Contact of cis-Golgi with ER exit sites executes cargo capture and delivery from the ER

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Protein transport from the endoplasmic reticulum (ER) to the Golgi apparatus is mediated by coat complex II (COPII) vesicles. It has been believed that COPII vesicles containing cargo are released from the ER exit sites (ERES) into the cytosol and then reach and fuse with the first post-ER compartment, cis-Golgi or ER-to-Golgi intermediate compartment (ERGIC). However, it still remains elusive how cargo loading to vesicles, vesicle budding, tethering and fusion are coordinated in vivo. Here we show, using extremely high speed and high resolution confocal microscopy, that the cis-Golgi in the budding yeast *Saccharomyces cerevisiae* approaches and contacts the ERES. The COPII coat cage then collapses and the cis-Golgi captures cargo. The cis-Golgi, thus loaded with cargo, then leaves the ERES. We propose that this ‘hug-and-kiss’ behaviour of cis-Golgi ensures efficient and targeted cargo transport from the ERES to cis-Golgi.

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The basic mechanism of endoplasmic reticulum (ER)-Golgi trafficking is conserved from yeast to higher animals and plants, but the number, size and dynamics of the ER exit sites (ERES) and Golgi vary across cell types and species. Most of the first post-ER compartments are located in the vicinity of the ERES. In C. elegans, plant cells and also the yeast P. pastoris, which have separate stacks of flattened cis-Golgi cisternae in the cytoplasm, the cis face of Golgi stacks show stable association with the ERES. In mammalian cells, Golgi stacks are clustered and connected with each other, forming a large Golgi ribbon in the perinuclear region. However, ER-to-Golgi intermediate compartment (ERGIC) is found near the ERES, which locate not only in the perinuclear region but also on the ER spreading throughout the cell.

The yeast S. cerevisiae presents a unique example. Specifically, individual Golgi cisternae, cis, medial and trans, do not stack but scatter in the cytoplasm. Our previous findings indicated that, among Golgi cisternae moving around in the cytoplasm, cis-Golgi has higher probability of staying in the vicinity of the ERES than trans-Golgi. To understand the significance of this behaviour of yeast Golgi, we employed an extremely high speed and high resolution microscopy technique, super-resolution confocal live imaging microscopy (SCLIM), that we recently developed.

ER-to-cis-Golgi and ER-to-ERGIC traffic has been believed to operate in a way that ER-derived COPII vesicles are once released into the cytosol and then get tethered to and fuse with neighbouring cis-Golgi or ERGIC. Our observations, including visualization of cargo, challenge this model. In S. cerevisiae, cis-Golgi approaches and contacts the ERES and, concomitantly with the collapse of COPII coats, captures cargo and then leaves the ERES.

Results

**cis-Golgi shows approach-and-contact actions toward the ERES.** We first conducted simultaneous dual-colour time-lapse observation of the ERES and cis- and trans-Golgi cisternae. ERES is defined as the place where COPII coat assembles, and cis-Golgi is defined as the place where COPII coat assembles, and cis-Golgi has higher probability of staying in the vicinity of the ERES than trans-Golgi. To understand the significance of this behaviour of yeast Golgi, we employed an extremely high speed and high resolution microscopy technique, super-resolution confocal live imaging microscopy (SCLIM), that we recently developed.

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**cis-Golgi is stuck at the ERES when secretion is compromised.** We next asked whether the 'hug-and-kiss' action of cis-Golgi to the ERES was affected when secretion was compromised. Glucose deprivation has been known to impair protein secretion in S. cerevisiae and cause slowing down of Golgi cisternal maturation and coalescence of the ERES. 3D images of glucose-deprived cells showed that the proportion of cis-Golgi cisternae associated with the ERES increased (26 ± 10% with glucose and 63 ± 23% without glucose (mean ± s.d.), whereas trans-Golgi remained mostly separate from the ERES (7 ± 8% with glucose and 4 ± 8% without glucose) (Fig. 2a,b). 3D time-lapse images of mRFP-Sed5 and Sec13-GFP in glucose-deprived cells demonstrated that cis-Golgi cisternae were largely stuck at the ERES and the Sec13-GFP signal did not decrease (Fig. 2c). Persistent contact between cis-Golgi and the ERES was also observed in cells treated with cycloheximide to block protein synthesis (Fig. 3).

**cis-Golgi is stuck at the ERES in tether-impaired cells.** Biochemical studies have revealed that the directionality of traffic from the ER to the Golgi is controlled by the sequential interactions between COPII coats and tethering factors. The tethering complex TRAPPi contains the activity as the guanine-nucleotide exchange factor for Ypt1 and binds to Sec23. TRAPPi activates Ypt1 thereby inducing binding between Ypt1 and its effector Usol, a long coiled-coil tethering protein localizing on the Golgi membrane. Subsequently, the protein kinase Hrr25 (ref. 21) localizing on the Golgi phosphorylates Sec23/24 complex, which facilitates shedding of COPII coats. This suggests that COPII coats are not removed from the membrane until reaching the Golgi membrane. We examined the relationship between the ERES and cis-Golgi in the usol1-1 cells, which lack the Usol tethering function at the restrictive temperature (37 °C). As shown in Fig. 4, under this condition, cis-Golgi stayed associated with the ERES (17 ± 12% at 25°C and 47 ± 16% at 37°C) (Fig. 4a,b) without invoking the decrease in Sec13-GFP signals (Fig. 4c). trans-Golgi was still freely moving in the cytoplasm (5 ± 6% at 25°C and 4 ± 5% at 37°C) (Fig. 4c). Taken together, these findings suggest that COPII coats are released only when normal functions of Usol and other transport machinery are fulfilled.

**Cargo is transiently loaded into the ERES.** Our data imply an intriguing possibility that cis-Golgi captures cargo at the ERES via the hug-and-kiss action. Live cell visualization of cargo transport along the secretory pathway had been a big challenge, but we succeeded in setting up a system to do so. We chose a GFP-tagged cargo Axl2-GFP, a transmembrane protein targeted to the plasma membrane of the bud tip, and expressed it under a heat-shock promoter (SSA1) in the temperature-sensitive sec31-1 mutant. The sec31-1 mutant is defective in the assembly of the Sec31 coat at the restrictive temperature (39°C). At this temperature, new synthesis of Axl2-GFP was induced but the COPII budding was...
blocked. Thus, the cargo Axl2-GFP accumulated in the ER of these cells (Fig. 5a, upper panel). On temperature shift-down to 25°C, the cells were restored from the COPII-budding block and Axl2-GFP, whose new synthesis was repressed, started export from the ER and went on to the Golgi (Fig. 5a, lower panel). Cargo loading into the ERES was observed in sec31-1 cells expressing heat-shock-induced Axl2-GFP and constitutive Sec13-mRFP (Fig. 5b). Two-dimensional time-lapse images showed that a small portion of cargo was loaded into the ERES on release from the COPII-budding block; the cargo signal then disappeared from the ERES coincidently with a decrease in the signal of Sec13-mRFP (Fig. 5b,c).

**cis-Golgi captures cargo at the ERES.** To visualize the behaviour of cargo in relation to ERES-to-Golgi transport, we next examined sec31-1 cells that constitutively expressed Mnn9-mCherry, a subunit of Golgi mannosyltransferase complex that localizes mainly in cis-Golgi tagged with mCherry, and the heat-shock-inducible Axl2-GFP. After the temperature shift to 25°C, Axl2-GFP was exported from the ER to the Mnn9-mCherry-labelled cis-Golgi (Fig. 6a, left panels). 2D and 3D time-lapse images of Mnn9-mCherry and Axl2-GFP demonstrate that cis-Golgi began to associate with a particular region of the ER, most probably the ERES, and captured cargo. Then the cis-Golgi left the ER together with the cargo (Fig. 6a,b, and Supplementary Movies 3–5).

**Discussion**

Our data have clearly demonstrated that yeast cis-Golgi contacts the ERES, by what we would call a hug-and-kiss action, and captures cargo during this contact (see Fig. 7 for our model). Our findings suggest that COPII vesicle do not necessarily have to be
released into the cytosol. They can be captured by cis-Golgi before the release. This mechanism appears to be much safer than forming many free COPII vesicles in the cytosol and also explains why COPII vesicles are not easily found in electron micrographs because COPII vesicles are not easily found in electron micrographs. Tethers such as the microtubule cytoskeletons probably do not play a role here, because cis-Golgi in S. cerevisiae moves independently of these cytoskeletons. Some matrix-like protein might stabilize the transport even in the species like S. cerevisiae that does not have highly organized Golgi structures.

Secretory cargoes vary in size; therefore, some cargo proteins, such as procollagen fibres in mammalian cells, are too large to load into conventional COPII vesicles, and oversized vesicles have been postulated to generate to accommodate secretion of such large cargoes. The oversized COPII vesicles may not have to be sealed and released if our proposed mechanism operates. A big question remains how the cis-Golgi finds the ERES. Actin and microtubule cytoskeletons probably do not play a role here, because cis-Golgi in S. cerevisiae moves independently of these cytoskeletons. Some matrix-like protein might stabilize the interaction between the ERES and the cis-Golgi. TFG protein of C. elegans and mammals is an interesting candidate, but yeast does not seem to have its close homologue. Tethers such as the actin and microtubule cytoskeletons probably do not play a role here, because cis-Golgi in S. cerevisiae moves independently of these cytoskeletons.
TRAPP complex and Uso1 are intriguing players that could guide the cis-Golgi to the ERES, and if so the word ‘hug’ would be appropriate. The functions of these tethers in hug-and-kiss actions will of course be a target of our future studies.

We previously demonstrated that yeast Golgi cisternae mature over time; the cis-Golgi changes its nature via medial-Golgi to trans-Golgi.3,31 An interesting question would be whether the cis-Golgi approaching the ERES goes into the maturation phase immediately after contact. In the glucose deprivation experiment (Fig. 3), a certain fraction of cis-Golgi was left unassociated with ERES, even though there are more ERES than cis-Golgi in number.8 This might be because there are two populations of cis-Golgi, one for capture of cargo from the ERES (hug-and-kiss) and the other for carbohydrate modification (maturation). The former cis-Golgi could be regarded as similar to the mammalian ERGIC. Recent studies on plant Golgi support this view.4,32 We have already made a substantial progress towards simultaneous multicolour observation at high-speed and super-high-resolution9 and will fully utilize it to address these and other unresolved questions by live cell imaging.
**Methods**

**Yeast strains, plasmids and culture conditions.** Yeast cells were grown in MCD medium (0.67% yeast nitrogen base without amino acids (Difco Laboratories), 0.5% casamino acids (Difco Laboratories) and 2% glucose) with appropriate supplements. For live imaging, cells were grown to a mid-log phase at 25°C.

Strains expressing fluorescent protein-tagged Sec13, Mnn9 were constructed by a PCR-based method using pFA6a plasmids as a template, which is described in the yeast GFP database at the University of California, San Francisco (primers 5’-TTTATGGAAGGAAAATCTTGAGGGTAAATGGGAACCCGCTGGTGAAGTTCATCAGCGGATCCCCGGGTTAATTAA-3’ and 5’-CTCATTTGCATTCTTTTTTCTTTTGAGATGTTTCATTTTAAATTCTTGATACTCTGAATTCGAGCTCGTTTAAAC-3’ for Sec13 and primers 5’-ATTTGGCTTACCAAACTATTTGGTTTATCACATAGAGGAAGAGAACCATCGGATCCCCGGGTTAATTTAA-3’ and 5’-CTTCAATTGATGTTTCATTTTAAATTCTTGATACTCTGAATTCGAGCTCGTTTAAAC-3’ for Mnn9). mRFP-Sed5 was expressed under the control of the TDH3 promoter on the low-copy plasmid pRS316 (ref. 34). Sec7-mRFP was expressed similarly except that the ADH1 promoter was used instead of the TDH3 promoter. Axl2-GFP was expressed under the control of a heat-shock promoter on the low-copy plasmid pRS316 or on the integrated plasmid pRS304.

**Fluorescence microscopy.** Cells were immobilized on glass slides using concanavalin A and imaged by SCLIM. SCLIM was developed by combining an Olympus model IX-71 inverted fluorescence microscope with a UPlanSApo 100NA 1.4 oil objective lens (Olympus, Japan), a high-speed and high-signal-to-noise-ratio spinning-disk confocal scanner (Yokogawa Electric, Japan), a custom-made spectroscopic unit, image intensifiers (Hamamatsu Photonics, Japan) equipped with a custom-made cooling system and two EM-CCD cameras (Hamamatsu Photonics, Japan) for green and red observation. For raising space resolution, a magnification lens (4 or 10) was put in the light path between the confocal scanner and the spectroscopic unit. For 3D images, we collected optical sections spaced 0.1μm apart in stacks by oscillating the objective lens vertically with custom-made piezo actuator (Yokogawa Electric) relative to sample planes. Typically, 80 optical sections could span whole cells. For 3D time-lapse images, we took sequential 7 or 15 optical sections around the centre of a cell.

**Figure 6 | cis-Golgi captures cargo at the ERES.** sec31-1 cells expressing heat-shock-inducible Axl2-GFP (cargo, green) and constitutive Mnn9-mCherry (cis-Golgi, red) were released from the COPII-budding block and then imaged. (a) Axl2-GFP is partly co-localized with cis-Golgi (left panels). Right panels show representative 2D time-lapse images of Axl2-GFP and cis-Golgi (boxed area). (b) 3D images were reconstructed from 15 optical slices 0.1μm apart around the centre of cells taken at 0.2-s intervals (left panels). Right panels show representative 3D time-lapse images of Axl2-GFP and cis-Golgi (boxed areas). Dashed circles, Axl2-GFP contained in cis-Golgi. Asterisks, Axl2-GFP captured by cis-Golgi. Arrowheads, Axl2-GFP leaving from the ER together with cis-Golgi. The lower panels are high-contrasted images to show the small amount of Axl2-GFP. Scale bar, 1μm (a,b).

**Figure 7 | A model of ER-Golgi cargo transport by a ‘hug-and-kiss’ action of cis-Golgi.** The ERES localizes on high-curvature ER domains such as on ER tubules or at the edge of ER sheets, and cargo protein (green) synthesized in the ER is loaded into the ERES. cis-Golgi (red large ellipse) performs an approach-and-contact (hug-and-kiss) action towards the ERES. During contact between COPII buds/vesicles and cis-Golgi, the COPII coat (blue line) cage collapses. cis-Golgi that received cargo then leaves the ERES.
software (Perkin Elmer, MA) was used to reconstruct and deconvolve 3D images via point-spread functions optimized for our spinning-disk confocal scanner. MetaMorph software (Molecular Devices, CA) was used for showing time-lapse images and for fluorescence signal change analysis. A thermo-controlled stage (Tokai Hit, Japan) was used to observe temperature-sensitive mutants.

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Author contributions

K.K. designed the study, performed all the experiments, analysed the data and wrote the manuscript. M.O. generated the yeast strains and plasmids. A.N. supervised the study.

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