Title
Nuclear rupture at sites of high curvature compromises retention of DNA repair factors.

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The nucleus is physically linked to the cytoskeleton, adhesions, and extracellular matrix—all of which sustain forces, but their relationships to DNA damage are obscure. We show that nuclear rupture with cytoplasmic mislocalization of multiple DNA repair factors correlates with high nuclear curvature imposed by an external probe or by cell attachment to either aligned collagen fibers or stiff matrix. Mislocalization is greatly enhanced by lamin A depletion, requires hours for nuclear reentry, and correlates with an increase in pan-nucleoplasmic foci of the DNA damage marker γH2AX. Excess DNA damage is rescued in ruptured nuclei by cooverexpression of multiple DNA repair factors as well as by soft matrix or inhibition of actomyosin tension. Increased contractility has the opposite effect, and stiff tumors with low lamin A indeed exhibit increased nuclear curvature, more frequent nuclear rupture, and excess DNA damage. Additional stresses likely play a role, but the data suggest high curvature promotes nuclear rupture, which compromises retention of DNA repair factors and favors sustained damage.

Introduction

Cytoskeletal forces applied to the ECM via adhesions are also exerted on the nucleus (Wang et al., 2009). Structural proteins generally sustain such forces up to the point at which some protein or its assembly breaks; even a microtubule well known for its rigidity bends and breaks under cytoskeletal forces (Odde et al., 1999). For the nucleus, lamin intermediate filament meshworks (Turgay et al., 2017) somehow stiffen the nucleus and sustain nuclear forces (Lammerding et al., 2007). Indeed, fibroblasts mutated or deficient in lamin A/C (lamin A hereafter) exhibit nuclear rupture with mislocalization of GFP–nuclear localization signal (NLS) constructs and GFP-tagged transcription factors, at least in sparse culture on standard rigid dishes (De Vos et al., 2011). Soft gels limit cell and nuclear spreading as well as stress fiber assembly (Swift et al., 2013; Buxboim et al., 2017), and soft gels rescue mislocalization of GFP-NLS, as does F-actin inhibition (Tamiello et al., 2013). Furthermore, an acetyl-transferase inhibitor that affects cytoskeleton assembly (Oh et al., 2017) and soft gels rescue mislocalization of GFP-NLS, as does F-actin inhibition (Tamiello et al., 2013).Senescence/apoptosis that is often associated with DNA damage is additionally rescued by culturing mutant cells on almost any type of ECM rather than on rigid plastic (Hernandez et al., 2010; de la Rosa et al., 2013).

Mechanisms of nuclear rupture and DNA damage nonetheless remain unclear.

DNA damage and repair are ongoing in the nucleus, with DNA repair factors diffusing to damage sites. DNA damage requires hours for repair to a basal level following acute exposure to damage-causing agents (Burma et al., 2001; Soubeyrand et al., 2010). However, multiple DNA repair factors (e.g., BRCA1) mislocalize to cytoplasm in breast tumors, and nuclear depletion associates with DNA damage and patient survival (Alshareeda et al., 2016). Breast tumors also have low lamin A (Caponi-chichi et al., 2011), as do several other cancers (Broers et al., 1993; Kaspi et al., 2017). Mislocalization of DNA repair factors such as 53BP1 can lead to their progressive loss (Nuciforo et al., 2007), with cytoplasmic degradation of the DNA repair complex Ku70–Ku80 also suggested in senescence of nonmalignant cells on rigid plastic (Seluanov et al., 2007). Nuclear rupture is one conceivable mechanism for mislocalization of DNA repair factors and a consequent excess of DNA damage. We hypothesized that such a process occurs as a result of high nuclear curvature, with rupture frequency increased by both intracellular and extracellular structural factors that include low levels of lamin A, high actomyosin stress, and stiff ECM.
**Results and discussion**

**High-curvature probes rapidly rupture nuclei without disrupting the plasma membrane.**

Nuclei in live U2OS osteosarcoma cells were probed with atomic force microscopy (AFM) tips of either medium or high curvature (4.5-µm sphere or pyramidal tip <0.1-µm diameter; Fig. 1, A and B). The force was held constant in a poly nano-Newton (nN) range similar to the contractile forces generated by cells (Saez et al., 2005). Nuclear factors that are known to be mobile within the nucleus were observed simultaneously with probing: these included YFP-NLS and GFP fusions of DNA repair factors 53BP1 and KU80 (Fig. 1 C). Sudden mislocalization to cytoplasm was frequently evident when probing with medium curvature tips after lamin A knockdown (Fig. S1 A), whereas WT nuclei required high-curvature tips (Fig. 1, A and B, bar graph). YFP or GFP signal filled the cytoplasm for minutes even after release of the AFM tip (Fig. 1 B, inset). The images suggest an intact plasma membrane and sustained selective rupture of the nuclear envelope; in contrast, changes in nuclear signal that reverse within seconds have been attributed to distortion of nuclear pores (Elosegui-Artola et al., 2017). Furthermore, lamin A is most abundant in stiff tissues such as bone (Swift et al., 2013), which is relevant to osteosarcoma, and our probing begins to suggest that lamin A protects against curvature-induced rupture.

Sites of nuclear rupture in interphase cells are marked by focal DNA binding of cytoplasmic cyclic GMP-AMP synthase (cGAS) upon cell migration through circular pores (Harding et al., 2017) or rectangular channels with high-curvature corners (Denais et al., 2016; Raab et al., 2016). Within nonmigrating cells probed in this study by AFM, mCherry-cGAS enriches within seconds where the tip indents the nucleus (Fig. 1 D), and the mobile DNA repair factor GFP-KU80 mislocalizes to cytoplasm simultaneously with cGAS accumulation under the probe tip. Total GFP and mCherry in the cell (bottom numbers) remain constant during the entire export, indicating plasma membrane integrity. (F) A gap in nuclear envelope GFP-LBR coincides spatiotemporally with cGAS accumulation under a probe tip, indicating nuclear envelope rupture. Bars, 10 µm. Arrows in A, B, and D–F point to site of AFM tip indentation of nucleus. All data are presented as mean ± SEM.
the lamin B meshwork after constricted migration (Buxboim et al., 2017). Under AFM tips, LBR gaps are evident at sites of cGAS nuclear entry (Fig. 1 F). LBR in ER and elsewhere was otherwise unaffected by high-curved perturbations.

Collagen fiber films promote nuclear curvature and DNA repair factor mislocalization

To test the hypothesis that nuclear curvature generated by a cell—rather than by external probes, pores, or channels—accelerates nuclear rupture with mislocalization of key nuclear factors, lamin A–knockdown cells were cultured on nanofilms of collagen I fibers that align as they self-assemble on mica (Fig. 2 A; Ivanovska et al., 2017). The U2OS cells and their nuclei elongate along fibers (Figs. 2 B and S1 B), appearing similar to other mesenchyme-derived cells in tendons (Docheva et al., 2005). The nanofilms are soft only in the sense that adjacent fibers are visibly pulled off of the nonadhesive mica after 6–12 h of cell adhesion; this limits lateral spreading of a cell and thus favors a high aspect ratio (AR; Fig. 2 B). Past research of lamin A–defective fibroblasts on soft gel substrates would predict no nuclear rupture (Tamiello et al., 2013).

Immunostaining for DNA repair factor KU70 showed cytoplasmic mislocalization within cells with high-AR nuclei (Fig. 2 C). Normalized maximum curvature increases with AR (Fig. 2 D) but is lower than ellipse theory at high AR, which suggests the nucleus resists strong bending (at the poles). Circularity is more similar in experiment and theory because circularity is less sensitive to the local differences at nuclear poles. Mislocalization of GFP-KU70 accompanied enrichment of mCherry-cGAS at nuclear pole(s) (Fig. 2 E, images). Furthermore, if cGAS foci were seen in a nucleus, then the same cell showed high cytoplasmic GFP-KU70 (Fig. 2 E, bar graph). Because KU70 dimerizes with KU80, similar mislocalization phenomena are expected for both (e.g., Fig. 1 D). Nuclear curvature imposed by cell-generated forces (that are also resisted by stiff ECM) thus increases the likelihood of focal nuclear rupture. Cell and nuclear responses to stiff ECM along the fiber direction, rather than lateral softness, are consistent with cells responding to other anisotropic substrates such as rigid glass hidden below a thin soft gel (Buxboim et al., 2017).

Excess DNA damage in cells correlates with mislocalized DNA repair factors

Partial knockdown of lamin A with shLMNA was stably achieved in A549 lung carcinoma cells, whereas shLMNA-treated U2OS cells showed a growth defect (Fig. S1 C). These cells grew normally (Fig. S1 D), but ~10–20% of nuclei exhibited KU80 mislocalization in cells on rigid coverslips (Fig. S1 E). A relationship with nuclear curvature was evident (even without collagen fiber films): the lamina was disrupted at nuclear poles with enrichment of the residual lamin A and depletion of lamin B (Fig. 3 A, i and ii). Anticorrelated distributions of lamin A and B occur at sites of nuclear rupture (also at nuclear poles) during constricted migration (Harada et al., 2014; Denais et al., 2016). Ruptured A549 nuclei were also more elongated, with lower nuclear circularity compared with nonruptured nuclei, while nuclear area was constant (Fig. 3 A, iii).

Figure 2. Aligned collagen fiber film promotes nuclear curvature and DNA repair factor mislocalization. (A) AFM image of long-range alignment of collagen fibrils. (B) Both immunofluorescence and AFM probing confirm cells elongate only along the collagen fibers. (C) Immunofluorescent image and quantitation of DNA repair factor KU70 in fixed cells after 6 h seeding. Ruptured nuclei (arrow in image) show high cytoplasmic/nuclear intensity ratio (plot) and are more elongated than nonruptured nuclei (low cytoplasmic/nuclear ratio) at 6 and 12 h. n > 15 cells per point in three experiments analyzing >100 cells per experiment. (D) As nuclear AR increases, normal maximum curvature increases, and circularity decreases. Our model predicts the trend. n = 46. (E) In nonruptured nuclei, cGAS is cytoplasmic, but cGAS enriches at high-curve sites of ruptured nuclei as validated by higher cytoplasmic/nuclear GFP-KU70 (bar graph). n > 10 cells. *, P < 0.05. Bars, 10 µm. All data are presented as mean ± SEM.
When lamin-A is low in cells that spread on rigid substrates, the nuclear envelope ruptures at high curvature sites.

A  A549 shLMNA (~70% knockdown)

DNA repair and DNA damage are ongoing (Fig. 1 C), and cell growth checkpoints require low damage. Immunostaining for the DNA damage marker γH2AX (Darzynkiewicz et al., 2011) together with KU80 indeed shows KU80 mislocalization tends to correlate with excess γH2AX foci in the shLMNA cells (Fig. 3, B and C). DNA damage foci were randomly distributed throughout the nucleoplasm rather than concentrated near sites of high-curvature lamina disruption (Fig. 3 B, images). Stable expression of GFP-LMNA rescued knockdown cells, and an electrophoretic comet assay for DNA damage confirmed the imaging (Fig. S1 F). Such a distribution is consistent with impeded repair of dispersed DNA damage.

Rescue of rupture-induced DNA damage by cooverexpression of DNA repair factors

Repair of DNA requires hours after acute damage by either radiation or drugs (Staszewski et al., 2008; Álvarez-Quilón et al., 2014). While GFP-NLS type constructs reenter nuclei quickly
(<1 h) after nuclear rupture and resealing (Hatch and Hetzer, 2016; Raab et al., 2016), DNA repair factors appear mislocalized more frequently, suggesting slower relocalization (Fig. S1 G). GFP-NLS is far smaller than many repair factors (e.g., 53BP1, BRCA1, or KU80–KU70 heterodimers ∼150–250 kD), and even passive permeation through nuclear pores decreases strongly with molecular size (Samudram et al., 2016). To study the kinetics of GFP-53BP1 mislocalization, U2OS cells on rigid coverslips were imaged live for ∼10 h. Decreased nuclear circularity and increased cell spreading are seen before nuclear envelope rupture (Fig. 3 D, right), suggesting rupture at a high-curvature fluctuation in a tensed nucleus. Deep knockdown of lamin A increased fluctuation ∼5–10-fold (to ∼10%) in both live and fixed samples (Fig. S1 H), and nuclear rupture caused rapid mislocalization (<15 min) with slow recovery back into the nucleus (∼6 h; Fig. 3 D). GFP-53BP1 loss to cytoplasm was accompanied by simultaneous mislocalization of endogenous DNA repair factors upon immunostaining after live-cell imaging (KU80 in Fig. S1 I), with simultaneous mislocalization observed for all DNA repair factors examined (BRCA1, KU70, RPA2, and BRCA2 in Fig. S2 A).

If mislocalization of key repair factors shifts the steady state toward excess DNA damage (Fig. 1 C), then their simultaneous overexpression within ruptured nuclei should rescue damage. KU80 and KU70 form heterodimers and repair DNA damage in a pathway orthogonal to that of BRCA1 (Gudmundsdottir and Ashworth, 2006; Chang et al., 2017), and partial knockdown of each repair factor results in excess DNA damage that is specifically rescued by repair factor overexpression without affecting a basal damage (Fig. S2 B). Expression plasmids for these three repair factors were thus pooled for cooverexpression (denoted GFP-3) in U2OS lamin A–knockdown cells (Fig. S2 C). GFP-53BP1 was used as a negative control (Ctl) because neither its overexpression nor si53BP1 affect DNA damage (Fig. S2 D). Rupture was assessed by cytoplasmic mislocalization of endogenous DNA repair factors or GFP fusions, and the latter transfections did not alter the ∼10% of ruptured cells in fixed cultures (Fig. 4 A). Even with rupture, nuclear GFP signal was intense relative to antibody staining for repair factors. For ruptured nuclei with cytoplasmic KU80 or GFP-53BP1, DNA damage was in equal excess, but GFP-3 rescued excess damage (Fig. 4 A). For nonruptured nuclei, GFP-3 had no effect on basal DNA damage (Fig. 4 A), and so the...
three DNA repair factors are not limiting except when the nucleus ruptures.

Rescue of nuclear rupture and DNA damage by soft ECM and myosin II inhibition

Since mislocalization of DNA repair factors results from nuclear envelope rupture that relates to curvature and forces on the nucleus, reduction of such nuclear stress should in principle rescue mislocalization of DNA repair factors and thereby rescue the excess DNA damage. Soft matrices minimize actomyosin contractility and cause cells and nuclei to be less spread with lower curvature and less lamin A (Swift et al., 2013; Buxboim et al., 2017) as confirmed for WT A549 (Fig. S2 E). A549 shLMNA cells on gels that are in the range of soft tissues (3 and 0.3 kPa; Swift et al., 2013) showed minimal nuclear rupture and DNA damage (Fig. 4 B) relative to cells on stiff gels (40 kPa mimics some fibrotic tissues; Dingal et al., 2015). Soft gels minimized actomyosin assembly and nuclear curvature as expected (Fig. 4 B, bar graph), and WT A549 cells showed little evidence of nuclear rupture or excess DNA damage.

Actomyosin tension drives spreading of a cell and its nucleus on rigid culture substrates, but myosin II relaxation by blebbistatin causes rapid cell and nuclear rounding (Fig. S2 F; Buxboim et al., 2014, 2017). Such inhibition again decreased cytoplasmic mislocalization of KU80 in shLMNA A549 cells to Ctl levels and also decreased DNA damage to levels similar to soft gels based on γH2AX foci and electrophoretic comet assays (Fig. 4 C). A small, persistent excess of DNA damage in knockdown cells after relaxing nuclear curvature (by soft ECM or myosin II inhibition) is consistent with partial loss of DNA repair factors in long-term, plastic-cultured shLMNA cells (BRCA1 and 53BP1, but not KU80; Fig. S2 G), although repair factor levels are unaffected by short-term perturbations (blebbistatin and gel cultures; Fig. S2, H and I). For 53BP1, protein degradation is faster in lamin A mutant cultures (Das et al., 2013); turnover could be favored by cytoplasmic mislocalization.

Overexpression of nonmuscle myosin II in WT cells with normal lamin A levels have the opposite effects of the above rescues. WT nuclei exhibited higher curvature, more cytoplasmic mislocalization of KU80, and more DNA damage by two assays (Fig. S3). Blebbistatin again rescued the effects. The results might explain why hyperactivation of nonmuscle myosin II (by depleting a repressive phosphatase) causes nuclear dysmorphia in standard cultures and increased DNA damage (Takaki et al., 2017). Lamin A defects are therefore unnecessary for nuclear disruption and DNA damage.

Solid tumors: High-curvature nuclei, mislocalized repair factors, and excess DNA damage

ECM stiffness can be high in solid tumors (Levental et al., 2009). We hypothesized that lamin A knockdown cells exhibit some of the processes in vivo as seen in cultures on rigid substrates. Lung tumors have low lamin A (Broers et al., 1993; Kaspi et al., 2017), and so lung-derived A549 Ctrl and shLMNA cells expressing tdTomato were engrafted in mice (Fig. 5 A) followed by resection of solid tumors for quantifying nuclear shape, KU80 mislocalization, and DNA damage. Anti-KU80 showed approximately threefold higher cytoplasmic immunostaining in shLMNA tumors than Ctrl tumors (Fig. 5, B and C). Immunostaining was human specific: mouse nuclei with intense DNA chromocenters showed no anti-KU80 (Fig. 5 B, open arrow). The fraction of shLMNA cells in vivo with mislocalized KU80 was slightly lower than the same cells in vitro on rigid substrates (Fig. 5 D). A549 tumors have a stiffness between that of soft gels and rigid plastic (Swift et al., 2013), which predicts an intermediate rupture level. Lower circularity of shLMNA nuclei in vivo (Fig. 5 D) is consistent with knockdown nuclei being more deformable (Harada et al., 2014). Electrophoretic comet assays showed more DNA damage in shLMNA tumors than Ctrl tumors (Fig. 5 E), and in vivo results are again lower than for the same knockdown cells on rigid substrates but are still higher than Ctrl’s (Fig. 4 C). While additional mechanisms might explain the excess DNA damage, repair factor mislocalization does associate clinically with increased DNA damage (Alshareeda et al., 2016).

In conclusion, nuclear rupture is shown in this study to occur with higher probability at high curvature sites under external probing forces and also under actomyosin forces when cells adhere to stiff ECM (Fig. 5 F). Additional mechanical stress on a strongly curved nucleus might also be important given that curvature does not necessarily equate to either strain or stress in a given structure. Indeed, lamin A behaves in situ as a viscous fluid (Harada et al., 2014), and imposing curvature on a fluid sheet will, over time, dissipate strain and stress, while imposing curvature on a solid sheet (e.g., lamin B in situ) or even lipid bilayers (i.e., liquid crystals) should increase strain, stress, and curvature energy. The latter is useful in understanding breakage of strongly bent microtubules (Odde et al., 1999). Regardless, for medium to high nuclear curvature, lamin A has a protective role under AFM probing (Fig. 1A) and in adherent cells: in particular, shLMNA and Ctrl A549 cells show no significant difference in circularity as a measure of nuclear curvature (Fig. S2 F), and yet nuclear rupture is favored with low lamin A (Fig. 3, A–C; Fig. 4, B and C; and Fig. 5). Nuclear mechanics is complex, but Gaussian curvature (product of two principal surface curvatures at a point) is likely of greater importance than mean curvature because rigid rectangular channels with acute angles cause rupture only in small channels (Denais et al., 2016; Raab et al., 2016). Inward curvature (e.g., AFM poking of Fig. 1) might also differ from outward curvature, but asymmetric plasma membranes with a subsurface cytoskeleton are known to rupture in either case when high forces are rapidly applied to thereby increase membrane tension (Hategan et al., 2003). This is all important to understanding how cell-extrinsic mechanisms such as ECM allow lamin A mutant mice, which typically die in weeks, to achieve a normal lifespan as mosaic mice of 50% normal cells and 50% mutant cells that is sustained in all tissues (de la Rosa et al., 2013). The excess DNA damage quantified in this study indeed shows unusual upstream contributions from ECM rigidity and actomyosin contractility via mechanisms involving curvature-induced lamina rupture and loss of DNA repair factors.

Materials and methods

Cell culture

U2OS human osteosarcoma cells and A549 human lung carcinoma cells were cultured in DMEM high-glucose medium and
Ham’s F12 medium (Gibco), respectively, supplemented with 10% FBS and 1% penicillin/streptomycin (MilliporeSigma). Cells were incubated at 37°C and 5% CO₂ as recommended by ATCC. For myosin II inhibition experiments, 20 µM blebbistatin (EMD Millipore) was added to cell culture for 24 h.

Immunostaining
Cells were fixed in 4% formaldehyde (MilliporeSigma) for 15 min followed by 15-min permeabilization by 0.5% Triton X-100 (MilliporeSigma), 30-min blocking by 5% BSA (MilliporeSigma), and overnight incubation in primary antibodies at 4°C. The antibodies used include lamin A/C (1:500; mouse; sc-7292; Santa Cruz Biotechnology, Inc.), lamin A/C (1:500; goat; sc-6215; Santa Cruz Biotechnology, Inc.), lamin B (1:500; goat; sc-6217; Santa Cruz Biotechnology, Inc.), γH2AX (1:500; mouse; 05–636–1; MilliporeSigma), 53BP1 (1:300; rabbit; NB100-304; Novus Biological), KU70 (1:500; mouse; sc-17789; Santa Cruz Biotechnology, Inc.), KU80 (1:500; rabbit; C48E7; Cell Signaling Technology), BRCA1 (1:500; mouse; sc-6954; Santa Cruz Biotechnology, Inc.), BRCA2 (1:500; mouse; sc-293185; Santa Cruz Biotechnology, Inc.), cGAS (1:500; rabbit; DiD3G; Cell Signaling Technology), and collagen I (1:500; mouse; C2456; MilliporeSigma). Finally, after 90 min incubation in secondary antibodies (1:500; donkey anti-mouse, -goat, or -rabbit; Thermo Fisher Scientific), the cells’ nuclei were stained with 8 µM Hoechst 33342 (Thermo Fisher Scientific) for 15 min. When used, 1 µg/ml phalloidin-TRITC (MilliporeSigma) was added to cells for 45 min just before Hoechst staining.

Imaging
Conventional epifluorescence images were taken using an Olympus IX71 microscope with a 40×/0.6 NA objective and a digital electron-multiplying charge-coupled device camera (Cascade 512B; Photometrics). Confocal imaging was done on a Leica TCS SP8 system with a 63×/1.4 NA oil-immersion objective.

Figure 5. Solid tumors: Low-lamin A cells show higher curvature nuclei, with increased KU80 mislocalization and DNA damage. (A) A549 Ctl and shLMNA subcutaneous xenografts in NSG mice are monitored by tdTomato fluorescence. Bar, 1 cm. (B) Confocal images of A549 Ctl and shLMNA tumors. Red arrows indicate cytoplasmic KU80. The white arrow indicates mouse nucleus with chromocenters. (C) Intensity profile of a region of interest in B shows cytoplasmic mislocalization of KU80. (D) Top: Cytoplasmic/nuclear KU80 is higher in A549 shLMNA tumors versus Ctl tumors. DNA stain sets baseline for cytoplasm. Middle: Cytoplasmic/nuclear mislocalization in tumors compare well with 2D cultures. Bottom: Nuclear circularity in shLMNA cells is lower than Ctl, while nuclear areas are similar. n = 10 images per tumor from three tumors per group; n > 100 cells for circularity and area. Dark and light gray asterisks indicate significance relative to Ctl. (E) Electrophoretic comet assay shows higher DNA damage level in shLMNA tumors compared with Ctl tumors (i) and in vitro shLMNA cultures (ii) treated with blebbistatin but lower than ones treated with DMSO (data from Fig. 4 C). In vitro data are normalized to A549 Ctl cells treated with DMSO. n = 5 tumors. *, P < 0.05. Bars, 10 µm. All data are presented as mean ± SEM. (F) Schematic of processes and factors that affect nuclear envelope rupture and excess DNA damage.
Live imaging
For live imaging, we used an EVOS FL Auto Imaging System with 10× or 20× object in normal culture conditions (37°C and 5% CO2; complete culture medium as specified above).

Alkaline comet assay
The assay was performed according to manufacturer instructions (Cell Biolabs). First, cells were detached, mixed with liquefied agarose at 37°C, deposited on a specially treated glass slide, and dried for 15 min at 4°C. Next, the glass slide, containing cells in agarose gel, was incubated in lysis buffer for 45 min and alkaline solution for 30 min. Electrophoresis was conducted at 300 mA for 30 min, and then the slide was washed with 70% ethanol and air dried overnight. Finally, DNA dye was applied for 15 min, and epifluorescence images were taken as described above.

Synthesis of soft and stiff polyacrylamide gels
Round glass coverslips (18 mm; Thermo Fisher Scientific) were cleaned in boiling ethanol and RCA solution (H2O2:H2O:5%NH4OH = 2:1:1 in volume) for 10 min each, and then they were functionalized in ATCS solution (chloroform with 0.1% allyltrichlorosilane [MilliporeSigma] and 0.1% trimethylamine [MilliporeSigma]) for 1 h. Fresh precursor solution for 0.3-kPa soft gels (3% acrylamide + 0.07% bis-arylamide in deionized water) and 40-kPa stiff gels (10% acrylamide + 0.3% bis-acrylamide in deionized water) were prepared. Then, 0.1% N,N,N′,N′-tetramethylethylenediamine (MilliporeSigma) and 1% ammonium persulphate (MilliporeSigma) were added to each precursor solution, and 20 µl of the resulting mixture was added to each coverslip to allow gel polymerization. To achieve collagen I coating, cross-linker sulfo-SAN PAH (10× or 20× object in normal culture conditions (37°C and 5% CO2; complete culture medium as specified above), and over the whole gel surface and photoactivated under 365 nm UV activation, and then collagen I solution (100 µg/ml in 50 mM Hepes; G-Biosciences) was applied on the gel surface and photoactivated under 365 nm UV light for 7 min. Excess sulfo-SANPAH was washed away following UV activation, and then collagen I solution (100 µg/ml in 50 mM Hepes) was applied overnight at RT with gentle shaking.

Transfection in U2OS cells
All siRNAs used in this study were purchased from GE Healthcare (ON-TARGETplus SMARTpool siBRCA1 L-003461-00, 5′-CAA CAUGGCCCAAGCAUCA-3′, 5′-CCCAAGGCGAAGAAU-3′, 5′-UGAUAAAGCUCCAGCAGGA-3′, and 5′-GAAGGACGUUUC AUCAUUC-3′; siKu80 L-010491-00, 5′-GCA UGG AUG UGA UUC GCA CGC ACUA-3′, 5′-UGA UAA AGC UCC AGC AGGA-3′, and 5′-GAA GGA GCU UUC GGA GGU AUA-3′, and 5′-GAA GGA GCU UUC GGA GGU AUA-3′; siKu80 L-010491-00, 5′-GCA UGG AUG UGA UUC GCA CGC ACUA-3′, 5′-UGA UAA AGC UCC AGC AGGA-3′, and 5′-GAA GGA GCU UUC GGA GGU AUA-3′, and 5′-GAA GGA GCU UUC GGA GGU AUA-3′; and nontargeting siRNA D-001810-10, 5′-UUGUUUACAGUGACUAUA-3′, 5′-UGGGUUACAGUGUGU GUGA-3′, 5′-UGGGUUACAGUGUUCAGUA-3′, and 5′-UGGGUU ACAGUGUUCAGUA-3′, except for siRBP1 (5′-UAUAACCCGUCUC CUGGCACUA-3′), which was a gift from R. Greenberg (University of Pennsylvania, Philadelphia, PA; Tang et al., 2013). We do not distinguish between lamin A and lamin C because mice seem equally viable expressing either lamin A or lamin C (Fong et al., 2006; Coffinier et al., 2010). GFP-BRCA1 (71116; Addgene) was a gift from D. Durocher (Lunenfeld-Tanenbaum Research Institute, Toronto, Canada); GFP-LBR was a gift from R.-H. Chen (Academia Sinica, Taipei, Taiwan); GFP-KU70 and GFP-KU80 were gifts from S.L. Ruleni (University of Sussex, Brighton, UK; Grundy et al., 2013); and GFP-53BP1 and mCherry-cGAS were gifts from R. Greenberg (Harding et al., 2017). GFP-LMNA (Swift et al., 2013), GFP-MIIA, GFP-MIIB, and GFP-MIIA-Y278F were all used in our prior research (Shin et al., 2011). Cells were passaged 24 h before transfection. A complex of siRNA oligonucleotides (25 nM) or GFPS (0.2–0.5 ng/ml) and 1 µg/ml Lipofectamine 2000 (Invitrogen; Thermo Fisher Scientific) was prepared according to the manufacturer’s instructions and then added for 3 d (siRNAs) or 24 h (GFPS) to cells in corresponding media supplemented with 10% FBS. GFP-3 in Fig. 4 consists of GFP-KU70, GFP-KU80, and GFP-BRCA1 (0.2–0.5 ng/ml each). All plasmids are confirmed to produce specific functional proteins by Western blotting (Fig. S2) or immunofluorescence (Figs. 1, 2, 3, 4, 5, S2, and S3). Knockdown or overexpression efficiency was determined by immunoblotting or immunofluorescence following standard methods.

Viral transduction
Human LMNA shRNA lentiviral transduction plasmids were purchased from MilliporeSigma (SHCLNG-NM_170707). We transfected these plasmids into bacteria, allowed them to replicate, and then harvested them using a QIAGEN Plasmid Maxi Kit. The resulting purified plasmids were sent to The Wistar Institute (Philadelphia, PA) to produce active lentiviral particles. We then made A549 shLMNA cells according to a standard transduction protocol: ~1,000,000 cells were seeded in a six-well plate with 2 ml culture medium and then treated for 2 d with active lentiviral particles and 8 µg/ml polybrene (MilliporeSigma). Stable A549 shLMNA clones were generated after puromycin selection. For the LMNA rescue experiments, shLMNA cells were transduced by GFP-LMNA. The LMNA knockdown and rescue efficiency were determined by immunoblotting. Transduction of U2OS was performed according to the same protocol outlined in this study. Unfortunately, shLMNA U2OS cells with low lamin A levels are not stable.

Immunoblotting
For each sample, ~1,500,000 cells were trypsinized, pelleted, resuspended in 1× LDS lysis buffer supplemented with 1% protease and 1% phosphatase inhibitors, and then sonicated (3 × 15 s pulses; intermediate power setting). After resting for 30 min on ice, samples were denatured at 80°C with 0.5% β-mercaptoethanol vol/vol for 10 min. Samples were then loaded onto a bis-Tris 4–12% gradient gel (equal loading volumes across all samples; usually 4 µl per 15-well), and electrophoresis was performed (100 V × 10 min; 160 V × 55 min). Finally, proteins were transferred from the gel onto a blotting membrane (iBlot; Thermo Fisher Scientific). After blocking in 5% nonfat vol/vol for 10 min, samples were probed with antibodies for 1 h, washed, and then treated for 5 min with horseradish peroxidase conjugate. Images were acquired on a Bio-Rad ChemiDoc XRS+ (Bio-Rad Laboratories, Hercules, CA). Band intensities were quantified using ImageJ (Schneider et al., 2012) to measure band intensities relative to local background intensities.

Image analysis
Image analysis was typically performed using ImageJ (Schneider et al., 2012). For Fig. 3 C, KU80 mislocalization in ruptured nuclei results in a high cytoplasmic/nuclear ratio based on Fig. 1.

Xia et al.
High curvature correlates with nuclear rupture

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Based on Fig. 2 B, such rupture is 5–10-fold more common for shLMNA cells than either Ctl cells or rescued cells (+GFP-LMNA) that rarely rupture. For the latter, we averaged all data together, whereas for shLMNA cells, we averaged in three bins based on cytoplasmic/nuclear ratio. Bins were defined as either low bin (with similar average and width as Ctl), medium bin (based on width of Ctl), or high bin (the remaining high points). Based on this binning, high KU80 mislocalization correlates with high γH2AX foci counts. Analysis reported in Fig. 5 B (cytoplasmic KU80; bottom) was done in MatLab (MathWorks). To generate the cytoplasmic KU80 images in Fig. 5 B, we uniformly subtracted the KU80 image background intensity from every pixel, and then we removed all signal at the nuclear region, with the bounds of the nucleus determined by Hoechst staining.

Establishment of A549 tumors in vivo
For each injection, ~1,000,000 cells were suspended in 100 µl ice-cold PBS and 25% Matrigel (BD) and injected subcutaneously into the flank of nonobese diabetic/severe combined immuno-deficient mice with null expression of interleukin-2 receptor γ chain (NSG mice). Mice were obtained from the University of Pennsylvania Stem Cell and Xenograft Core. All animal experiments were planned and performed according to Institute of Animal Care and Use Committee protocols.

In vivo tumor imaging
Mice were anesthetized via inhalation of isoflurane at 3 liters/min and maintained at 1.5 liters/min. Images were acquired using a PerkinElmer IVIS Spectrum with excitation and emission filters set at 535 nm and 580 nm, respectively, optimized for tdTomato imaging. Images of each face of the sagittal plane were taken to capture both left and right flanks. Mouse fur was soaked with ethanol to reduce autofluorescence before imaging.

In vitro tumor imaging
Tumors were isolated from mice and fixed in 4% formaldehyde overnight. Fixed samples were then permeabilized by 0.5% Triton X-100 (MilliporeSigma) for 15 min, blocked by 5% BSA (MilliporeSigma) for 30 min, and incubated at 4°C overnight in primary antibodies (listed above in the Immunostaining and Imaging sections). The next day, samples were exposed to secondary antibodies (1:500; Thermo Fisher Scientific) for 90 min, and, finally, were stained with 8 µM Hoechst 33342 (Thermo Fisher Scientific) for 15 min. In taking confocal scans, we excluded the outermost one to two cell layers of each tumor chunk because the peripheral cells are most likely to be physically damaged during sample preparation.

Nuclear indentation by AFM during fluorescent imaging
U2OS cells transfected with various GFP fusion constructs and siLMNA per above or YFP-NLS U2OS were replated on coverslips that rest on an inverted optical fluorescence microscope (Olympus IX81 with 40×/0.60 NA objective). Experiments were performed in closed liquid cell at temperature of ~29°C in DMEM high-glucose medium with 10% serum buffered at pH 7.4 with 25 mM Hepes to prevent cell death in the absence of CO₂ (Frigault et al., 2009). Cells were indented using two types of cantilever probes: (1) MSCT-AUHW (Bruker) cantilevers with nominal spring constant 0.03 N/m, nominal tip radius 10–40 nm, and nominal tip height 2.5–8 µm; and (2) bead probe cantilevers (Novascan) with nominal spring constant 0.06 N/m and a polystyrene beads of 4.5-µm diameter. Before each experiment, cantilever spring constants were calibrated via the thermal fluctuations method to determine indentation forces. Then, a fluorescent cell was chosen for probing by AFM based on expression pattern: for example, expression of GFP in the nucleus without detectable GFP in the cytoplasm even for high exposure times. The cantilever was positioned on the top of the nucleus, and the nucleus was locally compressed with forces of ~10–30 nN. When the cantilever deflection reached the predefined set point, the tip would dwell on the spot for a predefined time of 100 s before the cantilever was retracted and detached from the cell. Simultaneous fluorescent images were captured every 10 or 15 s for the entire probing cycle, including before force was applied and after the cantilever was retracted for several minutes. In multiple independent experiments that cumulated to 100s of pokes on nuclei inside cells, mCherry-cGAS reliably ruptured in and GFP-KU80 and YFP-NLS cells reliably ruptured out of a major fraction of nuclei (~50%; Fig. 1, A and B), whereas GFP-53BP1 was less reliable in some experiments for unknown reasons. In addition, when a cell was poke at a position other than the nucleus, plasma membrane rupture was never observed in terms of loss of cytoplasmic fluorescent, while poking over the nucleus rarely showed cytoplasmic rupture (~1%), consistent with nucleus-specific rupture.

Collagen films self-assembly and AFM imaging
Rat-tail collagen solution (CB354249; Corning; 200 µl at 0.03 mg/ml in 50 mM glycine buffer and 200 mM KCl, pH 9.2) was deposited on freshly cleaved mica disks at room temperature for self-assembly. After 30 min, liquid was gently aspirated, and disks were transferred to Dulbecco’s PBS. Collagen films were imaged in PBS at room temperature in tapping mode with NTEGRA (NT-MDT SI), with driving frequencies close to the resonance frequency of cantilevers with a nominal spring constant of 0.1 N/m.

Cell line verification
ATCC is a biological materials resource and standards organization with external accreditation from the International Organization for Standardization, and it provides cell line authentication test recommendations per Tech Bulletin number 8 (TB-0111-00-02; year 2010). This bulletin recommends five types of tests for the authentication of cell lines. (1) Cell morphology check by microscopy, (2) growth curve analysis, and (3) mycoplasma detection by DNA staining (for filaments or extracellular particulates) were conducted on all cell lines used in this study, and all cell lines maintained the expected morphology and standard growth rates with no mycoplasma detected. Two additional ATCC-recommended authentication tests were conducted on the A549 and U2OS cell lines as they were used in all of the main and supplemental
figures. (4) Species verification of A549 cells as human was done by RNA sequencing detection of uniquely human sequences, by mass spectrometry detection of uniquely human sequences, and by antibody targeting with anti-human antibodies. (5) Identity verification from DNA of A549 was confirmed using single nucleotide polymorphism (SNP) arrays (SNPAs). DNA was isolated by using the Blood and Cell Culture DNA Mini Kit (QIAGEN) per the manufacturer’s instructions, and DNA samples were sent to the Center for Applied Genomics Core in The Children’s Hospital of Philadelphia (Philadelphia, PA) for the SNPa HumanOmniExpress-24 Bead Chip Kit (Illumina), with >700,000 probes along the entire human genome. For each sample, the Genomics Core provided the data in the form of GenomeStudio files (Illumina). Chromosome copy number was analyzed in GenomeStudio with the cnvPartition plugin (Illumina). SNPa experiments also provided genotype data, which was used to give single nucleotide variant data. Genotyping in this Illumina system relied on the correlation between total intensity and intensity ratio of the two probes, one for CG and another for AT. These correlations were mapped to a standard clustering file (Illumina) to give the SNP calls. Consistent with ATCC’s karyotype analysis, SNPAs showed that the A549 cells are hypotriploid with a chromosome copy number of 66, including two copies of X and Y chromosomes and four copies of chromosome 17. Further consistent with ATCC’s descriptions of this cell line as Caucasian in origin, SNPAs analyses show that the A549 cells are mostly European in ancestry (~90%; Dodecad2.1 an ancestry lineage algorithm that has used mostly Illumina SNPa data; Alvey et al., 2017). Similarly, U2OS cells used in our laboratory contained ~65 chromosomes by SNPAs analysis, consistent with altered chromosomes reported by ATCC, and genomic drift in terms of copy number variation was low during standard culture as clonality of U2OS cells can be maintained after single-cell cloning and continuous culture for months (Irianto et al., 2017).

Determining nuclear envelope rupture
The ratio of a protein’s total cytoplasmic intensity to its total nuclear intensity (i.e., cytoplasmic/nuclear ratio) in immunofluorescence images of fixed cells is determined by the abundance of that particular protein as well as by antibody specificity. As a result, anti-53BP1 and anti-KU80 can generate different cytoplasmic/nuclear ratio values for nonruptured nuclei in images. Based on confocal images of anti-KU80, nonruptured nuclei have a cytoplasmic/nuclear ratio of <0.1 (Irianto et al., 2017). To confirm this finding and validate the cytoplasmic/nuclear ratio as an indicator of rupture, we validated with the cells in Fig. S1 E. As shown, when the cytoplasmic/nuclear ratio is >0.1, the nuclear intensity of KU80 is low compared with nearby cells, and KU80 fills the cytoplasm and becomes visible (Fig. S1 E, arrows). Thus, the cytoplasmic/nuclear ratio reflects rupture-induced changes in protein localization, providing confidence in this ratio as a method to detect nuclear envelope rupture. To be clear, all cytoplasmic/nuclear signal used in this study is the ratio of integrated intensity in the cytoplasm over integrated intensity in the nucleus.

High, medium, or low curvature
Constricted migration through 3-µm pores causes nuclear envelope rupture in most U2OS cells, whereas 8-µm pores cause nuclear rupture in only a small fraction of cells (Irianto et al., 2017). We therefore describe AFM probes <3 µm in diameter to be high curvature, 3-7 µm to be medium curvature, and >8 µm to be low curvature.

GFP-3 rescue experiment in Fig. 4 A
U2OS siLMNA cells were plated on rigid plastic and cultured overnight, and then they were transfected with either GFP-3 (consisting of GFP-KU70, GFP-KU80, and GFP-BRCA1) or GFP-53BP1. A third Ctrl sample was not transfected; we refer to these cells as nontreated (NT). After a 24-h transfection period, all cells were fixed and immunostained for γH2AX, and the NT sample was additionally immunostained for KU80. Cytoplasmic GFP signal (or KU80 signal in the NT case) was used to identify ruptured nuclei. Foci of γH2AX were counted for ruptured and nonruptured nuclei in all three samples.

Determining pole and side regions of nuclei
ImageJ was used to first fit an ellipse for a certain nucleus, and then the ellipse was split into three regions such that the total area of the two pole regions was approximately the same as the midbody region. For ruptured nuclei, the lamin A disruption on the nuclear envelope resided in either a pole region or the side, midbody region.

Circularity, curvature, and ellipse theory
Circularity describes how the shape varies from a circle. It is a property of the entire shape and is dimensionless (i.e., it does not depend on the size of the shape):

\[ C = \frac{4\pi A}{P^2}, \]  

and the perimeter of an ellipse is given by an elliptic integral is

\[ P_{\text{ellipse}} = \int_0^{2\pi} \sqrt{a^2 \sin^2 t + b^2 \cos^2 t} \, dt. \]  

Ramanujan’s approximation to the perimeter is

\[ P_{\text{ellipse}} \approx \pi \left( 3 a + \frac{3 a^2}{2 b} + \frac{15 a^3}{16 b^2} \right). \]
\[ P_{\text{rem}} = \pi(a + b) \left(1 + \frac{3\lambda}{10 + \sqrt{4 - 3\lambda}}\right), \lambda = \frac{(1 - \frac{b}{a})^2}{(1 + \frac{b}{a})^2}, \]  

so the circularity of an ellipse can be approximated as

\[ C_{\text{ellipse}} = \frac{4ab}{(a + b)^2 \left(1 + \frac{3\lambda}{10 + \sqrt{4 - 3\lambda}}\right)^2}, \]

or in terms of the ratio of the minor and major semi-axes, \( r = a/b, \)

\[ C_{\text{ellipse}} = \frac{4ab}{(r + 2 + \frac{1}{r})^2 \left(1 + \frac{3\lambda}{10 + \sqrt{4 - 3\lambda}}\right)^2}; \]

\[ \lambda = \frac{(1 - \frac{1}{r})^2}{(1 + \frac{1}{r})^2}. \]

### Curvature of an ellipse

The curvature of an ellipse defined by Eq. 3 is

\[ \kappa_{\text{ellipse}} = \frac{ab}{(a^2 + b^2)^{3/2}}. \]

The maximum curvature is

\[ \kappa_{\text{max}} = \frac{a}{b^2}, \]

(where we have assumed without loss of generality that \( a > b > 0). \) The curvature of a circle with the same area as the ellipse defined by Eq. 3 is

\[ \kappa_{\text{circle}} = \frac{1}{\sqrt{ab}}. \]

Let’s use \( \kappa_{\text{norm}} = \left(\kappa_{\text{max}}/\kappa_{\text{circle}}\right) \) as a normalized measure of the range in curvature. In terms of \( r = b/a, \) we have

\[ \kappa_{\text{norm}} = \frac{3}{2}r^2. \]

### Measured normalized maximum curvature

ImageJ was used to mark the contour of a nucleus, and then we drew a circle of radius \( R \) to be tangent to the pole of the nucleus with highest curvature. This gives the maximum curvature

\[ \kappa = \frac{1}{R}, \]

and a few nuclei \( (n > 3) \) were also analyzed using the ImageJ plugin “kappa curvature analysis” to confirm consistency with our fitted circle approach. We normalized by

\[ \kappa_{\text{circle}} = \frac{1}{R} = 1/\sqrt{A/\pi}, \]

where \( r \) is the radius of the circle with the same area as the nucleus and \( A \) is the area of the nucleus measured by ImageJ. Thus, \( \kappa_{\text{norm}} = \kappa/\kappa_{\text{circle}}. \)

### Statistical analyses

All statistical analyses were performed using Excel (2013; Microsoft). Unless otherwise noted, statistical comparisons were made by unpaired two-tailed Student’s \( t \) tests and were considered significant if \( P < 0.05. \) Unless mentioned, all plots show mean ± SEM. \( n \) indicates the number of samples, cells, or wells quantified in each experiment. Figure legends specify the exact meaning of \( n \) for each figure (Cumming et al., 2007). For figures presenting pooled data (Fig. 5 E), all numbers are normalized to Ctrl cells that do not show variations across experiments followed by unpaired two-tailed Student’s \( t \) tests.

### Online supplemental material

Fig. S1 shows how lamin A knockdown can affect cell growth as well as sustained but rapid mislocalization of multiple repair factors (unlike YFP-NLS) and DNA damage. Fig. S2 shows how many repair factors mislocalize, how their partial knockdown increases DNA damage, and how actomyosin inhibition favors rapid nuclear rounding without affecting repair factor levels. Fig. S3 shows how increased actomyosin stress can increase nuclear curvature as well as frequency of nuclear envelope rupture and DNA damage without affecting cell cycle.

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The authors declare no competing financial interests.

### Author contributions

Y. Xia, I.L. Ivanovska, and D.E. Discher designed experiments. A.J. Liu and R.A. Greenberg contributed key materials, methods, and discussion. Y. Xia, I.L. Ivanovska, K. Zhu, L. Smith, C.M. Alvey, J. Irianto, C.R. Pfeifer, S. Cho, J. Ji, A.J. Liu, S. Cho, and R.R. Bennett performed and analyzed experiments. Y. Xia, I.L. Ivanovska, and D.E. Discher wrote the manuscript.

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Xia et al.

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