Breaching the nuclear envelope in development and disease

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In eukaryotic cells the nuclear genome is enclosed by the nuclear envelope (NE). In metazoans, the NE breaks down in mitosis and it has been assumed that the physical barrier separating nucleoplasm and cytoplasm remains intact during the rest of the cell cycle and cell differentiation. However, recent studies suggest that nonmitotic NE remodeling plays a critical role in development, virus infection, laminopathies, and cancer. Although the mechanisms underlying these NE restructuring events are currently being defined, one common theme is activation of protein kinase C family members in the interphase nucleus to disrupt the nuclear lamina, demonstrating the importance of the lamina in maintaining nuclear integrity.

Introduction

The nuclear envelope (NE) in animal cells comprises three structures: the nuclear membrane, the nuclear pore complex (NPC), and the lamina. The nuclear membrane is divided into the inner nuclear membrane (INM) and outer nuclear membrane (ONM) based on protein content, but the membranes are contiguous with each other and with the ER. The nuclear membrane covers the chromatin and restricts nuclear–cytoplasmic trafficking to the NPCs. The NPCs extend through both the INM and ONM as well as the lamina (Schermelleh et al., 2008) and regulate the passage of macromolecules with molecular weights exceeding ~40 kD between the nucleus and the cytoplasm (Wente and Rout, 2010). The nuclear lamina is a dense meshwork of lamin filaments attached to the INM. The two major types of lamin proteins are the B-type, lamins B1 and B2, and the A-type, lamins A and C, which are different isoforms of the same gene (Dechat et al., 2010). The lamin proteins interact with transmembrane INM proteins, like LBR and Lap2, and chromatin-binding proteins, like BAF, at the nuclear periphery to form a stable network that supports the membrane and links the INM to the chromatin (Ellenberg et al., 1997; Moir et al., 2000; Wilson and Foisner, 2010). The expression of lamin and lamin-associated proteins varies widely between cell types, likely due to different requirements for nuclear mechanical stiffness and chromatin organization in cells with different functions (Burke and Stewart, 2013).

NE breakdown during mitosis has been the focus of many studies and is a dramatic example of endomembrane reorganization (Güttinger et al., 2009). Unexpectedly, however, it has been shown that the NE can also undergo extensive remodeling in interphase, despite the importance of nuclear compartmentalization for eukaryotic cell biology. At this time, four main types of nonmitotic NE remodeling have been characterized, and will be the focus of this review. First, NE budding has been identified as an export mechanism for large nuclear particles (see Fig. 1). In this process, INM-derived vesicles bud into the perinuclear space and fuse with the ONM to release enclosed nuclear contents into the cytoplasm with no obvious loss of nuclear integrity or cell viability. Lamina disruption is required for budding. Second, transient NE rupturing is characterized by a sudden loss of compartmentalization, causing mislocalization of both nuclear and cytoplasmic components, followed by the restoration of NE integrity without cell death (see Fig. 2, A and B). Third, NE collapse is similar to NE rupturing in that both involve a rapid loss of nuclear integrity associated with lamina gaps and chromatin herniation. However, the membrane does not repair, and instead ER tubules mislocalize to the chromatin (see Fig. 2 C). Fourth, two kinds of NE fusion have been described; (1) the ONM and INM fuse to make a channel through the NE to accommodate NPC insertion, and (2) the ONM and then INM of two separate nuclei fuse to make one contiguous nucleus (see Fig. 3). Thus, accumulating evidence suggests that much remains to be learned about the NE barrier and its remodeling during interphase in normal and diseased cells.

NE budding as an alternate mechanism for nuclear export

Passage through the nuclear pores is the main mechanism of transport between the cytoplasm and the nucleus. Even objects many times larger than the diameter of the NPC, such as certain messenger ribonucleoproteins (mRNPs; Grünwald et al., 2011),
exit the nucleus through the nuclear pores by undergoing a complex unfolding program (Mehlin et al., 1992; Kiseleva et al., 1998; Mor et al., 2010). However, the NPC is not the only avenue for nuclear export. Studies on herpes viruses, neutrophils, and neuromuscular junction formation have shown that large complexes, including mRNPs, can also exit the nucleus by budding through the NE.

The first indication of nuclear membrane–based export came from images of herpes virus capsids within the NE perinuclear space (Stackpole, 1969). Herpes viruses replicate their DNA and package it into capsids in the nucleus. These capsids are larger than the size of the NPC and thus must use an alternative route for nuclear egress, namely NE budding (Fig. 1 A). When the capsids form, they associate with viral protein complexes at the INM that recruit kinases to induce lamina disassembly (Muranyi et al., 2002; Park and Baines, 2006; Marschall et al., 2011) and drive vesiculation of the membrane (Klupp et al., 2007). Once in the perinuclear space, the capsid-containing vesicles then fuse with the ONM to release their cargo into the cytoplasm where the capsids undergo further maturation (Johnson and Baines, 2011). During this process the NE remains intact and cells remain viable.

Recent work from the Budnik group suggests that nuclear export by NE budding might also be important for normal development (Speese et al., 2012). During formation of the neuromuscular junction (NMJ) in Drosophila melanogaster, Wnt signaling mediates maturation of the synapse by promoting mRNP export from muscle cell nuclei (Speese et al., 2012). Unlike other mRNPs, which pass through the NPC, these NMJ-specific mRNPs exit the nucleus through the NE using a similar budding process as herpes viruses (Fig. 1 B; Speese et al., 2012). Because initiation of translation often follows mRNP unfolding at the nuclear pore (Mehlin et al., 1992), it is thought that this export pathway is required to prevent premature translation of mRNAs needed at the NMJ (Strambio-De-Castilla, 2013). Perinuclear vesicles have also been observed during both mouse and rabbit embryogenesis as well as in other developing organs in Drosophila (Gay, 1956; Hadek and Swift, 1962; Hochstrasser and Sedat, 1987; Szöllösi and Szöllösi, 1988), suggesting that NE budding may be routinely used for nuclear export during differentiation. Further research is needed to determine whether mRNP transport by NE budding is limited to early development and whether activation of the NE budding program can occur in all cell types. In addition, it is uncertain whether mRNPs targeted to the NE can also be exported through the NPC and how an export pathway is selected.

An alternative mechanism of NE-based export may be occurring in neutrophils in vivo. Neutrophils are short-lived immune cells that can form neutrophil extracellular traps (NETs), comprised of chromatin and anti-microbial proteins, to trap and kill bacteria (Brinkmann and Zychlinsky, 2012). In most cases NETs form after chromatin decondensation, NE breakdown, and cell lysis (Brinkmann et al., 2004; Fuchs et al., 2007). However a second type of NETosis has been identified where vesicles containing decondensed chromatin appear to bud off from the NE and fuse with the plasma membrane in the absence of NE breakdown or cell death (Pilsczek et al., 2010). This type of NETosis by NE remodeling appears to be important for pathogen containment in vivo (Yipp et al., 2012). In contrast to NE budding, images of this process suggest that the chromatin in the expanded perinuclear space is not in vesicles (Pilsczek et al., 2010). Thus, the chromatin may enter the perinuclear space by a different mechanism than INM budding.

Lamina disassembly during NE budding occurs by a mechanism similar to those that break down the lamina in mitosis and apoptosis. During both mitosis and apoptosis, kinases are activated that phosphorylate the lamins, causing the protein network to fall apart, or targeting the lamins for degradation by caspases (Cross et al., 2000). Members of the PKC kinase family have been shown to be important lamin kinases for mitosis and apoptosis (Hocevar et al., 1993; Goss et al., 1994; Thompson and Fields, 1996; Collas, 1999; Cross et al., 2000), as well as for INM vesicle formation around both mRNPs and herpes virus capsids (Muranyi et al., 2002; Park and Baines, 2006; Leach and Roller, 2010; Speese et al., 2012). Lamin protein phosphorylation by these kinases results in localized lamina disassembly at the site of NE budding (Muranyi et al., 2002; Park and Baines, 2006; Leach and Roller, 2010; Speese et al., 2012). Interestingly, different PKC family members are involved in herpes virus budding than in mRNP budding in Drosophila (Muranyi et al., 2002; Park and Baines, 2006; Milbradt et al., 2010; Speese et al., 2012), suggesting that different signaling pathways regulate lamina gap formation in different contexts. Mitotic activation of PKC results in global disassembly of lamina, thus an important question for interphase PKC activation is how lamina disassembly is restricted to sites of NE budding.

Another interesting question is how vesicle fusion events in the NE are mediated. Analysis of viral protein mutants suggests that vesicle fusion to the ONM is independent of INM envelopment and may be regulated by phosphorylation events.
NE budding as a transport mechanism appears to be limited to nuclear export; examples of cargo import by NE budding have not been identified. This could be due to biochemical asymmetries between INM and ONM that prevent vesicle budding from the ONM or block vesicle fusion to the INM. Consistent with this idea, parvoviruses, the only virus family known to bypass the NPCs for nuclear import, require transient disruption of the NE to access the nucleus. Parvoviruses are a family of small DNA viruses that includes adeno-associated virus. When parvovirus capsids reach the NE they generate large holes in the ONM, then in the INM, and finally in the lamina to allow capsid entry to the nucleus (Fig. 2 A; Cohen and Panté, 2005; Cohen et al., 2006; Porwal et al., 2013). In the absence of either membrane disruption or lamina disassembly, the virus fails to enter the nucleus (Porwal et al., 2013), indicating that breaking down both barriers is required for import. Nuclear integrity is lost a few minutes after infection, but in later stages the NE appears contiguous and compartmentalization is fully restored (Cohen et al., 2011). This indicates that even though parvoviruses generate large breaks in the membrane, the NE is still able to undergo repair.

During NE rupturing by parvovirus infection, as in NE budding of herpes viruses, PKCα activity is required to initiate lamina disruption (Porwal et al., 2013). However, NE rupturing in this system requires activation of an additional kinase, Cdk2, by PKCα and caspase-3 (Cohen et al., 2011; Porwal et al., 2013). Parvovirus infection induces much larger ruptures in the lamina than are observed during NE budding. Thus, additional kinase activation may be required to sustain more extensive lamin phosphorylation.

Infection with HIV can also cause dramatic NE instability. The HIV protein VPR is thought to modulate the cell environment to make it more favorable for viral replication (Andersen et al., 2008). However, VPR expression can induce repeated transient NE rupturing and loss of compartmentalization, causing mislocalization of nuclear and cytosolic cell cycle regulators, including wee-1 and cdc25 (de Noronha et al., 2001). NE rupturing is accompanied by prominent lamina gaps as well as large blebs of herniating chromatin, which are also present in cells infected with HIV (Fig. 2 B; de Noronha et al., 2001). VPR is dispensable for HIV infection (Zufferey et al., 1997), however, indicating that NE rupturing is not required for nuclear import of viral DNA. VPR expression can arrest the cell in G2 by induction of DNA damage (Roshal et al., 2003), and one model suggests that VPR-induced repetitive NE rupturing is the cause of this damage (Planelles and Benichou, 2009). However, it is also possible that by arresting cells in G2, VPR causes premature attachment of microtubules to the NE, resulting in membrane disruption. This would be consistent with the observation that microtubule interactions with the NE, which increase during G2, induce tears in the membrane (Beaudouin et al., 2002; Salina et al., 2002). Further work is needed to address how VPR expression causes lamina disruption, the frequency of NE rupturing in infections in vivo, and what the consequences of nuclear integrity loss are for viral infection.

**Figure 2. **Nuclear envelope rupturing and collapse. [A] Association of parvovirus capsids with the ONM causes breakdown of first the outer and then the inner nuclear membranes. Activation of PKC and Cdk kinases in the nucleus during this time forms large gaps in the lamina, allowing the capsids to enter the nucleoplasm and causing a loss of nuclear integrity. [B] When lamina organization is disrupted by changes in lamina proteins, patches of weak membrane form and chromatin can herniate. This membrane can undergo multiple rounds of NE rupturing and repair, causing mislocalization and entrapment of cytosolic and nuclear components. [C] Micronuclei have a high probability of NE rupturing but fail to undergo NE repair, causing a persistent loss of nuclear integrity. After rupturing, the chromatin forms aberrant associations with ER tubules in a process called NE collapse.
Nuclear envelope rupturing in laminopathies

Because lamina disruption is a general feature of NE remodeling, a significant rise in interphase NE dynamics is likely to be a feature of lamin-associated human diseases, known as laminopathies. Laminopathies are genetic diseases caused by mutations in lamin A/C or lamin-associated proteins that alter lamina organization and result in tissue-specific cell loss (Worman et al., 2010).

A common observation in cells expressing laminopathy mutant proteins is large gaps in the lamina where B-type lamins, NPCs, and other structural INM proteins are absent and large chromatin herniations appear to push out the weakened membrane (de Noronha et al., 2001; de Noronha et al., 1997; Vigouroux et al., 2001). These lamina gaps also occur when lamin B1 is misregulated (Vergnes et al., 2004; Vargas et al., 2012). Regardless of their origin, one consequence of lamina discontinuities is repeated nonlethal transient NE rupturing at these sites (De Vos et al., 2011; Vargas et al., 2012).

Transient NE rupturing causes both soluble proteins and cytoplasmic and nuclear organelles, like vesicles, mitochondria, and PML bodies, to become mislocalized. Although soluble proteins can be resorted to the correct compartment, mislocalized organelles become trapped when the NE repairs (De Vos et al., 2001; De Vos et al., 2011; Vargas et al., 2012). Thus, NE rupturing could be an important contributor to laminopathy pathology.

Direct evidence of NE rupturing in laminopathy cells has been limited to cultured cells, but indirect evidence of NE rupturing is present in fixed tissues. Gaps in the lamina have been observed in liver nuclei of mice lacking lamin A/C (Sullivan et al., 1999), and disruption of the nuclear membrane is apparent in muscle tissue in flies lacking B-type lamins (Lenz-Böhme et al., 1997). Biopsies from laminopathy patients show even clearer signs of NE rupturing. In post-mitotic cardiomyocytes, both disruption of the nuclear membrane and mislocalization of mitochondria to the nucleus have been observed by electron microscopy (Fidziańska et al., 2008; Gupta et al., 2010). Together, these data demonstrate that NE rupturing can occur in a variety of contexts in vivo.

Although mutation or loss of a lamin protein can induce NE rupturing, lamin depletion by itself is not sufficient to destabilize the membrane. Knockout of either both B-type lamin genes or all three lamin genes in embryonic stem cells does not cause an increase in lamina gaps or chromatin herniation (Kim et al., 2011, 2013). Recent modeling of chromatin herniation suggests that weak points in the membrane form when mismatched protein networks, like lamin B versus lamin A, generate tension on the NE (Funkhouser et al., 2013). Thus, loss of multiple lamins may be less problematic than misregulation of a single lamin because the NE is under less tension. Consistent with this hypothesis, the frequency of membrane rupturing in laminopathy cells decreases when the tension on the NE is reduced by growth on soft substrates (Coffinier et al., 2010; Tamiello et al., 2013).

This correlation is also observed in vivo; the most dramatic defects in NE integrity are found in muscle and heart tissue where the nuclei are under increased tension (Lenz-Böhme et al., 1997; Fidziańska et al., 2008; Gupta et al., 2010). Thus, additional factors, including the organization of the cytoplasmic cytoskeleton, likely determine the frequency of nuclear integrity loss in cells with lamina defects.

Tension in the cytoskeleton is transmitted to the nucleus via LINC complexes, which traverse the NE and connect the cytosolic cytoskeleton to the lamina, and defects in LINC complex members have been shown to affect nuclear structure (Tapley and Starr, 2013). Consistent with the hypothesis that increased tension increases NE defects in cells with an altered lamina, depleting the LINC complex member SUN-1 reduced chromatin herniation frequency in laminopathy cells (Chen et al., 2012). However, this was not true for all LINC complex members; interfering with nesprins increased nuclear defects and the severity of laminopathy symptoms (Kandert et al., 2007; Zhang et al., 2007; Puckelwartz et al., 2009, 2010). LINC complex proteins have additional roles in cellular organization and nuclear functions (Rothballer and Kutz, 2013b); thus, more research is required to determine the importance of LINC complex proteins in regulating NE integrity in laminopathy cells.

Recent work suggests that studying senescent cells may also provide valuable information about how lamin misregulation affects NE integrity. One hallmark of senescent cells is a significant decrease in lamin B1 levels (Shimi et al., 2011; Freund et al., 2012), accompanied by small chromatin herniations (Ivanov et al., 2013). Nuclei from senescent cells are more permeable after isolation (Ivanov et al., 2013), but the extent to which NE integrity is altered in these cells is unclear. Determining whether these nuclei are rupturing and, if not, what additional changes prevent this, could provide insight into the mechanism of NE rupturing.

Nuclear rupturing and collapse in cancer cells

Cancer cells often exhibit changes in nuclear morphology and lamin A/C expression similar to those that result in transient NE rupturing in laminopathies (Zink et al., 2004; Prokocimer et al., 2006). Consistent with this, examination of the NE in cultured cancer cell lines demonstrated that they have a significantly higher frequency of chromatin herniation and, unlike nontransformed cells, undergo NE rupturing. Similar to laminopathy cells, the frequency of NE rupturing can be increased by altering the structure of the lamina through decreasing lamin levels (Vargas et al., 2012). At this time, no marker has been developed to positively identify NE rupturing in fixed cancer tissues, but intranuclear mitochondria have been observed in leukemias and lymphomas (Brandes et al., 1965; Oliva et al., 1973), suggesting that NE rupturing and repair can occur in cancer cells in vivo.

Although changes in nuclear morphology are a gold standard for cancer diagnosis, very little is known about why disruption of the NE structure would benefit a cancer cell. It is known that altering the lamina can cause changes in heterochromatin formation and gene expression (Stewart et al., 2007), which could facilitate carcinogenesis (Prokocimer et al., 2006). In this model, NE rupturing would be a passive side effect of lamina disruption. However, NE rupturing could also promote cancer development. Distended chromatin at the site of NE rupturing can undergo changes in both chromatin structure and nuclear functioning (Shimi et al., 2008). In addition, regulation of gene expression might be compromised by mislocalization of transcription factors due to a repeated loss of compartmentalization (De Vos et al., 2011).
When NE rupturing is induced in primary nuclei, either through viral infection or disruption of the lamina, the NE is almost always repaired and compartmentalization restored (de Noronha et al., 2001; Cohen et al., 2011; De Vos et al., 2011; Vargas et al., 2012). However, this repair fails when NE rupturing occurs on micronuclei (MN). Recent work from our laboratory has shown that a large proportion of MN that form from chromosome missegregation rupture during interphase and fail to regain compartmentalization before mitosis. Similar to primary nucleus NE rupturing, MN rupturing also stems from lamin disorganization leading to lamina gaps (Hatch et al., 2013). But instead of resealing over the chromosome, the NE is replaced by ER tubules that invade the chromatin in a process we have termed “NE collapse” (Fig. 2 C).

Unlike NE rupturing in primary nuclei, membrane disruption in micronuclei has clear consequences for genomic instability. First, several nuclear functions important for maintaining chromosome integrity, including DNA damage repair and DNA replication, are impaired in intact micronuclei and abrogated by NE rupturing (Crasta et al., 2012; Hatch et al., 2013). Second, NE rupturing in MN can also trigger massive DNA damage (Hatch et al., 2013). The clustered DNA damage on chromosomes in collapsed MN, which likely arises from the sudden compaction of replicating DNA (Zhang et al., 2013), makes them an ideal substrate for chromothripsis (Crasta et al., 2012). In chromothripsis a single chromosome, or chromosome fragment, is shattered and then stitched together to form a highly rearranged chromosome (Stephens et al., 2011). Since its identification, evidence of chromothripsis has been found in a wide variety of cancers and is generally associated with poor outcomes (Kloostrerman et al., 2014). Although a causal link between MN rupturing and chromothripsis remains to be shown, loss of nuclear activity and accumulation of DNA damage is likely to cause significant changes in chromosome function, increasing the likelihood of aneuploidy.

Although NE rupturing in primary nuclei and MN are both rooted in lamina disorganization, it is unclear why MN develop lamina defects more often than primary nuclei. One possibility is that the higher curvature of the NE in MN can induce alterations in the lamina. However, MN with significantly different sizes have the same probability of NE rupturing (Hatch et al., 2013), suggesting that membrane curvature might not play a critical role. In addition, lamin B1 discontinuities in MN can already be observed in early G1 (Hatch et al., 2013), suggesting that problems with NE assembly could contribute to aberrant lamina formation. Lamina construction generally occurs late in NE formation after the membranes have sealed (Newport et al., 1990; Daigle et al., 2001; Haraguchi et al., 2008), although there are examples of lamins binding directly to the chromatin early in anaphase (Moir et al., 2000). Thus, misregulation of early steps in NE assembly could have downstream effects on lamina organization. One hypothesis is that because chromosomes have different overall chromatin properties, not all chromosomes are able to appropriately interact with NE proteins when separated from the main chromatin mass. Consistent with this, specific chromatin sequences or histone modifications are thought to facilitate protein recruitment during NE assembly (Güttlinger et al., 2009). Thus, further research on the interaction of NE assembly proteins with chromatin during mitotic exit could elucidate why MN have a high probability of NE rupturing.

**NE repair**

Although countless nuclear injection experiments in a variety of systems depended on the ability of the NE to repair after puncture, the mechanism by which the NE reseals is not understood. The transience of NE rupturing in the primary nucleus, even when membrane disruption is extensive (Cohen et al., 2006), demonstrates that the NE has a much larger capacity for repair than anticipated, but there is little information about potential mechanisms. It is likely that NE repair requires connectivity to the ER, as this connection is required for interphase NE expansion (D’Angelo et al., 2006; Anderson and Hetzer, 2007; Lu et al., 2011). Alternatively, NE repair could be initiated by interactions of INM proteins in the ER with the exposed chromatin, leading to membrane spreading, as occurs during post-mitotic NE closure (Anderson et al., 2009). Transmembrane proteins that localize to the chromatin early in NE assembly do aggregate on MN chromatin after NE collapse, as do ER tubules (Hatch et al., 2013). However, MN do not undergo NE repair, suggesting that this recruitment is not sufficient to reform the nuclear membrane. In contrast, INM proteins are largely depleted from the sites where NE rupturing and repair occurs in the primary nucleus (Sullivan et al., 1999; Vigouroux et al., 2001), suggesting that recruitment of NE assembly proteins to interphase chromatin may inhibit repair. Consistent with this idea, MN efficiently undergo post-mitotic NE assembly (Hatch et al., 2013), suggesting that the MN chromatin is competent for NE formation, but that this process is inhibited during interphase. Alternatively, changes in chromatin state after NE rupturing in MN, including compaction and loss of acetylation (Hatch et al., 2013), could alter the ability of INM proteins and ER membranes to productively interact with the MN chromatin. It will be important to determine whether similar chromatin changes are occurring during primary nucleus rupturing, and whether NE assembly proteins are important for primary nucleus repair to begin to understand this process.

**Membrane fusion events involved in NE reorganization**

Fusion of the INM and ONM occurs frequently in growing cells when new nuclear pores are assembled into the expanding nuclear membrane (D’Angelo et al., 2006). An early step in interphase NPC assembly is fusion of the ONM and INM to generate a channel where the NPC can go (Goldberg et al., 1997; Doucet et al., 2010). Nucleoporins then associate with the curved membrane channel to stabilize it and initiate pore assembly (Fig. 3 A; Rothballer and Kutay, 2013a; Smoyer and Jaspersen, 2014). Although the mechanism of nuclear membrane fusion is unclear, several proteins are known to function in the remodeling process. First, ER-shaping proteins, such as reticulons, have been shown to be required for membrane fusion before NPC and spindle pole body insertion in yeast (Dawson et al., 2009; Casey et al., 2012). In addition, several NE transmembrane proteins, including Sun-1 and members of the NPC, are required to shape the membrane channel, although it is unclear...
if these proteins initiate or stabilize membrane fusion (Rothballer and Kutay, 2013a; Smoyer and Jaspersen, 2014). Both NPC subcomplexes and transmembrane proteins have structural similarities to proteins involved in vesicle fusion (Devos et al., 2004, 2006; Brohawn et al., 2009; Rothballer and Kutay, 2013a), but it is unclear how related the two processes are. During NE budding, fusion must also occur between a vesicle derived from the INM and the ONM. Determining what INM proteins are retained in the vesicle and required for fusion may significantly enhance our understanding of ONM and INM fusion in general.

How two nuclei fuse was first addressed using yeast karyogamy as a model (Melloy et al., 2007, 2009; Ydenberg and Rose, 2008), and recent work has begun to elucidate nuclear fusion events in metazoans. In several species, including sea urchins, frogs, zebrafish, and rabbits, NE fusion is an important mechanism to maintain euploidy during embryogenesis. NE fusion occurs at two stages in these animals during early development: pronuclei fuse before the first mitotic division, and multiple nuclei are fused into a mononucleus during early cleavage divisions (Abrams et al., 2012). When the cytoplasm-to-nucleus ratio is very high, as in the oocyte, chromosomes can become separated during mitosis resulting in multinucleation, also called karyomere formation. Starting in telophase, these karyomeres fuse to form a mononucleus. Images of pronuclear fusion in sea urchin describe a model where proximity of the pronuclei causes mixing of the ONM followed by fusion of the INMs and mixing of the nuclear contents (Longo and Anderson, 1968). Although lamina disruption has not been observed, it is likely that some disassembly is required to permit expansion of the fusion pore (Lénárt and Ellenberg, 2003). Recent work in zebrafish identified brambleberry, an NE transmembrane protein, as an important protein for initiating NE fusion in both pronuclei and karyomeres (Fig. 3 B; Abrams et al., 2012). Depletion of brambleberry demonstrated that, although karyomeres can efficiently perform nuclear functions (Lemaitre et al., 1998), NE fusion is required for development, as embryos depleted of brambleberry arrested early in embryogenesis (Abrams et al., 2012). Further analysis of brambleberry interactors and functions will likely provide important insights into the mechanism of nuclear membrane fusion.

**Conclusion**

A brief survey of interphase NE remodeling and disruption events demonstrates that interphase NE dynamics are important in an increasing number of developmental and disease contexts. As the consequences of these events become clearer it will not only clarify the pathology of viral infections, laminopathies, and cancer, but could also impact new technology development. Currently, the parovirus AAV (adeno-associated virus) is being used as a delivery mechanism for gene therapy. Thus, understanding the consequences of NE rupturing from virus infection could be important to mitigate side effects of this treatment. In addition, one of the main problems in in vitro fertilization is the high frequency of aneuploidy in early divisions due to multinucleation (Chavez et al., 2012). An ability to initiate NE fusion in these cases may be able to increase the frequency of successful fertilizations. At this time, a number of questions remain about how remodeling in NE budding and fusion occurs and what the consequences of transient NE rupturing are for chromatin structure, gene expression, and other nuclear functions. The world of interphase NE dynamics is just beginning to be explored, but its importance in cell biology is already clear.

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