Proteomic characterization of isolated Arabidopsis clathrin-coated vesicles reveals evolutionarily conserved and plant-specific components

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Abstract

In eukaryotes, clathrin-coated vesicles (CCVs) facilitate the internalization of material from the cell surface as well as the movement of cargo in post-Golgi trafficking pathways. This diversity of functions is partially provided by multiple monomeric and multimeric clathrin adaptor complexes that provide compartment and cargo selectivity. The adaptor-protein assembly polypeptide-1 (AP-1) complex operates as part of the secretory pathway at the trans-Golgi network (TGN), while the AP-2 complex and the TPLATE complex jointly operate at the plasma membrane to execute clathrin-mediated endocytosis. Key to our further understanding of clathrin-mediated trafficking in plants will be the comprehensive identification and characterization of the network of evolutionarily conserved and plant-specific core and accessory machinery involved in the formation and targeting of CCVs. To facilitate these studies, we have analyzed the proteome of enriched TGN/early endosome-derived and endocytic CCVs isolated from dividing and expanding suspension-cultured Arabidopsis (Arabidopsis thaliana) cells. Tandem mass spectrometry analysis results were validated by differential chemical labeling experiments to identify proteins co-enriching with CCVs. Proteins enriched in CCVs included previously characterized CCV components and cargos such as the vacuolar sorting...
receptors in addition to conserved and plant-specific components whose function in clathrin-mediated trafficking has not been previously defined. Notably, in addition to AP-1 and AP-2, all subunits of the AP-4 complex, but not AP-3 or AP-5, were found to be in high abundance in the CCV proteome. The association of AP-4 with suspension-cultured Arabidopsis CCVs is further supported via additional biochemical data.

**Introduction**

Vesicle trafficking is critical for the exchange of materials between the various biochemically and functionally distinct compartments of the biosynthetic/secretory and endocytic pathways. In particular, the trafficking of vacuolar proteins and the polar localization of plasma membrane (PM) proteins is critical for nutrient uptake, pathogen response, and organismal homeostasis. Fundamental to the process of vesicle trafficking is the assembly of cytosolic coat protein complexes, which cluster cargo and generate the membrane curvature necessary for a budding vesicle (Bonifacino and Glick, 2004; Robinson, 2015). The distinctive geometric lattice that surrounds clathrin-coated vesicles (CCVs) is composed of clathrin triskelia, which in turn are composed of clathrin heavy chain (CHC) and clathrin light chain (CLC) subunits.

Since their initial discovery in metazoans and plants (Gray, 1961; Roth and Porter, 1964; Bonnett and Newcomb, 1966), CCVs have been demonstrated to function in endocytosis (Robinson, 2015; Reynolds et al., 2018) and post-Golgi trafficking (Orci et al., 1985; Hirst et al., 2012). The highly choreographed multi-step process of clathrin-mediated endocytosis (CME) requires clathrin and a large number of endocytic accessory proteins (EAPs) that function at specific sites on the PM (i.e. clathrin-coated pits (CCPs)) to mediate cargo recruitment, membrane invagination, and scission/severing of the nascent CCV from the PM and their uncoating prior to fusion with endosomes (Taylor et al., 2011; Kaksonen and Roux, 2018; Wu and Wu, 2021). Our current understanding of the complex network of proteins and lipids required for post-Golgi trafficking and endocytosis in plants largely comes from yeast and mammalian systems and the use of various biochemical, proteomic, genetic, and advanced quantitative live-cell imaging approaches. By comparison, mechanistic insight into clathrin-dependent membrane trafficking in plants remains limited.

CCVs facilitate multiple intracellular trafficking pathways via the action of adaptor protein complexes that are tasked with pathway-specific cargo recognition and clathrin recruitment at the PM and endomembrane compartments. The first clathrin adaptors to be identified were the assembly polypeptide (AP)-2 and AP-1 heterotetrameric complexes (Pearse and Robinson, 1984; Keen, 1987) that interact with protein cargo and specific phospholipids residing at the PM and the trans-Golgi Network (TGN), respectively (Robinson, 2015). These functions are largely conserved in plants, including the central role of AP-2 in endocytosis (Bashline...
et al., 2013; Di Rubbo et al., 2013; Fan et al., 2013; Kim et al., 2013) and of AP-1 in trafficking at TGN/early endosomes (TGN/EE) (Song et al., 2006; Park et al., 2013; Teh et al., 2013). The latter’s function is somewhat more complex as the plant TGN/EE appears to comprise numerous distinct sub-compartments with varying morphologies and functions in contrast to its metazoan counterpart (Dettmer et al., 2006; Viotti et al., 2010; Kang et al., 2011; Rosquete et al., 2018; Heinze et al., 2020; Shimizu et al., 2021). Moreover, an increasing number of studies have demonstrated that these core CCV trafficking proteins have evolved to accommodate processes critical for plant growth and development against the differing biophysical properties of plant cells, including cell wall formation, cytokinesis, and pathogen response (McMichael and Bednarek, 2013; Zhang et al., 2015; Gu et al., 2017; Ekanayake et al., 2019).

In addition to AP-2 and AP-1, three additional AP complexes, AP-3 (Dell’Angelica et al., 1997; Simpson et al., 1997; Zwikwia et al., 2011), AP-4 (Dell’Angelica et al., 1999; Hirst et al., 1999), and AP-5 (Hirst et al., 2013), mediate post-Golgi endomembrane trafficking in multicellular eukaryotes. However, relative to AP-1 and AP-2, their interaction with the clathrin machinery in mammalian and other systems is less defined. Proteomic analyses of mammalian CCVs suggest that of the five evolutionarily conserved adaptor complexes, only AP-1 and AP-2 were found to be associated with the clathrin coat (Blondeau et al., 2004; Borner et al., 2012). Recent super-resolution microscopy of plant cells showed a lack of colocalization between AP4 and clathrin (Shimizu et al., 2021). Lastly, the association of AP-5 with clathrin in metazoans and plants remains undetermined (Sanger et al., 2019).

Plants have shown evolutionary divergence in their key EAPs compared to other model systems including, for example, the TPLATE complex (TPC), which likely evolved in azoans and which is essential for plant CME (Gadeyne et al., 2014; Zwiewka et al., 2011; Narasimhan et al., 2020; Ekanayake et al., 2021). Undifferentiated suspension-cultured plant cells have been employed as a model system to examine the mechanistic details of their function(s) in CME (Konopka et al., 2011; Narasimhan et al., 2020; Ekanayake et al., 2021). Various plant uncoating factors have been identified, and the mechanistic details are under investigation (Robinson, 2015; Adamowski et al., 2018).

Trafficking factors including Rab and ADP-ribosylation factor (ARF)-related small GTPases (Jürgens et al., 2015; Lipatova et al., 2015; Takemoto et al., 2018), vesicle tethering complexes (Jürgens et al., 2015; Takemoto et al., 2018), and soluble N-ethylmaleimide-sensitive factor adaptor protein receptors (SNARE) proteins (Yun and Kwon, 2017) add complexity to the regulation of clathrin-mediated trafficking pathways.

Defining the protein complement of post-Golgi and endocytic CCVs will further enhance our understanding of the evolutionarily conserved and plant-specific core and accessory machinery involved in the formation and targeting of plant CCVs. Organelle proteomics is a useful tool for such studies as it can provide a global view of the protein content of a particular organelle, thereby placing known and un-annotated gene products in a functional context. Several studies have employed mass spectrometry (MS) to answer questions of mammalian CCV content including analyses of rat brain CCVs to examine their role in synaptic vesicle recycling (Blondeau et al., 2004; Takamori et al., 2006) and of those isolated from HeLa cells (Borner et al., 2006, 2012). In addition to the identification of previously unknown CCV components, the relative abundance of proteins across CCVs isolated from rat brain and liver has been examined to interrogate pathway-specific proteins such as AP-1 and AP-2 subunits (functioning in secretion and endocytosis, respectively) and the proportion of CCVs involved in post-Golgi and endocytic trafficking in different cell types (Girard et al., 2005). In plants, subcellular fractionation coupled with tandem MS has also been utilized to elucidate the protein content of endomembrane compartments including Arabidopsis Golgi cisternae (Parsons et al., 2012, 2019; Okekeogbu et al., 2019), vacuoles (Carter et al., 2004), syntaxin of plants 61 (SY6P1)-positive TGN compartments (Drakakaki et al., 2012), and multiple compartments of the Arabidopsis endomembrane system (Heard et al., 2015). Here, we report the proteomic assessment of CCVs isolated from undifferentiated Arabidopsis suspension cultured cells using tandem MS and quantitative immunoblotting to elucidate the ensemble of proteins underlying clathrin-mediated trafficking in plants and to better understand the similarities and differences among trafficking protein pathways within eukaryotes.

Results and discussion

CCV isolation

In plants, CCVs mediate trafficking of secretory and endocytic cargo necessary for cell expansion, cytokinesis, nutrient uptake, and pathogen immunity pathways underlying growth, morphogenesis, and defense in a variety of developmentally distinct cell types (Reynolds et al., 2018; Ekanayake et al., 2019). Undifferentiated suspension-cultured plant cells
offer, however, several benefits as a biological sample source for proteomic analysis of plant CCVs as they: (1) display high levels of division and expansion, processes that require a large flux of vesicular trafficking; (2) are easily scaled to provide large amounts of material; and (3) when grown under constant conditions, provide a population of cells uniform in cell type and development between biological replicates. With these features in mind, the CCVs used for proteomic analysis in this study were isolated from 3- to 4 day-old T87 suspension-cultured Arabidopsis cells (Axelos et al., 1992). In addition, the availability of the Arabidopsis genome sequence and in silico proteome (Arabidopsis Genome, 2000; Berardini et al., 2015) as well as the T87 transcriptome datasets (Stolc et al., 2005) facilitate the assessment of the enrichment or depletion of proteins of interest in the CCV proteome.

For proteomic analyses, plant CCVs were isolated under pH conditions that inhibit clathrin cage disassembly using a fractionation scheme that includes differential, rate-zonal centrifugation, and a final equilibrium deuterium/Ficoll gradient as described previously (Reynolds et al., 2014). Prior to MS analysis, the composition and quality of CCV preparations were assessed by morphological analysis using transmission electron microscopy (TEM) and immunoblotting for protein markers of the secretory and endocytic pathways. TEM analysis (Figure 1A) of the enriched CCV samples revealed that 80% ± 7.6% of the vesicles from four independent CCV preparations used for MS analysis displayed the characteristic geometry of clathrin coats with an average diameter of 75 ± 5.3 nm. Transmission electron micrographs of enriched CCVs at higher resolution show striking symmetry of the coat (Figure 1, B and C; Supplemental Figure S1).

**Figure 1** TEM of purified plant CCVs. A, Negative stain transmission electron micrographs of a typical CCV preparation. Uncoated vesicles are indicated by arrowheads and coated vesicles by arrows. B and C, Positive stain scanning transmission electron micrographs of CCVs. Scale bars (clockwise, starting top): 100, 20, and 50 nm.

CCV MS/MS sample preparation and analysis
To establish a comprehensive understanding of the protein composition of plant CCVs, two independent, parallel proteomic workflows were performed on enriched CCV protein samples (Supplemental Figure S2). In the first methodology, CCVs isolated from four independent preparations were resolved via one-dimensional sodium dodecyl-sulfate polyacrylamide gel electrophoresis (1D SDS-PAGE) before in-gel digestion of proteins by trypsin and subjection to liquid chromatography/tandem mass spectrometry (LC/MS-MS). To account for differences in protein sampling related to 1D SDS–PAGE sample fractionation and the use of detergents, we used a second methodology in which three independent CCV preparations were denatured in a urea buffer prior to trypsin digestion in-solution and separation of recovered peptides by LC/MS-MS.

Coomassie staining of a representative enriched CCV fraction separated by 1D SDS-PAGE is shown in Figure 2A. Gel slices 2, 6, and 8 contain bands of high protein abundance (intense Coomassie staining), which migrate at rates corresponding to the molecular weights of the clathrin coat proteins. To confirm the identity of the specific heavy or light chain clathrin isoforms in each polyacrylamide gel slice, we plotted the unique spectral counts as a percentage of unique spectral counts for that isoform across the entire SDS–PAGE gel and showed that the protein abundances in gel slices 2, 6, and 8 in Figure 2A corresponded to the presence of clathrin subunits (Figure 2B). Arabidopsis encodes two CHC isoforms, CHC1 and CHC2, of approximately the same mass (193 kDa) which share >90% sequence identity (Kitakura et al., 2011) and are transcribed at comparable levels in T87 suspension cultured cells (Stolc et al., 2005). In contrast to CHC, the three Arabidopsis CLC isoforms share...
only ~55% sequence identity (Wang et al., 2013) and vary in predicted mass (37.2, 28.8, and 29.1 kDa, CLC1-3, respectively). The spectral count and SDS–PAGE data confirmed that CHC contributes to the high protein abundance of gel slice 2 (predicted molecular weight of CHC1 and CHC2, 193 kDa), and that CLC contribute to the protein abundance of gel slices 6 and 8 (predicted molecular weight of CLC1, 37 kDa; of CLC2 and CLC3, 29 kDa). The identities of the other CCV-associated proteins in Figure 2A with abundances below the limits of detection by Coomassie staining were determined by proteomic and immunoblot analyses addressed below.

To define the CCV proteome, protein or protein group assignments were considered true if at least two unique peptides denoting the protein or protein group were found in at least two biological replicates based on a 1% false discovery rate (FDR) protein and peptide threshold (Supplemental Figures S2, B and C). Processed spectra from the first methodology were matched against the Arabidopsis protein sequence database using the Mascot searching algorithm (Perkins et al., 1999), while in the second methodology, MS/MS spectra were analyzed with the MaxQuant software package.

Following these criteria, protein assignments from the four independent CCV preparations analyzed by the first methodology (in which CCVs were separated by 1D SDS–PAGE) comprised a list of 3,745 proteins (Supplemental Data Set 1), the vast majority of which (~70%) were in relatively low abundance (<50 spectral counts across four replicates). Total spectral counts have previously been used as an approximation of overall protein abundance in biological samples for those proteins with a reasonably high (>10–20) total number of counts (Lundgren et al., 2010). Accordingly, we compared total counts over four independent CCV preparations without normalization as an approximation of overall protein abundance within CCVs and focused our analysis on those proteins with the highest associated spectral count totals. The proteomic data from CCVs fractionated by 1D SDS–PAGE and analyzed by tandem MS is presented in Supplemental Data Sets 1, 4, and 5. Protein assignments obtained from the second CCV methodology resulted in a list of 1,981 protein groups (Supplemental Data Set 2). Intensity-based absolute quantitation (iBAQ) values were used to sort protein groups within this dataset, as this method of label-free quantification has been previously judged to be a metric for protein abundances within biological samples and enables comparison of these abundances (Nagaraj et al., 2011; Schwanhäusser et al., 2011; Arike et al., 2012). The data from this second methodology are presented in Supplemental Data Sets 2 and 4.

Previous proteomic studies of CCVs purified from mammalian tissue have found that clathrin light and heavy chains predominantly exist in a stoichiometric 1:1 ratio. However, clathrin subunits were also found in nonstoichiometric ratios depending on the tissue and species source of the CCV sample (Blondeau et al., 2004; Girard et al., 2005; Borner et al., 2012). We used iBAQ levels to analyze the abundances of clathrin heavy and light chain subunits and determine the ratio of subunits within the triskelion independently of immunoblotting-based methods. We summed the iBAQ values for CHC and CLC subunits from each of three independent CCV preparations and calculated the ratio between these values as the measure of CLC:CHC per replicate of 1.24, 2.05, and 1.07 (Figure 2C; Supplemental Data Set 2). These values generally support a 1:1 ratio but suggest that, in some CCV preparations from plant cells, CLC subunits are in excess.
peptides, which results in identical peptides treated in this way differing by 4 Da. Accordingly, a quantitative ratio of an individual peptide’s abundance in the DFGL relative to the final enriched CCV preparation, as represented by spectral counts, can be derived.

DFGL and CCV fractions from two independent CCV preparations were separated by 1D SDS–PAGE prior to gel sectioning and in-gel digestion with trypsin. Peptides from both fractions were recovered and treated with dimethyl reagents as previously described (Boersema et al., 2009). In the second replicate, heavy (deuterated) and light labels applied to CCV and DFGL samples were swapped to control for potential discrepancies resulting in the identification of proteins that were enriched in \((\text{CCV}:\text{DFGL spectral count ratio} \geq 2.0)\) or depleted from \((\text{DFGL}:\text{CCV spectral count ratio} \geq 2.0)\) the purified CCVs (Figure 3; Supplemental Data Set 3). Cross-referencing these datasets yielded a core set of 781 proteins in the enriched CCV purification sample as detected by all three proteomic workflows, 213 (27%) of which had a CCV:DFGL spectral count ratio \(\geq 2.0\), that is, were enriched more than two-fold in the final preparatory step resulting in purified CCVs (Figure 3; Supplemental Data Set 3).

In previous studies, immunoblotting has been used to validate differential centrifugation as a means to purify CCVs from plant cells by confirming the depletion of markers of the endoplasmic reticulum (ER) and Golgi, as well as of other organelles including plastids, peroxisomes, and mitochondria, in addition to confirming the enrichment of CCV associated proteins (McMichael et al., 2013; Reynolds et al., 2014). However, the fold enrichment or depletion of these proteins relative to their initial abundance in the lysate \((S_{0.1})\) fraction has not been quantified. Here, we performed quantitative immunoblotting of equal amounts of protein from steps throughout the CCV purification process, including the DFGL and CCV fractions, to establish the fold enrichment and depletion of CCV mediated trafficking-associated and unassociated proteins, respectively, relative to the lysate (Figure 4). We then used the average fold enrichments between the DFGL and CCV fractions from quantitative immunoblotting of three independent CCV preparations (apart from immunoblotting for DRP2, where \(n = 2\)) to corroborate the values obtained by the dimethyl labeling proteomic workflow (Table 1).

The expected enrichment of CCV-associated proteins, such as clathrin coat proteins CHC and CLC2 and subunits of AP-1 and AP-2 complexes, as well as the cell plate, TGN, and putative CCV cargo marker, KNOLLE (Dhonukshe et al., 2010), is shown in Figure 4A. To demonstrate the removal of non-CCV trafficking-associated proteins from the purified CCVs, the depletion of cFBPase and SEC12 proteins, markers for the cytosol and ER, respectively, between the \(S_{0.1}\) fraction and final CCV sample, is shown in Figure 4B. We also quantitated the intensity of the proteins in Figure 4, A and B at the DFGL and CCV steps of the CCV purification scheme and compared these values to those obtained in the dimethyl labeling experiment (Table 1). The average fold enrichment
values between the CCV and DFGL samples for the proteins in Figure 4, A and B as determined by immunoblotting or dimethyl labeling are compared in Table 1, which shows that the general trends of enrichment of CCV-associated proteins and depletion of markers of subcellular compartments not associated with CCV-mediated trafficking away from the final CCV sample are consistent across both methods. These quantitative immunoblotting data support the use of the dimethyl labeling proteomic dataset (Supplemental Data Set 3) as a tool for researchers investigating proteins of interest and the strength of potential connections to CCV-mediated trafficking processes.

To assess the contributions of subcellular organelles to the complement of the 546 proteins detected by the first methodology that were depleted at least two-fold during the final CCV purification step, their predicted subcellular localizations were determined using the SUBcellular Arabidopsis consensus (SUBAcon) bioinformatics tool, an algorithm that integrates experimental fluorescent and proteomic data, as well as computational prediction algorithms to identify a likely protein location (Hooper et al., 2014) (Supplemental Data Set 5). The average fold depletion of each of the 546 proteins sorted by subcellular localization as identified by SUBAcon is shown in Figure 5A, and the contribution of each organelle to the abundance of spectral counts represented in the 546 protein depletion dataset is depicted in Figure 5B. Approximately 49% of depleted spectra were attributed to proteins associated with the cytoplasm and other organelles likely not directly participating in clathrin-mediated trafficking, such as the ER (Figure 5B). Demonstrating the effectiveness of the deuterium/Ficoll gradient, ~24% and 4% of all spectra corresponding to peptides more abundant in the DFGL relative to the final postcentrifugation CCV fraction (i.e. peptides indicating the

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**Table 1** Enrichment and depletion profiles of CCV-associated proteins and organelar markers

| Protein Name | Immunoblot | Dimethyl Labeling |
|--------------|------------|-------------------|
| CHC          | 5.15 ± 2.37 | 5.68 ± 1.48       |
| AP4E         | 3.14 ± 1.10 | 3.18 ± 0.60       |
| CLC2         | 2.07 ± 0.14 | 18.17 ± 13.70     |
| AP2M         | 4.33 ± 1.91 | 2.89              |
| AP1G         | 1.69 ± 0.53 | 5.28 ± 0.72       |
| KNOXE        | 1.29 ± 0.15 | 6.28 ± 4.87       |
| T-PLATE      | 0.88 ± 0.28 | 0.53              |
| AP2A         | 0.84 ± 0.30 | 2.62 ± 1.06       |
| cFBPase      | 0.77 ± 0.43 | –                 |
| SEC12        | 0.13 ± 0.06 | –                 |
| DRP1A        | 0.16 ± 0.02 | –                 |
| DRP1C        | 0.32 ± 0.34 | –                 |
| DRP2A        | 0.87 ± 0.59 | –                 |

The ratio of protein levels in the CCV fraction relative to levels in the DFGL fraction (without normalizing to levels in the S0.1 fraction) are presented alongside standard deviation about the mean. For average ratios derived by immunoblotting, n = 3 independent CCV preparations, except for DRP2 (n = 2). For dimethyl labeling values, a dash in the right-hand column indicates the protein was not detected in the labeling datasets. An absence of standard deviation about the mean in the right-hand column indicates the protein was present in only one of two replicates.
546 depleted proteins) were attributed to components of the ribosome and 26S proteasome, respectively.

Manual annotation of the 257 proteins that were enriched at least two-fold in the final CCV fraction relative to the deuterium/Ficoll-gradient load is depicted in Figure 5C. The fold enrichment and total spectral count are plotted for the enriched proteins sorted by manually assigned category relating to their roles in CCV trafficking. The fold enrichment and abundance of specific categories of CCV-associated proteins will be discussed further below.

Validation of CCV-associated proteins

Clathrin

Morphological and SDS–PAGE analyses demonstrated that CHC and CLC subunits were highly enriched in the CCV preparations (Figures 1, A and 2, A). Consistent with this, 41% of all spectra assigned to peptides corresponding to the 257 proteins that were enriched in the CCV fraction corresponded to the core clathrin coat components, namely CHC and CLC (Figure 5C). CHC1 and CHC2 were enriched six-fold.

Figure 5 Annotations of proteins identified by shotgun CCV proteomics that were more than two-fold depleted or enriched in the last stage of the CCV purification process. A, The average fold depletions of the 546 proteins that were more than two-fold depleted between the DFGL and CCV fractions and which overlapped with the CCV LC/MS–MS dataset deriving from 1D SDS–PAGE separation were plotted against the consensus subcellular localizations of the corresponding accession numbers predicted by the SUBA algorithm based on experimental and computational data. B, Proportion of total spectral counts across four independent CCV preparations for each > 2 × depleted protein also present in the CCV LC/MS–MS dataset deriving from 1D SDS–PAGE separation as categorized by SUBA annotated subcellular localization. C, The average fold enrichments of the 257 proteins that were more than two-fold enriched between the DFGL and CCV fractions and that overlapped with the CCV LC/MS–MS dataset deriving from 1D SDS–PAGE separation were plotted against the sum of the total spectral counts of each corresponding protein across four independent CCV preparations. Functional categorization of these proteins was manually annotated; the size and composition of each functional category are indicated below. The inset depicts the relative abundance of each functional category by summing and graphing the total spectral counts assigned to proteins within that functional category.
and five-fold, respectively, in CCVs compared to the DFGL (Table 1; Supplemental Data Set 4), which is comparable to their three-fold DFGL to CCV enrichment observed via quantitative immunoblotting (Figure 4, A and C). CLC1–3 were 8-, 18-, and 9-fold enriched in CCVs relative to the DFGL in dimethyl labeling experiments (Supplemental Data Set 4), while quantitative immunoblotting with αCLC2 antibody supported an enrichment for the CLC2 isoform in the final CCV fraction (Table 1; Figure 4, A and C).

The AP-1, AP-2, and TPC hetero-oligomeric adaptor protein complexes
Subunits of the previously characterized multimeric adaptor protein complexes AP-2, TPC, and AP-1 underlying endocytic and post-Golgi clathrin-dependent trafficking, respectively, were also identified in the suspension-cultured cell CCV proteome. All subunits, including large (A, B, and G), medium (M), and small (S) proteins, of the canonical, conserved heterotetrameric adaptin AP-1, and AP-2 complexes were detected in our datasets in high abundance, accounting for 16% of all spectra assigned to peptides enriched in CCVs in dimethyl labeling experiments (Supplemental Data Set 4). AP1G and AP2A were six- and three-fold more abundant in CCVs relative to the DFGL as observed by dimethyl labeling (Table 1; Supplemental Data Set 4). Consistent with this, AP1G was found to be enriched two-fold, and AP2A was found to be present in the final CCV fraction as determined by quantitative immunoblotting (Table 1; Figures 4, A and C), indicating that the enriched CCV preparations are a mixed population of post-Golgi and PM-derived CCVs. While both AP1G1 and AP1G2 homologs were detected in the dimethyl labeling replicates at an enrichment of about five-fold, only one of two AP2A homologs, AP2A2, was detected in the labeling studies at an enrichment of 2.6-fold, suggesting that AP2A1 is less abundant in the suspension-cultured cell CCV fraction. The medium and small subunits of the AP-1 and AP-2 complexes were enriched three-fold or greater as were the AP-1/2 B1 and B2 large subunits (six- and four-fold enrichment; Supplemental Data Set 4), which, similar to the case in Dictyostelium (Sosa et al., 2012) have been postulated to be interchangeably associated with AP-1 and AP-2 complexes in plants (Bassham et al., 2008). Efforts in recent years have established that the role of AP-1 in the trafficking of vacuolar cargo and clathrin recruitment in plant cells resembles that observed in yeast and mammalian systems. AP1M isoform mutants (ap1m1 and ap1m2) both show defects in trafficking of the soluble vacuolar protease precursor proaleurain (Song et al., 2006; Park et al., 2013). In addition, the AP-1 complex is critical for proper targeting of membrane-bound cargo to the tonoplast, at least partially via cytoplasmic sorting signals such as the N-terminal dileucine motif found in VACUOLAR ION TRANSPORTER1, which is mislocalized to the PM in ap1g mutants (Wang et al., 2014). TGN/EE integrity is, however, compromised in ap1 mutants, which manifests not only in defects in vacuolar protein transport but also in exocytic trafficking to the PM and cell plate, as well as CME (Park et al., 2013; Yan et al., 2021).

Recent characterization of the plant-specific TPC has revealed that the complex functions in endocytosis in concert with clathrin and AP-2 (Gadeyne et al., 2014; Bashline et al., 2015; Zhang et al., 2015; Wang et al., 2016). The two adapter complexes likely have overlapping as well as distinct functions (Gadeyne et al., 2014; Narasimhan et al., 2020; Johnson et al., 2021). Consistent with the former, both AP-2 and TPC bind clathrin and have been shown to interact with a common endocytic cargo protein, CES6 (Bashline et al., 2013; Sánchez-Rodríguez et al., 2018), one of three CESAs identified in the CCV proteome (Supplemental Data Set 4). However, compared to ap2 mutants, loss-of-function TPC subunit mutants display more severe biological phenotypic defects including pollen lethality (Van Damme et al., 2006; Gadeyne et al., 2014). Thus, given the essential nature of TPC in plant CME and the absence of homologs of most TPC subunits in yeast and metazoans, TPC is critical for functions unique to plants (Zhang et al., 2015). In our studies, MS analysis identified all eight core TPC subunits in the proteome derived from 1D SDS–PAGE separated CCVs (Supplemental Data Set 4). However, unlike the subunits of the heterotetrameric AP-1 and AP-2 complexes, which are all enriched in the CCV relative to DFGL fraction, the enrichment of the core TPC subunits was generally lower. Although the enrichment values of TPC subunits, TML and TASH3, were somewhat higher than those of the other subunits of TPC (e.g., TPLATE), no TPC subunit was strongly enriched in the last CCV purification step as detected by immunoblotting or dimethyl labeling (Table 1; Supplemental Data Set 4). Consistent with recent data from Johnson et al. showing that TPC is structurally more external to endocytic CCVs than AP-2, localizing around clathrin and AP2, and that TPC is loosely associated with purified CCV (Johnson et al., 2021), the EH domain-containing proteins EH1 and EH2 were present in the CCV proteome and similarly neither enriched alongside the vesicles as measured by the dimethyl labeling experiments (Supplemental Data Set 4). However, two ENTH-domain-containing TPC accessory components (AtECA4 and CAP1) were found in comparatively high abundance in CCVs (Supplemental Data Set 4). Given that ENTH proteins directly interact with membranes (Zouhar and Sauer, 2014), AtECA4 and CAP1 may be retained on CCVs to a larger extent than the core TPC. These data may explain why TPC components do not enrich to the degree of AP-2 subunits between the DFGL and CCV steps as measured by dimethyl labeling or immunoblotting, in that TPC subunits may have dissociated from the purified CCV. The abundance of TPC core and accessory proteins identified did not differ based on CCV sample preparation, including a similar enrichment of AtECA4 and CAP1 in CCVs relative to TPC core subunits (Supplemental Data Set 4).
The AP-4 adapter complex, but not AP-3 and AP-5, is associated with CCVs

In addition to the known clathrin-associated heterooligomeric adapter complexes, AP-1, AP-2, and TPC, the detection of all subunits of the less-studied AP-4 complex (AP4E, B, M, and S) and the abundance thereof at levels comparable to that of AP-1 and AP-2 are notable in the CCV proteome (Supplemental Data Set 4). The enrichment of the AP4E large subunit in CCV preparations was similar to that of AP2A and AP1G subunits (4.6-, 2.6-, and 5.6-fold CCV:DFGL ratios, respectively) as was the medium AP4M subunit relative to those corresponding to AP-2 and AP-1 (3.2-, 2.9-, and 5.9-fold, respectively) as determined by differential labeling experiments (Supplemental Data Set 4). In mammals, AP4M was not detected by immunoblotting in purified CCVs and the AP-4 complex was revealed to associate with non-CCVs near the TGN via immunogold electron microscopy (Hirst et al., 1999), suggesting AP-4 is not associated with clathrin in these organisms.

In plants, the four AP-4 subunits function together in a complex critical in trafficking to the protein storage vacuole (PSV). Similar to vacuolar sorting receptor (VSR) mutants vsr1, vsr3, and vsr4 (Zouhar et al., 2010), GREEN-FLUORESCENT SEED (GFS) loss-of-function AP-4 mutants gfs4, gfs5, gfs6, ap4e-1 corresponding to AP4B, AP4M, AP4S, and AP4E, respectively, mislocalize the PSV-targeted 12S globulin seed storage protein to the extracellular space (Fuji et al., 2016). Binding studies have demonstrated that the Arabidopsis AP4M subunit interacts with the cytoplasmic tail of VSR2 (Gershlick et al., 2014). Recently, evidence for interaction between AP4 subunits with DRPs and clathrin was shown via several independent approaches, including coimmunoprecipitations and yeast two-hybrid experiments (Fuji et al., 2016; Shimizu et al., 2021).

To further investigate the association of AP-4 subunits with plant CCVs, we generated a recombinant antibody against the C-terminal 22 amino acids of the AP4-E large subunit. The specificity of this antibody was confirmed by immunoblotting of total protein extracts prepared from subunit. The specificity of this antibody was confirmed by immunoblotting of total protein extracts prepared from Arabidopsis root tip cells (Fuji et al., 2016). Recently, GFP-tagged AP4M was shown to localize to the TGN/EE at sites distinct from AP1M2 (Shimizu et al., 2021).

Previous studies have indicated AP-4 functions at the TGN/EE (Hirst et al., 1999; Fuji et al., 2016; Shimizu et al., 2021). Consistent with this, functional GFP-tagged AP4M colocalizes with the TGN-resident SNARE mRFP-SYP43 and the endocytic tracer FM4–64 in Arabidopsis root tip cells (Fuji et al., 2016). Recently, GFP-tagged AP4M was shown to localize to the TGN/EE at sites distinct from AP1M2 (Shimizu et al., 2021). To further investigate the subcellular distribution of AP-4 and to determine whether it colocalizes with clathrin at the TGN/EE, we constructed lines that stably expressed N-terminal and C-terminal GFP- and RFP-tagged AP4E fusion proteins under control of the ubiquitin-10 promoter (Grefen et al., 2010). These constructs are functional in vivo as demonstrated by the rescue of the overall dwarf and abnormal growth phenotype observed in homozygous ap4e-1 and ap4e-2 plants (Supplemental Figure S4). Consistent with visualization of GFP-tagged AP4M (Fuji et al., 2016), ap4e-2 lines that expressed GFP-AP4E primarily displayed cytosolic and endomembrane subcellular localization with signal notably absent from the PM and tonoplast (Figure 6A).

We corroborated the subcellular localization of RFP-AP4E by pulse labeling, co-localization studies with the endocytic tracer dye, FM4–64. This dye has been used to distinguish plant endosomal compartments based on their spatial and temporal distribution of the dye upon internalization, for example, by labeling the TGN/EE within 2–6 min of internalization (Dettmer et al., 2006; Viotti et al., 2010). We observed co-localization between RFP-AP4E and FM4–64 (Pearson’s correlation coefficient [PCC] = 0.705 utilizing a Costes’ coefficient, 0.705; -value, 1.00; Pearson’s correlation coefficient, 0.37). Scale bars = 10 µm.

Figure 6 AP4E colocalizes with FM4–64 and clathrin at the TGN. A, ProUB10:GFP-AP4E expressed in homozygous ap4e-2 knockout lines has a primarily cytosolic and endomembrane localization in epidermal cells of root tips. B, TGN/EE labeled with FM4–64 after 6 min incubation co-localizes with ProUB10:RFP-AP4E in homozygous ap4e-2 knockouts (C); (D) Merge (Costes P-value, 1.00; Pearson’s correlation coefficient, 0.705). E, ProCLC2:CLC2-GFP and ProUB10:RFP-AP4E (F) colocalize at the TGN but not at the PM nor cell plate; (G) Merge (Costes P-value, 1.00; Pearson’s correlation coefficient, 0.37). Scale bars = 10 µm.
Clathrin accessory factors

In addition to hetero-oligomeric protein complexes, monomeric adapters, including members of the ENTH/ANTH/VHS domain-containing protein family and the Golgi-localized, gamma-ear-containing, ARF-binding (GGA) family, facilitate cargo recognition and vesicle formation at various points in the late endosome system. GGA proteins, which function in CCV formation at the TGN in yeast and mammals (Bonifacio, 2004), are absent from plants (Zouhar and Sauer, 2014). Arabidopsis contains 35 ENTH/ANTH/VHS domain-containing genes, 24 of which are putatively expressed in T87 suspension-cultured cells (Stolc et al., 2005). In addition to the TPC accessory proteins AtECA4 and CAP1 (see above), 13 other ENTH/ANTH/VHS-domain containing proteins were identified in the suspension-cultured cell CCV proteome (Supplemental Data Set 4) including the ENTH domain-containing monomeric clathrin adaptors EPSIN1 (EPS1) and EPS2 (Song et al., 2006; Lee et al., 2007; Collins et al., 2020) and the TGN-localized MODIFIED TRANSPORT TO THE VACUOLE 1 (MTV1; Sauer et al., 2013; Heinze et al., 2020).

EPS1 and MTV1 have previously been implicated in clathrin-mediated trafficking to the vacuole (Song et al., 2006; Sauer et al., 2013; Heinze et al., 2020). Consistent with additional roles in cargo trafficking to the PM, EPS1 modulates the PM abundance of the immune receptor FLAGELLIN SENSING2 and its co-receptor, BRI1-ASSOCIATED KINASE for effective defense responses (Collins et al., 2020). In our study, EPS1, EPS2, and MTV1 were enriched in CCVs relative to the DFGL in labeling experiments five-, five-, and seven-fold, respectively (Supplemental Data Set 4), consistent with data demonstrating their incorporation into CCVs (Sauer et al., 2013), biochemical interaction with clathrin and AP-1 (Song et al., 2006), and similar defects in the trafficking of the soluble vacuolar protease precursor proaleurain in eps1 and ap1m lines (Song et al., 2006; Park et al., 2013).

Mammalian and plant CME are also regulated by Eps15 homology domain-containing proteins, which function as membrane and/or protein adaptors (Chen et al., 1998; Bar et al., 2008; Schwiha and Korbe, 2020; Yperman et al., 2021). The Arabidopsis Eps15 homology domain-containing proteins, EHD1 and EHD2, were also found in the CCV proteome derived from both methodologies (Supplemental Data Set 4).

Several other proteins putatively functioning as trafficking adaptors were identified in the CCV proteome datasets including the sorting nexin homologs SNX1 and SNX2b (Supplemental Data Set 4). Mammalian sorting nexins interact with subunits of the endosomal retromer coat protein complex in a clathrin-independent fashion (McGough and Cullen, 2013) to mediate protein export from the early endosome to the TGN and PM by retrieving sorting receptors (e.g. the mannos-6-phosphate receptor) from the lysosome (Burd and Cullen, 2014). However, an interaction between the Arabidopsis retromer (VPS26a, 26b, 29, 35a, 35b, and 35c) and the three sorting nexin homologs (SNX1, 2a, and 2b) has yet to be shown, and conflicting evidence in the literature regarding SNX and retromer localization in plant cells at the TGN and MVB (reviewed in (Heucken and Ivanov, 2018; Robinson, 2018)) suggest multiple, independent roles for these complexes. Intriguingly, 12S globulin trafficking to the PSV is disrupted in snx mutants while that of 2S globulin remains apparently normal (Pourcher et al., 2010), a phenotype also observed in ap4 mutants (Fuji et al., 2016). Moreover, a yellow fluorescent protein (YFP) fusion with the Pisum sativum (pea) homolog of Arabidopsis VSR1 (BP80) colocalizes with AtSNX1 (Jailais et al., 2008) and trafficking of a similar BP80 reporter depends on SNX1 and
SNX2 function (Niemes et al., 2010). In contrast to SNX1 and SNX2b, components of the retromer core (VPS26A, VPS26B, VPS29, VPS35A, VPS35B, and VPS35C) were identified in the CCV proteome in low abundance (Supplemental Data Set 4), suggesting a possible role for SNX proteins in CCVs independent of retromer. Further studies are required to investigate the role of SNX proteins and a possible association with CCVs as an evolutionary divergence from the clathrin-independent nature of mammalian retromer/SNX function.

Several SH3 domain-containing proteins (SH3Ps) implicated in CCV trafficking were present in the suspension-cultured cell CCV proteome including the TASH3 subunit of the TPC, as well as SH3P1 and SH3P2, the latter of which is in high abundance and enriched in CCVs relative to DFGL fractions as measured by quantitative dimethyl labeling (eight-fold CCV:DFGL, Supplemental Data Set 4) and by immunoblotting (Nagel et al., 2017). Ubiquitylation of PM proteins serves as a posttranslational modification signaling for internalization and vacuolar degradation (Martins et al., 2015). SH3P2 localizes to the PM and functions as a ubiquitin-binding protein that may facilitate endosomal sorting complex required for transport (ESCRT) recognition of ubiquitylated cargo for subsequent degradation (Nagel et al., 2017). However, SH3P2 also localizes to the cell plate and to other clathrin-positive foci, suggesting this and other ubiquitin adaptors may function at other membranes beyond the PM.

Another set of clathrin accessory factors identified in the CCV proteome are Arabidopsis homologs of animal SCY1-LIKE2 proteins (SCYL2A and SCYL2B); SCYL2B is in high abundance and was found to co-enrich with CCVs by quantitative dimethyl labeling MS/MS experiments (15-fold CCV:DFGL ratio, Supplemental Data Set 4). In animals, SCYL2 binds clathrin (Duwel and Ungewickell, 2006), localizes to the Golgi, TGN, and other endosomes, and is incorporated into CCVs (Conner and Schmid, 2005; Borner et al., 2007). Although less is known about SCY1-LIKE2 proteins, which are postulated to function in uncoating of CCVs, with the protein being alternatively undetected in high abundance and was found to co-enrich with CCVs by quantitative dimethyl labeling MS/MS experiments (15-fold CCV:DFGL ratio, Supplemental Data Set 4). This parallel may reflect a functional coordination of ubiquitylated cargo for subsequent degradation (Nagel et al., 2017). However, SH3P2 also localizes to the cell plate and to other clathrin-positive foci, suggesting this and other ubiquitin adaptors may function at other membranes beyond the PM.

Another set of clathrin accessory factors identified in the CCV proteome are Arabidopsis homologs of animal SCY1-LIKE2 proteins (SCYL2A and SCYL2B); SCYL2B is in high abundance and was found to co-enrich with CCVs by quantitative dimethyl labeling MS/MS experiments (15-fold CCV:DFGL ratio, Supplemental Data Set 4). In animals, SCYL2 binds clathrin (Duwel and Ungewickell, 2006), localizes to the Golgi, TGN, and other endosomes, and is incorporated into CCVs (Conner and Schmid, 2005; Borner et al., 2007). Although less is known about SCY1-LIKE2 protein function in plants, SCYL2B was recently shown to localize to the TGN and interact with CHC as well as two related but functionally distinct TGN-associated SNAREs, VTI11 and VTI12, both of which were also identified in the CCV proteome (Supplemental Data Set 4; Jung et al., 2017). Taken together, these data suggest SCYL2A and SCYL2B function may reflect that of mammalian SCYL2 by mediating TGN CCV formation in Arabidopsis.

**DRPs**

Following cargo recognition and clathrin recruitment, the maturing CCV must separate from the PM prior to internalization. In metazoans, this is accomplished by the action of the dynamin GTPases, which oligomerize around the vesicle neck and utilize GTP hydrolysis to exert a constricting, twisting force, achieving scission (Antonny et al., 2016; Cheng et al., 2021). In plants, members of the DRP2 protein family, DRP2A and DRP2B, which are most closely related to mammalian dynamin (Backues et al., 2010; Smith et al., 2014), and the plant-specific DRP1 family members DRP1A and DRP1C (Collings et al., 2008; Mravec et al., 2011; Smith et al., 2014; Ekanayake et al., 2021) function together in cargo trafficking via CME (Ekanayake et al., 2021). DRP2 and DRP1 family members also biochemically interact with TPC and AP-2 (Gadeyne et al., 2014) and localize to PM CCPs (Konopka et al., 2008; Fujimoto et al., 2010; Wang et al., 2020).

Despite their clear association with PM CCPs (Konopka et al., 2008; Konopka and Bednarek, 2008; Fujimoto et al., 2010), DRP proteins were only detected at low levels in the total CCV proteome datasets (Table 1; Supplemental Data Set 4). Consistent with this, DRP1A and DRP1C show significant depletion in the CCV fraction relative to the DFGL as determined via quantitative immunoblotting (Figure 4, B and C; Table 1). This apparent discrepancy may reflect a short residency of DRP proteins at the PM wherein DRPs are recruited, accomplish their function(s), and dissociate from the newly formed CCV. The former hypothesis is supported by high-temporal-resolution imaging data showing recruitment of DRP2B, DRP1C, and DRP1A to endocytic foci at the PM simultaneously with or shortly after clathrin before their simultaneous disappearance, potentially reflecting a function in CCV maturation and/or scission (Konopka et al., 2008; Konopka and Bednarek, 2008; Fujimoto et al., 2010). The brief tenure of DRPs at nascent CCVs may also be reflected in the putative interaction between DRPs and the TASH3 subunits of the TPC (Gadeyne et al., 2014), which, along with other core TPC components, were likewise detected at relatively low abundance in the suspension-cultured cell CCV proteome (Supplemental Data Set 4). This parallel may reflect a functional coordination of recruitment and dissociation between DRPs and TPC in endocytic CCV formation. Alternatively, we cannot rule out the low abundance of DRP peptides in our dataset may be due to poor retention of the DRPs during the CCV isolation. In contrast, the DnaJ-related AUXILIN-LIKE1 and AUXILIN-LIKE2 proteins, which are postulated to function in uncoating of CCVs (Lam et al., 2001; Adamowski et al., 2018), were readily identified in the suspension-cell cultured CCV proteome (Figure 5C; Supplemental Data Set 4), possibly due to the conditions used in vesicle preparation (pH 6.4) that block clathrin coat disassembly but permit association of the AUXILIN-LIKE proteins with CCVs (Reynolds et al., 2014). Furthermore, recent data suggest that uncoating of CCVs in plants may not occur immediately after scission but as the CCV is trafficked away from the PM (Narasimhan et al., 2020).

Our results appear consistent with those observed in mammalian and yeast systems including the apparent uncertainty of dynamin association with isolated mammalian CCVs, with the protein being alternatively undetected (Blondeau et al., 2004) or observed with high confidence (Borner et al., 2006) in different preparations. However, the precise role(s) of members of the DRP2 and DRP1 protein...
families in CCV maturation and/or scission remains to be determined.

Vacuolar protein trafficking

Plant VSRs bind soluble vacuolar proteins including hydrolytic enzymes and vacuolar storage proteins (e.g. 12S globulin and 2S albumin) in the lumen of the secretory pathway through recognition of sorting motifs of vacuolar ligands, sequestering these cargo from others that are secreted or targeted toward the PM (Shimada et al., 2003). The primary mode of VSR function in plants has been postulated to reflect that of the mammalian mannose 6-phosphate receptor (MPR), which binds and traffics soluble lysosomal hydrolases via CCVs to the LE. The increasingly acidic nature of the maturing endosome triggers pH-dependent dissociation of its ligand, resulting in MPR recycling back to the TGN via retromer-mediated trafficking (Braulke and Bonifacino, 2009), at which the pH falls within the pH range for optimal MPR-ligand binding established by in vitro experiments (Tong and Kornfeld, 1989). However, increasing evidence indicates that VSR-mediated vacuolar protein trafficking is distinct in plants (Robinson and Neuhaus, 2016). In one instance of divergence from the mammalian MPR sorting model, plant VSR–ligand interaction appears to be initiated in the ER and cis-Golgi rather than the TGN/EE (daSilva et al., 2005; Gershlick et al., 2014; Künzl et al., 2016). Furthermore, the pH of the plant TGN (5.5) is lower than that of other compartments involved in VSR trafficking (Luo et al., 2015), ostensibly marking it as the site of ligand dissociation. Thus, the newly liberated VSRs must be recycled from the TGN back to the ER. In line with this hypothesis, a recent study has demonstrated the retrograde movement of nano-body tagged VSR from the TGN/EE to the early secretory pathway in tobacco (Nicotiana tabacum) protoplasts (Fruholz et al., 2018). Therefore, in addition to the AP/CCV-mediated anterograde trafficking of VSRs, alternative proposals have questioned whether VSR-laden CCVs carry soluble vacuolar cargo directly or are instead part of a recycling mechanism responsible for bringing VSRs back to the early secretory pathway after cargo dissociation in the TGN (reviewed in Kang and Hwang, 2014; Robinson and Neuhaus, 2016).

Previous analyses of isolated plant CCVs have identified abundant VSR proteins, consistent with a role for CCVs as VSR carriers (Mascalux et al., 2005; De Marcos Lousa et al., 2012). Indeed, the first plant VSR to be discovered (BP80) was initially detected in CCVs isolated from pea (P. sativum) cotyledons (Kirsch et al., 1994), and VSRs have been shown to co-fractionate with clathrin and vacuolar cargos during CCV preparations (Jolliffe et al., 2004). Consistent with this, five VSR proteins were identified in the suspension-cultured cell CCV proteome, of which three (VSR1, 3, and 4) were in highest abundance (Figure 5C, Supplemental Data Set 4).

Adaptor protein 1 and 4 complexes mediate the interaction of clathrin with VSR-ligand complexes, facilitating clustering of vacuolar cargo and vesicular formation necessary for vacuolar protein trafficking. The function of the AP-1 complex as an adaptor of vacuolar trafficking in plants reflects that of its mammalian/yeast counterparts in lysosomal/vacuolar trafficking. Recent work in Saccharomyces cerevisiae has demonstrated AP-1 is critical for retrograde recycling of cargo from mature to earlier Golgi cisternae (Papanikou et al., 2015; Day et al., 2018; Casler et al., 2019), suggesting that retrograde trafficking of VSRs and other cargo from the TGN and/or late Golgi to the early secretory pathway in plants may be mediated by AP-dependent CCVs. Moreover, binding of plant VSRs to AP1M is dependent on a cytoplasmic motif and is required for proper trafficking and maturation of the 12S globulin precursor seed storage protein (Gershlick et al., 2014; Fuji et al., 2016). Another member of the Arabidopsis VSR family, VSR4, interacts with AP1M2 partly through a complex cargo-recognition sequence. Mutations of this AP1M2 interaction motif result in VSR4 mislocalization to the PM and increased residency at the tonoplast, suggesting that this motif mediates both anterograde and retrograde VSR4 transport (Nishimura et al., 2016).

As noted above, mutants of AP-4 subunits mislocalize PSV cargo, a phenotype also observed in vsr and AP-1 subunit mutants (Zouhar et al., 2010; Park et al., 2013; Fuji et al., 2016), raising questions regarding the cargo and destination specificity of AP-1 and AP-4 dependent vacuolar trafficking. Pathway specific accessory proteins may aid in differentiating AP-1 and AP-4 function, such as the Arabidopsis homolog of the mammalian AP-4 accessory protein tepsin, MTV1, which has been shown to both bind clathrin and co-enrich with isolated CCVs as measured by immunoblotting and electron microscopy techniques (Borner et al., 2012; Sauer et al., 2013; Heimze et al., 2020). We have also found the AP4 interactor, MTV1, to be highly abundant in the Arabidopsis suspension-cultured cell CCV proteome and enriched in CCV pools relative to DFGL (seven-fold enrichment, Supplemental Data Set 4). Recent studies present a genetic interaction between AP4 and MTV1 and demonstrate that MTV1 functions in vacuolar trafficking (Heinz et al., 2020). Furthermore, the presence of EPS1, AP-1, AP-4, VTI11, and VSR1, 3, and 4 in the CCV data Set (Figure 5C; Supplemental Data Set 4) considered alongside available data regarding the vacuolar trafficking defects of ap1, ap4, and vsr mutants suggest that AP-1 and AP-4 facilitate anterograde, and possibly retrograde, CCV-mediated trafficking of VSRs to some degree.

Regulators of clathrin-mediated trafficking

Phospholipid metabolism

Intriguingly, one of the most abundant CCV proteome components identified was the phosphoinositide phosphatase SAC9, which is thought to regulate levels of phosphatidylinositol 4,5-bisphosphate [PI(4,5)P2] (Williams et al., 2005; Vollmer et al., 2011). In mammals, PI(4,5)P2 in the inner leaflet of the PM mediates CCP formation (Antonescu et al., 2011) through interactions with clathrin adaptor and accessory proteins including subunits of the AP-2 complex.
(AP2A, AP2B, and AP2M) (Jackson et al., 2010), epsin (Itoh et al., 2001), and dynamin (Vallis et al., 1999). The importance of these interactions in plant CME remains to be demonstrated, though PI(4,5)P₂ accumulates at sites of high membrane flux, like the growing pollen tube tip (Kost et al., 1999; Yao et al., 1999) and at the apex of expanding root hairs (Braun et al., 1999) and is incorporated into CCVs (Zhao et al., 2010). Moreover, mutants lacking the CCV-associated enzymes involved in the synthesis of PI(4,5)P₂, phosphatidylinositol 4-phosphate 5-kinasé PIP5K1 and PIP5K, show fewer CCPs at the PM of root epidermal cells (Ischebeck et al., 2013). Studies in plants have demonstrated that PI(4,5)P₂ recruits the endocytic accessory factor AP2M to the PM and interacts with the EH1 subunit of the TPLATE complex, suggesting that this phospholipid plays a role in the regulation of clathrin-mediated endocytosis (Doumane et al., 2021; Yperman et al., 2021). The presence of SAC9 in the CCV proteome (Figure 5C; Supplemental Data Set 4), suggests that PI(4,5)P₂ turnover in mature CCVs may play an important role in the regulation of CCV trafficking.

**Small GTPases**

Rabs and ARF GTPases regulate vesicle trafficking by modulating between GTP and GDP (active and inactive) bound states, which govern interactions with downstream effector proteins. Arabidopsis maintains 57 Rabs and 27 ARFs, of which the former can be grouped into eight distinct clades corresponding to mammalian Rab groups mediating different trafficking pathways (Rutherford and Moore, 2002). Of the numerous Rab identified in our study, members of the RabA1, D2, and E1 families were most abundant (Figure 5C; Supplemental Data Set 4). RabA proteins are most closely related to mammalian Rab11, which regulates endosomal trafficking and mediates TGN-PM trafficking in plants (Nielsen et al., 2008; Li et al., 2017; Zhou et al., 2020). In addition, members of the RabE family, which are orthologs of the Sec4p/Rab8 GTPase family, mediate exocytosis in yeast and mammalian cells (Rutherford and Moore, 2002) and regulate post-Golgi trafficking to the PM and to the cell plate during cytokinesis (Speth et al., 2009; Ahn et al., 2013; Orr et al., 2021). RabD2 proteins are related to mammalian Rab15, which participate in the early secretory pathway. In plants, however, RabD2 proteins localize to the Golgi/TGN (Pinheiro et al., 2009) and mediate post-Golgi trafficking at certain endosomes (Drakakagi et al., 2012).

Guanine nucleotide exchange factors (GEFs) activate GTPases by catalyzing the exchange of GDP for GTP. Evidence has shown STOMATAL CYTOKINESIS DEFECTIVE1 (SCD1) and SCD2 proteins participate in a complex that interacts with RabE1s in a nucleotide-dependent manner as well as subunits of the exocyst (Mayers et al., 2017), suggesting the SCD complex may function as a RabGEF in exocytosis. Both SCD1 and SCD2 enrich with the purification of CCVs by immunoblot analysis (McMichael et al., 2013) and were detected, albeit at low levels, in both unlabeled CCV proteomic datasets (Supplemental Data Set 4). Despite the presence of SCD1 and SCD2 in enriched CCVs and impaired internalization and post-Golgi trafficking defects of scl mutants, it is not clear whether the SCD complex directly functions in endocytosis or functions in recycling endocytic machinery to the PM.

The Arabidopsis suspension-cultured cell CCV proteome contains several additional structural and regulatory components of the exocyst tethering complex including SEC6, SEC15b, SEC10, EXO84B, and EXO70A1 (Figure 5C; Supplemental Data Set 4). Mammalian SEC15, SEC10, and EXO84 interact with vesicle-bound small GTPases to facilitate secretory trafficking from the TGN while EXO70 localizes to target membranes (Wu and Guo, 2015). Exocyst function in Arabidopsis appears to be conserved, including putative subcellular localizations at sites with high rates of vesicle fusion, for example, EXO70A1 and EXO84B at the CP (Fendrych et al., 2010).

In addition to GEFs, which initiate GTPase activation, GTPase activating proteins (GAPs) promoting Rab GTP hydrolysis to terminate GTPase signaling are likewise critical for controlling vesicle trafficking. The ARF-GAP NEVERSHED/AGDS/MTV4, which is required for vacuolar protein trafficking (Sauer et al., 2013), was previously demonstrated to co-enrich with CCVs using immunoblotting and immunoEM techniques. In agreement, this ARF-GAP is well represented in the CCV proteome (Supplemental Data Set 4).

**SNAREs**

In this study, we identified PM- and endosome-localized SNAREs in the CCV proteome (da Silva Conceicao et al., 1997; Bassham et al., 2000; Suwastika et al., 2008; Ebine et al., 2011; Eisenach et al., 2012; Uemura et al., 2012; Ichikawa et al., 2014). SNAREs that localized to the PM, such as SYP121, VAMP722, SYP132, and SYP71, were enriched seven-, five-, eight-, and six-fold, respectively, in the CCV fraction. Endosomal SNAREs, SYP61, SYP41, VT112, VAMP727, SYP21, and SYP43, were enriched 6-, 13-, 7-, 5-, 2-, and 9-fold, respectively (Figure 5C; Supplemental Data Set 4). The cell plate/cytokinesis-specific syntaxin and putative CCV cargo KNOLLE (KN) (Lauber et al., 1997; Boutté et al., 2010) was notably enriched in CCVs in dimethyl labeling (six-fold) and modestly enriched in quantitative immunoblotting (1.3-fold) experiments (Figure 4; Table 1; Supplemental Data Set 4). In contrast, SNAREs localized to compartments likely not engaged in CCV trafficking, such as the tonoplast protein VAMP713 (Takemoto et al., 2018) and the Golgi-localized VAMP714 (Uemura et al., 2005), were depleted in CCVs five- and three-fold relative to the DFGL, respectively (Supplemental Data Set 4).

We also identified several homologs of mammalian trafficking regulators involved in vesicle fusion in the CCV proteome, including PROTON ATPASE TRANSLOCATION CONTROL 1 (PATROL1), which contains the Munc13 MUN domain (Figure 4C; Supplemental Data Set 4). While the
mammalian Munc13 interacts with the SNARE syntaxin-1 to prime synaptic vesicles for fusion with the PM (Ma et al., 2011), less is known of MUN-domain containing protein function in plants. PATROL1 appears to modulate the delivery of the PM H+ -ATPase AHA1 to the PM possibly through interactions with the exocyst complex and localizes to endosomes and dynamic foci at the cell cortex (Hashimoto-Sugimoto et al., 2013; Higaki et al., 2014; Zhu et al., 2018). The presence of both PATROL1 and AHA1 in the CCV proteome (Figure 5C; Supplemental Data Set 4) suggests PATROL may regulate the trafficking of CCVs containing AHA1 through interactions with its MUN domain. Another SNARE interacting protein, VPS45, was detected in abundance in the CCV proteome and found to enrich in CCVs in differential labeling experiments (five-fold CCV:DFGL enrichment, Supplemental Data Set 4). VPS45 is a Munc18 protein that binds and regulates the TGN SNARE complex composed of SYP41, SYP61, YKT61, and VT12, potentially to mediate fusion of PVC-derived vesicles at the TGN (Bassham et al., 2000; Zouhar et al., 2009). Given the presence of VPS45 and its cognate SNAREs (SYP41, SYP61, VT12, and YKT61) in the CCV proteome (Figure 5C; Supplemental Data Set 4), clathrin-mediated trafficking may play a role in the regulation/function of the complex, possibly by recycling individual SNAREs for subsequent fusion events.

Conclusion

Space constraints prohibit a full discussion of all protein groups identified in the suspension-cultured cell CCV proteome; instead, we have reported on a subset of actively discussed protein groups and provided the complete CCV proteome as a rich data reference and resource for future investigations of proteins putatively involved in clathrin-mediated trafficking. An important caveat to interpreting the biological significance of the suspension-cultured CCV proteome is that CCV composition reflects the tissue/cell type and developmental stage of the biological source material from which they are isolated. The data reported here reflect the content of CCVs isolated from undifferentiated, rapidly dividing, and expanding Arabidopsis suspension-cultured cells under conditions amenable to tissue culture. Future experiments probing CCV content in other cell and tissue types under varying conditions or in different genetic backgrounds might apply our isolation methodology (Reynolds et al., 2014) using seedlings as a sample source as recently described (Nagel et al., 2017; Mosesso et al., 2019).

Our proteomic data demonstrate that, in addition to the canonical clathrin coat adapters AP-2 and AP-2, the AP-4 complex is incorporated into plant CCVs in an apparent contrast to its function in metazoans (Mattera et al., 2015; Robinson, 2015). This result inspires numerous questions regarding the evolutionary divergence of plant trafficking proteins relative to other eukaryotes, as well as the identity of which pathway(s) might be mediated by AP-4 and clathrin. Current biochemical and genetic studies in the literature suggest AP-4 mediates the trafficking of specific cargos to the PSV (Gershlick et al., 2014; Fuji et al., 2016), though whether AP-4 function in CCVs directly facilitates the anterograde trafficking of soluble cargos and their cognate receptors (i.e., VSRS) or instead participates in the recycling thereof remains to be determined. Additional studies identifying the composition of AP-2, AP-1, and AP-4-positive CCVs are needed to better understand cargo specificity and identify the regulators governing formation, trafficking, and fusion of these vesicles. Nevertheless, the identification of AP-4 as a CCV-associated protein complex presented here, together with recent genetic and biochemical evidence, will contribute to future experiments focused on elucidating the mechanisms of AP-4-dependent post-Golgi trafficking to the PSV. The question remains: is the interaction between AP-4 and clathrin direct as is the case for AP-1 and AP-2? Coimmunoprecipitation experiments (Fuji et al., 2016; Shimizu et al., 2021) and AP-4 abundance in the CCV proteome suggest that it might be, but a conclusive demonstration of a direct biochemical interaction remains to be determined. Determining the role of clathrin in VSR sorting, given the recent evidence of an ER/cis-Golgi to TGN/EE VSR-dependent, vacuolar trafficking pathway, will be essential as plant endomembrane dynamics become better understood (Robinson and Pimpl, 2014; Robinson and Neuhaus, 2016).

In contrast to the evolutionarily conserved endocytic adaptor AP-2, subunits of the TPLATE complex, which is essential for CME, were not significantly enriched in the final enrichment step toward purified CCVs, suggesting that the TPLATE complex may dissociate more readily from CCVs following their budding from the PM relative to AP-2. This observation may be resolved by a recent study in which TPLATE dissociated from CCVs earlier than clathrin and which positioned the TPLATE complex on the periphery of a clathrin-coated structure at the PM (Johnson et al., 2021). The positioning of the TPLATE complex to the periphery of the budding vesicle will be of significant biological interest.

As discussed above, the enriched CCV proteomic datasets contain numerous proteins that are unlikely to be associated with CCVs (e.g., ribosomal subunits). The presence of a protein in a shotgun MS/MS dataset, especially those of low abundance, must be taken as a preliminary indication only of its presence and therefore requires subsequent confirmation. Conversely, the absence or low abundance of proteins of interest might also be explained by low intracellular concentrations, such as in the case of regulatory proteins like the SCD1 and SCD2 subunits of the SCD complex which, while in low abundance in the MS/MS dataset, are shown to coenrich with isolated CCVs via immunoblotting and other techniques. Nevertheless, our CCV proteomics data expand our understanding of the plant endomembrane compartments for subsequent efforts to investigate the plant endomembrane network and their physiological function.
Materials and methods

Plant materials and growth conditions
Seed stocks of Col-0 (CS70000) and Col-3 (CS708) were obtained from the Arabidopsis Biological Research Center (ABRC). Seed stocks of the ap4e-1 and ap4e-2 T-DNA insertion mutants (Fuji et al., 2016) were graciously provided by T. Shimada of Kyoto University. CLC2-GFP transgenic plant lines were generated as previously described (Konopka et al., 2008). Seeds were sterilized in 70% (v/v) ethanol with 0.1% (v/v) Triton X-10 for 5 min and in 90% (v/v) ethanol for 1 min prior to plating on 0.5 x strength MS media (Murashige and Skoog, 1962) containing 0.6% (w/v) agar. Seeds were stratified without light at 4°C for 3 days prior to growing under continuous light at 22°C (T8 fluorescent bulb of 120 µmol m–2 s–1). Plants grown on soil were transferred from plates at 7–14 days to Metro-Mix 360 (SunGro Horticulture) and grown at 22°C long days (16 h of light exposure using T12 fluorescent bulb of 160 µmol m–2 s–1). ap4e-1 and ap4e-2 mutants were genotyped using primers GR1, 2, and 5 and GR3, 4, and 5, respectively (sequences listed in next section).

Undifferentiated Arabidopsis T87 cells (Axelos et al., 1992) were maintained in MS media supplemented with 0.2-mg/L 2,4-Dichlorophenoxyacetic (2,4-D) acid and 1.32-mM KH2PO4 under continuous light (T12 fluorescent bulb of 160 µmol m–2 s–1) at 22°C on an orbital shaker at 140 RPM. Cells were passaged weekly by a 1:10 dilution of a 7-day-old suspension culture into fresh MS 0.2 mg/L 2,4-D media.

Primers
GR1 – SAIL866C01_LP – 5’-CATGGGTATTGATGGTCTTG
G-3’
GR2 – SAIL866C01_RP – AGACCAGAACAAGCTAAG
CACG
GR3 – SAIL60E03_LP – ATAGGCTTCGAATCGAAGGC
GR4 – SAIL60E03_RP – ATGCAGGTGGAATCGTACTTG
GR5 – SAIL_IB3 – TAGCATCTGAATTTCCAACCAATC
TGATA
GR6 – SB1859STOPsense – CAAAGATCTCTCGGCTG
GACCTCTCTCTTCA
GR7 – SB1859STOPanti – TGAAGAAGAGAGGTGCTCA
GCCGAGAGATCTTTTG
GR8 – SB1859del673sense – CAAAAACCGAGCCTTCGC
AAAGGTTCTCCATGA
GR9 – SB1859del673anti – CTCCATGGACCTTCTTG
CGAAGCCTGCTTTTTG

Plasmid construction and plant transformation
All oligonucleotide primers used in this study were synthesized by Integrated DNA Technologies. Transgenic plants were generated using the floral dip method (Clough and Bent, 1998) with the Agrobacterium tumefaciens strain EHA105. N-terminal and C-terminal GFP and RFP fusions with AP4E under control of the UBQ10 promoter were created in pUBN-Dest and pUBC-Dest vectors (Grefen et al., 2010) using Gateway cloning (Invitrogen, Waltham, MA, USA). The AP4E CDS in pLIC6, obtained from ABRC clone DKLAT1G31730, was moved into pDONR221 (Invitrogen) using the BP Clonase kit. Stop codon insertion and frame correction for pUBN constructs was accomplished with primer pairs GR6&7 and GR8&9, respectively, before using the LR Clonase kit (Invitrogen) to move respective AP4E constructs into pUBN and pUBC destination vectors.

T1 seedlings from pUBN/pUBC transformed plants were sown on soil and selected by spray application of a 120 µg/mL Glufosinate (Liberty Herbicide, Bayer Crop Sciences) water solution containing 0.05% (v/v) Silwet-77 surfactant 7, 9, 12, and 14 days after germination.

CCV enrichment and analysis
CCVs were isolated from undifferentiated Arabidopsis T87 cells as described (Reynolds et al., 2014). Total protein yield in the enriched CCV fraction was approximately 300–500 µg per biological replicate. Protein concentrations of individual fractions for subsequent immunoblotting analysis and total CCV yield were obtained using the Pierce 660 nm Protein Assay (Thermo Fisher Scientific, Waltham, MA, USA).

Immunoblotting was performed as described (McMichael et al., 2013). Information about generation of antibodies and concentrations used is described in Supplemental Table S1.

Anti-AP4E antibody generation
The C-terminal 22 amino acids of the Arabidopsis AP4 Epsilson subunit were cloned from the ABRC stock DKLAT1G31730 into the pAN4GST GST-expression vector and the resulting construct used for phage display as described (Blanc et al., 2014) by the Geneva Antibody Facility at the Université de Genève, Switzerland. Antibody specificity was tested by probing the total protein content of wild-type, ap4e-1, ap4e-2, and various transgenic plants expressing GFP and RFP fusions with AP4E (Supplemental Figure S3). Seedlings grown on plates as described for 7–14 days were flash frozen and homogenized via mortar and pestle under liquid nitrogen. Tissue was resuspended in 2X Laemmli buffer, quantified using the Pierce 660 nm Protein Assay (Thermo Scientific). About 23.5 µg of each sample were fractionated on an 11% Tris–HCl SDS–PAGE gel and analyzed via immunoblotting.

Light microscopy
All confocal imaging experiments were conducted on a Nikon A1R-Si+ microscope. Colocalization and localization observations were made in root tip epidermal cells of 5- to 7-day-old T3 generation seedlings grown on plates as described above. Seedlings were mounted for imaging in 0.5 x MS media. FM4–64 treated samples were incubated in 0.5 x MS containing 4–µM FM4–64 for 3 min and mounted in the same solution before imaging after 6 min total incubation.

Colocalization analysis was performed using the JACoP plugin (Bolte and Cordelieres, 2006) in the Fiji (Schindelin et al., 2012) distribution of ImageJ2 (Rueden et al., 2017). Images were processed to remove background (Rolling ball
50-pixel diameter) and cropped to relevant ROIs before analysis with JACoP utilizing 1000 Costes randomizations with a point spread function of two pixels and Costes’ automated thresholding.

**STEM imaging of purified CCVs**

Four microliters of purified CCV preparation (0.33 mg/μL) were applied to carbon-coated and glow discharged (2 min at 7 × 10⁻³ mbar) 300-mesh copper EM grids (Electron Microscopy Sciences; CF300-CU) and incubated for 4 min at room temperature. Excess solution was removed with blotting paper and samples immediately fixed by a 20-min incubation at room temperature with 2% glutaraldehyde (v/v) in PEM buffer (100-mM PIPES, 1-mM MgCl₂, 1-mM EGTA, pH 6.9). Samples were then washed with phosphate buffer (0.1 M, pH 7.4) and distilled water, followed by a 20-min incubation with 0.1% tannic acid (w/v) in MilliQ water. After three washes in water, the samples were incubated with 0.2% aqueous uranyl acetate for 30 min at room temperature and then washed 3 times with water before being dehydrated in a series of graded ethanol (10%, 20%, 40%, 60%, 80%, 96%, and 100% for 2 min each) and two washes with hexamethyldisilane (99%). Dried samples were then coated with 3-nm platinum and 4-nm carbon using an ACE600 coating device (Leica Microsystems, Wetzlar, Germany). The sample grids were imaged using a JEOL JEM2800 scanning/transmission electron microscope at 200 kV.

**LC/MS-MS of CCVs separated by 1D SDS-PAGE**

Enriched CCV fractions were resolved via 1D SDS-PAGE on a 4%–15% Tris–HCl gradient gel (BioRad, Hercules, CA, USA; cat# 161-1158) at a constant 200 V for ~90 min. Gels were stained with Coomassie R250 and cut into ~10 bands. Gel bands were digested in-gel as described (Shevchenko et al., 1996) with modifications. Briefly, gel bands were dehydrated using 100% acetonitrile and incubated with 10-mM dithiothreitol in 100-mM ammonium bicarbonate, pH ~8, at 56°C for 45 min, dehydrated again and incubated in the dark with 50-mM iodoacetamide in 100-mM ammonium bicarbonate for 20 min. Gel bands were then washed with ammonium bicarbonate and dehydrated again. Sequencing grade modified trypsin was prepared to 0.01 μg/μL in 50-mM ammonium bicarbonate and ~50 μL of this was added to each gel band so that the gel was completely submerged. Bands were then incubated at 37°C overnight. Peptides were extracted from the gel by water bath sonication in a solution of 60% acetonitrile/1.0% trifluoroacetic acid and vacuum dried to ~2 μL.

Enriched DFGL or CCV fractions for dimethyl labeling experiments were prepared in the same manner as the other CCV samples up to peptide extraction and vacuum drying except that a 12.5% Tris–HCl SDS polyacrylamide gel was used. Each peptide sample was then re-suspended in 100-mM Triethylammonium and labeled in solution with dimethyl reagents (light label – C₂H₆, medium label – C₂H₂D₄) according to Boersema et al. (2009). Aliquots of DFGL and CCV samples labeled with light and heavy formaldehyde were analyzed by MS to measure the labeling efficiency, which was found to be >95% for each replicate. After labeling, peptides were purified using solid phase extraction tips (OMIX, www.varian.com). Same slice samples from each condition were combined and dried to ~2 μL. Peptides were then re-suspended in 2% acetonitrile/0.1%TFA to 25 μL. From this, 10 μL was automatically injected by a Waters nanoAcquity Sample Manager (www.waters.com) and loaded for 5 min onto a Waters Symmetry C18 peptide trap (5 μm, 180 μm × 20 mm) at 4 μL/min in 5% acetonitrile/0.1%Formic Acid. The bound peptides were then eluted onto a MICROM Bioresources (www.michrom.com) 0.1 × 150 mm column packed with 3 μm, 200A Magic C18AQ material over 90 min with a gradient of 5% B to 35% B in 77 min, ramping to 90%B at 79 min, holding for 1 min and returning to 5%B at 80.1 min for the remainder of the analysis using a Waters nanoAcquity UPLC (Solvent A = 99.9% Water/0.1% Formic Acid, Solvent B = 99.9% Acetonitrile/0.1% Formic Acid) with an initial flow rate of 1 μL/min.

Peptides were then re-suspended in 2% acetonitrile/0.1%TFA to 20 μL. From this, 10 μL was automatically injected by a Thermo (www.thermo.com) EASYnLC onto a Thermo Acclaim PepMap RSLC 0.075 mm × 150 mm C18 column and eluted at 250 nL over 90 min with a gradient of 5%B to 30%B in 79 min, ramping to 100%B at 80 min and held at 100% B for the duration of the run (Solvent A = 99.9% Water/0.1% Formic Acid, Solvent B = 99.9% Acetonitrile/0.1% Formic Acid). Eluted peptides were analyzed as follows: CCV I: peptides were sprayed into a hybrid ThermoFisher LTQ FT-ICR Ultra mass spectrometer with both ion cyclotron resonance (ICR) and LTQ linear ion trap mass analyzers and a Michrom ADVANCE nanospray source with survey scans taken in the FT (25,000 resolution determined at m/z 400) and the top five ions in each survey scan then subjected to automatic low energy collision-induced dissociation (CID) in the LTQ; CCV II: peptides were sprayed into a ThermoFisher LTQ Linear Ion trap mass spectrometer using a Michrom ADVANCE nanospray source with the top eight ions in each survey scan are then subjected to low energy CID in a data dependent manner; CCV III and CCV IV: peptides were sprayed into a ThermoFisher Q-Exactive mass spectrometer using a FlexSpray spray ion source with survey scans taken in the Orbi trap (70,000 resolution, determined at m/z 200) and the top twelve ions in each survey scan then subjected to automatic higher energy CID (HCD at 25%) with fragment spectra acquired at 17,500 resolution; Dimethyl Labeling Samples: peptides were sprayed into ThermoFisher LTQ FT-ICR Ultra mass spectrometer using a Michrom ADVANCE nanospray source with intensity measurements of the two diMe forms made in the FT-ICR portion of the instrument and peptide fragmentation performed in the linear ion trap. The resulting MS/MS spectra from CCV replicates were converted to peak lists using Mascot Distiller, version 2.4.3.3 (www.matrixscience.com) and searched against a custom database that included all
entries in the TAIR10 protein sequence database (downloaded from www.arabidopsis.org), appended with common laboratory contaminants (downloaded from www.thegpm.org, cRAP project), using the Mascot searching algorithm, version 2.4, with the following parameters: two allowed missed tryptic cleavages, fixed modification of carbamidomethyl cysteine, variable modification of oxidation of methionine, peptide tolerance ±5 p.p.m., MS/MS tolerance 0.50 Da (Replicates I and II) and 0.015 Da (Replicates III and IV). FDRs were calculated against a randomized database search. The Mascot output was then analyzed using Scaffold, version 4.8.4 (Proteome Software Inc., Portland, OR, USA) to probabilistically validate protein identifications. Assignments validated using the Scaffold 1% FDR protein threshold, containing two unique peptides, meeting the 1% FDR peptide threshold were considered true. Assignments matching these criteria from four distinct biological replicates (CCV enrichments) were cross-referenced to eliminate duplicates and filtered to exclude proteins not present in at least two biological replicates, generating a list of 3,745 protein assignments.

Quantitation of labeled MS peaks from dimethyl labeling samples and processing of the resulting MS/MS spectra to peak lists was done using MaxQuant3, version 1.2.2.5 (Cox and Mann, 2008). Peak lists were searched against the TAIR10 protein sequence database, downloaded from www.arabidopsis.org and appended with common laboratory contaminants using the Andromeda search algorithm within MaxQuant with the following parameters: two allowed missed tryptic cleavages, fixed modification of carbamidomethyl cysteine, variable modification of oxidation of methionine and acetylation of protein N-termini, DimethLys0, DimethNter0; DimethLys4, DimethNter4, peptide tolerance ±6 p.p.m., MS/MS tolerance 20 p.p.m., protein and peptide FDR filter set to 1% (Cox et al., 2011). A total of 1,109 unique proteins were identified between both replicates, 957 of which were identified in the LC/MS–MS dataset corresponding to CCVs digested in gel across two or more biological replicates.

Overlaps between proteomic datasets were identified by matching accession numbers between the 3,745 accession numbers in column 1 of Supplemental Data Set 1, the accession numbers of the 1,109 proteins identified in at least one of two dimethyl replicates (Supplemental Data Set 3), and the first accession number (corresponding to iBAQ values listed) of the majority accession numbers (identifiers of all sequences containing at least half of the peptides of the leading sequence) within each protein group (n = 1,981) in the alternative methodology CCV LC/MS–MS dataset (Supplemental Data Set 4). Overlaps between proteomic datasets were visualized using the application BioVenn (Hulsen et al., 2008).

Protein localization of the 546 more than two-fold depleted proteins between the DFGL and CCV purification steps (Supplemental Data Set 3) was determined by submission of the corresponding accession numbers (Supplemental Data Set 5) to the SUBcellular Arabidopsis (SUBA) consensus algorithm (Hooper et al., 2014).

LC/MS–MS of CCVs digested in solution
About 5 μg of 1 μg/μl isolated CCVs were lysed in 8 M urea, 20 mM HEPES pH 8.0, by repeatedly pipetting up and down. Proteins in each sample were reduced by adding 15 mM dithiothreitol and incubating the samples for 30 min at 55°C. Proteins were alkylated by addition of 30 mM iodoacetamide for 15 min at room temperature in the dark. The samples were diluted with 20 mM HEPES pH 8.0 to a urea concentration of 2 M and the proteins were digested with 4 μl Trypsin/LysC (Promega V5073: 20 μg + 80 μl 50-MM acetic acid) for 4 h at 37°C and boosted with an extra 2-μl Trypsin/LysC (Promega V5073: 20 μg + 80 μl 50-MM acetic acid) overnight at 37°C. Peptides were then purified on a OMIX C18 pipette tip (Agilent, Santa Clara, CA, USA).

Purified peptides were re-dissolved in 25-μl loading solvent A (0.1% trifluoroacetic acid in water/acetonic (98:2, v/v)) and 5 μl was injected for LC–MS/MS analysis on an Ultimate 3000 RSLCnano ProFlow system in-line connected to a Q Exactive HF mass spectrometer (Thermo). Trapping was performed at 10 μl/min for 4 min in loading solvent A on a 20-mm trapping column (made in-house, 100-μm inner diameter (I.D.), 5 μm beads, C18 Reprosil-HD, Dr Maisch, Germany) and the sample was loaded on a 200-mm analytical column (made in-house, 75 μm ID, 1.9-μm beads C18 Reprosil-HD, Dr Maisch). Peptides were eluted by a nonlinear gradient from 2% to 55% MS solvent B (0.1% FA in water/acetonicitrile (2:8, v/v)) over 175 min at a constant flow rate of 250 nL/min, reaching 99% MS solvent B after 200 min, followed by a 10 min wash with 99% MS solvent B and re-equilibration with MS solvent A (0.1% FA in water). The column temperature was kept constant at 50°C in a column oven (Butterfly, Phoenix S&T). The mass spectrometer was operated in data-dependent mode, automatically switching between MS and MS/MS acquisition for the 16 most abundant ion peaks per MS spectrum. Full-scan MS spectra (375–1,500 m/z) were acquired at a resolution of 60,000 in the orbitrap analyzer after accumulation to a target value of 3E6. The 16 most intense ions above a threshold value of 1.3E4 were isolated for fragmentation at a normalized collision energy of 28% after filling the trap at a target value of 1E5 for a maximum of 80 ms. MS/MS spectra (200–2,000 m/z) were acquired at a resolution of 15,000 in the orbitrap analyzer.

Data were analyzed with MaxQuant (version 1.6.10.43) using the built in Andromeda search engine with default settings, including an FDR set at 1% on both the peptide and protein level. Spectra were searched against the Araport11plus database, consisting of the Araport11_genes.2016.06.pep.fasta downloaded from www.arabidopsis.org, extended with sequences of all types of possible contaminants in proteomics experiments in general. These contaminants include the cRAP protein sequences, a list of proteins commonly found in proteomics experiments,
which are present either by accident or by unavoidable contamination of protein samples (The Global Proteome Machine, http://www.thegpm.org/crap/). In addition, commonly used tag sequences and typical contaminants, such as sequences from frequently used resins and proteases, were added. The Arabort1plus database contains in total 49,057 sequence entries. The mass tolerance for precursor and fragment ions was set to 4.5 and 20 p.p.m., respectively, including matching between runs and FDR set at 1% on PSM, peptide, and protein level. Enzyme specificity was set as C-terminal to arginine and lysine (trypsin), also allowing cleavage at arginine/lysine-proline bonds with a maximum of two missed cleavages. Variable modifications were set to oxidation of methionine residues and acetylation of protein N-terminals. Proteins were quantified by the MaxLFQ algorithm integrated in the MaxQuant software. Only protein groups with at least one unique peptide and with peptides identified in at least two of three biological replicates were retained, yielding a list of 1,981 protein groups (Supplemental Data Set 4).

Accession numbers
Accession numbers corresponding to proteins identified in this study can be found in Supplemental Data Sets 1–3. Files corresponding to raw proteomics data, as well as results, search, and peak list files can be accessed at the MassIVE data repository (MassIVE, University of California at San Diego) using the identifiers, PXD026180 or at doi:10.25345/C50F9D. A summary of the methods and protocols used, as well as gel images and supplementary files, can also be accessed here.

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

Supplemental Figure S1. TEM of CCVs (supports Figure 1).
Supplemental Figure S2. Schematics illustrating protocol for clathrin coated vesicle purification and workflows detailing the CCV proteome (Supports Figure 3).
Supplemental Figure S3. AP4E antibody is specific for the AP4E subunit (supports Figure 4).
Supplemental Figure S4. ProUB10GFP-AP4E is functional in vivo (supports Figure 6).
Supplemental Table S1. Antibodies used in this study.
Supplemental Data Set 1. Merged CCV proteomic datasets.
Supplemental Data Set 2. LC/MS–MS data corresponding to CCVs digested in solution.
Supplemental Data Set 3. CCV dimethyl labeling LC/MS–MS data.
Supplemental Data Set 4. LC/MS–MS data corresponding to discussed proteins.
Supplemental Data Set 5. Predicted protein localization data for proteins more than two-fold depleted between DFGL and CCV fractions.

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References
Adamowski M, Narasimhan M, Kania U, Glanc M, De Jaeger G, Friml J (2018) A functional study of AUXILIN-LIKE1 and 2, two putative clathrin uncoating factors in Arabidopsis. Plant Cell 30: 700–716
Ahn CS, Han JA, Pai HS (2013) Characterization of in vivo functions of Nicotiana benthamiana RabE1. Planta 237: 161–172
Antonescu CN, Aguet F, Danuser G, Schmid SL (2011) Phosphatidylinositol-(4,5)-bisphosphate regulates clathrin-coated pit initiation, stabilization, and size. Mol Biol Cell 22: 2588–2600
Antonny B, Burd C, De Camilli P, Chen E, Daumke O, Faelber K, Ford M, Frolov VA, Frost A, Hinshaw JE, et al. (2016). Membrane fission by dynamin: what we know and what we need to know. EMBO J 35: 2270–2284
Arabidopsis Genome I (2000) Analysis of the genome sequence of the flowering plant Arabidopsis thaliana. Nature 408: 796–815
Arike L, Valgepea K, Peil L, Nahku R, Adamberg K, Vilu R (2012) Comparison and applications of label-free absolute proteome quantification methods on Escherichia coli. J Proteomics 75: 5437–5448
Axelos M, Curie C, Mazzolini L, Bardet C, Lescure B (1992) A protocol for transient gene-expression in Arabidopsis-thaliana protoplasts isolated from cell-suspension cultures. Plant Physiol Biochem 30: 123–128
Backues SK, Korasick DA, Heese A, Bednarek SY (2010) The Arabidopsis dynamin-related protein2 family is essential for gametophyte development. Plant Cell 22: 3218–3231
Drakaki G, van de Ven W, Pan S, Miao Y, Wang J, Keinath NF, Weatherly B, Jiang L, Schumacher K, Hicks G, et al. (2012) Isolation and proteomic analysis of the SY6P61 compartment reveal its role in exocytic trafficking in Arabidopsis. Cell Res 22: 413–424

Dwul M, Ungewickell EJ (2006) Clathrin-dependent association of CVAK104 with endosomes and the trans-Golgi network. Mol Biol Cell 17: 4513–4525

Ebne K, Fujimoto M, Okatani Y, Nishiyama T, Goh T, Ito E, Danioiu T, Nishitani A, Uemura T, Sato MH, et al. (2011) A membrane trafficking pathway regulated by the plant-specific RAB GTPase ARA6. Nat Cell Biol 13: 853–859

Eisenach C, Chen ZH, Greven C, Blatt MR (2012) The trafficking protein SYPI21 of Arabidopsis connects programmed stomatal closure and K(+) channel activity with vegetative growth. Plant J 69: 241–251

Eknakyake G, LaMontagne E, Heese A (2019) Never walk alone: clathrin-coated vesicle (CCV) components in plant immunity. Annu Rev Physiol Pathol 57: 307–409

Eknakyake G, Smith J, Jones K, Stiers H, Robinson S, LaMontagne E, Kostos P, Cornish P, Bednarek S, Heese A (2021) DYNAMIN-RELATED PROTEIN DRP2B functions with DRP2B in plant growth, flag22-immune responses, and endocytosis. Plant Physiol 185: 1986–2002

Fan L, Hao H, Xue Y, Zhang L, Song K, Ding Z, Botella MA, Wang H, Lin J (2013) Dynamic analysis of Arabidopsis AP2 sigma subunit reveals a key role in clathrin-mediated endocytosis and plant development. Development 140: 3826–3837

Fendrych M, Synk L, Pecenková T, Toupalová H, Cole R, Drdová E, Nebesárová J, Sedínová M, Hala M, Fowler JE, et al. (2020) The Arabidopsis exocytosis complex is involved in cytokinesis and cell plate maturation. Plant Cell 22: 3053–3065

Feng Q, Song S, Yu S, Wang J, Li S, Zhang Y (2017) Adaptor protein-3-dependent vacuolar trafficking involves a subpopulation of COPII and HOPS tethering proteins. Plant Physiol 174: 1609–1620

Feraru E, Paciorek T, Feraru Ml, Zwiewka M, De Groodt R, De Rycke R, Kleine-Vehn J, Friml J, Borner GH, Robinson M, et al. (2010) The Arabidopsis dynamin-related proteins DRP2B and DRP1A participate together in clathrin-coated vesicle (CCV) components in plant immunity. Mol Biol Cell 21: e42004411

Fröhlich A, Sampathkumar A, McFarlane H, Sauer M (2020) A Munc13-like protein in Arabidopsis mediates H+-ATPase transmembrane trafficking. J Cell Sci 133: jcs203695

Heuze L, Freimuth N, Rößling A, Hahnke R, Riebschläger S, Fröhlich A, Sampathkumar A, McFarlane H, Sauer M (2020) EPSIN1 and MTV1 define functionally overlapping but molecularly distinct trans-Golgi network subdomains in Arabidopsis. Proc Natl Acad Sci USA 117: 25880–25889

Heinkel N, Ivanov R (2018) The retromer, sorting nexins and the plant endomembrane protein trafficking. J Cell Sci 131: jcs203695

Hashimoto-Sugimoto M, Igaki K, Iba K, Hasezawa S (2014) Dynamics and environmental responses of PATROL1 in Arabidopsis subsidiary cells. Plant Cell Physiol 55: 773–780

Hirst J, Barlow LD, Francisco GC, Sahlen达尔 DA, Seaman MN, Dacks JB, Robinson MS (2011) The fifth adaptor protein complex. PLoS Biol 9: e1001170

Hirst J, Borner GH, Antrobus R, Pedersen AA, Hydén NA, Sahlen达尔 DA, Robinson MS (2012). Distinct and overlapping roles for AP-1 and GGAs revealed by the “knocksideways” system.Curr Biol 22: 1711–1716

Hirst J, Bright NA, Rous B, Robinson MS (1999) Characterization of a fourth adaptor-related protein complex. Mol Biol Cell 10: 2787–2802

Hirst J, Irving C, Borner GH (2013) Adaptor complex proteins AP-4 and AP-5: new players in endosomal trafficking and progressive spastic paraplegia. Traffic 14: 153–164

Hirst J, Izhak D, Antrobus R, Borner G, Robinson M (2018) Role of the AP-5 adaptor protein complex in late endosome-to-Golgi retrieval. PLoS Biol 16: e2004411

Hirst J, Schlacht A, Nortić JP, Traynor D, Bloomfield G, Antrobus R, Kay RR, Dacks JB, Robinson MS (2014) Characterization of TSET, an ancient and widespread membrane trafficking complex. Elife 3: e02866

Hopper C, Tanz S, Castlejohn I, Vacher M, Small I, Millar AH (2014) SUBACon: a consensus algorithm for unifying the subcellular localization data of the Arabidopsis proteome. Bioinformatics 30: 3356–3364

Hulsen T, de Vlieg J, Alkema W (2008) BioVenn - a web application for the comparison and visualization of biological lists using area-proportional Venn diagrams. BMC Genomics 9: 1–6

Ichikawa M, Hirano T, Enami K, Fusetier T, Kato N, Kwon C, Voigt B, Schulze-Lefert P, Baluska F, Sato MH (2014) Syntaxin of plant proteins SYPI23 and SYPI32 mediate root hair tip growth in Arabidopsis thaliana. Plant Cell Physiol 55: 790–800

Ischebeck T, Werner S, Krishnamoorthy P, Lerche J, Meijón M, Stenzel I, Löfke C, Wiessner T, Im YJ, Perera IY, et al. (2013) Phosphatidylinositol 4,5-bisphosphate influences PIN polarization by controlling clathrin-mediated membrane trafficking in Arabidopsis. Plant Cell 25: 4894–4911
Itoh T, Koshiba S, Kigawa T, Kikuchi A, Yokoyama S, Takenawa T (2001) Role of the ENTH domain in phosphatidylinositol-4,5-bisphosphate binding and endocytosis. Science 291: 1047–1051

Jackson LP, Kelly BT, McCoy AJ, Gaffry T, James LC, Collins BM, Honing S, Evans PR, Owen DJ (2010) A large-scale conformational change couples membrane recruitment to cargo binding in the AP2 clathrin adaptor complex. Cell 141: 1220–1229

Jailais Y, Fobis-Loisy I, Miége C, Gaude T (2008) Evidence for a sorting endosome in Arabidopsis root cells. Plant J 53: 237–247

Johnson A, Dahhan D, Gnyliukh N, Kaufmann W, Zelden V, Costanzo T, Mahou P, Hrtyan M, Wang J, Aguilera-Servin J, et al. (2021) The TPLATE complex mediates membrane bending during plant clathrin-mediated endocytosis. Proc Natl Acad Sci USA 118: e2113046118

Jolliffe NA, Brown JC, Neumann U, Vicre M, Bachi A, Hawes C, Ceriotti A, Roberts LM, Frigerio L (2004) Transport of ricin and 2S albumin precursors to the storage vacuoles of ricinocarpus endosperm involves the Golgi and VSR-like receptors. Plant J 39: 821–833

Jung JY, Lee DW, Ryu SB, Hwang I, Schachtman DP (2017) SCYL2 genes are involved in clathrin-mediated vesicle trafficking and essential for plant growth. Plant Physiol 175: 194–209

Jürgens G, Park M, Richter S, Touihri S, Krause C, El Kasmi F, Johnson A, Dahhan D, Gnyliukh N, Kaufmann W, Zelden V, Constantino T, Hrtyan M, Wang J, Aguilera-Servin J, et al. (2021) The TPLATE complex mediates membrane bending during plant clathrin-mediated endocytosis. Proc Natl Acad Sci USA 118: e2113046118

Kaksonen M, Roux A (2018) Mechanisms of clathrin-mediated endocytosis. Nat Rev Mol Cell Biol 19: 313–326

Kang BH, Nielsen E, Preuss ML, Mastronarde D, Staeelin LA (2011) Electron tomography of RabA4b- and PI-4Kbeta1-labeled trans Golgi network compartments in Arabidopsis. Traffic 12: 313–329

Kang H, Hwang I (2014) Vacular sorting receptor-mediated trafficking of soluble vacuolar proteins in plant cells. Plants (Basel) 3: 392–408

Kee JH (1987) Clathrin assembly proteins: affinity purification and a model for coat assembly. J Cell Biol 105: 1899–1998

Kim SY, Xu ZY, Song K, Kim DH, Kang H, Reichardt I, Sohn Ej, Friml J, Jürgens G, Hwang I (2013) Adapter protein complex 2-mediated endocytosis is crucial for male reproductive organ development in Arabidopsis. Plant Cell 25: 2970–2985

Kirsch T, Paris N, Butler JM, Beever L, Rogers JC (1994) Purification and initial characterization of a potential plant vacuolar targeting receptor. Proc Natl Acad Sci USA 91: 3403–3407

Kitakura S, Vanneste S, Robert S, Löfke C, Teichmann T, Tanaka H, Friml J (2011) Clathrin mediates endocytosis and polar distribution of PIN auxin transporters in Arabidopsis. Plant Cell 23: 1920–1931

Konopka CA, Backues SK, Bednarek SY (2008) Dynamics of Arabidopsis dynamin-related protein 1C and a clathrin light chain at the plasma membrane. Plant Cell 20: 1363–1380

Konopka CA, Bednarek SY (2008) Comparison of the dynamics and functional redundancy of the Arabidopsis dynamin-related isoforms DRP1A and DRP1C during plant development. Plant Physiol 147: 1590–1602

Kost B, Lemichez E, Spielhofer P, Hong Y, Tolias K, Carpenter C, Chua NH (1999) Rac homologues and compartmentalized phosphatidylinositol 4, 5-bisphosphate act in a common pathway to regulate polar pollen tube growth. J Cell Biol 145: 317–330

Künzl F, Führholz S, Fäßler F, Li B, Pimpl P (2016) Receptor-mediated sorting of soluble vacuolar proteins ends at the trans-Golgi network/early endosome. Nat Plants 2: 16017

Lam BC, Sage TL, Bianchi F, Blumwald E (2001) Role of SH3 domain-containing proteins in clathrin-mediated vesicle trafficking in Arabidopsis. Plant Cell 13: 2499–2512

Lauber MH, Waizenegger I, Steinmann T, Schwarz H, Mayer U, Hwang I, Lukowitz W, Jürgens G (1997) The Arabidopsis KNOLLE protein is a cytokinesis-specific syntaxin. J Cell Biol 139: 1485–1493

Lee GJ, Kim H, Kang H, Jang M, Lee DW, Lee S, Hwang I (2007). EpsinR2 interacts with clathrin, adaptor protein-3, ActVIT12, and phosphatidylinositol-3-phosphate. Implications for EpsinR2 function in protein trafficking in plant cells. Plant Physiol 143: 1561–1575

Li R, Rodriguez-Furlan C, Wang J, van de Ven W, Gao T, Raikhel N, Hicks G (2017) Different endomembrane trafficking pathways establish apical and basal polarities. Plant Cell 29: 90–108

Lipatova Z, Hain AU, Nazarky VY, Segev N (2015) Ypt/Rab GTases: principles learned from yeast. Crit Rev Biochem Mol Biol 50: 203–211

Lundgren DH, Hwang SI, Wu L, Han DK (2010) Role of spectral counting in quantitative proteomics. Expert Rev Proteomics 7: 39–53

Luo Y, Scholl S, Doering A, Zhang Y, Irani NG, Rubbo SD, Neumetzler L, Krishnamoorthy P, Van Houtte I, Myle E, et al. (2015) V-ATPase activity in the TGN/EE is required for exocytosis and recycling in Arabidopsis. Nat Plants 1: 15094

Ma C, Li W, Xu Y, Rizo J (2011) Munc13 mediates the transition from the closed syntaxin-Munc18 complex to the SNARE complex. Nat Struct Mol Biol 18: 542–549

Martins S, Dohmann EM, Cayrel A, Johnson A, Fischer W, Pojer F, Satiat-Jeunemaitre B, Jaillais Y, Chory J, Geldner N, et al. (2015) Internalization and vacuolar targeting of the brassinosteroid hormone receptor BRI1 are regulated by ubiquitination. Nat Commun 6: 6151

Masclaux FG, Galaud JP, Pont-Lezica R (2005) The riddle of the plant vacuolar sorting receptors. Protoplasma 226: 103–108

Mattera R, Guardia CM, Sidhu SS, Bonifacino JS (2015) Bivalent motif-ear interactions mediate the association of the accessor protein tspesin with the AP-4 adaptor complex. J Biol Chem 290: 30736–30749

Mayers JR, Hu T, Wang C, Cárdenas JJ, Tan Y, Pan J, Bednarek SY (2017) SCD1 and SCD2 form a complex that functions with the exocyst and RabE1 in exocytosis and cytokinesis. Plant Cell 29: 2610–2625

McCough JJ, Cullen PJ (2013) Clathrin is not required for SNX-BAR-retromer-mediated carrier formation. J Cell Sci 126: 45–52

McMichael CM, Bednarek SY (2013) Cytoskeletal and membrane dynamics during higher plant cytokinesis. New Phytol 197: 1039–1057

McMichael CM, Reynolds GD, Koch LM, Wang C, Jiang N, Nadeau J, Sack FD, Gelderman MB, Pan J, Bednarek SY (2013) Mediation of clathrin-dependent trafficking during cytokinesis and cell expansion by Arabidopsis stomatal cytokinesis defective proteins. Plant Cell 25: 3910–3925

Mosseno N, Blaske T, Nagel MK, Laumann M, Isono E (2019) Preparation of clathrin-coated vesicles from Arabidopsis thaliana seedlings. Front Plant Sci 9: 1972

Mravec J, Petrášek J, Li N, Boeren S, Karlova R, Kitakura S, Parezoňová M, Naramoto S, Nodzynski T, Dhonukshe P, et al. (2011) Cell plate restricted association of DRP1A and PIN proteins is required for cell polarity establishment in Arabidopsis. Curr Biol 21: 1055–1060

Murashige T, Skoog F (1962) A revised medium for rapid growth and bio assays with tobacco tissue cultures. Physiol Plant 15: 473–497

Nagaraj N, Wisniewski J, Geiger T, Cox J, Kircher M, Kelso J, Pääbo S, Mann M (2011) Deep proteome and transcriptome mapping of a human cancer cell line. Mol Syst Biol 7: 1–8

Nagel M, Kalinowska K, Vogel K, Reynolds GD, Wu Z, Anzenberger F, Ichikawa M, Tsutsumi C, Sato MH, Kuster B, et al. (2017) Arabidopsis SH3P2 is an ubiquitin-binding protein that functions together with ESCRT-I and the deubiquitylating enzyme AMSH3. Proc Natl Acad Sci USA 114: E7197–E7204

Narasimhan M, Johnson A, Prizak R, Kaufmann W, Tan S, Casillas-Perez B, Friml J (2020) Evolutionarily unique mechanistic
framework of clathrin-mediated endocytosis in plants. eLife 9: 1–30

Nielsen E, Cheung AY, Ueda T (2008) The regulatory RAB and ARF GTPases for vesicular trafficking. Plant Physiol 147: 1516–1526

Niemes S, Langhans M, Viotti C, Scheuring D, San Wan Yan M, Jiang L, Hillmer S, Robinson DG, Pimpl P (2010) Retromer recycles vacuolar sorting receptors from the trans-Golgi network. Plant J 61: 107–121

Nishimura K, Matsunami E, Yoshida S, Kohata S, Yamauchi J, Jisaka M, Nagaya T, Yokota K, Nakagawa T (2016) The tyrosine-sorting motif of the vacuolar sorting receptor VSR4 from Arabidopsis thaliana, which is involved in the interaction between VSR4 and AP1M2, μ1-adaptin type 2 of clathrin adaptor complex 1 subunits, participates in the post-Golgi sorting of VSR4. Biosci Biotechnol Biochem 80: 694–705

Okekeogbu IO, Pattathil S, González Fernández-Niño SM, Aryan UK, Penning BW, Lao J, Heazlewood J, Hahn MG, McCann MC, Carpita NC (2019) Glycome and proteome components of Golgi membranes are common between two angiosperms with distinct cell-wall structures. Plant Cell 31: 1094–1112

Orci L, Ravazzola M, Amherdt M, Louvard D, Perrelet A (1985) Clathrin-immunoreactive sites in the Golgi apparatus are concentrated at the trans pole in polypeptide hormone-secreting cells. Proc Natl Acad Sci USA 82: 5385–5389

Orr F, Fury F, Warner E, Agar E, Garbarino J, Cabral S, Dubuque M, Butt A, Munson M, Vidal L (2021) Rab-E and its interaction with myosin XI are essential for polarised cell growth. New Phytologist 229: 1924–1936

Papanikou E, Day KJ, Austin J, Glick BS (2015). COPI selectively internalizes its cargo. Mol Biol Cell 16: 791–801

Park M, Song K, Reichardt I, Kim H, Mayer U, Stierhof YD, Jiang L, Hillmer S, Robinson DG, Pimpl P (2010) Retromer recycles vacuolar sorting receptors from the trans-Golgi network. Plant J 61: 107–121

Reynolds GD, August B, Bednarek SY (2014) Preparation of enriched plant clathrin-coated vesicles by differential and density gradient centrifugation. Methods Mol Biol 1209: 163–177

Reynolds GD, Wang C, Pan J, Bednarek SY (2018) Introroads into internalization: five years of endocytic exploration. Plant Physiol 176: 208–218

Robinson DG (2018) Retromer and VSR recycling: a red herring? Plant Physiol 176: 483–484

Robinson DG, Neuhaus JM (2016) Receptor-mediated sorting of soluble vacuolar proteins: myths, facts, and a new model. J Exp Bot 67: 4435–4449

Robinson DG, Pimpl P (2014) Receptor-mediated transport of vacuolar proteins: a critical analysis and a new model. Protoplasma 251: 247–264

Robinson MS (2015). Forty years of clathrin-coated vesicles. Traffic 16: 1210–1238

Rosqueta MR, Davis DJ, Drakakaki G (2018) The plant trans-Golgi network: not just a matter of distinction. Plant Physiol 176: 187–198

Roth TF, Porter KR (1964) Yolk protein uptake in the oocyte of the mosquito Aedes aegypti. L. J Cell Biol 20: 313–332

Rueden CT, Schindelin J, Hiner MC, DeZonia BE, Walter AE, Arena ET, Eliceiri KW (2017) ImageJ2: imageJ for the next generation of scientific image data. BMC Bioinformatics 18: 529

Rutherford S, Moore I (2002). The Arabidopsis Rab GTPase family: another enigma variation. Curr Opin Plant Biol 5: 518–528

Sánchez-Rodríguez C, Shi Y, Kesten C, Zang D, Sancho-Andrés IA, Ivanov A, Lampugnani ER, Sklodowski K, Fujimoto M, Nakano A, et al. (2018). The cellulose synthases are cargo of the TPLATE adaptor complex. Mol Plant 11: 346–349

Sanger A, Hirst J, Davies AK, Robinson MS (2019) Adaptor protein complexes and disease at a glance. J Cell Sci 132: 1–2

Sauer M, Delgadillo MO, Zouhar J, Reynolds GD, Pennington JG, Jiang L, Liljegren SJ, Stierhof YD, Dejaegere G, Otegui MS, et al. (2013). MT1V and MT4V encode plant-specific ENTH and ARF GAP proteins that mediate clathrin-dependent trafficking of vacuolar cargo from the trans-Golgi network. Plant Cell 25: 2217–2235

Schindelin J, Arganda-Carreras I, Frise E, Kaynig V, Longair M, Pietzsch T, Preibisch S, Rueden C, Saalfeld S, Schmid B, et al. (2012). Fiji: an open-source platform for biological-image analysis. Nat Methods 9: 676–682

Schwanhäusser B, Busse D, Li N, Dittmar G, Schuchhardt J, Wolf J, Chen W, Selbach M (2011) Global quantification of mammalian gene expression control. Nature 473: 337–342

Schwihla M, Korbel B (2020) The beginning of the end: initial steps in the degradation of plasma membrane proteins. Front Plant Sci 11: 680

Shevchenko A, Wilm M, Vorm O, Mann M (1996) Mass spectrometric sequencing of proteins silver-stained polyacrylamide gels. Anal Chem 68: 850–858

Shimada T, Fuji K, Tamura K, Kondos M, Nishimura M, Harashima I (2003) Vascular sorting receptor for seed storage proteins in Arabidopsis thaliana. Plant Physiol 131: 16095–16100

Shimizu Y, Takagi J, Ito E, Ito Y, Ebine K, Komatsu Y, Goto Y, Yamada M, Tanaka T, Takeda T, Ito Y, Ebine K, Komatsu Y, Goto Y (2004) Plant cytokinesis requires dopamine beta-hydroxylase. Plant Physiol 136: 918–929

Shishido T, Matsuoka Y, Sugimoto N, Ohashi K, Furutani T, Oda T, Ikebe Y, Inoue T, Kato T, et al. (2013) The clathrin-coated vesicles are involved in the degradation of plasma membrane proteins. Front Plant Sci 11: 680

Shevchenko A, Wilm M, Vorm O, Mann M (1996) Mass spectrometric sequencing of proteins silver-stained polyacrylamide gels. Anal Chem 68: 850–858

Shimada T, Fuji K, Tamura K, Kondo M, Nishimura M, Harashima I (2003) Vascular sorting receptor for seed storage proteins in Arabidopsis thaliana. Proc Natl Acad Sci USA 100: 16095–16100

Shimizu Y, Takagi J, Ito E, Ito Y, Ebine K, Komatsu Y, Goto Y, Sato M, Toyooka K, Ueda T, et al. (2021) Cargo sorting zones in the trans-Golgi network visualized by super-resolution confocal live imaging microscopy in plants. Nat Commun 12: 1–14

Simpson F, Peden AA, Christopoulou L, Robinson MS (1997) Characterization of the adaptor-related protein complex, AP-3. J Cell Biol 137: 835–845

Smith JM, Leslie ME, Robinson SJ, Kurasick DA, Zhang T, Backues SK, Cornish PV, Koo AJ, Bednarek SY, Heese A (2014) Loss of Arabidopsis thaliana clathrin subunit VLS1 reveals separation of innate immune signaling pathways. PLoS Pathog 10: e1004578

Song J, Lee MH, Lee GJ, Yoo CM, Hwang I (2006) Arabidopsis EPSIN1 plays an important role in vacuolar trafficking of soluble cargo proteins in plant cells via interactions with clathrin, AP-1, VTI11, and VSR1. Plant Cell 18: 2258–2274
Proteomic analysis of Arabidopsis CCVs

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Sosa RT, Weber MM, Wen Y, O’Halloran TJ (2012) A single beta adaptin contributes to AP1 and AP2 complexes and clathrin function in Dictyostelium. Traffic 13: 305–316

Speth EB, Imboden L, Hauck P, He SY (2009) Subcellular localization and functional analysis of the Arabidopsis GTPase RabE. Plant Physiol 149: 1824–1837

Stolc V, Samanta MP, Tongprisit W, Sethi H, Liang S, Nelson DC, Hegeman A, Nelson C, Rancour D, Bednarek S, et al. (2005) Identification of transcribed sequences in Arabidopsis thaliana by using high-resolution genome tile arrays. Proc Natl Acad Sci USA 102: 4453–4458

Suwastika IN, Uemura T, Shinya T, Sato MH, Takeyasu K (2008) SYP71, a plant-specific Qc-SNARE protein, reveals dual localization to the plasma membrane and the endoplasmic reticulum in Arabidopsis. Cell Struct Funct 33: 185–192

Takamori S, Holt M, Stenius K, Leemke E, Gronborg M, Riedel D, Urlaub H, Schenck S, Brügger B, Ringler P, et al. (2006) Molecular anatomy of a trafficking organelle. Cell 127: 831–846

Takemoto K, Ebine K, Askanji JC, Krüger F, Gonzalez ZA, Ito E, Goh T, Schumacher K, Nakano A, Ueda T (2018) Distinct sets of tetherying complexes, SNARE complexes, and Rab GTPases mediate membrane fusion at the vacuole in Arabidopsis. Proc Natl Acad Sci USA 115: E2457–E2466

Taylor M, Perraid D, Merrifield C (2011) A high precision survey of the molecular dynamics of mammalian clathrin-mediated endocytosis. PLoS Biol 9: e1000604

Teh OK, Shimono Y, Shirakawa M, Fukao Y, Tamura K, Shimada T (2013) The AP-1 mu adaptin is required for KNOLLE localization at the cell plate to mediate cytokinesis in Arabidopsis. Plant Cell Physiol 43: 838–847

Tong P, Kornfeld S (1989) Ligand interactions of the cation-dependent mannose 6-phosphate receptor: comparison with the cation-independent mannose 6-phosphate receptor. J Biol Chem 264: 7970–7975

Uemura T, Kim H, Saito C, Ebine K, Ueda T, Schulze-Lefert P, Nakano A (2012) QA-SNAREs localized to the trans-Golgi network regulate multiple transport pathways and extracellular disease resistance in plants. Proc Natl Acad Sci USA 109: 1784–1789

Uemura T, Sato MH, Takeyasu K (2005) The longin domain regulates subcellular targeting of VAMP7 in Arabidopsis thaliana. FEBS Lett 579: 2842–2846

Vallis Y, Wigge P, Marks B, Evans PR, McMahon HT (1999) Importance of the pleckstrin homology domain of dynamin in clathrin-mediated endocytosis. Curr Biol 9: 257–260

Van Damme D, Coutuer S, De Rycke R, Bouget F, Inzé D, Geelen D (2006) Somatic cytokinesis and pollen maturation in Arabidopsis depend on TPLATE, which has domains similar to coat proteins. Plant Cell 18: 3502–3518

Viotti C, Bubek J, Stierhof YD, Krebs M, Langhans M, van den Berg W, van Dongen W, Richter S, Geldner N, Takano J, et al. (2010). Endocytic and secretory traffic in Arabidopsis merge in the trans-Golgi network/early endosome, an independent and highly dynamic organelle. Plant Cell 22: 1344–1357

Vollmer AH, Youseff NN, DeWald DB (2011) Unique cell wall abnormality in the putative phosphoinositide phosphatase mutant AtSAC9. Planta 234: 993–1005

Wang C, Hu T, Yan X, Meng T, Wang Y, Wang Q, Zhang X, Gu Y, Sánchez-Rodriguez C, Gadeyne A, et al. (2016). Differential regulation of clathrin and its adaptor proteins during membrane recruitment for endocytosis. Plant Physiol 171: 215–229

Wang C, Yan X, Chen Q, Jiang N, Fu W, Ma B, Liu J, Li C, Bednarek SY, Pan J (2013) Clathrin light chains regulate clathrin-mediated trafficking, auxin signaling, and development in Arabidopsis. Plant Cell 25: 499–516

Wang J, MYlE E, Johnson A, Besbrugge N, De Jaeger G, Friml J, Ple Skot R, Van Damme D (2020) High temporal resolution reveals simultaneous plasma membrane recruitment of TPLATE complex subunits. Plant Physiol 183: 986–997

Wang J, Yperman K, Grones P, Jiang Q, Dragwidge J, MYlE E, Mor E, Nolf J, Eeckhout D, De Jaeger G, et al. (2021) Conditional de-stabilization of the TPLATE complex impairs endocytic internalization. Proc Natl Acad Sci USA 118: e2023456118 https://doi.org/10.1073/pnas.2023456118

Wang X, Cai Y, Wang H, Zeng Y, Zhuang X, Li B, Jiang L (2014) Trans-Golgi network-located AP1 gamma adaptins mediate dileucine motif-directed vacuolar targeting in Arabidopsis. Plant Cell 26: 4102–4118

Williams ME, Torabinjed AJ, Cohick E, Parker K, Drake EJ, Thompson JE, Horter M, Dewald DB (2005) Mutations in the Arabidopsis phosphoinositide phosphatase gene SAC9 lead to overaccumulation of PtdIns(4,5)P2 and constitutive expression of the stress-response pathway. Plant Physiol 138: 686–700

Wolfenstetter S, Wirsching P, Dotzauer D, Schneider S, Sauer N (2012) Routes to the tonoplast: the sorting of tonoplast transporters in Arabidopsis mesophyll protoplasts. Plant Cell 24: 215–232

Wu B, Guo W (2015) The exocyst at a glance. J Cell Sci 128: 2957–2964

Wu M, Wu X (2021) A kinetic view of clathrin assembly and endocytic cargo sorting. Curr Opin Cell Biol 71: 130–138

Yan X, Wang Y, Xu M, Dahhan D, Liu C, Zhang Y, Lin J, Bednarek S, Pan J (2021) Cross-talk between clathrin-dependent post-Golgi trafficking and clathrin-mediated endocytosis in Arabidopsis root cells. Plant Cell 33: 3057–3075

Yao L, Jannmy P, Frigeri LG, Han W, Fujita J, Kawakami Y, Apgar JR, Kawakami T (1999). Pleckstrin homology domains interact with filamentous actin. J Biol Chem 274: 19752–19761

Yperman K, Papageorgiou A, Merreron R, De Munck S, Bloch Y, Eeckhout D, Jiang Q, Tack P, Grigoryan R, Evangelidis T, et al. (2021) Distinct EH domains of the endocytic TPLATE complex confer lipid and protein binding. Nat Comm 12: 1–11

Yun HS, Kwon C (2017) Vesicle trafficking in plant immunity. Curr Opin Plant Biol 40: 34–42

Zhang Y, Persson S, Hirst J, Robinson MS, Van Damme D, Sanchez-Rodriguez C (2015). Change your TPLATE, change your fate: cmae and beyond. Trends Plant Sci 20: 41–48

Zhao Y, Yan A, Feijo JA, Furutani M, Takenawa T, Huang I, Fu Y, Yang Z (2010) Phosphoinositides regulate clathrin-dependent endocytosis at the tip of pollen tubes in Arabidopsis and tobacco. Plant Cell 22: 4031–4044

Zhou Y, Yang Z, Niu Y, Fan T, Qian D, Luo C, Shi Y, Li S, An L, Xiang Y (2020) The tip-localized phosphatidylinerine established by Arabidopsis ALA3 is crucial for Rab GTPase-mediated vesicle trafficking and pollen tube growth. Plant Cell 32: 3170–3187

Zhu X, Li S, Pan S, Xin X, Gu Y (2018) CS11, PATROL1, and exocyst complex cooperate in delivery of cellulose synthase complexes to the plasma membrane. Proc Natl Acad Sci USA 115: E3578–E3587

Zouhar J, Muñoz A, Rojo E (2010). Functional specialization within the vacuolar sorting receptor family: VSR1, VSR3 and VSR4 sort vacuolar storage cargo in seeds and vegetative tissues. Plant J 64: 577–588

Zouhar J, Rojo E, Bassham DC (2009). AtVPS45 is a positive regulator of the SYP41/SYP61/VTI12 SNARE complex involved in trafficking of vacuolar cargo. Plant Physiol 149: 1668–1678

Zouhar J, Sauer M (2014) Helping hands for budding prospects: ENTH/ANTH/VHS accessory proteins in endocytosis, vacuolar transport, and secretion. Plant Cell 26: 4232–4244

Zwiewka M, Feraru E, Möller B, Hwang I, Feraru MI, Klein-Vehn J, Weijers D, Friml J (2011) The AP-3 adaptor complex is required for vacuolar function in Arabidopsis. Cell Res 21: 1711–1722
