Transient nuclear envelope rupturing during interphase in human cancer cells

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Keywords:
- nuclear envelope
- nuclear permeability barrier
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- genomic instability
- cancer

Abbreviations:
- DMs, double minute chromosomes
- EM, electron microscopy
- ER, endoplasmic reticulum
- GFP, green fluorescent protein
- INM, inner nuclear membrane
- IF, immunofluorescence
- NE, nuclear envelope
- NEBD, nuclear envelope breakdown
- NEF, nuclear envelope formation
- NERDI, nuclear envelope rupture during interphase
- NPC, nuclear pore complex
- ONM, outer nuclear membrane
- TEM, transmission electron microscopy

Neoplastic cells are often characterized by specific morphological abnormalities of the nuclear envelope (NE), which have been used for cancer diagnosis for more than a century. The NE is a double phospholipid bilayer that encapsulates the nuclear genome, regulates all nuclear trafficking of RNAs and proteins and prevents the passive diffusion of macromolecules between the nucleoplasm and the cytoplasm. Whether there is a consequence to the proper functioning of the cell and loss of structural integrity of the nucleus remains unclear. Using live cell imaging, we characterize a phenomenon wherein nuclei of several proliferating human cancer cell lines become temporarily ruptured during interphase. Strikingly, NE rupturing was associated with the mislocalization of nucleoplasmic and cytoplasmic proteins and, in the most extreme cases, the entrapment of cytoplasmic organelles in the nuclear interior. In addition, we observed the formation of micronuclei-like structures during interphase and the movement of chromatin out of the nuclear space. The frequency of these NE rupturing events was higher in cells in which the nuclear lamina, a network of intermediate filaments providing mechanical support to the NE, was not properly formed. Our data uncover the existence of a NE instability that has the potential to change the genomic landscape of cancer cells.

Introduction

The nuclear envelope (NE) is a physical membrane barrier that separates the nucleus from the cytoplasm. It fulfills at least two essential functions in eukaryotic cells: first it regulates the movement of molecules between the nucleus and the cytoplasm by active, signal-dependent transport via aqueous channels that are formed by the nuclear pore complexes (NPCs), and second it creates a permeability barrier that prevents the passive diffusion of molecules larger than ~40 kDa across the NE. An intact nuclear permeability barrier is generally considered to be a prerequisite for nuclear transport and to be critical for proper cell compartmentalization.

Morphologically and structurally abnormal nuclei are frequently observed in cancer cells. Morphometric criteria such as NE invaginations, extrusions and lobes are routinely used in the clinic for cancer diagnostics and prognosis, and in some cases, karyometric features were found to be more appropriate than biomarkers to predict metastases. Despite the clinical relevance of aberrant NE morphology, it remains unclear why such changes to the NE are more prevalent in cancer cells, and if these characteristic morphological features of the NE contribute to cell transformation and tumor formation.
envelope breakdown (NEBD) occurs only at the onset of mitosis and facilitates the equal segregation of the genome and other cellular components into two daughter cells. NEBD is initiated by a series of phosphorylation events that trigger the breakdown of the NPCs and lamina and is followed by the retraction of the NE during late anaphase/telephase (Fig. S1A).

To test whether non-mitotic NE rupturing can also be visualized using this reporter, we transfected a human U2OS osteosarcoma cell line expressing GFP-NLS with a FLAGtagged version of the HIV-1 protein Vpr, which has been shown to rupture the NE during interphase leading to cell cycle arrest in G2. Non-mitotic NE rupturing can also be visualized using this reporter, we transfected a human U2OS osteosarcoma cell line expressing GFP-NLS with a FLAGtagged version of the HIV-1 protein Vpr, which has been shown to rupture the NE during interphase leading to cell cycle arrest in G2. To test whether non-mitotic NE rupturing can also be visualized using this reporter, we transfected a human U2OS osteosarcoma cell line expressing GFP-NLS with a FLAGtagged version of the HIV-1 protein Vpr, which has been shown to rupture the NE during interphase leading to cell cycle arrest in G2.

Having established a reliable assay to monitor NE integrity throughout the cell cycle, we transfected and monitored the localization of GFP-NLS in U2OS cells over a period of at least 36 h. Images were acquired with a 3 min time interval to ensure that NE integrity was observed with sufficient temporal resolution throughout the cell cycle. We observed that in a late subset of U2OS cells (~8% of cells in a population) GFP-NLS transiently appeared in the cytoplasm concomitant with a decrease in nuclear signal and in the absence of mitotic division (Fig. 1A, Vid. S1).

To determine the kinetics of interphase NE ruptures we measured the fluorescence intensity of the GFP-NLS in the nucleoplasm and cytoplasm over the course of this event. The loss of nuclear integrity was extremely rapid with virtually complete equilibration of nuclear and cytoplasmic GFP intensity occurring within a single frame (~3 min) (Vid. S1). This was followed by a slower recovery period on a timescale similar to that of post-mitotic NE reformation.

In order to examine the dynamics of individual spilling events we employed curve fitting algorithms to interphase NE rupture events imaged with high (30 sec) temporal resolution. To calculate the rate of recovery after NERDI events, we measured the nuclear and cytoplasmic intensity of GFP-NLS and fit these intensities to a segmented regression where the intact nuclei were fit to a plateau constant, spilling events were fit to a linear intensities to a segmented regression where the intact nuclei were fit to a plateau constant, spilling events were fit to a linear regression and recovery events were fit to a sigmoidal curve (mean R^2 = 0.97) (Fig. 1B). We hypothesized that recovery after NERDI fits well to a sigmoidal curve rather than an exponential curve because nuclear import likely begins before ruptures are fully closed. This analysis gave recovery half-times of ~6 min

Figure 1 (See next page). Nuclear envelope rupture during interphase. (A) U2OS cells transiently transfected with GFP-NLS and imaged every 3 min for 36 h show transient interphase rupture of the NE followed by recovery of GFP-NLS into the nucleus. (B) Dynamics of a rupture event. U2OS cells expressing GFP-NLS were imaged every 30 s to capture NERDI in high temporal resolution. GFP intensity was normalized by setting the maximum and minimum intensity for each cell to 1 and 0, respectively. Curve fittings of individual interphase NE ruptures were plotted (lines) along with raw data (points). Data was fit using the equation: Y = IF(X > X0, Y0, Ymax) where: X0 is the point of inflection between the plateau and the spilling event, Y0 is the plateau value, X1 is the initial point of recovery, S is the slope of spilling, Bottom is the lower plateau for recovery. Top is the upper plateau of recovery, Log50 is the point of 50% recovery, and HillSlope is the linear rate of recovery.

(C) Representative images of HeLa cervical and SJSA osteosarcoma cancer cell lines demonstrating spilling in diverse cancer cell types. (D) U2OS cells transiently transfected with GFP-NLS and imaged every 3 min show localized nuclear deformation and cytoplasmic GFP signal originating from the site of deformation.

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although some cells seemed to struggle to repair the NE taking up to 9 min to reach the same 50% fluorescence recovery point (Fig. 1B).

It is important to note that the frequency of NE rupturing during interphase (NERDI) did not increase with laser excitation intensity, exposure time, or expression level of the GFP<sup>+</sup>NLS

![Figure 1. For figure legend, see page 89.](image-url)
reporter (data not shown). Thus, it is unlikely that interphase loss of NE integrity is a result of experimental design or imaging. In addition, the disruption of the NE barrier did not result in apoptosis, as indicated by the ability of the cells to re-acumulate their nuclear contents, persist in culture, and go on to complete cell division and cytokinesis (Fig. 1A, right panels; Vids. S2 and S3).

To determine whether NERDI is specific to U2OS cells, we expressed and observed the dynamics of a nuclear reporter in two additional cell lines, the human H4a (cervical carcinoma) and SJSA (osteosarcoma) cell lines. We observed transient mislocalization of the reporter to the cytoplasm during interphase in both of these cell lines (Fig. 1C), indicating that NERDI is not specific to U2OS cells and affects cancer cell lines arising from divergent tissues. Importantly as a control, in primary human fibroblast IMR90 cells we observed less than 1% of cells exhibiting interphase NE rupture (Fig. 2A). Our results from the single primary cell line are limited but suggest that NERDI may be more common in malignant cells and could correlate with the well-described alterations in NE structure in cancer cells.

Analyzing still images of NERDI in U2OS cells revealed that nuclear ruptures initiate from localized deformations of the NE (Fig. 1D, large arrows; Vid. S4). These NE herniations expand and eventually rupture, with the observed efflux of GFP-NLS appearing initially in the cytoplasmic region proximal to the site of NE herniation before rapidly diffusing throughout the cell body (Fig. 1D, small arrows; Vid. S4). This observation suggests that local perturbations in nuclear integrity originate from large structural changes in the NE.

Knockdown of lamins increases the frequency of NE rupturing during interphase. Since aberrations in the nuclear lamina have been shown to affect the mechanical properties of the NE,7,8,9 and since the NE herniations observed prior to rupture are reminiscent of structures seen in cells lacking either lamin B1 or lamins A and C,10,11 we reasoned that altered lamin organization might be facilitating interphase nuclear rupturing in U2OS cells. To test this idea, we treated U2OS cells with either a scrambled siRNA or a cocktail of siRNAs directed against the three nuclear lamin genes, lamin A/C, B1, and B2, that effectively targets each lamin gene, lamin A/C, B1, and B2, that effectively targets each of the three lamins in individual cells (Fig. S2A). We found that transfection efficiency limited the population-averaged reduction in lamin protein levels to ~20% by protein gel blot (Fig. S2B), and that this reduction led to a statistically significant (p < 0.05) increase in the frequency of nuclear rupture events compared with control siRNA (Fig. 2A). In contrast, cells knocked down for the NPC components Nup93 or Nup107, or the INM proteins LBR and Lap2β in combination, had no statistically significant increase in NERDI frequency (p = 0.21, 0.94, and 0.65 respectively) (Fig. 2A). These results further support our model that NERDI is not due to changes in the passive diffusion limit, but rather to large-scale disruptions of the nuclear. Nuclear pore defects as a contributing factor to NERDI can be surmised by the rapid and mass movement of GFP-NLS into the cytoplasm, which suggests a large temporary tear in the NE rather than a continual pore-associated leakiness.

Additional evidence for lamins normally functioning to prevent NERDI came from results in transfection microscopy (TEM) performed on U2OS cells stably expressing an shRNA against lamin B1 that resulted in a similar frequency (~25%) of NERDI as the triple siRNA cocktail. In lamin B1 shRNA cells stained with tannic acid to enhance the nuclear lamina (Fig. 2B, solid arrows) nuclear herniations similar to those seen in live-imaging just prior to NE rupture are clearly visible (Fig. 2B, enlargements). Of note, areas where the NE is distended have a marked absence of electron density (Fig. 2B, open arrows), suggesting that these areas, which are prone to rupture, are deficient in lamina assembly. The interior of the NE herniations often exhibit a gradation in staining by TEM with darker nucleoplasm-like material near the nuclear interior blending with lighter cytoplasm-like material in the distal area of the herniation (Fig. 2B), as would be expected if the nuclear and cytoplasmic compartments had previously mixed at this location.

In addition to the increased NERDI frequency, the recovery time of cells treated with the triple lamin knock-down significantly increased (p < 0.05) from an average of ~6 min for control cells to ~12 min for triple lamin siRNA (Fig. 2C and D), indicating that lamins both prevent nuclear rupturing and promote the repair of NE rupturing. The ability of lamin reduction to increase the frequency of NERDI, as well as to increase the time required to recover from such defects further supports the idea that the observed phenomenon is a consequence of altered nuclear structure.

Our observations that NERDI increases with reduced lamin expression are particularly interesting since reduced and absent

Figure 2 (See next page). Reduced lamin levels accentuate nuclear ruptures. (A) Frequency of NERDI in U2OS cells after treatment with two rounds of knock down by siRNA directed against: the three lamin genes, LBR and Lap2β, Nup93, or Nup107, compared with reporter only (mock), scrambled siRNA, or non-transfected U2OS controls (p = 0.02 LaminKd vs Scr or siRNA). Cells were imaged for a period of 56 h and frequencies represent the proportion of cells that experience an interphase NE rupture at least once over the course of the experiment. (B) Transmission electron microscopy (TEM) of U2OS stably reduced for lamin B1 expression by shRNA and stained with tannic acid to enhance lamin visualization (solid arrows). Characteristic NE herniations exhibit reduced tannic acid staining (open arrow). (C) Dynamics of a rupture in U2OS cells treated with lamin siRNA. U2OS cells transfected with GFP-NLS and three lamin siRNA pool were imaged every 30 s to capture NERDI in high temporal resolution. Curve fittings, as in Figure 1B, of individual interphase NE ruptures are plotted (lines) along with raw data (points) and show the dynamics of the event. (D) Recovery half-lives of ruptures were obtained from each curve and averaged for control and three lamin siRNA treated cells with measured half-lives of ~6 and ~12 min, respectively. (E) Left: U2OS cells stably reduced for lamin B1 expression by shRNA and expressing the GFP-NLS reporter were transiently transacted with either mCherry alone or human lamin B2 tagged with mCherry (mCherry-LmnB2). Frequency of NERDI was analyzed in transfected cells imaged for 36 h. Cells with mCherry-LmnB2 aggregates were excluded from analysis. Right: Average number of spills per cell was determined for cells transfected with either mCherry or mCherry-LmnB2 and imaged for 36 h. For both, n ≥ 340 cells over two experiments. Error bars are standard error and the difference in percent cells with spilling nuclei is significant (p ≤ 0.01) by Student’s t-test.
expression of nuclear lamins has previously been reported in tumors tissue. Interestingly, we found that lamin levels were not uniform in U2OS cells; immunofluorescence using antibodies against lamin B1 and B2 revealed significant differences in lamin staining within a single field of cells (Fig. S2C). This variation, which is also present in various cell cycle phases of U2OS cells synchronized by double thymidine block (data not shown), could underlie the differences in spilling frequency we observe in normal U2OS cells. Likewise, consistent with other reports,53,54 we found that lamin protein levels also vary widely between different normal and cancer cell lines as determined by protein gel blot (Fig. S2D). Of note, several breast cancer lines exhibited lower levels of lamin expression, including those classified as particularly invasive, compared with the non-transformed MCF-10A line that retains breast tissue specific differentiation characteristics (Fig. S2D). The lines with the most reduced lamin levels were not conducive to our live-imaging approach and so the presence of interphase nuclear rupturing in them remains uncertain. Because depletion of lamin has been shown to have diverse effects on nuclear organization and functions,55,56 we wanted to ensure the phenotype we observed was the result of changes in NE structure and not downstream effects on gene expression or chromatin reorganization. To do this, we overexpressed the other B-type lamin, lamin B2, in U2OS cells depleted of lamin B1 and observed NERDI frequency. Depletion of lamin B1 alone by shRNA gave a robust knock down of lamin B1 in individual cells (Fig. S3A) and an overall reduction in lamin B1 levels by protein gel blot (Fig. S3B). Lamin B2 functions similarly to lamin B1 in structuring the NE,37,38 and our expression construct localized as expected in control U2OS cells and without disrupting endogenous lamin B1 (Fig. S2E). While lamin B2 localizes to the NE similar to lamin B1, it is unlikely to have the same protein interactions, as lamin B2 null mice have distinct phenotypes.49 We reasoned that expression of lamin B2 might compensate for the structural deficits of the lamin B1 depletion. To test this, U2OS cells stably depleted of lamin B1 were transfected with either mCherry-lamin B2 or mCherry alone. Expression of mCherry-lamin B2 was able to significantly decrease (p = 0.0059) the percentage of cells exhibiting NERDI, as well as decrease the average number of times each rupturing cell ruptured (Fig. 2E), suggesting that it is the structural function of the lamins that normally maintains interphase nuclear integrity, and not their functions in interphase nuclear organization or transcription. Interphase NE rupturing causes mislocalization of cellular components. Having established that our GFP-reporter is mislocalized to the cytoplasm during transient NE rupturing, we next wondered whether endogenous nucleoplasmic proteins would also be present in the cytoplasm during NE rupturing. Since our reporter for nuclear integrity is a soluble protein, we postulated that other soluble nuclear factors might be released into the cytoplasm during an interphase rupture event. We first characterized the localization of elf4aIII, a member of the DEAD-box family of RNA helicases and a soluble nuclear factor that is part of the exon junction complex loaded onto mRNAs inside the nucleus.57-59 We observed elf4aIII localizing in the nucleus and NE herniations during interphase by immunofluorescence in U2OS cells stably expressing an shRNA against lamin B1 (Fig. 3A, top panels arrows). In cells with ruptured NEs, we identified by cytoplasmic GFP-NLS localization and the absence of DNA condensation (Fig. 3A, bottom panels arrows), elf4aIII was mislocalized to the cytoplasm (Fig. 3A), indicating that endogenous proteins are misplaced from the nucleus during NERDI. We also observed an increase in diffuse nuclear tubulin staining concomitant with NERDI (Fig. 3A, middle panels solid vs. open arrows), suggesting that the loss of the permeability barrier across the NE occurs in both directions. In order to confirm altered localization of cytoplasmic proteins during NERDI and that loss in nuclear integrity is bidirectional, we stained fixed U2OS cells expressing the lamin B1 shRNA with antibodies against UPF1, a cytoplasmic mRNA factor that is recruited upon recognition of a stop codon by the translation machinery.60 Immunofluorescence imaging of U2OS cells stably reduced for lamin B1 to increase the frequency of NERDI showed that UPF1 is present within the interphase nucleus when nuclei undergo interphase rupture, as evidenced by GFP-NLS presence in the cytoplasm (Fig. 3B, top panels arrows). Again interphase rupture is distinguished from mitotic rupture by comparison to mitotic cells where the chromatin is clearly condensed and the mitotic spindle is visible (Fig. 3B, bottom panels arrows). Although both cytoplasmic and nuclear proteins, specifically those involved in mRNA processing, are mislocalized during transient rupturing, it is plausible that their aberrant localization is corrected either by nuclear import/export machinery or during the next division cycle. In all observed cases their mislocalization coincided with a loss of nuclear integrity, as determined by efflux of GFP-NLS from the nucleus. Therefore, it is likely that NERDI associated mislocalization of protein factors is recoverable as long as such factors carry appropriate localization signals, or in the worst case scenario factors could be reapportioned during the next mitotic cycle.
Figure 3. For figure legend, see page 93.
Since peripheral chromatin is in close proximity to the NE via interactions with the lamina and INM proteins, we wondered whether NERDI, which likely results in the disruption of these interactions, could result in the loss of genomic material from the nucleus. To test this possibility, we expressed H2B-mCherry, which localizes exclusively to chromatin throughout the cell cycle, and GFP-NLS in U2OS cells in which the three lamins had been depleted. Analysis of still images of an interphase NE rupture event clearly shows the presence of H2B-mCherry within the NE deformations, as indicated by co-localization with GFP-NLS (Fig. 3C, left panels arrows). The presence of DNA in NE herniations is also supported by the observation of Hoechst labeling of these structures in immunofluorescent images of U2OS cells stably reduced for lamin B1 expression (see Fig. 3A).

The NE deformation eventually ruptures, as indicated by the diffuse cytoplasmic GFP-NLS signal, and notably during this event H2B-mCherry also extends beyond the pre-rupture nuclear boundary, but with a more limited range (Fig. 3C, bottom panels arrows). The localization pattern of the H2B-mCherry signal is inconsistent with it being soluble and freely diffusible during the rupture (compare with GFP-NLS, Fig. 3C middle panel), and instead indicates that NERDI alters chromatin organization at the site of rupturing. After repair of the NE, indicated by nuclear accumulation of GFP-NLS, H2B-mCherry remains segregated within an extra-nuclear body (Fig. 3C, right panels arrows). Genomic instability is a hallmark of cancer and our data showing a persistent presence of chromatin outside the normal nuclear boundary make it plausible for premature rupturing of the NE to contribute to chromosome aberrations that accumulate over time in cancer cells. Our live-imaging results frequently show these extra-nuclear particles moving great distances from the post-rupture repaired NE (Fig. S3). If these particles contain genomic information, the potential for mutagenesis is virtually certain.

NE rupturing causes temporary loss of cellular compartmentalization. We next asked whether mislocalization of cytosolic components during NERDI is limited to soluble proteins by examining organelle localization during NERDI. We examined mitochondria localization by expressing pTurboRFP-mito in U2OS cells depleted of lamin B1 and expressing the integrity reporter GFP-NLS. We were able to clearly observe mitochondria present in the nucleus of cells, as indicated by 3D reconstruction (Fig. 4A, Vid. S5). In addition, cytosolic components within the nucleus were visible in knockdown cells by TEM (Fig. 4B, i and ii). These structures do not represent nuclear invaginations since they lack the double membrane with ribosome decoration that characterizes the NE (Fig. 4B, iii). Although we cannot exclude the possibility that some of these organelles are trapped during NE reformation after mitosis, it is unlikely that this is the case as nuclear envelope formation membrane recruitment to chromatin happens during a highly compacted chromatin state with ER membrane tubules being recruited to and flattening directly on the chromatin surface. To that end, attempts to physically drag mitochondria into the nucleus during NEF by tagging mitochondrial membrane proteins with chromatin binding domains failed to cause nuclear entrapment (data not shown). Our observations of mitochondria and other cytoplasmic organelles in cells with a high frequency of NERDI are consistent with results from a recent study of laminopathy cells that also found intranuclear mitochondria associated with interphase NE rupturing. The movement of organelles into the nucleus could have severe consequences for the cell since structures of this size are likely to be trapped inside the nucleus for the duration of interphase.

**Discussion**

The NE partitions the eukaryotic cell into two compartments between which there is a highly regulated exchange of proteins, nucleic acids, and cellular activities. With the delineation of specific cellular processes to distinct and separate spaces, spatial regulation is able to add to the complex series of pathways that control normal cellular physiology. Nuclear-cytoplasmic transport, which occurs through the nuclear pore complex, is an important aspect of normal cell function, and defects in this process have been reported in human genetic diseases and in divergent types of cancer. These defects can occur in the signal-transduction pathways that regulate the transfer of factors such as p53 and β-catenin in and out of the nucleus, or in the general nuclear export and import machinery itself. Our results show that the most dramatic example of compromised spatial identity in the cell may well be NERDI. We describe a phenomenon in cancer cells where the interphase NE transiently ruptures, mixing nuclear and cytoplasmic components during NERDI. We examined organelle localization during NERDI. We observed organelles present in the nucleus of cells, as indicated by 3D reconstruction (Fig. 4A, Vid. S5). In addition, cytosolic components within the nucleus were visible in knockdown cells by TEM (Fig. 4B, i and ii). These structures do not represent nuclear invaginations since they lack the double membrane with ribosome decoration that characterizes the NE (Fig. 4B, iii). Although we cannot exclude the possibility that some of these organelles are trapped during NE reformation after mitosis, it is unlikely that this is the case as nuclear envelope formation membrane recruitment to chromatin happens during a highly compacted chromatin state with ER membrane tubules being recruited to and flattening directly on the chromatin surface. To that end, attempts to physically drag mitochondria into the nucleus during NEF by tagging mitochondrial membrane proteins with chromatin binding domains failed to cause nuclear entrapment (data not shown). Our observations of mitochondria and other cytoplasmic organelles in cells with a high frequency of NERDI are consistent with results from a recent study of laminopathy cells that also found intranuclear mitochondria associated with interphase NE rupturing. The movement of organelles into the nucleus could have severe consequences for the cell since structures of this size are likely to be trapped inside the nucleus for the duration of interphase.
Figure 4. Consequences of interphase nuclear rupture. (A) U2OS cells expressing GFP-NLS (green) and pTurboRFP-mito (red), and stained for DNA with Hoechst (blue) show nuclear mitochondria in confocal slice (top), maximum intensity projection (middle) and 3D reconstruction with nuclear mitochondria indicated (arrows). (B) TEM of U2OS cells stably reduced for lamin B1 expression by shRNA showing cytoplasmic bodies enclosed within the nucleus (i and ii) contrasted from nuclear invaginations with characteristic NE double membrane and ribosome decoration (iii). (C) Proposed model for Nuclear Envelope Rupture During Interphase (NERDI) in cancer cells. Reduced lamin expression leads to a weakened NE that distends outward, eventually rupturing with a mixing of nuclear and cytoplasmic components.
was that the NE breaks down only during mitosis, when the majority of transcription is halted and when chromatin is in a highly compacted state. Realizing that, at least in some of the most frequently used cell lines, the NE transiently ruptures in a small subset of cells may lead to a new evaluation of previous results. Interphase NE rupturing suggests that additional levels of nuclear-cytoplasmic communications exist and may bring new insights to transcriptional regulation and gene expression studies or other work involving factors with regulated movement between the nuclear and cytoplasmic compartments. In addition to its role in mediating signal-dependent, active nuclear transport, a long-proposed cellular function for the evolution of the NE is to form a protective shield around the nuclear genome and thereby prevent the direct contact between cytoplasmic proteins, organelles, and metabolic by-products and the nuclear DNA. Our findings that organelles and cytoplasmic components can enter the interphase nucleus suggest that, in some cancer cells, eukaryotic cell organization is compromised and may result in insult to the genome. The presence of such components in the nucleus during interphase, when DNA is replicated and transcriptional programs are executed, could be a source of DNA damage that has not yet been appreciated. Cellular organelles such as mitochondria generate a high number of reactive oxygen species that, without the proper machinery present to neutralize them, could induce mutagenesis. Of note, the presence of cytoplasmic structures, including mitochondria, vacuoles and Golgi fragments inside the nuclei of neoplastic cells has been observed repeatedly by EM45 without a concrete explanation or mechanism for their presence. Furthermore, ‘nuclear mitochondria’ were observed in lymphoid tumors over three decades ago,46 suggesting that this phenomenon might be relevant for tumor formation in vivo. Our results provide a cellular biological explanation for the presence of these ‘nuclear cytoplasmic bodies.’

Why does NE rupturing in interphase occur in cancer cells but not, or very infrequently, in primary cells? One explanation could be that the mechanical properties of the NE in cancer cells are different from non-cancer cells. For instance, loss of lamin expression or altered lamin structure is often found in cancer cells, including leukemia and lymphoma.47-49 colon cancer,50 prostate cancer,51 gastric cancer52 and lung cancer.24,52 Our data, along with results from other labs and studies of the affect of lamin loss on nuclear elastic properties, indicate that low lamin levels or defects in chromatin organization and gene regulation.44 Since the organization of the NE at a level that has not previously been reported in refs. 18 and 20).

Materials and Methods

Cell culture. U2OS, HeLa, SJSA and MCF7 cell lines were cultured according to standard tissue culture practices and maintained in the logarithmic phase of growth in DMEM (CellGo) supplemented with 10% fetal bovine serum (HyClone), penicillin, and streptomycin. IMR90 cell line was cultured in DMEM with Glutamax (Gibco) supplemented with 20% fetal bovine serum (HyClone) and non-essential amino acids (CellGo).

siRNA transfection. Cells were transfected twice at 2d and 4d prior to analysis using 0.6 μl Lipofectamine 2000 (Invitrogen) with 25-50 nmol of the siRNA oligos: Lamin A/C (UCG UGU UC3/UCU GGA AGU CCA GTT), lamin B1 (CCU CCU CTT), scrambled (UAG ACA CCA UGC ACA AUC CTT), LBR, Lap2b, Nup93, and Nup 107 (sequences previously reported in refs. 18 and 20).

Expression constructs. GFP-NLS was constructed using the Gateway system (Invitrogen) to insert a sequence of tandem EGFPs and the NLS (PPKKRV) from the SV40 large T antigen into the N-terminal cycle3-GFP containing vector, pCDNA6/DEST35. FLG-Vpr was a generous gift of Warner Greene.
For fixed imaging, cells were grown on glass coverslips and images captured with a Leica SP2 scanning confocal microscope using SimplePCI software (Compix) or on a Zeiss/Yokagawa Scientific) and images captured with an EM CCD (Hamamatsu).

Antibodies used in this study are: rabbit anti-a-eIF4AIII and rabbit anti-a-UPF1 were kind gifts from Jens Lykke-Andersen, UC San Diego, Alex Fluor 488, goat anti-a-mouse Alexa Fluor 647, goat anti-a-rabbit Alexa Fluor 568, goat anti-a-mouse HMEC lysates were a generous gift from Clodagh O'Shea, the Salk Institute, La Jolla, CA USA, mouse anti-a-lamin A (Sigma), goat anti-a-lamin B1 (Santa Cruz), mouse anti-a-lamin B2 (Abcam), rabbit anti-a-FLAG (Cell Signaling), and mouse anti-a-tubulin (Sigma). Secondary antibodies are, from Invitrogen: goat anti-a-mouse Alexa Fluor 488, donkey anti-a-goat Alexa Fluor 488, goat anti-mouse Alexa Fluor 647, goat anti-a-rabbit Alexa Fluor 568, goat anti-a-mouse Alexa Fluor 647, goat anti-a-rabbit Alexa Fluor 647, and donkey anti-a-goat Alexa Fluor 647, and from Li-Cor: goat anti-a-mouse IRDye 680, goat anti-a-rabbit IRDye 680, donkey anti-a-goat IRDye 680, donkey anti-a-rabbit IRDye 800, donkey anti-a-goat IRDye 800.

Protein gel blotting and whole cell lysates. Whole cell lysates were collected at 70–90% confluency by washing twice in PBS, scraping in lysis buffer, and protein concentration normalized using BCA Protein Assay Kit (Pierce). MDAMB, HCC and HMEC lysates were a generous gift from Clodagh O'Shea, the Salk Institute, La Jolla, CA USA. Protein gel blotting was performed using the indicated primary and secondary antibodies. Blots were analyzed on the La-Gor Odyssey system and processed using Photoshop CS5 extended (Adobe).

Live and confocal imaging. Live-imaging was performed in 8 well μ-slide chambers (iBidi) on either a Yokagawa spinning disk built around a Leica DMIRE2 inverted confocal microscope with a 20x air or 63x 1.4NA oil immersion objective at 37°C maintained by a CO2 enriched air stream incubator (Solent Scientific) and images captured with an EM CCD (Hamamatsu) using SimplePCI software (Compix) or on a Zeiss/Yokagawa spinning disk inverted confocal microscope with a 20x air or 63x 1.4NA oil immersion objective at 37°C maintained by a CO2 enriched air stream incubator (Pecon) and images captured with an EM CCD (Hamamatsu) using AxioVision software (Zeiss).

For fixed imaging, cells were grown on glass coverslips and images were acquired on either a Leica SP2 scanning confocal microscope with a 63x 1.4NA oil immersion objective with LCS software (Leica), or on a Zeiss LSM 710 scanning confocal microscope with a 63x 1.4NA oil immersion objective with Zen software (Zeiss). Fluorochromes and stains used in this study are EGFP, channel3GFP, mCherry, Alexa Fluor 488, Alexa Fluor 568, Alexa Fluor 647, Hoechst 33342 (Molecular Probes), and MitoTracker Red CMXRs (Invitrogen).

Electron microscopy. Cells were grown in 35mm plastic culture dishes were fixed using the protocol of Gilula et al. (1978). The cells were fixed in 2.5% glutaraldehyde in 0.1M Na cacodylate buffer (pH7.3), buffer washed and fixed in 1% osmium tetroxide in 0.1M Na cacodylate buffer. They were subsequently treated with 0.5% tannic acid followed by 1% sodium sulfate in cacodylate buffer and then dehydrated in graded ethanol series. The cells were cleared in HPMAs (2-hydroxypropyl methacrylate: Ladd Research) and embedded in LX112 resin. Following overnight polymerization at 60°C, small pieces of resin were attached to blank blocks using SuperGlue (Scotch). Thin sections (70nm) were cut on a Reichert Ultracut E (Leica) using a diamond knife (Diatome, Electron Microscopy Sciences), mounted on parlodion coated, copper, slot grids and stained in uranyl acetate and lead citrate. Sections were examined at 80KV on a Philips CM100-TEM (FEI) and data documented on an MegaView III CCD camera (Olympus Soft Imaging Solutions).

Image processing and data analysis. Images were analyzed and processed for display using Photoshop CS5 extended (Adobe). Spilling frequency was determined using a MatLab (Mathworks) nucleic counting algorithm to count total number of cells, movies were then analyzed manually, frame by frame, for interphase nuclear ruptures. The number of times each cell ruptured was also tabulated and the average of this number used to determine the frequency of ruptures within individual cells. Statistics were performed using either Prism 5 (GraphPad) or Excel 2011 (Microsoft). Curve fitting of rupture dynamics was done using Prism 5 (GraphPad) as described in the text.

3D reconstruction. Optimized confocal z-series were acquired on a Zeiss LSM 710 scanning confocal microscope and assembled into 3D surfaces for each channel by absolute intensity and with thresholding adjusted so that generated surfaces are matched to fluorescence signal using Imaris (BitPlane).

Disclosure of Potential Conflict of Interest
No potential conflict of interest was disclosed.

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Supplemental Material
Supplemental material may be downloaded here: http://www.landesbioscience.com/journals/nucleus/article/18954/
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