Haploinsufficiency of the Sec7 Guanine Nucleotide Exchange Factor Gea1 Impairs Septation in Fission Yeast

Alan M. Eckler1*, Caroline Wilder1*, Antonio Castanon1, Veronica M. Ferris1, Rachael A. Lamere1, Benjamin A. Perrin1, Ross Pearlman1, Blaise White1, Clifton Byrd1, Nicholas Ludvik1, Nona Nichols1, Kristen Poole-Sumrall1, Elizabeth Sztui2, Melanie L. Styers1*

1 Department of Biology, Birmingham-Southern College, Birmingham, Alabama, United States of America, 2 Department of Cell, Developmental, and Integrative Biology, University of Alabama at Birmingham, Birmingham, Alabama, United States of America

Abstract

Membrane trafficking is essential to eukaryotic life and is controlled by a complex network of proteins that regulate movement of proteins and lipids between organelles. The GBF1/GEA family of Guanine nucleotide Exchange Factors (GEFs) regulates trafficking between the endoplasmic reticulum and Golgi by catalyzing the exchange of GDP for GTP on ADP Ribosylation Factors (Arfs). Activated Arfs recruit coat protein complex 1 (COP-I) to form vesicles that ferry cargo between these organelles. To further explore the function of the GBF1/GEA family, we have characterized a fission yeast mutant lacking one copy of the essential gene gea1 (gea1+/−), the Schizosaccharomyces pombe ortholog of GBF1. The haploinsufficient gea1+/− strain was shown to be sensitive to the GBF1 inhibitor brefeldin A (BFA) and was rescued from BFA sensitivity by gea1p overexpression. No overt defects in localization of arf1p or arf6p were observed in gea1+/− cells, but the fission yeast homolog of the COP-I cargo sac1 was mislocalized, consistent with impaired COP-I trafficking. Although Golgi morphology appeared normal, a slight increase in vacuolar size was observed in the gea1+/− mutant strain. Importantly, gea1+/− cells exhibited dramatic cytokinesis-related defects, including disorganized contractile rings, an increased septation index, and alterations in septum morphology. Septation defects appear to result from altered secretion of enzymes required for septum dynamics, as decreased secretion of eng1p, a β-glucanase required for septum breakdown, was observed in gea1+/− cells, and overexpression of eng1p suppressed the increased septation phenotype. These observations implicate gea1 in regulation of septum breakdown and establish S. pombe as a model system to explore GBF1/GEA function in cytokinesis.

Introduction

Membrane trafficking and protein secretion are essential for maintaining cellular life and underlie many fundamental cellular processes, including cell signaling, nutrient uptake, waste processing, and deposition of the extracellular matrix [1–7]. Membrane trafficking collectively refers to the vesicle-mediated movement of proteins and lipids between different cellular membranes [8,9]. As all membrane and secreted proteins are synthesized in the rough endoplasmic reticulum (ER), proper sorting and transport of these proteins is necessary to ensure that they reach the appropriate destinations for their functions [10]. Hence, cellular life has evolved to develop complex machinery to regulate protein sorting and formation of transport vesicles that carry membrane and secreted proteins throughout the cell.

Vesicle formation within the secretory pathway is regulated by ADP-Ribosylation Factors (Arfs) [11–14], small GTPases that oscillate between an active, GTP-bound form and an inactive, GDP-bound form [15–17]. Activated Arfs recruit coat proteins to ERGIC (ER-Golgi intermediate compartment), Golgi, and post-Golgi membranes [18–22]. These coat proteins drive vesicle formation and promote selection and packaging of the appropriate cargoes into vesicles [23]. Thus, Arf activation drives the formation of transport vesicles that deliver cargo proteins to target membranes. Arf activation is tightly regulated through the action of Guanine nucleotide Exchange Factors (GEFs) and GTPase Activating Proteins (GAPs). GEFs catalyze the exchange of GDP for GTP on Arfs to promote Arf activation [24,25], whereas GAPs inactivate Arfs through activation of their intrinsic GTPase activity [26,27].

Arf activation is catalyzed by the Sec7 family of Arf GEFs, named after their founding member S. cerevisiae SEC7 [28–30]. All members of this family possess a highly-conserved Sec7 domain, which catalyzes GDP/GTP exchange on Arfs [30,31]. Although 9 distinct classes of Sec7 family Arf GEFs have been identified based on phylogeny [28], only two subfamilies have been shown to be inhibited by the fungal metabolite Brefeldin A (BFA), a well-characterized inhibitor of protein secretion [25,31–36]. Of these two families, the GBF1/GEA family has been shown to localize to early secretory compartments, including the ERGIC and Golgi [33,37], while the SEC7/BIG family has been shown to localize to...
both secretory and endocytic compartments, including the trans-Golgi network (TGN) and endosomes [38–42].

Characterization of GBF1/GEA family members has provided significant insight into the physiological roles of this family. The GBF1/GEA family has been shown to be required for membrane recruitment of coat protein complex-I (COP-I) in budding yeast and mammalian cells [33,37,43,44]. COP-I facilitates retrograde traffic from the Golgi to the ER, as well as anterograde traffic from the ERGIC to the cis-Golgi [45–49]. *H. sapiens* GBF1 has been shown to activate ARF1, ARF4, and ARF5 and to reside in the ERGIC and Golgi [50,51]. In mammalian cells, siRNA-mediated depletion of GBF1 or expression of the GBF1 dominant-negative mutant E794K results in tubulation or fragmentation of the Golgi and ERGIC and inhibition of protein secretion [33,39,50]. GBF1 has also been implicated in post-Golgi trafficking through interactions with the Golgi-localized, gamma-ear-containing, Arf-binding (GGA) coat proteins [52]. In *Saccharomyces cerevisiae*, loss or mutation of the GBF1 family members *gea1* and *gea2* result in defects in ER-Golgi and intra-Golgi transport, alterations in actin morphology, and impaired autophagy [37,43,53,54]. Mutations in the *Drosophila melanogaster* GBF1 homolog *Gartenzweig* (Gar) results in defects in tubulogenesis as a result of impaired formation of polarized epithelia, implicating the GBF1/GEA family in maintenance of cellular polarity [53]. Together, these observations indicate that GBF1 plays important roles in ERGIC and Golgi homeostasis, as well as potential roles in cellular polarization.

In addition to its roles in organellar trafficking, GBF1/GEA family members have also been implicated in regulation of the cell cycle. Depletion of mammalian GBF1 results in cell cycle arrest at the G0/G1 phase and is thought to be associated with the unfolded protein response, ultimately inducing apoptosis [56]. GBF1 activity is also regulated in a cell cycle-specific manner through phosphorylation by the cyclin B–cyclin-dependent kinase 1 (CDK1) complex [57]. Loss of *gea1* and *gea2* activity in *S. cerevisiae* causes defects in polarity of the actin cytoskeleton and budding at 37°C, resulting in the formation of multiple buds [53]. However, despite these observations, the precise mechanisms that underlie the role of GBF1/GEA family members in regulation of the cell cycle remain largely unexplored.

The goal of this study was to characterize the function of *gea1*, the fission yeast ortholog of GBF1. Because deletion of *S. pombe* *gea1* gene is lethal, the present study was performed using the haploinsufficient heterozygote strain Gea1+/- (see Methods). Because the GBF1/GEA family has been shown to be required for membrane recruitment of COP-I, we expected that GBF1 would be involved in the regulation of COP-I recruitment to the Golgi, as well as in the regulation of the cell cycle. However, we found that overexpression of *gea1* resulted in the formation of multiple buds [53], suggesting that GBF1 has a role in the regulation of cell cycle progression. To further investigate the role of GBF1 in cell cycle progression, we performed a reverse-transcriptase-PCR (RT-PCR) analysis of cell cycle-related genes.

Reverse-transcriptase-PCR (RT-PCR)

RT-PCR was performed to analyze levels of the *gea1* mRNA in wild-type and *gea1*+/- cells. RNA was purified from yeast cells using the RNEasy Mini kit in combination with the RNase-Free DNase set from Qiagen (Valencia, CA). RT-PCR was conducted using the Qiagen OneStep RT-PCR Kit according to the manufacturer's instructions. Briefly, reactions contained 1X RT-PCR buffer, 0.6 μM *gea1*-specific or 28S rRNA-specific primers (see Table 2), 500 ng total RNA, 0.4 mM dNTPs, 1X Q-solution, and RT-PCR enzyme mix. Reactions were incubated in a Biometra T3 Thermocycler under the following conditions: 1 cycle of 50°C for 35 min; 1 cycle of 95°C for 15 min; 30 cycles of 94°C for 45 sec, 55°C for 45 sec, and 72°C for 1 min; followed by a final extension at 72°C for 10 min.

The aforementioned plasmids and PCR products were transformed into wild-type and *gea1*+/- yeast using the lithium acetate method, as previously described [62]. Transformants were selected on EMM containing appropriate supplements. Transformants expressing *gea1*-GFP were confirmed by PCR analysis using the primer pairs listed in Table 2 (*eng1*-CasF and CassIntR; CassIntF and *eng1*-CasR).

Materials and Methods

Strains and growth conditions

A list of strains used in this study is shown in Table 1. All strains were derived from the sp286 wild-type strain and the isogenic *geo1+/−* strain, which were purchased from Bioneer (Alameda, CA). Cells were cultured in Yeast Extract plus Supplements (YES; MP Biomedical, Solon, OH) or Edinburgh Minimal Media (EMM; MP Biomedical) containing appropriate nutritional supplements. Deletion mutants were selected by growth on YES media containing 200 μg/mL G418. Cells were cultured at 30°C unless otherwise indicated.

Plasmids and DNA manipulations

The pFA6A-GFP-ura4MX6 plasmid was a kind gift from Eishi Noguchi (Drexel University College of Medicine, Philadelphia, PA). The pREP4X and pREP4X-engl plasmids, which express eng1p under control of the *urn1*+ promoter, were a kind gift from Carlos Vazquez de Aldana (Universidad de Salamanca, Salamanca, Spain). The pDUAL-YFH1-Gea1 vector and pDUAL-YFH1-Gea2 plasmids expressing gea1-YFP, arfl-YFP, arfl6-YFP, sac11-YFP, *SPBC1.95.03*, and sac12-YFP (SPBC3C7.01c) under control of the full-strength *urn1*+ promoter were purchased from the Riken Biosource Center DNA Bank (Ibaraki, Japan, deposited by M. Yoshida [58–60]).

The polymerase chain reaction (PCR) was used to amplify DNA fragments containing the GFP-ura4 cassette from pFA6A-GFP-ura4MX6 as previously described [61,62]. Primers containing regions of the *gea1* (gea1-GFPi and gea1-GFPc) and *engl* (engl-GFPi and engl-GFPc) genes are listed in Table 2. Primers for gea1 were described previously [63]. PCR reactions contained 1X Phusion® GC Buffer, 1 nM primers, the pFA6A-GFP-ura4MX6 template, 0.4 mM dNTPs, and Phusion® polymerase (Thermo Fisher Scientific, Inc., Waltham, MA). Reactions were incubated in a Biometra T3 Thermocycler under the following conditions: 1 cycle of 98°C for 1 min; 30 cycles of 98°C for 10 sec, 60°C for 15 sec, and 72°C for 2 min; followed by a final extension at 72°C for 10 min.

The aforementioned plasmids and PCR products were transformed into wild-type and *gea1*+/- yeast using the lithium acetate method, as previously described [62]. Transformants were selected on EMM containing appropriate supplements. Transformants expressing *eng1*-GFP were confirmed by PCR analysis using the primer pairs listed in Table 2 (*eng1*-CasF and CassIntR; CassIntF and *eng1*-CasR).

Reverse-transcriptase-PCR (RT-PCR)

RT-PCR was performed to analyze levels of the *gea1* mRNA in wild-type and *gea1*+/- cells. RNA was purified from yeast cells using the RNEasy Mini kit in combination with the RNase-Free DNase set from Qiagen (Valencia, CA). RT-PCR was conducted using the Qiagen OneStep RT-PCR Kit according to the manufacturer’s instructions. Briefly, reactions contained 1X RT-PCR buffer, 0.6 μM *gea1*-specific or 28S rRNA-specific primers (see Table 2), 500 ng total RNA, 0.4 mM dNTPs, 1X Q-solution, and RT-PCR enzyme mix. Reactions were incubated in a Biometra T3 Thermocycler under the following conditions: 1 cycle of 50°C for 35 min; 1 cycle of 95°C for 15 min; 30 cycles of 94°C for 45 sec, 55°C for 45 sec, and 72°C for 1 min; followed by a final extension at 72°C for 10 min. The resulting products were resolved on 0.4% agarose gels and ethidium bromide-stained.
bands were visualized using a BioDocIt system (UVP, LLC; Upland, CA). Band intensities were quantified using Image J (NIH, Bethesda, MD). The intensities of gea1 bands were normalized to the corresponding 28S rRNA bands. Results are reported as percent of the wild-type sample.

### Spot assays

Wild-type cells, gea1+/− cells, and gea1+/− cells transformed with pDUAL-YFH1c-gea1 were cultured overnight in YES media, and the density of the resulting cultures was measured by monitoring the absorbance at 600 nm. Equivalent numbers of each cell type were subjected to a 10-fold serial dilution, and 5 μL of each concentration was spotted on plates containing the following media: YES, YES containing 2 mM valproic acid (VPA). The plates were then incubated at 25°C, 30°C, or 37°C for 3–7 days prior to imaging.

### BFA dose-response assay

To quantify BFA sensitivity of wild-type cells, gea1+/− cells, and gea1+/− cells transformed with pDUAL-YFH1c-gea1 or the corresponding empty vector, overnight cultures of each strain were diluted to a final concentration of 5×10^5 cells/mL in fresh YES media and incubated at 30°C for 20 minutes. The samples were then washed once with ice-cold medium, followed by resuspension in fresh YES media containing 32 μM FM4-64 (Invitrogen Molecular Probes). The cells were then incubated for 30 min at 30°C. The samples were then washed three times with ice-cold medium, followed by incubation in fresh medium at 30°C for 30 min. The samples were then imaged as described below.

### FM4-64 uptake and vacuolar fusion

Overnight cultures of wild-type and gea1+/− cells were resuspended in fresh YES media containing 32 μM FM4-64 (Invitrogen Molecular Probes). The cells were then incubated at 30°C for 20 minutes. The cells were then washed and resuspended in fresh YES media and incubated at 30°C for 30 min (for FM4-64 staining) or in dH2O for 90 min (to assay vacuolar fusion) as previously described [64,65]. Following this incubation, the cells were washed once with H2O, followed by resuspension in EMM

---

### Table 1. Yeast strains used in this study.

| Strain name | Genotype | Source |
|-------------|----------|--------|
| Wild-type   | ade6-M210/ade6-M216 ura4-D18/ura4-D18 leu1-32/leu1-32 h+/h+ | Bioneer |
| gea1+/−     | ade1-A::kanMX4/geo1 ade6-M210/ade6-M216 ura4-D18/ura4-D18 leu1-32/leu1-32 h+/h+ | Bioneer |
| Wild-type + gea1-GFP | ade6-M210/ade6-M216 ura4-D18/ura4-D18 leu1-32/leu1-32 h+/h+ gea1-GFP::ura4 | This study |
| gea1+/− + gea1-GFP | ade1-A::kanMX4/geo1 ade6-M210/ade6-M216 ura4-D18/ura4-D18 leu1-32/leu1-32 h+/h+ gea1-GFP::ura4 | This study |
| Wild-type + eng1-GFP | ade1-A::kanMX4/geo1 ade6-M210/ade6-M216 ura4-D18/ura4-D18 leu1-32/leu1-32 h+/h+ eng1-GFP::ura4 | This study |
| gea1+/− + eng1-GFP | ade1-A::kanMX4/geo1 ade6-M210/ade6-M216 ura4-D18/ura4-D18 leu1-32/leu1-32 h+/h+ eng1-GFP::ura4 | This study |

---

### Table 2. Primers used in this study.

| Primer name | Sequence (5′→3′) |
|-------------|------------------|
| gea1 forward | TGCCGAGAGCATGACACTGAGC |
| gea1 reverse | CCAACAGGCGCAACCTTTCGGC |
| 28S forward | TGAGAAGGGATGTTGGACCTGCTT |
| 28S reverse | ATTCGGCTCACAACCCCTTTCGGC |
| gea1-GFPF | GACTTTAATATATCAACAAACGAGGCCAATAAGAACAAACACTAATAAATCTCCTCACAACCGACTGTCGGATCCCGCGTTAAATTA |
| gea1-GFPr | CAATGAGCATATATGAAATAATGATGCTCCCTTAAATCCCTCTACAAGAAGAAATAAGAGGAAGAGAGAAAAATGGAAGAGGGAAAAATGGAAGGGAATAAAAGTCCCGCCCGGTTAATTA |
| eng1-GFPF | GCTTGTGGTATAGTGCTCTATGACTCTTCTTATACCTACGGTGTTGGTGCTGTGCTGCGATCCCGGGTTAATTA |
| eng1-GFPr | TATCCCCAAAGGGGCTTCAAGTTGAGTGCTGACGTTCCAGCCACAGGTTGTAATATGATGAGAGGTAATAAAAGTCCCGCCCGGTTAATTA |
| eng1-cassF | ACTGCAACCGAGCTTGTAT |
| CassIntR | GCATCACCTTCAACCCCTTCC |
| CassIntF | TCACCAGTCGCAAAATATACACA |
| eng1-cassR | AGTCTAAAGGTTTCACATCCAGTGT |

---

**DOI**: 10.1371/journal.pone.0056807.t001

**DOI**: 10.1371/journal.pone.0056807.t002
for imaging. Imaging and quantification of vacuolar size was performed as described below.

**Alexa 568-phalloidin/DAPI staining**

Alexa 568-phalloidin staining was performed as previously described [66]. Briefly, wild-type and *gea1+/−* cells were fixed by incubating in pre-warmed 4% paraformaldehyde at 30°C for 1 h. The cells were then washed three times in PEM buffer (0.1 M Na Pipes, pH 6.8, 1 mM EGTA, 1 mM MgCl₂), followed by extraction with 1% Triton X-100 in PEM for 30 s. After 3 additional washes with PEM, the cells were stained with 0.06 units/μL Alexa 568-phalloidin in PEM for 30 min at 37°C. The cells were then washed three times with PEM and resuspended in PEM containing 1 μg/mL DAPI (4',6-diamidino-2-phenylindole; Sigma Aldrich, St. Louis, MO). Samples were imaged by fluorescence microscopy, as described below.

**Calcofluor white staining**

To fix cells for Alexa 568-phalloidin staining, cells from overnight cultures in YES media were resuspended in 4% Paraformaldehyde (PFA) in phosphate-buffered saline (PBS) and incubated for 10 min at 30°C. The cells were then washed two times with PBS prior to further staining. Fixed wild-type and *gea1+/−* cells were then resuspended in mounting media (50% glycerol in PBS) containing 50 μg/mL calcofluor white (Sigma-Aldrich; St. Louis, MO) and incubated for 10 min at 30°C. The cells were washed with PBS and resuspended in mounting media prior to imaging and quantification as described below.

**Fluorescence microscopy and quantification**

Single color fluorescence images or dual actin/DAPI images were captured using a Zeiss Axioskop 2 fluorescence microscope. Dual color images of YFP and BODIPY® TR C₃- ceramide were captured using a Perkin Elmer ERS 6FE spinning disk confocal microscope, and images were processed and analyzed in Volocity version 5.3 software (Perkin Elmer, Shelton, CT). Imaging experiments were repeated a minimum of three times using independent cultures. For each experiment, 10–15 images were captured per slide. Both fluorescence and differential interference contrast (DIC) or phase-contrast images were captured for comparison.

For quantification of sac11-YFP and sac12-YFP localization, cells were manually scored as having Golgi localization (punctate distribution throughout the cytoplasm), ER localization (cell cortex and nuclear envelope), or mixed.

Vacuole size was quantified using Image J. Briefly, the pixel area of the largest vacuole in each cell was measured using the software program. The number of cells with vacuolar areas encompassed by each of the stated thresholds was analyzed. The percentage of cells with a given area compared to the total number of cells was then calculated for each threshold and plotted using SigmaPlot (Systat Software, Inc.; San Jose, CA).

For cell size measurements, phase-contrast images were captured from three independent wild-type and *gea1+/−* cultures, as described above. Cell size was analyzed by comparing the length of each cell to that of a micrometer. Results are reported as the percentage of wild-type control cells.

For septum quantification, the number of septated cells was calculated as a percentage of the total number of cells. Each septated cell was scored as normal or abnormal. Cells with abnormal septa included those that contained septa that were not positioned in the center of the cell, those that had septa that were not perpendicular to the length of the cell, those that had misshapen septa, such as forked or abnormally thick septa, and those that contained multiple septa.

To quantify *eng1-GFP* localization, wild-type and *gea1+/−* cells carrying an integrated GFP cassette at the endogenous *eng1* locus were cultured in EMM media lacking uracil. Images (n = 30 per culture) were captured and processed using identical settings. Septated cells were identified on DIC images and were scored based on whether *eng1-GFP* was visible at the septum.

**Transmission electron microscopy**

Wild-type and *gea1+/−* cells were prepared for transmission electron microscopy as described in [67]. Briefly, cells were washed three times with dH₂O, prior to fixation in 2.5% glutaraldehyde in 0.1 M phosphate buffer, pH 7.2, for 2 h at 4°C. The fixed cells were then washed in 0.1 M phosphate buffer, pH 7.2, prior to post-fixation in 3% potassium permanganate at RT for 90 min. Cells were then washed with distilled water at RT and stained with 2% uranyl acetate in distilled water at 4°C for 30 min. The cells were dehydrated with alcohols in a graduated series and embedded in Epon 812 resin. Ultrathin sections were stained with uranyl acetate and lead citrate and were then observed with an FEI-12 Spirit transmission electron microscope (FEI; Hillsboro, Oregon) at an accelerating voltage of 80 kV.

**1,3-β-Glucanase activity assay**

β-glucanase activity was measured as described below in a manner similar to that previously described [68].

**Collection of media and preparation of cell lysates**

Briefly, a volume of overnight cell culture in YES media containing 1×10⁶ wild-type or *gea1+/−* cells was collected and subjected to centrifugation at 1000× g for 5 min. The resulting supernatant was collected, and secreted proteins were concentrated using a centrifugal filter unit with a molecular weight cut-off (MWCO) of 30000 Da (Millipore; Billerica, MA), which was subjected to centrifugation at 4000× g for 5 min at 4°C. The remaining solution was then dialyzed against 50 mM acetate buffer, pH 5.5, using a Slidealay dialysis cassette (MWCO, 2000 Da) overnight at 4°C. The volumes of the resulting samples were then equalized. The pelleted cells were resuspended in 50 mM acetate buffer, pH 5.5, and lysed by vortexing in the presence of glass beads (0.5 mm) for 5 minutes at 4°C.

**1,3-β-Glucanase Activity Assay.** Samples containing secreted proteins and cell lysates were incubated with 0.25% laminarin [β-(1,3)-glucan] in 50 mM acetate buffer, pH 5.5, for 1 h at 30°C. The amount of reducing sugars released by the reaction was then assayed using the Somogyi-Nelson method as previously described [69,70].

**Quantitative analysis of acid phosphatase secretion**

Secretion of acid phosphatase activity was assayed as previously described with slight modifications [67]. Briefly, equal numbers of wild-type and *gea1+/−* cells cultured in YES media were washed and diluted in 10 mL EMM with or without 40 μg/mL BFA and incubated with agitation at 30°C. At 0 h, 1 h, 2 h, 3 h, and 4 h, the OD₆⁰₀ of the culture was measured to estimate the number of cells, and a sample was collected for acid phosphatase secretion analysis. The sample was immediately subjected to centrifugation at 25000× g for 1 min, and the supernatant, containing secreted acid phosphatase, was stored at 4°C until all samples were collected. Acid phosphatase activity was then assayed by incubating each sample with an equal volume of substrate solution (2 mM p-nitrophenyl phosphate, 0.1 M sodium acetate, pH 4.0) for 5 min at 30°C. The reaction was stopped by the addition of a volume of 1M NaOH. Phosphatase activity was then quantified by
measuring the absorbance of each reaction at 405 nm. Phosphatase activity (OD405) was normalized to cell density (OD600) to control for differences in cell growth.

Western blot analysis
Wild-type and gea1+/− cells were harvested from 50 mL cultures by centrifugation and washed with dH2O. The cells were then resuspended in modified TEG buffer (40 mM Tris-HCl, pH 7.5; 1 mM EDTA; 10% glycerol; 0.1% NP-40; 150 mM NaCl) containing 1 mM PMSF and 1X Complete EDTA-free protease inhibitors (Roche; Basel, Switzerland). The cells were lysed by repeated agitation with acid-washed glass beads (Sigma Aldrich). Approximately 50 μg of each cell lysate was then resolved on a 4–20% SDS-PAGE gel (Biorad; Hercules, CA). The proteins were transferred to a polyvinylidene fluoride (PVDF) membrane (Millipore), which was blocked in 5% milk in Tris-buffered saline containing 0.1% Tween-20 (TBST). The membrane was probed with the following primary antibodies: rabbit anti-GFP (Ab2920; Abcam; Cambridge, MA; 1:2000) and mouse anti-β-actin (mAbcam 8224; Abcam; 1:600). After washing with TBST, the membrane was incubated with Goat anti-mouse horse radish peroxidase (HRP) or Goat anti-rabbit HRP secondary antibodies (Sigma, 1:10000). Bound antibodies were detected using enhanced chemiluminescence Western blotting substrate (Pierce; Rockford, IL). Band intensity was quantified using Image J, and the intensity of the GFP bands was normalized to that of β-actin to control for equal loading.

Bioinformatic and statistical analyses
Bioinformatic analyses were performed in the Biology Workbench 3.2 (San Diego Supercomputer Center, San Diego, CA). Homologs of H. sapiens GBF1 were identified using the BLASTp function, searching relevant databases. Alignments were performed using the CLUSTALW algorithm.

All statistical analyses were performed on a minimum of three independent experiments. Results are reported as mean ± standard deviation (SD) or mean ± standard error of the mean (SEM), as indicated. Significant differences were analyzed using the Student’s t-test, and p-values less than 0.05 were deemed statistically significant.

Results
S. pombe gea1 belongs to the BFA-sensitive GBF1/GEA Arf-GEF family
The GBF1/GEA family of Arf-GEFs is highly conserved, with homologs in all eukaryotes, ranging from plants to fungi to animals (Fig. 1A). Vertebrates, including humans, rats, mice, chickens, and zebrafish, all possess a single ortholog of GBF1. In contrast, the plant Arabidopsis thaliana possesses three GBF1 homologs (GNOM, Gnom-L1, and Gnom-L2). Unlike S. cerevisiae, which possesses two orthologs of GBF1 (gea1 and gea2), S. pombe possesses a single GBF1 ortholog, similar to vertebrates, flies, and nematodes. Previous studies in budding yeast have shown that gea1 and gea2 can functionally compensate for one another [37], complicating functional analyses. Therefore, we chose the fission yeast Schizosaccharomyces pombe as a model to characterize GBF1/GEA function.

As complete loss of gea1 is lethal in S. pombe, we used heterozygote gea1+/− cells as a model to analyze gea1 function. To confirm that loss of one copy of gea1 results in decreased levels of gea1 mRNA, we performed the reverse transcriptase-polymerase chain reaction (RT-PCR) on RNA purified from wild-type and gea1+/− cells using primers specific to gea1 and the 21S rRNA (Fig. 1B, inset). Quantification of RT-PCR results and normalization to expression of the 21S rRNA indicated that expression of the gea1 mRNA in gea1+/− cells was 46% of wild-type (Fig. 1B), demonstrating that loss of one copy of gea1 was sufficient to induce haploinsufficiency in gea1 expression.

Our phylogenetic analysis suggests that gea1 belongs to the GBF1/GEA family of Arf GEFs. Importantly, H. sapiens GBF1 was first identified based on the BFA resistance phenotype observed in cells overexpressing GBF1 [51], and the activity of budding yeast gea1p has been shown to be sensitive to BFA [30]. BFA is a fungal metabolite that stabilizes an abortive complex between a subset of Sec7 family GEFs and ARFs [36]. Treatment of cells with BFA blocks protein secretion, causes Golgi fragmentation and accumulation of Golgi proteins in the ER, and leads to cell death at high concentrations [71]. These effects have been shown to be due to Arf-GEF inhibition [36]. Based on our phylogenetic analysis, we hypothesized that GBF1 and gea1 are orthologs and that BFA treatment would inhibit gea1p activity, resulting in sensitivity to the drug in gea1+/− cells. To test this prediction, wild-type and gea1+/− cells were plated on YES medium containing 10 μg/mL BFA and incubated at 30°C, 25°C, or 37°C for 3–5 days prior to imaging. Gea1+/− cells exhibited BFA sensitivity at all temperatures analyzed (Fig. 1C), indicating that gea1p is a functional member of the GBF1/GEA family. Interestingly, these results also revealed that gea1+/− cells were cold-resistant, suggesting that slower growth at lower temperatures may allow the mutant to overcome or cell cycle defects associated with decreased expression of gea1.

In addition to BFA sensitivity, we also analyzed the sensitivity of gea1+/− cells to the immunosuppressant FK506 and the anti-covulsive agent valproic acid (VPA), drugs previously shown to cause growth sensitivity in a number of S. pombe membrane trafficking mutants [67,72–74]. Gea1+/− cells exhibited slight sensitivity to FK506 and VPA, although FK506 sensitivity was reversed at 25°C (Fig. 1C). These results are consistent with a defect in membrane transport in gea1+/− cells.

To confirm that the BFA sensitivity phenotype was due to decreased expression of gea1p, we created a “rescued” gea1+/− strain by transforming the strain with a vector driving expression of gea1-YFP. Examination of gea1+/− cells with low level overexpression of gea1-YFP by confocal fluorescence microscopy revealed that gea1-YFP was found in small punctate dots in the cytoplasm that colocalized with the Golgi-specific dye BODIPY TR C2-ceramide (Fig. 2A; [75,76]). Some aggregation was observed with higher expression (data not shown). Punctate cytoplasmic localization was also observed for gea1-GFP under control of its endogenous promoter (Figure 2B). This localization pattern is consistent with that observed for the human GBF1 protein, which also localizes to the Golgi [33]. Importantly, overexpression of gea1-YFP restored the growth of gea1+/− cells in the presence of BFA, both in a spot assay (Fig. 2C) and in a quantitative dose-response assay (Fig. 2D). Consistent with this observation, overexpression of gea1-YFP in the wild-type strain resulted in BFA resistance (Fig. 2D), similar to previous studies of GBF1 in mammalian cells [51]. These results confirm that BFA sensitivity of gea1+/− cells is indeed due to loss of gea1 expression and establish gea1p as a homolog of the mammalian GBF1 protein.

Gea1 haploinsufficiency affects COP-I-dependent transport
The GBF1/GEA family of Arf-GEFs has previously been shown to regulate COP-I-mediated trafficking through activation of Arfs [33,37,43,51]. Therefore, we analyzed Arf localization in S. pombe.

PLOS ONE | www.plosone.org 5 February 2013 | Volume 8 | Issue 2 | e56807
Fission yeast possess two arf homologs, termed arf1 and arf6. Sequence analysis has revealed arf1 to belong to the class I/II family of Arfs, which localize to the Golgi and endosomes and exhibit BFA-sensitive activation [33,77]. In contrast, arf6 belongs to the class III Arfs, which localize to the plasma membrane and are not sensitive to BFA [77,78]. Wild-type and gea1+/− cells were transformed with plasmids overexpressing YFP-arf1p or YFP-arf6p. In wild-type cells, Arf1p localized to punctate structures in the cytoplasm (Fig. 3A). These structures exhibited slight overlap with the Golgi-specific dye BODIPY® TR C5-ceramide (Supplemental Figure S1A, arrowheads). Limited colocalization with the Golgi was not unexpected, as class I/II arfs have also been shown to be recruited to the endosomes and trans-Golgi network by the Sec7/BIG family of Arf-GEFs [38,79–81]. In contrast, arf6p localized to the cell surface and the membranes surrounding the septum. No overt changes were observed in arf1p or arf6p localization, although septum structure appeared altered in the gea1+/− cells. However, subtle changes in arf localization, such as partial changes in distribution between the cytoplasm and membrane compartments, could not be excluded.

GBF1 activity is required for recruitment of COP-I to Golgi membranes [33,51]. Therefore, we next tested whether COP-I-dependent trafficking pathways were operational in gea1+/− cells by analyzing localization of sac1 homologs. Sac1 is a lipid phosphatase that exhibits specificity for phosphatidylinositol-4-phosphate (P4P) [49] COP-I has been shown to be required for its retention in the ER, and human sac1 mutants incapable of binding COP-I accumulate in the Golgi due to inhibition of retrograde transport [49]. Sequence analysis revealed two potential homologs for S. pombe sac1, SPBC19F5.03 and SPAC3C7.01c, which we termed sac11 and sac12, respectively. Overexpression of sac11p-YFP in wild-type cells revealed that sac11p was found at the cell cortex and surrounding the nucleus, consistent with ER localization (Fig. 3B). In contrast, sac12p-YFP localized to punctate spots that colocalized with a Golgi marker in wild-type cells (Fig. 3B and Supplemental Figure S1B). These observations suggest that sac11p is orthologous to the COP-I cargo, mammalian sac1. In gea1+/− cells sac11-YFP was found in punctate structures that colocalized with the Golgi marker BODIPY® TR C5-ceramide, while Sac12p localized to the cytoplasm (Fig. 3C). These data suggest that recycling of sac11p from the ER to the Golgi is selectively impaired in gea1+/− cells, consistent with impaired COP-I activity.

Figure 1. S. pombe gea1 belongs to the BFA-sensitive GBF1/GEA Arf-GEF family. A. Protein sequences of GBF1 homologs in Schizosaccharomyces pombe (Sp), Saccharomyces cerevisiae (Sc), Rattus norvegicus (Rn), Mus musculus (Mm), Homo sapiens (Hs), Gallus gallus (Gg) Danio rerio (Dr), Drosophila melanogaster (Dm), Caenorhabditis elegans (Ce), Arabidopsis thaliana (At) and Dictyostelium discoideum (Do) were selected based on sequence homology and aligned. B. Total RNA purified from wild-type and gea1+/− cells was subjected to RT-PCR using primers specific for gea1 and the 28S rRNA (inset). Intensities of the resulting bands were quantified and are presented as the mean ± SD (n = 3). C. Wild-type and gea1+/− S. pombe cells were subjected to 10-fold serial dilution and spotted on YES media, YES and the 28S rRNA (inset). Intensities of the resulting bands were quantified and are presented as the mean ± SD (n = 3). C. Wild-type and gea1+/− S. pombe cells were subjected to RT-PCR using primers specific for gea1 and the 28S rRNA (inset). Intensities of the resulting bands were quantified and are presented as the mean ± SD (n = 3). C. Wild-type and gea1+/− S. pombe cells were subjected to RT-PCR using primers specific for gea1 and the 28S rRNA (inset). Intensities of the resulting bands were quantified and are presented as the mean ± SD (n = 3). C. Wild-type and gea1+/− S. pombe cells were subjected to RT-PCR using primers specific for gea1 and the 28S rRNA (inset). Intensities of the resulting bands were quantified and are presented as the mean ± SD (n = 3). C. Wild-type and gea1+/− S. pombe cells were subjected to RT-PCR using primers specific for gea1 and the 28S rRNA (inset). Intensities of the resulting bands were quantified and are presented as the mean ± SD (n = 3). C. Wild-type and gea1+/− S. pombe cells were subjected to RT-PCR using primers specific for gea1 and the 28S rRNA (inset). Intensities of the resulting bands were quantified and are presented as the mean ± SD (n = 3). C. Wild-type and gea1+/− S. pombe cells were subjected to RT-PCR using primers specific for gea1 and the 28S rRNA (inset). Intensities of the resulting bands were quantified and are presented as the mean ± SD (n = 3). C. Wild-type and gea1+/− S. pombe cells were subjected to RT-PCR using primers specific for gea1 and the 28S rRNA (inset). Intensities of the resulting bands were quantified and are presented as the mean ± SD (n = 3).
Figure 2. Sensitivity of gea1Δ/Δ cells to BFA can be rescued by overexpression of gea1p. A. Gea1+/− cells were transformed with pDUAL-YFH1c-gea1 to drive expression of gea1-YFP. Spinning-disc confocal fluorescence microscopy revealed that gea1-YFP localized to punctate structures in the cytoplasm that colocalized with the Golgi-specific stain BODIPY® TR C5-Ceramide. Scale bar, 14 μM. B. Wild-type and gea1Δ/− cells expressing endogenous gea1 tagged with GFP were imaged by fluorescence microscopy. The gea1-GFP protein localizes to punctate cytoplasmic structures. Scale bar, 10 μM. C. Wild-type, gea1+/−, and gea1Δ/− cells transformed with pDUAL-YFH1c-gea1 (gea1+/− + gea1-YFP) cells were subjected to 10-fold serial dilution and spotted on YES media and YES +10 μg/mL BFA. Plates were incubated at 30°C for 3 days. D. Equal numbers (5×10⁶ cells) of wild-type, wild-type (WT) + pDUAL-YFH1c (empty vector, WT + EV), wild-type + pDUAL-YFH1c-gea1 (WT + gea1), gea1+/−, gea1Δ/− + pDUAL-YFH1c,
and gea1+/− cells were stained with the Golgi-specific dye BODIPY FL C5-ceramide [75,82]. Ceramide staining revealed no overt differences in the punctate Golgi structures (Fig. 4A and Supplemental Figure S1). Results from transmission electron microscopy also suggested that Golgi membranes remain intact in gea1+/− cells (Fig. 4C).

Morphology of vacuoles and other endocytic organelles was assessed by uptake of the fluorescent dye FM4-64. Importantly, uptake of FM4-64 is dependent upon a competent endocytic pathway [64], and both wild-type and gea1+/− cells successfully took up the dye, indicating that endocytosis was not inhibited by loss of gea1. However, analysis of vacuoles stained with FM4-64 revealed a slight increase in vacuolar size (Fig. 4A). To quantify this increase in vacuolar size, the pixel area associated with the largest vacuole in each cell was analyzed. This analysis revealed a trend towards larger vacuoles in gea1+/− cells when compared to wild-type cells (Fig. 4B). We next analyzed whether increased vacuolar size in gea1+/− cells was associated with changes in vacuolar fusion. Incubation of fission yeast in a hypotonic solution results in an increase in vacuolar fusion [65]. Therefore, we stained wild-type and gea1+/− cells with FM4-64, followed by incubation in H2O for 90 minutes to induce fusion. No changes in vacuolar fusion were observed in gea1+/− cells compared to wild-type cells, suggesting that the enlarged vacuoles were competent for fusion (Fig. 4D).

**Gea1+/− cells exhibit defects in cytokinesis and septation**

Microscopic analysis of gea1+/− cells revealed a large number of septated cells and potential defects in septum structure (see Fig. 3A). Based on these observations, we hypothesized that gea1+/−

Figure 3. COP-I-dependent transport is impaired in gea1+/− cells. A. Wild-type and gea1+/− cells were transformed with pDUAL-YFH1c-arfl or pDUAL-YFH1c-arfl6 and imaged by fluorescence microscopy. Scale bar, 10 μM. B. Wild-type and gea1+/− cells were transformed with pDUAL-YFH1c-sac11 (SPBC19F5.03) or pDUAL-YFH1c-sac12 (SPAC3C7.01c) and imaged by fluorescence microscopy. Scale bar, 10 μM. C. Localization of sac11-YFP in wild-type (n = 207) and gea1+/− cells (n = 110) was scored as ER (surrounding the cell cortex and nuclear envelope), Golgi (punctate in the cytoplasm), or mixed. Sac11-YFP was predominately found in the ER in wild-type cells and in the Golgi and mixed in gea1+/− cells. D. Localization of sac12-YFP in wild-type (n = 204) and gea1+/− cells (n = 147) was scored as ER, Golgi, or mixed. Sac11-YFP was predominately found in the Golgi in both wild-type and gea1+/− cells. Error bars represent the mean ± SD from 3 independent experiments.

doi:10.1371/journal.pone.0056807.g003
2 cells have defects in the cell cycle, likely in assembly of the contractile actomyosin ring and/or septum. Cell cycle delays in *S. pombe* are associated with an increase in cellular length [83]. Therefore, to assess whether the cell cycle might be altered in *gea1*+/− cells, we measured the length of wild-type and *gea1*+/− cells. The mutant cells were approximately 40% longer than the wild-type cells, consistent with a cell cycle delay (Fig. 5A).

To determine whether the cell cycle delay was associated with impaired cytokinesis, we stained wild-type and *gea1*+/− cells with Alexa 568-phalloidin, which specifically binds to F-actin, labeling actin patches and actomyosin rings [84]. In wild-type cells, actin staining was clearly visible in contractile rings located in the medial portion of binucleate cells and at the cell tips (Fig. 5B). Actin patches were also visible at the tips of *gea1*+/− cells, although they appeared slightly less organized than in the wild-type cells. However, the most notable difference between the two strains was in the medial contractile ring structures in the *gea1*+/− cells. Although some normal contractile rings were present (Figure 5B, asterisk), many of the contractile rings appeared to be comprised of extended networks of medial actin patches, in some cases surrounding an abnormal septum (Figure 5B, arrowheads). Quantification of these results revealed that although the numbers of medial actin ring structures were similar in wild-type and *gea1*+/− cells, the haploinsufficient cells exhibited a significant increase

---

**Figure 4. Organellar morphology in *gea1*+/− cells.** A. Wild-type and *gea1*+/− cells were stained with 5 μM BODIPY FL C5-ceramide to label the Golgi or with 32 μM FM4-64 to label the vacuole. Staining was visualized by fluorescence microscopy. Scale bar, 10 μM. B. The pixel area associated with the largest vacuole was measured using Image J for individual wild-type (n = 262) and *gea1*+/− (n = 225) cells. The percentage of cells with vacuolar sizes of the indicated ranges was plotted using SigmaPlot. C. Wild-type (A) and *gea1*+/− cells (B) were subjected to transmission electron microscopy to visualize membranous structures. Representative images show flat ribbon-like structures consistent with Golgi membranes (insets) that appear similar in wild-type and *gea1*+/− cells. N, nucleus. Scale bar, 500 nm. D. Cells were labeled with FM4-64, followed by incubation in H2O for 90 min to induce vacuolar fusion. Scale bar, 10 μM.

doi:10.1371/journal.pone.0056807.g004
in the number of disorganized, abnormal rings compared to the wild-type cells (Figure 5C).

Defects in actomyosin ring assembly led us to analyze septum morphology in gea1+/− cells. In S. pombe, the septum is a cell wall-related structure that forms between cells undergoing cytokinesis and is subsequently degraded to separate the two new cells [85–87]. Staining with calcofluor revealed that gea1+/− cells exhibit a significantly higher septation index and have an increased number of abnormal septa (Fig. 6A). Morphological abnormalities observed in septated cells included the presence of multiple septa per cell and/or septa that were forked, mislocalized, or abnormally thick (Fig. 6A). Ultrastructural observation of these cells by transmission electron microscopy confirmed the presence of altered septum structures (Fig. 6C).

Secretion of eng1p is selectively impaired in gea1+/− cells

Production and degradation of the septum require intact membrane transport pathways [88–93], and enzymes that make and degrade the septum are secreted in a cell cycle-regulated manner [94,95]. These observations suggest a potential role for gea1p in transport of these enzymes to the septum. To determine whether defects in septation were due to generalized inhibition of secretion in gea1+/− cells, we analyzed secretion of acid phosphatase, a highly secreted protein in fission yeast [96]. Levels of acid phosphatase secretion were similar between wild-type and gea1+/− cells (Fig. 7A), indicating that general secretion was not impaired by loss of one copy of gea1. Treatment of wild-type cells with 40 μg/mL BFA, a dose previously shown to inhibit secretion in S. pombe [97], resulted in substantial inhibition of acid phosphatase secretion, which was even greater in gea1+/− cells (Fig. 7A). Although the BFA hypersensitivity phenotype suggests that general protein secretion may be partially regulated by gea1, the single remaining copy of gea1 is clearly sufficient to support secretion of at least some proteins in gea1+/− cells.

As we observed no defects in general protein secretion in gea1+/− cells, we next tested whether specific defects in transport of enzymes relevant to septation were present in gea1+/− cells. We chose eng1p, a β-glucanase involved in degradation of the primary septum, as a model enzyme based on previous evidence that eng1p is secreted, requiring the function of the exosome complex and rho4 [63,89,91]. Loss or altered secretion of eng1p results in increases in the septation index and in the number of septa per cell [63,89,91], similar to the morphological defects we observed in gea1+/− cells. Importantly, secretion of eng1p differs significantly from that of acid phosphatase because eng1p is both produced and secreted in a cell cycle-specific manner [63,89,91]. To analyze eng1p secretion in gea1+/− cells, we performed a β-glucanase activity assay, which has previously been established as a highly specific measure of β-glucanase levels [63]. Eng1p activity levels were similar in lysates from wild-type and gea1+/− cells, indicating similar levels of expression (Fig. 7B). However, substantially less eng1p activity was observed in the medium from gea1+/− cells (Fig. 7B), suggesting a defect in secretion. To assess whether decreased secretion of eng1p was accompanied by mislocalization of the protein, we tagged endogenous eng1 with GFP. Importantly,
Western blot analysis revealed no significant difference in eng1-GFP expression between the wild-type and \textit{gea1}+/- strains ($p = 0.70$), consistent with enzymatic assays (compare Fig. 7A and 7E). As previously described, eng1-GFP localized to the septum in wild-type cells ([63] and Fig. 7C). A total of 94% of septated wild-type cells exhibited localization of eng1-GFP to the septum. In contrast, in \textit{gea1}+/- cells, levels of eng1p were decreased or absent from the septa (Fig. 7C), with only 59% of septated \textit{gea1}+/- cells having detectable levels of eng1-GFP at the septum. These results indicate that loss of one copy of \textit{gea1} is sufficient to impair secretion of specific cargo proteins required for cytokinesis. Importantly, overexpression of eng1p rescued the excessive septation defect in the \textit{gea1}+/- strain (Figure 7D), confirming that \textit{gea1} is a member of the GBF1/GEA family of Arf GEFs. Additionally, a fission yeast homolog of the COP-I cargo sac1 was mislocalized from the ER to the Golgi in \textit{gea1}+/- cells, consistent with a role for \textit{gea1} in COP-I-dependent transport, similar to its mammalian counterpart [33,49].

Comparisons with data from other fission yeast membrane trafficking mutants are consistent with a role for \textit{gea1} in membrane transport. \textit{Gea1}+/- cells exhibit slight sensitivity to FK506 and valproic acid (Fig. 1C), similar to strains with mutations or deletions in the subunits of the AP-1 coat complex, and the GTPases \textit{rho3}, \textit{ypt3}, and \textit{ryh1} [67,72–74]. However, although \textit{gea1} mutants exhibit some phenotypic overlap with these mutants with respect to FK506 and VPA sensitivity and defects in septation, some of the phenotypic aspects associated with \textit{gea1} haploinsufficiency appear to be quite different. For example, we found that vacuolar size was slightly increased and secretion of acid phosphatase was not impaired in \textit{gea1}+/- cells (Fig. 4B, 7A). In contrast, deletion of the \textit{atp6} subunit of AP-1 was shown to result in increased vacuolar fragmentation and decreased acid phosphatase.

**Discussion**

In this study, we have established \textit{S. pombe} as a model system to analyze Arf-GEF function. Unlike \textit{S. cerevisiae}, \textit{S. pombe} possesses only one ortholog of GBF1, similar to vertebrates (Fig. 1A). Importantly, we have shown that \textit{gea1}+/- cells exhibit sensitivity to the GBF1 inhibitor BFA (Fig. 1C, 2C, 2D) and that overexpression of \textit{gea1}-YFP results in BFA resistance (Fig. 2D), confirming that \textit{gea1} is a member of the GBF1/GEA family of Arf GEFs. Additionally, a fission yeast homolog of the COP-I cargo sac1 was mislocalized from the ER to the Golgi in \textit{gea1}+/- cells, consistent with a role for \textit{gea1} in COP-I-dependent transport, similar to its mammalian counterpart [33,49].

**Figure 6.** \textit{Gea1}+/- cells exhibit alterations in septum number and morphology. \textit{A.} Wild-type and \textit{gea1}+/- cells were stained with calcofluor white to visualize septa and imaged by fluorescence microscopy. Scale bar, 10 \textmu{}m. \textit{B.} Quantification of \textit{A.} Cells were scored as having abnormal septa if multiple septa were present, or if the septum was mislocalized, abnormally thick, or forked. Error bars represent mean ± SD from 3 independent experiments. \textit{C.} Wild-type and \textit{gea1}+/- cells were subjected to transmission electron microscopy to visualize septum defects. Representative images of multi-septated cells and septa with morphological abnormalities are shown. Scale bar, 1 \textmu{}m.

doi:10.1371/journal.pone.0056807.g006
We also found that gea1 haploinsufficiency resulted in cold resistance and that FK506 sensitivity could be reversed at lower temperatures (Fig. 1C), differing from previous studies of other membrane trafficking mutants [67,72–74]. FK506 is an inhibitor of calcineurin, and inhibition or loss of calcineurin activity has been shown to impair both septation and membrane transport in fission yeast [67,72–74,98]. We speculate that slower growth at colder temperatures may allow sufficient time for delayed trafficking of enzymes required for septation in haploinsufficient gea1+/− cells, potentially underlying reversed sensitivity to FK506. Together, these observations suggest that, although gea1 is appears to be important for membrane trafficking and septation in S. pombe, this role may be distinct from that of previously characterized membrane trafficking mutants in fission yeast.

As mentioned previously, we observed no defects in secretion of acid phosphatase in gea1+/− cells (Fig. 7A), despite the observation that treatment with BFA has been shown to block protein secretion in fission yeast ([97] and Fig. 7A). However, previous studies have shown that impaired function of GBF1/GEA family members in both mammalian cells and budding yeast causes cargo-specific defects in protein secretion. Secretion of transmembrane, but not soluble, cargoes is decreased in mammalian cells depleted of GBF1 [50], and secretion of only a subset of cargoes is decreased in budding yeast cells with mutations in gea1 [37]. Together, these results suggest that the GBF1/GEA family members, which are evolutionarily conserved, may have unique roles in regulating specific trafficking pathways.

**Figure 7. Secretion of the β-glucanase eng1p is selectively inhibited in gea1+/− cells.** A. Equal numbers of wild-type and gea1+/− S. pombe cells were used to inoculate 10 mL of EMM media. At the indicated times, an aliquot of the media was subjected to spectrophotometric determination of secreted acid phosphatase activity by monitoring cleavage of the substrate p-nitrophenyl phosphate at 405 nm. Acid phosphatase activity was normalized to cell density, determined by monitoring absorbance at 600 nm (OD600). Treatment with 40 μg/mL BFA to inhibit secretion served as a negative control. B. β-glucanase activity in cell lysates and in media was measured by quantifying release of reducing sugars from the substrate laminarin as previously described [68]. Normal levels of β-glucanase activity were detected in wild-type and gea1+/− cell lysates, but β-glucanase activity was decreased in the media from gea1+/− cells. Error bars represent the mean ± SD from 3 independent experiments. *p<0.02 C. Wild-type and gea1+/− cells expressing eng1p tagged with GFP under control of the endogenous promoter were imaged by fluorescence microscopy. Localization of eng1-GFP to the septum was substantially decreased in gea1+/− cells. Arrows indicate septated cells with no visible eng1-GFP at the septum. Scale bar, 10 μM. D. Gea1+/− cells were transformed with either the empty pREP4X vector or pREP4X carrying eng1. Wild-type and both gea1+/− strains were then stained with calcofluor, and the percentage of cells with septa were counted for each culture. Overexpression of eng1p rescued the excessive septation defect of gea1+/− cells. Error bars represent the mean ± SD from 3 independent experiments. *p<0.02 compared to wild-type. ***p<0.05 compared to gea1+/- + pREP4X. E. Lysates from wild-type and gea1+/− cells in which endogenous eng1 was tagged with GFP (shown in C) were subjected to immunoblot analysis using antibodies directed against GFP and β-actin. No significant difference in eng1p-GFP levels was observed between the wild-type and gea1+/− cells (p = 0.70).

DOI: 10.1371/journal.pone.0056807.g007
family may selectively regulate secretion of specific cargo proteins, as opposed to total protein secretion.

In addition to impaired secretion of transmembrane cargo proteins, >90% depletion of GBF1 in mammalian cells has been shown to result in tubulation and fragmentation of the Golgi and decreased recruitment of Arf to the membrane [50]. However, in the *gea1/−* model, no overt defects in Golgi structure or Arf membrane recruitment were observed (Fig. 3A, 4A, 4C). Several differences between *S. pombe* *gea1/−* cells and mammalian GBF1-depleted cells may underlie these inconsistencies. First, *gea1/−* cells retain 50% of *gea* expression (Fig. 1B), which may be sufficient to perform the normal housekeeping activities of this protein in fission yeast. Second, the Golgi architecture differs substantially between yeast and mammalian cells. In fission yeast, the Golgi is present as multiple polarized mini-stacks, in contrast to the single Golgi ribbon present in mammalian cells [99]. This “fragmented” nature may reflect differences in Golgi biogenesis between yeast and mammalian cells and may also hinder observations of subtle changes in Golgi structure or Arf recruitment. Third, fission yeast have only one class I/II Arf (arf1p, Fig. 3A), but they possess three Arf-GEFs predicted to localize to the Golgi and/or endosomes, gea1p, sec71p, and sec72p. In contrast, the mammalian homologs of these Arf GEFs (GBF1, BIG1, and BIG2) exhibit distinct specificities for the four Golgi-localized Arfs present in mammalian cells [50,51,100]. Therefore, there may be functional redundancies between some of the fission yeast Arf-GEFs that are not present in mammalian cells with respect to maintenance of Golgi structure, Arf recruitment, and general secretion.

Analysis of the *gea1/−* mutant has uncovered a novel role for *gea* in regulation of vacuolar size. Consistent with the localization of arf1p to both the Golgi and the vacuole, we observed a slight increase in vacuolar size in *gea1/−* cells (Fig. 4A, B). Interestingly, mutations in *S. cerevisiae* *gea1* and *gea2* have previously been shown to cause a slight increase in vacuolar fragmentation, consistent with a role for this family of proteins in vacuolar homeostasis in yeast [37]. However, the precise mechanisms underlying this phenotype remain unclear. Alterations in trafficking of ion channels to vacuolar membranes could alter the osmotic pressure of the vacuole, resulting in swollen vacuoles. Alternatively, improper delivery of factors required for vacuolar fusion and fusion could dysregulate the precise balance between fusion and fission required to maintain vacuolar size.

The major defects observed in *gea1/−* cells were associated with cytokinesis and septation. We observed disorganization of contractile actomyosin ring (CAR) structures (Fig. 5) and cells with multiple septa, mislocalized septa, and malformed septa (Fig. 6). Cytokinesis is a tightly regulated process in *S. pombe*, governed by signaling of kinases of the septation initiation network [85,101]. The septum forms just behind the CAR, and septum formation is tightly linked to CAR assembly [102]. Septum formation requires polarized delivery of glucan synthases, such as bgslp, that synthesize the new cell wall material [103,104] and glucanases, such as eng1lp and arg1lp, that rapidly break down the septum to separate the two new daughter cells [63,94]. Defects in delivery of these enzymes to the site of septum assembly and breakdown result in impaired cell division, similar to that observed in *gea1/−* cells (Fig. 6). As an initial approach to determine whether secretion of cell cycle-specific enzymes was impaired in *gea1/−* cells, we chose to examine eng1lp secretion. Eng1lp has been shown to be required for dissolution of the primary septum, and defects in secretion of eng1lp have been shown to affect septation in fission yeast [63,89,91]. Specifically, mutation of *rho4* and genes belonging to the exocyst complex, a part of the machinery required for vesicle fusion, have been shown to decrease secretion of eng1lp, resulting in impaired septum breakdown and septation defects overlapping those we observed in *gea1/−* cells [89,91]. Furthermore, the cell cycle-specific expression pattern of eng1lp has been well-characterized and is demonstrated to peak specifically during septation [63]. Our results demonstrate that secretion of eng1lp is impaired in *gea1/−* cells. Although eng1 activity was similar in cell lysates from wild-type and *gea1/−* cells, secretion of eng1lp activity from *gea1/−* cells was approximately 20% of the wild-type (Fig. 7B). We also observed a substantial decrease in localization of eng1-GFP to the septum in *gea1/−* cells. Furthermore, similar to rho4A mutants [89], overexpression of eng1lp was able to suppress the excessive septation defect in *gea1/−* cells (Fig. 7D). These results suggest that, although eng1lp is produced, only a small fraction of eng1lp reaches the septum during cytokinesis. We hypothesize that the mislocalized eng1lp is likely degraded due to lack of export.

Characterization of *gea* function in *S. pombe* may provide insight into a conserved network of proteins that connect membrane traffic and cytokinesis in other eukaryotes. In the present study, we observed defects in trafficking of sac1lp in *gea1/−* cells (Fig. 3B, C). Importantly, the *S. cerevisiae* homolog of sac1lp has been shown to play a role in septation, suggesting that this pathway may be conserved in yeast [105]. The secretory pathway also appears to play an important role in cytokinesis in higher eukaryotes. In *C. elegans*, treatment of embryos with Brefeldin A results in regression of the cleavage furrow, suggesting that secretion is required for completion of cytokinesis [106]. Secretion also occurs at the cleavage furrow in sea urchin embryos late in cytokinesis and is independent of constriction of the cleavage furrow [107].

Characterizing the network of proteins that work with *gea* to drive septation in fission yeast may also shed light on these pathways in mammalian cells. For example, defects in contractile actomyosin ring assembly and septum structure similar to those found in the *gea1/−* mutant have been observed in cells expressing mutants of pik1p, a fission yeast phosphatidylinositol-4-kinase (PI4K) [100]. These overlapping phenotypes suggest that *gea* and pik1 may act in the same pathway to control cytokinesis and septum biogenesis. In mammalian cells, PI4K type IIIa has been shown to be required for Rab1-mediated recruitment of GBF1 to Golgi membranes [109]. Additionally, mammalian PI4KIIib has been shown to colocalize with GBF1 during hepatitis C virus replication and, like GBF1, to be required for viral replication [110]. Together these results suggest that protein networks connecting GBF1/GEA family members and PI4K family members may be conserved in higher eukaryotes.

Together, our observations suggest a novel role for *gea* in polarized, cell cycle-specific secretion. The selective inhibition of polarized secretion of eng1lp to the septum suggests that the GBF1/GEA family may play previously unappreciated roles in cell cycle progression. GBF1 has previously been implicated in regulation of the cell cycle, as depletion of GBF1 results in cell cycle arrest at the G0/G1 phase and is associated with induction of ER unfolded protein response, ultimately inducing apoptosis [56]. GBF1 activity has also been shown to be regulated in a cell cycle-specific manner through phosphorylation by the cyclin B-cyclin dependent kinase 1 (CDK1) complex [57]. In budding yeast and in *Drosophila*, loss of gea1 and *gaz* activity have been associated with polarity defects, suggesting a role in polarized secretion [35,55,111]. Therefore, future studies to examine the secretion of other enzymes required for septum formation and of polarity factors required for CAR positioning in *gea1/−* cells may help to identify a network of interactions required for proper cell cycle.
progression and polarity determination in fission yeast and other organisms.

In summary, our data supports the hypothesis that gea1p plays an important role in cytokinesis in S. pombe by regulating the trafficking of key components required for the septation. These studies shed light on a novel role for the GBF1/GEA family of Arf-GEFs and establish S. pombe as a model to explore GBF1/GEA function.

Supporting Information

Figure S1 Analysis of Golgi localization. A. Wild-type and gen1Δ/Δ cells were transformed with pDUAL-YFH1c-arf1 and stained with the Golgi-specific stain BODIPY® TR C2-ceramide. Arf1-YFP exhibited limited colocalization with BODIPY® TR C2-ceramide. B. Wild-type and gen1Δ/Δ cells transformed with pDUAL-YFH1c-sac1/1 were stained with BODIPY® TR C2-ceramide. Sac11-YFP localized to the Golgi in gen1Δ/Δ cells, but not in wild-type cells. C. Wild-type cells transformed with pDUAL-YFH1c-sac1/2 and stained with BODIPY® TR C2-ceramide showed that sac12-YFP exhibited Golgi localization as expected. Scale bars, 14 μM.

Acknowledgments

The authors thank Amber O'Connor and Brad Yoder of the University of Alabama at Birmingham Department for Cell Biology for assistance with spinning disc confocal microscopy. We thank Ed Phillips at Birmingham High Resolution Imaging Facility for his assistance with electron microscopy sample preparation and imaging. We would also like to thank Isaac Delgado of Birmingham-Southern College for her assistance and technical expertise.

Author Contributions

Conceived and designed the experiments: AME AC VMF RP CW BW CB NL NN KPS ES MLS. Performed the experiments: AME AC VMF RAL RP CW BW CB NL NN KPS MLS. Analyzed the data: AME AC VMF RAL RP CW BW CB NL NN KPS ES MLS. Contributed reagents/materials/analysis tools: ES MLS. Wrote the paper: AME VMF RP CW KPS ES MLS. Edited and approved of the final manuscript: AME AC VMF RP CW BW CB NL NN KPS ES MLS.

References

1. Goldstein J, Brown MS (1976) The LDL pathway in human fibroblasts: a receptor-mediated mechanism for the regulation of cholesterol metabolism. Curr Top Cell Regul 11: 147–181.
2. Mayinger P (2011) Signaling at the Golgi. Cold Spring Harb Perspect Biol 3: 1–14.
3. McPherson PS, Kay BK, Hussain NK (2001) Signaling on the endocytic pathway. Traffic 2: 373–384.
4. Hirst J, Robinson MS (1998) Clathrin and adaptors. Biochim Biophys Acta 1404: 173–193.
5. Holloway ZG, Grabski R, Szul T, Syers ML, Coventry JA, et al. (2007) Activation of ARF-ribosylation factor catalyses biogenesis of the ATP7A-containing trans-Golgi network compartment and its Cu-induced trafficking. Am J Physiol Cell Physiol 293: C1753–1767.
6. Lippincott-Schwartz J, Roberts TH, Hirschberg K (2000) Secretory protein trafficking and organelle dynamics in living cells. Annu Rev Cell Dev Biol 16: 557–589.
7. Reggiori F, Klionsky DJ (2002) Autophagy in the eukaryotic cell. Eukaryot Cell 1: 11–21.
8. Bonifacino JS, Glick BS (2004) The mechanisms of vesicle budding and fusion. Cell 116: 153–166.
9. Rothman JE (1996) The protein machinery of vesicle budding and fusion. Science 270: 1090–1098.
10. Sabatini DD, Kreibich G, Morimoto T, Adesnik M (1982) Mechanisms for the regulation of the size of coated vesicles. Proc Natl Acad Sci U S A 79: 1238–1242.
11. Boman AL, Kahn RA (2000) A family of ARF guanine nucleotide exchange factors at the Golgi complex: GBF1 and GBF2. Mol Biol Cell 11: 1241–1255.
12. Boman AL, Kahn RA (1995) Arf proteins: the membrane traffic police? Trends Biochem Sci 20: 147–150.
13. Balch WE, Kahn RA, Schwanger R (1992) ARF-ribosylation factor is required for vesicular trafficking between the endoplasmic reticulum and the cis-Golgi compartment. J Biol Chem 267: 13053–13061.
14. Regazzi R, Ulrich S, Kahn RA, Wohllien CB (1991) Redistribution of ARF-ribosylation factor during stimulation of permeabilized cells with GTP anaologues. Biochim Biophys Acta 759 (Pt 5): 639–644.
15. Lippincott-Schwartz J, Cole NR, Donaldson JG (1998) Building a secretory apparatus: role of ARF1/COP1 in Golgi biogenesis and maintenance. Histochem Cell Biol 109: 449–462.
16. Donaldson JG, Kahn RA, Lippincott-Schwartz J, Klausner RD (1991) Binding of ARF and beta-COP to Golgi membranes: possible regulation by a trimeric G protein. Science 254: 1187–1199.
17. Lippincott-Schwartz J, Cole NR, Donaldson JG (1998) Building a secretory apparatus: role of ARF1/COP1 in Golgi biogenesis and maintenance. Histochem Cell Biol 109: 449–462.
18. Donaldson JG, Kahn RA, Lippincott-Schwartz J, Klausner RD (1991) Binding of ARF and beta-COP to Golgi membranes: possible regulation by a trimeric G protein. Science 254: 1187–1199.
19. Serafini T, Orci L, Amherdt M, Brunner M, Kahn RA, et al. (1991) ARF-ribosylation factor is a subunit of the coat of Golgi-derived COP-coated vesicles: a novel role for a GTP-binding protein. Cell 67: 239–253.
20. Teash LM, Ostrom JA, Kornfeld S (1993) Biochemical dissection of ARF1 recruitment onto Golgi membranes. J Cell Biol 123: 561–573.
21. Teash LM, Ostrom JA, Kornfeld S (1993) Biochemical dissection of ARF1 recruitment onto Golgi membranes. J Cell Biol 123: 561–573.
22. Teash LM, Ostrom JA, Kornfeld S (1993) Biochemical dissection of ARF1 recruitment onto Golgi membranes. J Cell Biol 123: 561–573.
23. Teash LM, Ostrom JA, Kornfeld S (1993) Biochemical dissection of ARF1 recruitment onto Golgi membranes. J Cell Biol 123: 561–573.
BG1s are required for assembly and maintenance of the Golgi stack and trans-Golgi network, respectively. Mol Biol Cell 19: 523–535.

39. Zhao X, Lasell TK, Melanson P (2002) Localization of large ADP-ribosylation factor-guanine nucleotide exchange factors to different Golgi compartments: evidence for distinct functions in protein traffic. Mol Biol Cell 13: 119–133.

40. Shintosato C, Wajgur S, Wakahara M, Uchihaya Y, Nakayama K (2002) Dominant-negative mutant of BIG2, an ARF guanine nucleotide exchange factor, specifically affects membrane trafficking from the trans-Golgi network through inhibition of membrane association. Yeast 18: 1-12.

41. Morohashi Y, Balklava Z, Ball M, Hughes H, Lowe M (2010) Phosphorylation regulates septation in fission yeast. J Bacteriol 192: 2213–2223.

42. Achstetter T, Franzusoff A, Field C, Schekman R (1988) SEC7 encodes an ADP-ribosylation factor in yeast. J Cell Biol 107: 1-CF.

43. Shintosato C, Yoshida Y, Kawamoto K, Takatsu H, Nakayama K (2002) Overexpression of an ADP-ribosylation factor-guanine nucleotide exchange factor, BIG, uncouples budding from membrane association. Yeast 18: 1419-1427.

44. Kawamoto K, Yoshida Y, Tamaki H, Torii N, Shintosato C, et al. (2000) The ADP-ribosylation factor-nucleotide exchange factors Gea1p and Gea2p have overlapping, but not redundant functions in retrograde transport from the Golgi to the endoplasmic reticulum. Mol Biol Cell 12: 1035–1045.

45. Kawamoto K, Yoshida Y, Tamaki H, Torii N, Shintosato C, et al. (2002) GBF1, a guanine nucleotide exchange factor for ADP-ribosylation factors, is localized to the cis-Golgi and involved in membrane association of the COP I coat. Traffic 3: 463–473.

46. Styne ML, O’Connor AK, Grabowski R, Cockett-Bocca E, Nasel E (2008) Deletion of beta-COP reveals a role for COP I in compartmentalization of secretory compartments and in biosynthetic transport of carbohydrate. In: Am J Physiol Cell Physiol 294: C485–1947.

47. Shinohara T, Takeda K, Hasegawa H, Tagawa T, Yamashita H, et al. (1998) Identification of a novel ADP-ribose acceptor and anterograde-cargo-rich domains in endoplasmic reticulum-to-Golgi transport complexes. Curr Biol 9: 821–824.

48. Abo M, Banjoh M, Rosse T, Balch WE (1995) Sequential coupling between COP I and COP I coat vesicles in endoplasmic reticulum to Golgi transport. J Cell Biol 131: 873–893.

49. Lee MC, Miller EA, Goldberg J, Orci L, Schekman R (2004) Bi-directional transport between the ER and Golgi. Annu Rev Cell Dev Biol 20: 87–123.

50. Rohde HL, Cheong FY, Konrad G, Pauk H, Mayinger P, et al. (2003) The human phosphathosphorylase SAC1 interacts with the coatomer I complex. J Biol Chem 278: 52089–52099.

51. Nasel T, Grabowski R, Lyoos S, Morohashia Y, Shresthap S, et al. (2007) Dissection of the function of the guanine nucleotide exchange factor GBF1 in Golgi biogenesis and protein trafficking. J Cell Sci 120: 3929–3940.

52. Claude A, Zhao BP, Kuziemsky CE, Dahan S, Berger SJ, et al. (1999) GBF1: A novel Golgi-associated BFA-resistant guanine nucleotide exchange factor that displays specificity for ADP-ribosylation factor 5. J Cell Biol 145: 71–84.

53. Zakrzewska E, Perron M, Laroche A, Pallotta D (2003) A role for GEA1 and BIG1 in membrane trafficking. Mol Biol Cell 13: 765–778.

54. Nefedov S, McCormick PJ (2007) The Arf GEF GBF1 is required for GGA coat proteins localization and membrane tubulation. J Biol Chem 277: 9086–9097.

55. Lefrancois S, McCormick PJ (2007) The Arf GEF GBF1 is required for protein trafficking through functional interaction with adaptin in Fission Yeast. J Cell Sci 120: 1015–1028.

56. Miyazaki K, Kono T, Kita K, Katsura K, Takegawa K, et al. (2007) Valproic acid affects membrane trafficking and cell-wall integrity in fission yeast. Genetics 175: 1655–1675.

57. Pagano RE, Martin OC, Kang HC, Haugland RP (1995) A novel fluorescent ceramide analogue for studying membrane traffic in animal cells: accumulation at the Golgi apparatus results in altered spectral properties of the sphingolipid precursor. J Cell Biol 111: 1267–1279.

58. Natter K, Leitner P, Fasching A, Wolinski H, McCraith S, et al. (2005) The spatiotemporal organization of lipid synthesis in the yeast Saccharomyces cerevisiae derived from large scale green fluorescent protein tagging and high resolution microscopy. Mol Cell Proteomics 4: 662–672.

59. Ji L, Kelly WG, Logsdon JM Jr, Schurko AM, Harle BD, et al. (2004) Functional genomic analysis of the ADP-ribosylation factor family of GTPases: phosphorylation among diverse eukaryotes and function in C. elegans. FASEB J 18: 3834–3850.

60. Casnaghi MM, Whitney JA, Carroll K, Zhang C, Boman AL, et al. (1996) Intracellular distribution of Arf proteins in mammalian cells: Arf6 is uniquely localized to the plasma membrane. J Biol Chem 271: 21767–21774.

61. Kondo Y, Hanai Y, Nakai W, Kato Y, Nakayama K, et al. (2012) ARF1 and ARF3 Are Required for the Integrity of Recycling Endosomes and the Recycling Pathway. Cell Struct Funct 37: 141–154.

62. Roel F, Stephens DJ (2010) Specific functions of BIG1 and BIG2 in endomembrane organization. PLoS One 5: e10809.

63. Ishizaki R, Shin HW, Mitsuhashi H, Nakayama K (2000) Redundant roles of BIG1 and BIG2, guanine-nucleotide exchange factors for ARF-guanine nucleotide exchange factors, in membrane traffic. Mol Biol Cell 11: 1527–1535.

64. Vida TA, Emer SD (1995) A new viral stain for visualizing vacuolar membrane dynamics and endocytosis in yeast. J Cell Biol 128: 779–792.

65. Bone N, Miller JB, Toda T, Armstrong J (1998) Regulated vacuole fusion and fission in Schizosaccharomyces pombe: an osmotic response dependent on membrane streaming. Curr Biol 8: 135–144.

66. Sawin KE, Nurse P (1998) Regulation of cell polarity by microtubules in fission yeast. J Cell Biol 142: 457–471.

67. Kita A, Sugura R, Shoji H, He Y, Deng L, et al. (2004) Loss of Apm1, the multiprotein exocyst complex subunit, in the fission yeast Schizosaccharomyces pombe affects cell separation execution and endocytosis in fission yeast. Mol Biol Cell 15: 2920–2931.

68. San Segundo P, Correa J, Vazquez de Aldana CR, del Rey F (1995) SNG1, a gene encoding a sporulation-specific 1.3-beta-glucanase in Saccharomyces cerevisiae. J Bacteriol 175: 3832–3837.

69. Somogyi M (1952) Notes on sugar determination. J Biol Chem 195: 19–23.

70. Nelson MJ (1957) Colorimetric analysis of sugars. Methods Enzymol 3: 85–86.

71. Klausner RD, Donaldson JG, Lippincott-Schwartz J (1992) Brefeldin A: inside of the control of membrane traffic and organellar structure. J Cell Biol 116: 1071–1080.

72. Kita A, Li C, Yu Y, Umeda N, Diao A, et al. (2011) Role of the Small GTPase Rho5 in Golgi/Endosome trafficking through functional interaction with adaptin in Fission Yeast. Mol Biol Cell 14: 1015–1028.

73. Miyazaki K, Kuno T, Kita K, Katsura K, Takegawa K, et al. (2007) Valproic acid affects membrane trafficking and cell-wall integrity in fission yeast. Genetics 175: 1655–1675.

74. Matsuyama A, Shirai A, Yashiroda Y, Kamata A, Horinouchi S, et al. (2004) Evidence for distinct functions in protein traffic of the trans-Golgi network through inhibition of membrane association. Yeast 18: 1419–1427.

75. Martin-Cuadrado AB, Duenas E, Sipiczki M, Vazquez de Aldana CR, del Rey F (2003) The endo-beta-1,3-glucanase engl1 is required for dissolution of the primary septum during cell separation in Schizosaccharomyces pombe. J Cell Sci 116: 1683–1690.
94. Dekker N, Speijer D, Grun CH, van den Berg M, de Haan A, et al. (2004) Role of the alpha-glucanase Agn1p in fission-yeast cell separation. Mol Biol Cell 15: 3903–3914.
95. Cortes JC, Ishiguro J, Dura A, Ribas JC (2002) Localization of the (1,3)beta-D-glucan synthase catalytic subunit homologue Bgs1p/Cps1p from fission yeast suggests that it is involved in septation, polarized growth, mating, spore wall formation and spore germination. J Cell Sci 115: 4081–4096.
96. Mitchison JM, Croam J (1969) Linear synthesis of sucrase and phosphatases during the cell cycle of Schizosaccharomyces pombe. J Cell Sci 5: 373–391.
97. Turi TG, Webster P, Rose JK (1994) Brefeldin A sensitivity and resistance in Schizosaccharomyces pombe. Isolation of multiple genes conferring resistance. J Biol Chem 269: 24229–24236.
98. Yoshida T, Toda T, Yanagida M (1994) A calcineurin-like gene ppb1+ in fission yeast: mutant defects in cytokinesis, cell polarity, mating and spindle pole body positioning. J Cell Sci 107 (Pt 7): 1725–1735.
99. Suda Y, Nakano A (2011) The Yeast Golgi Apparatus. Traffic.
100. Togawa A, Morinaga N, Ogasawara M, Moss J, Vaughan M (1999) Purification and cloning of a brefeldin A-inhibited guanine nucleotide-exchange protein for ADP-ribosylation factors. J Biol Chem 274: 12308–12315.
101. Krapp A, Simanis V (2006) An overview of the fission yeast septation initiation network (SIN). Biochem Soc Trans 36: 411–415.
102. Geiss KL, Simanis V (1997) The control of septum formation in fission yeast. Genetics 145: 2939–2951.
103. Liu J, Wang H, McCollum D, Balasubramanian MK (1999) Dec1lp/Cps1lp, a 1,3-beta-glucan synthase subunit, is essential for division septum assembly in Schizosaccharomyces pombe. Genetics 153: 1193–1203.
104. Cortes JC, Kontoni M, Martins IM, Munoz J, Moreno MB, et al. (2007) The (1,3)beta-D-glucan synthase subunit Bgs1p is responsible for the fission yeast primary septum formation. Mol Microbiol 65: 201–217.
105. Tahirovic S, Schorr M, Then A, Berger J, Schwarz H, et al. (2003) Role for lipid signaling and the cell integrity MAP kinase cascade in yeast septum biogenesis. Curr Genet 45: 71–78.
106. Skop AR, Bergmann D, Mohler WA, White JK (2003) Completion of cytokinesis in C. elegans requires a brefeldin A-sensitive membrane accumulation at the cleavage furrow apex. Curr Biol 11: 735–746.
107. Shuster CB, Burgess DR (2002) Targeted new membrane addition in the cleavage furrow is a late, separate event in cytokinesis. Proc Natl Acad Sci U S A 99: 3633–3638.
108. Park JS, Steinbach SK, Desautels M, Hemmingsen SM (2009) Essential role for Schizosaccharomyces pombe pkl1 in septation. PLoS One 4: e6179.
109. Dumaresq-Doiron K, Savard MF, Akam S, Costantino S, Lefrancois S (2010) The phosphatidylinositol 4-kinase PI4KIIIalpha is required for the recruitment of GBF1 to Golgi membranes. J Cell Sci 123: 2273–2280.
110. Zhang L, Hong Z, Lin W, Shao RX, Goto K, et al. (2012) ARF1 and GBF1 generate a PI(4)P-enriched environment supportive of hepatitis C virus replication. PLoS One 7: e32135.
111. Soul T, Burgess J, Jeon M, Zinn K, Marques G, et al. (2011) The Gark Sec7 domain guanine nucleotide exchange factor for Arf regulates salivary gland development in Drosophila. Cell Logist 1: 69–76.