Fluorescence fluctuation spectroscopy reveals differential SUN protein oligomerization in living cells

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\textbf{ABSTRACT} Linker-of-nucleoskeleton-and-cytoskeleton (LINC) complexes are conserved molecular bridges within the nuclear envelope that mediate mechanical force transmission into the nucleoplasm. The core of a LINC complex is formed by a transluminal interaction between the outer and inner nuclear membrane KASH and SUN proteins, respectively. Mammals encode six KASH proteins and five SUN proteins. Recently, KASH proteins were shown to bind to the domain interfaces of trimeric SUN2 proteins in vitro. However, neither the existence of SUN2 trimers in living cells nor the extent to which other SUN proteins conform to this assembly state have been tested experimentally. Here we extend the application of fluorescence fluctuation spectroscopy to quantify SUN protein oligomerization in the nuclear envelopes of living cells. Using this approach, we demonstrate for the first time that SUN2 trimerizes in vivo and we demonstrate that the in vivo oligomerization of SUN1 is not limited to a trimer. In addition, we provide evidence to support the existence of potential regulators of SUN protein oligomerization in the nuclear envelope. The differential SUN protein oligomerization illustrated here suggests that SUN proteins may have evolved to form different assembly states in order to participate in diverse mechanotransduction events.

\textbf{INTRODUCTION} The nuclear envelope (NE) is a subdomain of the endoplasmic reticulum (ER) that encloses the genome and delineates the nuclear compartment in eukaryotic cells (Kite, 1913). It is defined by concentric inner and outer nuclear membranes (INM and ONM, respectively) separated by an \~{}30–50 nm perinuclear space (PNS) that is contiguous with the ER lumen (Watson, 1955). The INM contains a unique set of proteins that interact with the underlying nuclear lamina and chromatin, while the ONM is an extension of the ER (Burke and Stewart, 2014). INM–ONM fusion generates fenestrations throughout the NE that are occupied by nuclear pore complexes (Watson, 1959).

Though most nuclear–cytoplasmic communication occurs via nuclear pore complexes (Knockenhauer and Schwartz, 2016), it can also be mechanical in nature, such that forces generated by the cytoskeleton are transmitted across the NE into the nucleoplasm by LINC complexes (Alam et al., 2014). These conserved NE-spanning molecular bridges are critical for cell division, DNA damage repair, meiotic chromosome pairing, mechanoregulation of gene expression, and nuclear positioning (Meinke and Schirmer, 2015). Under-scoring their importance is a growing list of genetic mutations in LINC complex proteins associated with human diseases including ataxia and muscular dystrophy (Horn, 2014).

LINC complexes are composed of the ONM Klarsicht/ANC-1/Syne homology (KASH) proteins and the INM Sad1/UNC-84 (SUN) proteins (Crisp et al., 2006). KASH proteins are defined by a C-terminal KASH domain, which includes a transmembrane domain followed by the \~{}10–32-residue luminal KASH peptide (Starr and Han, 2002).
The divergent N-termini of KASH proteins extend into the cytoplasm, where they engage the cytoskeleton and signaling molecules (Luxton and Starr, 2014). SUN proteins are identified by their KASH-binding luminal C-terminal SUN domain (Malone et al., 1999), whereas their N-termini interact with A-type lamins, chromatin, and other INM proteins within the nucleoplasm (Chang et al., 2015).

Mammals encode two major SUN proteins, SUN1 and SUN2, which are widely expressed in somatic cells (Meinke and Schirmer, 2015). Consistent with their high level of sequence similarity (e.g., mouse SUN1 and SUN2 share 65% identity), SUN1 and SUN2 perform redundant functions during the DNA damage response (Lei et al., 2012) and radial neuronal migration in the developing mouse cerebral cortex and hippocampus (Zhang et al., 2009), as well as synaptic nuclear anchorage in mouse skeletal muscle (Lei et al., 2009). These redundancies may be due to the ability of both SUN1 and SUN2 to interact promiscuously with the KASH peptide of several KASH proteins including nesprin-1, -2, and -3 (Stewart-Hutchinson et al., 2008). Nevertheless, examples of SUN protein function specificity also exist. For example, SUN1 is differentially required for meiotic chromosome pairing (Ding et al., 2007; Horn et al., 2013) and nuclear pore complex insertion and distribution (Lu et al., 2008; Talamas and Hetzer, 2011). However, the molecular mechanisms underlying the redundant and specific functions of SUN1 and SUN2 remain unclear.

Recent in vitro studies reveal that SUN2 trimerizes due to the presence of a coiled coil (CC)–containing helical region within its luminal domain (Nie et al., 2016; Sosa et al., 2012; Zhou et al., 2012). SUN2 oligomerization is essential for KASH binding, which is further stabilized by an intermolecular disulfide bond formed between conserved cysteine residues in the SUN domain and KASH peptide (Sosa et al., 2012). Despite these important mechanistic advances, the in vivo relevance of SUN protein trimerization remains unclear due to the lack of suitable methods for measuring protein assembly states within the NE. Here, we sought to address this deficiency by extending the application of FFS to quantify protein–protein interactions in the NE in living cells.

RESULTS AND DISCUSSION

Quantifying NE protein–protein interactions in living cells

FFS characterizes fluctuating fluorescence signals generated by fluorescently labeled proteins passing through the optical observation volume of a confocal or two-photon microscope (Slaughter and Li, 2010). Subsequent brightness analysis of these fluorescence fluctuations provides quantitative information about the stoichiometry of the labeled proteins (Chen et al., 2003). We recently combined FFS with z-scans to quantify protein–protein interactions at the plasma membrane and in thin cell sections (Macdonald et al., 2010; Smith et al., 2014). Given the ~30–50-nm thickness of the PNS, we explored the use of FFS and z-scans as a method of quantifying protein–protein interactions within the NE. For simplicity, we refer to the INM, ONM, and PNS collectively as the NE for the remainder of this work.

A z-scan consists of an axial scan of the two-photon spot through a cell expressing a fluorescently labeled protein (Figure 1A; Macdonald et al., 2010). The resulting axial intensity profile or “z-scan” characterizes the subcellular distribution of the labeled proteins. Thus, a labeled NE protein produces a z-scan with two peaks along the trajectory, which correspond to signals generated within the ventral and dorsal NE. To quantify the fluorescence contributions from the NE and the nucleoplasm, z-scans were analyzed as previously described (Smith et al., 2014, 2015). FFS data were collected with the two-photon spot repositioned at either NE.

We initially tested the feasibility of FFS and brightness analysis within the NE by measuring the normalized brightness (b) of EGFP in this subcellular compartment. EGFP was targeted to the ER lumina/PNS by adding the signal sequence (SS) from a luminal protein, torsinA (Goodchild and Dauer, 2004, Figure 1B). The b indicates the average oligomeric state of an EGFP-tagged protein (Chen et al., 2010); that is, a monomer and a dimer correspond to a b of 1 and 2, respectively. To confirm that brightness accurately reflects stoichiometry within the NE, we measured the b of a dimeric EGFP construct (SS-EGFP2; Chen et al., 2003). Following expression, both constructs localized to the ER/NE (Figure 1C). z-scans and FFS data for these constructs were collected from expressing cells, followed by the calculation of b and number concentration (N), which is the average number of EGFPs within the observation volume. Further details on b and N can be found in Materials and Methods.

Independent of N, the mean and standard deviation of b from SS-EGFP- or SS-EGFP2–expressing cells was 1.06 ± 0.14 and 1.98 ± 0.18, respectively (Figure 1D). These results are, within experimental uncertainty, consistent with SS-EGFP being monomeric and SS-EGFP2 dimeric within the NE. This work establishes FFS and brightness analysis as a powerful method for probing the in vivo biochemical and biophysical behavior of NE proteins within their native cellular environment.

SUN2 oligomerization in the NE

Next, we sought to detect in vivo SUN2 trimerization using FFS and brightness analysis. Because quantitative brightness analysis of FFS data requires labeled proteins to be mobile (Hur et al., 2014) and full-length EGFP-tagged SUN2 was shown to be highly

![FIGURE 1: FFS and brightness analysis in the NE. (A) Identification of the dorsal (NE0) and ventral (NE1) NEs in a cell expressing EGFP-tagged NE proteins by z-scan FFS. Fluorescence intensity fluctuations are measured at either NE. (B) Constructs used in this figure. (C) Representative epifluorescence images of U2OS cells expressing the indicated constructs. Scale bar: 5 μm. (D) Brightness analysis of the cells described in C. Each data point represents the average b measured in a single cell.](image-url)
Unlike either SS-EGFP-SUN2
520–731 or SS-EGFP-SUN2
595–731, the b values obtained for SS-EGFP-SUN2
702–913 remained close to one and do not increase with concentration (Figure 2E). These results demonstrate SUN2 trimerization within the NE of living cells and that the SUN domain is not sufficient for this oligomerization, consistent with previously reported in vitro studies (Sosa et al., 2012; Zhou et al., 2012).

While SUN2
261–731 contains both types of CC, SUN2
520–731 possesses only the noncanonical CC. Despite this difference, both constructs displayed similar N-dependent oligomerization within the NE (Figure 2, C and D) suggesting that the noncanonical CC may be sufficient for SUN2 trimerization, in agreement with the use of a similar human SUN2 construct to solve the crystal structure of SUN2 trimers (Sosa et al., 2012).

**SUN1 oligomerization in the NE**

To provide insight into the conservation of SUN protein trimerization, we next investigated the oligomerization of SUN1 within the NE. Like SUN2, EGFP-tagged full-length SUN1 was shown by FRAP to be immobile (Östlund et al., 2009). Consequently, we generated an SS-EGFP-tagged construct encoding the entire luminal domain of SUN1 (SS-EGFP-SUN1
457–931) (Figure 3A). Owing to the lack of structural information available for SUN1, we limited our analysis to this construct, which was analogous to SS-EGFP-SUN2
520–731 (SS-EGFP-SUN1
702–913), and another that encodes the SUN1 domain (SS-EGFP-SUN1
1777–1913) (Figure 3A). All three constructs localized to the peripheral ER and NE (Figure 3B).

The b of SS-EGFP-SUN1
457–931 increased linearly over the entire range of N (Figure 3C), which prohibited the fitting of the data to a binding curve and the estimation of the stoichiometry of this construct. Nevertheless, the highest b value measured for SS-EGFP-SUN1
457–931 was ~5, indicating the presence of higher-order oligomeric states than those observed for SS-EGFP-SUN2
261–731. In contrast to what was observed for SS-EGFP-SUN2
520–731, the b of SS-EGFP-SUN1
702–913 did not increase appreciably above 1 over the range of measured N (Figure 3D). Finally, a lack of oligomerization similar to what was observed for SS-EGFP-SUN2
595–731, was also reflected by the b values obtained for the SUN domain encoding the SS-EGFP-SUN1
1777–1913 construct (Figure 3E). These results suggest that trimers are not the limiting assembly state for all SUN proteins, which is in agreement with a previous report of the existence of immobile macromolecular assemblies of SUN1 within the NE composed of dimers and tetramers (Lu et al., 2008).

**SUN protein oligomerization in the cytoplasm**

Given the requirement of SUN2 trimerization for KASH-binding (Sosa et al., 2012), SUN protein oligomerization may represent an important target for the regulation of LINC complex assembly. In fact, a recent report shows that the two canonical CCs of SUN2 display distinct oligomeric states, the modulation of which regulates the ability of SUN2 to interact with KASH peptides (Nie et al., 2016). However, the mechanisms responsible for regulating SUN2 oligomerization within the NE remain unknown. As an initial step toward defining these mechanisms, we quantified the oligomerization of constructs encoding EGFP-tagged luminal domains of SUN1 or SUN2 in the heterologous subcellular environment of the cytoplasm.
Cyttoplasmic expression of these constructs was achieved by removing the SS (Figure 4, A and B). FFS experiments were performed in the cytoplasm of these cells as previously described (Macdonald et al., 2010, 2013). The \( b \) of EGFP-SUN2\(^{261–731} \) increased with \( N \) and appeared to reach a limiting value of 3 at \( N > 600 \) (Figure 4C), which agrees with the estimated stoichiometry of the analogou structure in the NE. A fit of the cytosolic \( b \) data measured for cytoplasmic EGFP-SUN2\(^{261–731} \) to a monomer/dimer/trimer binding curve is shown with an estimated stoichiometry of 3 at \( N > 600 \) (Figure 4C). Because \( K_{\text{DT}} > K_{\text{MD}} \), the dimer population is negligible and a monomer/trimer equilibrium is sufficient to describe the \( b \) binding curve for EGFP-SUN2\(^{261–731} \) (Supplemental Figure S1A).

Only the cytoplasmic observation volume is experimentally obtainable (Macdonald et al., 2013). Thus, we computed an estimated NE observation volume to express \( N \) values in the NE as an approximate equivalent cytoplasmic \( N \) by accounting for the volume difference (see Materials and Methods). Applying this procedure converts the trimeric NE \( b \) binding curve (Figure 2C) to a predicted \( b \) binding curve in the cytoplasm (Figure 4C). A comparison of the trimeric NE \( b \) binding curve and the predicted cytoplasmic binding curve demonstrates that the observed \( b \) increase with \( N \) in the cytoplasm is significantly more pronounced than was observed in the NE (see figure legends for more details). In contrast, both EGFP-SUN2\(^{250–731} \) and EGFP-SUN2\(^{250–731} \) appeared to be monomeric, as their \( b \) values remained near 1 (Figure 4, E and F). These results reveal that the oligomerization of EGFP-tagged SS-EGFP-SUN2\(^{261–731} \) and SUN2\(^{250–731} \) is sensitive to as yet unidentified environmental factors. In addition, differences in the behavior of the SUN1\(^{457–913} \) constructs in the cytoplasm relative to the NE were also observed. While the \( b \) of SS-EGFP-SUN1\(^{457–913} \) and EGFP-SUN1\(^{457–913} \) rises with increasing \( N \), the \( b \) increase of EGFP-SUN1\(^{457–913} \) within the cytoplasm slows at higher concentrations, unlike the linear increase with \( N \) observed in the NE (Figure 4F; see figure legend for more details). However, the \( b \) of EGFP-tagged SUN1\(^{457–913} \) rises much faster in the cytoplasm than on the predicted \( b \) curve, which was converted from the NE \( b \) binding reaction (Figure 3C). Because it did not display appreciable oligomerization within the NE, we did not measure the oligomerization of SS-EGFP-SUN1\(^{457–913} \) in the cytoplasm.

A more pronounced \( N \)-dependent \( b \) increase for EGFP-tagged SUN2\(^{261–731} \) and SUN1\(^{457–913} \) was observed in the cytoplasm than in the NE (Figure 4, C and F). A potential explanation for these results could be the presence of unlabeled endogenous SUN proteins within the NE that compete with these labeled SUN constructs, leading to a reduction in \( b \). This competition leads to an apparent reduction in the measured binding affinity (Chen and Mueller, 2007), which potentially explains the shift of the binding curve of EGFP-tagged SS-EGFP-SUN2\(^{261–731} \) and SUN2\(^{1457–913} \) to lower \( N \). Unlike the binding affinity, the saturating value of \( b \) remains unchanged and is approached after the exogenously expressed EGFP-tagged protein concentration exceeds the endogenous concentration as well as the \( K_b \) value, thereby identifying the stoichiometry of EGFP-tagged protein complexes (Chen and Mueller, 2007).

To directly verify that the reported brightness values for the SS-EGFP-tagged luminal domains of SUN1 and SUN2 (Figures 2C and 3C) are not lowered by the endogenous population, we performed additional measurements of both SS-EGFP-SUN2\(^{261–731} \) and SS-EGFP-SUN1\(^{457–913} \) in U2OS cells expressing a short hairpin RNA (shRNA), which efficiently depleted either endogenous SUN1 or SUN2, or a noncoding (NC) control shRNA (Supplemental Figure S2A). SS-EGFP-SUN2\(^{261–731} \) was previously observed to approach the saturating value \( b = 3 \) for values of \( 100 < N < 200 \) (Figure 2C). Therefore, we selected SUN2 shRNA-expressing cells with concentrations of SS-EGFP-SUN2\(^{261–731} \) in this range, measured \( b \), and then calculated the median and quartile values (Supplemental Figure S2B). Comparing \( b \) in the absence of shRNA with \( b \) in the presence of either NC or SUN2 shRNA resulted in \( p \) values of 0.88 and 0.85, respectively. These results suggest that the \( b \) measured for SS-EGFP-SUN2\(^{261–731} \) is unaffected by the absence or presence of endogenous SUN2, and thus the reported \( b \) values accurately reflect the average stoichiometry of SS-EGFP-SUN2\(^{261–731} \) at concentrations of \( 100 < N < 200 \).
FIGURE 4: SUN1 and SUN2 oligomerization in the cytoplasm. (A) Constructs used in this figure. (B) Representative epifluorescence images of U2OS cells expressing the indicated constructs. Scale bar, 5 μm. (C–G) Plots of b vs. N for the indicated constructs. The data in C were fitted to a trimeric binding model (solid blue line), which is shown in D and E (dashed blue line) with K_{D0} = 8000 (60 μM) ± 4000 and K_{MT} = 0.3 (0.002 μM) ± 0.2. The data in F were fitted to a monomer/trimer/hexamer binding model (solid blue line) with K_{MT} = 100 (0.7 μM) ± 60 and a trimer–hexamer dissociation coefficient K_{TH} = 1500 (10 μM) ± 400, which is then shown in G (dashed blue line). Estimated binding curves (dashed red lines) for the data obtained in the NE for the indicated constructs are presented in C and F by converting N from the NE to its cytoplasmic value.

Similarly, we compared b measurements in the presence and absence of NC or SUN1-depleting shRNA (Supplemental Figure S3A) over the range 100 < N < 200. As with SS-EGFP-SUN2\textsuperscript{520-731}, we measured b and determined the median and quartiles. The b measurements performed on SS-EGFP-SUN1\textsuperscript{457-913} in the presence of NC or SUN1 shRNA resulted in p values of 0.78 and 0.61, respectively, indicating that the measured brightness is unaffected by the absence or presence of endogenous SUN1 (Supplemental Figure S3B). Taken together with our observations of SS-EGFP-SUN2\textsuperscript{520-731}, these results demonstrate that the presence of endogenously expressed unlabeled SUN proteins has a negligible impact on our reported brightness at the higher concentrations we measured. Furthermore, the loss of SUN2\textsuperscript{520-731} oligomerization in the cytoplasmic environment as compared with the NE (Figures 2D and 4D) cannot be caused by endogenous competition in the NE, since such competition can only lower the brightness in the NE environment.

Thus, our data imply the existence of unidentified regulators of SUN protein oligomerization within the NE, which may be chemical in nature. For instance, the contiguous ER lumen and PNS have a high calcium concentration and an oxidizing environment that favors the formation of disulfide bonds (Elggaard and Helenius, 2003). Because the conserved cation loop in the SUN domain of SUN2 is required for KASH binding, which also requires SUN2 trimerization (Sosa et al., 2012), we anticipate that SUN protein oligomerization may be sensitive to changes in the concentration of calcium within the PNS. In addition, SUN1 oligomerization involves interchain disulfide bonds, which leads us to speculate that SUN protein oligomerization may be influenced by the redox potential of the ER lumen/PNS (Elggaard and Helenius, 2003). Alternatively, luminal proteins such as the AAA+ ATPase torsinA, which was recently shown to localize to and be required for the assembly of transmembrane actin-associated (TAN) lines in migrating fibroblasts (Saunders et al., 2017), may structurally regulate SUN protein oligomerization.

Models of SUN1 and SUN2 oligomerization
We propose that SUN2 monomers are in equilibrium with SUN2 trimers in the NE, with no evidence for a significantly populated dimeric state (Figure 5A). Currently, we cannot distinguish between two models of SUN1 oligomerization that are not mutually exclusive (Figure 5, B’ and B”). In the first, SUN1 oligomerizes via a monomer/dimer/tetramer reaction (Figure 5B’). In the second, SUN1 oligomerizes via a monomer/trimer/hexamer reaction (Figure 5B”). Both reactions would ultimately lead to the assembly of higher-order SUN1 oligomers through progressive oligomerization (i.e., monomer to trimer to hexamer to n-mer). The second model is favored based on recently published computational modeling results from the Mofrad laboratory, which suggest that, like SUN2, SUN1 is capable of forming stable homo-trimers (Jahed et al., 2018). However, they found that unlike SUN2 homo-trimers, SUN1 homo-trimers were able to form lateral complexes via the association of their SUN domains. This model is consistent with our observation of b values in excess of trimers for the case of SS-EGFP-SUN1\textsuperscript{457-913}. Future work will be needed to carefully explore the modeling predictions put forward by the Mofrad laboratory.

The ability of SUN1 to form higher-order oligomers than SUN2 may be related to its ability to form rings around NE-associated meiotic telomeres (Horn et al., 2013) and to localize to nuclear pore
complexes (Liu et al., 2007). These SUN1-specific localizations may explain the differential requirement for SUN1 during meiotic chromosome pairing and DNA double-stranded break repair (Ding et al., 2007; Horn et al., 2013; Lottersberger et al., 2015) as well as nuclear pore complex insertion and distribution (Lu et al., 2008, Talamas and Hetzer, 2011). Moreover, SUN2 trimerization may be required for actin-dependent nuclear movement, as demonstrated by the specific recruitment of SUN2 to TAN lines in migrating fibroblasts (Luxton et al., 2010). Future efforts aimed at further understanding the mechanisms of the differential oligomerization of SUN1 and SUN2 will provide important insights into LINC complex-dependent mechatronduction and nuclear–cytoplasmic communication.

MATERIALS AND METHODS

Antibodies

Anti-EFPP mouse monoclonal antibody (mAb) MAB3580 was purchased from Sigma-Aldrich and was used at a dilution of 1:200 for immunofluorescence. Anti-SUN1 (ab74758) and -SUN2 (ab87036) were used at a 1:200 dilution to validate the shRNA-mediated depletion of SUN1 or SUN2 by immunofluorescence. Secondary antibodies were from two different sources. From Jackson ImmunoResearch Laboratories, we purchased goat anti-mouse secondary antibodies conjugated to Alexa Fluor 488 or rhodamine. From ThermoFisher Scientific, we purchased goat anti-rabbit secondary antibodies conjugated to Dylight 488 or 561. All secondary antibodies were used at a 1:200 dilution for immunofluorescence.

Reagents

4′,6-Diamidino-2-phenylindole (DAPI) was purchased from ThermoFisher Scientific. Restriction enzymes were either purchased from New England Biolabs (NEB) or Promega. Phusion DNA polymerase, T4 DNA ligase, and T4 PolyNucleotide Kinase (PNK) were also purchased from NEB. All other chemicals were from Sigma-Aldrich unless otherwise specified. Wizard SV Gel and the PCR Clean-Up System were from Promega. The GeneJet Plasmid Midiprep Kit was from ThermoFisher Scientific.

Cell culture

U2OS cells obtained from the American Type Culture collection were cultured using standard sterile technique in DMEM with 10% fetal bovine serum (FBS) from Hyclone Laboratories.

DNA constructs

NC (TR30015), SUN1 (TF300647B/FI302582), and SUN2 (TF300646A/FI302577) HuSH-29 shRNA constructs in pRFP-C-RS were purchased from OriGene Technologies. The SS-EFPP and SS-EFPP2 constructs were generated as follows, using a previously described human SS-EFPP-torsinA construct (Goodchild and Dauer, 2004). EFPP from SS-EFPP-torsinA was PCR-amplified using the primers SS-EFPP-F and SS-EFPP-R (Table 1), which contain 5′ Nhel and EcoRI cut sites, respectively. The PCR product was purified and digested alongside SS-EFPP-torsinA with Nhel and EcoRI. Following gel purification, the digested PCR product and plasmid were ligated together to create SS-EFPP. To generate SS-EFPP2, EFPP was amplified from SS-EFPP-torsinA using the primers SS-EFPP2-F and SS-EFPP2-R (Table 1), which contain 5′ Nhel and EcoRI cut sites, respectively. In addition, SS-EFPP2-F encodes a 10-amino acid linker (GHGTGSGTS) following the BsrGI site, while SS-EFPP2-R encodes a mutated BsrGI site that disrupts the 3′ BsrGI present in EFPP. The resulting PCR product was then purified and digested beside SS-EFPP with BsrGI and Apal. The digested PCR product and plasmid were purified and ligated to make SS-EFPP2.

Previously described EFPP-tagged full-length mouse SUN1 and SUN2 constructs (Luxton et al., 2010) were used as templates for the generation of the SS-EFPP-tagged luminal SUN1 and SUN2 constructs. To create SS-EFPP-SUN1261–731, the sequence encoding amino acids 261–731 was PCR-amplified from EFPP-SUN2 using the SS-EFPP-SUN1261–731-F and SS-EFPP2-SUN1261–731-R primer pair (Table 1), which contain 5′ BsrGI and EcoRI cut sites, respectively. In addition, SS-EFPP-SUN1261–731-F encodes a 10-amino acid linker (GHGTGSGTS) following the BsrGI site. The PCR product was purified and digested alongside SS-EFPP with BsrGI and EcoRI. Following gel purification, the digested PCR product and plasmid were ligated together to create SS-EFPP-SUN262–731. SS-EFPP-SUN262–731 and SS-EFPP-SUN295–731 were both generated via kinase, ligase, DpnI treatment where 2 µl PCR product was treated with T4 ligase, T4 PNK, and DpnI in T4 ligase buffer in a 20-µL reaction for 20 min at room temperature. The forward primers used to create SS-EFPP-SUN1262–731 and SS-EFPP-SUN295–731 were SS-EFPP-SUN1262–731-F and SS-EFPP-SUN295–731-F, respectively (Table 1). The same reverse primer, SS-EFPP-SUN29R (Table 1), was used for both SS-EFPP-SUN1262–731 and SS-EFPP-SUN295–731.

To create SS-EFPP-SUN1457–913, the sequence encoding amino acids 457–913 was PCR-amplified from EFPP-SUN1 using the primers SS-EFPP-SUN1457–913-F and SS-EFPP-SUN1457–913-R (Table 1), which contain 5′ BsrGI and EcoRI cut sites, respectively. SS-EFPP-SUN1457–913-F also encodes a 10-amino acid linker (GHGTGSGTS) following the BsrGI site. The PCR product was purified and digested beside SS-EFPP with BsrGI and EcoRI. Following gel purification, the digested PCR product and plasmid were ligated together to create SS-EFPP-SUN1457–913. SS-EFPP-SUN1457–913 and SS-EFPP-SUN177–913 were generated via kinase, ligase, DpnI method as described above. The forward primers used to create SS-EFPP-SUN1457–913 and SS-EFPP-SUN177–913 were SS-EFPP-SUN1702–931-F and SS-EFPP-SUN2771–913-F, respectively (Table 1). The same reverse primer, SS-EFPP-SUN1A-R (Table 1), was used for both SS-EFPP-SUN1702–931 and SS-EFPP-SUN177–913.

The cytoplasmic EFPP-tagged SUN1 and SUN2 constructs were generated via kinase, ligase, DpnI reactions as follows. EFPP-SUN2620–731 and EFPP-SUN2451–731 were made first using the primers SS2A-F and SSA-R (Table 1). Kinase, ligase, DpnI treatments were used to make EFPP-SUN2620–731 and EFPP-SUN295–731. The forward primers used to create EFPP-SUN2620–731 and EFPP-SUN295–731 were SS-EFPP-SUN2620–731-F and SS-EFPP-SUN295–731-F, respectively (Table 1). The same reverse primer, SS-EFPP-SUN1A-R (Table 1), was used for both SS-EFPP-SUN2620–731 and SS-EFPP-SUN295–731.
and SS-EGFP-SUN1\(^{520-731}\)-F, respectively (Table 1). The same reverse primer, SS-EGFP-SUN2\(^{520-731}\)-R (Table 1), was used for both EGFP-SUN\(^{261-731}\) and EGFP-SUN\(^{595-731}\). EGFP-SUN\(^{702-913}\) was generated via kinase, ligase, DpnI treatment from EGFP-SUN\(^{1457-913}\) using the primers SS-EGFP-SUN1\(^{520-731}\)-F and SS-EGFP-SUN1\(^{777-913}\)-R (Table 1).

### Transfections

Transient transfections of cDNA and shRNA constructs were performed using GenJet from SignaGen Laboratories or Lipofectamine LTX from Invitrogen according to the manufacturers’ instructions. Measurements were performed 24 or 48 h after transfection for FFS experiments in the absence or presence of shRNA, respectively. Immediately before measurement, the growth medium was replaced with Dulbecco’s phosphate-buffered saline (PBS) with calcium and magnesium from Biowhittaker. Brightness measurements in the presence of shRNA were only performed on cells expressing turboRFP.

### Fixed- and live-cell epifluorescence microscopy

All fixed-cell imaging was performed on an Eclipse Ni-E microscope driven by NIS-Elements software using a 40x/1.30 NA Plan Fluor oil immersion objective lens (Nikon Instruments, Melville, NY), a SOLA solid state white-light excitation subsystem (Lumencor), and a CoolSNAP ES2 CCD camera (Photometrics). A custom DAPI filter set (Jena, Germany) and driven by SlideBook 6.0 from 3I. All live cell epifluorescence images were acquired with a Zeiss Plan-Apo chrom 63x/1.4NA oil objective, a Sutter DG4 light source, and a BrightLine Sedat filter set optimized for DAPI, FITC, TRITC, and CY5 from Semrock was used.

**FFS measurements**

The instrumental setup has been described previously (Hur and Mueller, 2015). When measurements were performed in the presence of shRNA, a dichroic mirror with a center wavelength of 515 nm (515DCLPXR; Chroma Technology) was used to split the emission light of EGFP and turboRFP into “green” and “red” detection channels, and an additional short-pass filter centered at 512 nm (FF01-512/SP; Semrock) was added to the green channel to remove any reflected light from turboRFP. All analysis steps were performed with programs written in Research Systems IDL 8.3. The experimental two-photon spot or point-spread function (PSF) of the two-photon microscope was described using a modified squared Gaussian–Lorentzian (mGL) model (Macdonald et al., 2010). A z-scan calibration procedure was performed as previously described to determine the radial and axial beam waist (\(w_0\) and \(z_0\)) and the axial decay parameter (\(\gamma\)) (Macdonald et al., 2010), which resulted in \(z_0 = 1.02 \pm 0.1 \mu\text{m}\), \(y = 2.30 \pm 0.3\), and \(w_0 = 0.45 \pm 0.05 \mu\text{m}\). An initial z-scan passing through the nucleus of the cell generated a z-scan with \#1.5 glass coverslip bottoms from In Vitro Scientific. Cells were then washed twice with live imaging media composed of HBSS ( Gibco, Invitrogen) containing essential and nonessential MEM amino acids (Invitrogen), 2.5 g/l glucose, 2 mM glutamine, 1 mM sodium pyruvate, and 20 mM HEPES (pH 7.4) and transferred to a 37°C Okolab full-enclosure incubator (Ottaviano, Italy) with temperature control attached to a Intelligent Imaging Innovations (3I) Marianas 200 Microscopy Workstation built on a Zeiss AxioObserver Z.1 stand (Jena, Germany) and driven by SlideBook 6.0 from 3I. All live cell epifluorescence images were acquired with a Zeiss Plan-Apochromat 63x/1.4NA oil objective, a Sutter DG4 light source, and a Photometrics CoolSnap HQ2 CCD. A BrightLine Sedat filter set optimized for DAPI, FITC, TRITC, and CY5 from Semrock was used.

## Table 1: Primers used to generate the constructs used in this paper.

| Primer name | DNA sequence | 5’ RE site |
|-------------|--------------|-----------|
| SS-EGFP-F   | GTGGCTAGCGTGAGCAAGGGCGAGGAG | Nhe |
| SS-EGFP-R   | GACTGACCTTAGCATCTGCTAGCTCATG | EcoRI |
| SS-EGFP\(_2\)-F | GCTGTACAGGGCCAGGGCCAGGGTCTACAGGGGAGGGGGGTCGAGCAAGGGC | BsrGI |
| SS-EGFP\(_2\)-R | AACGGGCCGCTGCAATCATGACTTTACTTATACAGCGTCATGCC | Apal |
| SS-EGFP-SUN1\(^{457-913}\)-F | GCTGTACAGGGCCAGGGCCAGGGTCTACAGGGGAGGGGGGTCGAGCAAGGGC | BsrGI |
| SS-EGFP-SUN1\(^{457-913}\)-R | GATTTCTACTGATGGCTGTCCCTCG | EcoRI |
| SS-EGFP-SUN1\(^{702-913}\)-F | ACATCGGGATTATGTGTC | — |
| SS-EGFP-SUN1\(^{777-913}\)-F | TGTTACTTCTACAGTACC | — |
| SS-EGFP-SUN2\(^{261-731}\)-F | GCTGTACAGGGCCAGGGCCAGGGTCTACAGGGGAGGGGGGTCGAGCAAGGGC | BsrGI |
| SS-EGFP-SUN2\(^{261-731}\)-R | TTTGGAATCCCTAGTGCGAGGCTCTC | EcoRI |
| SS-EGFP-SUN2\(^{520-731}\)-F | TTTGGAATCCCTAGTGCGAGGCTCTC | — |
| SS-EGFP-SUN2\(^{595-731}\)-F | TTTGGAATCCCTAGTGCGAGGCTCTC | — |
| SS-EGFP-SUN\(_\Delta\)-R | CCGGCTCCCTGAGACC | — |
| SSA-F | GTGGCTAGCGTGAGCAAGGGCGAGGAG | Nhe |
| SSA\(_\Delta\)-R | CATGGATCGCCGCTCGGTACC | — |

The F or R in the primer name refers to forward or reverse, respectively. Restriction enzyme (RE) cut sites are underlined. The sequence encoding the linker is bolded.
fluorescence intensity profile that was fitted using a previously described model (Smith et al., 2014, 2015) to identify the fluorescence contributions from the ventral and dorsal NEs as well as the nucleoplasm. Cells with an intensity fraction of >90% of fluorescence from the dorsal and ventral NEs were selected for further FFS measurements to ensure that the influence of fluorescence from non-NE sources on the analysis is negligible (Smith et al., 2014). An FFS measurement was conducted for 60 s and a sampling time of 7 of 50 µs with the PSF centered on the ventral or dorsal NE. The collected photon counts were analyzed to extract Mandel’s Q factor, \( Q = (\Delta F)^2/\langle F \rangle - 1 \), which is a measure of the variance of the fluorescence signal with respect to the mean (Sanchez-Andres et al., 2005; Hur and Mueller, 2015; Hennen et al., 2017). Q and the brightness \( \lambda \), which is the photon count rate of a single molecule, are proportional to one another, \( Q = \gamma \lambda T \) (Sanchez-Andres et al., 2005). The gamma factor \( \gamma \) depends on the PSF and the sample geometry (Smith et al., 2014). Given that the NE is significantly thinner (~30–50 nm; Franke et al., 1981) than the two-photon spot produced by our microscope (~1 µm), the NE acts as an infinitely thin “delta (δ)-layer” (Macdonald et al., 2010). For an mGL PSF centered on a δ-layer, \( \gamma \) is 0.5 (Macdonald et al., 2010). Additional FFS experiments performed in the nuclei of cells expressing EGFP serve to establish the raw brightness \( \gamma_{\text{EGFP}} \) as previously described (Chen et al., 2003). The normalized brightness \( b \) is defined by \( b = \gamma_{\text{EGFP}}/\gamma \) a dimensionless number that describes the average oligomeric state of the labeled protein (Macdonald et al., 2013, 2014). For example, a homodimeric complex leads to \( b = 2 \). Brightness analysis for FFS experiments in the cytoplasm are performed as described above, but with an adjusted \( \gamma \) to reflect the different geometry. The value of \( \gamma_2 \) has been determined using a z-scan FFS as described earlier (Macdonald et al., 2010).

The number concentration \( N \) represents the average number of labeled protein monomers in the observation volume. Because the brightness of an individual EGFP protein is given by \( \gamma_{\text{EGFP}} \) the time-averaged fluorescence intensity of a measurement is proportional to the number concentration, \( \langle F \rangle = \lambda_{\text{EGFP}}N \). We experimentally calculate \( N \) by dividing the average intensity \( \langle F \rangle \) by \( \gamma_{\text{EGFP}} \) (Chen et al., 2003). This procedure is valid for FFS experiments in the cytoplasm as well as at the NE. The observation volume \( V_O \) is given by the overlap between the two-photon PSF and the fluorescent sample. Converting the number concentration into a molar concentration is achieved by \( c = N_{\text{Av}}(V_O N_A) \), where \( N_A \) is Avogadro’s number. The volume \( V_O \) of a cytoplasmic FFS experiment is measured using a previously published procedure (Macdonald et al., 2010). In contrast, the volume \( V_{\text{NE}} \), and therefore the labeled protein concentration of FFS experiments at the NE, cannot be determined experimentally. However, molar concentrations in the NE can be estimated as described below.

**Brightness modeling**

A monomer/dimer/trimer equilibrium reaction was used to model \( b \) as a function of \( N \) assuming 3 molecules of monomers, dimers, and trimers (\( N_1, N_2, N_3 \)) were selected by the reactions \( A + A \rightleftharpoons A_2 \) and \( A + A_2 \rightleftharpoons A_3 \) with the dissociation coefficients \( K_{D_M} \) and \( K_{D_T} \), respectively. By definition, the normalized brightness of an n-mer is given by \( b = n \). The brightness of this mixture of species is

\[
b = \frac{1}{3!} \sum_{i=1}^{3} b^i N_i / \sum_{i=1}^{3} b N_i
\]

The total number of monomeric proteins in the observation volume is \( A_0 = A + 2A_2 + 3A_3 \). The same analysis was performed for other binding equilibrium models, such as the monomer/trimer reaction \( 3A \rightleftharpoons A_3 \) with a dissociation coefficient defined by \( K_{D_T} = [A]^3 / A_3 \). A detailed description of the modeling is found in the Supplemental Materials. Fitting of the experimental data to binding models was accomplished using bootstrapping (Efron, 1992). Confidence intervals of the estimated parameters were also determined from the bootstrap algorithm.

The observation volume \( V_{\text{NE}} \) of NE measurements is small compared to the observation volume \( V_O \) of cytoplasmic FFS experiments, which is reflected in the measured \( N \). To facilitate the comparison of \( b \) changes with concentration of a protein in both compartments, it is useful to translate between the measured \( N \) in both environments. This is achieved by the molar concentration, defined by \( c = N_{\text{Av}} / (V_O^{\text{NE}} N_A) = N_{\text{NE}} / (V_{\text{NE}}^{\text{NE}} N_A) \), which demonstrates that the values of \( N \) in the cytoplasm and the NE are proportional to one another, \( N_{\text{NE}} = N_{\text{NE}} N_{\text{NE}} / V_{\text{NE}}^{\text{NE}} N_{\text{NE}} \). While the observation volume at the NE cannot be measured, it can be modeled as the product of the cross-sectional area of the PSF and the thickness \( h \) of the NE layer.

\[
V_{\text{NE}} = \frac{\pi w^2}{4} h
\]

Given previously published measurements of NE thickness (Franke et al., 1981), we assume that the NE has an average thickness of \( h = 40 \text{ nm} \), which leads to a volume of \( 6.9 \times 10^{-3} \text{ fl} \). This value is 34-fold smaller than the observation volume in the cytoplasm. Thus, the multiplication of \( N \) in the NE by 34 compensates for the difference in observation volume and determines the equivalent cytoplasmic \( N \). The observation volume in the cytoplasm with a fully embedded PSF was determined to be 0.23 fl. These values served to calculate molar dissociation coefficients, which are quoted to one significant digit to reflect the uncertainties of the estimate.

**Immunofluorescence**

Cells grown on #1.5 coverslips were fixed in −20°C methanol as previously described (Saunders et al., 2017). Coverslips were mounted on slides using Fluoromount purchased from Thermo-Fisher Scientific.

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