Spatial control of phospholipid flux restricts endoplasmic reticulum sheet formation to allow nuclear envelope breakdown

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The nuclear envelope is a subdomain of the endoplasmic reticulum (ER). Here we characterize CNEP-1 (CTD [C-terminal domain] nuclear envelope phosphatase-1), a nuclear envelope-enriched activator of the ER-associated phosphatidic acid phosphatase lipin that promotes synthesis of major membrane phospholipids over phosphatidylinositol (PI). CNEP-1 inhibition led to ectopic ER sheets in the vicinity of the nucleus that encased the nuclear envelope and interfered with nuclear envelope breakdown (NEBD) during cell division. Reducing PI synthesis suppressed these phenotypes, indicating that CNEP-1 spatially regulates phospholipid flux, biasing it away from PI production in the vicinity of the nuclear envelope to prevent excess ER sheet formation and NEBD defects.

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The endoplasmic reticulum (ER) is composed of sheets and tubules bounded by a membrane bilayer that faces the cytoplasm on one side and an internal lumen on the other. The ER is partitioned into subdomains specialized for different functions. The nuclear envelope subdomain is a spherical sheet perforated by nuclear pores that encases the chromatin. The outer membrane and lumen of the nuclear envelope are directly contiguous with other ER structural elements, whereas passage of proteins to the inner nuclear membrane (INM) is gated by the nuclear pores (Hetzer 2010; English and Voeltz 2013). The chromatin-facing INM is associated with the nuclear lamina, a dense filament meshwork that provides structural support (Hetzer 2010; Gerace and Huber 2012). The ER also serves as a platform for de novo phospholipid synthesis (Fagone and Jackowski 2009; Lagace and Ridgway 2013). A central player is the conserved phosphatase lipin, which converts phosphatidic acid (PA) to diacylglycerol [DAG] [Carman and Han 2006; Siniossoglou 2013]. In metazoans, lipin is at a branching point in the phospholipid synthesis pathway; the major building blocks of membrane bilayers [phosphatidylcholine [PC] and phosphatidylethanolamine [PE]) are synthesized from the lipid product DAG, whereas phosphatidylinositol (PI) is synthesized from the lipin substrate PA [Carman and Han 2006; Lagace and Ridgway 2013; Siniossoglou 2013]. PI is converted to phosphoinositides [PIPs] when transported to other organelles that have specific PIP kinases and phosphatases [Balla 2013]. Within the ER, PC and PE make up >70% of total phospholipid, whereas PI makes up <10% [van Meer et al. 2008]. In metazoans, decreased lipin activity is predicted to shift phospholipid flux away from production of DAG and the major membrane phospholipids PC and PE toward the production of PI. This is in contrast to budding yeast, where a pathway that is not present in metazoans [the CDP–DAG pathway] (Supplemental Fig. S1A) converts PA to major membrane phospholipids [Carman and Han 2011; Siniossoglou 2013]. Thus, lipin inhibition in budding yeast leads to an increase in all phospholipids and an abnormal expansion of the nuclear envelope that does not occur in metazoans [Han et al. 2006; Siniossoglou 2013].

Understanding lipin in metazoans is motivated by the fact that it is mutated in a number of human metabolic diseases and in a mouse model for lipodystrophy [Reue 2009]. Lipin is recruited to the ER through a peripheral membrane-binding domain; however, soluble lipin also exists that transcriptionally coactivates genes involved in lipid metabolism [Siniossoglou 2013]. Lipin activity and localization are regulated via multisite phosphorylation by Cdk1 and mTORC1 [Carman and Han 2011; Siniossoglou 2013], which negatively regulate lipin activity [Eaton et al. 2013]. The conserved C-terminal domain (CTD) phosphatase CTDNEP1 [formerly Dullard] is an integral membrane protein enriched on the nuclear envelope that is predicted to activate lipin by dephosphorylating it [Siniossoglou et al. 1998; Kim et al. 2007; Han et al. 2012; Eaton et al. 2013].

Lipin inhibition in HeLa cells and Caenorhabditis elegans embryos slows nuclear envelope disassembly during mitosis [Golden et al. 2009; Gorjánész and Mattaj 2009; Mall et al. 2012]. This has been proposed to result from a reduction in DAG levels, which in turn could decrease the activity of DAG-dependent protein kinase Cs that phosphorylate lamins to promote their disassembly.
Here, we investigate the role of lipin in nuclear envelope disassembly in the C. elegans embryo by characterizing the effects of inhibiting its nuclear envelope-localized regulator, CTDNEP1 (CNEP-1 [CTD nuclear envelope phosphatase-1] in C. elegans). CNEP-1 inhibition led to ectopic ER sheets that encased the nuclear envelope in an extra ER layer and interfered with nuclear envelope disassembly. CNEP-1 inhibition also led to elevated PI levels, and reducing PI synthesis suppressed ectopic ER sheet formation and nuclear envelope disassembly defects in the cnep-1 mutant. We conclude that CNEP-1 spatially controls lipin-dependent phospholipid flux to limit PI levels and restrict ER sheet formation in the vicinity of the nuclear envelope.

Results and Discussion

Inhibition of the C. elegans homolog of the lipin activator CTDNEP1, called CNEP-1, slows nuclear envelope disassembly during the first division of the C. elegans embryo (Han et al. 2012), a phenotype also observed following partial inhibition of lipin [LPIN-1 in C. elegans] (Golden et al. 2009; Gorjánész and Mattaj 2009; Han et al. 2012). To investigate how CNEP-1 promotes nuclear envelope disassembly, we generated transgenes expressing GFP fusions with lipin and wild-type or phosphatase-defective [PD] CNEP-1 [CNEP-1 WT or CNEP-1 PD] (Fig. 1B; Supplemental Fig. S1B). In contrast to GFP::LPIN-1, which localized throughout the ER (Fig. 1B; Gorjánész and Mattaj 2009), CNEP-1::GFP was specifically enriched on the nuclear envelope [Fig. 1B]. Consistent with its central role in phospholipid synthesis [Fig. 1A], injection of dsRNA targeting lipin into adult worms led to the arrested cessation of embryo production [Fig. 1B]. In contrast, deletion of the cnep-1 gene [cnep-1 Δ] [Supplemental Fig. S1C] led to only a modest reduction in brood size [Fig. 1B]. The localization of CNEP-1 to the nuclear envelope, combined with the reduced severity of the cnep-1 phenotype, suggested that CNEP-1 activates a pool of lipin at the nuclear envelope to serve a specific function.

To characterize the effects of the cnep-1 deletion on nuclear envelope dynamics, we filmed embryos expressing a GFP fusion with the INM protein LEM-2 [Galy et al. 2003]. After fertilization, the maternal and paternal pronuclei form, increase in size, and migrate toward each other, meeting in the middle as the embryo enters mitosis [Fig. 1C]. The nuclear envelopes become permeable, and microtubules penetrate to interact with the chromosomes. The nuclear envelopes continue to disassemble as the embryo progresses into anaphase, and the scission of maternal and paternal nuclear envelopes allows chromosome mixing approximately coincident with anaphase onset [Fig. 1C,E]. When the nuclear envelopes reform in telophase, a single nucleus forms in each daughter cell [Fig. 1C,D]. Perturbations that slow nuclear envelope disassembly result in a failure of scission [Audhya et al. 2007]. In this case, two nuclei form in each daughter cell—one containing the maternal chromosomes, and one containing the paternal chromosomes [“twinned” nuclei] [Fig. 1C,D]. Delayed scission can also result in a less severe phenotype characterized by misshapen, “oblong” nuclei [Fig. 1D].

In control embryos, 100% of nuclei at the two-cell stage exhibit a normal spherical morphology. In contrast, 76% of two-cell stage nuclei in cnep-1Δ embryos and 100% following partial LPIN-1 depletion were twinned or oblong, indicative of delayed nuclear envelope scission.

The two-cell stage nuclear morphology defect in cnep-1Δ embryos was rescued by the wild-type, but not the PD, cnep-1 transgene [Fig. 1D]; both transgenic proteins were expressed at similar levels [Supplemental Fig. S1B] and localized to the nuclear envelope [Fig. 1B; Supplemental Fig. S1D]. Thus, the phosphatase activity of CNEP-1 is essential for its role in nuclear envelope disassembly. Consistent with the idea that CNEP-1 functions by dephosphorylating LPIN-1, levels of slower-migrating phosphorylated LPIN-1 species were increased in cnep-1Δ worms and restored to control levels by the wild-type, but

Figure 1. The lipin activator CNEP-1 concentrates on the nuclear envelope and is required for timely nuclear envelope disassembly during mitosis. (A) Schematic showing lipin at the branching point between the synthesis of major membrane phospholipids and PI. (B) Fluorescence confocal images of embryos expressing the ER lumen marker mCherry::SP12 along with GFP::LPIN-1 [top] [n > 10 embryos] or mCherry::SP12 and CNEP-1::GFP in a background deleted for the endogenous cnep-1 gene [bottom] [n > 10 embryos]. Graphs plot normalized brood size. Error bars are SEM. (N) Number of worms; (n) total number of embryos. (C) Schematics highlight scission of the maternal and paternal pronuclear envelopes [green] during chromosome [red] segregation in the first division of control C. elegans embryos and failure of scission in embryos exhibiting a nuclear envelope breakdown [NEBD] defect. (D, left) Inverted contrast fluorescence confocal images of two-cell stage embryos expressing GFP::LEM-2 illustrate nuclear morphology phenotypes. [Right] Graph plotting frequency of abnormal nuclear phenotypes for the indicated conditions. (E) Fluorescence confocal images of control and cnep-1Δ embryos expressing GFP::LEM-2 and mCherry::histine. Times are in seconds relative to anaphase onset. Yellow arrowheads mark the site of nuclear envelope scission in wild type. (F) Immunoblots of lysates from control and cnep-1Δ worms [top] or cnep-1Δ worms harboring the wild-type [WT] or PD transgenes [bottom] probed for LPIN-1. Bars, 10 μm.
not the PD, *cnep-1* transgene [Fig. 1F]. In contrast to the effect of the *cnep-1* deletion on nuclear envelope scission, no defects were observed in nuclear envelope expansion following fertilization or in nuclear envelope permeabilization during mitotic entry [Supplemental Fig. S1E,F].

To understand how the defect in nuclear envelope disassembly arises in *cnep-1* mutants, we used transmission electron microscopy (TEM) after high-pressure freezing and freeze substitution to examine the nuclear envelope at different stages of mitosis. In control embryos, prophase nuclei (judged by chromosome condensation state) were encased in a single double-membrane sheet [Fig. 2A]. This was also the case for prometaphase–metaphase nuclei, although short isolated regions were observed where two stacked lumens could be detected along the nuclear envelope periphery [Fig. 2B, dark-red arrow]. In *cnep-1∆* embryos, short regions with two stacked lumens were observed in prophase nuclei [Fig. 2A], and prometaphase–metaphase nuclei appeared encased in a second ER layer, two parallel lumens could be traced along the majority of the nuclear periphery [Fig. 2B; Supplemental Fig. S2]. This result suggests that the failure of nuclear envelope scission in *cnep-1∆* embryos is due to wrapping of the nuclear envelope in an extra ER layer, which places eight membrane bilayers and four lumens between the maternal and paternal genomes (Fig. 2B, bottom inset).

Wrapping of the nuclear envelope in an extra ER layer in the absence of CNEP-1 is likely due to reduced activation of lipin in the vicinity of the nuclear envelope. A reduction in lipin activity could exert such an effect by either reducing levels of its product, DAG, which in turn may reduce levels of the major membrane phospholipids PE and PC, or increasing levels of its substrate, phosphatidic acid (PA), which in turn may elevate production of PI [Fig. 3A]. To distinguish between these possibilities, we directly measured phospholipid content by metabolically labeling control and *cnep-1∆* worms and analyzing isolating lipids by thin-layer chromatography. This analysis revealed that PI and PA levels were increased in *cnep-1∆* worms, with PI levels being elevated ~2.5-fold compared with controls, in contrast, PE and PC levels were not significantly altered [Fig. 3B].

To assess the relationship of the alteration in lipid levels to the *cnep-1∆* phenotype, we analyzed the consequences of depleting other enzymes in the phospholipid biosynthetic pathway [Fig. 3A] on nuclear envelope disassembly. Partial depletion of the germline choline/ethanolamine transferase CEPT-2 (choline/ethanolamine phosphotransferase-2), which acts downstream from lipin to convert DAG into PE/PC, phenocopied *cnep-1* inhibition [Fig. 3C]. Like inhibition of CNEP-1 and lipin, CEPT-2 inhibition is expected to bias biosynthetic flux toward PI. To test the prediction that elevated PI levels are the origin of the nuclear disassembly defect, we individually depleted the two enzymes required to convert PA to PI: CDGS-1 (CDP-DAG synthase-1), which converts PA to CDP-DAG, and PSY-1 (PI synthase-1), which converts CDP-DAG to PI. Neither depletion affected two-cell stage nuclear morphology [Fig. 3C]. However, depletion of either PSY-1 or CDGS-1 significantly rescued the nuclear morphology defects resulting from inhibition of *cnep-1* or partial inhibition of *lipin-1* [Fig. 3C; Supplemental Movie S1]. In contrast, neither depletion rescued the nuclear envelope disassembly defects observed following depletion of the transmembrane nucleoporin NPP-12 [the *C. elegans* gp210 homolog] [Fig. 3C; Audhya et al. 2007; Galy et al. 2008], indicating that the suppression is specific for the lipin pathway. Biochemical analysis revealed that *cdgs-1* inhibition reduces the elevation of PI, but not of PA, in *cnep-1∆* worms [Fig. 3D], consistent with the fact that CDGS-1 is between PA and PI in the lipid biosynthetic pathway [Fig. 3A]. The suppression of the *cnep-1* and *lipin-1* phenotypes by inhibition of the enzymes that convert PA to PI, together with the direct biochemical analysis of lipid content, strongly suggest that the defect in nuclear envelope disassembly following CNEP-1 inhibition is caused by an increase in PI levels.

These findings argue that restricting PI synthesis in the vicinity of the nuclear envelope is important to prevent defects in nuclear envelope disassembly. The nuclear envelope disassembly defect resulting from *cnep-1* and *lipin-1* inhibition also appears to be distinct from that resulting from depletion of NPP-12, a transmembrane nucleoporin that controls lamin disassembly and INM protein dispersal [Audhya et al. 2007; Galy et al. 2008]. Consistent with the idea that inhibition of CNEP-1 and NPP-12 disrupts nuclear envelope disassembly via distinct means, CNEP-1 inhibition slowed lamin disassembly and LEM-2 dispersal but to a reduced extent compared

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**Figure 2.** Mitotic nuclei in *cnep-1∆* mutants are wrapped in an additional ER layer. (A,B) Transmission electron micrographs of the nuclear regions of prophase [top] and prometaphase/metaphase [bottom] high-pressure-frozen embryos (eight to 16 cells) at the indicated mitotic stages (*n* = 15 mitotic cells for control, *n* = 11 mitotic cells for *cnep-1∆*). Magnified images of boxed regions are paired with micrographs pseudocolored to highlight the nuclear envelope/ER lumens [red] that separate the cytoplasm [yellow] and nucleus [blue]. Cineched-in regions along the nuclear envelope in A are where nuclear pore complexes are inserted. Arrows [dark red] point to isolated regions where two adjacent lumens are detected. One-hundred percent of *cnep-1∆* nuclear envelopes during late stages of NEBD (*n* = 7), as compared with none in control (*n* = 6), displayed an additional double-membrane sheet encircling the nuclear envelope. Bars, 1 μm.
with NPP-12 depletion (Supplemental Fig. S3). In addition, simultaneous inhibition of CNEP-1 along with NPP-12 led to a greatly enhanced defect in nuclear envelope disassembly. In embryos in which CNEP-1 or NPP-12 alone was depleted by RNAi or in embryos homozygous for deletions of the *cnep-1* or *npp-12* genes, twinned nuclei were often observed at the two-cell stage, but cytokinesis was always successful. In contrast, simultaneous inhibition of *cnep-1* and *npp-12* led to a severe defect in which cytokinesis failed in 50% of cases due to the inability of the cytokinetic furrow to cut through the remnants of the nuclear envelope (Fig. 3E; Supplemental Movie S2). Like the defects in nuclear envelope disassembly in the *cnep-1*Δ mutant, the cytokinesis defect in the double-deletion mutant was rescued by CDGS-1 depletion (Fig. 3E). We conclude that increased PI synthesis caused by CNEP-1 inhibition leads to wrapping of nuclei in an extra ER layer that slows nuclear envelope disassembly.

To identify the source of the extra ER layer that encases the nuclear envelope in *cnep-1*Δ embryos, we analyzed ER morphology in embryos expressing a GFP-tagged ER lumen marker (GFP::SP12). The ER is normally relatively evenly distributed in interphase embryos and then condenses to form patches as embryos enter mitosis [Poteryaev et al. 2005; Audhya et al. 2007]. ER morphology was noticeably altered in *cnep-1*Δ embryos with prominent ER patches apparent prior to the onset of mitosis (Fig. 4A);
similar defects were seen following partial inhibition of lipin-1 or cept-2 [Fig. 4A; Supplemental Fig. S4A]. Similar to our analysis of nuclear morphology, depletion of either of the enzymes required to convert PA to PI (PI3Y-1 or CDCS-1) did not lead to any detectable alteration in ER morphology on its own [Fig. 4A; Supplemental Fig. S4A]; however, both inhibitions rescued the ER morphology defect resulting from deletion of cnep-1 or partial inhibition of lipin-1 [Fig. 4A; Supplemental Fig. S4; Supplemental Movie S3]. In our fluorescence images, ER patches were often enriched near pronuclei [Fig. 4A, Supplemental Fig. S4A]. Consistent with the idea that the ER patches detected at the light level are composed of ER sheets, examination of nuclei in cnep-1Δ embryos by TEM revealed a substantial increase in the number of ER sheets near the nuclear envelope [Fig. 4B]. Unlike control embryos, where very few streaks of reduced electron density (consistent with an ER sheet in cross-section) were detected, many such sheets were present near nuclei in cnep-1Δ embryos. We therefore propose that the extra ER layer that encases the nucleus following permeabilization in cnep-1Δ embryos is composed of perforated ER sheets.

Wrapping of the nuclear envelope in an extra ER layer in cnep-1Δ embryos suggests that CNEP-1 prevents ER sheets from stacking onto the nuclear envelope by ensuring that the PI concentration on the nuclear envelope and ER membranes in the vicinity of the nucleus remains low (model in Fig. 4C). As the nuclear envelope and ER are a single contiguous membrane system, we expect that phospholipids diffuse between the ER and nuclear envelope while at the same time being acted on by localized biosynthetic enzymes that bias their distribution. In the absence of CNEP-1, higher PI levels would cause the ER near the nuclear envelope to form an extensive network of sheets that encases the permeabilized nuclear envelope, slowing its disassembly [Fig. 4C]. The fact that the second ER layer in cnep-1Δ embryos only encases the nuclear envelope following its permeabilization suggests the existence of a mechanism that prevents ER sheets from stacking onto the nuclear envelope during interphase. This mechanism may be inactivated during mitosis when the nuclear envelope has been shown to invaginate and thus transiently form stacked membrane segments to aid in nuclear envelope disassembly [Calarco et al. 1972; Galy et al. 2008]. It is also interesting that the nuclear envelope is uniformly encased in a single additional ER layer, suggesting that whereas stacking of ER sheets onto the nuclear envelope is permitted, stacking of ER sheets onto each other is not.

The mechanism that we propose to explain how lipin/CNEP-1 inhibition delays nuclear envelope disassembly is significantly different from the previously proposed mechanism, based on work in mammalian cells, in which lipin inhibition was suggested to decrease DAG-induced activation of protein kinase Cs, thereby slowing lamin phosphorylation and disassembly [Mall et al. 2012]. Future work will be needed to determine whether this discrepancy is due to a difference between systems or whether a mechanism similar to the one that we propose also underlies the nuclear envelope disassembly defect in mammalian cells.

The idea that PI synthesis is spatially regulated to control ER structure is consistent with recent work showing that phospholipid synthesis enzymes, including phosphatidylinositol synthase, are enriched in a Rab10-dependent ER subdomain and that this compartmentalization is important to prevent the excessive formation of ER sheets [Kim et al. 2011; English and Voeltz 2012]. How PI levels control the extent of ER sheet formation and whether the relevant lipid is PI itself or another PI-derived PIP is important future questions. Asymmetric distribution of lipids with curvature-favoring structures between the two membrane monolayers in a bilayer can generate membrane curvature. However, high curvature is thought to require levels of compositional asymmetry that are unlikely in vivo [Shibata et al. 2009]. This, combined with recent work characterizing membrane-associated proteins that shape ER tubules and sheets, has led to the conclusion that proteins are the main players in controlling ER structure [Shibata et al. 2009; Goyal and Blackstone 2013]. It is therefore attractive to think that higher PI concentrations could promote sheet formation by modulating the function of ER-shaping proteins; testing this hypothesis will be an important goal of future work.

Materials and methods

C. elegans strains and RNA-mediated interference

C. elegans strains and dsRNAs are listed in the Supplemental Material. L4 or young adult hermaphrodites were injected with dsRNA and incubated for 12–48 h at 20°C before dissection to obtain embryos for filming [for details, see the Supplemental Material].

Immunoblotting

Affinity-purified antibodies against amino acids 25–161 of LPIN-1 were generated using standard procedures. Antibody signals were detected using an HRP-conjugated secondary antibody (GE Healthcare) and the WesternBright Sirius detection system (Advansta).

Light microscopy

Embryos were imaged at 20°C with a 60×1.4 NA Plan Apochromat lens on an Andor Revolution XD Confocal System (Andor Technology) mounted on an inverted microscope (TE2000-E, Nikon) equipped with a confocal scanner unit (CSU-10, Yokogawa), solid-state 100-mW lasers, and a high-resolution interline CCD camera (Clara or iXon, Andor Technology). Images of the ER lumen marker were acquired using an inverted Zeiss Axio Observer Z1 system with a Yokogawa spinning-disk confocal head (CSU-X1), a 63×1.4 NA Plan Apochromat objective, and a QuantEM:512SC EMCCD camera (Photometrics).

Phospholipid analysis

Worms were labeled overnight with 1 mM/mL phosphorus-32 orthophosphate (PerkinElmer), and lipids were extracted following standard protocols (Bligh and Dyer 1959). Total phospholipids were spotted in triplicate, separated by thin-layer chromatography (60:35:10 solution of chloroform:methanol:20% methylamine), and detected by exposing onto film (for details, see the Supplemental Material).

High-pressure freezing and electron microscopy

Whole animals were high-pressure-frozen and freeze-substituted. Samples were embedded, and images of post-stained ultrathin sections were taken using an FEI Tecnai Spirit TEM at 80 kV (Fig. 2) or an FEI Tecnai 12 TEM operated at 100 kV (Fig. 4). For details, see the Supplemental Material.

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