Mammalian microtubule P-body dynamics are mediated by nesprin-1

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Nesprins are a multi-isomeric family of spectrin-repeat (SR) proteins, predominantly known as nuclear envelope scaffolds. However, isoforms that function beyond the nuclear envelope remain poorly examined. Here, we characterize p50Nesp1, a 50-kD isoform that localizes to processing bodies (PBs), where it acts as a microtubule-associated protein capable of linking mRNP complexes to microtubules. Overexpression of dominant-negative p50Nesp1 caused Rck/p54, but not GW182, displacement from microtubules, resulting in reduced PB movement and cross talk with stress granules (SGs). These cells disassembled canonical SGs induced by sodium arsenite, but not those induced by hydrogen peroxide, leading to cell death and revealing PB-microtubule attachment is required for hydrogen peroxide-induced SG anti-apoptotic functions. Furthermore, p50Nesp1 was required for miRNA-mediated silencing and interacted with core miRISC silencers Ago2 and Rck/p54 in an RNA-dependent manner and with GW182 in a microtubule-dependent manner. These data identify p50Nesp1 as a multi-functional PB component and microtubule scaffold necessary for RNA granule dynamics and provides evidence for PB and SG micro-heterogeneity.

Introduction

Nesprins are a family of cellular scaffolds and linkers, composed of spectrin repeats (SRs) and a C-terminal nuclear envelope (NE) targeting KASH (Klarsicht/ANC-1/Syne homology) transmembrane domain (Zhang et al., 2001, 2005; Warren et al., 2005; Mellad et al., 2011; Rajgor and Shanahan, 2013). To date, four nesprin proteins have been identified, encoded by separate genes and capable of generating multiple tissue-specific isoforms. The full-length gene products of nesprin-1 and -2 contain a pair of N-terminal calponin homology domains that bind F-actin (Zhang et al., 2002). Nesprin-3 interacts with plectin, a cytoskeletal cross-linker that associates nesprin-3 with intermediate filaments (Wilhelmsen et al., 2005). Nesprin-4 interacts with Kif5B, a subunit of kinesin-1, and functions in nuclear migration and cell polarity (Roux et al., 2009; Horn et al., 2013). At the NE, nesprins form high-order structures called the linker of the nucleoskeleton and cytoskeleton (LINC) complex (Crisp et al., 2006; Stewart-Hutchinson et al., 2008; Mellad et al., 2011), which connects the nuclear lamina to the cytoskeleton.

In addition to nuclear–cytoskeletal coupling, scaffolding roles for nesprin-1 and -2 have been identified beyond the NE for KASH-less isoforms. The nesprin-1 isoform GSRP-56 localizes to the Golgi and regulates its structure (Kobayashi et al., 2006), whereas KASH-less nesprin-2 scaffolds ERK1/2 complexes in promyelocytic leukemia bodies and regulates vascular smooth muscle cell (VSMC) proliferation (Bernardi and Pandolfi, 2007; Warren et al., 2010). Other KASH-less isoforms include Drop-1 and CPG2, which are down-regulated in cancers and required for synaptic plasticity, respectively (Cottrell et al., 2004; Marmé et al., 2008).

Recently, we identified additional KASH-less tissue-specific nesprin-1 and -2 isoforms, suggesting new intracellular scaffolding functions for nesprins (Rajgor et al., 2012). In this report, we show that one of these, p50Nesp1, localizes and interacts with a family of RNA-binding proteins in processing bodies (P-bodies/ PBs). PBs are dynamic, nonmembranous domains containing non-translating mRNAs and proteins involved in post-transcriptional...
processes, including mRNA decapping (e.g., Dcp1a/2, Lsm1-7), mRNA degradation (e.g., XRN1), nonsense-mediated decay (e.g., hUPF1, hSMG5/7), translational repression (e.g., Rck/p54, eIF4E-T), and miRNA-mediated gene silencing (e.g., Argonautes, GW182, Rck/p54; Eulalio et al., 2007a; Moser and Fritzler, 2010). The presence of mRNA species in PBs, including mRNA decay intermediates and miRNAs, suggests they are likely to be functional entities (Sheth and Parker, 2003; Nathans et al., 2009; Castilla-Llorente et al., 2012). Additionally, mRNAs within PBs are capable of being released and translated onto polysomes (Brengues et al., 2005; Balagopal and Parker, 2009). Thus, PBs are likely to function as post-transcriptional regulatory hubs by acting as reservoirs for nontranslating mRNAs. Furthermore, PBs are anchored to microtubules (MTs) and are capable of moving within the cytosol (Aizer et al., 2008; Lindsay and McCaffrey, 2011).

During stress, related RNA stress granules (SGs) form (Anderson and Kedersha, 2008, 2009). SGs are composed of collapsed translation initiation complexes and RNA-binding proteins involved in several aspects of cellular metabolism. Their formation is thought to be essential for cell survival, as they accommodate transcripts for housekeeping proteins during stress to facilitate the preferential translation of proteins and repair enzymes required to deal with the insult (Arimoto et al., 2008). The frequent association observed between SGs and PBs is thought to mediate an exchange process where transcripts can be transferred to PBs for degradation or storage (Kedersha et al., 2005). Although PBs and SGs are structurally distinct, eIF4E, XRN1, and TTP are present in PBs in unstressed cells and capable of relocating to SGs under stress, suggesting bidirectional exchange occurs (Kedersha et al., 2005).

Mechanisms surrounding PB formation, disassembly, and dynamic properties related to SG association remain poorly characterized. In this study, we show p50Nesp1 connects PBs to MTs by acting as a microtubule-associated protein (MAP). Moreover, the dynamic properties of PB–MT attachment via p50Nesp1 are investigated with respect to PB motion and their ability to cross talk with SGs. Additionally, we demonstrate that p50Nesp1 interacts with core miRNA-induced silencing complex (miRISC) components and is required for miRNA function and PB stability.

**Results**

**Non-NE nesprin-1 variants**

A polyclonal antibody, pAbN4, was generated against a peptide sequence in SR49 of nesprin-1, which is present in multiple KASH-less isoforms that display nucleolar and cytoplasmic localizations when ectopically expressed in cells (Fig. 1 A; Rajgor et al., 2012).

Immunofluorescence microscopy using pAbN4 on U2OS cells, human dermal fibroblasts (HDFs), and VSMCs detected nuclear structures that colocalized with the nucleolar marker fibrillarin and recognized one or more nucleoli per cell (Fig. 1 B and Fig. S1, A and B). pAbN4 also exhibited some diffuse cytoplasmic staining, but most obviously detected multiple cytoplasmic foci that varied in size and number (Fig. 1 B); U2OS cells had 5 or more foci, VSMCs rarely expressed more than 4, and HDFs had the greatest variation with some cells foci-less and others displaying up to 20 (Fig. 1 C).

NE localization was not observed in these cell lines, suggesting they do not express large nesprin-1 KASH domain variants. However, nuclear rims were detected in C2C12 myoblasts, showing pAbN4 is capable of detecting large KASH domain variants containing SR49. C2C12 cells had no nucleolar staining but had a large number of cytoplasmic foci (Fig. S1 C).

Although nucleolar and diffuse cytoplasmic stainings were expected based on exogenous p31Nesp1, p23Nesp1, and p12Nesp1 localizing to the nucleolus in HDFs and cytosol in U2OS cells (Rajgor et al., 2012), cytoplasmic foci staining was not. Peptide blocking pAbN4 diminished all nucleolar and cytosolic staining, verifying pAbN4 specificity (Fig. 1 D). In addition, a second polyclonal, pAbN5, created to another peptide sequence in SR49 also labeled multiple cytoplasmic foci and the cytosol. Unlike pAbN4 it labeled the nuclear matrix and not the nucleolus, with all localizations lost by peptide blocking (Fig. 1 D). Taken together, these data suggested that these cytoplasmic foci contained a bona fide nesprin-1 isoform.

**Nesprin-1 localizes to mRNA processing bodies**

Transmission immunoelectron microscopy was performed to identify the nature of the cytoplasmic foci. Gold labeling was apparent around the pericentriolar material and also localized to multiple electron-dense, nonmembranous foci surrounding the centrosome in HDFs (Fig. 2 A)—all features characteristic of P-bodies/PBs (Eystathioy et al., 2002).

pAbN4 cytoplasmic foci colocalized with the core RISC (RNA-induced silencing complex) regulator GW182, translational regulator RFP-Rck/p54, and de-capping cofactors Hedls and Dcp1a-YFP within PBs of U2OS cells (Fig. 2 B). Likewise, the cytoplasmic foci detected with pAbN5 also colocalized with Hedls (Fig. 2 C), and those detected in C2C12 cells colocalized with GW182 (Fig. S1 D).

Immunostaining for eIF3-η, which labels SGs in arsenite-stressed cells, often juxtaposed with the nesprin-1 foci, a typical phenomenon observed between PBs and SGs (Fig. 2 D) (Kedersha et al., 2005).

**p50Nesp1 associates with Dcp1a, Rck/p54, and Ago2 complexes, but not GW182**

To identify components associated with nesprin-1 in PBs, pAbN4 immune complexes from U2OS cells were probed for PB proteins. Dcp1a, Rck/p54, and Ago2 were detected with nesprin-1; however, GW182 was absent. Similarly, pAbN5 also precipitated Dcp1a and Ago2 (unpublished data). All interactions were abolished when lysates were predigested with RNase A, indicating nesprin-1 complexes are bridged by RNA species (Fig. 3 A). When reverse co-immunoprecipitations (co-IPs) were performed, Dcp1a, Rck/p54, and Ago2 precipitated a 50-kD nesprin-1 isoform corresponding to p50Nesp1. GW182 immune complexes did not contain any nesprin-1 isoforms detected by pAbN4 (Fig. 3 B). Immune complexes could not be probed with pAbN5, as this antibody was unsuitable for immunoblotting (unpublished data).
**p50Nesp1** GST fusion constructs were created to map binding sites for its associated PB proteins. Full-length p50Nesp1, its first two SRs (p50NT), last two SRs (p50CT) and the four individual SRs (p50SR1, p50SR2, p50SR3, and p50SR4) were GST tagged and incubated with lysates from U2OS. Rck/p54, Dcp1a, and Ago2 pulled down with full-length GST-p50Nesp1 and GST-p50NT. More specifically, GST-p50SR1 pulled down Rck/p54 and Ago2 but not Dcp1a, which associated with GST-p50SR2 instead. Consistent with our co-IPs, GW182 did not interact with any GST constructs (Fig. 3 C).
In addition, immune complexes from Flag-p50Nesp1–transfected U2OS cells also contained Ago2, Dcp1a, and Rck/p54, but not GW182. Flag-p50NT was also capable of weakly co-precipitating Ago2, Dcp1a, and Rck/p54 (Fig. 3 D). However, a truncated isoform, p31 Nesp1, which contains the SRs present in p50NT, was unable to co-immunoprecipitate any of these proteins, suggesting these interactions were specific to p50Nesp1.

p50Nesp1 localizes to PBs and microtubules

To examine the effects of ectopic p50Nesp1 on PBs, Flag-p50Nesp1 was expressed in multiple cell lines. In VSMCs, which express a small number of PBs (Fig. 1 C), Flag-p50Nesp1 localized to and induced PB formation. However, in U2OS cells and HDFs, but not VSMCs, Flag-p50Nesp1 formed bundled structures reminiscent of cytoskeletal filaments. These structures did not contain F-actin or intermediate filaments, as no colocalization with phalloidin or vimentin was apparent (Fig. S2). However, the bundles strongly colocalized with α-tubulin, suggesting p50Nesp1 may be a microtubule-associated protein (MAP) capable of direct interactions (Fig. 4 A). To examine this possibility, we performed in vitro MT cosedimentation assays. Full-length p50Nesp1 and p50CT co-pelleted with polymerized MTs after ultra-centrifugation, but were not detected in pellets when MTs were absent. The SRs in p50NT remained in the supernatant after ultracentrifugation in the presence and absence of MTs, ruling out MT interaction with these SRs. Interestingly, SR3 and SR4 failed to pellet with MTs independently, indicating the MT binding of p50Nesp1 requires both the C-terminal SRs in tandem (Fig. 4 B).

In both U2OS cells and VSMCs, Flag-p50CT localized to MTs in a similar fashion to full-length p50Nesp1 in U2OS cells, confirming MT association (Fig. 4 C). However, Flag-SR1, -SR2, -SR3, -SR4, and -p50NT all displayed diffusive cytoplasmic localizations in U2OS cells (Fig. 4 D).

p50Nesp1 recruits PBs to MTs

By interacting with PB proteins through its N-terminal SRs and MTs via its C-terminal SRs, we hypothesized p50Nesp1 as a PB–MT linker. To explore this possibility, we examined Rck/p54 and GW182 PBs in U2OS cells, confirming MT association (Fig. 4 C). However, Flag-SR1, -SR2, -SR3, -SR4, and -p50NT all displayed diffusive cytoplasmic localizations in U2OS cells (Fig. 4 D).

To demonstrate Rck/p54 recruitment and displacement from MTs was not an artifact of MT bundling, we examined Rck/p54 in U2OS cells transfected with prostate-derived sterile 20-like kinase 1α (PSK1α), a kinase which promotes similar MT bundling (Mitsopoulos et al., 2003). In these cells, Rck/p54 PBs colocalized strongly with PSK1α-induced MTs, confirming that displacement of Rck/p54 PBs from bundled MTs was a p50CT-specific event (Fig. S3).
Rck/p54 expression appeared to be enhanced in Flag-p50Nesp1 cells compared with neighboring untransfected cells and Flag-p50CT cells, whereas GW182 expression appeared to be enhanced in both. Immunoblotting confirmed Rck/p54, Dcp1a, and Ago2 were more abundant in Flag-p50Nesp1 cells compared with Flag-p50CT and Flag-control (empty flag vector), whereas GW182 expression was higher in both Flag-p50CT and Flag-p50Nesp1 compared with Flag-control. α-Tubulin levels were elevated in both Flag-p50CT- and Flag-p50Nesp1-transfected cells, indicative of MT stabilization (Fig. 5 C).

**p50Nesp1−GW182 interaction is MT dependent**

GW182 recruitment to Flag-p50Nesp1 and Flag-p50CT bundles suggested that GW182−p50Nesp1 association maybe MT dependent. Our binding studies were performed at 4°C and therefore MTs were likely to be depolymerized. Therefore, we re-performed co-IPs and GST pull-downs in U2OS cells pretreated with paclitaxel (taxol) for 24 h and co-IP buffer supplemented with taxol. Under these conditions, GW182 purified with pAbN4 immune complexes and also pulled down with GST-p50Nesp1.
Figure 4. p50\textsuperscript{Nesp1} localizes to PBs and to MTs. (a) Flag-p50\textsuperscript{Nesp1} localized to and induced PB formation in VSMCs. Arrows point to colocalized foci shown in inset. In U2OS cells and HDFs, Flag-p50\textsuperscript{Nesp1} localized to and bundled MTs. (b) GST-p50\textsuperscript{Nesp1} and GST-p50\textsuperscript{CT} co-pellet with MTs in vitro. In the absence of MTs, all constructs remained in the supernatant. (c) Flag-p50\textsuperscript{CT} localizes to MTs in U2OS cells and VSMCs. (d) Flag-SR1, -SR2, -SR3, -SR4, and -p50\textsuperscript{NT} were cytoplasmic in U2OS cells. Bars: (main panels) 10 µm; (a, insets) 2 µm.
and the MT-binding SRs in GST-p50CT (Fig. 5, D and E). When pAbN4 immune complexes were isolated in a similar manner in the presence of nocodazole to U2OS cells, GW182–nesprin interactions were abolished (Fig. 5 D), but not nesprin–1–Rck/p54. –Dcp1a, or –Ago2 interactions, suggesting p50\textsuperscript{Nesp1} and GW182 interact in an MT-dependent manner.

p50\textsuperscript{Nesp1}, MT scaffolds mediate PB motion

As Flag-p50CT was able to displace Rck/p54 from MTs in U2OS cells, PB motion was examined. Real-time tracking of Dcp1a-YFP in U2OS cells cotransfected either with Flag-p50\textsuperscript{Nesp1}, Flag-p50CT, or Flag-control was performed. Dcp1a-YFP PBs were traced for 2 min using time-lapse microscopy. The area covered by each PB was determined and used as a measure to determine motion. As described previously, PBs mainly localized in spatially confined regions with restricted motion (Aizer et al., 2008). Flag-control (n = 84) and Flag-p50\textsuperscript{Nesp1} (n = 262) expressing cells covered areas of 0.0798 µm\textsuperscript{2} and 0.0799 µm\textsuperscript{2}, respectively. However, cells expressing Flag-p50CT (n = 116) covered significantly reduced areas averaging 0.0514 µm\textsuperscript{2}, comparable coverage to Flag-control cells treated with nocodazole (n = 429) at 0.0501 µm\textsuperscript{2} (Fig. 5 F).

To confirm that U2OS cells coexpressing Dcp1a-YFP and the respective Flag-protein were being recorded, culture dishes were stained for anti-Flag after filming. Every cell expressing Dcp1a-YFP also expressed Flag-p50\textsuperscript{Nesp1} or Flag-p50CT, allowing easy selection for cotransfected cells. Interestingly, Flag-p50\textsuperscript{Nesp1} no longer localized to MTs when cotransfected with Dcp1a-YFP and instead colocalized with it in PBs. However, Flag-p50CT localized to MTs with Dcp1a-YFP remaining unattached (Fig. 5 G). These data suggest that expression levels of proteins capable of binding to SR1+2 of p50\textsuperscript{Nesp1} may determine whether ectopic p50\textsuperscript{Nesp1} localizes to PBs or MTs.

To determine if the hampered PB–SG dynamics in Flag-p50CT cells was specific for H\textsubscript{2}O\textsubscript{2}-induced stress, we re-performed these experiments in cells stressed with 0.5 mM sodium arsenite (SA) for 1 h. The number of PB–SG connections as well as β-globin localization was similar to that seen by cells treated with H\textsubscript{2}O\textsubscript{2} (Fig. 6, D–F). However, SGs disassembled in both Flag-control and Flag-p50CT transfected cells when allowed to recover in normal media (Fig. 6 D). In agreement with a previous report, these experiments suggest H\textsubscript{2}O\textsubscript{2}-induced SGs differ from those induced by SA, and suggest there may be heterogeneity in the dynamic properties of SGs induced by different stressors (Emara et al., 2012).

To examine the dynamics of mRNA sorting and SGs in cells where the binding proteins associated with SR1+2 of p50\textsuperscript{Nesp1} remained attached to MTs, we performed these experiments in cells transfected with Flag-p50\textsuperscript{Nesp1}. SGs induced by H\textsubscript{2}O\textsubscript{2} or SA were frequently associated with, or were in close proximity to, the bundled MTs detected by Rck/p54 staining (Fig. 6 G). These cells contained small amounts of the reporter mRNA in SGs, with the majority colocalizing with Rck/p54 on MTs, consistent with the localization of the reporter with Rck/p54 PBs in Flag-control cells. The presence of β-globin scattered along the MTs made quantification difficult. Therefore, to measure the efficiency of mRNA sorting within these cells, we quantified the number of SGs containing the β-globin reporter relative to empty ones in H\textsubscript{2}O\textsubscript{2}- and SA-treated cells. In Flag-control cells, ~20% of SGs contained the reporter, whereas in Flag-p50\textsuperscript{Nesp1} significantly more (~30%) contained the reporter. However, β-globin mRNA was detected in ~90% of SGs in Flag-p50CT cells, a significant increase compared with both Flag-control and Flag-p50\textsuperscript{Nesp1} (Fig. 6, H and I). These findings indicate that there are changes in mRNA distribution in cells overexpressing p50\textsuperscript{Nesp1} and p50CT; however, in Flag-p50\textsuperscript{Nesp1} cells mRNA can be retained in complexes containing Rck/p54 and p50\textsuperscript{Nesp1} binding partners, whereas in Flag-p50CT cells, where PBs are displaced from MTs, mRNA cannot be distributed into Rck/p54 PBs. Like Flag-control cells, H\textsubscript{2}O\textsubscript{2}- and SA-induced SGs disassembled during recovery in Flag-p50\textsuperscript{Nesp1} cells, indicating that retention of PBs onto MTs has no effect on SG disassembly. (Fig. 6 G).

H\textsubscript{2}O\textsubscript{2} triggers cell death in Flag-p50CT-expressing cells

We performed 3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide (MTT) assays to determine whether the SG disassembly defects induced by Flag-p50CT in H\textsubscript{2}O\textsubscript{2}-treated cells caused toxicity. After applying H\textsubscript{2}O\textsubscript{2} or SA onto cells for 1 h, MTT readings were performed hourly for up to 8 h in recovering cells. Mild cellular toxicity was observed 4–5 h into
Figure 5. p50Nesp1 links PBs to microtubules (MTs) and regulates PB motion. (a) Flag-p50\textsuperscript{Nesp1} recruits Rck/p54 to MTs in U2OS cells. Flag-p50CT displaces Rck/p54 PBs from MTs. Arrows point to region shown in inset. (b) Flag-p50\textsuperscript{Nesp1} and Flag-p50CT recruit GW182 to MTs. (c) Western blot showing levels of PB components and tubulin in U2OS cells transfected with Flag-p50\textsuperscript{Nesp1}, -p50CT, and -control (empty flag vector). GAPDH served as a loading control.
recovery in Flag-p50CT cells treated with H₂O₂ and more severe toxicity was observed after 7 h, and this was confirmed by time-lapse video microscopy that showed cell death. By contrast, no toxicity or death was seen in Flag-p50CT cells stressed with SA. Similarly, Flag-control or Flag-p50Nesp1-transfected cells did not display any toxic symptoms after H₂O₂ or SA treatment (Fig. 7, A and B).

p50Nesp1 knock-down eliminates PBs

A panel of siRNAs was designed to examine PBs in p50Nesp1-depleted cells. si-83 and si-90 were created to target the first and last coding exons of p50Nesp1, respectively. si-136 was created toward the C terminus of the nesprin-1 giant to target KASH isoforms and serve as a negative control (Fig. 8 A). To validate the efficiency of these siRNAs, we measured nesprin-1 knock-down using pAbN4 and a previously described monoclonal antibody generated to SRs near the KASH domain of nesprin-1, to determine NE isoforms after H₂O₂ or SA treatment (Fig. 7, A and B).

si-83 and si-90 knocked down p50Nesp1, p31Nesp1, 144Nesp1, and an unknown ∼75-kD variant detected by pAbN4. Additionally, si-90 eliminated p41Nesp1, whereas the nontargeting si-83 had no effect on its expression. si-136 had no effect on the expression levels of any nesprin-1 isoform detected by immunoblotting with pAbN4 (Fig. 8 B). si-136 reduced NE staining in U2OS cells detected by the C-terminal monoclonal, indicating it is capable of knocking down KASH-containing isoforms. However, si-83 did not reduce NE staining. Taken together, these data suggest that U2OS cells do not express large nesprin-1 KASH isoforms containing SR49, supporting the absence of NE staining in U2OS cells by pAbN4 (Fig. S5).

We used pAbN4, GW182, and Rck/p54 staining to assess PBs in nesprin-1–depleted U2OS cells (Fig. 8, C and D). RNAi with si-136 showed no difference in PB phenotype to controls, indicating KASH isoforms have no involvement in PB structures. In contrast, si-83 and si-90 both eliminated nesprin-1, GW182, and Rck/p54 foci in up to 80% of cells (Fig. 8 E). Nucleolar knockdown was not observed by any of the siRNAs, suggesting nucleolar staining by pAbN4 may be artifactual.

To identify which SRs were central for PB structure, we created si-83–resistant Flag-p50Nesp1 and Flag-p50NT constructs to perform rescue experiments. Flag-p50Nesp1 recruited Rck/p54 and GW182 to MTs in U2OS cells transfected with si-83. Interestingly, Flag-p50NT rescue reformed Rck/p54 and GW182 PBs, even though the construct localized diffusively within the cytosol rather than directly to PBs (Fig. 8 F). Flag-p50Nesp1 or Flag-p50NT plasmids containing the correct si-83 seed region were not detected in si-83-transfected cells (unpublished data). In Flag-p50CT rescue cells, GW182 was recruited to MTs; however, Rck/p54 PBs were not detected (Fig. 8 F).

Focusing on miR-7a

The loss of macroscopic PBs does not necessarily imply loss of function of miRNA-processing pathways (Chu and Rana, 2006; Eulalio et al., 2007b). p50Nesp1 binding partners Rck/p54, GW182, and Ago2 are core components of the miRISC. Therefore, we used the Let-7a miRISC luciferase reporter assay to assess if p50Nesp1 is required for miRNA function (Chu and Rana, 2006; Lytle et al., 2007; Johnston et al., 2010). The psi-CHECK2-lent-7X3 vector encoding luciferase contains 3x Let-7 miRNA-binding sites within its 3′ UTR, and when transfected into cells expressing endogenous Let-7, luciferase expression is translationally repressed. However, knockdown of proteins involved in the miRISC pathway reduces silencing of the reporter and enhances luciferase expression. Knockdown of p50Nesp1 using si-83 or si-90, but not si-136, significantly reduced miRISC activity in U2OS cells. Rck/p54 knockdown served as a positive control (Fig. 9 A). Similar effects on miRISC activity and PB number were also observed when these experiments were performed in VSMCs (unpublished data).

Flag-p50Nesp1 was able to rescue miRISC function in si-83–transfected cells, confirming miRISC attenuation was due to p50Nesp1 knockdown and not depletion of other nesprin-1 isoforms. Furthermore, a similar rescue was achieved by Flag-p50NT, but not by the MT binding Flag-p50CT or the individual four SRs, implying SR1+2 together make up the scaffold for translationally repressed mRNPs (Fig. 9, B and C). However, p31Nesp1, which also contains the p50NT SRs, was unable to rescue miRISC activity (unpublished data).

Finally, to determine whether p50Nesp1 had silencing capabilities independent of miRNAs, like GW182, we tethered V5-AN-p50Nesp1 to a Renilla luciferase reporter with boxB elements in its 3′ UTR. However, no significant changes in luciferase activity were observed (Fig. 9 D). Therefore, it appears p50Nesp1 cannot directly induce translational repression and may require association with GW182, Ago2, Rck/p54, and other proteins along with miRNAs to facilitate silencing.

Discussion

Here, we characterized a novel KASH-less nesprin-1 isoform, p50Nesp1, which localizes to PBs and links them to MTs. p50Nesp1 is important for PB dynamic motion and ability to associate and cross talk with SGs, and is the first SR-containing protein found to localize to PBs and act as a core miRISC component.

p50Nesp1 localizes and interacts with PB proteins and MTs

The cytoplasmic foci detected by pAbN4 and N5 colocalized with PB but not SG proteins. We identified p50Nesp1 enriched in mRNPs with Rck/p54, Ago2, and Dcp1a, and mapped the binding
Figure 6. **p50Nesp1** is required for PB–SG association. (a) Rck/p54 PBs (blue) and PABP-1 SGs (red) in Flag-control and Flag-p50CT–transfected U2OS cells treated with H_2O_2 and allowed to recover. Orange arrows point to β-globin mRNA (green) in SGs and purple arrows in PBs. (b) Flag-p50CT cells (n = 300) had significantly fewer cells showing one or more PB–SG connection relative to Flag-control cells (n = 300). (c) Flag-control cells (n = 300) accumulated significantly more β-globin mRNA in PBs, whereas Flag-p50CT cells (n = 300) had significantly more reporter in SGs. (d–f) As in a–c, respectively, but with SA. ***, P < 0.001, Student’s t test. (g) Rck/p54, PABP-1, and β-globin in Flag-p50Nesp1 cells treated with H_2O_2 or SA and allowed to recover. Orange arrows point to β-globin in SGs and purple arrows to microtubules. (h) Percentage of H_2O_2-induced SGs containing β-globin reporter. (i) As in h, but with SA-induced SGs. ***, P < 0.001; *, P < 0.05; one-way ANOVA, Bonferroni post-hoc test. Bars: (main panels) 10 µm; (a, d, and g, insets) 2 µm.
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region of these proteins to SR1+2 of p50Nesp1 (Fig. 10). These interactions with p50Nesp1 were RNA dependent; however, we do not know if RNA can interact directly with SR1+2 or whether they are bridged by other RNA-binding proteins to p50Nesp1.

In U2OS cells, Flag-p50Nesp1 promoted MT bundling and was able to bind MTs in vitro through SR3+4 (p50CT). By interacting with PB components via SR1+2 and possessing MT-binding domains in SR3+4, we show p50Nesp1 is a PB–MT linker. In support of this, Flag-p50Nesp1 recruited Rck/p54 to bundled MTs, whereas Flag-p50CT de-coupled Rck/p54 from MTs by acting as a dominant negative.

Interestingly, no GW182 was initially detected as binding to p50Nesp1. However, GW182 localized to MTs in Flag-p50Nesp1- and Flag-p50CT–transfected U2OS cells, suggesting the two proteins may interact in an MT-dependent manner. Indeed, GW182 was detected in pAb4 co-IPs and with GST-p50Nesp1 and GST-p50CT in the presence of taxol.

Unlike U2OS cells, in VSMCs, which express fewer PBs, Flag-p50Nesp1 localized to and induced PB formation. These different localizations were not simply due to differences in expression levels of the construct between the cell lines, as U2OS cells expressing low levels of Flag-p50Nesp1 also showed MT localization (unpublished data). Rather, we provide evidence that the differences in localization may be due to the abundance of other PB components associated with p50Nesp1. We showed that VSMCs express more Dcp1a, Ago2, and Rck/p54, but not GW182, compared with U2OS cells, and that Flag-p50Nesp1 co-expressed with Dcp1a-YFP in U2OS cells preferentially colocalized with Dcp1a-YFP PBs and not to MTs. This contrasted with Flag-p50CT, which remained attached to MTs when co-expressed with Dcp1a-YFP. Furthermore, although Flag-p50CT also localized to MTs in VSMCs, Flag-p50Nesp1 relocalized from PBs to MTs when Dcp1a RNAi was performed. Taken together, these data suggest that the abundance and/or stoichiometry of proteins bound to SR1+2 of p50Nesp1 influence whether it localizes to PBs or MTs. We also observed that Flag-p50Nesp1 overexpression in U2OS increased levels of Dcp1a and other N-terminal binding partners, suggesting that a feedback mechanism regulates

Figure 7. Flag-p50CT promotes cell death in response to H2O2. (a) Cells were treated with H2O2 or SA for 1 h to induce SGs, followed by hourly MTT readings during recovery. 0 h is MTT reading before stress. (b) Time-lapse microscopy demonstrates lethality of Flag-p50CT in U2OS cells ~8 h after H2O2 stress relief, but not after SA. Bars, 10 µm.
Figure 8. **p50Ne split knockdown eliminates PBs.** (a) Schematic depicting siRNA oligos relative to nesprin-1 giant and p50Ne split. (b) Immunoblotting confirmed efficiency of si-83 and si-90 RNAi. (c) si-83- and si-90-mediated p50Ne split RNAi eliminates pAbN4 and GW182 PBs in U2OS cells. (d) Quantification of PB depletion in U2OS cells (n = 600 cells). *, P < 0.001; one-way ANOVA, Dunnett’s post-hoc test. (f) Flag-p50Ne split, Flag-p50NT, and Flag-p50CT rescues in si-83-transfected cells. Bars, 10 µM.
Nesprin-1 is a novel microtubule P-body scaffold

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Nesprin-1 and nesprin-3 have been shown to dimerize in vitro (Djinovic-Carugo et al., 2002; Mislow et al., 2002; Ketema et al., 2007). In addition, protein regulator of cytokinesis 1 (PRC1, also known as MAP65) interacts directly with MTs via its SRs to bundle anti-parallel filaments by forming homodimers produced from opposing centrioles (Subramanian et al., 2010). Whether p50Nesp1 can cross-link MT filaments into bundles by forming such homodimers is worth further exploration.

p50Nesp1 is a core miRISC protein

We identified p50Nesp1 as a scaffold for miRISC complexes based on its ability to interact with Rck/p54, Ago2, and GW182. p50Nesp1 knockdown attenuated miRISC activity, leading to PB elimination in U2OS cells. Importantly, VSMCs also showed attenuated miRISC activity and reduced PB numbers when the abundance of these binding partners. This is also supported by the observation that Flag-p50CT did not increase levels of the binding partners. Unfortunately, due to the poor transfection efficiency of VSMCs, we were unable to test this further and future studies using constructs cloned into viral vectors are needed. Whether additional proteins also operate within this complex to regulate PB versus MT association also needs to be determined.

We showed the SRs in p50CT directly interact with MTs in tandem but also appear to promote MT bundling. This bundling is potentially due to recruitment of MT cross-linkers and further screening of p50Nesp1-binding partners may identify such factors. As high levels of Dcp1a appeared to attenuate MT bundling, proteins bound to the SRs in p50NT may play a role. Alternatively, bundling may be achieved by oligomerization of p50Nesp1. Dimerization of SR proteins is not uncommon and both nesprin-1α and nesprin-3α have been shown to dimerize in vitro (Djinovic-Carugo et al., 2002; Mislow et al., 2002; Ketema et al., 2007). In addition, protein regulator of cytokinesis 1 (PRC1, also known as MAP65) interacts directly with MTs via its SRs to bundle anti-parallel filaments by forming homodimers produced from opposing centrioles (Subramanian et al., 2010). Whether p50Nesp1 can cross-link MT filaments into bundles by forming such homodimers is worth further exploration.

Figure 9. p50Nesp1 is required for miRISC function. (a) Cells depleted of p50Nesp1 have attenuated miRISC function. si-Rck/p54 served as a positive control. (b) Flag-p50Nesp1 and Flag-p50NT rescued si-83 miRISC in U2OS cells. (c) Immunoblotting shows expression levels of nesprin constructs in RNAi cells. (d) Tethering AN-V5-p50Nesp1 to 5BoxB sites present in the 3′ UTR of Renilla had no effect on silencing. AN-V5-Rck/p54 served as positive control. Immunoblots show expression levels of p50Nesp1 and Rck/p54.
transfected with si-83 and si-90, but not si-136, suggesting both cell lines have similar p50Nesp1 miRISC functionality (unpublished data). Rescue experiments in U2OS cells demonstrated these effects were most likely caused by p50Nesp1 knockdown and not other isoforms targeted by the siRNAs. p50Nesp1 rescue restored miRNA function; however, p31Nesp1, which was knocked down by si-83 and si-90 RNAi and did not interact with PB components, was unable to rescue miRISC despite containing p50NT SRs. Although we cannot rule out the possibility that another larger unidentified isoform also depleted by both siRNAs plays a role in PB biology, this evidence suggests that each nesprin-1 isoform has unique localization and binding properties that potentially restrict its functionality to a particular process.

p50Nesp1 rescue recruited Rck/p54 and GW182 to MTs, demonstrating PB components recruited to MTs have functional miRISCs. p50CT rescue did not restore miRISC activity; however, it was able to restore GW182 to MTs, indicating GW182 bound to MTs alone cannot function in silencing. Likewise, Flag-p50CT could not attenuate miRISC activity in control cells, suggesting p50Nesp1 links to MTs are not critical for miRISC function. Although the dynamics of p50Nesp1–GW182 interactions on MTs require further investigation, we speculate that MT filaments may transiently bring GW182 into close proximity with the mRNPs bound to SR1+2 to mediate miRNA-mediated gene silencing. However, p50NT was also able to rescue miRISC activity to the same extent as full-length p50Nesp1, and interestingly, restored Rck/p54 and GW182 PBs without localizing to PBs itself. Potentially, small amounts of p50NT may be present within the restored PBs and account for miRISC rescue. Alternatively, silencing may occur in the cytosol rather than PBs, as miRISC silencing has been shown to occur in the absence of PBs (Chu and Rana, 2006; Eulalio et al., 2007b). The reestablishment of GW182 was surprising, considering it is unable to interact with p50NT. It is plausible the revival of Rck/p54 (and likely Ago2) may change the organization or conformations of mRNPs within the newly formed PBs, for example, by allowing GW182 to form direct interactions with Rck/p54 and Ago2 attached to the SRs of p50NT.
Interestingly, p50Nesp1 alone was unable to induce silencing when tethered to a reporter mRNA construct, suggesting it needs to associate with GW182 complexes to induce silencing. These associations could be transient and mediated by MTs; however, the factors regulating such interactions remain unclear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear. For example, GW182 may interact independently with MTs via other mechanisms such as Myosin-Va (see below) or clear.
ataxia type 2 and interacts with Rck/p54 and localizes to both PBs and SGs (Affaitati et al., 2001; Nonhoff et al., 2007). Therefore, disruption of nesprin-1 MT scaffolds promoting defects in PB assembly, motion, silencing functions, and SG cross talk in the neuronal system could be a novel causal occurrence promoting neurological diseases.

Materials and methods

Plasmids
Flag-p50Nesp1 and Flag-p31Nesp1 were previously cloned by our laboratory (Affaitati et al., 2010). pRK5-Myc-PSK1a was a gift from J. Morris (King’s College London, London, England, UK; Minshall et al., 2010). pDsRed2-C1-Rck/p54 was provided by G. Hutvagner (University of Dundee, Scotland, UK; Johnston et al., 2002). pEYFP-Dcp1, pEF-7B-MS2bs, and pEGFP-MS2 were gifts from N. Standart (Cambridge University, Cambridge, England, UK; Minshall et al., 2007). GW182 was a gift from M. Fritzler (University of Calgary, Alberta, Canada; Eystathioy et al., 2002). pDsRed2-C1-Rck/p54 was provided by G. Hutvagner (University of Dundee, Scotland, UK; Johnston et al., 2010), prK5-Myc-PSK1a was a gift from J. Morris (King’s College London, London, England, UK; Mitsopoulos et al., 2003).

Tissue culture
Human dermal fibroblasts (HDFs), osteosarcoma cells (U2OS), and C2C12 myoblasts were passaged after reaching 70% confluency and maintained in DMEM complete media (Sigma-Aldrich) supplemented with 10 U/ml penicillin, 10 mg/ml streptomycin, 200 µg/ml G418, and 10% FBS. Primary or SV40 VSMCs were cultured in medium M199 (Sigma-Aldrich) supplemented in 10 U/ml penicillin, 10 mg/ml streptomycin, 200 µg/ml G418, and 20% FBS. When appropriate, cells were treated with 10 µg/ml nocodazole, 10 µg/ml cycloheximide, 500 µM sodium arsenite, or 2 µM H2O2 for 1 h.

Generation of nesprin-1 pAbN4 and pAbN5
Polyclonal rabbit antibodies to human nesprin-1 N4 and N5 were generated against synthetic polypeptides EGNQGLGLKPLAKGGK (Affaitati et al., 2001; Nonhoff et al., 2007). Therefore, disruption of nesprin-1 MT scaffolds promoting defects in PB assembly, motion, silencing functions, and SG cross talk in the neuronal system could be a novel causal occurrence promoting neurological diseases.

Immunoblotting
Whole-cell lysates were prepared by scraping PBS-washed cells in ice-cold co-IP buffer (10 mM Tris, pH 7.4, 150 mM KCl, 1 mM EDTA, 1% Triton, and protease inhibitor cocktail). Lysed cells were centrifuged at 13,000 g for 20 min to obtain the supernatant lysate, and up to 20 µg of lysate were resolved on a 10% SDSPAGE gel and transferred to PVDF using a semi-dry apparatus. The membranes were blotted with the antibodies indicated in each figure, and bands were visualized using the ECL Western blotting substrate (Thermo Fisher Scientific). Membranes were incubated with the following primary antibodies at a 1:1,000 dilution: rabbit polyclonal pAbN4, rabbit polyclonal pAbN5, mouse monoclonal anti-Rck/p54 (ab54611; Abcam), mouse monoclonal anti-Dcp1a (ab57654; Abcam), rabbit polyclonal anti-Rck/p54 (A300461A; Bethyl Laboratories, Inc.), mouse monoclonal anti-Flag (F3165; Sigma-Aldrich), and anti-GAPDH (sc-32238; Santa Cruz Biotechnology, Inc.). Mouse monoclonal anti-β-actin (A-9452; Sigma-Aldrich), mouse anti-Ago2 (ab57113; Abcam), and goat polyclonal anti-GST (ab66613; Abcam) were used at 1:10,000. Secondary antibodies conjugated to HRP were obtained from GE Healthcare and used at 1:10,000 dilutions.

Let-7 miRISC luciferase assay
12,000 U2OS cells were plated into each well of a 24-well plate and cultured under standard growth conditions overnight. The following day, three wells per experimental condition were transfected with 1 µl nesprin or Rck/p54 scrambled siRNA oligo (all at 200 pM stock solutions) using HiPerFect (Qiagen) transfection reagent. 72 h after siRNA transfection, 1 µg of psiCHECK2-lent-7X3 or psiCHECK2-lent-7X3m was cotransfected with Flag constructs into each well using SuperFect (Qiagen) transfection reagent. Luciferase readings were taken 24 h after plasmid transfection using the Dual-Luciferase Reporter assay (Promega). Renilla luciferase activity was normalized to Firefly luciferase activity to control for transfection efficiency (both transfected off the psi-CHECK2 plasmid).

Plasmid transfections
Plasmids were transfected into U2OS cells, HDFs, and VSMCs using SuperFect transfection reagent (Qiagen) according to the manufacturer’s instructions. Cells were transfected onto coverslips or MatTek glass-bottom dishes and fixed for immunostaining or used for time-lapse microscopy 24 h after transfection, respectively.

Tethering assay
50,000 U2OS cells were plated per well of a 6-well plate and transfected with 1 µg phRL-Glo380x8 (Promega) and 50 ng pG3l using Superfect transfection reagent the following day. 48 h after transfection, luciferase readings were taken using the Dual-Luciferase Reporter assay system.

Immunofluorescence microscopy
Cells were fixed for 5 min in 3.7% PFA (Sigma-Aldrich) at 1:1,000, mouse anti-fibrillarin (ab4566; Abcam) at 1:200, mouse anti-S6K used to detected Hedls (sc-9362; Santa Cruz Biotechnology, Inc.) at 1:200, mouse monoclonal anti-Flag (F3165; Sigma-Aldrich) at 1:200, mAb anti-53BP1 (ab163052; Abcam), and rabbit rabbit anti-Dcp1a (ab47811; Abcam). Secondary antibodies conjugated to HRP were obtained from GE Healthcare and used at 1:10,000 dilutions.

Whole-cell lysates were prepared by scraping PBS-washed cells in ice-cold co-IP buffer (10 mM Tris, pH 7.4, 150 mM KCl, 1 mM EDTA, 1% Triton, and protease inhibitor cocktail). Lysed cells were centrifuged at 13,050 g for 20 min to obtain the supernatant lysate, and up to 20 µg of lysate were resolved on a 10% SDS-PAGE gel and transferred to PVDF using a semi-dry apparatus. The membranes were blotted with the antibodies indicated in each figure, and bands were visualized using the ECL Western blotting substrate (Thermo Fisher Scientific). Membranes were incubated with the following primary antibodies at a 1:1,000 dilution: rabbit polyclonal pAbN4, rabbit polyclonal pAbN5, mouse monoclonal anti-Rck/p54 (ab54611; Abcam), mouse monoclonal anti-Dcp1a (ab57654; Abcam), rabbit polyclonal anti-Rck/p54 (A300461A; Bethyl Laboratories, Inc.), mouse monoclonal anti-Flag (F3165; Sigma-Aldrich), and anti-GAPDH (sc-365602; Santa Cruz Biotechnology, Inc.). Mouse monoclonal anti-β-actin (A-9452; Sigma-Aldrich), mouse anti-Ago2 (F1804; Sigma-Aldrich) at 1:1,000, mouse anti-fibrillarin (ab4566; Abcam) at 1:200, mouse anti-S6K used to detected Hedls (sc-9362; Santa Cruz Biotechnology, Inc.) at 1:200, mouse monoclonal anti-Flag (F3165; Sigma-Aldrich) at 1:200, mAb anti-53BP1 (ab163052; Abcam), and rabbit rabbit anti-Dcp1a (ab47811; Abcam). Secondary antibodies conjugated to HRP were obtained from GE Healthcare and used at 1:10,000 dilutions.
objects were separated by giving Volocity a size guide of 5 µm^2. Dcp1a-pixel and subject to a 3 × 3-pixel Gaussian filter to remove noise. Touching using a 4-µm local contrast adjustment. Resultant objects were dilated by 1 of whole image intensity, with a minimum object area of 0.5 µm^2, and involved in Volocity using the “fast” (noniterative) method. PBs were detected ages were acquired every 200 ms for 2 min using a cooled CCD camera cation of 64. Volocity was used to control time-lapse acquisition and im-
ate constructs in phenol red–free DMEM and imaged on a microscope
washed three times in PBS, post-fixed in 2% glutaraldehyde for 1 h, passed
antibody (diluted 1:200) at room temperature for 30 min. Subsequently,
and washed with PBS before being mounted onto slides using Mowiol
cretion for 2 h using 0.2 mM IPTG (Sigma-Aldrich) at 30°C. Purification of the proteins was performed according to the GE Healthcare protocol using glutathione-Sepharose 4B beads (GE Healthcare). Pull-downs were performed from U2OS whole-cell lysates prepared as described above. 200 µg of protein lysates were incubated with 30 µl beads overnight at 4°C with rotation. Bound proteins were eluted into protein loading buffer and immuno-
vaspsembled, a tubulin buffer containing taxol was added to the microtubules to maintain their stability for several hours at RT. Next, 5 µg of GST-purified proteins was performed according to the GE Healthcare protocol using glutathione-Sepharose 4B beads (GE Healthcare). Pull-downs were performed from U2OS whole-cell lysates prepared as described above. 200 µg of protein lysates were incubated with 30 µl beads overnight at 4°C with rotation. Bound proteins were eluted into protein loading buffer and immuno-
nate constructs in 6-well plates and imaged on the same microscope and
washed three times with co-IP buffer, boiled in loading buffer, and analy-
used by immunofluorescence microscopy.

Time-lapse microscopy
For PB tracking, U2OS cells were transfected as described with appropri-
ate constructs in phenal red–free DMEM and imaged on a microscope (model H7000; Olympus) with a 40x LCUPlanFLN/0.6 NA objective air objective, and acquisition was controlled using Volocity software (PerkinElmer).

Peptide blocking
For peptide-blocking experiments, blocking peptides were designed to the antibody epitope of interest and synthesized by Peptide Protein Research Ltd. 10 µg of peptide was incubated with every 1 µg of primary antibody diluted in 1% BSA blocking buffer. For pAbN4 peptide blocking, the pAbN5 peptide served as a control and vice versa. The mixture was incubated with constant rotation for 1 h at RT and then incubated with fixed cells on coverslips for 1 h at RT. The ability of the peptide to block staining was determined by immunofluorescence microscopy.

Transmission immunoelectron microscopy
Cells were seeded onto 35-mm culture dishes 24 h before harvest. At the time of harvest, the cells were fixed in 100% methanol at 4°C for 10 min. The preparation was then washed in PBS and incubated with the pAbN4 antibody (diluted 1:200) at room temperature for 30 min. Subsequently, the preparations were washed three times with PBS and then incubated for 40 min with protein A gold (Polysciences). The preparations were then washed three times in PBS, post-fixed in 2% glutaraldehyde for 1 h, passed through a graded ethanol series, and embedded in Epon 812 (Poly-
sciences). After polymerization, the material was sectioned in an ultrami-
Putative constructs in 6-well plates and imaged on the same microscope and

Microtubule binding assay
In vitro microtubule-binding assays were performed using the Microtubule Binding Protein Spin-Down Assay Kit (BK029; Cytoskeletal, Inc.). GST-purified proteins were prepared as described above and incubated with polymer-
ized microtubules in an Eppendorf tube according to the manufacturer’s instructions. In brief, microtubules were assembled in Eppendorfs by incu-

Online supplementary methods
Figs. 5 (A and B), 6 (A, D, and G), and 8 (C and F) are available on the JCB DataViewer for observation under higher magnification.

Online supplemental material
Fig. S1 a shows that the nuclear structures detected with pAbN4 in U2OS cells are nucleoli, based on its colocalization with nucleolar marker fibrilla-
in U2OS cells. Fig. S1 b shows that the nuclear structures detected with pAbN4 in HDFs are nucleoli, based on its colocalization with nucleolar marker fibrillarin in HDFs. Fig. S1 c shows that pAbN4 labeled nuclear rims as well as cytoplasmic foci in C2C12 myoblasts, suggesting the anti-
body can detect large nesprin-1 KASH nuclear envelope variants that con-
tain SR49. Fig. S1 d shows the cytoplasmic foci in C2C12 myoblasts are

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also P-bodies, based on their colocalizations with GW/P bodies. Fig. S5 shows that si-136, but not si-83, knocks down nuclear envelope staining observed in U2OS cells by pAbN4. Online supplemental material is available at http://www.jcb.org/cgi/content/abstract/205/4/474.


