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Acute Depletion of Diacylglycerol from the Cis-Golgi Affects Localised Nuclear Envelope Morphology During Mitosis

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Abstract

Dysregulation of nuclear envelope (NE) assembly results in various cancers; for example, renal and some lung carcinomas ensue due to NE malformation. The NE is a dynamic membrane compartment and its completion during mitosis, is a highly regulated process but the detailed mechanism still remains incompletely understood. Previous studies have found isolated diacylglycerol (DAG) containing vesicles are essential for completing the fusion of NE in non-somatic cells. We investigated the impact of DAG depletion from cis-Golgi in mammalian cells on NE reassembly. Using advanced electron microscopy, we observed an enriched DAG population of vesicles at the vicinity of the NE gaps of telophase mammalian cells. We applied a miniSOG–C1-domain tag that localized DAG-enriched vesicles at the perinuclear region, which suggested the existence of NE fusogenic vesicles. We quantified the impact of Golgi-DAG depletion by measuring the in situ NE rim curvature of the re-forming NE. The rim curvature in these cells was significantly reduced compared with controls, which indicated a localized defect in NE morphology. Our novel results demonstrate the significance of the role of DAG from cis-Golgi for the regulation of NE assembly.
Introduction

The nuclear envelope (NE) is a sub-compartment of the endoplasmic reticulum (ER) and it breaks down and reforms during an open mitosis. However, the detailed molecular mechanism regarding its reassembly remains to be determined. Although envelopment of the chromatin by the ER is one of the NE reassembly models (1) and contact between ER cisternae and chromosomes has been shown to initiate mammalian NE reassembly at anaphase (2), these descriptions do not explain the necessity of membrane fusion or the possible requirement of correctly formed curved regions for the localisation of the nuclear pore complexes (NPCs).

Proteins play an important role in membrane fusion and morphology (3-6) but these mechanisms also require lipids. Membrane fusion can be facilitated in a protein-free context (7, 8) meaning lipids are critical effectors during the NE fusion.

To explain membrane fusions in NE completion, the vesicle fusion model provides an additional mechanism (9). Membrane fusion as well as membrane regions with high curvature such as the NE rims are promoted by the presence of negatively curved lipids, where diacylglycerol (DAG) destabilises the localised membrane morphology (10-12). The most curved region of the NE, which exists between the inner and outer nuclear membranes where nuclear pore complexes are recruited, is referred to as the nuclear envelope rim.

As mentioned above, neutral lipids such as DAG, which have a high negative spontaneous curvature, locally destabilise the lamellar bilayer to induce membrane fusion (13, 14). The understanding of how this lipid behaves during membrane fusion has mostly resulted from in vitro or reconstitution experiments (10, 11, 15) and its complex behaviour has not been directly examined in live cells or model organisms.
Our group has isolated a population of membrane vesicles known as MV1, a distinctive membrane compartment in vivo, with atypical polyphosphoinositide compositions, (16), as one of the critical NE precursors of echinoderm male pronuclei. MV1 has elevated levels of phosphoinositides, a lipid-modifying enzyme PLCγ and its upstream regulator src family kinase 1 (SFK1) (16, 17). These three proteo-lipid components are the fusion “tool-kit” responsible for localised production of DAG.

In mammals, DAG localises at the NE, ER and Golgi (18). Using a rapalogue based heterodimer system (19, 20), we have demonstrated the structural role of DAG in mammalian NE reassembly where localised depletion of DAG at the NE and ER results in failure of NE reassembly and abnormal ER morphology (18). The abnormal ER morphology and fragmented NE from these experiments resulted in taking our experiments forward to determine whether localised highly curved regions, such as the rim curvature of the NE would be affected by the acute and inducible depletion of DAG. To examine this we set out to investigate the impact of DAG depletion from the cis-Golgi, a reservoir of DAG in mammalian cells, on mammalian NE reassembly using the rapalogue dimerisation approach.

Here we show that locally depleted DAG from the cis-Golgi impacts the NE rim curvature of the reforming NE. The rim curvature in these cells is significantly reduced compared to the controls. We measured the in situ rim curvature and the pore diameters to illustrate the localised defect in NE morphology. These membraneous defects may result in affecting the proper nuclear pore insertions. Our results demonstrate the significance of the role of DAG from the cis-Golgi for the regulation of nuclear envelope assembly.
Materials and Methods

Cell line and maintenance

HeLa cell line was obtained from the American Type Culture Collection (ATCC), ATCC#: CCL2. The cells were grown in DMEM containing 10% fetal bovine serum (FBS, Gibco), supplemented with penicillin (100 U/ml) and streptomycin (100 µg/ml) and incubated at 37 °C in 10% CO₂.

Constructs

GRASP65-GFP and its non-Golgi-targeting mutation GRASP65.G2A-GFP were gifts from Prof. Yan-Zhuang Wang (University of Michigan). GRASP65-GFP-2FKBP and GRASP65.G2A-GFP-2FKBP were created by adding a 2FKBP domain into the given constructs at its C terminus with flanking BsrGI sites. The GRASP65, GFP tag and 2FKBP rapalogue-binding domain, were made into such sequence so that the constructs can successfully localise to the Golgi and recruit DGKεK. GFP-2FKBP-LBR, RFP-Flag-FRB-DGKεK and RFP-Flag-FRB-DGKεK.D434N were created as described in (18).

Confocal imaging

Cells were grown on gridded glass coverslips in 3.5-cm dishes (MatTek Corporation) and double transfected with the rapalogue dimerisation constructs described above. Cells were transfected with 0.5 µg DNA of each construct using Lipofectamine LTX and PLUS reagent (Invitrogen) in OPTIMEM medium (Gibco BRL) as recommended by the manufacturer. The cells were left overnight in the transfection mix in antibiotic-free medium before replacing it with fresh medium. Experiments were performed 36–48 hours post transfection. The transfected cells were stained with ER tracker Blue-White DPX (Invitrogen) at a final concentration of 1 µM according to manufacturer's protocol before the rapalogue experiment (18, 19). Cells were incubated in a 5 L/h
humidified chamber with 10% CO2 adapted to a Zeiss confocal microscope (Zeiss LSM 710). Images were acquired prior to and after the recruitment of DAG kinase to the targeted membrane compartment, ~30min – 1h after addition of 500 nM rapalogue heterodimeriser (Clontech). Cells were then imaged every 30–45min at low pixel resolution (512 x 512 with a 0.6x zoom) from interphase to late anaphase to limit laser damage. A high pixel resolution (1024 x 1024 with a 2x zoom) series of images was also acquired from late anaphase to cytokinesis. Each experiment was performed at least three times. All images were treated the same manner; i.e only minor adjustments of brightness and contrast were applied to every pixel.

**Correlative light and electron microscopy (CLEM)**

Live cells were followed to the required stage of mitosis using confocal microscopy. Cells were then fixed in 4% EM grade formaldehyde (TAAB) in 0.1 M phosphate buffer (PB) pH 7.4 to halt the cell cycle prior to re-imaging for brightfield and high magnification fluorescence signals. Cells were subsequently located using the grid number of the gridded glass coverslip (MatTek). Secondary fixation was performed in 1.5% glutaraldehyde/2% formaldehyde in 0.1 M PB for 30–60min. After fixation, coverslips were carefully removed from the MatTek dishes and washed several times in 0.1 M PB.

For TEM, the cells were post-fixed in 1.5% potassium ferricyanide/1% osmium tetroxide for one hour, before rinsing in PB, and incubating in 1% tannic acid in 0.05 M PB for 45 minutes to enhance membrane contrast. After a brief rinse in 1% sodium sulphate in 0.05 M PB, the coverslips were washed twice in distilled water, dehydrated through an ascending series of ethanol to 100% prior to infiltration with Epoxy resin and polymerisation overnight at 60°C. The coverslips were removed
from the resin blocks by plunging briefly into liquid nitrogen. The cells of interest were identified by correlating the grid and cell pattern on the surface of the block with previously acquired confocal images. The area of interest was cut from the block and further trimmed by hand using a single edged razor blade to form a small trapezoid block face for serial ultrathin sectioning. Using a diamond knife, serial ultrathin sections of 70 nm thicknesses were cut through the entire extent of the cells of interest (80 to 140 sections) and collected on 1.5% formvar-coated single slot grids. The sections were counterstained with lead citrate to further enhance contrast prior to viewing in the electron microscope (FEI Tecnai G2 Spirit BioTWIN with Gatan Orius CCD camera). Serial images were stacked and aligned, and the NE, ER and centrioles and vesicles were manually segmented using Amira (FEI), based on their electron density and morphological features. Circular membrane structures with similar x, y and z diameter were segmented as vesicles. Movies were created from 2D tiff stacks using Quicktime Player 7 Pro and compressed using Stomp (Shinywhitebox Ltd).

For serial blockface scanning electron microscopy (SBF SEM), the fixed cells were processed following the method of the National Centre for Microscopy and Imaging Research (21), which impregnates the sample with high concentrations of heavy metals to introduce maximal contrast and conductivity when viewed in the SEM. The cell of interest was identified by correlation of grid reference with previously acquired confocal images; this area was trimmed to a small trapezoid, excised from the resin block, and attached to a SBF SEM specimen holder using conductive epoxy resin (Circuitworks CW2400). Prior to commencement of a SBF SEM imaging run, the sample was coated with a 2-nm-layer of platinum to further enhance conductivity.
SBF SEM data was collected using a 3View2XP (Gatan, Pleasanton, CA) attached to a Sigma VP SEM (Zeiss, Cambridge). To relocate the cell of interest in the SEM, an overview was first acquired at 5 kV, sufficient to penetrate the platinum coating and generate an image of the underlying sample. Inverted backscattered electron images were then acquired through the entire extent of the cell of interest at a resolution of 8,192 x 8,192 pixels (horizontal frame width of 36.74 µm; pixel size of 4.5 nm) using a 2 µs dwell time and 50 nm slice thickness. The SEM was operated in variable pressure mode at 5 to 10 Pa, with high current mode active, 20µm aperture, an accelerating voltage of 2 kV, and an indicated magnification of 7,000. Typically, around 400 slices were necessary to image an entire cell, representing a total volume of approximately 27,000 µm³.

As data was collected in variable pressure mode, only minor adjustments in image alignment were needed, particularly where the field of view was altered in order to track the cell of interest.

**Electron tomography**

For electron tomography, samples were prepared as detailed above but 200-nm-thick serial sections were collected through the entire extent of the cells of interest. Tomograms were acquired from the 200 nm sections at targeted regions in the reforming nuclear envelope – either at specific gaps, or where collections of vesicles were evident in close proximity to the reforming envelope. Images were collected at 1-degree intervals across a maximal tilt range of ±70°, with 0.79 nm width per pixel for 2,048 x 2,048 pixels, with a per pixel resolution of 0.79 nm. Tomograms were processed with IMOD (22), using patch tracking for alignment, and simultaneous iterative reconstruction technique (SIRT) for volume reconstruction. The reconstructed volume was exported as a series of 2D tiff images, and the NE and
adjacent membranous structures, including vesicles, were manually segmented and reconstructed using Amira (Visage Imaging, Berlin). Movies were created from the 2D tiff stacks using Quicktime Player 7 Pro, and compressed using Stomp (Shinywhitebox Ltd). It is of note that one cannot negate embedding artefacts enough to have a definitive measurement of curvature and so cryo-microscopy is preferred; however, the cryo-tomography required is extremely technically challenging and cannot currently be performed for rare correlative events due to complexity of sample preparation and difficulties in targeting specific cells through the cryo workflow. It should be noted that if this methodology were to be used for a larger study, or under varying imaging conditions, there would be a justification to verify these trends in a case-by-case manner.

**Segmentation of ultrastructure**

Serial micrographs were stacked and aligned using Amira (Visage Imaging, Berlin). The pixels of NE, ER, centrioles and vesicles were manually traced based on their electron density and morphological features on the micrograph. Circular membrane structures with similar xy and z diameter were segmented as vesicles in the TEM analysis of the telophase cell. Discontinuities of the reforming NE around the chromatin were segmented as the NE gaps.

**MiniSOG photo-oxidation**

Cells were transfected with GFP-PKCeC1aC1b-SOG and light microscopy performed to identify cells of interest prior to initial fixation with 4% formaldehyde in 0.1 M PB, and secondary fixation with 2.5% glutaraldehyde in 0.1 M PB. The cells were washed in 0.1 M PB and incubated in blocking buffer (50 mM Glycine, 10 mM KCN, 20mM aminotriazole in 0.1 M PB) for 30 minutes. The dishes were then loaded into a customised cooling chamber to maintain the samples at ~4°C. The lid of the chamber
was connected to an oxygen supply at a rate of 2.5 l/min. The cells were then incubated in diaminobenzidine solution (5.4 mg in 10 ml 0.1 M PB) and photo-oxidised for 30 min using a Hg lamp, after which they were further fixed on ice with 2% formaldehyde/1.5% glutaraldehyde in 0.1 M PB for 30 minutes, and post-fixed with 0.5% osmium tetroxide in 0.1 M PB for 30 minutes before embedding as described above for CLEM. The photo-oxidised cells were relocated using the MatTek grid coordinates, and serially sectioned for examination using TEM.

**NE Rim Curvature Measurements**

Blocks, stained as per the serial block face method, were instead serial sectioned at 300 nm onto pioloform coated slot grids. 15 nm gold particles (Aurion) were drop cast for 5 min on both sides with the excess blotted off. Micrographs were acquired through the cell and two areas selected on each of the daughter cells. Single tilt tomograms were taken of the selected areas using a 200 KeV transmission electron microscope (T20, FEI) with a LaB₆ filament and a 4k x 4k Eagle (FEI) bottom mounted camera. The tilt series was taken, using the FEI software, at 2° intervals between ±60° where possible, at a nominal magnification of 14,500x, giving pixel sizes of 0.74 x 0.74 nm². The tilt series were reconstructed using IMOD (22), subsequent image analysis using Fiji (23), and statistical analysis using Prism (GraphPad).

Tortuosity of the nuclear membranes could not be analysed quantifiably from the 3D tilt series reconstructions, due to the volumetric field of view. Instead, if the membrane curvature has been disrupted, a requirement for pathological tortuosity, it would be expected to be evident at the tight curvatures between the inner and outer
membranes defined here as Rim Diameter to be consistent with modelling of the ER (24).

To achieve an estimate of any altered curvature it was decided to quantify the nuclear pores. Two measures were taken (Figure 4): The diameter of the pore and the distance between the inner and outer nuclear membrane, its Rim Diameter. Initially to achieve good membrane contrast the 3D reconstructed image stacks were reduced, by binning by a factor of four (leaving $3 \times 3 \times 3 \text{ nm}^3$ voxels in $x$-$y$ and $z$ directions), and dynamic range to 8-bit after suitable windowing. Three-dimensional image clips were taken from every nuclear pore completely enclosed in the reconstruction as described by Figure 4. Each clip was rotated such that the membranes are parallel to the $x$-axis, then rotated around the $x$-axis by $10^\circ$ increments until the pore appeared by eye as near to a circle as possible. From this point the clip was rotated back by $90^\circ$ making the pore perpendicular to the viewed plan. The width and rim diameter were then measured at a depth of the widest diameter of the pore by measuring the distance between centres of the dark bilayer membranes where they just become parallel.
Results

The presence of vesicles in the vicinity of the reforming NE

In our previous work on the echinoderm model, we showed that MV1, a vesicular compartment enriched in PLC\(\gamma\) and phosphoinositides, was recruited to each stage of membrane fusion, required for NE and zygote formation (17). Based on these results we predicted that vesicles enriched in phosphoinositides or their derivative DAG may also play a role in NE assemble in mammalian cells.

To demonstrate the presence of vesicles that may participate in NE completion at late mitotic stages, we investigated the ultrastructure of telophase cells by transmission electron microscopy (TEM). Figure 1a and Figure 1c show that multiple gaps existed at the NE of the examined telophase cells, and vesicles were detected in the proximity of these gaps. Electron tomography (ET) confirmed these membrane structures occupied a spherical space in the virtual volume of the tomogram (Figure 1c and Supplementary Movie 1) implying the presence of vesicles as opposed to tubules. Some vesicles were very close to the edge of the reforming NE as indicated by the reconstructed 3D model (Figure 1d). However, there were numerous vesicles in a telophase cell. A comprehensive 3D reconstruction revealed at least 14,000 vesicles in one of the daughter cells at this mitotic stage (Figure 1b). Therefore, we focused on studying the vesicles that were close (500 nm) to the reforming NE since physical contact is required for membrane fusion.

The number of vesicles is significantly higher at the large gaps

To understand how vesicles distributed around the reforming NE of a telophase cell, we compared the concentration of vesicles at the regions of gaps and regions without gaps (Figure 2). A gap was defined by the shortest distance between the two converging edges of the reforming NE in the \(x\)-direction multiplied by its thickness in...
the z-direction, not exceeding 140 nm. The size range of the gaps varied within three orders of magnitude, therefore, we grouped the gaps into small (5,800 to 9,900 nm²), medium (11,000 to 82,000 nm²) and large (100,00 to 350,000 nm²), based on their vertical area (x-z) in the micrographs (Figure 2a). A cylindrical volume was defined as the “vicinity” of different regions of interest and the volume has a radius of 1 µm so that it did not exceed the boundary of the cell. For a gap, one of its ends was selected as the centre of the cylindrical volume (Figure 2b). We compared the concentration of the vesicles in the "vicinity", with the same volume at different regions. The vesicle concentration at the large-gap-regions was significantly higher than that in other regions, whereas at medium-gap-regions and small-gap-regions the vesicle concentration reduced in relation to the gap size (Figure 2c). This might imply that the vesicles in the large-gap-regions were being utilised to induce membrane fusion of the NE fragments. The mean diameter of all the vesicles in the close field was 57±0.5 nm (Figure 2d).

**DAG-enriched vesicles in the proximity of the NE**

To identify DAG-enriched vesicles we exploited the C1a-C1b domain of PKCε and its non-binding mutant C1aC1b-W264G. This domain specifically recognises the subcellular localisation of DAG. To illustrate the DAG specificity the non-binding mutant was utilised (Supplementary Figure 2). Moreover, to enable the direct recognition of these vesicles at the EM level, at the vicinity of the NE, a miniSOG (mini Singlet Oxygen Generator) approach was utilised. The conventional methods (e.g. immunogold) for TEM were inappropriate since the permeabilisation step extracted the lipids of interest. The miniSOG tag (25) provided TEM contrast, via photo-oxidation. This step released oxygen to polymerise 3,3'-diaminobenzidine
DAB) at the targeted structures which attracted preferential osmium staining. Figure 3a shows that many vesicles were labelled with GFP-PKCεC1aC1b-SOG reaction products in an interphase cell at the perinuclear region. The mean size of the miniSOG-labelled vesicles was greater due to the formation of the DAB polymer (Figure 3b). Hence this demonstrated that the C1a-C1b domain has recognised the DAG-enriched vesicles. If the C1 domain did not bind to DAG the size of the vesicles would remain unchanged. We suggest that these DAG-enriched vesicles are present at interphase for the maintenance of the NE morphology.

**Golgi-DAG depletion by the rapalogue dimerisation system**

To investigate the localised and inducible impact of Golgi-DAG depletion on NE reassembly, during mitosis, we used the rapalogue (rapamycin analogue) dimerisation system and the Golgi-targeting domain of GRASP65 (Golgi reassembly stacking protein 65), a cis-Golgi marker expected to be co-recruited with DAGKinaseε (DGKeK), providing spatial specificity. The C1 domain of DAGkinase specifically targets DAG [18].

Cells were transfected with the GRASP-GFP-2FKBP and RFP-Flag-FRB-DGKeK. Confocal images were acquired prior to and after the recruitment of DGKeK to the targeted membrane compartment, ~30min to 1h after addition of 500 nM rapalogue heterodimeriser. Cells were imaged every 30–45min from interphase to late anaphase to limit laser damage. A series of high-resolution images were also acquired from late anaphase to cytokinesis. They were fixed after reaching late telophase where the NE should be fully assembled.

DAG can be observed at the Golgi in addition to the NE and ER (Figure 4a). Figure 4b shows that the rapalogue construct GRASP-GFP-2FKBP has a similar localisation compared to the original construct GRASP-GFP and the reported non-Golgi-binding
mutation (G2A) indicated that the GRASP domain is Golgi-specific. Figure 4c indicates the successful recruitment of DGKεK to the Golgi when rapalogue was added.

To examine the influence of DAG depletion on the formation of the NE at the late mitotic stages, the cells were fixed at late telophase/cytokinesis. Figure 5 shows the interventions by the rapalogue dimerisation system imaged by fluorescence microscopy. DGKεK and its catalytically inactive mutant DGKεK.D434N were recruited to the Golgi upon rapalogue addition (Figure 5a-b). ER tracker Blue-White DPX functions as the marker to determine if the NE was reformed. In both Golgi-DAG depleted (DGKεK recruited) and Golgi-DAG unperturbed conditions (DGKεK.D434N recruited), signals of the ER tracker were observed around the chromatin. Unlike the ER/NE-DAG depletion (Figure 5c), where the NE was totally fragmented at the fluorescence microscopy level (18), an obvious change here was not perceived. This was expected, as the depletion of DAG from the ER would affect prominently the NE morphology as NE is a sub-unit of the ER.

To observe whether a minor fragmentation occurred at the electronic resolution, the complete 3D NE ultrastructure in Golgi-DAG depleted and Golgi-DAG unperturbed cells was examined by serial block face scanning electron microscopy (SBF SEM), and again only a mild or no NE fragmentation was observed in both conditions (Supplementary Figure 1).

We suggest that Golgi-DAG depletion was insufficient to cause a general NE morphological defect. However, this would not rule out the effect Golgi-DAG depletion may have on the finer structures of the NE. We therefore, postulated that there would be localised morphological defects at high curvature regions of the NE where DAG may have a localised role in maintaining these high curvature regions.
Golgi-DAG depletion causes decrease in nuclear envelope rim curvature

To test this postulation we measured, using a higher resolution imaging technique, the most curved region of the NE, which exists between the inner and outer nuclear membranes where nuclear pore complexes are recruited. This region is referred to as the nuclear envelope rim. The NE rim curvature was utilised to determine the effect of the Golgi-DAG depletion on localised membrane curvature (24).

Four experimental conditions were compared: DAGKinase with rapalogue (DKGεK+Rap), DAGkinase mutant with rapalogue (D434N+Rap), DAGkinase without rapalogue (DKGεK-Rap) and untransfected cells. From these four conditions, nuclear pores from 21 tomograms, were sampled evenly across the NE of telophase cells and measurements were acquired under these conditions. Beam induced mass loss causes anisotropic section shrinkage usually thought of as approximately 30% in thickness and 2-8% in x-y. To ensure that our measurements were not biased by the anisotropy of section shrinkage in ET, samples with the most pores from four tomograms (DKGεK+Rap) were initially tested for dependence on the pore’s angle in the sample. We determined that there was no significant dependence (p=0.089) on the rotated angle to rim curvature measurement but there was a dependence (p=0.046) on pore diameter to the angle (Pearson, data not shown). This is likely a low dependence due to the hard resin used and not using excessive beam intensity. Nevertheless, the trend in the measurements was well below the innate variation and therefore considered as insignificant for further analysis.
The rim diameters measured from where the inner and outer membranes become parallel were quantified (Figure 6a-b) and we determined that the four groups did not all fit a normal distribution. Therefore, a Kruskal-Wallis with Dunn’s multiple comparison test was performed. There was an indication that the DKGεK+Rap rim diameter measurements had a bimodal distribution. The rim diameter was significantly different only in comparison with DKGεK+Rap indicating the principle of our measurement criteria was valid and useful. This is indicative of the alteration of membrane curvature in the localised absence of DAG.

It was observed, but not quantified, that the DKGεK+Rap seemed to have an anomalous rim diameter change, with both the rim curvatures and the pores not in the plane of the envelope as normally observed.

The alignment of nuclear pores at the rim of the NE can be a parameter for NE defects if the analysis were performed on a larger scale. The rim diameters were converted into curvature (κ) by κ = 2/RD. The median ± interquartile range/2 for κ_untransfected = (0.077±0.010) nm⁻¹ but was significantly reduced (p<0.0001) for the Golgi-DAG depleted cells where κ_DKGεK+ Rap = (0.060±0.014) nm⁻¹. This demonstrated that without the DAG from the Golgi a tight localised curvature was not formed.

The pore diameter measurements (Figure 6c) fitted a normal distribution. Therefore, a one-way ANOVA analysis with Tukey’s multiple comparisons test was performed to determine the statistical significance. The curvature of the pore ring is far less than the rim diameter and only the DKGεK-Rap gave a significant difference in pore diameter (Figure 6c). It is difficult to propose what may have caused this. Perhaps it was due to experimental rather than biochemical differences.

The four tomograms were taken in the same session under identical conditions marking the highly significant difference of the rim curvatures within DKGεK+Rap. It
was unlikely that the other conditions produced different measurements due to an artefact as the conditions were kept as similar as possible.

In summary, this new quantitative method of measurements provides \textit{in situ} membrane curvature values. It is by using such a quantitative method that we demonstrated, a new concept that the tight rim curvature of the NE requires DAG that is targeted from the Golgi to the gaps during NE formation.
Discussion

Membrane vesicles have been isolated as NE precursors from several non-mammalian cell-free assays decades ago (26-28). These vesicles have been shown to constitute a specific cellular compartment enriched in membrane fusion machinery in situ (16, 17). Nevertheless, the existence of vesicular NE precursor in mammals is unknown. Using TEM and electron tomography, we have detected vesicles in the vicinity of the reforming NE in telophase cells (Figure 1), and the concentration of these vesicles is significantly higher in the large-gap-regions than in regions of smaller gaps and regions without gaps. Such observation suggests that vesicles play a role at the gaps for NE completion (Figure 2). Moreover, it is unlikely that completion of the NE is solely contributed by the ER cisternae.

In the echinoderm NE assembly assay, Golgi-derived vesicles are required for recruiting the ER and completing the NE of male pronuclei through membrane fusion (29). In mammals, there is evidence for the presence of DAG (30) and its precursor PtdIns(4,5)P₂ (18, 31) at the Golgi. A recent study has shown that PLCγ1, which hydrolyses PtdIns(4,5)P₂ to DAG, participates in Golgi-DAG production (32). Furthermore, Golgi membranes (vesicles and cisternae) are detected in the perinuclear region during late mitotic stages (33, 34). These findings are reminiscent of the fusion machinery-enriched vesicles needed for NE reassembly in vitro (17) and the perinuclear vesicles detected in this study.

We applied GFP-miniSOG to visualise DAG in fluorescence and electron microscopy. This is the first study to access the usefulness of miniSOG constructs in labelling small structures such as vesicles. Our results indicate the presence of DAG-enriched vesicles in the cytoplasm of interphase cells, and some of these vesicles localised in the perinuclear region (Figure 3). We suggest that these vesicles are required for the
maintenance of the nuclear envelope morphology and perhaps would participate in NE completion in later mitotic stages.

When we recruited high levels of DGKεK to the Golgi the newly formed NE was not grossly fragmented and the quantification of the number of gaps created due to the lack of DAG at the Golgi was inconclusive. The catalytically inactive DGKεK did not cause any observable phenotype (Supplementary Figure 1).

To test whether the lack of DAG at the Golgi affected a more definite structure of the NE we developed a novel quantitative method for measuring in situ curvature of the NE rim curvature. The results showed that the rim curvature in the absence of DAG from the Golgi was significantly lower than in its presence. The catalytically inactive DGKεK did not affect the rim curvature. The curvature of the NE pore is far less than the NE rim diameter so one would not expect it to be as effected by DKGεK alteration and this is born out in the results. We have demonstrated a suitable quantitative methodology for determining curvature alteration, which in turn could be linked to the pore complex binding to the NE rim. Our current methodology has limitations due to dehydration artifact in the EM processing causing more tortuous membranes. The main result is that measurements have more variation so if anything would make significance harder to achieve. This error could be reduced if necessary by cryo-techniques such as high pressure freezing and freeze substitution of the aqueous component.

Our investigations have now provided evidence for the regulatory role(s) of both NE-(18) and Golgi-DAG during mitosis. A follow-up question is how are these vesicles targeted to the gaps in such a manner that would deliver the required localised DAG for maintaining the appropriate NE rim curvature.
The presence of chromatin-targeting proteins, such as LBR, in NE precursors, has been reported (35, 36), and this could provide a mechanism. However, the perinuclear vesicles have a small diameter of ≈ 55 nm as well as constrained negative curvature and therefore they may not have sufficient surface for binding receptors. We speculate that membrane tethering, recruitment of membrane to a specific location by peripheral membrane proteins (tethers), is a mechanism to recruit vesicles to the gaps. Many tethers are originated from the Golgi (37), among them the p115 that has been reported as a general fusion factor interacting with other Golgi-proteins such as giantin, GM130 and GRASP65 (38, 39). The trafficking direction and mechanism of p115-dependent tethering remains controversial but it is notable that p115 has a capacity bind to various targets (40). Interestingly, p115 does not bind to mitotic Golgi (41), giving us a hint that p115 may interact with NE/ER proteins and recruit Golgi vesicles to the reforming NE. Moreover, membrane curvature sensors may assist the recruitment of vesicles to the gaps. For instance, munc13 has a C1 domain targeting DAG and has been reported as regulating membrane fusion in different compartments (42-44).

Overall, in this study we have developed a quantitative methodology that has permitted us to address the importance of the DAG localised at the cis-Golgi and how it may be implicated in maintaining and “fine-tuning” the morphology of the nuclear envelope in mammalian mitosis.
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Contributor Roles

GC designed, analysed and performed the experiments
KPA: Designed and took 3D-TEM data for rim curvature quantification.
KM and AA: Data analysis for robustness of rim curvature measurements
LH: Performed ‘control’ HeLa cell experiments for rim curvature measurements
JM: Took 3D-TEM data for rim curvature measurements.
CP assisted with miniSOG experiments, ET, reconstruction and segmentation of data.
MCD prepared cells for SBF SEM, and advised on confocal experiments and analysis.
LC: collected the SBF SEM data.
BL: supervised the whole project.
BL, GC, LC and KA wrote the manuscript.

All authors contributed intellectually to manuscript preparation.
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Figure 1

(a) Unmodified electron micrograph

(b) 3D model (ER excluded)

(c) Telophase cell

1. Identification of gap
2. Identification of gap (magnified)

(d) Z-direction progression of electron tomogram

- 3D model
  - 0°, 60°, 120°, 180°, 240°, 300°
Figure 1: Detection of perinuclear vesicles by TEM and electron tomography

(a) A HeLa cell was fixed at telophase (inset), cut into serial sections and prepared for TEM analysis. Cellular structures and gaps were traced manually (segmentation) in different colours: NE (red), ER (cyan), centriole (purple), vesicles (yellow), gaps (green). Top and bottom panels depict the unmodified and segmented electron micrographs respectively. The white arrow indicates the vesicles close to a NE gap. (b) A 3D model displaying all segmented structures described in (a) except the ER. A large number of vesicles (n = 14,000) was detected in the telophase cell. (c) Top panel: electron micrographs from a 200 nm-thick section taken through a telophase HeLa cell. The identified gap (yellow arrow) was analysed by electron tomography to produce a tomogram with a thickness of 163 nm. Bottom panel: snapshots of the unmodified and segmented tomogram along the z-axis, with the reforming NE segmented in red and vesicles segmented in yellow. White arrows indicate virtual space occupied by the vesicles. N: nucleus, C: cytoplasm. (d) 3D model reconstructed from the segmented tomogram at different angles. Scale bar, as indicated.
Figure 2: Quantitative analysis of gaps and vesicles at the NE of telophase cells

(a) There were 76 gaps at the NE of the cell described in Figure 1a. The gaps were categorised based on their size. Small (n = 22), medium (n = 46) and large gaps (n = 8) differed in their area by an order of magnitude. (b) A circular sampling field with a radius of 1 µm was drawn, the centre of the sampling field is either an end point of a gap or a random point at the NE as a control (n = 10). In the field, the number of vesicles and their distance from the centre were measured. The field was subdivided into the close and far field as indicated. (c) The number of vesicles per area analysed by one way-ANOVA. The number of vesicles in both close and far field of the large gaps was significantly higher than that in other regions. Asterisk scale: * P ≤ 0.05, ** P ≤ 0.01, *** P ≤ 0.001, **** P ≤ 0.0001. (d) The diameter of vesicles plotted into a histogram.
Figure 3: DAG-enriched vesicles at the NE of interphase cells
(a) Electron micrographs of an interphase HeLa cell transfected with the DAG reporter (GFP-C1aC1b-SOG) for 16 h. The transfected cell was fixed and photo-oxidised in the presence of DAB to enhance local contrast during EM sample preparation. Red boxes in dash corresponding to the magnified regions of interest (ROI). In the electron micrographs, numerous vesicles had enhanced electron density and were distinguished from the background EM contrast (yellow arrows). N, nucleus; C, cytoplasm. Scale bars, as indicated.
(b) The diameter of miniSOG labelled vesicles in the proximity of the NE (within a radius of 1 µm, as described in Figure 2b) was measured as described in Figure 2d and plotted into a histogram. The mean diameter of these vesicles was 71 nm and most of the vesicles (n = 18) fell into the 65 and 70 nm-group.
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Figure 4

original constructs

rapalogue
dimerisation tools
Figure 4: The rapalogue dimerisation system that targets Golgi

(a) DAG detection by GFP-PKCεC14C15. (b) Localisation of the original GRASP65 constructs and its rapalogue dimerisation unit-conjugated counterparts. GRASP65.G2A is a non-Golgi-binding mutation. (c) Left: diagram of the rapalogue dimerisation constructs that target DAG at the Golgi. Upon addition of rapalogue, RFP-Flag-FRB-DGKεK (DGKεK), a lipid-modifying enzyme, is expected to be recruited to GRASP-GFP-2FKBP (Golgi) through the formation of an RFP-FKBP heterodimer so that DAG at the Golgi will be converted to phosphatidic acid (PtdOH). Right: confocal images showing the successful recruitment of DGKεK (magenta) to the Golgi (green). Yellow arrows indicate the Golgi localisation of both constructs 2h post rapalogue addition.
Figure 5

Golgi-DAG depleted

Golgi-DAG unperturbed

NE-DAG depleted
Figure 5: Morphology of Golgi-DAG depleted cell at light microscopy level

HeLa cells were double transfected with different rapalogue dimerisation constructs, stained with the ER tracker, treated with rapalogue and followed through mitosis before fixation at cytokinesis. Yellow arrows indicate the recruitment of the lipid-modifying enzymes to different cellular compartments. (a) DGKεK was recruited to the Golgi in the presence of rapalogue (Golgi-DAG depleted). The NE reformed around the chromatin as indicated by the ER tracker (white arrows) (b) A catalytically inactive DGKεK (DGKεK.D434N) was recruited to the Golgi upon rapalogue addition (Golgi-DAG unperturbed). The NE reformed around the chromatin (white arrows). (c) DGKεK was recruited to the NE in the presence of rapalogue (NE-DAG depleted). There was no obvious NE reformed.
Figure 6

Tomogram pore clip

Orientation along dotted line

Rotate until circular

Rotate 90° and measure envelope and pore diameter

C

Pore diameter (nm)

Untransfected
DKG8K - Rap
DKG8K + Rap
D434N + Rap

Untransfected
DKG8K - Rap
DKG8K + Rap
D434N + Rap
Figure 6: NE rim curvature is affected by the lack of DAG at the Golgi.

(a) The analysis sequence from a nuclear pore 3D image clip (see Supplementary Figure 2 for whole tomographic reconstruction). The image clip is rotated until the nuclear pore is parallel to the x-axis, i.e. along the red dotted line between the nucleus (Nuc) and cytoplasm (Cyt). ER is the endoplasmic reticulum. The image clip is then rotated around the x-axis until the most circular pore can be observed, followed by a 90° rotation back to achieve a pore perpendicular to the field of view. Rim diameters (i and ii) and pore diameter (iii) are measured and presented as in plot (b) and (c) respectively. (b) The rim diameters for control HeLa cells (Untransfected, 2 pairs of cells), DKGεK-transfected but without rapalogue (DKGεK-Rap, 1 pair of cells), DKGεK-transfected with rapalogue (DKGεK+Rap, 2 Pairs of Cells) and D434N transfected with rapalogue (D434N+Rap, 2 pairs of cells). The red lines indicate the median ± interquartile range. The analysis was Kruskal-Wallis test with Dunn’s multiple comparison test. (c) Pore diameters for the same pores analysed in (b). The red lines indicate the mean ± standard deviation. The analysis was one-way ANOVA with Tukey’s multiple comparisons test. For (b) and (c) * indicates significance (* P<0.05, *** P<0.001, **** P<0.0001), all other combinations were not significant.