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The ESCRT machinery counteracts Nesprin-2G-mediated mechanical forces during nuclear envelope repair

Graphical abstract

Highlights

- Cytoskeletal forces exerted on the nucleus can rupture its membrane
- BROX is recruited to sites of rupture by the ESCRT membrane remodeling machinery
- BROX ubiquitinates the LINC complex protein Nesprin-2G, targeting it for degradation
- BROX coordinates local relaxation of mechanical stress with membrane remodeling

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In brief

Nuclear envelope (NE) integrity is essential to protect the genome from damage. Wallis et al. investigate how the NE is repaired in case of rupture. The study demonstrates that BROX is locally recruited by ESCRTs and reduces cytoskeletal stress through its action on Nesprin-2G to help reseal the ruptured membrane.
The ESCRT machinery counteracts Nesprin-2G-mediated mechanical forces during nuclear envelope repair

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SUMMARY

Transient nuclear envelope ruptures during interphase (NERDI) occur due to cytoskeletal compressive forces at sites of weakened lamina, and delayed NERDI repair results in genomic instability. Nuclear envelope (NE) sealing is completed by endosomal sorting complex required for transport (ESCRT) machinery. A key unanswered question is how local compressive forces are counteracted to allow efficient membrane resealing. Here, we identify the ESCRT-associated protein BROX as a crucial factor required to accelerate repair of the NE. Critically, BROX binds Nesprin-2G, a component of the linker of nucleoskeleton and cytoskeleton complex (LINC). This interaction promotes Nesprin-2G ubiquitination and facilitates the relaxation of mechanical stress imposed by compressive actin fibers at the rupture site. Thus, BROX rebalances excessive cytoskeletal forces in cells experiencing NE instability to promote effective NERDI repair. Our results demonstrate that BROX coordinates mechanoregulation with membrane remodeling to ensure the maintenance of nuclear-cytoplasmic compartmentalization and genomic stability.

INTRODUCTION

The nuclear compartment is highly dynamic, and its integrity is constantly challenged by mechanical forces. The response to these forces is modulated by nuclear envelope (NE)-associated proteins including the linker of nucleoskeleton and cytoskeleton complex (LINC), which transmits forces between the cytoskeleton and the nucleus (Lee and Burke, 2018). Besides compression and deformation, the NE undergoes several remodeling events, most notably its complete disassembly and reassembly during mitosis (Güttinger et al., 2009). Transient NE ruptures during interphase (NERDI) also occur upon disruption of the NE organization and mechanical stresses generated by cytoskeletal forces (De Vos et al., 2011; Hatch and Hetzer, 2016; Takaki et al., 2017; Vargas et al., 2012), and failure to repair NERDIs contributes to genomic instability (Cho et al., 2019; Denais et al., 2016; Earle et al., 2020; Irianto et al., 2017; Raab et al., 2016; Xia et al., 2018). Therefore, a tight coordination between membrane remodeling and mechanical forces is critical to preserve NE integrity and protect the genome from damage.

NE ruptures expose the chromatin-associated protein barrier to autointegration factor (BAF) and inner nuclear membrane proteins such as LEMD2, which recruits endosomal sorting complex required for transport (ESCRT) machinery to remodel the damaged membranes (Halfmann et al., 2019; Penfield et al., 2018; Thaller et al., 2019; Young et al., 2020). NERDI repair is completed by local polymerization of ESCRT-III filaments and subsequent constrictive forces supplied by the AAA-ATPase VPS4 to seal the NE gap (Denais et al., 2016; Raab et al., 2016). Despite these advances in our understanding of the mechanisms underpinning NE repair, it remains unknown how local regulation of compressive forces at the NE is coordinated with membrane remodeling to re-establish nuclear compartmentalization.

To address this question, we investigated the function of BROX, a conserved Bro1 domain protein that binds ESCRT-III (Ichioka et al., 2008). Here, we identify BROX as a NE-associated...
Figure 1. BROX is required for efficient NERDI repair

(A) Recovery of NE integrity after NERDI in cells expressing mCherry-NLS alone or with GFP-L-BROX r and treated with control (siCTRL) or BROX-specific siRNA. siCTRL, n = 61; siBROX, n = 60; siBROX + GFP-L-BROX r, n = 56; p = 0.0225 (siCTRL-siBROX), p ≤ 0.05 (siCTRL-siBROX + WT).
factor that modulates the biomechanical properties of the nuclear compartment. We demonstrate that BROX counteracts compressive cytoskeletal forces at the NE through the interaction with Nesprin-2G, thus coordinating membrane remodeling and modulation of mechanical forces to facilitate efficient NERDI repair.

RESULTS

BROX is recruited to NERDI events and is required for efficient repair

To directly assess the integrity of the nuclear compartment in cells lacking BROX, HT1080 cells stably expressing mCherry fused to a nuclear localization signal (mCherry-NLS) were treated with BROX-specific siRNA oligos and monitored using time-lapse microscopy. In this system, cytoplasmic accumulation of mCherry-NLS indicates NE rupture, and the time to restore the basal nucleo-cytoplasmic ratio of mCherry-NLS correlates with repair time (Figures 1A and 1B; Video S1). BROX depletion significantly delayed mCherry-NLS nuclear re-accumulation after NERDI as compared with control, while stable expression of siRNA-resistant BROX (GFP-L-BROX) restored repair times to control levels (Figures 1A–1C and S1A; Video S1). Importantly, GFP-L-BROX transiently accumulated at rupture sites, as denoted by DNA herniations and co-localization with mCherry-BAF foci (Denais et al., 2016; Halfmann et al., 2018; Young et al., 2020), supporting a direct role of BROX in repair function.

BROX regulates NE morphology and compressive actin cytoskeletal forces at the NE

To better understand the role of BROX in NERDI repair, steady-state changes in the nuclei of cells lacking BROX were studied. BROX depletion increased the proportion of cells showing compressive nuclear actin cables (Figures 2A, 2B, S2A, and S2B) and NE invaginations (Figures S2E–S2H). 3D reconstructions of cells stably expressing GFP-Emerin (Figure 2C) and transmission electron microscopy (TEM) of HT1080BROX(H2AX)-positive foci (Figures 1K and 1L; Video S3), BROX cells also exhibited increased levels of 53BP1 and γ-H2AX foci (Figures 1I and 1J), and similar DNA damage was observed on a second clone (HT1080BROX(C2)) (Figures S1G and S1H), strengthening previous observations that NE repair defects increase DNA damage (Cho et al., 2019; Denais et al., 2016; Irianto et al., 2017; Raab et al., 2016; Xia et al., 2018). This phenotype could be rescued by re-expression of GFP-L-BROX in HT1080BROX(C2) (Figures 1K and 1L).

We then explored the role of upstream regulators of BROX function. CHMP7 triggers the assembly of ESCRT-III subunits such as CHMP4B at NE fenestrations (Vietri et al., 2015; Olmos et al., 2016). While CHMP7 depletion prevented enrichment of GFP-L-BROX(WT) at rupture sites (Figures 1K and 1L; Video S3), BROX depletion had no effect on CHMP4B-L-GFP recruitment (Figures S1I and S1J; Video S3), demonstrating that BROX is recruited to the damaged NE by the CHMP7/ESCRT-III axis.

(B) Representative images of the quantifications shown in (A). Scale bar, 20 μm.

(C) Quantification of the recovery times for events in (A). **p = 0.0039, ***p > 0.05. Median and quartiles are shown in red and blue, respectively.

(D) Quantification of mCherry-BAF and GFP-L-BROX/WT recruitment to the NE in cells experiencing NERDI (n = 7).

(E and F) Representative images of the quantifications shown in (D) and (G), respectively. Scale bars, 20 μm (E) and 10 μm (F).

(G) Recovery of NE integrity after NERDI in cells expressing mCherry-NLS and GFP-L-BROX/WT (n = 17) or GFP-L-BROX(C408S) mutant (n = 15).

(H) Percentage of BROX-decorated NERDIs. WT, n = 71; C408S, n = 49; p < 0.0001.

(I) Quantification of 53BP1 and γH2AX-positive foci. n = 147 for each condition; ****p < 0.0001, NSp > 0.05. Median and quartiles are shown in red and blue, respectively, and cell lysates were examined by blotting with the indicated antibodies.

(J) Representative images of the quantifications shown in (I). Arrowheads show DNA damage foci. Scale bar, 15 μm.

(K) Representative images of the quantifications shown in (L). Scale bar, 5 μm.

(L) Quantification of BROX-decorated NERDIs in siRNA-treated cells co-expressing mCherry-NLS and GFP-L-BROX migrating through 4-μm constrictions. Cell lysates were examined by blotting with the indicated antibodies. siCTRL n = 65, siCHMP7 n = 59; p < 0.0001.

See also Figure S1
Figure 2. BROX regulates compressive forces at the NE

(A) Quantification of actin cables in HeLa cells expressing GFP-Lap2β treated with the indicated siRNA and stained with SiR-actin. Cell lysates were examined by blotting with the specified antibodies. siCTRL, n = 124; siBROX, n = 104; p = 0.0008.

(B) Representative images of the quantifications shown in (A). Scale bar, 10 μm.

(C) 3D reconstruction of the nucleus of a representative HT1080δBROX cell expressing GFP-Emerin. Scale bar, 5 μm.

(D) TEM of a representative HT1080δBROX cell. Scale bar, 5 μm. Insets show magnifications of the selected areas. Scale bars, (Di) 1 μm, (Dii) 500 nm.

(E) Quantification of intranuclear tube-like structures (INTs) in wild type (WT; n = 91) or BROX-depleted (δBROX; n = 63) cells imaged by TEM. p < 0.0001.

(F) Quantification of INTs in cells expressing mCherry-Emerin and imaged by confocal microscopy. Rescue experiments were performed by adding GFP-L-BROXr. WT, n = 157: δBROX, n = 109; δBROX + GFP-L-BROXr, n = 154; p < 0.0001. Cell lysates were examined by blotting with the indicated antibodies.

(G) Representative image of the quantifications shown in (F). Arrowheads show INTs. Scale bar, 5 μm.

(H) Percentage of NE ruptures in WT (n = 38) and δBROX (n = 22) cells under constant force measured with force-clamp AFM. p ≤ 0.0001.

(I) Quantification of the force required to break through the nuclear (NM, left graph) or plasma membrane (PM, right graph) in HT1080 WT, δBROX, or lamina/C-knockout (δLMNA) cells. WT-NM, n = 141; δBROX-NM, n = 173; δLMNA-NM, n = 58; WT-PM, n = 97; δBROX-PM, n = 94; ****p < 0.0001, ***p > 0.05.

(J) Analysis of NERDI frequency in HT1080 cells expressing mCherry-NLS alone or with GFP-L-BROXr and treated with the indicated siRNA. Cell lysates were examined by blotting with the specified antibodies. siCTRL, n = 1,019; siBROX, n = 914; siBROX + GFP-L-BROXr, n = 662; *p = 0.0113, N.Sp > 0.05.

See also Figure S2.
Figure 3. BROX/Nesprin-2G interaction is needed for the release of compressive forces at the NE and rupture repair
(A and B) Pull-down experiments using recombinant GST-tagged wild-type (WT) or mutant (L350A, H204A) BROX as bait and lysates from 293T cells transfected with the indicated GFP constructs as preys. Eluates were analyzed by blotting with α-GFP antibody and band intensity is plotted normalized to BROX WT signal in (B) *p = 0.0192, **p = 0.0036, ***p = 0.001, N.S. p > 0.05.
(C) Electrostatic surface representations of BROX Bro1 domain showing residues needed for binding to Nesprin-2G (red), CHMP5 (yellow), or both CHMP4B and CHMP5 (green) and conserved residues among BROX proteins in metazoan.
(D) Representative images of the quantifications shown in (E) and (F). Arrowhead highlights BROX recruitment to site of rupture. Scale bar, 10 μm.
prone to rupture and excess DNA damage (Xia et al., 2018, 2019). Therefore, extreme NE bending may contribute to the NE ruptures in BROX-depleted cells.

**The interaction between BROX and Nesprin-2G is required to release compressive forces at the NE**

A genome-wide yeast two-hybrid screen subsequently revealed a potential interaction between BROX and Nesprin-2, a component of the LINC complex (Lee and Burke, 2018). All Nesprin-2 fragments identified contained spectrin repeats 29 to 33(SR29–33), which only appear in giant Nesprin-2(Nesprin-2G) (Rajgor and Shanahan, 2013). BROX interaction with the Nesprin-2G SR29–33 region (Nesprin-2GSR29–33) was first confirmed by yeast two-hybrid (Figure S3A). We then engineered human GFP-tagged mini-Nesprin-2G (hmN2G), a functional reporter for Nesprin-2G (Luxtton et al., 2010), by adding the SR29–33 region. The resulting construct, GFP-hmN2GSR29–33 (Figure S3B), bound GST-BROX in a co-precipitation assay (Figures 3A and 3B), thus confirming the BROX/Nesprin-2G interaction in the NE native environment.

Next, we used existing BROX crystal structures to map the interaction with Nesprin-2G (Mu et al., 2012; Zhai et al., 2011). We performed alanine substitution of exposed residues (Figure 3C) and tested the resulting mutants by yeast two-hybrid (Figure S3A). A BROX residue essential for CHMP5 binding (H204) (Mu et al., 2012) was dispensable for the interaction with Nesprin-2GSR29–33 (Figure S3A), thus indicating that CHMP5 and Nesprin-2G bind overlapping but distinct surfaces on BROX, which are evolutionarily conserved. Accordingly, mutation of L350 or I354 rendered BROX unable to bind Nesprin-2GSR29–33 without altering its ability to interact with ESCR-TIII (Figures 3A and S3A). Co-precipitation experiments confirmed the specific loss of binding between GFP-hmN2GSR29–33 and BROX L350A, while GFP-hmN2GSR29–33 interaction with BROX H204A was retained (Figures 3A and 3B). Therefore, these studies identified BROX L350A as a specific tool to determine whether loss of Nesprin-2G binding impacts BROX function. Crucially, replacement of endogenous BROX with GFP-BROXL350A abolished BROX recruitment to sites of NE rupture and resulted in impaired NE resealing (Figures 3D–3F, S3C–S3E; Videos S4 and S5), thereby abolishing the phenomenon observed in BROX-depleted cells (HT1080BROXGFP-L-BROXrL350A) (Figure 3G), indicating that BROX-Nesprin-2G interaction regulates the biomechanical properties of the nucleus.

LINC-complex-mediated nuclear compression contributes to NE ruptures (Hatch and Hetzer, 2016). Therefore, we reasoned that BROX may help reduce the mechanical stress on the nucleus by counteracting Nesprin-2G-mediated compressive forces. This model predicts that the increase on compressive forces observed in nuclei of BROX-depleted cells may be reduced upon Nesprin-2G depletion. Consistently, silencing of Nesprin-2G in HT1080BROX cells reverted the increased number of INTs, NE invaginations and nuclear actin cables to wild-type levels (Figures 3H, 3I, and S3F–S3J). As a control for specificity, depletion of Nesprin-2G in wild-type HT1080 cells did not alter the NE compressive forces, as indicated by the number of actin cables observed in this condition (Figure 3I). Importantly, the frequency of NERDI events in BROX-depleted cells was reduced to control levels by co-depleting Nesprin-2G. Moreover, BROX function was dispensable in the context of NE rupture when Nesprin-2G levels were reduced (Figures 3J and 3K; Video S6). Therefore, BROX function in NE homeostasis and repair is largely restricted to the counteraction of compressive forces imposed by Nesprin-2G dysregulation.

**Nesprin-2G ubiquitination state and localization at compression sites is mediated by BROX**

We observed that GFP-hmN2GSR29–33 exhibited increased ubiquitination levels compared with GFP-hmN2G (Figure S4A), suggesting that BROX may counteract excess Nesprin-2G activity by ubiquitination. This model was supported by the specific increase of GFP-hmN2GSR29–33 ubiquitination upon FLAG-BROX/WT overexpression, while this effect was not induced by FLAG-BROXL350A or FLAG-BROXH204A (Figure 4A). Critically, although GFP-L-BROXH204A was still recruited to sites of rupture, replacement of endogenous BROX with this mutant resulted in impaired NE repair (Figures 4B, S3E, S4B, and S4C; Video S4), thus providing a strong correlation between BROX NE repair activity and Nesprin-2G ubiquitination.

The p97 AAA + ATPase complex together with the ubiquitin/proteasome system is involved in turnover of NE components (Buchwalter et al., 2019; Tsai et al., 2016). Treatment with the p97 inhibitor EeyarestatinI resulted in the accumulation of ubiquitinated GFP-hmN2GSR29–33 (Figure 4C), while inhibition of ER mannosidase with kifunensine had no significant effect, suggesting that ubiquitinated Nesprin-2G is targeted for ER-associated degradation (ERAD) without entering the ER lumen (Fagioli...
and Sitia, 2001; Wang et al., 2008). We then reasoned that BROX may regulate compressive forces at the NE by controlling steady-state levels of Nesprin-2G. In agreement with this model, stably expressed GFP-hmN2GSR29–33 was upregulated upon BROX depletion (Figure S4D), and this resulted in the accumulation of INTs decorated with Nesprin-2G (Figures 4D and S4E).

To further test our model, we developed a fluorescent sensor to visualize ubiquitin in live cells by fusing the high-affinity ubiquitin-binding SOUBA domain from UBAP1 (Agromayor et al., 2012) to GFP (GFP-SOUBA) (Figures S4F and S4G). To validate this sensor, ESCRT-mediated degradation of ubiquitinated cargo was inhibited by depleting VPS4, resulting in large cytosolic vesicular compartments decorated by GFP-SOUBA that were reminiscent of enlarged endosomal compartments (Doyotte et al., 2005) (Figure S4G). GFP-SOUBA clusters also appeared in cells treated with siRNA against UFD1, a p97 cofactor needed for ERAD-mediated proteosomal degradation (Meyer et al., 2002) (Figure S4G). We therefore hypothesized that UFD1 depletion could be used to capture transient ubiquitination events during NERDI. Accordingly, a striking perinuclear accumulation of GFP-SOUBA was observed during the time preceding a NERDI event, and this was abrogated in BROX-depleted cells (Figures 4E and 4F; Video S7). Similarly, the perinuclear GFP-SOUBA signal was reduced by Nesprin-2G co-depletion (Figure S4H), supporting BROX-dependent ubiquitination of Nesprin-2G in this context. Furthermore, perinuclear GFP-SOUBA enrichment was restricted to cells experiencing NERDIs (Figure S4I), indicating that steady-state levels of BROX at the NE (Figures S1B and S1C) induce ubiquitination in unstable nuclei, which will later rupture. Importantly, UFD1 depletion did not increase NERDI frequency (Figure S4J). GFP-SOUBA accumulation around the nucleus was associated with progressive rounding of the nucleus before a rupture event (Figure 4E; Video S7), and this change was less marked in BROX-depleted cells (Figure 4G). Increases in nuclear roundness correlate with inhibition of actomyosin contractility (Takaki et al., 2017), thus suggesting that BROX rebalances the forces exerted by the cytoskeleton on the nucleus in cells experiencing NE instability. In further support for this model, BROX-depleted cells migrating through narrow constrictions showed an increased number of GFP-hmN2GSR29–33fibers at sites of compression (Figures 4H and 4I; Video S8). These fibers are comparable to the structures observed at the surface of the nucleus in cells that have a GFP-tag added to the endogenous Nesprin-2G gene, which are mechanically linked with actin (Davidson et al., 2020). Moreover, the observed increase in compressive forces correlated with the inability of HT1080MbroX cells to repair their nuclear ruptures and with higher death frequency (Figure 4J), a phenotype that resembles the migration-induced apoptosis observed in cells with decreased levels of LaminA (Harada et al., 2014).

**DISCUSSION**

Our study demonstrates that BROX regulates NE homeostasis by restricting the cytoskeletal forces imposed by Nesprin-2G. BROX silencing destabilizes the delicate mechanical balance of the NE, favoring the prevalence of actin-driven compressive forces. This mechanical dysregulation increases the susceptibility of BROX-silenced cells to suffer NERDI and impairs the efficient repair of these ruptures, ultimately compromising genome stability. Intriguingly, haploinsufficiency of BROX is associated with familial non-medullary thyroid cancer (Pasquali et al., 2021); thus, it is conceivable that NE instability associated with BROX deficiencies may contribute to tumor cell invasion (Nader et al., 2021).

We propose a model (Figure 4K) whereby BROX is recruited to the sites of NE rupture by the CHMP7/ESCRT-III membrane repair machinery to trigger the ubiquitination of Nesprin-2G. As consequence, the LINC complex may uncouple from the cytoskeleton at sites of actin-mediated compression during NE resealing, facilitating the final closure of wounded nuclear membranes. In the absence of BROX, dysregulated Nesprin-2G activity may increase local actin-driven compression at the rupture site, generating tensile forces that pull the ruptured membranes apart. Thus, the lack of coordination between membrane

![Figure 4. BROX regulates Nesprin-2G protein levels at compression sites](image-url)
remodeling and mechanical force regulation delays NERDI repair, exposing the genome to cytoplasmic nucleases that cause genetic instability. Giant Nesprin proteins can function at the NE independently of LINC complexes by maintaining cellular connectivity across the cytoplasm (Hao et al., 2021), a function that may not be fully recapitulated by the mini-nesprin approach used in our study. Also, a clear low GFP-Nesprin signal is visible in the cytoplasm of genetically engineered cells (Davidson et al., 2020), suggesting that BROX may further regulate the overall mechanical integrity of the cell in a LINC-independent manner by stabilizing the distal cytosolic nesprin-based meshwork, which could indirectly function as a buffer against forces exerted on the nucleus.

HT1080ΔBROX cells stably expressing GFP-hmN2GSR29-33 show lower post-migration survival than parental cells expressing the same construct when migrating through narrow constrictions, suggesting a protective role of BROX in migration stresses. It has been previously shown that high contractility conditions stabilize Lamin A to shield the genome from DNA damage by retaining repair factors in the nucleus (Cho et al., 2019; Xia et al., 2019). Whether similar DNA protection mechanisms are activated in the absence of BROX needs further investigation. Of note, HT1080 cells stably expressing GFP-hmN2GSR29-33 show increased cell death when migrating through constricted microchannels as compared with standard 2D culture, indicating that stable overexpression of the mini-Nesprin-2G construct could be altering the cell’s mechanical stiffness and susceptibility to stress by perhaps modulating endogenous LINC complexes.

Recent work has proposed that the nucleus functions as a ruler that senses changes in cell shape as consequence of externally applied mechanical stimuli (Lomakin et al., 2020; Venturini et al., 2020). The unfolding of NE invaginations and wrinkles by external forces triggers calcium-dependent activation of cytosolic phospholipase A2 (cPLA2). This mechanism induces cortical myosin II contractility, allowing the cells to rapidly tune their cytoskeletal network in response to changes in environmental conditions. The aberrant nuclear morphology in BROX-silenced cells suggests that the BROX-Nesprin-2G interaction could play a major role in mechanosensing by regulating nuclear shape, perhaps modulating the calcium-dependent activation of cPLA2 and/or the induction of cortical myosin II contractility.

Limitations of the study
We have depleted BROX from cells using two complementary approaches, CRISPR and siRNA. Of the two knockout clones selected for our studies, HT1080ΔBROX-C1 has a higher DNA content than the parental HT1080 cell line, while HT1080ΔBROX-C2 has the same DNA content. Both clones show similar defects in NE morphology and DNA damage that can be rescued by re-expressing BROX and recapitulate the phenotypes observed upon siRNA-mediated BROX depletion, showing that DNA content does not influence the magnitude of the phenotypes observed. However, although the reduced breakthrough forces of the nuclear membrane observed in HT1080ΔBROX-C1 cells can be rescued by re-expressing BROX, we cannot fully exclude that DNA content influences the nuclear mechanics probed by the AFM tip, as this has only been tested with HT1080ΔBROX-C1 cells.

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SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.devcel.2021.10.022.

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AUTHOR CONTRIBUTIONS
S.S.W., L.N.V., E.O., J.M.-S., and M.A. conceived the study. E.I. and S.G.-M. designed, performed, and analyzed AFM experiments. M.A.C. and R.A.F. designed, performed, and analyzed electron microscopy experiments. S.S.W., L.N.V., E.O., and M.A.C.-G. generated tools and performed and analyzed data from all other experiments. G.R.K. and M.F. helped with the design and analysis of experiments using microchannels. M.A. and J.M.-S. analyzed data and wrote the manuscript with assistance from all other authors.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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# STAR METHODS

## KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Antibodies          |        |            |
| Rabbit polyclonal anti-BROX | Atlas Antibodies | Cat# HPA031445; RRID: AB_10602697 |
| Mouse monoclonal anti-HSP90αβ | Santa Cruz Biotechnology | Cat# sc-13119; RRID: AB_675659 |
| Rabbit polyclonal anti-CHMP7 | Proteintech | Cat# 16424-1-AP; RRID: AB_2079500 |
| Mouse monoclonal anti-GFP (clones 7.1 and 13.1) | Roche | Cat# 11814460001; RRID: AB_390913 |
| Rabbit polyclonal anti-HA | Antibodies Online | Cat# ab16048; RRID: AB_443298 |
| Rabbit polyclonal anti-UFD1L | Proteintech | Cat# 10615-1-AP; RRID: AB_2213944 |
| Rabbit polyclonal anti-53BP1 | Novus Biologicals | Cat# NB100-304; RRID: AB_10003037 |
| Rabbit monoclonal anti-Phospho-Histone H2A.X (Ser139) (20E3) | Cell Signalling | Cat#9718 |
| Rabbit polyclonal anti-Lamin B1 | Abcam | Cat# ab16048; RRID: AB_443298 |
| Secondary donkey anti-rabbit Alexa Fluor 555 | ThermoFisher Scientific | Cat# A-31572; RRID: AB_162543 |
| Secondary goat anti-mouse (H+L DyLight™ 680 4X PEG Conjugate) | Cell Signalling Technology | Cat# 5470; RRID: AB_10696895 |
| Secondary goat anti-mouse (H+L DyLight™ 800 4X PEG Conjugate) | Cell Signalling Technology | Cat# 5257; RRID: AB_10693543 |
| Secondary goat anti-rabbit (H+L DyLight™ 680 4X PEG Conjugate) | Cell Signalling Technology | Cat# 5151; RRID: AB_10697505 |
| Secondary goat anti-rabbit (H+L DyLight™ 800 4X PEG Conjugate) | Cell Signalling Technology | Cat# 7076; RRID: AB_330924 |
| Secondary goat anti-mouse (HRP-linked) | Cell Signalling Technology | Cat# 7074; RRID: AB_2099233 |
| Secondary goat anti-rabbit (HRP-linked) | Cell Signalling Technology | Cat# 7074; RRID: AB_2099233 |
| Bacterial and virus strains |        |            |
| BL21 | New England Biolabs | C2530H |
| Chemicals, peptides, and recombinant proteins |        |            |
| Polyethyleneimine | Polysciences | Cat# 23966-1 |
| Polybrene | Merck Millipore | Cat# TR-1003-G |
| Puromycin | Sigma-Aldrich | Cat# P8833 |
| G418 | Thermo Fisher Scientific | Cat# 10131035 |
| Lipofectamine 3000 | Thermo Fisher Scientific | Cat# L3000008 |
| RNAiMAX | Thermo Fisher Scientific | Cat# 13778150 |
| Dharmafect-1 | Horizon Discovery | Cat# T-2001-02 |
| Chlorophenol red-B-D-galactopyranoside (CPRG) | Sigma-Aldrich | Cat# 10884308001 |
| Reversine | Sigma-Aldrich | Cat# R3904 |
| SiR-actin | Spirochrome | Cat# SC-001 |
| Hoechst 33258 | Sigma-Aldrich | Cat# 861450 |
| Isopropyl β-D-thiogalactopyranoside (IPTG) | Generon | Cat# IB0168-25G |
| Glutathione sepharose 4B beads | GE Healthcare Life Sciences | Cat# 17075601 |
| complete protease inhibitor cocktail | Sigma-Aldrich | Cat# 00000011697498001 |
| GFP-Trap M | ChromoTek | Cat# GTM-20 |
| Eeyarestatin I | Sigma-Aldrich | Cat# E1286 |
| Kifunensine | Sigma-Aldrich | Cat# K1140 |
| Epoxy resin | TAAB | Cat# T028 |
| Uranylless stain | TAAB | Cat# S474 |
| Reynolds lead citrate | TAAB | Cat# L037 |
| Prolong Diamond Antifade Mountant | Thermo Fisher Scientific | Cat# P36965 |

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Continued

### REAGENT or RESOURCE SOURCE IDENTIFIER

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| (S)-3'-amino Blebbistatin | Cambridge Bioscience Ltd | Cat# CAY24170 |
| Y-27632 dihydrochloride | Sigma-Aldrich | Cat# Y0503 |

### Critical Commercial Assays

| Critical commercial assays | SOURCE | IDENTIFIER |
|---------------------------|--------|------------|
| ABI High-Capacity cDNA Reverse Transcription Kit | Thermo Fisher Scientific | Cat# 4368814 |

### Experimental models: cell lines

| Experimental models: cell lines | SOURCE | IDENTIFIER |
|---------------------------------|--------|------------|
| HT1080                           | ATCC   | Cat# CRL-7951; RRID: CVCL_0317 |
| HeLa                            | ATCC   | Cat# CRM-CCL-2; RRID: CVCL_0030 |
| HEK 293T                         | ATCC   | Cat# CRL-3216; RRID: CVCL_0063 |
| HT1080 LaminA/C                  | De Vos laboratory (Antwerp University) | N/A |

### Experimental models: organisms/strains

| Experimental models: organisms/strains | SOURCE | IDENTIFIER |
|----------------------------------------|--------|------------|
| Y190 Yeast Strain                      | Harper et al., 1993 | N/A |

### Oligonucleotides

| Oligonucleotides | SOURCE | IDENTIFIER |
|------------------|--------|------------|
| siNT: ON-TARGETplus Non-targeting Control siRNA | Horizon Discovery (Dharmacon) | D-001810-01-50 |
| siBROX: Hs_C1orf58_3 FlexiTube siRNA | Qiagen | SIO4234300 |
| siCHMP7: 5'-GGGAGAAG AUUGUGAAGUUDdTdT-3' | Ventimiglia et al., 2018 | Horizon Discovery (Dharmacon) Custom siRNA |
| siNesprin-2-1: 5'-GAGUGUGCG GAGGAAACUAAU-3' | Jayo et al., 2016 | Horizon Discovery (Dharmacon) Custom siRNA |
| siNesprin-2-2: 5'-GAGGAAGAG GUGCAUGUUAU-3' | Jayo et al., 2016 | Horizon Discovery (Dharmacon) Custom siRNA |
| siUFD-1: 5'-GGGAGAAG AUUGUGAAGUUDdTdT-3' | Olmos et al., 2015 | Horizon Discovery (Dharmacon) Custom siRNA |

### Recombinant DNA

| Recombinant DNA | SOURCE | IDENTIFIER |
|-----------------|--------|------------|
| pCMS28 mCherry-NLS | Ventimiglia et al, 2018 | N/A |
| pCMS28 mCherry-BAF | This paper | N/A |
| pNG72 GFP-L-BROXr | This paper | N/A |
| pNG72 GFP-L-BROXr C408S | This paper | N/A |
| pCMS28 mCherry-Emerin | Ventimiglia et al, 2018 | N/A |
| pNG72 CHMP4B-L-GFP | Ventimiglia et al, 2018 | N/A |
| YFP-LAP2β       | Carlton et al., 2012 | N/A |
| pNG72 GFP-Emerin | This paper | N/A |
| pNG72 GFP-NLS | Ventimiglia et al, 2018 | N/A |
| pGEX Empty       | Agromayor et al. 2009 | N/A |
| pGEX BROX WT     | This paper | N/A |
| pGEX BROX L350A  | This paper | N/A |
| pGEX BROX H204A  | This paper | N/A |
| pCR3.1 GFP-CHMP4B | Ventimiglia et al, 2018 | N/A |
| pCR3.1 GFP-empty | Ventimiglia et al, 2018 | N/A |
| pCR3.1 hmN2αSP29-33 | This paper | N/A |
| pNG72 GFP-L-BROXr L350A | This paper | N/A |
| KT7 BROX WT | This paper | N/A |
| KT7 BROX E137A | This paper | N/A |
| KT7 BROX H144A | This paper | N/A |
| KT7 BROX R145A | This paper | N/A |
| KT7 BROX H204A | This paper | N/A |
| KT7 BROX L208A | This paper | N/A |
| KT7 BROX L208/212D | This paper | N/A |

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RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Monica Agromayor (monica.agromayor@kcl.ac.uk).

Materials availability
Plasmids and cell lines generated in this study are available from the lead contact upon request.

Data and code availability
- All data reported in this paper will be shared by the lead contact upon request
- This paper does not report original code
- Any additional information required to reanalyse the data reported in this work paper is available from the lead contact upon request

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Cell lines
Cells were cultured at 37°C and 5% humidified CO₂. All cells were grown in Dulbecco’s Modified Eagle Medium (DMEM, Gibco, MA, USA) supplemented with 10% fetal bovine serum (FBS) (Sigma-Aldrich, MO, USA) and 20μg/mL gentamicin (Life Technologies MA, USA). Cells were regularly tested for the presence of mycoplasma. HT1080LMNA cells were a kind gift from Professor Winnock De Vos (Antwerp University, Belgium)(Robijns et al., 2016).

METHOD DETAILS

Plasmids
Wild-type BROX was amplified by polymerase chain reaction (PCR) from cDNA (GenBank: NM_144695.3). BROX point mutations and siRNA resistant forms of BROX were generated using PCR site-directed mutagenesis. BROX constructs containing a 25 nm flexible linker (GGGGSx13) and human mini-Nesprin-2G (hmN2G) were created by gene synthesis (GeneWiz). hmN2G constructs were

Continued

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| KT7 BROX L212D      | This paper | N/A        |
| KT7 BROX K322A      | This paper | N/A        |
| KT7 BROX Y348A      | This paper | N/A        |
| KT7 BROX Y348A      | This paper | N/A        |
| KT7 BROX L350A      | This paper | N/A        |
| KT7 BROX I354A      | This paper | N/A        |
| HB18 CHMP4B         | Martin-Serrano et al, 2003 | N/A        |
| HB18 CHMP6 C-terminal truncation | This paper | N/A        |
| HB18 Nesprin-2      | This paper | N/A        |
| pNG72 GFP-SOUBA     | This paper | N/A        |
| HA-Ubiquitin        | Agromayor and Martin-Serrano, 2006 | N/A        |
| pCR3.1 FLAG-BROX WT | This paper | N/A        |
| pCR3.1 FLAG-BROX H204A | This paper | N/A        |
| pCR3.1 GFP-hmN2G    | This paper | N/A        |
| pCR3.1 FLAG-BROX L350A | This paper | N/A        |

Software and algorithms
- FIJI https://fiji.sc/ N/A
- Prism 9 GraphPad N/A
- Imaris 9 Bitplane N/A
- AutoQuant X3 Media Cybernetics N/A
- UCSF Chimera https://www.cgl.ucsf.edu/chimera/ N/A
designed to match previously described mini-Nesprin-2G constructs utilising the mouse nesprin-2 sequence (Luxton et al., 2010). hmN2G contains residues 3-482, two flexible linkers (GGGGSx2) flanking a NotI site followed by residues 6562-6907. hmN2G adds the BROX binding region encompassing residues 1860-3790. The C-terminal fragment of human CHMP5 (residues 149–219) was amplified by PCR. Plasmids that express GAL4-BROX fusion proteins were derived by insertion of BROX encoding sequences into pGBK7 (Clontech, Palo Alto, California). Plasmids that express CHMP4B, CHMP5149–219 and hmN2G in yeast, fused to a VP16 activation domain and an HA tag, were derived by insertion of PCR products into pVP16/HA (Bogerd et al., 1993). The plasmid containing an HA-tagged form of ubiquitin has been described elsewhere (Treier et al., 1994). The list of plasmids used in this study is detailed in the key resources table.

**Generation of stable cell lines**

HT1080 and HeLa cells stably expressing fluorescently tagged fusion proteins were generated using MLV-based retroviruses as described previously (Ventimiglia et al., 2018). Briefly, 293T cells were transfected with 200 ng of pHIT-VSVG, 900 ng of MLV-GagPol, and 900 ng of the pCMS28 or pNG72 retroviral packaging vectors using polyethylenimine (PEI; Polysciences, Germany). Viral-containing supernatants were collected 48 hr later, mixed with 8 μg/ml polychrere (Merck Millipore, Germany) before being used to transduce target cells. Selection with puromycin (200 ng/ml, Sigma-Aldrich) or G418 (500 μg/ml, Invitrogen, ThermoFisher, MA, USA) was applied 48 hr later, and cells were passaged under continual selection. Cells stably expressing CHMP4B-L-GFP, mCherry-Emerin, mCherry-NLS, H2B-mCherry and YFP-Lap2β, have been described previously (Ventimiglia et al., 2018).

**Generation of HT1080BROX cell line**

Guide RNAs targeting the sites located in the 5’ and 3’ ends of the BROX gene locus were designed using the Zhang Lab website (http://crispr.mit.edu) and cloned into px330-U6-Chimeric_BB-CBh-hSpCas9 plasmid (Addgene plasmid #42230). The sequences are as follows (5’F-CACCGTTTTGTCTATAGAAAACATC; 5’R- AAACGATGTTTTCTATAGACAAAAC; 3’F-CACCGTCAAACCTCAA; 3’R- AAACGATGTTTTCTATAGACAAAAC; 3’R- AAACGATGTTTTCTATAGACAAAAC). HT1080 cells were co-transfected with both guide RNA containing plasmids and a pCR3.1-GFP plasmid (2.5 μg of each guide RNA plasmid and 25 ng of pCR3.1-GFP) using Lipofectamine 3000 (Invitrogen). After 48 hours, single cells expressing GFP were sorted using FACS (BD FACS Aria III, BD, NJ, USA) and recovered in DMEM supplemented with 20% FCS to facilitate cell growth. Successful clones were then analysed for BROX knockout by western blotting to confirm the loss of the endogenous protein.

**Transfections**

For transient expression of GFP-, HA- or FLAG-tagged proteins in 293T or HT1080 cells, transfections were performed using PEI. Cells were grown on 6-well plates for 24 hours to reach 70% confluency before transfection with the appropriate plasmids (to a maximum of 2 μg DNA) using 16 μg PEI. Cells were washed after 16 hours and harvested 36 hours post transfection.

For siRNA transfections, HT1080 were plated and transfected concurrently with siRNA to a final concentration of 10 nM using RNAiMAX (Invitrogen) according to manufacturer’s instructions. 48 hours post transfection the cells received a second round of siRNA at the same concentration. Cells were either imaged or fixed 16h post transfection. For partial depletion of Nesprin-2, HT1080 cells were co-transfected once with 5 nM each of custom Nesprin-2 specific oligos Nesprin-2.1 and Nesprin-2.2 for final concentration of 10 nM. HeLa cells were transfected with 100 nM of the corresponding siRNA using Dharmafect-1 (Dharmacon RNA technologies, CO, USA) according to manufacturer’s instructions. A list of siRNAs used is provided in the key resources table. Protein depletion was confirmed by western blotting or qPCR as described below.

**Western blotting**

Samples were denatured in Laemmi buffer before resolving via SDS-PAGE. Proteins were then transferred onto 0.2 μm nitrocellulose membranes (Protran, GE Healthcare) probed with the indicated primary and secondary antibodies diluted in 1% milk. A list of antibodies used is provided in the key resources table. For HRP-conjugated secondary antibodies, membranes were further treated with Amersham ECL Prime western blotting detection reagent (GE Healthcare). Membranes were visualised using Li-Cor Odyssey Infrared scanner and software.

**qPCR**

HT1080 cells were plated onto 24-well plates at a density of 3x10^4 and transfected with siRNA as described above. 24 hours post transfection, cells were harvested for RNA extraction using the RNeasy Mini Kit (Qiagen) following manufacturer’s instructions. In brief, cells were directly lysed in the plate by addition of the lysis buffer and homogenized by syringing. RNA was then purified by passing through spin columns and washing. Purified RNA then underwent reverse transcription PCR using the ABI High Capacity cDNA Reverse Transcription Kit (ThermoFisher) following manufacturer’s instructions and using the following thermocycler settings: 25 °C 10 min; 37 °C 120 min; 85 °C 5 min; 4 °C hold. qPCR was performed using the TaqMan Fast Advanced Master Mix kit according to manufacturer’s instructions with the following probes; nesprin-2 (Hs00396027_m1) (TaqMan Gene EX Assays, ThermoFisher), GAPDH (Hs99999905_m1) (TaqMan Gene EX Assays, ThermoFisher).
Yeast-two hybrid assays

**Genome-wide screen by Hybrigenics**

Full-length BROX cloned in pB27 (N-LexA-bait-C fusion) was used in an ULTImate Y2H screen against the Human Placenta_RP5 complementary DNA Gal4-activating domain-fusion library (Hybrigenics, Paris, France). Prey fragments of positive clones were amplified by PCR and sequenced at their 5’ and 3’ junctions and the resulting sequences were used to identify corresponding interacting proteins in the GenBank database via a fully automated procedure. Hybrigenics automatically computed through their algorithms a predicted biological score (PBS) for each interaction to assess the reliability of the interaction and ranked the results, depending on the confidence of binding in four categories A to D (A having the highest confidence of binding).

**Yeast two-hybrid alanine scanning**

Yeast cells (Y190)(Harper et al., 1993) were transformed with the pGBK7 and pVP16/HA-derived plasmids described above and grown for 3 days at 30°C on Sabouraud agar (SD, Sigma-Aldrich) supplemented with Leucine-Tryptophan drop-out medium (Sigma-Aldrich) to select for positive co-transformants. Protein-protein interaction was screened by β-galactosidase activity using chlorophenol red-β-D-galactopyranoside (CPRG, Roche, Switzerland). Data is represented as the mean ± SEM of three technical repeats from one representative experiment out of three independent assays and β-gal activities are expressed as absorbance units (A540 nm); the background activity in these assays (using pGBK7 and pVP16/HA transformants) is approximately 0.01 absorbance units.

**Analysis of BROX conservation and 3D modelling**

Electrostatic surface representations of BROX Bro1 domain are based on the previously described structure (PDB: 3ULY)(Mu et al., 2012) The analysis of BROX conservation across multiple species was performed using the ConSurf server(Ashkenazy et al., 2016).

3D surface modelling of BROX was achieved with UCSFChimerau(Petter et al., 2004; Sanner et al., 1996) sing the previously described structure.

**Mechanical perturbation of cells using an AFM**

HT1080 wild-type or HT1080*BROX cells stably expressing GFP-NLS and H2B-mCherry were seeded on glass-bottom dishes (WillCo Wells) and starved for 24 hours. Experiments were performed using a Bruker BioScope Resolve AFM. Silicon nitride cantilevers, with a nominal spring constant of 0.6 N/m, with four-sided pyramidal tips, with semi-included front, back, and side angles of 35°, nominal tip radius of 20 nm, and minimal tip height of 3.2 μm, were used (MLCT-Bio-DC Tip F, Bruker). These probes are specifically designed to have a low thermal sensitivity to minimize the thermal influence of fluorescent excitation. The spring constant was calibrated by thermal tuning in MilliQ water with the simple harmonic oscillator model fit in the NanoScope. During acquisition, cells were maintained at 37°C with a combination of stage and objective heaters, with the medium buffered with 50 mM HEPES. The Resolve AFM is incorporated with a Nikon Eclipse Ti inverted epifluorescence microscope with a 60x NA 1.40 oil immersion objective and a CoolLED illumination source. Cell nuclei were identified, and a constant force (100 nN, using z-feedback control) was applied perpendicular to the cell nucleus for 5 minutes. GFP-NLS expression was used to monitor nuclear envelope integrity, which was confirmed by nuclear localization of this fluorescent signal. Images were acquired every 10 seconds for 10 minutes using an Andor iXon Ultra 888. To avoid mechanical noise of filter cube turret rotation, an OpoSpt II Bypass Image Splitter (Cairn Research), was implemented, allowing the simultaneous acquisition of two different optical wavelengths images (GFP-NLS and H2B- mCherry) on either side of the camera sensor.

**Breakthrough forces experiments**

Force spectroscopy measurements on cells using high-indenting forces were previously described (Atilla-Gokcumen et al., 2014). Briefly, cells were plated on cell culture glass-bottom dishes (WillCo Wells) 24h prior the AFM experiments. Experiments were performed using a commercial Bruker BioScope Resolve AFM coupled with a Nikon Eclipse Ti inverted epifluorescence microscope. Cells were maintained at 37°C with a combination of stage and objective heaters and the medium was buffered with 50 mM HEPES.

MLCT tip F, k = 0.6 N/m, Bruker AXS, Karlsruhe, Germany, were calibrated as described before, by thermal tuning in MilliQ water with the simple harmonic oscillator model fit in the NanoScope. Optical microscopy was used to localise the AFM tip on the top of the cell cytoplasm or nucleus. To avoid overlapping regions from neighbouring cells, only isolated cells were indentated. Cells were indentated at a constant velocity of 1 μm sec-1 until the culture dish substrate was encountered (applying forces up to 120 nN).

The penetration and rupture of individual membranes within the cell by the AFM probe were observed in the discontinuity of the force versus extension trace and fingerprinted by a series of breakthrough jumps (Yokokawa et al., 2008). Traces showing two breakthrough events indicate the probe has indentated the plasma membrane, whereas traces showing more than four breakthrough jumps indicate that the probe has punctured both the plasma membrane and the NE.

**Coprecipitation assays**

**Recombinant protein pulldowns**

Competent BL21.DE3 E. coli bacteria were transformed with pGEX vectors expressing WT BROX or BROX mutants and grown in 250 mL lysogeny broth (LB) media (Sigma-Aldrich). Isopropyl β-D-thiogalactopyranoside (IPTG, Generon, UK) was added after reaching an OD600 of 0.8 to a final concentration of 0.5 mM and left overnight at 16°C to induce protein expression. Purification of recombinant proteins was performed at 4°C. Bacteria was then collected and resuspended in GEX-lysis buffer (50 mM Tris,
HT1080 cells stably expressing GFP-hmN2GSR29-33 were transfected with the corresponding siRNA as indicated above. 24 hours after the second round of siRNA transfection, cells were resuspended in FACs buffer (1% FCS in PBS) and analysed for GFP expression using a BD FACSCanto II flow cytometer (BD). Flow-cytometry data was analysed with FlowJo software (TreeStar).

**Transmission Electron Microscopy (TEM)**

For transmission electron microscopy (TEM) analysis, cells were growth on glass coverslips at a confluency of 10x10⁴ cells and fixed overnight at 4°C in 1 ml of lysis buffer (50 mM Tris, pH 7.4, 150 mM NaCl, 5 mM EDTA, 5% glycerol, 1% Triton X-100 and a protease inhibitor cocktail (complete mini-EDTA free). The lysates were sonicated and centrifugated at 15,000 rpm for 10 minutes at 4°C and the clarified supernatant were incubated with anti-GFP coupled magnetic microparticles (GFP-Trap, ChromoTek, Germany) for 2 hours. The microparticles were washed four times with wash buffer (50 mM Tris, pH 7.4, 150 mM NaCl, 5 mM EDTA, 5% glycerol, and 0.1% Triton X-100). Bound proteins were eluted in 60 μL of Laemmli buffer, boiled and analyzed with western blot. Eeyarestatin I (2 μg/ml, Sigma-Aldrich) or Kifunensine (10 μM, Sigma-Aldrich) were added 16 hours before lysis when required.

**Immunofluorescence**

HT1080 cells were grown on coverslips and fixed for 20 minutes with methanol at -20°C or 4%PFA at room temperature, depending on primary antibody used. Cells were blocked in PBS containing 1% FBS for 1 hour followed by incubation with primary antibodies for 3 hours. After washing, cells were then stained with Alexa Fluor-conjugated secondary antibodies (Thermo Fisher Scientific). In conjunction, DNA was stained with Hoechst 33258 (Sigma-Aldrich). Coverslips were finally mounted onto slides using ProLong Diamond Antifade Mountant (Thermo Fisher Scientific).

Fixed cells were imaged using either a Nikon Eclipse Ti wide-field inverted microscope equipped with a CoolSnap HQ2 CCD camera (Photometrics, AZ, USA) or a Nikon Eclipse Ti inverted CSU-X1 spinning disk confocal microscope (Nikon, Japan) equipped with an Andor Zyla VSC-10899 (Andor, UK).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

**Image analysis**

All microscopy analysis was done blind.

**NERDI analysis in unconstrained migration**

HT1080 cells stably co-expressing mCherry-NLS and the corresponding GFP-fused BROX proteins were grown on glass bottomed 24-well plates (Eppendorf, Germany) and transfected with the appropriate siRNA. An hour before imaging, cells were treated with Hoechst 33258 (1:10000) to stain DNA and consistently track nuclei in successive time frames. Cells were imaged lived for 16 hours using a 60x oil-immersion objective (NA 1.4) equipped Nikon Eclipse Ti inverted CSU-X1 spinning disk confocal microscope attached to an environmental chamber. Images were acquired every 5 minutes using an Andor Zyla VSC-10899 camera with 2x2 binning.

NERDI events were identified as abrupt drops in mCherry-NLS nuclear fluorescence intensity. Because cytoplasmic accumulation of mCherry-NLS indicates NE rupture, and repair is indicated by its re-accumulation in the nucleus, mCherry signal was quantified over time by drawing a small region of interest (ROI) both in the nucleus and cytoplasm and intensities were extracted using Fiji (Schindelin et al., 2012). The ratio of both signal intensities, expressed as arbitrary units (a.u.), was used to evaluate the rupture repair times. This considers differences in expression levels of mCherry-NLS for each cell. These values were then normalized to the
mCherry-NLS intensity of the frame prior to nucleus rupture and the first drop in mCherry-NLS intensity (rupture start) set to \( T=0 \). Recovery time was calculated by counting the number of frames it takes for the nucleo/cytoplasmic ratio to reach at least 0.9 a.u. and multiplying by the image acquisition frequency. To increase the frequency of NERDI events in this setting, cells were treated 16 hours before imaging with 0.1 \( \mu \)M of the MPS1 kinase inhibitor reverterine (Sigma-Aldrich), which causes abnormalities in chromosome segregation during mitosis that produce chromosome bridges and predisposes the NE to rupture (Maciejowski et al., 2015).

**NERDI analysis using confinement microchannels**

HT1080 or HT1080\( ^{\text{BROX}} \) cells stably co-expressing mCherry-NLS and the corresponding GFP-tagged proteins were seeded into 6-well plates and transfected with the appropriated siRNA. After 48 hours, cells were resuspended in 60 \( \mu \)L DMEM and seeded in RTV 615 clear silicone (Momentive Performance, NY, USA) microchannels bearing 4 \( \mu \)m constrictions previously mounted on 35mm glass-bottom plates (Ibidi, Germany). Fresh media was added to just cover the bottom of the channels. 6 hours later, fresh media supplemented with Hoechst 33258 (1:1000) to stain DNA was added to fully submerge the channels for 1 hour before imaging live for 16 hours using a 60x oil-immersion objective (NA 1.4) equipped Nikon Eclipse Ti inverted CSU-X1 spinning disk confocal microscope with attached environmental chamber. Z-series (1.5 \( \mu \)m step size) images were acquired every 5 minutes using an Andor Zyla VSC-10899 camera. For NERDI analysis, rupture events were identified as abrupt drops in mCherry-NLS nuclear fluorescence intensity in cells migrating through constrictions and were analysed as described above.

**Quantification of 53BP1 and \( \gamma \)H2AX foci**

HT1080 WT, \( \delta \)BROX or \( \delta \)BROX GFP-L-BROX\(^{\text{X}} \) cells were grown on coverslips, fixed and stained with with either an \( \alpha \)-53BP1 or an \( \alpha \)-\( \gamma \)H2AX antibody and Hoescht as described above. Maximum intensity projects of whole cell Z-stack images were used to identify nuclei by automated thresholding of the Hoescht signal before establishing a threshold for foci using the parental HT1080 cells. The number of foci per nucleus were automatically counted using the ‘Analyse Particles’ tool of FIJI (Schindelin et al., 2012) with a minimum particle size of 0.1.

**Quantification of actin cables**

HeLa, HT1080 and HT1080\( ^{\text{BROX}} \) cells stably expressing YFP-Lap2\( ^{\text{B}} \) were plated onto glass-bottomed 24-well plates at a density of 2.5x10\(^4\) cells per well. Cells were transfected with the appropriate siRNA as described above. Before imaging, cells were incubated with silicon-rhodamine actin (Spirochrome, CO, USA) at a final concentration of 100 nM for 60 minutes. For myosin-II inhibition experiments, cells were treated with 50 \( \mu \)M (S)-3’-amino Blebbistatin (Cambridge Bioscience Ltd) or 50 \( \mu \)M Y27632 (Sigma-Aldrich) for 1 hour before addition of Sir-actin. Z-series (0.3 \( \mu \)m step size) were captured live using either a Nikon Eclipse Ti wide-field inverted microscope equipped with a CoolSnap HQ2 CCD camera (for HeLa cells); or a Nikon Ti-eclipse inverted CSU-X1 spinning disk confocal microscope using an Andor Zyla VSC-10899 camera. Images were blinded before analysis and fields with sparse or isolated cells were chosen to aid in identifying individual cells. Actin cables were defined as linear arrays of actin crossing the dorsal surface of the nucleus and coinciding with a Lap2\( ^{\text{B}} \) positive groove.

**Quantification of NE invaginations and intranuclear tubules (INTs)**

For analysis of NE morphology, cells were plated on coverslips, fixed, and stained with an \( \alpha \)-Lamin B1 antibody as described above. Coverslips were imaged blind and presence of NE invaginations was quantified by eye based on the presence of lamina folds at the NE periphery that were present in multiple z-stacks.

For quantification of INTs, HT1080 and HT1080\( ^{\text{BROX}} \) cells stably expressing either GFP-Emerin or mCherry-Emerin plus GFP-L-BROX\(^{\text{X}} \) constructs were plated onto 24-well glass bottomed plates at a density of 2.5x10\(^4\), treated with the appropriate siRNA as described above, and imaged live. The entire volume of the cells was captured as a Z-series with 0.1 \( \mu \)m step size using a 100x oil-immersion objective (NA 1.45) equipped Nikon Eclipse Ti inverted CSU-X1 spinning disk confocal microscope with attached environmental chamber and an Andor Zyla VSC-10899 camera. INTs were quantified using the volumetric view in the Nikon Advanced Research Program (4.1) (Nikon). 3D reconstructions were performed in Imaris (Bitplane, UK) using the surfaces function.

**Quantification of fluorescence intensity profiles**

HT1080 cells stably co-expressing mCherry-Emerin along with GFP-L-BROX\( ^{\text{X}} \) WT or C408S were seeded on glass bottomed 24-well plates and transfected with BROX-specific siRNA oligos. Live cells were imaged with a 100x oil-immersion objective equipped (NA 1.45) Nikon Eclipse Ti CSU-X1 inverted spinning disk confocal microscope using an Andor Zyla VSC-10899 camera. The entire volume of the cells was captured as a Z-series with 0.1 \( \mu \)m step size before deconvolution using AutoQuant X3 Deconvolution Software (Media Cybernetics, MD, USA). mCherry-Emerin and GFP-L-BROX\(^{\text{X}} \) fluorescence was quantified using Nikon Advanced Research Program (4.1) by drawing a line through the nuclear envelope and plotting the data in Microsoft Office Excel (Microsoft, WA, USA).

**Analysis of perinuclear GFP-SOUBA accumulation and nuclear rounding quantification**

HT1080 cells stably co-expressing mCherry-NLS and GFP-SOUBA were grown, transfected and imaged as described above. Changes in nuclear roundness over time were quantified using FIJI (Schindelin et al., 2012) In brief, Hoeschst 33258 positive nuclei were used to generate a nucleus specific region of interest (ROI) which could then be measured for roundness. This step could then be repeated for each timeframe for the duration of the movie.
Quantification of hmN2G\textsuperscript{SR29-33} cables and cell death

HT1080 and HT1080\textsuperscript{BROX} cells stably expressing GFP-hmN2G\textsuperscript{SR29-33} and mCh-NLS were seeded in RTV 615 clear silicone (Motive Performance, NY, USA) microchannels bearing 4\textmu m constrictions and imaged live as described above. For the quantification of hmN2G\textsuperscript{SR29-33} cables, accumulation of GFP signal was identified in all z-stacks as cells migrated through the constriction and nuclear rupture was determined based on nuclear mCherry-NLS signal loss. Cell death was determined visually by morphological changes such as appearance of nuclear fragments, loss of nuclear compartmentalisation and cytoplasm shrinking. Dead cells were counted if they had migrated through the channel and died within 1 hour of exiting the constriction. Panel 4I includes cells that die in panel 4J.

Statistical analysis

Graphs and statistical tests were made using GraphPad Prism 9 (GraphPad, CA, USA). In all cases, error bars are mean ±SEM. Statistical tests used were two-tailed unpaired t-tests, Mann-Whitney test or one-way ANOVA with Tukey post-hoc multiple comparisons test where appropriate. All experiments are representative of at least 3 biological replicates. Details of statistical significance and information about sample size are included in their respective figure legends. N.S. p>0.05 represents no significant difference. Statistical difference is shown as *p<0.05, **p<0.01, ***p<0.001 and ****p<0.0001. For curves shown in Figures 1A, 1G, 3F, 3K, and 4B, T=0 marks the rupture start and the significance compared to the control was calculated at 35 min using a two-tailed unpaired t test.
Supplemental information

The ESCRT machinery counteracts
Nesprin-2G-mediated mechanical
forces during nuclear envelope repair

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Figure S1: BROX requires farnesylation for NE localisation and does not recruit CHMP4B to NE rupture sites. Related to Figure 1.

(A) Trace analysis of representative individual ruptures of similar size from siCTRL and siBROX treated cells and included in Figure1A.

(B-C) Analysis of BROX localisation in live HT1080 cells expressing mCherry-Emerin and GFP-L-BROX\(^\text{R}\) wild-type (WT) or non-farnesylated mutant (C408S) (B) Deconvolved images corresponding to the quantifications shown in (C). White lines mark the region of the NE where the fluorescence intensity profile was quantified. Scale bar, 10µm. (C) Fluorescence intensity profiles of HT1080 GFP-L-BROX\(^\text{R}\) WT (top) or C408S (bottom) cells.

(D-E) Time-lapse analysis of HT1080 cells expressing mCherry-NLS and GFP-L-BROX\(^\text{R}\) WT or C408S migrating through 4µm constrictions after treatment with siBROX. Hoechst was added to stain DNA. (D) Representative images of the quantifications shown in (E). Scale bar, 10µm. (E) Percentage of BROX-decorated NERDIs. GFP-L-BROX\(^\text{R}\) WT n=80; GFP-L-BROX\(^\text{R}\) C408S n=81; p<0.0001. Cell lysates were examined by blotting with the indicated antibodies.

(F) Recovery kinetics of NE integrity after NERDI in HT1080 wild-type (WT; n=29) or BROX-depleted (δBROX; n=46) cells stably expressing mCherry-NLS. Cell lysates were examined by blotting with the indicated antibodies.

(G-H) Analysis of DNA damage in HT1080 wild-type (WT) and BROX-depleted clone2 (δBROX C2) cells stained with α-53BP1 or α-γH2AX antibodies and Hoechst. (G) Quantification of 53BP1 and γH2AX -positive foci. The samples used for HT1080 WT were the same as in Figure 1I. WT n=147; HT1080\(^\delta\)BROXC2 n=147; p<0.0001. Median and quartiles are shown in red and blue respectively and cell lysates were examined by blotting with the indicated antibodies. (H) Representative images of the quantifications shown in (G). Arrowheads show DNA damage foci. Scale bar, 15µm.

(I-J) Time-lapse analysis of HT1080 cells expressing mCherry-NLS and CHMP4B-L-GFP. Cells were transfected with indicated siRNAs and imaged while migrating through 4µm constrictions. Hoechst was added to stain DNA. (I) Percentage of CHMP4B-decorated NERDI. siCTRL n=238; siCHMP7 n=81; siBROX n=137; ****p<0.0001, N.Sp>0.05. Cell lysates were analysed by blotting with the indicated antibodies. (J) Representative images of the quantification shown in (I). Scale bar, 10µm.
Figure S2: NE phenotypes resulting from BROX depletion can be rescued by re-expression of GFP-L-BROX'. Related to Figure 2.

(A-D) Analysis of actin cables in HT1080 wild-type (WT) or BROX-depleted (δBROX) stably expressing GFP-Lap2β and stained with SiR-actin. (A) Percentage of cells with nuclear actin cables. WT: n=160; δBROX n=116; p=0.0186. (B) Representative images of the quantifications shown in (A). Arrowhead denotes an actin cable. Scale bar, 5µm. (C-D) Percentage of cells with nuclear actin cables in cells treated with Y27632 or blebbistatin. siCTRL+H2O: n=675; siBROX+H2O n=611; siCTRL+Y27632: n=577; siBROX+Y27632 n=576; siCTRL+DMSO: n=680; siBROX+DMSO n=670; siCTRL+Blebb: n=570; siBROX+Blebb n=656; *p=0.0464, **p= 0.0068, ****p<0.0001

(E-F) Analysis of nuclear morphology in HT1080 cells stably expressing either GFP (GFP-ø) or GFP-L-BROX' after treatment with the indicated siRNAs and staining with α-Lamin B1 antibody. (E) Quantification of percentage of cells with NE invaginations. Non-transfected n=559; GFP-ø+siCTRL n=487; GFP-ø+siBROX n=523; GFP-L-BROX'+siCTRL n=667; GFP-L-BROX'+siBROX n=721; **p=0.0031, ***p=0.0007, N.S p>0.05. Cell lysates were analysed by blotting with the indicated antibodies. (F) Representative images of quantifications shown in (E). Arrowheads indicate NE invaginations. Scale bars, 10µm.

(G-H) Analysis of nuclear morphology in HT1080 WT and BROX-depleted cells (δBROX C1 and δBROX C2) stably expressing either GFP-ø or GFP-L-BROX' after staining with α-Lamin B1 antibody. (G) Quantification of percentage of cells with NE invaginations. WT+GFP-ø n=246; δBROX C1+GFP-ø n=283; δBROX C1+GFP-L-BROX' n=199, δBROX C2+GFP-ø n=291; δBROX C2+GFP-L-BROX' n=270; ****p<0.0001, ***p<0.001, N.S p>0.05. Cell lysates were analysed by blotting with the indicated antibodies. (H) Representative images of quantifications shown in (G). Arrowheads indicate nuclear NE invaginations. Scale bar, 10µm.

(I) Typical force-extension curves conducted on a live cell when only the plasma membrane (PM; top) or also the nuclear membrane (NM; bottom) are indented. Rupture of the plasma membrane (top inset) is hallmarked by two consecutive discontinuities in the contact region of the force-extension trace, while NE rupture can be fingerprinted by the presence of up to six discontinuities, corresponding to the rupture of the plasma membrane (two) in addition to the four nuclear membranes (bottom inset).

(J-K) Scatterplot of the breakthrough force versus the cell height in the PM or the NM. Histograms display the distribution of breakthrough forces corresponding to the rupture of the PM (J) or the NM (K). The colour-code corresponds to the type of cell analysed: WT (black), δBROX (green) and δLMNA (red). Number of traces are the same as in Figure 2I.
Figure S3: Characterisation of BROX interactions and dependence on Nesprin-2G interaction for localisation during NERDI. Related to Figure 3.

(A) BROX was tested for interactions with CHMP4B (left graph), CHMP5149-219 (middle graph) or Nesprin-2GSR29-33 (right graph) by yeast two-hybrid assays.

(B) Diagram schematic of human mini-Nesprin-2G (hmN2G) constructs used. hmN2G is based on the previously published mN2G but uses the human sequence of Nesprin-2G. hmN2GSR29-33 contains the BROX interacting region identified in the yeast-two hybrid screen flanked by flexible linkers.

(C-D) Time-lapse analysis of HT1080 cells expressing mCherry-NLS and GFP-L-BROX\(^r\) wild-type (WT) or L350A mutant cells migrating through 4\(\mu\)m constrictions. Hoechst was added to stain DNA. (C) Representative images of quantifications shown in (D). Scale bar, 10\(\mu\)m. (D) Percentage of BROX-decorated NERDI. GFP-L-BROX\(^r\) WT n=80; GFP-L-BROX\(^r\) L350A n=113; \(p=0.0058\). The dataset corresponding to GFP-L-BROX\(^r\) WT is the same used in the quantification shown in Figure S1E. Cell lysates were examined by blotting with the indicated antibodies.

(E) Quantification of the rupture recovery times in HT1080 cells expressing mCherry-NLS and GFP-L-BROX\(^r\) wild-type (WT) or mutant forms (H204A, L350A, C408S); ***\(p=0.0007\), **\(p=0.0067\), *\(p=0.0391\).

(F-G) Analysis of nuclear morphology in HT1080 wild-type (WT) or BROX-depleted (\(\delta\)BROX) cells transfected with indicated siRNAs and stained with \(\alpha\)-Lamin B1 antibody. (F) Percentage of cells with NE invaginations. WT+siCTRL n=245; WT+siNesprin-2 n=168; \(\delta\)BROX+siCTRL n=176; \(\delta\)BROX+siNesprin-2 n=193; ****\(p<0.0001\), ***\(p<0.001\). Cell lysates were analysed by blotting with the indicated antibodies. (G) Representative images of quantifications shown in (F). Arrowheads indicate NE invaginations. Scale bar, 20\(\mu\)m.

(H) HT1080 cells transfected with either control or Nesprin-2-specific siRNA were analysed for Nesprin-2 mRNA levels by qPCR. Results were normalised to GAPDH mRNA levels.

(I) Representative images of quantifications in Figure 3H. Arrowheads indicate INTs. Scale bar, 10\(\mu\)m.

(J) Representative images of quantifications in Figure 3I. Arrowhead indicates an actin cable. Scale bar, 10\(\mu\)m.
Figure S4: BROX-CHMP5 interaction is not required for BROX recruitment to rupture sites and GFP-SOUBA perinuclear rings form in cells that will undergo NERDI. Related to Figure 4.

(A) GFP pull-down experiments of 293T transiently co-expressing HA-Ubiquitin together with GFP-hmN2G$_{SR29-33}$ or GFP-hmN2G. Eluted fractions were examined by Western blot using the indicated antibodies and the ratio between HA and GFP bands intensity in the eluted fractions was plotted. $p=0.0036$

(B-C) Time-lapse analysis of HT1080 cells expressing mCherry-NLS and GFP-L-BROX$^c$ wild-type (WT) or mutant (H204). Hoechst was added to stain DNA. (B) Representative images of the quantifications shown in (C). Scale bar, 5µm. Arrowheads highlight site of rupture (C) Percentage of BROX-decorated NERDI. WT n=38; H204A n=59 cells; $p=0.0244$. The movies used to quantify recruitment for WT conditions were the same as the ones used for the recruitment quantification shown in Figure 3D. Cell lysates were examined by blotting with the indicated antibodies.

(D) FACS analysis of HT1080 cells stably expressing GFP-empty (GFP-ø) or GFP-hmN2G$_{SR29-33}$ after transfection with indicated siRNAs. Left: GFP fluorescence intensity profiles corresponding to one representative experiment. Right: quantification of GFP mean fluorescence intensity (MFI). *$p<0.05$, \textit{N.S} $p>0.05$.

(E) Quantitation of the number of INTs decorated with Nesprin-2G per cell in HT1080 and HT1080$^δ$BROX cells stably expressing GFP-hmN2G$_{SR29-33}$. WT n=16; $δ$BROX n=10; $p<0.0001$.

(F) Analysis of GFP-SOUBA expression in HT1080 cells stably expressing GFP-ø or GFP-SOUBA. Cell lysates were analysed by blotting with indicated antibodies.

(G-J) Live imaging of HT1080 cells co-expressing GFP-SOUBA and mCherry-NLS. (G) Analysis of GFP-SOUBA localisation in cells transfected with the indicated siRNAs. Scale bars, 10µm. (H) percentage of cells with perinuclear GFP-SOUBA accumulation. siCTRL n=588; siNesprin2 n=437; siUFD1 n=504; siNesprin2+siUFD1 n=468; **$p=0.025$, \textit{N.S} $p>0.05$. (I) Analysis of GFP-SOUBA perinuclear accumulation in cells undergoing NERDI. SOUBA-negative n=1857; SOUBA-positive n=88; $p<0.0001$. (J) Percentage of cells showing NERDI after transfection with indicated siRNA. siCTRL n=2016; siUFD1 n=1910, $p=0.9309$. 
