TFG facilitates outer coat disassembly on COPII transport carriers to promote tethering and fusion with ER–Golgi intermediate compartments

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Edited by Jennifer Lippincott-Schwartz, Howard Hughes Medical Institute, Ashburn, VA, and approved July 31, 2017 (received for review June 2, 2017)

The conserved coat protein complex II (COPII) mediates the initial steps of secretory protein trafficking by assembling onto subdomains of the endoplasmic reticulum (ER) in two layers to generate cargo-laden transport carriers that ultimately fuse with an adjacent ER–Golgi intermediate compartment (ERGIC). Here, we demonstrate that Trk-fused gene (TFG) binds directly to the inner layer of the COPII coat. Specifically, the TFG C terminus interacts with Sec23 through a shared interface with the outer COPII coat and the cargo receptor Tango1/cTAGES. Our findings indicate that TFG binding to Sec23 outcompetes other associations in a concentration-dependent manner and ultimately promotes outer coat disassembly. Additionally, we demonstrate that TFG tethers vesicles harboring the inner COPII coat, which contributes to their clustering between the ER and ERGIC in cells. Together, our studies define a mechanism by which COPII transport carriers are retained locally at the ER/ERGIC interface after outer coat disassembly, which is a prerequisite for fusion with ERGIC membranes.

COPII | Trk-fused gene | tether | coat disassembly | endoplasmic reticulum

In most metazoan systems, cargoes transported from the endoplasmic reticulum (ER) must traverse a vesicular-tubular cluster of membranes known as the “ER–Golgi intermediate compartment” (ERGIC) en route to the Golgi apparatus (1–5). This process involves the function of two distinct coat complexes (6–9). The conserved coat protein complex II (COPII) assembles at ER subdomains and mediates transport to the ERGIC (5, 10–12), whereas the COPII complex directs transport from the ERGIC to the Golgi as well as retrograde transport to the ER (5, 13–15). In both cases, the coats define the architecture of the transport carriers but must eventually disassemble to enable membrane fusion at the target compartment (16, 17). In the case of COPII, which minimally consists of seven core subunits and an ADP ribosylation factor (Arf)-type GTPase (18, 19), uncoating is promoted by association with target membrane tethers such as the Dsl1 complex, which interferes with COPII subunit interactions, and stimulation of GTP hydrolysis on Arf by members of the Arf guanine nucleotide-activating protein (GAP) (ArafGAP) family (20–23). In contrast, mechanisms that direct COPII uncoating have been more controversial. The COPII coat consists of two layers, an inner adaptor layer composed of the GTPase Sar1 and Sec23–Sec24 heterodimers and an outer lattice-like cage made up of Sec13–Sec31 heterotetramers (7, 24, 25). Initial studies suggested that Sar1 is required for COPII coat disassembly after transport carriers undergo scission from the ER (26–29). However, subsequent in vitro studies contended that Sar1 is rapidly removed from COPII carriers immediately following their scission at the ER (14, 30). More recently, phosphorylation of COPII subunits at ERGIC and Golgi membranes has been suggested to promote uncoating, but in vitro studies have failed to support this idea, and the consequences of posttranslational modifications on COPII coat stability remain unclear (14, 31–33). Related to COPII tethering by the Dsl1 complex, the transport protein particle (TRAPP) complex tethers COPII transport carriers at the ERGIC and co-Golgi (14, 20, 30, 34). However, unlike the function of Dsl1, biochemical studies suggest that TRAPP associates with Sec23 on a site that overlaps with its binding domain for Sar1, which is not predicted to disrupt inner or outer coat integrity. Moreover, recent work suggests that TRAPP actually promotes outer COPII coat assembly during the formation of transport carriers (35). Thus, it remains unknown how the COPII coat disassembles before fusion at the ERGIC.

Several additional regulators of COPII-mediated trafficking have also been described recently, some of which directly impact coat stability. Members of the Sec16 family function at ER subdomains to inhibit Sar1 GTPase activity and facilitate COPII coat assembly (36–38). In a related manner, the cargo receptors Tango1 and cTAGES associate directly with Sec23 at the ER to promote the formation of elongated COPII-coated tubules (39, 40). Ultimately, outer coat assembly has been suggested to out-compete Tango1/cTAGES and Sec16 interactions with the inner coat to enable fission of transport carriers in a manner dependent on Sar1 GTP hydrolysis (40, 41).

Significance

The endoplasmic reticulum (ER) serves as a platform for the packaging of most secretory proteins into conserved coat protein complex II (COPII)-coated transport carriers destined for ER–Golgi intermediate compartments (ERGIC) in animal cells. In this work, we demonstrate that Trk-fused gene (TFG), a protein implicated in multiple neurodegenerative diseases and oncogenesis, functions in this pathway by interacting directly with the COPII protein Sec23. Specifically, we show that TFG outcompetes interactions between the inner and outer layers of the COPII coat, indicating that TFG promotes the uncoating process after transport carriers undergo scission from the ER. Moreover, we demonstrate that TFG simultaneously captures and concentrates COPII transport carriers at the ER/ERGIC interface to enable the rapid movement of secretory cargoes to the ERGIC.
Following release from the ER, COPII transport carriers are restricted to the ER/ERGIC interface (42, 43). Our previous studies indicated that this distribution is regulated by Trk-fused gene (TFG), a homo-oligomeric protein complex that assembles into a meshwork and colocalizes with COPII carriers in cells (44, 45). Here, we demonstrate that TFG associates directly with Sec23 and exhibits the ability to outcompete interactions between layers of the COPII coat. Our data are most consistent with a model in which TFG promotes the release of the outer COPII coat and restricts diffusion of carriers harboring the inner coat until tethering occurs at ERGIC membranes. In doing so, TFG functions to maintain the integrity of the early secretory pathway.

Results

TFG Facilitates the Export of Conventional Cargoes from the ER. We demonstrated previously that TFG plays a conserved role in regulating the export of diverse cargoes, including SNAREs, Golgi enzymes, and other biosynthetic transmembrane proteins of all sizes, from the ER (44, 45). Additionally, we found that penetrant depletion of TFG disrupts early secretory pathway integrity, leading to the accumulation of inner and outer COPII-coated transport carriers away from the ER/ERGIC interface (44, 46). A recent report similarly demonstrated that TFG promotes organization of ER exit sites but suggested a specialized function for TFG in the export of large cargoes, such as procollagens (47). To test directly the contribution of TFG to COPII-dependent cargo transport, we used a reconstructed system in which permeabilized cells lacking cytosol were supplemented with purified recombinant components (10, 11, 48). As described previously, the addition of COPII subunits was sufficient to promote the formation of transport carriers harboring small, native cargoes, including the lectin ERGIC-53 and the R-SNARE Sec22B, but not ribophorin, an ER resident protein (Fig. 1A and Fig. S1A). The addition of TFG further stimulated the release of ERGIC-53- and Sec22B-containing transport carriers (Fig. S1B) but exhibited a negligible effect on procollagen (PC1) secretion (Fig. 1A and Fig. S1A), indicating that the role of TFG in the early secretory pathway is unlikely to be specific for large cargoes.

We therefore questioned whether the modest effects of TFG inhibition on the export of small artificial cargoes reported recently (47) could be a consequence of insufficient protein depletion. To address this possibility, we first defined conditions under which TFG was partially (∼44%) and more fully (∼77%) depleted (Fig. 1C). We then examined the distribution of COPII carriers relative to Sec16A, a marker of ER subdomains at which COPII coat assembly occurs (50). When TFG was partially depleted, COPII structures that were modestly reduced in intensity remained juxtaposed to sites marked by Sec16A (Fig. 1D and Fig. S1C), consistent with recently published findings (47). However, upon more penetrant inhibition of TFG, as determined by immunofluorescence studies (Fig. S1D), we observed a dramatic increase in the number of COPII and ERGIC-53 labeled structures, which were no longer juxtaposed to Sec16A (Fig. 1D and E and Fig. S1E). Notably, under all conditions, Sec16A-labeled sites continued to exhibit juxtaposed Sec24A, indicating that ER subdomains continue to produce COPII transport carriers in the absence of TFG (Fig. S1F).

Since human tissue-culture cells undergo apoptosis rapidly after penetrant depletion of TFG (44), we turned to the Caenorhabditis elegans germline as a model system, where we previously demonstrated an ability to achieve greater than 95% inhibition of TFG expression (45). Examination of both large (E-cadherin; >3,000 amino acids) and small (eight different transmembrane SNAREs and the minimal transmembrane domain of mannosidase II) cargoes demonstrated a uniform defect in secretion from the ER following TFG depletion (Fig. S1G). Taken together, these data indicate that TFG plays a general role in the normal trafficking of conventional COPII carriers.

Dysregulation of COPII function in rodent models results in varying phenotypes ranging from early embryo lethality to more modest impacts on physiology due to redundancies in subunit expression (51, 52). If TFG functioned only in the secretion of large cargoes such as procollagens, we would predict that its deletion phenotype would be similar to that of Tangle1-knockout animals, which exhibit several developmental abnormalities but are born in normal Mendelian frequencies (53). Using CRISPR-mediated genome editing, we isolated heterozygous rats harboring a 44-bp deletion within the first coding exon of TFG, which results in an early frame shift and produces only the first 21 amino acids of native TFG (Fig. S1H). After outcrossing one of the founders six times, we mated heterozygous animals and genotyped 114 progeny that were born from five distinct mating pairs. In total, we obtained 39 wild-type animals, 75 heterozygous animals, and no homozygous-null...
animals (Table S1). We also performed timed pregnancies and isolated embryos at E9.5 and E14. Genotyping revealed that none of the recovered embryos were homozygous-null (Table S1). Together, these data suggest that loss of TFG results in early embryo lethality in rats before E9.5, which is more consistent with a role for TFG in general protein secretion.

TFG Associates Directly with the Inner COPII Coat Protein Sec23. To investigate potential mechanisms by which TFG regulates COPII-mediated protein transport, we conducted a series of immunoprecipitations using TFG antibodies and freshly prepared rat liver cytosol and two distinct human cell lines (RPE1 and HeLa). These studies revealed specific interactions with Sec23 but not other components of the COPII machinery (Fig. 2A and Table S2). Additionally, we conducted a directed yeast two-hybrid screen to identify TFG-interacting partners in the early secretory pathway (Table S3). This approach again revealed a specific association between TFG and the inner COPII coat protein Sec23 (Fig. 2B). Consistent with these findings, recombinant GST-tagged TFG, but not GST alone, was also capable of recovering native Sec23 from rat liver cytosol (Fig. S2A).

To delineate the region of TFG necessary for its association with Sec23, we again used a yeast two-hybrid approach. Our findings indicated that the C-terminal half of TFG (amino acids 194–400), which we demonstrated previously to be intrinsically disordered (44), was necessary and sufficient to bind to Sec23 (Fig. 2C). To determine whether the interaction was direct, we purified recombinant forms of Sec23 and the TFG C terminus and examined their elution profiles separately and in combination following gel filtration chromatography. Individually, Sec23 exhibited a Stokes radius of 2.4 nm, whereas the TFG C terminus exhibited a Stokes radius of 3.5 nm (Fig. 2D). In contrast, when the proteins were mixed, they coeluted with an average Stokes radius of 3.6 nm, indicating that they formed a complex (Fig. 2D and E). To determine the molecular mass of the complex, we used a combination of size-exclusion chromatography and multiangle light scattering (Fig. S2 B–E). These data demonstrated that Sec23 and the TFG C terminus form a 127.8 (±1.4)-kDa complex, consistent with a 1:1 association (Fig. S2E).

The Localization of TFG Is Governed by Its Association with COPII-Coated Transport Carriers. The C-terminal portion of TFG is poorly conserved across metazoans based on amino acid alignment algorithms (Fig. S2F). However, its overall amino acid content varies relatively little from C. elegans (≈30% proline, 17% glycine, 13% glutamine, 10% alanine, and 8% serine) to humans (≈20% proline, 10% glycine, 21% glutamine, 11% alanine, and 9% serine). To determine the specific region of TFG necessary for binding to Sec23, we began truncating TFG from its C-terminal end and measuring the impact on Sec23 binding using the yeast two-hybrid system. Removal of as little as 10 amino acids from the C terminus of TFG had a dramatic impact on Sec23 binding, and deletion of 20 amino acids reduced binding to background levels (Fig. 3A). These findings were confirmed using recombinant forms of TFG in pull-down experiments from rat liver cytosol (Fig. 3B and Fig. S3A). By contrast, deletion of the entire C-terminal half of TFG with the exception of the last 22 amino acids (i.e., deletion of amino acids 194–378) preserved its ability to interact with Sec23 (Fig. 3A). These data define the minimal domain of TFG that is necessary for its association with Sec23 (Fig. S2G), although its proline-rich domain (PRD) may also contribute to binding, based on previous work examining PRDs from Sec31 and Tango1/cTAGE5 (40).

We previously demonstrated that TFG localizes to the ER/ERGIC interface together with COPII-coated transport carriers (44, 45). To determine whether its interaction with Sec23 is necessary, we depleted the endogenous form of TFG using a siRNA targeting its 3′ UTR and replaced it with various TFG transgenes that we predict would alter Sec23 binding in cells (Fig. 3 C and D). To maintain tight control over relative transgene expression (Fig. S3 B and C), we inserted a single copy of each at the adeno-associated virus insertion site.

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Fig. 2. TFG interacts directly with the inner COPII coat subunit Sec23. (A) Immunoprecipitations using IgG or antibodies directed against TFG were conducted using rat liver cytosol, separated by SDS-PAGE, and immunoblotted using the indicated antibodies (n = 3 in each condition). Asterisks highlight nonspecific bands recognized by Sec23 and Sec13 antibodies. (B) Yeast coexpressing plasmids encoding TFG (bait fusion) and several unique prey constructs were plated (10-fold dilutions, left to right) on either selective (−Ura, −Leu, −His) or histidine-supplemented medium for 48 h (n = 3). The GTPase-deficient form of Sar1B (H79G) was used in these experiments. (C) Yeast coexpressing plasmids encoding Sec23A (bait fusion) and prey constructs encoding distinct regions of TFG were plated as described in A (n = 3). Coexpression of bait and prey constructs encoding TFG fusions was used as a positive control, and an empty prey construct was used for a negative control. (D) Purified C. elegans Sec23 and TFG C-terminal domain (CTD), amino acids 196–486, were separated individually and as a mixture by gel filtration chromatography, and specific fractions were analyzed by SDS-PAGE followed by Coomassie staining (n = 3 in each condition). Stokes radii were calculated based on the elution profile of known standards. MW, molecular weight marker. (E) Retention times of C. elegans TFG (amino acids 196–486) and full-length Sec23 (individually and as a mixture) were plotted based on densitometry after gel filtration chromatography and SDS-PAGE analysis of fractions. AU, arbitrary units.
tegration site 1 (AAVS1) safe harbor locus using TALEN-mediated genome editing. All transgenes were placed under control of the Tet-On system and were followed by an internal ribosome entry site (IRES) RFP cassette to monitor expression levels quantitatively by fluorescence microscopy (Fig. 3D and Fig. S3C). Replacement of endogenous TFG with a full-length, untagged transgene did not affect its colocalization with COPII carriers (Fig. 3D). In contrast, transgenes lacking either 10 or 20 amino acids from the TFG C terminus exhibited reduced colocalization with COPII-labeled structures (Fig. 3D–F and Fig. S3D and E). Additionally, overexpression of TFG was shown previously to generate large, potentially phase-separated regions of the cytoplasm capable of titrating COPII transport carriers (44). However, overexpression of truncated TFG (amino acids 1–380) abolished this ability (Fig. S3F). Truncated isoforms of TFG were also incapable of promoting COPII carrier budding in vitro, unlike the full-length protein (Fig. S3 F and G). Together, these data are consistent with the idea that an interaction with Sec23 is necessary for TFG to assemble at the ER/ERGIC interface and facilitate COPII-mediated trafficking.

To confirm a role for COPII transport carriers in directing TFG distribution, we examined the impact of inhibiting inner COPII coat assembly. In mammals, two largely redundant isoforms of Sec23 are expressed (Sec23A and Sec23B). Using CRISPR-mediated genome editing, we generated three independent RPE1 cell lines lacking the Sec23B isoform (Fig. 4A and Fig. S4A–E), which enabled us to achieve a dramatic reduction in COPII accumulation at ER subdomains upon siRNA-mediated depletion of Sec23A (Fig. 4B and C and Fig. S4F). Under these conditions, we found that TFG was distributed diffusely throughout the cytoplasm and largely failed to accumulate near Sec16A-labeled subdomains at the ER (Fig. 4A, D, and E and Fig. S4A–C, F, and G), even though Sec16A levels were dramatically elevated (Fig. S4F). These data indicate that Sec23 is the major binding partner of TFG in the early secretory pathway, although other COPII-associated factors may...
also contribute to its recruitment (45), and that the presence of COPII transport carriers is required for TFG assembly at the ER/ERGIC interface.

**TFG Competes with the Outer COPII Coat for Inner Coat Binding.** Based on our studies and previous work from others, Sec23 appears to integrate the actions of multiple regulators of COPII-mediated transport via its ability to associate with numerous factors including Sar1, Sec24, Sec31, Sec16, TRAPPC3, p125A, members of the TANGO1 cargo receptor family, and, now, TFG (10, 14, 36, 39, 54–56). Some of these interactions have been shown to be competitive, suggesting that unique associations with Sec23 occur sequentially as transport carriers form, undergo scission from the ER, and tether to ERGIC membranes (14, 40). We used confocal and stimulated emission depletion (STED) microscopy to define the relative distribution of TFG with several of these factors, based on the availability of validated antibodies. Consistent with our previous work, TFG was juxtaposed to Sec16, a marker of ER subdomains that produce COPII transport carriers, and colocalized with Sec23, Sec24, and Sec31, coat proteins on the carriers that accumulate at the ER/ERGIC interface under steady-state conditions (Fig. 5A and B and Fig. S5A and B). Notably, based on the analysis of numerous points of colocalization, TFG exhibited more extensive overlap with inner COPII subunits than with outer COPII subunits (Fig. 5A and B and Fig. S5A). Tango1 colocalized with the Sar1 exchange factor Sec12 on ER subdomains and was juxtaposed to TFG, consistent with the idea that Tango1 associates...
with COPII carriers as they initially form but fails to be incorporated into the carriers when they leave the ER (Fig. 5C and Fig. S5C and D). Together, our data suggest that TFG binds to COPII transport carriers subsequent to their scission from ER subdomains.

Both Sec31 and TANGO1/cTAGE5 exploit PRDs to associate with a common motif on Sec23 (40). Given the presence of a PRD in TFG, we investigated the possibility that it interacts with Sec23 in a manner that would compete with Sec31 and/or TANGO1. We first purified a recombinant form of Sec31 fused to GST, which included its PRD and a neighboring fragment previously shown to stimulate Sec23-mediated GTP hydrolysis on Sar1 (29), and immobilized it on glutathione agarose. In contrast to GST alone, Sec32-Sec24 heterodimers were retained on beads harboring GST-Sec31 (Fig. 6A). We then repeated the assay in the presence of either a twofold or fivefold molar excess of the soluble recombinant TFG C terminus and found that Sec23-Sec24 retention by GST-Sec31 was significantly reduced in both cases (Fig. 6A and Fig. S6A). These data suggest that TFG disrupts the association between the inner and outer layers of the COPII coat. In contrast, a truncated form of TFG lacking the final 20 residues implicated in Sec23 binding failed to compete with Sec31 (Fig. 6B). In parallel, we also tested the ability of the TFG C terminus to compete with the TANGO1 PRD for Sec23 binding.

Again, we found that TFG could successfully outcompete Tango1 in retaining Sec23-Sec32 heterodimers, consistent with current models indicating that Tango1 is efficiently removed from COPII transport carriers upon their scission from the ER (Fig. S6B).

To determine the relative affinities of TFG and Sec31 for Sec23, we performed a reciprocal competition assay in which a GST fusion to the TFG C terminus was immobilized on glutathione agarose. Sec23-Sec24 heterodimers bound to resin containing GST-TFG but not to GST alone (Fig. 6C). The addition of a twofold and fivefold molar excess of soluble Sec31 resulted in decreased Sec23-Sec24 retention on the resin, suggesting that Sec31 can similarly compete with TFG for Sec23 binding (Fig. 6C and Fig. S6C). However, based on the high concentration of TFG that accumulates at the ER/ERGIC interface (44, 45) combined with its ability to form octameric complexes (44) with high avidity for Sec23, our findings are consistent with a model in which TFG actively facilitates dissociation of the outer COPII coat.

Sec31 was previously demonstrated to stimulate Sec23-mediated GTP hydrolysis on Sar1 (27). Since TFG and Sec31 exhibit an overlapping binding site on Sec23, we examined whether TFG also plays a role in modulating Sec23 activity, which may promote inner coat disassembly. We used two distinct assays to monitor GTP hydrolysis on Sar1 following the addition of Sec23-Sec24 in

![Figure 6. TFG outcompetes Sec31A for Sec23A binding.](image-url)
liposomes (liposomes 3). 486, harboring a polyhistidine tag) on
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Upper C Sar1 and TFG tethers COPII-coated liposomes. (elegans 3) were maintained at 4 °C, and
Fig. S7 = and C terminus (amino acids 464), and centrifuged. (Right) The lipidosome pellet was resuspended with buffer containing 0.2% Triton X-100, and nitro-
benzoxadiazole (NBD) fluorescence was quantified to determine the degree of tethering. AU, arbitrary units. (C, Upper) Quantification of NBD fluorescence (F) obtained from sedimented liposomes in the pres-
ence of different concentrations of the full-length or truncated TFG C terminus (graphical plots; n = 3). *P < 0.05 (compared with control), calculated using a paired t
test. (Lower) The presence of Sec23 (10% total) and TFG (100% total) with sedimented liposomes was confirmed by SDS/PAGE analysis followed by Coomassie staining. (D) Liposome aggregation was measured (absorbance: 405 nm) over time. Light liposomes harboring Sar1GTP alone or Sar1GTP with Sec23 (COPII) were incubated with heavy liposomes in the presence or absence of TFG (30 nM). Reactions (n = 3) were maintained at 4 °C, and samples were taken at the indicated time points. (E) Microcontact printing was used to generate bio-
tinylated surfaces to which FITC-streptavidin was bound. The C. elegans TFG C terminus (amino acids 196–486) fused to the streptavidin-binding peptide was incubated with these regions and subsequently
bound. The presence of Sec23 GAP activity (Fig. S7A). These data suggest that TFG is unlikely to facilitate inner COPII coat disassembly at the ER/ERGIC interface.

The TFG C Terminus Is Sufficient to Capture COPII-Coated Carriers. In the absence of TFG, transport carriers harboring both the inner and outer layers of the COPII coat accumulate throughout cells (also see Fig. 1) (44). These data suggest that TFG functions both to promote outer coat dissociation and to retain COPII carriers at the ER/ERGIC interface. To directly test whether TFG is capable of capturing COPII-coated vesicles, we de-
veloped an in vitro assay using two types of synthetic liposomes (Fig. 7A) (57). One set of “heavy” liposomes (~250 nm in di-
meter) was generated in the presence of sucrose to enable facile
sedimentation upon low-speed centrifugation and contained lipids that bind polyhistidine-tagged proteins (DOGS-NTA-Ni) with high affinity. Another set of “light” liposomes (~100 nm in diameter) was generated using a mixture of lipids, which enable assembly of the inner COPII coat. A polyhistidine-tagged form of the recombinant TFG C terminus was bound to the heavy liposomes and mixed with COPII-coated light liposomes (Fig. 7B). Upon sedimentation specifically in the presence of TFG, we re-
covered both types of liposomes, indicating that the TFG C ter-
minus was sufficient to bind inner COPII-coated vesicles (Fig. 7C).

Consistent with these findings, the presence of the TFG C terminus on vesicles was sufficient to promote the aggregation of COPII-
coated vesicles, as determined by spectrophotometric measure-
ments (Fig. 7D). Importantly, a truncated version of the TFG C terminus lacking the last 20 amino acids necessary for efficient Sec23 binding exhibited a reduced ability to mediate cosedi-
mentation of COPII-coated light liposomes (Fig. 7B and C).

To confirm a role for TFG in capturing COPII transport carriers, we generated clusters of its C terminus on glass coverslips using a microcontact printing approach (58, 59). The TFG-enriched domains were exposed to synthetic liposomes containing rhodamine-labeled lipids and were imaged using confocal microscopy. Only when the synthetic liposomes were precoated with the inner COPII proteins were they retained on the artificial TFG microdomains (Fig. 7E and Fig. S7B). To-
gether, our data support a model in which TFG facilitates outer
coating of COPII-coated light liposomes (Fig. 7B, C, and D)
and Sec23 or
Sec23 (COPII) were incubated with
TFG is capable of capturing COPII-coated vesicles, we de-

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Fig. 7. TFG tethers COPII-coated liposomes. (A) Diagram depicting the liposome-tethering assay highlighting the presence of C. elegans TFG (amino acids 196–486, harboring a polyhistidine tag) on heavy liposomes containing DOGS-NTA(Ni) (gray) and COPII components (C. elegans Sar1GTP and Sec23) bound to light liposomes (yellow). (B, Left) Light liposomes harboring COPII (Sar1GTP, 2 μM; Sec23, 250 nM) were mixed with heavy liposomes (1:1 ratio, 1 nM total lipids), bound to varying amounts (0, 10, 30, and 100 nM) of full-length TFG C terminus (amino acids 196–486) or truncated TFG C terminus (amino acids 196–464), and centrifuged. (Right) The liposome pellet was resuspended with buffer containing 0.2% Triton X-100, and nitro-
benzoxadiazole (NBD) fluorescence was quantified to determine the degree of tethering. AU, arbitrary units. (C, Upper) Quantification of NBD fluorescence (F) obtained from sedimented liposomes in the pres-
ence of different concentrations of the full-length or truncated TFG C terminus (graphical plots; n = 3). *P < 0.05 (compared with control), calculated using a paired t

test. (Lower) The presence of Sec23 (10% total) and TFG (100% total) with sedimented liposomes was confirmed by SDS/PAGE analysis followed by Coomassie staining. (D) Liposome aggregation was measured (absorbance: 405 nm) over time. Light liposomes harboring Sar1GTP alone or Sar1GTP with Sec23 (COPII) were incubated with heavy liposomes in the presence or absence of TFG (30 nM). Reactions (n = 3) were maintained at 4 °C, and samples were taken at the indicated time points. (E) Microcontact printing was used to generate bio-
tinylated surfaces to which FITC-streptavidin was bound. The C. elegans TFG C terminus (amino acids 196–486) fused to the streptavidin-binding peptide was incubated with these regions and subsequently
exposed to liposomes containing rhodamine-PE, which were coated with either Sar1GTP and Sec23 or Sar1GTP alone. Surfaces were inverted onto glass cov-
erslips and imaged using confocal microscopy (n = 3). Arrows indicate TFG clusters that recruit COPII-coated liposomes. (Scale bar, 25 μm.)
COP II coat dissociation but local retention of inner COP II-coated transport carriers at the ER/ERGIC interface.

Discussion

In general, current models depicting the ER/ERGIC interface fail to account for the spatial and temporal relationships between COP II coat disassembly and the local retention of transport carriers before target membrane fusion (14, 30, 34, 47, 60). Based largely on work conducted in yeast, which lack ERGIC membranes, a stepwise transport mechanism has been proposed that relies on sequential interactions between the inner COP II coat protein Sec23 and three distinct binding partners: Sar1, TRAPP3, and casein kinase 16 (14, 61). However, precisely how transport carriers traverse the 300- to 500-nm distance between the ER and ERGIC has remained unclear (43). Our data support a concerted tethering model in which TFG promotes outer COP II coat dissociation after transport carriers leave the ER. Simultaneously, TFG functions to retain inner COP II-coated carriers at the ER/ERGIC interface. By concentrating transport carriers with only partial coats, TFG generates a platform that facilitates TRAPP complex-mediated homotypic fusion and nascent ERGIC formation (Fig. 8).

The minimal COP II machinery required to generate cargo-laden, ER-derived transport carriers has been defined for several years, but regulators of this pathway continue to emerge, some of which are expressed only in metazoan organisms, which have evolved more complex systems to control cargo efflux in response to environmental and developmental cues (62, 63). Biochemical and genetic screens in human cells, Drosophila, and C. elegans have identified several factors that play important roles at the ER in cargo selection, exit site organization, and modulating secretory capacity, but components that facilitate movement of COP II-coated carriers between the ER and ERGIC have been more challenging to define (55, 64–67). Microtubules and other cytoskeletal elements are absent within this region, but, nevertheless, most COP II-coated carriers cluster within the ER/ERGIC interface, suggesting the presence of an underlying meshwork that links the organelles to form an integrated secretory unit (44, 68).

In Drosophila, Tango1 plays an integral role in organizing the early secretory pathway by forming micron-sized ring structures that encircle COP II-coated carriers and the cis-Golgi (55). Upon Tango1 inhibition, exit sites on the ER become uncoupled from the Golgi, and both conventional and unconventional (large cargo) secretion are negatively impacted. Depletion of TFG in C. elegans similarly blocks general secretion and uncouples exit sites on the ER from ERGIC membranes in human cells (44, 45). Notably, whereas mammals express both Tango1 and TFG, Drosophila lack a TFG ortholog, and C. elegans lack Tango1-like receptors, suggesting that these components evolved separately to regulate early secretory pathway organization in unique ways (69). Consistent with this idea, Tango1 and TFG are spatially restricted from one another in human cells (Fig. 5C). Current evidence suggests that Tango1 acts early to organize exit sites of the ER and enable procollagen export (40, 70, 71), whereas TFG functions to initiate dissociation of the outer COP II coat and restrict diffusion of inner COP II-coated carriers at the ER/ERGIC interface.

A recent study suggested that TFG acts specifically to regulate the transport of large cargoes (47). However, our findings are inconsistent with this idea. In particular, it is difficult to reconcile how the association between TFG and Sec23 would be restricted to large transport carriers. Additionally, early embryo lethality observed upon TFG knockout in rodents is inconsistent with the more subtle impacts of deleting receptors known to regulate the secretion of large cargoes (53, 55). In contrast to our studies, the prior report failed to use immunofluorescence to determine the degree to which TFG was depleted in cells, raising the distinct possibility that TFG was only partially reduced (47). In this scenario, the specific impact on collagen secretion may have been indirectly caused by disruption of exit site organization as a result of long-term partial depletion of TFG, which could disproportionately affect the export of large cargoes over others.

The ability of TFG to tether inner COP II-coated carriers is highly reminiscent of the functions proposed for synapsin and synuclein in synaptic vesicle clustering (72–76). All three proteins self-associate and harbor disordered regions composed largely of proline, glutamine, glycine, and serine, which form relatively weak but multivalent interactions with vesicle-associated proteins (77–79). In particular, TFG assembles into octameric ring structures (44), with intrinsically disordered C-terminal domains that interact directly with Sec23 on COP II carriers. By tethering neighboring transport carriers, TFG may facilitate a liquid-phase separation at the ER/ERGIC interface (80). Consistent with this idea, blocking COP II transport disrupts the ability of TFG to assemble at exit sites on the ER. Through phase separation, TFG would establish a nonmembranous compartment, which promotes the tight clustering of transport carriers. Synaptic vesicles are readily exchanged in and out of liquid-phase clusters, allowing unimpeded fusion with the plasma membrane when appropriately stimulated (81). Similarly, upon inner coat disassembly, COP II transport carriers would escape the TFG liquid phase and fuse homotypically or heterotypically with juxtaposed ERGIC membranes, potentially in a manner dependent on the TRAPP complex (30). Based on previous work, TRAPP3 and casein kinase 16 outcompete Sar1 for Sec23 binding, which may...
promote inner COPII coat disassembly in vivo and expose SNAREs necessary for membrane fusion (14).

Materials and Methods
All rodent experiments were conducted in the Sprague-Dawley background (Envigo) and were conducted after approval by the University of Wisconsin-Madison Institutional Animal Care and Use Committee.

Recombinant Protein Expression and Purification and Yeast Two-Hybrid Studies. Recombinant proteins were purified following expression in BL21 (E. coli T7, Sigma) (82) or baculovirus-mediated expression in Sf9 cells (83). Protein purification was conducted using glutathione agarose beads (for GST fusion) or nickel- affinity resin (for polyhistidine-tagged proteins), followed by ion exchange chromatography and gel filtration. For Sar1 isoforms, the GST tag was removed using PreScission protease in the presence of 500 μM GDP overnight to facilitate nucleotide exchange. Expression and purification of other COPII proteins and TFG were carried out as described previously (44).

For yeast two-hybrid studies, all bait and prey constructs were sequenced verified. Yeast transformed with the indicated bait and target plasmids were grown in medium lacking uracil and leucine overnight at 30°C with shaking, diluted to an OD600 of 0.25 in medium lacking amino acids, and spotted onto selective medium. Plates were incubated at 30°C for 48 h and then were imaged.

Synthetic Liposome Generation, Immunoblotting, and Immunoprecipitations. Liposomes were generated as described previously (84); detailed composition can be found in SI Materials and Methods. Immunoblotting of extracts was performed as described previously (85) using antibodies described in SI Materials and Methods. For immunoprecipitation studies, antibodies were covalently linked to protein A beads (Bio-Rad). Rat liver cytosol (25 mg protein) ~1 mL or cell-line extracts were added to the beads and incubated at 4°C for 2 h. The beads were washed, and bound proteins were eluted in 100 mM glycine (pH 2.6). Following TCA precipitation, samples were examined by solution mass spectrometry or were separated by SDS/PAGE for immunoblot analysis (45).

CRISPR-Mediated Genome Editing, TALEN Expression System, and siRNA-Mediated Depletion. A guide RNA (gRNA) (a gift from David Ginsburg, University of Michigan, Ann Arbor, MI) directed to exon 1 of Sec23B together with Cas9-GFP were transfected into RPE1 cells using FuGENE HD (Promega). Cells were single-cell sorted using FACS, and clones were examined by immunoblot analysis. Sanger sequencing was used to confirm the presence of mutations that led to frameshift and truncation of Sec23B. To generate clonal doxycycline-inducible cell lines to express TFG transgenes, we used a TALEN-mediated targeting system, which incorporated the transgenes at the AAVS1 locus (86). A puromycin-resistant donor cassette encoding various forms of TFG followed by an IRES and a cDNA encoding RFP was used in all cases. Depletion studies were carried out using siRNAs described in SI Materials and Methods.

Immunofluorescence, Imaging Studies, and Antibodies. For all immunofluorescence experiments, RPE1 cells were grown as described previously (44) and were processed as detailed in SI Materials and Methods. Confocal imaging was conducted on a Nikon Eclipse TiE swept-field confocal microscope using a 60× objective and a CoolSNAP HQ2 CCD camera. Acquisition parameters were controlled by Nikon Elements software, and image analysis was conducted using IMaris software. Immunofluorescence of fixed cells was performed as described previously (87). Briefly, 20–30 Z sections at 0.2-μm steps were acquired (depending on sample thickness). Colocalization and signal intensity analysis were performed using IMaris software. Specified areas of cells located distally from the perinuclear Golgi were selected, and thresholds for size and intensity were set for each channel respectively, as described in SI Materials and Methods. STED microscopy was performed on a Leica TCS SP8 3× superresolution microscope. Images were collected using a 775-nm depletion laser for all dyes used (Alexa 555, Alexa 594, and Alexa 660) and were processed using Huygens deconvolution software. The FWHM of the STED point-spread function was determined to be 90 nm, based on the imaging of diffraction-limited fluorescent beads.

For all doxycycline-inducible expression studies, wild-type or the indicated TALEN-edited RPE1 cells were plated on glass coverslips and treated with doxycycline (3 ng/mL) and either control or TFG 3′ UTR siRNA (10 pmol per well). For all Sec23A-knockdown experiments, either wild-type or Sec23B-knockout cell lines were plated on glass coverslips at 7.5% confluency and were treated with the appropriate Sec23 siRNAs (10 pmol total per well) for 96 h. C. elegans strains used in this study were derived from the Bristol strain N2. With the exception of the strain expressing HMR-1:GFP from its endogenous locus (88), all transgenic animals were created by biolistic transfection (89), and the transgenes were under the control of PIE1 regulatory sequences to enable germline expression (90, 91).

GST Pull-Down Experiments, COPII Budding Reactions, and Liposome-Tethering Assays. Fifty micrograms of the indicated GST-fusion protein was incubated with 50 μL of fresh glutathione resin in 50 mM Heps (pH 7.6), 100 mM NaCl, 1 mM EDTA, and 1 mM DTT. After a 15-min incubation, the resin was washed and incubated with 1.5 mg of protein from cleared rat liver cytosol, prepared as described previously (92), at 4°C for 1 h. After extensive washing, bound proteins were eluted in 1× PBS (pH 8.0) and 30 mM reduced glutathione to a final volume of 900 μL and were precipitated with TCA. The precipitate was resuspended in sample buffer and separated by SDS/PAGE. All GST competition experiments were conducted as described previously (40) and are detailed in SI Materials and Methods.

In vitro budding reactions to analyze the formation of large transport carriers harboring procollagen were conducted as described previously (49). Light liposomes for tethering reactions were extruded through a 0.05-μm filter and were clarified by centrifugation at 16,163 × g for 15 min. Heavy liposomes were reconstituted with Sar1 buffer supplemented with 0.75 μM sucrose and were extruded through a 0.4-μm filter. Liposome-tethering assays and liposome-aggregation experiments (determined by measuring absorbance at 405 nm) were based on previous work (57, 53) and are described in SI Materials and Methods.

ACKNOWLEDGMENTS. We thank Alma Seito and Ashley Hutchinson for technical support in baculovirus expression, Pietro De Camilli for useful discussions, and David Melville for help with technical aspects of the COPII budding reaction. This work was supported in part by NIH Grants GM110576 (to J.R.Y.), P41 GM103533 (to L.R.V.), American Cancer Society Grant 123268-RSG-12-139-01-CSM (to A.A.), Brain Research Foundation Grant BRFSG-2015-03 (to A.A.), the University of Wisconsin Carbone Cancer Center Grant P30 CA04520, and the University of Wisconsin Institute for Clinical and Translational Research Grant UL1TR000427. The Structural Genomics Consortium (SGC) is a registered charity (no. 1097737) that receives funds from numerous sources (www.thesgc.org/about/partners).


