Cross-talk between clathrin-dependent post-Golgi trafficking and clathrin-mediated endocytosis in Arabidopsis root cells

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Abstract

Coupling of post-Golgi and endocytic membrane transport ensures that the flow of materials to/from the plasma membrane (PM) is properly balanced. The mechanisms underlying the coordinated trafficking of PM proteins in plants, however, are not well understood. In plant cells, clathrin and its adaptor protein complexes, AP-2 and the TPLATE complex (TPC) at the PM, and AP-1 at the trans-Golgi network/early endosome (TGN/EE), function in clathrin-mediated endocytosis (CME) and post-Golgi trafficking. Here, we utilized mutants with defects in clathrin-dependent post-Golgi trafficking and CME, in combination with other cytological and pharmacological approaches, to further investigate the machinery behind the coordination of protein delivery and recycling to/from the TGN/EE and PM in Arabidopsis (Arabidopsis thaliana) root cells. In mutants with defective AP-2-/TPC-dependent CME, we determined that clathrin and AP-1 recruitment to the TGN/EE as well as exocytosis are significantly impaired. Likewise, defects in AP-1-dependent post-Golgi trafficking and pharmacological inhibition of exocytosis resulted in the reduced association of clathrin and AP-1 recruitment to the TGN/EE and a reduction in the internalization of cargoes via CME. Together, these results suggest that post-Golgi trafficking and CME are coupled via modulation of clathrin and adaptor protein complex recruitment to the TGN/EE and PM.

Introduction

Post-Golgi secretory and endocytic vesicle trafficking pathways control the delivery and retrieval of proteins and membranes from the cell surface and thereby regulate cell growth and numerous biochemical processes at the plasma membrane (PM) essential for the interaction and exchange of information between a cell and its environment. Clathrin, an evolutionarily conserved vesicle coat protein complex, which localizes to the PM and trans-Golgi network/early
endosome (TGN/EE) (Konopka et al., 2008; Ito et al., 2012), plays a critical role in post-Golgi exocytic/endocytic routes that control the abundance and activity of PM proteins (Lam et al., 2007; McMahon and Boucrot, 2011; Kirchhausen et al., 2014; Zouhar and Sauer, 2014; Kukulski et al., 2016; Kaksonen and Roux, 2018; Reynolds et al., 2018). Binding of clathrin to the PM or TGN/EE is dependent upon membrane compartment-specific adaptor protein complexes that, together with clathrin, recruit additional accessory proteins necessary for the formation, cargo recognition, budding, and release of cargo-containing clathrin-coated vesicles (CCVs) (McMahon and Boucrot, 2011; Paez Valencia et al., 2016; Kaksonen and Roux, 2018). Clathrin-mediated endocytosis (CME) in plants is dependent upon various endocytic accessory proteins, including two types of adaptor protein complexes: the evolutionarily conserved heterotetrameric complex ADAPTOR PROTEIN2 (AP-2, consisting of AP2α, AP2β, AP2μ, and AP2σ), and the evolutionarily ancient heterooctameric TPLATE complex (TPC) (Kim et al., 2013; Yamaoka et al., 2013; Gadeyne et al., 2014; Zhang et al., 2015). Following budding and release from the PM, cargo-containing plant CCVs uncoat in close proximity to the TGN/EE, rather than immediately after scission from the PM (Narasimhan et al., 2020). Within the TGN/EE, PM proteins are either recycled back to the PM or targeted through intermediate/late endosomes (termed multivesicular bodies; MVBs) for vacuolar degradation. In addition to PM cargos, endosomes also receive secretory proteins translated de novo and fated for the vacuole.

The ADAPTOR PROTEIN1 complex (AP-1, consisting of AP1γ, AP1β, AP1μ, and AP1σ) functions in clathrin-dependent sorting and trafficking between the TGN and endosomes (Castillon et al., 2018; Martzoukou et al., 2018). However, AP-1 has also been implicated in the polarized sorting of PM proteins in epithelial and neuronal cells (Gravotta et al., 2012, 2019; Li et al., 2016) as well as intra-Golgi recycling of secretory cargoes and Golgi-resident proteins (Papanikou et al., 2015; Day et al., 2018; Casler et al., 2019).

In plants, morphometric studies have estimated that 75% of the total membrane incorporated into the PM of an expanding cell or cell plate during cytokinesis is recycled by endocytosis (Otegui and Staehelin, 2000; Otegui et al., 2001). Therefore, the coordination of exocytic and endocytic trafficking is critical to regulate cell surface area as well as PM protein abundance and activity (Zhang et al., 2019). Components of the endocytic machinery, including clathrin, are necessary in plants and animals for the process of exocytosis (Jaiswal et al., 2009; Kawasaki et al., 2011; Larson et al., 2017). Furthermore, disruption of the exocyst complex or the function of SNAREs (exocytic soluble N-ethylmaleimide-sensitive factor attachment protein receptors), which are required for secretory vesicle tethering and fusion with the PM, respectively, inhibits endocytosis (Wu et al., 2005; Jose et al., 2015; Boehm et al., 2017; Larson et al., 2017). However, the mechanisms that coordinate the activities of the post-Golgi exocytic and endocytic trafficking are poorly understood.

We previously demonstrated that loss or partial loss of function of the CME adaptor protein complexes, AP-2 or
TPC, results in a decrease in the levels of TGN/EE-associated clathrin (Wang et al., 2016a), suggesting that post-Golgi trafficking is modulated by CME. In this study, we show that defects in post-Golgi trafficking or CME affect the recruitment of clathrin and its accessory proteins to the TGN/EE and PM, thereby providing further evidence that the processes of exocytosis and endocytosis are coordinated/coupled in plant cells.

Results

Clathrin localization to the TGN/EE depends on AP-1

To examine whether the function of AP-1 is necessary for clathrin localization to the TGN/EE, as observed in other systems (Burgess et al., 2011; Daboussi et al., 2012; Martzoukou et al., 2018), we established that subunits of the AP-1 complex and clathrin colocalize. As expected based on previous studies of the localization (Park et al., 2013; Teh et al., 2013; Wang et al., 2013a) and biochemical analysis of the AP-1 complex subunit composition (Teh et al., 2013), AP1μ2 tagged with red fluorescent protein (AP1μ2-RFP) and AP1σ2 subunits tagged with green fluorescent protein (AP1σ2-GFP) colocalized in Arabidopsis root cells at the TGN/EE (Supplemental Figure S1, A) as well as with GFP- or monomeric Kusabira-Orange fluorescent-tagged clathrin light chain 2 (CLC2-GFP or CLC2-mKO) (Supplemental Figure S1, B and C).

Next, we examined whether AP-1 is required for clathrin recruitment to the TGN/EE. For these studies, we used the previously characterized loss-of-function Arabidopsis ap1μ2 (hapless13) mutant (Park et al., 2013; Wang et al., 2013a, 2016b) as well as the ap1σ1-1 and ap1σ2-1 T-DNA insertion mutant lines, in which the expression of genes encoding AP1σ subunit isoforms, AP1σ1 and AP1σ2 is disrupted (Supplemental Figure S2), ap1σ-1 and ap1σ-2 seedlings, as well as seedlings with a single functional copy of either AP1σ1 or AP1σ2 (ap1σ1-1/ap1σ1, ap1σ2-1 and ap1σ-1 ap1σ-2/ap1σ2) exhibited no discernable phenotypes. By contrast, homozygous ap1σ1-1 ap1σ2-1 double mutant seedlings displayed a reduction in growth on half-strength Murashige and Skoog (MS) agar plates (Supplemental Figure S2, C) and loss of viability on soil after 2–3 weeks (Supplemental Figure S2, D), similar to the phenotype of ap1μ2 null mutants (Park et al., 2013; Teh et al., 2013; Wang et al., 2013a). The observed developmental defects in ap1σ1-1 ap1σ2-1 were rescued by the introduction of a 35Spro:AP1σ2-GFP transgene (Supplemental Figure S2, E). These data indicate the functional redundancy of AP1σ1 and AP1σ2 in plant growth and development.

As detected by quantitative immunofluorescence (IF) microscopy using affinity-purified anti-CLC1 and anti-CHC antibodies (Wang et al., 2013b; Supplemental Table S2), loss of AP1μ2, or AP1σ1 and AP1σ2, resulted in a ~40%–60% reduction in the levels of TGN/EE-associated clathrin (Figure 1, A–D). Likewise, live-cell imaging showed that fluorescence intensity at the TGN/EE derived from the Arabidopsis CHC isoforms, CHC1 and CHC2, and the three CLC isoforms, CLC1, CLC2, and CLC3, tagged with GFP, decreased by ~55%–60% (CHCs) and ~50%–60% (CLCs) in ap1μ2 relative to the WT (Supplemental Figure S3, A–O).

Given that homozygous ap1μ2 and ap1σ1 ap1σ2 double mutant seedlings exhibited major defects in growth and development (Supplemental Figure S2), we examined whether clathrin localization is also affected in less severe ap-1 RNA interference (RNAi) lines. For these studies, we used previously described transgenic lines in which the levels of AP1μ2 transcript are reduced by the expression of an AP1μ2-RNAi construct under the control of the INNER NO OUTER (INO) promoter (INOpro:AP1μ2-RNAi) (Wang et al., 2016b).

Although INOpro:AP1μ2-RNAi was previously reported to specifically downregulate the levels of AP1μ2 mRNA in the outer integuments of developing ovules (Wang et al., 2016b), we identified two independent INOpro:AP1μ2-RNAi transgenic lines (AP1μ2-RNAi-1 and AP1μ2-RNAi-2) that showed a ~60% reduction in the expression of AP1μ2 mRNA relative to WT (Supplemental Figure S4, A). Although not as severe as homozygous ap1μ2 and ap1σ1 ap1σ2 double mutants, the AP1μ2-RNAi-1 and AP1μ2-RNAi-2 lines displayed defects in root growth (Park et al., 2013; Teh et al., 2013; Wang et al., 2013a; Supplemental Figure S4, B and C). Nevertheless, similar to homozygous ap1μ2 and ap1σ1 ap1σ2 double mutants, downregulation of AP1μ2 mRNA levels resulted in a significant loss of TGN/EE-associated GFP-tagged CLC3 and CHC2 in root cells (Supplemental Figures S4, D–G). In addition to the reduction in TGN/EE-associated clathrin, in our homozygous ap1μ2 mutant lines (Figure 1, A–D and Supplemental Figures S3, A–O, S4, D–G). Further analyses of the effects of loss of AP-1 on CME are described below.

Loss of AP-1 causes differential perturbation of TGN/EE-resident protein localization

The loss of TGN/EE-associated clathrin suggests that TGN/EE organization and/or function may be altered in ap-1 mutants. In agreement, the ultrastructure of the Golgi apparatus was previously shown to be altered in ap1μ2 mutant cells (Park et al., 2013). Therefore, to further probe the integrity of the Golgi apparatus in ap-1 mutants, we investigated the localization of the TGN/EE-resident proteins vacuolar ATPase subunit a1 (VHAa1), SYNTAXIN 41 (SYP41) and SYP61, which are required for post-Golgi trafficking, tagged with the fluorescent proteins RFP (VHAa1-RFP), GFP (SYP41-GFP), and cyan fluorescent protein (CFP-SYP61) (Uemura et al., 2004; Dettmer et al., 2006; Robert et al., 2008; Drakakaki et al., 2012).
Moreover, treatment of ap1μ2 mutants with the vesicle trafficking inhibitor brefeldin A (BFA), which causes aggregation of TGN/EE membranes in plant cells (Geldner et al., 2003; Lam et al., 2009), resulted in differential accumulation of VHAa1-RFP, SYP41-GFP, and SYP61-CFP in BFA bodies. While we observed no detectable differences in the formation of BFA bodies containing VHAa1-RFP in WT or ap1μ2 mutant root cells treated with 50 μM BFA for 10 min (Figure 2, E–G), we did notice a significant reduction in the size of SYP41-GFP and SYP61-CFP labeled BFA bodies in ap1μ2 mutant cells relative to WT (Figure 2, L–N, S–U). These results demonstrate that localization of the TGN/EE-resident proteins SYP41 and SYP61, but not that of VHAa1, is affected in ap1μ2 mutants, suggesting that loss of AP-1 has differential effects on the localization of TGN/EE-resident proteins and/or the organization of this compartment.

Disruption of clathrin or AP-1 impairs post-Golgi trafficking to the PM

The TGN/EE is involved in post-Golgi trafficking of soluble and membrane cargoes to the vacuole and PM (Lam et al., 2007; Gendre et al., 2015). To address whether the loss of TGN/EE-associated clathrin or AP-1 affected exocytosis, we assessed the intracellular trafficking and delivery of the soluble secreted marker sec-GFP (Zheng et al., 2004) to the apoplast in the root cells of mutants defective for subunits of clathrin or the AP-1 complex. Consistent with the secretion defects previously observed in chc1-2 and chc1-3 (Larson et al., 2017) and ap1μ2 mutants (Park et al., 2013), sec-GFP accumulated within intracellular compartments in chc1, chc2, chc1/CHC1 chc2, and chc2 chc3 mutants as well as in ap1μ2 and ap1σ1-1 ap1σ2-1 double mutants (Figure 3, A–C, F–L). Notably, the levels of intracellular sec-GFP fluorescence signal in mutants defective for clathrin or AP-1 subunits were similar to those observed in WT seedlings treated with endosidin 2 (ES2), an inhibitor of exocytosis (Zhang et al., 2016; Figure 3, D–F).

We previously showed that clathrin is necessary for the recycling of internalized PM proteins to the PM (Wang et al., 2013b). Therefore, we examined whether AP-1 was required for the recycling of the PM marker proteins auxin efflux transporter PIN-FORMED 2 (PIN2-GFP) and RARE COLD-INDUCIBLE 2A (RCI2A-GFP). To monitor recycling, WT and mutant root cells expressing PIN2-GFP or RCI2A-GFP were treated for 60 min with the reversible trafficking inhibitor BFA, in the presence of the protein synthesis inhibitor cycloheximide (CHX) to entrap the PM internalized marker proteins in BFA bodies. Following BFA washout, we analyzed the loss of BFA body-associated PIN2-GFP or RCI2A-GFP by confocal microscopy and quantitative image analysis. As shown in Supplemental Figure S5, recycling to the PM of PIN2-GFP and RCI2A-GFP was reduced in the
ap1μ2 and ap1σ1-1 ap1σ2-1 mutants, relative to the WT. These results together suggest that clathrin and AP-1 are necessary for exocytic trafficking of secreted and recycling of endocytosed proteins to the PM.

Defects in AP-1 affect endocytosis

In addition to defects in secretion, ap1μ2 mutants have also been reported to exhibit delayed internalization of the lipophilic PM tracer dye FM4-64 (Teh et al., 2013), suggesting that AP-1 is required for endocytosis. However, other studies have reported contradictory results, including no significant defects in internalization (Park et al., 2013) and increased FM dye internalization (Wang et al., 2013a) in ap1μ2 mutants versus WT. Therefore, we sought to reevaluate whether AP-1 function is necessary for endocytosis of FM4-64.

To assess FM4-64 uptake in WT and ap-1-mutant cells, we initially loaded the dye into the PM of seedling root cells under low temperature conditions (4 °C), which inhibit CME (Bolte et al., 2004; Dhonukshe et al., 2007). Prior to shifting the cells to 23 °C, the initial FM4-64 fluorescence signal intensity was identical at the PM of WT, ap1μ2, and ap1σ1-1 ap1σ2-1 root epidermal cells, with no detectable intracellular accumulation of the dye (Figure 4, A and B). Following a 6-min incubation at 23 °C during which CME is active, intracellular FM4-64-labeled puncta became readily apparent in WT root cells (Figure 4, C and D). By contrast, the ratio of intracellular/PM FM4-64 signal was significantly lower in ap1μ2, and ap1σ1-1 ap1σ2-1 mutant root cells relative to WT (Figure 4, C and D). Internalized FM4-64 in both WT and ap1μ2 root cells predominantly colocalized with CLC2-GFP (Supplemental Figure S6), suggesting that loss of AP-1 function reduces FM4-64 endocytosis, but not its delivery to the TGN/EE. Similarly, partial downregulation of AP1 function through use of the AP1 RNAi-1 and AP1 RNAi-2 lines resulted in a reduction of FM4-64 internalization relative to WT, although not as severely as in ap1μ2-mutant lines (Supplemental Figure S4, H–K).

Disruption of AP-1 or exocytosis affected clathrin and CME accessory protein localization to the PM

Consistent with the reduction in FM4-64 internalization seen in ap-1 mutants (Figure 4, A–D and Supplemental Figure S4, H–K), loss of AP-1 function resulted in a decrease in the levels of CHC and CLC isoforms at the PM (Figure 1, A–D and
roots mock-treated (DMSO; D) or treated with ES2 (E) for 4 h. G–L, Sec-GFP secretion in the WT (G) and an individual root. **

Figure 3 Impaired secretion of Sec-GFP in ap-1, clc, chc mutants and ES2-treated roots. A–F, Sec-GFP secretion in WT (A) and ap-1 (B, C) roots or in WT roots mock-treated (DMSO; D) or treated with ES2 (E) for 4 h. G–L, Sec-GFP secretion in the WT (G) and clc2 clc3 double mutant (H), chc1-2 mutant (I), chc2-2 mutant (J), and chc1/CHC1 chc2 mutant (K) roots. F and L, Boxplot representation of Sec-GFP fluorescent signal intensities (in arbitrary units, a.u.) in the roots (F, n = 32, 12, 14, 16, and 15 measurements, respectively; L, n = 30, 29, 30, 27, and 24 measurements, respectively). Each data point represents an individual root. **P < 0.001, ***P < 0.0001 (Student’s t test; compared with the WT or the mock). Bars = 50 µm.

Supplemental Figures S3, A–O, S4, D–G). Therefore, we examined whether the PM localization of endocytosis accessory proteins was also affected in loss-of-function ap-1 mutants by quantitative IF microscopy. Accordingly, we conducted immunolocalization using affinity-purified anti-AP2µ, anti-AP2σ (Wang et al., 2016a), anti-TPLATE, and anti-TML antibodies (see Methods and Supplemental Table S2 for description of antibodies). The specificity of anti-TPLATE/TML antibodies was confirmed by IF analysis (Supplemental Figure S7, A) on seedling roots in which the levels of TPLATE and TML mRNA were lowered through expression of the estradiol-inducible artificial microRNA (amiR) constructs amiR

To further examine how loss of AP-1 affects the PM localization of clathrin and endocytic accessory factors at the PM (Figure 1 and Supplemental Figures S3, A–O, S4, D–G, L–O), the densities of foci containing CLC3-GFP (Figure 5, A and B; regular/small foci; ~46% reduction), AP2µ-YFP (Figure 5, C and D; ~38% reduction), and TML-YFP (Figure 5, E and F; ~51% reduction) were visibly reduced in the ap1µ2 mutant relative to the WT. However, the frequency distribution and average lifetime of PM-associated CLC3-GFP, AP2µ-YFP, and TML-YFP foci in ap1µ2 were similar to the WT (Figure 5, G–O and Supplemental Movies S1–S3). The number of CLC3-GFP-labeled TGN/EE, which appear as large bright structures when they enter the cortical TIRF imaging field (Supplemental Movie S1), was also significantly reduced (Figure 5, A and B; large structure), likely due to the decreased levels of TGN/EE-associated clathrin in the ap1µ2 mutant relative to WT cells (Figure 1, A–D and Supplemental Figures S3, A–O, S4, D–G).

We corroborated the reduction in the levels of membrane-associated clathrin and clathrin-associated factors in ap1µ2 relative to WT seedlings by quantitative immunoblot analysis. We detected no significant differences in the abundance of CLC1-3, CHC1-2, AP2µ, or AP2σ in total protein extracts from ap1µ2 and WT seedlings (Figure 6). Likewise, the levels of TPLATE, TML, DRP1A, and DRP1C were similar in protein lysates from WT and ap1µ2 seedlings accumulating GFP/YFP-tagged fusions of these proteins (Figure 6 and Supplemental Figure S8, G–I). The specificity of the anti-GFP antibody is shown in Supplemental Figure S7, B. Loss of AP1µ2, however, resulted in a decrease in microsomal membrane-associated clathrin and clathrin accessory proteins, with a corresponding increase in the levels of
these proteins in the soluble protein fraction (Figure 6 and Supplemental Figure S8, G–I). We used the fractionation of the integral endoplasmic reticulum membrane marker protein sterol methyltransferase1 (SMT1) (Boutté et al., 2010) as a control to assess the samples enriched in soluble and microsomal membrane proteins from WT and ap1μ2 seedlings (Supplemental Figure S7, C). Furthermore, RT-qPCR showed that loss of AP1μ2 does not alter the expression of genes encoding clathrin subunits or clathrin accessory factors (Supplemental Figure S9). Taken together, these results suggest that the reduction in the levels of PM- and TGN/EE-associated clathrin and its accessory proteins in ap-1 mutants is likely due to defects in their membrane recruitment rather than changes in its expression levels and/or its degradation.

**Figure 4** AP-1 disruption or exocytosis/PM recycling inhibitors impair FM4-64 internalization in root cells. A–D, Confocal microscopy analysis of FM4-64 internalization in WT and ap-1 meristem epidermal root cells. Seedling roots were labeled with dye for 1 min on ice and either directly imaged (0 min) or after 6 min incubation at 23 °C. B, D, Quantitative analysis of the relative ratios of the intracellular to PM fluorescent signal of FM4-64 labeled WT and ap-1 mutant root cells after incubation for 0 or 6 min at 23 °C (B, n = 158–264 cells from 7 to 12 roots each; D, n = 129–178 cells from 13 to 18 roots each). E–N, FM4-64 uptake in WT roots either mock-treated (DMSO) or treated with ConcA, ES2, and ES16, respectively. I and N, Quantitative analysis of the relative ratios of the intracellular to PM fluorescent signal of FM4-64-labeled WT root cells after incubation for 0 or 6 min at 23 °C (I, n = 203–267 cells from 12 to 14 roots each; N, n = 226–281 cells from 12 to 15 roots each). The seedlings were pretreated with DMSO or exocytotic inhibitors for 30 min, followed by labeling with DMSO/exocytotic inhibitors and FM4-64 for 1 min on ice before immediate or delayed imaging. Arrows and arrowheads show representative FM4-64-labeled vesicles or PM-bound FM4-64, respectively. Shown are means ± SD. ***P < 0.0001 (Student’s t test; compared with the WT or the mock). Bars = 10 μm.
Figure 5  The density, but not the lifetime of PM-associated clathrin, AP-2, and TPC foci is reduced in ap-1 mutants. A–F, TIRF microscopy analysis of PM-associated foci of CLC3-GFP (A, B), AP2\(\mu\)-YFP (C, D), and TML-YFP (E, F) in WT and \(ap1\mu2\). B, D, and F, Relative densities of PM-associated foci (B, left/small foci, \(n = 63–85\) cells from 14 to 23 roots each; right/large structure, \(n = 46–50\) cells from 14 to 23 roots each; D, \(n = 112–183\) cells from 22 to 46 roots each; F, \(n = 93–101\) cells from 25 to 36 roots each). G, J, and M, Kymographs of PM-associated foci in WT and \(ap1\mu2\). H, K, and N, Frequency distribution of lifetimes for PM-associated foci in WT and \(ap1\mu2\). I, L, and O, Mean lifetimes of PM-associated foci in WT and \(ap1\mu2\) (H and I, \(n = 1,860–2,022\) foci from 12 to 13 roots each; K and L, \(n = 1,738–1,821\) foci from 10 to 12 roots each; N and O, \(n = 1,668–1,894\) foci from 11 to 13 roots each). White and black arrows indicate CLC3-GFP-labeled TGN/EE and time direction, respectively. Shown are means ± so. ***\(P < 0.0001\) (Student’s t test; compared with the WT). Bars = 5 (A, C, and E) or 2 \(\mu\)m (G, J, and M).
Pharmacological inhibitors of exocytosis cause defects in endocytosis and the recruitment of CME proteins to the PM

The results described above indicate that both exocytosis and endocytosis are impaired in ap-1 mutants, suggesting that these trafficking pathways are interconnected. To further investigate whether CME is more broadly dependent on exocytosis, we examined the effects of inhibitors that block post-Golgi/endosomal and exocytic trafficking on CME in plant cells. However, in contrast to ap-1 mutants that displayed altered TGN/EE-resident protein localization (Figure 2), there was no discernable difference between the subcellular distribution or fluorescence signal intensity of TGN/EE-associated VHAa1-RFP, SYP41-GFP, and SYP61-CFP in seedlings treated with ConcA (5 μM), ES2 (100 μM), or ES16 (200 μM) relative to mock controls (Supplemental Figure S10). Nevertheless, similar to mutants defective in AP-1 function, the internalization of FM4-64 in WT root cells treated with ConcA, ES2, or ES16 significantly decreased compared with the mock control (Figure 4, E–N).

Impaired internalization of FM4-64 with ConcA, ES2, and ES16 was associated with a decrease in the levels of clathrin at TGN/EE and PM as well as a reduction in clathrin accessory proteins at the PM (Supplemental Figures S11, S12). In root cells of WT seedlings treated with ConcA (5 μM), ES2 (40 μM), or ES16 (20 μM) for 60 min, we observed a reduction in the levels of PM-associated AP2μ-YFP/AP2σ-GFP and TPLATE-GFP/TML-YFP (Supplemental Figure S12) as well as the loss of PM- and TGN/EE-associated CLC3/-CHC2-GFP (Supplemental Figure S11), relative to the mock control. These results are supported by quantitative immunoblot analysis confirming that the levels of membrane-associated clathrin and AP-2/TPC, as well as DRP1A and DRP1C, are reduced in seedlings treated with inhibitors of exocytosis/PM recycling relative to the mock control (Supplemental Figure S13). Together, these data support a model wherein inhibition of exocytic trafficking due to the loss of AP-1 function or treatment with small molecule inhibitors of exocytosis causes impaired CME.

Inhibition of CME impairs exocytosis

To address the reciprocal question of whether inhibition of CME affects post-Golgi trafficking, we examined trafficking of PIN2-GFP and RC12A-GFP from BFA bodies in root epidermal cells lacking AP-2 or TPC subunits. Previous studies have shown that PIN2-GFP and RC12A-GFP recycling from the TGN/EE to the PM is dependent on clathrin function (Kitakura et al., 2011; Wang et al., 2013b). Similarly, the trafficking of internalized PIN2-GFP (Figure 7, A–D) and RC12A-GFP (Supplemental Figure S14, A–D) from BFA bodies following BFA removal in AP-2-deficient lines, ap2μ-1 (Wang et al., 2016a) and ap2σ-1 (Fan et al., 2013) was slower than that of WT. Likewise, estradiol-induced downregulation of TPLATE and TML transcript levels in amiR-TPLATE and amiR-TML lines (Gadeyne et al., 2014) resulted in inhibited recycling of internalized PIN2-GFP (Figure 7, E–H) and RC12A-GFP (Supplemental Figure S14, E–H) to the PM.
In addition to affecting PM protein recycling, inhibition of CME reduced the secretion of de novo-synthesized proteins. Analysis of ap-2 mutants (ap2μ-1 and ap2σ-1; Figure 8, A–D) as well as of TPC-deficient lines (amiR-TPLATE and amiR-TML in the presence of 20 μM estradiol; Figure 8, G–K), accumulating the secretory reporter sec-GFP, showed a marked increase in intracellular GFP fluorescence relative to the WT control and the mock control (Figure 8, A, G, and I), respectively. Sec-GFP secretion was not affected in WT cells treated with estradiol (20 μM; Figure 8, F), indicating that the observed inhibition of sec-GFP trafficking in the amiR-TPLATE and amiR-TML lines is specific to the inhibition of TPLATE and TML transcript levels.

Interference of CME impairs AP-1 recruitment to the TGN/EE
As shown previously, defects in the endocytic adaptor complexes AP-2 or TPC result in the loss of PM- and TGN/EE-associated clathrin (Wang et al., 2016a). To examine if AP-2 or TPC functions are likewise required for AP-1 recruitment to the TGN/EE, we analyzed the association of fluorescent protein-tagged AP1μ2 (AP1μ2-RFP) and AP1σ2 subunits...
(AP1σ2-GFP) with the TGN/EE in WT, ap-2, and tpc mutant lines (Figure 9). By live-cell imaging, we observed a reduction of TGN/EE-associated levels of AP1μ2-RFP and AP1σ2-GFP in loss-of-function ap2μ-1 (~53% and ~56% reduction, respectively) and ap2σ-1 (~68% and ~66% reduction, respectively) mutant lines relative to WT (Figure 9, A–D). Similarly, downregulation of TPLATE and TML transcript levels in estradiol-treated amiR-TPLATE and amiR-TML seedlings (Gadeyne et al., 2014; Wang et al., 2016a) resulted in a loss of TGN/EE-associated clathrin and AP-1 subunits in ap-2 and tpc mutants (Wang et al., 2016a; Supplemental Figure S15, A and C).
To confirm that downregulation of the levels of TPLATE and TML is associated with the loss of TGN/EE-associated AP-1, we performed time course analyses of TPLATE and TML mRNA levels (Supplemental Figure S15, D) and TGN/EE localization of AP1μ2-RFP and AP1σ2-GFP (Supplemental Figure S16) following estradiol treatment of amiR-TPLATE and amiR-TML lines. Relative to mock (dimethyl sulfoxide, DMSO)-treated amiR-TPLATE and amiR-TML seedlings, there was a significant decrease in TPLATE and TML transcript levels after 8, 16, and 24 h of treatment with 20 μM estradiol (Supplemental Figure S15, D). Likewise, we observed a progressive depletion of TGN/EE-associated AP1μ2-RFP and AP1σ2-GFP in estradiol-treated amiR-TPLATE and amiR-TML seedling root cells over the same time course (Supplemental Figure S16, A). Immunoblot analysis of soluble- and membrane-enriched subcellular fractions confirmed that ap2-1 loss of function mutants or downregulation of TPC subunit expression results in the loss of associated microsomal membranes and a corresponding increase in the levels of soluble CLC1 (Supplemental Figure S17, A and B), AP1μ2-RFP (Supplemental Figure S17, C and D), and AP1σ2-GFP (Supplemental Figure S17, E and F) relative to the WT. The specificity of the anti-RFP and anti-GFP antibodies for AP1μ2-RFP and AP1σ2-GFP, respectively, is shown in Supplemental Figure S7, B. By contrast, defects in AP-2 or TPC subunits did not alter the TGN/EE localization of the TGN/EE-resident marker proteins VHAa1-RFP, SYP41-GFP, or SYP61-CFP (Supplemental Figure S18).

To further explore the mechanism of how loss of AP-2/TPC subunits affects TGN/EE recruitment of clathrin and AP-1 subunits, we analyzed the dynamics of CLC3-GFP and AP1σ2-GFP association with the TGN/EE in the ap2σ-1 mutant and the amiR-TPLATE line using fluorescence recovery after photobleaching (FRAP; 60 and 120 min after photobleaching all fluorescence signals in cells of interest; for 5–6 cells/layer and 3–4 cell layers). The fluorescent signal recovery at the TGN/EE of CLC3-GFP and AP1σ2-GFP was much slower in ap2σ-1 or amiR-TPLATE than in the WT or the mock control (Figure 10). These results together suggest that loss of AP-2/TPC function impairs TGN/EE recruitment of clathrin and AP-1 subunits.

Discussion

Exocytosis and endocytosis are key to controlling the protein and lipid composition of the PM. Coupling between these trafficking pathways is therefore likely critical for the establishment and maintenance of signaling across the PM, as well as the activities of proteins involved in diverse processes at the cell surface, including nutrient uptake, cell wall biogenesis, and pathogen defense.

Rapid compensatory endocytosis in neurons and other animal cells that undergo large-scale regulated exocytosis has been well established (Watanabe et al., 2013; Houy et al., 2015; Gómez-Elías et al., 2020). Work in a variety of systems including yeast (Saccharomyces cerevisiae), Metazoa, Trypanosoma brucei, and plants has also suggested links between the machinery involved in constitutive exocytic vesicle transport, docking, and fusion with the PM and CME (Riezman, 1985; Gurunathan et al., 2000; Sommer et al., 2005; Jose et al., 2015; Johansen et al., 2016; Boehm et al., 2017; Larson et al., 2017; Ravikumar et al., 2018). In Arabidopsis, mutants defective for exocytosis also display an inhibition of endocytosis, as observed in mutants lacking subunits of the STOMATAL CYTOKINESIS DESTRUCTIVE (SCD) complex, which are required for activation of post-Golgi vesicle-associated Rab1 GTPases, and in the secretary SNARE mutant syp121 (McMichael et al., 2013; Larson et al., 2017; Mayers et al., 2017). Likewise, loss of RabH1B, a Golgi-localized Rab GTPase that functions in the trafficking of CELLULOSE SYNTHASE 6 (CESA6) to the PM, affects both the exocytosis and endocytosis of CESA6 (He et al., 2018). In addition, the cytoskeleton has been demonstrated to function in the coordination of membrane transport to and from the PM (Stamnes, 2002; Meunier and Gutiérrez, 2016; Zhang et al., 2019). For example, RICE MORPHOLOGY DETERMINANT (RMD), an actin-organizing protein, is required for exocytic and endocytic trafficking necessary for root growth (Li et al., 2014). The results presented here and in the recent study by Larson et al. (2017) indicate that additional mechanisms involving the coordinated regulation of the recruitment of clathrin and its accessory factors to the PM and TGN/EE are important for modulating exocytic and endocytic membrane transport.

Clathrin and AP-1 are required for post-Golgi trafficking in plant cells

In mammalian cells, yeast, fungi, and Toxoplasma gondii, clathrin and AP-1 functions are necessary for post-Golgi trafficking from the TGN to endosomes and in the polarized trafficking of some PM proteins (Gall et al., 2002; Jaiswal et al., 2009; Pieperhoff et al., 2013; Castillon et al., 2018; Martzoukou et al., 2018; Casler et al., 2019; Gravotta et al., 2019). Similarly, in plant cells, post-Golgi exocytic and vacuolar protein trafficking are dependent upon clathrin and AP-1 (Song et al., 2006; Park et al., 2013; Teh et al., 2013; Wang et al., 2013a, 2013b; Larson et al., 2017; Shimizu et al., 2021; Figure 3 and Supplemental Figure S5). As shown in Figure 1 and Supplemental Figures S1, S3, AP-1 is required to recruit clathrin to the TGN/EE, which serves as a major hub for the sorting and trafficking of newly synthesized proteins and endocytic cargoes to the PM and vacuole.

However, it remains to be determined whether clathrin and AP-1 function directly in the formation of clathrin-coated transport vesicles at the TGN/EE for the delivery of proteins to the vacuole and/or PM in plant cells. Alternatively, clathrin and AP-1 may be required for the maintenance of TGN/EE structure and function (Pantazopoulou and Glick, 2019), and thus loss of either clathrin or AP-1 would result in general defects in biosynthetic and endocytic cargo sorting and trafficking to/from the TGN/EE. Our results revealed that loss of function of AP1μ2 results in the significant reduction of clathrin as well as AP1σ2 recruitment to the TGN/EE (Supplemental Figure...
Furthermore, *ap-1* mutants displayed differential changes in the localization/retention of TGN/EE-resident proteins (Figure 2). Taken together, these results provide further evidence of a relationship between clathrin- and AP-1-dependent protein trafficking and the integrity of the TGN/EE (Park et al., 2013; Wang et al., 2013a). Further studies are required to understand the molecular function of clathrin and AP-1 at the TGN/EE in plants. Nevertheless, our data demonstrate that disruption in post-Golgi trafficking due to loss of TGN/EE-associated clathrin or AP-1 results in the inhibition of CME.

**Coordinated regulation of exocytic and endocytic vesicle trafficking**

Loss of AP-1 or treatment of WT seedlings with exocytosis inhibitors resulted in defects in the uptake of FM4-64 (Figure 4 and Supplemental Figure S4, H–K) and the membrane recruitment of clathrin and endocytic accessory
proteins (Figure 6 and Supplemental Figures S8, S13). However, disruption of AP-1 function and/or exocytosis did not alter the lifetimes of clathrin/AP-2/TPC foci on the PM (Figure 5 and Supplemental Movies S1–S3), suggesting that exocytosis modulates initiation but not the dynamics of CCV formation and budding during CME.

Similar to the manner in which defects in clathrin/AP-1-dependent post-Golgi trafficking negatively affect CME, disruption of CME in ap-2 or tpc mutants was associated with a reduction in the levels of TGN/EE-associated clathrin and AP-1 subunits (Wang et al., 2016a; Figure 9 and Supplemental Figure S16) as well as impaired exocytic trafficking/secretion (Figures 7, 8). Collectively, these findings indicate that the association of clathrin and its accessory factors with the TGN/EE (AP-1/clathrin) and PM (AP-2/TPC/clathrin/DRP) is coordinately regulated, thereby coupling the processes of post-Golgi and endocytic trafficking in root cells. However, the signaling mechanisms by which clathrin-dependent exocytosis and CME are coupled remain to be identified.

The proper balance and maintenance of exocytosis and endocytic membrane transport is crucial for plant growth, development, and responses to various environmental cues (Zhang et al., 2019). During cytokinesis and cell expansion, the polarized delivery of endomembrane-derived vesicles to the cell plate and PM, respectively, is balanced by the retrieval of membrane and proteins (Samuels and Bisalputra, 1999; Otegui and Staelen, 2000; Ketelaar et al., 2008). Discordance between exocytosis and endocytosis causes abnormal cell growth and morphogenesis in plants. For example, Arabidopsis chc mutants display defects in exocytosis and endocytosis and thereby develop aberrant cotyledons and embryos (Kitakura et al., 2011; Larson et al., 2017). Recent studies have also demonstrated that homeostasis of exocytosis and endocytosis is necessary for pollen tube tip growth, cold tolerance, stomatal movements, and osmotic stress response in plants (Deng et al., 2015; Zwiewka et al., 2015; Feng et al., 2016; Larson et al., 2017). The coupling of exocytosis and endocytosis may therefore be a critical point of regulation by developmental and environmental signals. Indeed, auxin and salicylic acid have been previously shown to rapidly (within 5 min of treatment) cause the dissociation of clathrin from both the TGN/EE and PM (Wang et al., 2013b, 2016a). However, the molecular mechanisms involved in the coordination of the recruitment of clathrin and its accessory factors to the PM and TGN/EE remain to be determined.

Materials and methods

Plant materials and growth conditions

The following transgenic lines and mutants were used in this study: transgenic lines RPSSAproCHC1-GFP (Dejonghe et al., 2016), RPSSAproCHC2-GFP (Ortiz-Moreaa et al., 2016), 3SSproCLC1-GFP (Wang et al., 2013b), CLC2proCLC2-GFP (Konopka et al., 2008), CLC2proCLC2-mKO (Ito et al., 2012), CLC3proCLC3-GFP (Dejonghe et al., 2016), AP1proAP1-GFP (Wang et al., 2013a), 3SSproAP1-GFP (this study), AP2proAP2-GFP (Bashline et al., 2013),AP2proAP2-GFP (Fan et al., 2013), 3SSproTPLATE-GFP (Van Damme et al., 2006), TMLproTML-YFP (Gadeyne et al., 2014), 3SSproDRP1A-GFP (Konopka and Bednarek, 2008), DRP1proDRP1GFP (Konopka et al., 2008), PIN2proPIN2-GFP (Xu and Scheres, 2005), 3SSproRCI2A-GFP (Cutler et al., 2000), 3SSprosec-GFP (Zheng et al., 2004), VHAa1proVHAa1-RFP (Dettmer et al., 2006), 3SSproSYP41-GFP (Uemura et al., 2004), and SYPSIproSYP61-GFP (Robert et al., 2008) and estradiol-inducible ESTRproamiR-TPLATE and amiR-TML lines (Gadeyne et al., 2014); clathrin mutants clc2-1 (SALK_016049; Wang et al., 2013b), clc3-1 (CS100219; Wang et al., 2013b), chc2-1 (SALK_103252 from ABRC), chc2-2 (SALK_042321 from ABRC), ap1-1 (SALK_145719 from ABRC), ap1-2-1 (SALK_049615C from ABRC), ap1-2 (CS16318 from ABRC), ap2-1 (SALK_141555; Fan et al., 2013), and ap2-1-1 (SALK_083693C from ABRC; Wang et al., 2016a).

The construct of 3SSproAP1-GFP was generated using PCR, restriction digestion, and ligation into transformation vectors and consequently transformed into the WT accession Columbia-0 (Col-0). Homozygous mutant lines, including chc (Wang et al., 2013b), ap-1 (Supplemental Table S1), ap-2 (Fan et al., 2013; Wang et al., 2016a), and tpc (Gadeyne et al., 2014) were isolated and/or identified by PCR- and RT-PCR- or ESTR-based assays, respectively. The clc2-1 clc3-1, chc1-2 chc2-2, and ap1-1-1 ap1-2-1 double mutants were generated by crossing and verified by genotyping PCR. Fluorescently tagged marker lines were crossed into the chc, ap-1, ap-2, and tpc mutants, and homozygous lines were confirmed based on their mutant phenotypes, genotyping PCR (Supplemental Table S1), and fluorescence.

Seeds were surface sterilized and stratified for 3 days at 4 °C in the dark and then sown onto half-strength MS medium with 1.5% (w/v) agar containing 20 mM ConcA, 20 mM AS2, 20 mM AS16, 20 mM CHX, and 20 mM ESTR. The resulting seedlings were grown vertically for 5 days; subsequent liquid treatment, 16-h light/8-h dark photoperiod, and 80% relative humidity provided by a mixture of cool [3,000K] and daylight. The following chemical solutions and treatments were applied: ConcA (5 mM), ES2 (40 mM), ES16 (10 mM), CHX (50 mM), and ESTR (20 mM) for stock solutions. Unless otherwise specified, all reagents were purchased from Sigma–Aldrich. DMSO was used to dissolve BFA (50 mM), ConcA (5 mM), ES2 (40 mM), ES16 (10 mM), CHX (50 mM), and ESTR (20 mM) for stock solutions. Unless otherwise indicated in the text, final working concentrations were 5 μM for ConcA, 20 μM for ES16 and ESTR, 40 μM for ES2, and 50 μM for CHX and BFA.

All pretreatment and posttreatment time durations are indicated in the text. All pretreatments and treatments were performed as previously described (Wang et al., 2016a). For induction of amiR-TPLATE and amiR-TML, surface-sterilized and stratified seeds were sown onto half-strength MS medium with 1.5% (w/v) agar containing 20 μM ESTR and then grown vertically for 5 days; subsequent liquid treatments also contained 20 μM ESTR. The resulting seedlings were used in live-cell imaging or IF analysis.
Polyclonal antibodies

Polyclonal antibodies (anti-TPLATE and anti-TML) were raised in rabbits using synthesized peptides for each protein (see Supplemental Table S2) coupled with keyhole limpet hemocyanin containing an additional N-terminal Cys (Huabio). Antibodies were affinity purified using immobilized peptide affinity columns, and their specificity was verified as shown in Supplemental Figure S7, A.

IF and live-cell confocal microscopy

Immunolocalization analysis was performed as previously described (Wang et al., 2013b, 2016a). All primary antibodies (Supplemental Table S2) used for immunolocalization were detected using Cy3-labeled anti-rabbit secondary antibodies (1:100 dilution; Sigma–Aldrich Catalog No. C2306). Images were captured using a confocal laser scanning microscope (Leica TCS SP5 AOBS). For imaging Cy3, the 543-nm line of the helium/neon laser was used for excitation, and emission was detected from 550 to 570 nm. For live-cell imaging of CFP, GFP, and YFP, the 458-, 488-, and 514-nm lines of the argon laser were used for excitation, and emission was detected from 460 to 500, from 496 to 532, and from 520 to 560 nm, respectively. For mKO and RFP imaging, the 543- and 594-nm lines of the helium/neon laser were used for excitation, and emission was detected from 560 to 600 and from 612 to 666 nm, respectively. For FM4-64 staining imaging, the 594-nm line of the helium/neon laser was used for excitation, and emission was detected from 651 to 761 nm. For quantitative measurement of fluorescence, all parameters (laser, pinhole, and gain settings) of the confocal microscope were identical among different treatments or genotypes.

To quantify the intensities of fluorescence signals at the PM or intracellular compartments, digital images of root epidermal cells within the division/transition zone were analyzed using ImageJ software (http://rsb.info.nih.gov/ij/; Schindelin et al., 2015). The intracellular signal intensity within individual cells was defined as a region of interest (ROI) subtending the PM. Nuclear regions present within the plane of focus were excluded from the intracellular ROI. To quantify the level of PM signal intensity, ROIs that encompassed the fluorescence signal of the entire cell cortex were assigned. For quantitative analysis of the recycling of internalized PM proteins entrapped in BFA bodies prior to and after BFA washout, non-BFA background signal was subtracted from the intracellular ROI before calculating the ratio of intracellular to PM fluorescence signal intensities.

Localization of TGN/EE-resident proteins after BFA treatment was examined by determining the relative BFA body area per cell. To quantify the relative BFA body area per cell, individual BFA bodies within a single optical section of an epidermal root cell were defined as ROIs and total area of the ROIs divided by the number of BFA bodies per cell was determined using ImageJ.

Confocal FRAP analysis

FRAP assays for GFP-fused proteins were performed on a spinning disk confocal microscope (Zeiss Cell Observer). A ROI (dashed boxes in Figure 10; approximately 5–6 cells/layer) selected with the zoom function was bleached for 1 min at full laser power until reaching the background levels of roots that did not accumulate GFP-fused proteins by scanning. GFP fluorescence was bleached throughout 3–4 cell layers, as monitored by scanning above and below the cells of interest as previously described (Grebe et al., 2003). For quantification of GFP fluorescence intensities before (pre-bleach) or after bleaching (after-bleach), the fluorescent signal intensities of the ROIs were measured by ImageJ. To compare the rates of GFP fluorescent signal recovery between the WT versus ap2α-1 cells or the mock versus estradiol treatments in amir-TPLATE, the fluorescent signal intensities before and 0-min after bleaching were set to 100% and 0%, respectively (Grebe et al., 2003).

Live-cell TIRF microscopy

Seedlings grown vertically were mounted on a glass slide in liquid half-strength MS medium and observed under a TIRF system with an Olympus IX-83 microscope equipped with a 100× oil-immersion objective (numerical aperture 1.49; Olympus). GFP-/YFP-fused proteins were excited with the 488-nm laser line. Time-lapse series images were taken at up to 250 images per sequence and acquired with 200-ms exposure time. Dynamic behaviors including foci densities and lifetimes of fluorescently labeled proteins of interest at the PM were tracked using the MATLAB Graphical User Interface program as previously described (Xue et al., 2018). The kymographs were generated using ImageJ.

Membrane-associated protein isolation and immunoblot analysis

Preparation of total, soluble, and microsomal membrane subcellular protein fractions was as described previously (Abas et al., 2006; Wang et al., 2016a). For immunoblot analysis, the antibodies including anti-CLC1, -CLC2, -CLC3, -CHC, -AP2μL, -AP2α (Supplemental Table S2), -GFP (1:1,000 dilution; TransGen Biotech Catalog No. HT801-01), anti-RFP (1:1,000 dilution; MBL Biotech Catalog No. M204-3), and anti-SMT1 (1:1,000 dilution; Agrisera Catalog No. AS07266) were used. Band fluorescent signal intensities were quantified as described previously (Wang et al., 2016a). Coomassie Brilliant Blue (CBB) staining served as a loading control to calculate the relative band fluorescent signal intensities (Wang et al., 2016a). All primary antibodies used in immunoblotting were detected using anti-rabbit or -mouse secondary antibodies (1:50,000 dilution; Huabio Catalog No. HA1019 or HA1009) conjugated to horseradish peroxidase and detected by a chemiluminescent-enhanced substrate kit (Thermo Scientific).

RT-PCR and RT-qPCR assays

The total RNA was isolated using the RNeasy Plant Mini Kit (Qiagen). First-strand cDNA was synthesized with the SuperScript III First-Strand Synthesis System (Invitrogen). The cDNA templates were used in PCR amplification for AP1α1 (28 cycles), AP1α2 (28 cycles), and ACTIN 2 (26
cycles; as internal control) transcripts with the gene-specific primers (Supplemental Table S1). The RT-qPCR assay was performed with gene-specific primers (Supplemental Table S3) as described previously (Wang et al., 2016a).

Accession numbers

Sequence data from this article can be found in the Arabidopsis Genome Initiative under the following accession numbers: CLC1 (At2g20760), CLC2 (At2g40060), CLC3 (At3g1890), CHC1 (At3g11130), CHC2 (At3g08530), AP1α1 (At4g35410), AP1α2 (At2g17380), AP1μ2 (At1g60780), AP2α (At1g47830), AP2μ (At5g46630), TPLATE (At3g01780), TML (At5g7460), PIN2 (At5g57090), RCI2A (At3g05880), DRP1A (At5g42080), DRP1C (At1g14830), VHAa1 (At2g28520), SYP41 (At5g26980), SYP61 (At1g28490), UBIQUITIN 7 (At2g35635), and ACTIN 2 (At3g18780).

Supplemental data

The following materials are available in the online version of this article.

Supplemental Figure S1. Co-localization analysis of AP-1 subunits and clathrin light chain, CLC2.

Supplemental Figure S2. Molecular and phenotypic characterization of ap1α1 and ap1α2.

Supplemental Figure S3. GFP-tagged CLC, CHC, and AP1α2 membrane association in ap1μ2.

Supplemental Figure S4. Developmental and endocytic defects in ap1μ2.

Supplemental Figure S5. Reduced recycling to the PM of PIN2-GFP and RCI2A-GFP in ap-1.

Supplemental Figure S6. Co-localization analysis of internalized FM-64 and clathrin at the TGN/EE in ap-1.

Supplemental Figure S7. Analysis of the specificity of antibodies used for immunodetection studies.

Supplemental Figure S8. Reduction in levels of PM-associated DRP1-GFP in ap-1.

Supplemental Figure S9. RT-qPCR analysis of transcriptional levels of endogenous and exogenous core components of CME machinery in ap1μ2.

Supplemental Figure S10. Effects of exocytosis/PM recycling inhibitors on TGN/EE marker localization.

Supplemental Figure S11. Inhibitors of exocytosis/PM recycling cause a reduction in levels of membrane-associated clathrin.

Supplemental Figure S12. Inhibitors of exocytosis/PM recycling decrease the levels of PM-associated AP-2/TPC subunits.

Supplemental Figure S13. Immunoblot analysis of effects of exocytosis/PM recycling inhibitors on membrane recruitment of clathrin, AP-2/TPC, and DRP1.

Supplemental Figure S14. Reduced recycling to the PM of RCI2A-GFP in ap-2 and tpc.

Supplemental Figure S15. RT-qPCR analysis of transcriptional levels of AP-1 and clathrin in ap-2 and tpc.

Supplemental Figure S16. Downregulation of TPLATE or TML mRNA expression reduced the levels of TGN/EE-associated AP-1.

Supplemental Figure S17. Membrane recruitment of clathrin and AP-1 is reduced in ap-2 and tpc mutant seedlings.

Supplemental Figure S18. Localization of TGN/EE markers in ap-2 and tpc.

Supplemental Data Set S1. Summary of statistical analyses.

Supplemental Table S1. PCR primer sequences for genotyping, RT-PCR, and cloning.

Supplemental Table S2. Information for antibodies.

Supplemental Table S3. RT-qPCR primer sequences.

Supplemental Movie S1. TIRF imaging of CLC3-GFP in WT and ap1μ2.

Supplemental Movie S2. TIRF imaging of AP2μ-YFP in WT and ap1μ2.

Supplemental Movie S3. TIRF imaging of TML-YFP in WT and ap1μ2.

Acknowledgments

We dedicate this manuscript in memory of our mentor, friend, and colleague, Prof. Jianwei Pan. His passion for science, kindness, and generosity remains as an inspiration to us all. We are grateful to Yang Zhao, Liping Guan, Yahu Gao, and Haiyan Li (Core Facility for Life Science Research, Lanzhou University) as well as Yaning Cui and Yucong Zhou for their technical assistance, and the Arabidopsis Biological Resource Center (ABRC) for seed stocks.

Funding

This work was supported by grants to J.P. from the National Natural Science Foundation of China (Nos. 91754104, 31820103008, and 31670283); grants to S.Y.B. from the National Science Foundation (Nos. 1121998 and 1614915), and grants to X.Y. from the Fundamental Research Funds for the Central Universities (lzujbky-2019-it13 and lzujbky-2020-kb05).

Conflict of interest statement. The authors declare no conflict of interest.

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