We observed that the cell population in which DNBs with the FLAG signals were detected was markedly decreased in ΔRRM1, ΔRRM2, and ΔPR mutants (Supplemental Figure S6). In particular, ΔRRM2 was diffusely distributed throughout the nucleoplasm, while its small fraction was still detectable in the DNB (Supplemental Figure S6). We further investigated the rescue activity of each HNRNPL mutant for DNB formation in the HNRNPL-depleted cells. As shown in Figure 4B, WT, ΔGR, ΔRRM3, and ΔRRM4 had rescue activities that led to DNB formation in ∼40% of cells; however, ΔRRM1 and ΔRRM2 attenuated the rescue activity to <20%. A co-IP assay of DBC1 with the FLAG-HNRNPL mutants revealed that neither ΔRRM1 nor ΔRRM2 of HNRNPL succeeded in interacting with DBC1 (Figure 4C), suggesting that the HNRNPL-DBC1 interaction via either RRM1 or RRM2 is required for the rescue activity of HNRNPL during DNB formation.

Meanwhile, ΔPR almost completely abolished rescue ability (Figure 4B). In contrast to ΔRRM1 and ΔRRM2, ΔPR was able to localize itself in DNBs (Supplemental Figure S6) and subsequently interacted with DBC1 but lacked rescue activity (Figure 4C). The PR has been reported as a part of the multiple IDR, and as an intriguing...
RRMs and PR of HNRNPL promote in vitro droplet formation

Recent studies revealed that some NBs exhibited phase-separated liquid-like properties that are likely caused by demixing induced by the IDRs present in the RBP components (Banani et al., 2017; Uversky, 2017; Fox et al., 2018). To investigate whether the HNRNPL contributes to phase separation, we performed in vitro analysis using the recombinant HNRNPL protein fused with a maltose-binding protein (MBP-HNRNPL WT, a series of deletion mutants lacking the annotated domains, and PRmut3) (Supplemental Figure S8A). DNB concentrations of HNRNPL were estimated at 13.7 μM, and the in vitro protein concentration of HNRNPL was 3.3 μM (see Materials and Methods), suggesting that the HNRNPL concentration used in our in vitro experiment was within the level of HNRNPL in DNBS. We examined whether the TEV-cleaved HNRNPL proteins could form droplets under different salt conditions. The results were recorded by three different methods including sedimentation assay, turbidity assay, and microscopic observation (Supplemental Figure S8, B and C). In a sedimentation assay, TEV-cleaved HNRNPL WT, ΔGR, and PRmut3, was significantly decreased in the supernatant by reducing the NaCl concentrations, and ∆RRM3 and ∆RRM4 were nearly absent in the supernatant after centrifugation even at 150 mM NaCl, whereas ∆RRM1 and ∆RRM2 levels were unchanged by NaCl concentration and ∆PR declined slightly in 50 mM NaCl (Figure 5A) and Supplemental Figure S8D). On the other hand, TEV-cleaved MBP was unaffected by different concentrations of NaCl (Figure 5A and Supplemental Figure S8D). In a turbidity assay, the turbidity of HNRNPL WT, ΔGR, ∆RRM3, ∆RRM4, and PRmut3 was decreased upon increasing salt concentration, while that of ∆RRM1, ∆RRM2, and ∆PR was significantly decreased in comparison to WT (Figure 5B and Supplemental Figure S8E). We also observed the turbid solutions of HNRNPL protein using microscopy (Figure 5C and Supplemental Figure S8F). HNRNPL WT, ΔGR, ∆RRM3, ∆RRM4, and PRmut3 formed dynamic liquid droplets, which increased in number by decreasing salt concentration (50–150 mM NaCl), whereas ∆RRM1, ∆RRM2, and ∆PR formed liquid droplets in 50 mM NaCl but failed in higher salt concentrations. In ∆RRM2, the formation of aggregates was detected at each concentration (Figure 5C), suggesting that this causes the reduction of the HNRNPL band in the sedimentation assay regardless of the salt concentration (Figure 5A). These results revealed that RRM1, RRM2, and PR of HNRNPL promote droplet formation in vitro.

Thus, our results indicate that the HNRNPL RRM1, RRM2, and PR contribute to both DNB assembly in vivo and droplet formation in vitro. These data suggest that the HNRNPL RRM1, RRM2, and PR-mediated phase separation would be, at least in part, the driving force of DNB assembly (Figure 6B).

DISCUSSION

In this study, we demonstrated that each of the two types of DBC1-containing NBs, DNB in HCT116 and NIH3T3 cells and SNB in HeLa cells, requires distinct RNA polymerase transcripts for its formation and maintenance. Considering the RNase-sensitive features of both DNB and SNB, these putative transcripts likely act as arcRNAs of the NBs (Figure 6A). Because the IncRNA expression tends to be cell type specific (Cabili et al., 2011; Djebali et al., 2012), cell type-specific
Identification of the essential DNB components. Three DNB components were reciprocally knocked down by RNAi in HCT116 and NIH3T3 cells. Both efficacy and specificity of knockdowns of each protein by RNAi were detected by Western blotting (A: HCT116 cells, C: NIH3T3 cells). The molecular mass marker (kDa) is shown on the left. Each protein was detected by IF (B: HCT116 cells, D: NIH3T3 cells). Arrowheads indicate DNBs. The cell populations (%) in which each protein signals is observed are shown in B and D (100 cells, n = 5). Bar, 10 μm.

Either DBC1 or HNRNPL knockdown resulted in disappearance of DNBs in HCT116 and NIH3T3 cells (Figure 3). In HeLa cells, on the other hand, DBC1 knockdown did not affect either the integrity of SNB or SNB localization of HNRNPL. Besides, HNRNPL knockdown resulted in the separation of SNBs into the Sam68 substructure and the DBC1 substructure (Mannen et al., 2016). These results suggest that HNRNPL plays different roles in the formation of DNB compared with the formation of the DBC1 substructure in SNB (Figure 6A). The HNRNPL mutant lacking either RRMI or RRMR lost both the rescue activity for DNB formation and the interaction with DBC1 (Figure 4, B and C), suggesting that RRMI and RRMR cooperatively contribute to form the DNB through the interaction either with DBC1 or with arcRNA of the DNB (Figure 6B). On the other hand, HNRNPL mutants such as the ΔPR mutant and the PR substitution mutants (PRmut1-3) mostly lacked the rescue activity for DNB formation (Figure 4B), suggesting that PR of HNRNPL is required for DNB formation (Figure 6B). In addition, in vitro analysis of HNRNPL's RRMI, RRMR, and PR indicated the likelihood of droplet formation through a homomeric HNRNPL interaction, whereas PR substitution mutants (PRmut3) were unaffected for this ability (Figure 5). These results point out that the RRMI, RRMR, and PR of HNRNPL possibly contribute to the assembly of phase-separated DNBs through either a homomeric interaction with HNRNPL or a heteromeric interaction between HNRNPL and other DNB components. Our observation on the 1,6-HD sensitivity behavior of DNB also supports these arguments.

The biological roles of DNB remain elusive. It was reported that DBC1 directly interacts with SIRT1 to inhibit SIRT1 activity (Kim et al., 2008; Zhao et al., 2008). Indeed, our co-IP experiment detected the interaction of DBC1 with SIRT1; however, we failed to detect localization of SIRT1 in DNBs (Supplemental Figure S4, C and D), suggesting that DBC1 interacts with SIRT1 out of DNB and that DNB is not involved in the regulation of SIRT1 function. HNRNPL knockdown affects the early processing of 18S rRNAs, such as the increased abundance of 34S RNAs and the decreased abundance of both 26S and 18S-E RNAs (Tafforeau et al., 2013). DNB including HNRNPL may regulate the optimal pre-rRNA processing events at the perinucleolus. To further understand the function of the DNB, it is of crucial importance to identify their arcRNAs. Functional analyses of arcRNAs will elucidate the mechanism underlying both the formation and the dynamics of DNBs as well as their biological functions. Detailed analyses of DNBs will reveal the nature of the mechanism underlying the functions of arcRNAs in the formation of arcRNA-dependent NBs.

MATERIALS AND METHODS
Request a protocol through Bio-protocol.
**Plasmid construction**

The pcDNAs/FRT/TO-DBC1-3xFLAG plasmid was generated by inserting the PCR-amplified DBC1-3xFLAG sequence into the pcDNAs/FRT/TO vector (Thermo Fisher Scientific) between the EcoRV and Xhol sites. The pcDNA6/TR-IREs-puro plasmid was constructed by ligating the PCR-amplified sequence containing a pUC origin to the TetR gene of pcDNA6/TR (Thermo Fisher Scientific) with the segment of IRES to the puromycin-resistance gene of pCAGGS-FLPe (Gene Bridges) between the EcoRI and PstI sites. The HNRNPL-deletion mutants were constructed as described.
FIGURE 5: The RRM and PR of HNRNPL contribute to phase separation. (A) Sedimentation assay with the indicated salt concentrations of HNRNPL. The presence or the absence of TEV protease is shown above the panel (“+” or “−”). The molecular mass marker (kDa) is shown on the left. (B) Turbidity assay with the indicated NaCl concentrations of HNRNPL. Mean of five independent experiments, ±SD. (C) Imaging of the indicated salt concentrations of HNRNPL. Bar, 20 μm.
The mutated residues of HNRNPL 
PRmut1 were P335A, P336A, P337A, and P338A, while those of 
HNRNPL PRmut2 were P366A, P367A, P368A, P369A, P370A, 
P371A, P372A, and P373A, and those of HNRNPL PRmut3 were 
P335A, P336A, P337A, P338A, P366A, P367A, P368A, P369A, 
P370A, P371A, P372A, and P373A. The HNRNPL PR mutants were generated by 
site-directed mutagenesis using PCR. The 
pMAL_TEV-HNRNPL plasmid was generated by inserting the PCR-amplified 
HNRNPL sequence into the pMAL_TEV vec-
tor (Yoshizawa et al., 2018) between the 
Hin-dIII and 
BamHI sites. The primers used are 
listed in Supplemental Table S1.

Cell culture
We generated HCT116/TR_DBC1-3xFLAG 
cells using the Flp-In System (Thermo Fisher 
Scientific). First, we generated the doxycycline-induced HCT116/FRT stable cell lines, 
HCT116/FRT/TR, that express the tetracy-
cline repressor (TR) from pcDNA6/TR_IRES-
puro. The pcDNA6/TR-IRES-puro plasmids 
were transfected using Lipofectamine 3000 
(Thermo Fisher Scientific) into the HCT116/ 
FRT cells (Satoh et al., 2014) and selected 
with 2 μg/ml puromycin, and single clones 
of HCT116/FRT/TR cells were isolated. 
HCT116/TR_DBC1-3xFLAG cells were pre-
pared using the isolated HCT116/FRT/TR 
by the Flp-In System and cultured at 37°C 
with 5% CO₂ in DMEM/10% fetal bovine se-
rum (FBS), supplemented with 2 μg/ml pu-
romycin and 200 μg/ml hygromycin B. 
HCT116, NIH3T3, and HeLa cells were cul-
tured in DMEM/10% FBS at 37°C with 5% 
CO₂. The cells were treated with CX5461 
(2 μM, AdooQ Bioscience), actinomycin D 
(0.03 μg/ml or 0.3 μg/ml, Wako) and DRB 
(100 μM, TCI) for 4 h or 1,6-HD and 2,5-HD 
(2.5%, 5% or 7.5%, Sigma-Aldrich) for 5 min, 
respectively.

RNase treatment of cells
The RNase treatment was performed as de-
scribed previously (Mannen et al., 2016). 
Briefly, the cells were seeded onto round, 
12-mm-diameter coverslips of 24-well 
plates, rinsed in phosphate-buffered saline 
(PBS), and then rinsed in a permeabilization 
buffer (20 mM Tris-HCl, pH 7.4, 5 mM 
MgCl₂, 0.5 mM EGTA (ethylene glycol 
tetraacetic acid), and cOmplete, Mini, 
EDTA-free protease inhibitor cocktail 
[Sigma-Aldrich]). Subsequently, cells were 
permeabilized for 10 min at room tempera-
ture (RT) in the permeabilization buffer con-
taining 2% Tween-20 and then rinsed once 
with the sole permeabilization buffer. The 
permeabilized cells were incubated with 
RNase A (Nacalai Tesque; 1 μg/ml prepared 
in PBS) for 20 min at RT. After the RNase 
treatment, cells were rinsed with PBS and fixed with 4% paraformal-
dehyde prepared in PBS at RT for 10 min. The fixed cells were per-
meabilized with 0.5% Triton X-100 prepared in PBS for 15 min, 
rinsed, and blocked with 1% bovine serum albumin (BSA) prepared 
in PBS containing 0.1% Tween-20 (PBST) for 1 h. The slides were
incubated at 4°C overnight with primary antibodies (diluted in PB containing 1% BSA) against specific proteins. Unbound antibodies were removed by three 10-min washes with PBST. The slides were then incubated with Alexa-conjugated secondary antibodies (Thermo Fisher Scientific) for 1 h at RT, washed, and mounted with Fluoro-KEEPER Antifade Reagent (Nacalai Tesque). Immunostained cells were examined using a confocal laser scanning microscope (FV1000D; Olympus). The antibodies used are listed in Supplemental Table S1.

**Plasmid transfection**

For microscopic observation, cells were seeded onto round, 12-mm-diameter coverslips of 24-well plates and transfected with plasmids using Lipofectamine 3000 (Thermo Fisher Scientific) according to the manufacturer’s instructions. The cells were usually fixed 24 h after transfection. For both small interfering RNA (siRNA) transfection and plasmid rescue experiments, cells were seeded onto round, 12-mm-diameter coverslips of 24-well plates and transfected with siRNA using Lipofectamine RNAi MAX (Thermo Fisher Scientific) according to the manufacturer’s instructions. Twenty-four hours after siRNA transfection, cells were transfected with plasmids using Lipofectamine 3000 (Thermo Fisher Scientific) according to the manufacturer’s instructions. The cells were usually fixed 24 h after transfection.

**RNA interference**

Stealth siRNAs were purchased from Thermo Fisher Scientific. Cells were grown in six-well plates and transfected with siRNAs (33 nM, final concentration) using the Lipofectamine RNAiMAX reagent (Thermo Fisher Scientific) according to the manufacturer’s instructions. After 24 h, the cells were trypsinized and seeded into six-well plates for the preparation of proteins or onto round, 12-mm-diameter coverslips of 24-well plates for the immunocytochemical experiments. The cells were cultured for 24 h before harvesting. The siRNA Control was purchased from Thermo Fisher Scientific (12935300). The siRNAs used are listed in Supplemental Table S1.

**RNA-FISH**

The RNA probes were synthesized using either T7 or SP6 RNA polymerase using a digoxigenin (DIG) RNA labeling kit (Roche). Linearized plasmids (1 μg) containing an IGS fragment were used as templates for transcription. RNA-FISH was performed as described previously (Mito et al., 2016). Briefly, the cells were seeded onto a multichamber culture slide, washed with PBS, and fixed with 4% paraformaldehyde prepared in PBS at RT for 10 min. The fixed cells were treated with 0.2 N HCl for 20 min and then with 3 mg/ml proteinase K at 37°C for 5 min. The slides were incubated with a prehybridization solution (2× SSC [sodium saline citrate], 1× Denhardt’s solution, 50% formamide, 10 mM EDTA, pH 8.0, 100 μg/ml yeast tRNA, and 0.01% Tween-20) at 55°C for 2 h. The prehybridized slides were then incubated with a hybridization solution (made of the prehybridization solution supplemented with 5% dextran sulfate and 2 μg/ml DIG-labeled RNA probe) at 55°C for 16 h. After hybridization, the slides were washed twice with a preheated wash buffer (2× SSC, 50% formamide, and 0.01% Tween-20) at 55°C for 30 min, while the excessive RNA probes were digested by incubation with 10 μg/ml RNase A prepared in NTET buffer (10 mM Tris-HCl, pH 8.0, 1 mM EDTA, 500 mM NaCl, and 0.1% Tween-20) at 37°C for 1 h. The slides were then washed once with buffer A (2× SSC and 0.01% Tween-20) at 55°C for 30 min and twice with buffer B (0.1x SSC and 0.01% Tween-20) at 55°C for 30 min. For detection, the slides were washed with Tris-buffered saline containing 0.1% Tween-20 (TBST), incubated with a blocking solution (3% BSA prepared in TBST) at RT for 1 h, and subsequently incubated with anti-DIG antibodies diluted in the blocking solution at 4°C overnight. Unbound antibodies were removed by three 15-min washes in TBST. The slides were then incubated with Alexa-conjugated secondary antibodies diluted in the blocking solution for 1 h at RT. After washing, the slides were mounted with Vectashield (Vector Laboratories) containing 4′,6-diamidino-2-phenylindole (DAPI). Fluorescence images were visualized at RT on a microscope (FV1000D; Olympus). The primers and antibodies used are listed in Supplemental Table S1.

**Immunofluorescence**

Cells were seeded onto round, 12-mm-diameter coverslips and fixed with 4% paraformaldehyde prepared in PBS at RT for 10 min. The fixed cells were permeabilized with 0.5% Triton X-100 prepared in PBS for 15 min, rinsed, and blocked with 1% BSA prepared in PBS containing 0.1% Tween-20 (PBST) for 1 h. The slides were incubated at 4°C overnight with primary antibodies (diluted in PBST containing 1% BSA) against specific proteins. Unbound antibodies were removed by three 10-min washes with PBST. The slides were then incubated with Alexa-conjugated secondary antibodies for 1 h at RT, washed, and mounted with Fluoro-KEEPER Antifade Reagent (Nacalai Tesque). Immunostained cells were examined using a confocal laser scanning microscope (FV1000D; Olympus). The antibodies used are listed in Supplemental Table S1.

**Immunoblotting**

Cells were lysed in a lysis buffer (50 mM Tris-HCl, pH 8.0, 150 mM NaCl, 0.5% IGEPAL CA630, 1 mM dithiothreitol [DTT], and complete EDTA-free protease inhibitor [Sigma-Aldrich]) and then disrupted by three pulses of sonication for 10 s. The cell extracts were cleared by centrifugation, and the protein concentration was determined using the Bradford method. A one-fifth volume of the 5x SDS sample buffer was added, and the samples were heated before the SDS–PAGE separation. After fractionation, the proteins were transferred to Immobilon-P Transfer membranes (Merck) by electrobting. The antibodies used are listed in Supplemental Table S1.

**Immunoprecipitation**

HCT116 cells were suspended in the lysis buffer for 10 min on ice and then disrupted by three pulses of sonication for 10 s. The resulting cell extracts were cleared by centrifugation at 20,400 × g for 10 min. The supernatant containing the HCT116 cells’ extract was mixed with the anti-DYKDDDDK magnetic agarose (Thermo Fisher Scientific) and rotated at 4°C overnight. The beads were finally washed five times with lysis buffer. For the RNase treatment, the beads were washed three times and treated with RNase A (1 μg/ml) at 4°C for 3 h or left untreated. The beads were then washed two times with the lysis buffer. The IP samples were eluted from the beads by 500 μg/ml 3xFLAG peptide (Protein Ark) at 4°C for 1 h.

**Mass spectrometric analysis**

Peptide mixtures for mass spectrometric analysis were prepared as described previously (Kanayama et al., 2017). Proteins in IP samples were precipitated with 10% trichloroacetic acid, resuspended in 20 μl 100 mM Tris-HCl buffer, pH 8.8, containing 6 M urea, and reduced by adding 2 μl 50 mM Tris-(2-carboxymethyl)phosphine and incubating at 60°C for 1 h. The generated free sulfhydryl groups were alkylated with 1 μl 20 mM methyl methanethiosulfonate by incubation at RT for 10 min. Proteins were digested by 1 μg of lysyl endopeptidase (Wako) at 37°C overnight. The resulting peptide mixtures were desalted with C18 Empore Disks (3M) and subjected to Liquid Chromatograph (LC)-MS/MS analysis as described earlier.
TEV-treated HNRNPL and NaCl were mixed into a buffer containing 50 mM Tris-HCl, pH 7.4, 10% (vol/vol) glycerol, and 2 mM DTT (50, 75, 100, 125, or 150 mM, final salt concentration). HNRNPL was subsequently diluted to a final concentration of 3.3 μM. Dilutions of HNRNPL supplemented with salt in the indicated concentrations were centrifuged for 10 min at 20,400 × g at 25°C. The supernatant was separated by SDS–PAGE and stained with Coomassie Brilliant Blue.

Turbidity assay

For turbidity analysis of HNRNPL, the absorbance of 3.3 μM TEV-treated HNRNPL proteins at the indicated final salt concentrations of 50, 75, 100, 125, or 150 mM (dissolved in a buffer containing 50 mM Tris-HCl, pH 7.4, 10% [vol/vol] glycerol, and 2 mM DTT) was measured at 400 nm using a SH-1000 plate reader (CORONA ELECTRIC). Values were normalized to a 50 mM NaCl concentration of HNRNPL to determine the relative turbidity value.

Imaging of the turbid HNRNPL solution

For imaging experiments, 3.3 μM TEV-treated HNRNPL proteins at the indicated final salt concentrations of 50, 75, 100, 125, or 150 mM (dissolved in a buffer containing 50 mM Tris-HCl, pH 7.4, 10% [vol/vol] glycerol, 2 mM DTT) were dropped onto individual wells of 96-well Optical-Bottom Plates (Thermo Fisher Scientific) and observed by microscope (Eclipse Ti2, Nikon) at RT.

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Estimation of the concentrations of HNRNPL in DNB

The cellular abundance levels of proteins, measured via MS-based quantitative proteomics studies, were obtained from the PaxDb (Wang et al., 2015). The cell line–integrated abundance values retrieved from PaxDb were converted into concentrations using the following formula:

\[ C = \frac{k \times A}{N_A} \]

(Milo, 2013), where \( k = 3 \times 10^6 \) proteins/fl, the Avogadro constant \( N_A = 6.02 \times 10^{23} \) molecules/mol, and A is the abundance. The intracellular concentration of HNRNPL was estimated at 2.0 μM. Our cell fractionation experiment determined that the intracellular distribution of HNRNPL in nuclei and cytoplasm was 4:1 (Supplemental Figure S9). In addition, the nuclear-to-cytoplasmic volume (N/C) ratio of HCT116 was approximately 0.18 (Ganguly et al., 2016). Consequently, the nuclear and cytoplasmic concentrations of HNRNPL were estimated to be 5.5 and 1.4 μM, respectively. The nucleoplasm and DNB volumes as well as the proportion of HNRNPL protein present within the DNB were estimated from the IF intensity profile over different Z positions (Z-stack) using cellSens Dimension software (Olympus). The ratio of nucleoplasm volume to DNB volume was 629:1, whereas the ratio of the proportion of HNRNPL protein present in the nucleoplasm to that in the DNB was 0.4:1:0. Consequently, the nucleoplasmic and DNB concentrations of HNRNPL were estimated at 5.5 and 13.7 μM, respectively.

Recombinant protein expression and purification

All recombinant proteins were expressed individually in Rosetta 2 (DE3) Escherichia coli cells (Novagen; induced with 0.5 mM isopropyl-β-D-thiogalactoside [IPTG] for 16 h at 18°C). Bacteria expressing MBP-HNRNPL proteins were lysed using a cell homogenizer (QSonica) in a lysis buffer (50 mM Tris-HCl, pH 7.4, 500 mM NaCl, 10% [vol/vol] glycerol, 2 mM DTT). MBP-HNRNPL proteins were purified by affinity chromatography using amylose resin (NEB), eluted with the lysis buffer containing 20 mM maltose, and purified by gel filtration chromatography using a HiLoad 16/600 Superdex 200 pg column (GE Healthcare). The purified proteins were freshly frozen in liquid nitrogen and stored at −80°C. Protein concentrations were determined by absorbance at 280 nm using their extinction coefficients predicted by the ProtParam tool. MBP-HNRNPL proteins (10 μM) (dissolved in a buffer containing 50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 10% [vol/vol] glycerol, 2 mM DTT) were treated with Tobacco Etch Virus (TEV) protease (Sigma-Aldrich; 25 μg/ml TEV, final concentration) in reaction volumes of 400 μl for 3 h at 30°C.

Sedimentation assay

For sedimentation analysis of HNRNPL, 10 μM purified and TEV-treated HNRNPL and NaCl were mixed into a buffer containing 50 mM Tris-HCl, pH 7.4, 10% (vol/vol) glycerol, and 2 mM DTT (50, 75, 100, 125, or 150 mM, final salt concentration). HNRNPL was subsequently diluted to a final concentration of 3.3 μM. Dilutions of HNRNPL supplemented with salt in the indicated concentrations were centrifuged for 10 min at 20,400 × g at 25°C. The supernatant was separated by SDS–PAGE and stained with Coomassie Brilliant Blue.

(Tohsato et al., 2012). The peptides were loaded on a frit-less Mightysil C18 column, washed with 2% acetonitrile in 0.1% formic acid, and eluted using the DiNa Al automatic system (KYA TECH) at a flow rate of 0.2 μl/min and the following elution gradient: 0%–50% solvent B (80% acetonitrile in 0.1% formic acid) in solvent A (2% acetonitrile in 0.1% formic acid) from 0 to the 195th min, 50%–100% solvent B in solvent A from the 195th to the 210th min, and 100% solvent B from the 210th to the 220th min. MS/MS analysis was performed by using a Q-TOF hybrid mass spectrometer (QSTAR Elite; AB Sciex). Protein identification was performed using ProteinPilot software 2.0 (AB Sciex) against the human UniProt database (version 2015_11) as described by Tohsato et al. (2012).
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