CC2D1B Coordinates ESCRT-III Activity during the Mitotic Reformation of the Nuclear Envelope

Graphical Abstract

Highlights

- CC2D1B localizes at the intersection between NE and microtubules at the time of nuclear resealing
- CC2D1B inhibits the polymerization of ESCRT-III in vitro
- CC2D1B licenses ESCRT-III filaments once ER is fully deposited on chromatin disks
- Reduced severing of spindle microtubules during NE resealing in CC2D1B-depleted cells

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In Brief

A faithful reformation of the nuclear envelope after mitosis is critical to ensure the proper compartmentalization of the genetic material. Ventimiglia et al. identified CC2D1B as a key regulator of this process to ensure properly timed ESCRT-III machinery recruitment and licensing of ESCRT-III filaments for nuclear envelope resealing.
CC2D1B Coordinates ESCRT-III Activity during the Mitotic Reformation of the Nuclear Envelope

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SUMMARY

The coordinated reformation of the nuclear envelope (NE) after mitosis re-establishes the structural integrity and the functionality of the nuclear compartment. The endosomal sorting complex required for transport (ESCRT) machinery, a membrane remodeling pathway that is highly conserved in eukaryotes, has been recently involved in NE resealing by mediating the annular fusion of the nuclear membrane (NM). We show here that CC2D1B, a regulator of ESCRT polymerization, is required to re-establish the nuclear compartmentalization by coordinating endoplasmic reticulum (ER) membrane deposition around chromatin disks with ESCRT-III recruitment to the reforming NE. Accordingly, CC2D1B determines the spatiotemporal distribution of the CHMP7-ESCRT-III axis during NE reformation. Crucially, in CC2D1B-depleted cells, ESCRT activity is uncoupled from Spastin-mediated severing of spindle microtubules, resulting in persisting microtubules that compromise nuclear morphology. Therefore, we reveal CC2D1B as an essential regulatory factor that licenses the formation of ESCRT-III polymers to ensure the orderly reformation of the NE.

INTRODUCTION

Metazoan cells disassemble their nuclear envelope (NE) during prophase to undergo an open mitosis (Champion et al., 2017). The reassembly of the NE at anaphase is a highly regulated process that begins when endoplasmic reticulum (ER) membranes are delivered to the rims of naked chromatin disks (Anderson and Hetzer, 2007; Lu et al., 2011). The final steps of NE reformation comprise the severing of the spindle microtubules that persist in the newly formed nuclear compartment and the sealing of the resulting gaps in the nuclear membrane (NM) mediated by the endosomal sorting complex required for transport (ESCRT) machinery (Olmos and Carlton, 2016; Scourfield and Martin-Serrano, 2017; Vietri et al., 2016). The ESCRT-associated protein CHMP7 works as a site-specific adapter that interacts with ER membranes to reach the fenestrations left by the expanding NM around the spindle microtubules (Gu et al., 2017; Olmos et al., 2016; Vietri et al., 2015). The C-terminal region of CHMP7 is thought to trigger the polymerization of the ESCRT-III components CHMP4B and CHMP2A to seal the NE. These growing ESCRT-III filaments then recruit the AAA ATPase Spastin, a microtubule severing enzyme, thus ensuring tight coordination between spindle microtubule removal and membrane remodeling. Lastly, a second ESCRT-associated AAA ATPase, VPS4, mediates the disassembly of ESCRT-III filaments and promotes the fusion of the NM, resulting in the final sealing of the NE and the re-establishment of nuclear integrity and functionality (Olmos et al., 2015, 2016; Vietri et al., 2015). A wider role for ESCRT proteins in the maintenance of nuclear integrity has been recently established, as this pathway is required to repair ruptures in the NE arising from cell migration through narrow spaces (Denais et al., 2016; Raab et al., 2016).

Despite advances in our understanding of the main factors involved in the mitotic reformation of the NE, the mechanisms determining its spatiotemporal coordination remain elusive (Schellhaus et al., 2016; Webster and Lusk, 2016). In this work, we have explored the function of coiled-coil and C2 domain-containing protein B (CC2D1B), a poorly characterized member of the Lgd/CC2D1 family of proteins that also includes the closely related human CC2D1A and its D. melanogaster ortholog Lgd. These proteins are characterized by the presence of four tandem DM14 domains that interact with the CHMP4 family of ESCRT-III proteins (Martinelli et al., 2012; Troost et al., 2012; Usami et al., 2012) and one C-terminal C2 domain that mediates their binding to membrane lipids (Drsusheimer et al., 2015; Gallagher and Knoblich, 2006). In D. melanogaster, Lgd regulates Notch signaling through the modulation of intracellular trafficking (Childress et al., 2006; Gallagher and Knoblich, 2006; Jäckel and Klein, 2006). Furthermore, the loss of Lgd function drives the over-proliferation of the imaginal discs, a phenotype that can be rescued by the expression of human CC2D1A or CC2D1B,
indicating a functional overlap between these proteins (Drusenheimer et al., 2015). This redundancy was also observed in mammalian models, as double CC2D1A/CC2D1B knockout mice died early during embryonic development (Zamarbide et al., 2018), but the deletion of CC2D1A or CC2D1B genes alone does not induce developmental defects (Drusenheimer et al., 2015; Zamarbide et al., 2018). However, non-redundant roles of these proteins are indicated by the perinatal death of CC2D1A-deficient mice (Al-Tawashi et al., 2012; Drusenheimer et al., 2015; Zamarbide et al., 2018).

It has been described that CC2D1A and Lgd inhibit the in vitro polymerization of ESCRT-III by binding to the multimerization domain of CHMP4B (Martinelli et al., 2012; McMillan et al., 2017), thus suggesting a key role for the Lgd/CC2D1 family of proteins in the regulation of ESCRT activities. Here, we show that CC2D1B interacts with CHMP7 and, in dividing cells, is recruited to the reforming NE in a CHMP7-dependent manner. Furthermore, CC2D1B depletion results in the loss of coordination between ER deposition at chromatin disks and ESCRT-III recruitment to the sites of NE sealing. Critically, the premature recruitment of ESCRT-III to the reforming NE is paralleled with the aberrant recruitment of Spastin, which results in persisting spindle microtubules that compromise the morphology of the NE. Therefore, we reveal CC2D1B as a key regulatory factor that coordinates ESCRT-III recruitment and microtubule severing with ER deposition during the mitotic reformation of the NE. This work exposes the unexpected critical role of ESCRT-III deregulation in cellular processes involving membrane remodeling.

RESULTS

CC2D1B Binds CHMP7 and Is Required for NE Resealing

The closely related CC2D1A/B proteins contain a predicted membrane-binding C2 domain and four DM14 domains that autonomously bind the core region of the CHMP4 family of ESCRT-III proteins (Martinelli et al., 2012; Usami et al., 2012) (Figure 1A). We hypothesized that CC2D1B may also interact with the C-terminal domain of CHMP7 that shows a significant similarity with CHMP4B (Hori et al., 2006). As shown in Figure 1B, GST-CC2D1B binds to HA-CHMP7, HA-CHMP4B, and HA-CHMP2A, whereas no association with HA-CHMP3 was detected. Importantly, the interaction with CC2D1B was retained by the C-terminal region of CHMP7 corresponding to the ESCRT-III-like domain but not by the CHMP7 N-terminal ER-binding domain (Figures 1A and 1C). Furthermore, we were able to detect a triple interaction between CC2D1B, CHMP4B, and CHMP7 in pull-down experiments using cells co-expressing GST-CHMP4B, HA-CHMP7, and YFP-CC2D1B (Figure S1A). Further mapping experiments showed that an HA-fused CC2D1B fragment containing only the DM14 domains (HA-CC2D1B residues 162-594) interacts with GST-CHMP4B and GST-CHMP7 (Figures S1B–S1D), whereas the deletion of the fourth DM14 domain (HA-CC2D1B residues 162-493) specifically abrogates the interaction with GST-CHMP7 (Figure S1D) without affecting the binding to GST-CHMP4B (Figure S1C). Accordingly, HA-CHMP7 was able to bind a fragment of CC2D1B containing only the fourth DM14 domain (GST-CC2D1B residues 528-594; Figure S1E). Altogether, these results indicate that CC2D1B binds to CHMP7 specifically through its DM14 4 domain, whereas the binding to CHMP4B maps to a more extended region that also includes DM14 1–3, thus providing a likely mechanism for the simultaneous binding of CC2D1B to CHMP4B and CHMP7.

The CC2D1B/CHMP7 interaction prompted us to analyze the potential function of CC2D1B in the mitotic sealing of the NE, observing that the silencing of CC2D1B delayed the recovery of nuclear GFP-NLS after cell division without altering the rate of nuclear expansion (Figures 1D–1F and S1F; Video S1). Furthermore, an increase in the proportion of cells with aberrant nuclear morphologies was also observed (Figures 1G and 1H). It has been described that these two phenotypes are associated with defects in NE reformation (Asencio et al., 2012; Olmos et al., 2015, 2016). Importantly, the effect of CC2D1B silencing on nuclear morphology was also observed in the immortalized human retina pigmented epithelial cell line RPE-1 (Figures S1G–S1I), as well as in primary human dermal fibroblast (HDF; Figures S1J–S1L) and primary mouse adult fibroblast (MAF; Figures S1M–S1O). Altogether, these results suggest that CC2D1B is required for the integrity and functionality of the nuclear compartment.

The CRISPR/Cas9 system was subsequently used to delete the CC2D1B locus in HCT116 cells. The resulting cells (HCT116<sup>CC2D1B</sup>) showed an increased proportion of nuclei with aberrant morphology compared to the parental cells (Figures 2A–2C). Importantly, this phenotype was rescued upon expression of a GFP-CC2D1B fusion (HCT116<sup>CC2D1B/GFP-CC2D1B</sup> cells), allowing us to discard off-target effects in HCT116<sup>CC2D1B</sup> cells. The full rescue activity of GFP-CC2D1B

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**Figure 1.** CC2D1B Is Required to Maintain NE Integrity and Functionality

(A) Schematic representation of CC2D1B and CHMP7.
(B) GST pull-down experiments of 293T cells transiently co-expressing GST-CC2D1B together with HA-tagged CHMP2A, CHMP3, CHMP4B, or CHMP7.
(C) GST pull-down experiments of 293T cells transiently co-expressing GST-CC2D1B together with HA-tagged CHMP7 full-length, CHMP7 Nt (residues 1–238), or CHMP7 Ct (residues 238–453).
(D–F) Time-lapse analysis of HCT116 cells stably co-expressing GFP-NLS and H2B-mCherry and transfected with control, CHMP7 or CC2D1B siRNAs. As the recovery of nuclear integrity after cell division was previously described to be dependent on CHMP7 expression (Olmos et al., 2016), CHMP7 silencing was used to validate our experiments.
(E) Representation of the nucleo-cytoplasmic GFP-NLS ratio over time. Mean ± SEM; siControl n = 29; siCC2D1B n = 35, p = 0.0003; siCHMP7 n = 22, p = 0.0402. Significance compared with the control was calculated at 90 min using a two-tailed unpaired t test.
(F) Representative frames corresponding to time-lapse images of siControl (top panels) or siCC2D1B (bottom panels) treated cells. Scale bars, 10 μm.

(G and H) Analysis of nuclear morphology in siRNA transfected HCT116 cells. (G) Percentage of cells showing an aberrant nuclear morphology. Mean ± SEM; siControl n = 2571; siCC2D1B n = 2040, p = 0.0102; siCHMP7 n = 1887, p = 0.0024. Significance of NE invaginations compared to the control was calculated using a two-tailed unpaired t test.
(H) Representative images corresponding to the quantifications shown in (G). Scale bars, 10 μm.

See also Figure S1 and Video S1.
Figure 2. CC2D1B Is Recruited to the Reforming NE in a CHMP7-Dependent Way

(A–C) Characterization of HCT116<sup>ΔCC2D1B</sup> cells. (A) Total cell lysates corresponding to HCT116 Control, HCT116<sup>ΔCC2D1B</sup>, HCT116<sup>ΔCC2D1B</sup>/GFP and HCT116<sup>ΔCC2D1B</sup>/GFP-CC2D1B were analyzed by WB using an anti-CC2D1B antibody. (B) Analysis of nuclear morphology in HCT116<sup>ΔCC2D1B</sup> cells. Mean ± SEM; (legend continued on next page)
also highlighted that expression of this fusion in HCT116\textsubscript{CC2D1B} cells provides a relevant system to determine the localization of CC2D1B in the context of mitotic NE reformation. Then, we analyzed HCT116\textsubscript{CC2D1B}/GFP-CC2D1B cells by time-lapse spinning disk confocal microscopy, observing that GFP-CC2D1B transiently accumulated in the perinuclear region during telophase (Figure 2D; Video S2). Remarkably, this enrichment was abrogated upon depletion of CHMP7 (Figures 2E and 2F; Video S2) in agreement with the CHMP7/CC2D1B interaction described above. The localization of GFP-CC2D1B on the reforming NE was further confirmed in cells co-expressing the NE marker lamina-associated polypeptide 2\textsubscript{l} fused to mCherry (mCherry-LAP2\textsubscript{b}) (Figure 2G; Video S3). The co-localization of GFP-CC2D1B with mCherry-tubulin was then explored, observing that CC2D1B decorates spindle microtubules, frequently marking the site of subsequent microtubule severing (Figure 2H; Video S4). As the intersection between the NE and spindle microtubules defines the membrane gaps sealed by ESCRT-III (Vietri et al., 2015), our results suggest that CC2D1B may be part of the ESCRT-associated machinery recruited by CHMP7 to seal the NE during mitosis.

**The Role of CC2D1B on NE Reformation Depends on Its Ability to Bind Membrane Lipids**

We then explored functional determinants shared by the Lgd/CC2D1 family, thereby performing structural studies of the highly conserved C-terminal region of the Drosophila ortholog Lgd (Figure S2A). The structure revealed an \(\alpha\)-helical N-terminal domain that contains a \(\sim\)50 Å long helical hairpin, followed by a long coil structure arranged anti-parallel to the hairpin which connects to the C2 domain at amino acid 664 (Figures 3B and 3C). The C2 fold ends at amino acid 796 and the remaining C-terminal region (residues 797–815) intertwines with the N-terminal helical domain, thereby stabilizing it (Figures 3B and S2B). The C2 domain of Lgd forms a classical \(\beta\)-sandwich structure with two four-stranded \(\beta\) sheets that classifies as a class II C2 domain with N and C termini oriented at the bottom of the sandwich and opposite to the top loops (Figure 3C), similar to the topology of the phospholipase C-\(\beta\)1 C2 domain (Essen et al., 1996). The Lgd C2 domain is atypical as two short \(\alpha\) helices are inserted into the loops connecting \(\beta\) strands 5 with 6 (\(\alpha3\)) and \(\beta\)-7 with 8 (\(\alpha2\)). These \(\alpha\) helices are conserved in human CC2D1A/B orthologs (Figure S2A), suggesting that these regions are functionally important. The convex part of this C2 domain is rigidly solvent exposed. Intriguingly, the Lgd helical hairpin is structurally similar to the Lgd DM14 3 domain with an root-mean-square deviation (RMSD) of 2.6 Å (Figure S2C), although its orientation toward the C2 domain would sterically prevent CHMP4B/shrub binding (McMillan et al., 2017). The hairpin shows further significant structural similarity with ESCRT-III CHMP3, CHMP4, and Ist1 (Bajorek et al., 2009; Martinelli et al., 2012; Muzio et al., 2006; Xiao et al., 2009) with RMSD’s of 3.0–3.1 Å on \(\sim\)70 equivalent Cx atoms, perhaps suggesting further associations with the ESCRT machinery.

The C2 fold described above suggested a role of the Lgd C terminus in lipid binding, a notion consistent with the previously described interaction of Lgd with phosphoinositides (Gallagher and Knoblich, 2006). Therefore, we performed a stringent pull-down assay to determine the binding of Lgd (residues 358–816) to a phosphoinositide panel, observing a weak but significant interaction with phosphatidylinositol 4,5-biphosphate (PIns(4,5)\(\beta\)) (Figure S2D). This result is supported by our structural data as the C2 domain of Lgd shows strong similarities with the C2 domains of protein kinase C\(\alpha\) (PKC\(\alpha\)) and synaptotagmin (Figure S2E) that bind to PIns(4,5)\(\beta\) through a basic cluster that makes direct contacts with the phosphates of the inositide ring (Guerrero-Valero et al., 2009; Guille `n et al., 2013). Interestingly, this cluster is conserved in Lgd (residues R689, K701, K703) as well as in eukaryotic orthologs including human CC2D1B (Figures S2A and S2E), which is consistent with the ability of CC2D1A and CC2D1B to associate with membrane pellets in cell fractioning experiments (Drusenheimer et al., 2015).

These structural and biochemical studies provided the basis to assess the functional contribution of the C2 domain of CC2D1B to the mitotic sealing of the NE. Therefore, a triple mutation (R721A, K734A, K736A) disrupting the putative phosphoinositide binding site in CC2D1B (Figure 3D) was engineered, and the resulting mutant (CC2D1B-C2M) was fused to GFP and stably expressed in HCT116\textsubscript{CC2D1B} cells. Importantly, GFP-CC2D1B Wt and GFP-CC2D1B-C2M were expressed at comparable levels in a cellular background that does not express endogenous CC2D1B (Figure 3E; lanes 3 and 4). The time-lapse analysis of these cells indicated that GFP-CC2D1B-C2M recruitment to the reforming NE was abrogated (Figure 3F, bottom panels, and Video S5). Furthermore, we detected PIns(4,5)\(\beta\) foci colocalizing with CHMP4B-L-GFP on the NE of dividing cells, supporting a lipid-mediated recruitment of CC2D1B to the rescaling NE (Figures 3G and S2F). Remarkably, the mislocalization of GFP-CC2D1B-C2M correlated with its inability to rescue the rate of mCherry-NLS nuclear recovery after mitosis (Figure 3H), whereas GFP-CC2D1B Wt showed full rescue activity in this assay. Accordingly, GFP-CC2D1B-C2M failed to lower the high proportion of HCT116\textsubscript{CC2D1B} cells with aberrant nuclear morphology (Figures 3I and S2G). These results indicate that the recruitment of CC2D1B to the reforming NE depends on its ability to bind membrane lipids and suggest that this recruitment is essential to re-establish the nuclear compartmentalization after cell division.
CC2D1B Determines the Timely Recruitment of ESCRT Proteins to the Resealing NE

The localization of CC2D1B on the reforming NE prompted us to investigate whether this protein may be involved in the coordination of ESCRT activity during nuclear reformation. To analyze the dynamics of ESCRT-III subunits during this process, we fused CHMP4B or CHMP2A to a 25-nm flexible linker (Kim et al., 2016) followed by GFP. The resulting CHMP4B-L-GFP and CHMP2A-L-GFP constructs were expressed at sub-physiological levels in HCT116 cells, and the initial characterization indicated that these fusions showed the expected localization and activity as they localized at the midbody (Figure S3A); their expression did not delay midbody abscission time (Figure S3B); they were recruited to the reforming NE in a CHMP7-dependent manner (Figures S3C–S3F; Videos S6 and S7), and they relocalized from a diffuse cytoplasmic pattern to the surface of aberrant endosomes upon depletion of VPS4A/B (Figure S3G). Moreover, pull-down analysis showed the specific association of endogenous CC2D1B with CHMP4B-L-GFP although not with CHMP2A-L-GFP (Figure S3H). Thus, cells stably expressing CHMP4B-L-GFP or CHMP2A-L-GFP are suitable models to analyze the role of CC2D1B in the dynamic localization of ESCRT-III during NE reformation.

We next developed an imaging-based semi-quantitative approach to determine the dynamics of ESCRT recruitment to the reforming NE (Figures 4A–4D; Videos S8, S9, and S10). Notably, the recruitment of ESCRT-III subunits was tightly coordinated in control cells, as GFP-CHMP7 and CHMP4B-L-GFP signals peaked at 370 s after cleavage furrow ingression, while the CHMP2A-L-GFP signal peaked closely at 290 s (Figure 4D, top panel). Strikingly, CC2D1B depletion resulted in the premature and asynchronous recruitment of ESCRTs (Figure 4D, bottom panel, and S4A), as GFP-CHMP7 recruitment occurred more than 4 min earlier in CC2D1B-depleted cells, and the accumulation of GFP-CHMP7, CHMP4B-L-GFP, and CHMP2A-L-GFP at the NE overlapped with the beginning of cleavage furrow ingression. Furthermore, the coordination between CC2D1B and CHMP2A was lost in CC2D1B-depleted cells, as evidenced by the 2-min difference between the peak recruitment of GFP-CHMP7 (100 s) and CHMP4B-L-GFP (220 s). To validate the use of the furrow ingression as a suitable time marker for the quantification of ESCRT-III recruitment, we analyzed the effect of CC2D1B depletion on the time gap between anaphase onset and the beginning of furrow ingression, observing that the silencing of CC2D1B increases this gap by 1 min when compared to control cells (Figures S4B–S4D; Video S11). However, although this result suggests a modest effect of CC2D1B depletion on mitosis timing, this difference is not sufficient to completely explain the early recruitment of ESCRT-III in CC2D1B-silenced cells. The premature recruitment of ESCRT subunits to the NE was also confirmed in cells stained for endogenous CHMP2A (Figure 4E). As the distance between both chromosomes sets increases during anaphase, the midzone length was used as a surrogate marker for the timing of CHMP2A recruitment. We observed that cells with CHMP2A at chromatin disks showed a shorter midzone length upon depletion of CC2D1B (Figure 4F), thus supporting the notion that CHMP2A recruitment to the reforming NE occurs earlier in these cells.

We next quantified the area under the fluorescence recruitment curves of GFP-CHMP7, CHMP4B-L-GFP, and CHMP2A-L-GFP as an indication of the total amount of ESCRT proteins recruited to the resealing NE. We observed that the silencing of CC2D1B had no effect on the total amount of GFP-CHMP7, CHMP4B-L-GFP, or CHMP2A-L-GFP fluorescence recruited (Figure 4E). As quantification of total fluorescence recruitment does not allow the discrimination between parameters as the number of ESCRT foci recruited and their persistence, we analyzed these parameters individually. We observed that the silencing of CC2D1B does not affect the number of GFP-CHMP7, CHMP4B-L-GFP, or CHMP2A-L-GFP foci recruited to the resealing NE (Figure 4F). Finally, we observed that, although the depletion of CC2D1B does not significantly alter the persistence time of GFP-CHMP7 and CHMP4B-L-GFP fluorescence on the NE, it increases the persistence time of CHMP2A-L-GFP (Figure 4G), perhaps suggesting the abortive formation of CHMP2A-containing polymers. Altogether, these
results indicate that the silencing of CC2D1B not only affects the timing of ESCRT-III recruitment to the resealing NE but could also have consequences on the quality of ESCRT-III polymers, suggesting that this machinery could be unable to fulfill its function in CC2D1B-depleted cells.

We then investigated whether CC2D1B regulates ESCRT-III polymerization using an in vitro assay previously described (Martellini et al., 2012). Briefly, a C-terminal truncation of CHMP4B fused to MBP (MBP-CHMP4B<sub>AC-AlkB</sub>) produces monomers (top fractions) and CHMP4B filamentous polymers (bottom fraction) (Pires et al., 2009) when analyzed by sucrose gradient centrifugation. Upon cleavage of monomeric MBP-CHMP4B<sub>AC-AlkB</sub> by tobacco etch virus (TEV) protease, CHMP4B becomes predominantly polymeric, whereas MBP stays in the upper monomeric fractions. Importantly, cleavage of monomeric MBP-CHMP4B<sub>AC-AlkB</sub> in the presence of a fragment of CC2D1B (residues 317–558) containing the third DM14 domain prevented the accumulation of CHMP4B<sub>AC-AlkB</sub> in the polymeric form (Figure 4G). These data suggest that CC2D1B inhibits CHMP4B polymerization in vitro, supporting a model whereby ESCRT-III polymerization is directly regulated by CC2D1B during the resealing of the NE.

To gain more mechanistic insights into the role of CC2D1B on the recruitment of ESCRT-III to the reforming NE, we compared the timing of GFP-CC2D1B and CHMP4B-L-GFP recruitment. We observed that, although both proteins are recruited roughly at the same time, CC2D1B disappeared from the reforming NE approximately 5 min before CHMP4B (Figure 4H), suggesting that CC2D1B could be specifically involved in the regulation of the early stages of ESCRT polymerization during NE reformation. We also asked whether the ability of CC2D1B to bind membrane phospholipids could play a role in the timing of ESCRT-III recruitment. Then, we analyzed the recruitment of CHMP2A-L-mCherry in HCT116 cells co-expressing GFP-CC2D1B Wt (Figure S4H; Video S12) or C2M (Figure S4I; Video S12), observing that the timing of CHMP2A-L-mCherry recruitment is not significantly different in these cell lines (Figure 4I). Thus, the timely recruitment of ESCRT-III to the NE is independent of the ability of CC2D1B to bind membrane phospholipids.

CC2D1B Coordinates NM Deposition with ESCRT-III Recruitment to the NE

To exclude general perturbations during early stages of NE reformation as consequence of CC2D1B silencing, we analyzed ER deposition around chromatin in HeLa cells co-expressing H2B-mCherry and the ER membrane-associated protein Sec61β fused to YFP (Figures 5A and S5A; Video S13). Although CC2D1B depletion resulted in the premature recruitment of CHMP4B-L-GFP (Figures 5B, 5C, bottom panel, and S5B; Video S13), in line with our results using HCT116 cells, no effect on the dynamics of NM deposition was observed (Figure 5A, 5C, top panel, and S5A). Notably, a tight coordination between NE reformation and ESCRT-III recruitment was evident in control cells, as CHMP4B-L-GFP peak recruitment (Figure 5C, bottom panel, black curve) occurred once ER deposition reached a plateau (Figure 5C, top panel, black line), which indicated that NE reformation is completed. In contrast, the peak recruitment of CHMP4B-L-GFP (Figure 5C, bottom panel, red curve) overlapped with the NE deposition phase (Figure 5C, top panel, red line) in CC2D1B-depleted cells, supporting the hypothesis of a premature recruitment of ESCRT-III in cells lacking CC2D1B expression.

To confirm these observations, we analyzed the dynamics of NE deposition and ESCRT-III recruitment by super resolution time-lapse microscopy using HCT116 cells stably co-expressing CHMP4B-L-GFP and the NE marker Emerin fused to mCherry (Figures 5D, 5E, and S5C–S5E; Video S13), observing that in control cells the recruitment of CHMP4B to the resealing NE began when mCherry-Emerin covered a high percentage of the nuclear perimeter (median value = 89.61%). In contrast, CC2D1B-depleted cells showed CHMP4B puncta decorating the NE in earlier stages of nuclear reformation, when the resealing NE covered a lower percentage of nuclear perimeter (median value = 72.6%). We observed a similar result in HCT116 cells expressing GFP-CHMP2A-L-mCherry in the perinuclear area.
stably co-expressing CHMP4B-L-GFP and mCherry-Sec61β (Figure S5F; Video S13). In control cells, CHMP4B was recruited to the resealing NE once this structure was completely formed around the nascent nucleus. On the other hand, in cells silenced for CC2D1B expression, CHMP4B was recruited in earlier stages of ER deposition when the reforming NM was not yet fully
enclosing the nuclear compartment, thus supporting our quantitative data. Altogether, these results confirm that CC2D1B ensures that ESCRT-III is recruited to the reforming NE at the right time, once chromatin is fully enclosed by the NE.

**CC2D1B Silencing Impairs the Severing of Spindle Microtubules**

NE resealing requires the severing of microtubules at the intersection between the mitotic spindle and the NE by the ESCRT-recruited ATPase Spastin (Vietri et al., 2015). The role of CC2D1B on ESCRT-III recruitment to the reforming NE prompted us to investigate whether this protein could also be involved in Spastin recruitment. We first observed that CC2D1B silencing induced the premature recruitment of GFP-Spastin M87 to the NE (Figures 6A, middle panels, and S6C, bottom panels; Video S14), and, as expected, this recruitment was ablated in CHMP7-depleted cells (Figure 6A, right panels, and S6C, bottom panels; Video S14). Furthermore, CC2D1B silencing induced a strong reduction in the total amount of GFP-Spastin fluorescence recruited to the NE (Figure 6D). These results suggest that the asynchronous recruitment of ESCRT-III subunits to the reforming NE in CC2D1B-depleted cells impairs the downstream recruitment of Spastin, and, consequently, the ability of these cells to sever the spindle microtubules may be compromised.

We next studied the kinetics of microtubule disassembly at the nascent nuclear compartment using HCT116 cells co-expressing mCherry-tubulin and CHMP2A-L-GFP. In agreement with previous reports (Vietri et al., 2015), CHMP2A accumulation preceded microtubule disassembly in control cells (Figure 6E, top panels; Video S15). Conversely, in Spastin-depleted cells, dense microtubule bundles connecting the midbody with the spindle pole persisted even at late stages of cell division, when CHMP2A-L-GFP has already disappeared from the perinuclear region (Figure 6E, middle panels, and S6B, left panels; Video S15). These observations were further supported by a semi-quantitative approach to estimate the disassembly rate of nuclear microtubules by quantifying mCherry-tubulin fluorescence in the nucleus over time. The silencing of Spastin delayed spindle microtubule severing (Figure 6F), although it had a negligible effect on the recruitment dynamics of CHMP2A-L-GFP (Figure S6A). Interestingly, the dynamics of microtubules disassembly in control and Spastin-silenced cells diverge at the time that coincides with the peak recruitment of CHMP2A-L-GFP (black curve in Figure 6F), supporting the notion that ESCRT-III recruitment triggers the severing of spindle microtubules by Spastin. Critically, the silencing of CC2D1B recapitulated the phenotypes observed in Spastin-depleted cells, as shown by the persistence of spindle microtubules at late stages of cell division (Figure 6E, bottom panels, and S6B, right panels; Video S15) and the delayed rate of spindle microtubule disassembly (Figure 6H). Importantly, these effects paralleled with the premature recruitment of CHMP2A-L-GFP to the reforming NE (Figure S6A). Of note, the above effects on microtubule disassembly were specific for the nuclear compartment as depletion of either Spastin or CC2D1B had no effect on the cytoplasmic signal of mCherry-tubulin (Figures 6G and 6I). Taken together, our results suggest that CC2D1B depletion mimics the effect of Spastin silencing on the severing of nuclear spindle microtubules, impairing the ability of the cells to accurately disassemble the mitotic spindle and proceed with the proper sealing of the NE.

Lastly, we reasoned that the persistence of spindle microtubules beyond cell division could give rise to cells bearing microtubules spanning the nuclear compartment during interphase, therefore providing a plausible model to explain the nuclear aberrations observed in CC2D1B-depleted cells. Supporting this notion, we detected transnuclear microtubules in HCT116 cells stably expressing mCherry-tubulin (Figure 7A; Video S16), observing that their frequency dramatically increased from 6% in the control cells to 17.39% in CC2D1B-depleted cells and 15.54% in Spastin-silenced cells (Figure 7B). 3D reconstruction of these cells revealed that tunnels containing microtubules pierce the nucleus, exerting a deforming force on its poles and inducing deep invaginations on its surface (Figures 7C and 7D; Video S16), therefore preventing the nucleus from reaching a typical round shape (Figure S7A). Taken together, our data suggest that the uncoordinated recruitment of ESCRT-III and Spastin to the reforming NE in CC2D1B-depleted cells has not only a local effect on the resealing of the NE but also more profound implications permanently modifying the morphology of the nucleus and therefore compromising its integrity and functionality.

**DISCUSSION**

We show here that CC2D1B organizes the timely recruitment of ESCRT-III and associated proteins to the sealing gaps of the nuclear compartment. This local effect on the resealing of the NE but also more profound implications permanently modifying the morphology of the nucleus and therefore compromising its integrity and functionality.
A

Transnuclear microtubules

mCherry
Tubulin

Merge
Hoechst

B

Transnuclear microtubules (% of cells)

C

Front view

D

Top view

E

1) Control

2)

3)

1) ΔCC2D1B

2)

3)

CC2D1B CHMP4B CHMP7 Tubulin Spastin

(legend on next page)
reforming NE. We demonstrate that CC2D1B silencing results in the premature and unregulated polymerization of ESCRT-III components at the NM during mitosis. As result of this dysregulation, the ability of Spastin to clear the nucleus from spindle microtubules is reduced, which provokes deleterious effects on nuclear morphology and integrity.

We propose a model (Figure 7E) whereby CC2D1B could play a double role in the regulation of CHMP4B activity. On one hand, the N-terminal DM14 domains of CC2D1B could mediate the binding and stabilization of the monomeric cytoplasmic form of CHMP4B, preventing then its premature or non-productive polymerization. On the other hand, the ability of CC2D1B to interact with membrane phospholipids through its C-terminal C2 domain could mediate the precise delivery of CHMP4B monomers to the gaps in the resealing NE, where they would be able to polymerize and form constriction competent ESCRT-III filaments. A role for CC2D1B as a scaffold protein able to position CHMP4B at the required location is supported by the solved structure of Lgd that shows a rigid connection between the C-terminal C2 domain and the helical domain that bridges to the N-terminal DM14 domains. Interestingly, an Lgd fragment consisting only of the third DM14 and the C2 domains is sufficient for Notch regulation (McMillan et al., 2017), corroborating that the C2 domain has an important role in CHMP4 positioning. Our model is also supported by previous published data indicating that members of the Lgd/CC2D1 family of proteins are able to interact with cell membranes in vivo, as they associate to the insoluble membrane fractions in cell fractioning experiments (Drusenheimer et al., 2015; Gallagher and Knoblich, 2006). Our PIP pull-down experiments suggest that Lgd/CC2D1 C2 domains specifically interact with PIns(4,5)P2-enriched membranes, a result supported by the presence of PIns(4,5)P2 foci on the reformating NE, and by the structural similarities with other C2 domains known to interact with PIns(4,5)P2 (Guerrero-Valero et al., 2009; Guillén et al., 2013). The modeling of CC2D1B C2 domain binding to PIns(1,4,5)P3 suggests a further level of regulation in the activity of Lgd/CC2D1 proteins via lipid modification (Dumas et al., 2010). It has also been described that Lgd binds preferentially to PIns(3)P, PIns(4)P and PIns(5)P in PIP strip experiments, although a weaker interaction with PIns(3,4)P2, PIns(3,5)P2, and PIns(4,5)P2 was also detected (Gallagher and Knoblich, 2006). The apparent discrepancy with Gallagher et al. could be explained by the different methods used to analyze the weak interaction of Lgd with phospholipids.

Our data also suggest a regulatory interaction between CC2D1B and CHMP7. It has been shown that the third DM14 domain of CC2D1A and Lgd interacts preferentially with CHMP4B and shrub, respectively (Martinelli et al., 2012; McMillan et al., 2017), suggesting that other DM14 domains might be simultaneously available for CHMP7 interaction. Accordingly, our results indicate that the interaction of CC2D1B with CHMP4B and CHMP7 could be mediated by different DM14 domains. Thus, we hypothesize that the formation of a transient CHMP7-CC2D1B-CHMP4B complex at the gaps of the NE could be necessary for a fully productive polymerization of the ESCRT machinery. In this context, the cytoplasmic sequestration of CHMP4B by CC2D1B could keep CHMP7 away from the gaps in the NE until the membrane fenestrations reach a threshold size suitable for ESCRT polymerization. Conversely, in CC2D1B-depleted cells, CHMP7 and CHMP4B would be free to unrestrainedly interact with each other, triggering the premature and unregulated polymerization of ESCRT-III on the forming NE. A role for CC2D1B in the regulation of the cytoplasmic activity of CHMP4B is supported by our data indicating that although the C2M mutant form of CC2D1B is not recruited to the resealing NE, its expression has no effect on the recruitment time of ESCRT-III.

The recruitment of Spastin to the growing ESCRT-III filaments depends on its ability to bind Ist1 and CHMP1B (Agromayor et al., 2009; Reid et al., 2005; Yang et al., 2008). Thus, we speculate that aberrant ESCRT-III polymers unable to efficiently bind Spastin may be formed at the NE in the absence of CC2D1B. Alternatively, the bundles of spindle microtubules may be immature for severing at the time Spastin is recruited in CC2D1B-depleted cells. Accordingly, it has been proposed that the spatial and temporal regulation of microtubule dynamics during nuclear reformation is key to ensure the normal functionality of the nucleus (Xue et al., 2013). We show here that as consequence of the defective severing of microtubules during the reassembly of the nuclear compartment, bundles of microtubules persist through interphase, forming deep nuclear invaginations that entirely transsect the nucleus deforming its surface. As the

Figure 7. Persistent Transnuclear Microtubules Originate Deep Nuclear Invaginations in Interphase
(A) z stack series corresponding to an HCT116 cell stably co-expressing mCherry-tubulin with CHMP2A-L-GFP, transfected with CC2D1B siRNA, and incubated with Hoechst 33258 30 min before imaging to stain DNA. The signal corresponding to Hoechst 33258 is shown in green. The signal corresponding to CHMP2A-L-GFP is not shown. This cell was selected as an example of a nucleus showing transnuclear microtubule tunnels. Scale bar 10 µm.
(B) Frequency of transnuclear microtubule tunnels in HCT116 cells stably co-expressing mCherry-tubulin with CHMP2A-L-GFP and transfected with control, Spastin, or CC2D1B siRNAs. Mean ± SEM; sControl n = 1,083; siCC2D1B n = 850, p = 0.0145; siSpastin n = 1,075, p = 0.0007. Significance compared to the control was calculated using a two-tailed unpaired t test.
(C and D) 3D rendering of the cell shown in (A). (C) Front view corresponding to Hoechst 33258 channel. (D) Top view corresponding to Hoechst 33258 and mCherry-tubulin. 80% transparency was applied to Hoechst 33258 channel to facilitate the observation of transnuclear microtubule channels. Scale bars, 5 µm.
(E) Proposed model. Left panel: (1) CC2D1B binds to the monomeric cytoplasmic form of CHMP4B, hindering its association with CHMP7 and therefore preventing its premature recruitment to the NE. (2) By means of its ability to interact with CHMP7 and CHMP4B through its N-terminal DM14 domains and with membrane phospholipids through its C-terminal C2 domain, CC2D1B could function as a scaffold to position CHMP4B monomers close to the NM. Then, CC2D1B could mediate the organization of a transient CHMP7-CC2D1B-CHMP4B complex at the sealing gaps of the NE, facilitating the delivery of CHMP4B monomers to the growing ESCRT-III filament. (3) Spastin is recruited by the ESCRT-III to catalyze the severing of spindle microtubules, which facilitates the final sealing of the NE. Right panel: (1) in CC2D1B-silenced cells, CHMP4B monomers are free to associate to CHMP7 without restrictions, (2) triggering the premature polymerization of CHMP4B monomers on the NE. (3) These prematurely formed ESCRT filaments cannot be competent for Spastin recruitment and membrane constriction, which results in an impaired ability of the cells to sever spindle microtubules, leading to deleterious effects in nuclear integrity and functionality. See also Figure S7 and Video S16.
number of these transnuclear microtubule tunnels increases as consequence of the disruption of normal cell division, we speculate that the presence of these tunnels could have a deleterious effect on cell performance.

In summary, the evidence presented here reveals CC2D1B as a key player in the mitotic reformation of the NE, by licensing ESCRT-III polymerization after the ER-derived NM is fully deposited on chromatin disks. Our results also highlight the importance of ESCRT-associated proteins that negatively regulate ESCRT-III polymerization for the progress of cellular processes involving membrane remodeling and open the gate to the identification of similar mechanisms specifically implicated in other ESCRT-dependent processes besides the resealing of the NE.

STAR METHODS

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SUPPLEMENTAL INFORMATION

Supplemental Information includes 7 figures, 1 table, 1 data file, and 16 videos and can be found with this article online at https://doi.org/10.1016/j.devcel.2018.11.012.

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AUTHOR CONTRIBUTIONS

L.N.V. and J.M.-S. conceived and designed the project. W.W. designed and supervised the structural work. L.N.V., M.A.C.-G., A.C., and I.M.P. generated tools and performed functional experiments. N. Martinelli, P.M., and N. Miguet performed structural and biochemical experiments. Y.O. and J.G.C. performed immunofluorescence experiments and helped with the experimental design. L.N.V., M.A.C.-G., N. Martinelli, A.C., P.M., Y.O., J.G.C., W.W., and J.M.-S. analyzed data. L.N.V., J.M.-S., and W.W. wrote the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Rabbit polyclonal anti-CC2D1B (H-258) | Santa Cruz Biotechnology | Cat# sc-134634; RRID: AB_10611611 |
| Rabbit polyclonal anti-CHMP7 | Proteintech | Cat# 16424-1-AP; RRID: AB_2079500 |
| Mouse monoclonal anti-Spastin (Sp 3G11/1) | Santa Cruz Biotechnology | Cat# sc-53443; RRID: AB_785771 |
| Mouse monoclonal anti-GFP (clones 7.1 and 13.1) | Roche | Cat# 11814460001; RRID: AB_390913 |
| Rabbit polyclonal anti-HSP90α/β (H-114) | Santa Cruz Biotechnology | Cat# sc-7947; RRID: AB_2121235 |
| Mouse monoclonal anti-HSP90α/β (F-8) | Santa Cruz Biotechnology | Cat# sc-13119; RRID: AB_675659 |
| Mouse monoclonal anti-GST (B14) | Santa Cruz Biotechnology | Cat# sc-138; RRID: AB_627677 |
| Rabbit polyclonal anti-HA | Antibodies Online | Cat# ABIN100176; RRID: AB_10779560 |
| Rabbit polyclonal anti-Lamin B1 | Abcam | Cat# ab16048; RRID: AB_443298 |
| Chicken polyclonal anti-GFP | Abcam | Cat# ab13970; RRID: AB_300798 |
| Mouse monoclonal anti-alpha Tubulin | Abcam | Cat# ab7291; RRID: AB_2241126 |
| Rabbit polyclonal anti-CHMP2A | Proteintech | Cat# 10477-1-AP; RRID: AB_2079470 |
| Mouse monoclonal anti PtdIns(4,5)P$_2$ | Echelon | Cat# Z-P045; RRID: AB_427225 |
| Secondary donkey anti-rabbit Alexa Fluor 555 | ThermoFisher Scientific | Cat# A31572; RRID: AB_162543 |
| Secondary goat anti-chicken Alexa Fluor 488 | ThermoFisher Scientific | Cat# A11039; RRID: AB_142924 |
| Secondary goat anti-rabbit Alexa Fluor 488 | ThermoFisher Scientific | Cat# A11008; RRID: AB_143165 |
| Secondary donkey anti-mouse Alexa Fluor 594 | ThermoFisher Scientific | Cat# A21203; AB_141633 |
| Secondary goat anti-mouse (DyLight™ 680 Conjugate) | Cell Signaling Technology | Cat# 54705; RRID: AB_10696895 |
| Secondary goat anti-rabbit (DyLight™ 800 Conjugate) | Cell Signaling Technology | Cat# 5151; RRID: AB_10697505 |
| **Bacterial and Virus Strains** |        |            |
| BL21 | New England Biolabs | C2530H |
| **Chemicals, Peptides, and Recombinant Proteins** |        |            |
| cOmplete™ Protease Inhibitor Cocktail | Sigma-Aldrich | Cat#000000011697498001 |
| Lipofectamine RNAiMax | Life Technologies | Cat#13778075 |
| Polyethylenimine (PEI) | Polysciences | Cat# 23966-1 |
| DharmaFECT-1 | Dharmacon | Cat#T-2001 |
| Glutathione Sepharose 4B beads | GE Healthcare Life Sciences | Cat# 17075601 |
| GFP-Trap M | ChromoTek | Cat#GTM-20 |
| Prolong Diamond Antifade Mountant | ThermoFisher Scientific | Cat#P36965 |
| Hoechst 33258 | Sigma-Aldrich | Cat#861405 |
| PIP beads | Echelon | Cat#P-B00S |
| Ni-NTA Agarose | Qiagen | Cat#30210 |
| PEG 3500 Mme | Sigma | Cat#701750 |

| **Deposited Data** |        |            |
| Lgd (residues 550-816) structure | This paper | PDB: 6EI6 |

(Continued on next page)
## Continued

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Experimental Models: Cell Lines** | | |
| HCT116 | ATCC | Cat# CCL-247; RRID: CVCL_0291 |
| Hela | ATCC | Cat# CRM-CCL-2; RRID: CVCL_0030 |
| hTert-RPE-1 | ATCC | Cat# CRL-4000; RRID: CVCL_4388 |
| Human dermal fibroblasts (HDF) | Tanya Shaw laboratory (KCL) | N/A |
| Mouse adult fibroblasts (MAF) | Axel Behrens laboratory (Francis Crick Institute) | N/A |
| **Oligonucleotides** | | |
| CC2D1B siRNA 5’-GAUCAACUCUUCAAAAC UAAUU-3’ | Dharmacon | N/A |
| CHMP7 siRNA 5’-GGGAGAAGAUUGUGAAGUUUUU-3’ | Dharmacon | N/A |
| VPS4A siRNA 5’- CCGAGAAGCUGAAGGAUUA-3’ | Dharmacon | N/A |
| VPS4B siRNA 5’- CCAAAGAAGCUGAAGGAUUA-3’ | Dharmacon | N/A |
| Spastin siRNA | Qiagen | Cat# SIO2781219 |
| Mouse CC2D1B siRNA | Dharmacon | Cat# L-056827-00-0005 |
| **Recombinant DNA** | | |
| pNG72 GFP-CC2D1B Wt | This paper | N/A |
| pNG72 GFP-CC2D1B C2M | This paper | N/A |
| pNG72 GFP-CHMP3B | This paper | N/A |
| pNG72 CHMP4B-Linker-GFP | This paper | N/A |
| pNG72 CHMP2A-Linker-GFP | This paper | N/A |
| pNG72 GFP-NLS | This paper | N/A |
| pCMS28 mCherry-NLS | This paper | N/A |
| pCMS28 YFP-Sec61β | This paper | N/A |
| pCMS28 mCherry-Sec61β | This paper | N/A |
| pCMS28 mCherry-Lap2β (residues 244-454) | This paper | N/A |
| pCMS28 h2B-mCherry | This paper | N/A |
| pCMS28 GFP-CHMP7 | Olmos et al., 2016 | N/A |
| pCMS28 mCherry-Tubulin | Agromayor et al., 2009 | N/A |
| pCMS28 mCherry-Emerin | This paper | N/A |
| pCMS28 CHMP2A-Linker-mCherry | This paper | N/A |
| pCAG GST-CC2D1B | This paper | N/A |
| pCAG GST-CC2D1B (residues 528-594) | This paper | N/A |
| pCAG GST-CHMP4B | This paper | N/A |
| pCAG GST-CHMP7 | This paper | N/A |
| pCR3.1 HA-CHMP7 | This paper | N/A |
| pCR3.1 HA-CHMP4B | Martin-Serrano et al., 2003 | N/A |
| pCR3.1 HA-CHMP2A | Martin-Serrano et al., 2003 | N/A |
| pCR3.1 HA-CHMP7 Nt | This paper | N/A |
| pCR3.1 HA-CHMP7 Ct | This paper | N/A |
| pCR3.1 YFP-CC2D1B | This paper | N/A |
| pCR3.1 HA-CC2D1B (residues 162-493) | This paper | N/A |
| pCR3.1 HA-CC2D1B (residues 162-594) | This paper | N/A |
| pLGC2 | Vantourout et al., 2018 | N/A |
| pProEx-Htb Lgd (residues 550-816) | This paper | N/A |
| pProEx-Htb Lgd (residues 358-816) | This paper | N/A |
| pProEx-Htb CC2D1B (residues 317-558) | This paper | N/A |

(Continued on next page)
CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Juan Martin-Serrano (juan.martin_serrano@kcl.ac.uk).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Cell Lines
Male HCT116 cells, female Hela cells, 293T cells of indeterminate sex and MAF cells derived from female mice were cultured in Dulbecco’s Modified Eagle Medium (DMEM) containing 10% fetal calf serum (FCS) and gentamycin 20 &mu;g/ml. hTert-RPE-1 cells were cultured in Dulbecco’s Modified Eagle Medium (DMEM) containing 10% fetal calf serum (FCS), NEAA, hygromycin B 10 &mu;g/ml and gentamycin 20 &mu;g/ml. HDF were cultured in Dulbecco's Modified Eagle Medium (DMEM) containing 10% fetal calf serum (FCS) and penicillin-streptomycin (100 U/ml). Stable cell lines were generated using MLV-based retroviruses as described previously (Carlton and Martin-Serrano, 2007) and selected using puromycin (120 ng/ml for HCT116 or 200 ng/ml for Hela) or G418 (250 &mu;g/ml for HCT116 or 500 &mu;g/ml for Hela).

METHOD DETAILS

Retroviral Plasmids
Modified versions of pCMS28 (Carlton and Martin-Serrano, 2007) or pNG72 (Caballe et al., 2015) plasmids were used as retroviral packaging vectors. pCMS28 mCherry-Tubulin and pCMS28 GFP-CHMP7 constructs were previously described (Agromayor et al., 2009; Olmos et al., 2016). CC2D1B Wt, CC2D1B-C2M and Spastin M87 were cloned as N-terminal GFP fusions into pCMS28. CC2D1B Wt, CC2D1B-C2M and Spastin M87 were cloned as N-terminal GFP fusions into pCMS28. CHMP2A-Linker and CHMP4B-Linker were generated by gene synthesis (GeneArt, ThermoFisher) and cloned as C-terminal GFP fusions into pCMS28. The SV40 Large T antigen nuclear localization signal was cloned as N-terminal GFP fusions into pCMS28. Sec61b was cloned as N-terminal YFP or mCherry fusions into pCMS28. Lap2b residues 244-454 was cloned as a N-terminal mCherry fusion in pCMS28. Emerin was cloned as a N-terminal mCherry fusion in pCMS28. H2B-mCherry was cloned as a C-terminal mCherry fusion in pCMS28.

Transfections
For retroviral production or GST pulldowns, 293T cells were transfected using Polyethylenimine (PEI; Polysciences, Germany) and Hela cells were transfected using Dharmafect-1 (Dharmacon RNA Technologies) according to manufacturer's instructions. Custom siRNA against CC2D1B (5'-GAUCAACUCUUCAACUAAU-3'), CHMP7 (5'-GGGAGAAGAUUGUGAAGUUUUU-3'), siVP54A (5'-CCGAGAAGCAAGAAAGGAUUA-3') and siVP54B (5'-CCAAAGAAGCACUGAAAGA-3') were purchased from Dharmacon. siRNA against Spastin (Hs_SPAST_1, catalog no. SI02781219) was purchased from Qiagen. siRNA against mouse CC2D1B (ON-TARGETplus Mouse CC2D1B siRNA – SMARTpool, catalog no. L-056827-00-0005) was purchased from Dharmacon.

Generation of HCT116CC2D1B Cell Line
A pair of guide RNA targeting the adjacent regions to the CC2D1B locus were designed using the Zhang Lab website (http://crispr.mit.edu) and cloned into a pLG2C plasmid expressing GFP and Cas9 (kindly gifted by Dr. Pierre Vantourout, KCL). The sequences for the guide oligos were 5'- CAA CAG TGC TAC CCN TGG TA GTT TT-3' and 5'- AAG ACC TAC CAT GCT GGG TA GTT TT-3'. HCT116 cells were co-transfected with both guides using Lipofectamine 3000 (Invitrogen, Life Technologies). 48 hours later single cells expressing GFP were FACs sorted. HCT116 clones tested for CC2D1B expression by western blot.

GST Pulldowns
293T cells from 6-well plates were co-transfected with the corresponding pcAG-GST- and pcR3.1 HA-tagged constructs. 48 hours later, the cells were lysed at 4°C in 1 ml of lysis buffer containing 50 mM Tris, pH 7.4, 150 mM NaCl, 5 mM EDTA, 5% glycerol,
1% Triton X-100 and a protease inhibitor cocktail (complete mini-EDTA free, Sigma). The lysates were sonicated and centrifugated at 15,000 rpm for 10 minutes and the supernatant was incubated with glutathione-sepharose beads (Amersham Biosciences) for 3 hours at 4°C and washed three times with wash buffer containing 50 mM Tris, pH 7.4, 150 mM NaCl, 5 mM EDTA, 5% glycerol, and 0.1% Triton X-100. Bound proteins were eluted in 100 μl of Laemmli buffer, boiled and analyzed by western blot.

**GFP Immunoprecipitations**

HCT116 cell lines stably expressing CHMP4B-L-GFP or CHMP2A-L-GFP fusions were lysed at 4°C in 1 ml of lysis buffer as described above for GST pulldowns. Clarified lysates were incubated with anti-GFP coupled magnetic microparticles (GFP-Trap, ChromoTek) for 2 hours and the microparticles were washed four times with wash buffer. Bound proteins were eluted in 100 μl of Laemmli buffer, boiled and analyzed by western blot.

**Immunoblotting**

Samples were resolved by SDS-PAGE and proteins transferred to Nitrocellulose membranes. The corresponding primary and DyLight- or HRP-conjugated secondary antibodies were diluted in 1% milk. The membranes were scanned using a Li-Cor Odyssey infrared scanner and software system (Li-Cor Biosciences). The primary antibodies used were: rabbit CC2D1B (H-258, Santa Cruz Biotechnology), rabbit CHMP7 (16424-1-AP, Proteintech), mouse Spastin (Sp 3G11/1, Santa Cruz Biotechnology), mouse GFP clones 7.1 and 13.1 (11814460001, Roche), rabbit Hsp90 (H-114, Santa Cruz Biotechnology), mouse Hsp90 (F-8, Santa Cruz Biotechnology), mouse GST (B-14, Santa Cruz Biotechnology), rabbit HA (ABIN100176, Antibodies Online). The secondary antibodies used were: goat anti-mouse DyLight 680-conjugated (Cell Signaling), goat anti-rabbit DyLight 800-conjugated (Cell Signaling), goat anti-mouse HRP-conjugated (Cell Signaling), goat anti-rabbit HRP-conjugated (Cell Signaling). The raw western blots corresponding to the insets shown in the main and supplemental figures are shown in Data S1.

**Immunofluorescence**

Cells were grown on coverslips and fixed for 10 minutes with 4% PFA at 4°C or for 20 minutes with methanol at -20°C. Cells were blocked with 1% FCS in PBS, stained with the indicated primary antibodies, washed and stained with the corresponding Alexa-conjugated secondary antibodies (ThermoFisher Scientific). DNA was stained with Hoechst 33258. Coverslips were mounted in Prolong Diamond Antifade Mountant (ThermoFisher Scientific). The primary antibodies used were: rabbit Lamin B1 (ab16048, Abcam), chicken GFP (ab13970, Abcam), mouse tubulin DM1A (ab7291, Abcam), rabbit CHMP2A (10477-1-AP Proteintech), mouse PtdIns(4,5)P2 (Z-P045, Echelon). Fixed cells were imaged on a Nikon Ti-Eclipse wide-field inverted microscope equipped with a CoolSnap HQ2 CCD camera (Photometrics, Tucson, AZ).

**Spinning Disk Confocal Imaging**

**ESCRT-Related Proteins Recruitment to the NE**

Cells stably-expressing the corresponding GFP- or mCherry-tagged proteins were seeded on 24-wells glass bottom plates (Cellvis) and transfected with the corresponding siRNA. Cells were imaged 48 hours later using 100x oil-immersion objective equipped Nikon Eclipse Ti-E inverted microscope with attached environmental chamber. Images were acquired every 15 seconds for HCT116 cells or every 10 seconds for Hela cells using an Andor Neo sCMOS camera using 2x2 binning. The quantification of the time-lapse images was performed using Fiji (Schindelin et al., 2012). Briefly, GFP fluorescence around chromatin was selected by manually adjusting the threshold of the green channel. Then a binary mask was created and the total intensity (mean intensity x mask area) of GFP-tagged proteins was quantified for every time point. The curve corresponding to the recruitment of GFP-tagged proteins was represented as mean ± SEM and the beginning of the furrow ingress was set as t0. The area under the curves was quantified using Prism (GraphPad Software).

To compare the timing of GFP-CC2D1B versus CHMP4B-L-GFP recruitment, the first (T1) and the last (Tf) frames showing GFP accumulation in the perinuclear area were manually scored in every movie and the data were represented as 5-95 percentile boxes using Prism (GraphPad Software).

**Nuclear Membrane Deposition**

Hela cells stably co-expressing YFP-Sec61β and H2B-mCherry were seeded on 24-wells glass bottom plates (Cellvis), transfected with the corresponding siRNA and imaged in a Nikon Eclipse Ti-E inverted CSU-X1 spinning disk confocal microscope with attached environmental chamber. Images were acquired every 15 seconds for HCT116 cells or every 10 seconds for Hela cells using an Andor Neo sCMOS camera using 2x2 binning. The quantification of the time-lapse images was performed using Fiji (Schindelin et al., 2012). Briefly, GFP fluorescence around chromatin was selected by manually adjusting the threshold of the green channel. Then a binary mask was created and the total intensity (mean intensity x mask area) of GFP-tagged proteins was quantified for every time point. The curve corresponding to the recruitment of GFP-tagged proteins was represented as mean ± SEM and the beginning of the furrow ingress was set as t0. The area under the curves was quantified using Prism (GraphPad Software).

To compare the timing of GFP-CC2D1B versus CHMP4B-L-GFP recruitment, the first (T1) and the last (Tf) frames showing GFP accumulation in the perinuclear area were manually scored in every movie and the data were represented as 5-95 percentile boxes using Prism (GraphPad Software).

**Spindle Microtubules Depolymerisation**

HCT116 cells stably co-expressing mCherry-Tubulin and CHMP2A-L-GFP were seeded on 24-wells glass bottom plates (Cellvis), transfected with the corresponding siRNA and imaged in a Nikon Eclipse Ti-E inverted CSU-X1 spinning disk confocal microscope using the same conditions as described above. mCherry-Tubulin fluorescence was quantified in Fiji (Schindelin et al., 2012) by drawing a line through the nuclear spindle microtubules or the cytoplasm in every time-lapse image. The t0 was set at the beginning of furrow ingress and fluorescence values were normalized to t0 and represented as mean ± SEM.
Analysis of Transnuclear Microtubules Tunnels

Cells stably co-expressing mCherry-Tubulin and CHMP2A-L-GFP were seeded on 24-wells glass bottom plates (Cellvis) and transfected with the corresponding siRNA. Cells were incubated with Hoechst 33258 to stain the DNA 30 minutes before imaging. Z-stacks of images separated by 100 nm were acquired using a 100x oil-immersion objective equipped Nikon Eclipse Ti-E inverted CSU-X1 spinning disk confocal microscope with attached environmental chamber and connected to Andor Neo sCMOS camera without binning. The Z-stacks were used to quantify the percentage of cells showing transnuclear microtubule tunnels.

For 3D reconstruction of nuclei, cells were treated and imaged as above, Z-stacks were deconvoluted using AutoQuant X3 Deconvolution Software (Media Cybernetics) and 3D renderizations were performed using Imaris software (Bitplane).

Super Resolution Microscopy Imaging

Cells stably co-expressing CHMP4B-linker-GFP and mCherry-Emerin were seeded on 24-wells glass bottom plates (Cellvis) and transfected with the corresponding siRNA. Cells were imaged 48 hours later using 100x oil-immersion objective equipped Nikon Eclipse Ti-2. Inverted spinning disk confocal microscope with a VT-iSIM scan head (VisiTech International) and an environmental chamber. Images were acquired every 15 seconds using a Hamamatsu Orca-Flash4.0 camera. The images were deconvolved using Nikon software.

To quantify the percentage of NE closure at CHMP4B Ti, we selected the first frame of the time-lapse acquisition showing CHMP4B-L-GFP recruitment to the resealing NE (CHMP4B Ti). Next, we used Nikon Software to calculate the nuclear perimeter (NP) by manually drawing a line around the nucleus. Then, we quantified the extent of NE reformation (NER) by manually drawing a line on the mCherry-Emerin signal. Finally, we calculated the ratio NER/NP for every image and refer to this ratio as percentage of NE closure at CHMP4B Ti.

Wide Field Microscopy Imaging

For the analysis of nuclear GFP- or mCherry-NLS fluorescence recovery, cells stably expressing GFP-NLS and H2B-mCherry were transfected twice with the corresponding siRNA and imaged 6 hours after the second transfection. Images were acquired every 5 minutes using a 40x oil-immersion objective equipped Nikon Ti-Eclipse wide-field inverted microscope with attached environmental chamber. Z-Stacks of 3 slices with a separation of 0.9 μm were acquired using an ORCA-Flash 4.0 camera (Hamamatsu). For the quantification of the nuclear to cytoplasmic ratio of GFP-NLS, a square ROI was manually drawn and nuclear and cytoplasmic GFP fluorescence, as well as background fluorescence, were quantified for every time point using Fiji (Schindelin et al., 2012). For nuclear expansion rate quantification, the area corresponding to the H2B-mCherry channel was scored in every frame using a thresholding method in Fiji. For the analysis of nuclear mCherry-NLS fluorescence recovery the same method was followed but cells stably expressing mCherry-NLS were incubated with Hoechst 33258 to stain the DNA 30 minutes before imaging.

For the quantification of anaphase onset and furrow ingression, cells stably expressing GFP-NLS and H2B-mCherry were transfected twice with the corresponding siRNA and imaged 6 hours after the second transfection. Images were acquired every 1 minute using a 40x oil-immersion objective equipped Nikon Ti-Eclipse wide-field inverted microscope with attached environmental chamber and connected to an ORCA-Flash 4.0 camera (Hamamatsu).

The quantification of midbodies abscession time in HCT116 cells stably co-expressing mCherry-Tubulin with CHMP2A-L-GFP or CHMP4B-L-GFP was performed as previously described (Caballe et al., 2015).

Protein Expression and Purification

cDNA encoding Lgd (residues 550-816), Lgd (residues 358-816) and CC2D1B (residues 317-558) were cloned into expression vector pProEx-Htb (Invitrogen) and protein expression was performed in BL21 E. coli cells. Cells were grown at 37 °C to an OD of 0.6 and protein expression was induced with 1mM IPTG for 3 h.

Bacteria expressing Lgd (residues 550-816) were lysed by sonication in buffer A (50 mM Tris pH 7.5, 150 mM NaCl, 5 mM β-mercaptoethanol, 20 mM imidazole) including EDTA-free protease inhibitors (Sigma), pelleted by centrifugation and the soluble fraction was loaded onto a Ni-NTA (Qiagen) resin. The column was washed with buffer A supplemented with 1 M NaCl and subsequently with buffer B (50 mM Tris pH 7.4, 150 mM NaCl, 50 mM imidazole). The protein was eluted in buffer C (50 mM Tris pH 7.4, 150 mM NaCl, 250 mM imidazole). The His-tag was removed with TEV protease at a ratio of 2 μg for 1 mg of protein overnight at 4 °C. The processed protein was dialyzed against buffer A, loaded on a Ni-NTA resin and the flow-through collected. Lgd (residues 550-816) was further purified on a Superdex 75 column (GE Healthcare) in buffer D (2 mM Tris pH 7.5, 150 mM NaCl, 5 mM β-mercaptoethanol). The same protocol was applied to purify CC2D1B (residues 317-558) and modified to purify Lgd (residues 358-816). After elution from the second Ni-NTA column, Lgd (residues 358-816) was dialyzed against buffer E (25 mM Na-citrate pH 6, 100 mM NaCl, 5 mM β-mercaptoethanol) and further purified on a Mono S column (GE Healthcare). The final purification step included gel filtration on a Superdex 200 column (GE Healthcare) in buffer E.

PIP Pull-Down Experiment

40-μl slurries of eight different phosphoinositide coated beads and control beads (Echelon, Inc) were washed once in 500 μl of binding buffer (10 mM HEPES pH 7.5, 300 mM NaCl, 1% (v/v) Nonidet P-40). After centrifugation, beads were incubated with 15 μg of purified Lgd (residues 358-816), rotating for 4 h at 4 °C. Beads were pelleted by centrifugation and the supernatant was collected as the unbound fraction. The beads were washed five times in 500 μl of binding buffer and Lgd (residues 358-816) was eluted by
boiling the beads for 5 min in 20 μl of SDS–PAGE sample loading buffer. Fractions corresponding to unbound and bound proteins were analyzed by SDS-PAGE and bands were detected by Coomassie Blue Staining.

**Sedimentation Experiments**

The sucrose gradient centrifugation experiments were performed as described (Martinelli et al., 2012). Briefly the indicated proteins were separated on sucrose gradients in HBS buffer (0.01 M Heps, pH 7.4, 0.15 M NaCl), by overlaying sucrose solutions of 60% (65 μl), 40%, 30%, 20% and 5% (85 μl each). Centrifugation was performed in a Beckman SW55 rotor at 40,000 rpm for 6h at 4°C. Fractions from the gradients were analysed on a 15% SDS-PAGE and bands were detected with Coomassie Blue staining.

**Crystallization and Structure Solution of Lgd (550-816)**

Lgd (residues 550-816) was concentrated to 5 mg/ml and crystals were obtained by the vapor diffusion method in hanging drops mixing equal volumes of complex and reservoir solution (0.1 M Bis-Tris pH 6.5, 15-25% PEG 3500, 200 mM ammonium sulfate). The crystal was cryo-cooled at 100 K in reservoir buffer containing 25% (v/v) glycerol. The selenomethionine substituted Lgd (residues 550-816) was crystallized under the same conditions. A complete native dataset and SAD data set were collected at the ESRF (Grenoble, France) beam line ID14-4. Data were processed and scaled with MOSFLM (Battye et al., 2011) and SCALA (Evans, 2006). The crystals belong to space group P21 with 2 molecules per asymmetric unit. Although the asymmetric unit contains two protomers, which dimerize via their helical N-terminal domains with a relatively large interface covering 964 Å², MALLS and analytical ultracentrifugation demonstrated a monomeric state of Lgd (550-816) in solution (data not shown). The unit cell dimensions are a=87.81 Å, b=54.22 Å, c=97.26 Å (β=98.19) for the native data set and a=87.08 Å, b=53.77 Å, c=97.26 Å (β=99.49) for the SAD data set. The structure was solved by the single anomalous dispersion method employing the data set collected at the peak wavelength (0.9795 Å) and the SAS protocol of Auto-Rickshaw (Panikkar et al., 2005). Briefly, heavy atom sites were localized using the program SHELXD (Schneider and Sheldrick, 2002) and the correct hand for the substructure was determined using the programs ABS (Hao, 2004) and SHELXE (Sheldrick, 2002). Initial phases were calculated after density modification using the program SHELXE (Sheldrick, 2002) and they were improved by density modification and NCS averaging using the program DM (Cowtan et al., 2011). An initial partial model was built with ARP/wARP (Morris et al., 2004) and BUCANEER (Cowtan, 2006) and refinement with REFMAC (Mursudov et al., 1997) using data to 2.9 Å resolution (R factor of 0.29 and R free of 0.34). The incomplete model was subsequently employed as search model for molecular replacement with Phaser (McCoy et al., 2007) and the 2.4 Å native data set. Model building was completed manually with Coot (Emsley et al., 2010) and the model was refined with REFMAC (Mursudov et al., 1997) and Phenix (Adams et al., 2010) to an R factor of 0.2067 and R free of 0.2459. The model contains chain A residues 575 to 815 and chain B residues 575 to 815. N-terminal residues 550 to 576 were disordered in both protomers and 89.79/7.92 % of the residues are within the most favored and allowed regions of a Ramachandran plot (CCP4, 1994). Molecular graphics figures were generated with PyMOL (W. DeLano; http://www.pymol.org) and sequence alignments with the program ESPript (Gouet et al., 1999).

Modelling of the corresponding CC2D1B fragment was performed with the SWISS-MODEL protein structure homology-modelling server (Biasini et al., 2014). Docking of Ins(1,4,5)P3 to the CC2D1B structural model was performed with the SwissDock web server (Grosdidier et al., 2011).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

All graphs and statistical test were performed using Prism (GraphPad Software). Statistical significance was tested using two-tailed unpaired t-test. All statistical information for quantitative datasets is shown in the corresponding figure legend.

**DATA AND SOFTWARE AVAILABILITY**

Co-ordinates and structure factures corresponding to Lgd (residues 550-816) have been deposited in the Protein Data Bank with accession ID 6EI6.