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Journal research area most appropriate for the paper: Cell Biology
Dynamic Response of Prevacuolar Compartments to Brefeldin A in Plant Cells

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This work was supported by grants from the Research Grants Council of Hong Kong (CUHK4156/01M, CUHK4260/02M, CUHK4307/03M and CUHK4580/05M), NSF of China (30529001) and CUHK Scheme C to L. Jiang.

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**ABSTRACT**

Little is known about the dynamics and molecular components of plant prevacuolar compartments (PVCs) in the secretory pathway. Using transgenic tobacco BY-2 cells expressing membrane-anchored yellow fluorescent protein (YFP) reporters marking Golgi or PVCs, we have recently demonstrated that PVCs are mobile multivesicular bodies (MVBs) defined by vacuolar sorting receptor (VSR) proteins. Here we demonstrate that Golgi and PVCs have different sensitivity in response to Brefeldin A (BFA) treatment in living tobacco BY-2 cells. BFA at low concentrations (5 to 10 µg mL\(^{-1}\)) induced YFP-marked Golgi stacks to form both ER-Golgi hybrid structures and BFA-induced aggregates, but had little effect on YFP-marked PVCs in transgenic BY-2 cells at both confocal and immunoEM levels. However, BFA at high concentrations (50 to 100 µg mL\(^{-1}\)) caused both YFP-marked Golgi stacks and PVCs to form aggregates in a dosage-dependent and time-dependent manner. Normal Golgi or PVC signals can be recovered upon removal of BFA from the culture media. Confocal immunofluorescence and immunoEM studies with specific organelle markers further demonstrate that the PVC aggregates are distinct but physically associated with Golgi aggregates in BFA-treated cells and that PVCs might lose their internal vesicle structures at high BFA concentration. In addition, VSR-marked PVCs in root tip cells of tobacco, pea, mungbean and *Arabidopsis* upon BFA treatment are also induced to form similar aggregates. Thus, we have demonstrated that BFA’s effects are not limited to ER and Golgi but extended to PVC in the endomembrane system, which might provide a quick tool for distinguishing Golgi from PVC for its identification and characterization, as well as a possible new tool in studying PVC-mediated protein traffic in plant cells.
Introduction

All eukaryotic cells have secretory and endocytic pathways that are composed of several functionally distinct membrane-bounded compartments each with characteristic proteins, which can be used as markers to define these compartments. Prevacuolar compartments or late endosomes are intermediate organelles where secretory and endocytic traffic leading to vacuole or lysosome merges (Bethke and Jones, 2000; Jiang and Rogers, 2003; Lam et al., 2005; Lemmon and Traub, 2000; Maxfield and McGraw, 2004). These prevacuolar or late endosomal compartments characteristically possess internal microvesicles and are thus called “multivesicular endosomes or bodies” (MVB). In animal cells these microvesicles appear to originate in the early or recycling endosomes (Parton et al., 1992), and have a different protein composition to the limiting membrane (Griffiths et al., 1990; Kobayashi et al., 1998). The microvesicles and the soluble content of these MVBs are most likely delivered into the interior of the lysosomal compartment via direct fusion (Luzio et al., 2000; Katzmann et al., 2002).

In spite of the obviously important role of PVCs in mediating protein traffic to vacuoles in the secretory and endocytic pathways, the identification and characterization of plant PVCs, both functionally and morphologically, have been challenging due to the complexity of the plant vacuolar system, the existence of multiple transport pathways leading to distinct vacuoles, and the lack of good markers (antibody and reporter) to be used to define or identify PVCs (Jiang and Rogers, 1998; Bethke and Jones, 2000; Jiang and Rogers, 2003; Lam et al., 2005). Putative plant PVCs were identified using antibodies specific for the Arabidopsis syntaxin Pep12p and the pea vacuolar sorting receptor (VSR) BP-80 (Bassham et al., 1995; Conceição et al., 1997; Paris et al., 1997). Studies on the relative distribution of VSR proteins between Golgi stacks and PVCs in various plant cells demonstrated that VSR proteins largely separated from Golgi markers and thus must be concentrated on PVCs under steady state conditions (Li et al., 2002). In addition, a chimeric BP-80 reporter containing the BP-80 transmembrane domain (TMD) and cytoplasmic tail (CT) colocalized with endogenous VSR
proteins when the reporter was expressed in transgenic tobacco culture cells (Jiang and Rogers, 1998). Therefore, both VSR proteins and the BP-80 reporter can be used as markers to define and identify plant PVCs.

Multivesicular body type organelles have been described some time ago in plant cells (Robinson et al., 2000). Typically, MVBs in plant cells are somewhat smaller than a Golgi stack and have an osmiophilic plaque somewhere on their surface, a feature shared by multivesicular endosomes in mammalian cells (Raposo et al., 2001; van Dam and Stoorvogel, 2002). Conclusive proof that plant PVCs are MVBs was provided by a recent study (Tse et al., 2004) that demonstrated the presence of VSR proteins in MVBs by immunogold labeling of tobacco BY-2 cell sections, where VSR labeling was restricted to the boundary membrane rather than the internal vesicles of the MVBs, suggesting that only a small proportion of the VSRs that reach the PVC at any one time are destined for degradation. Further confirmation that MVBs are PVCs and are distinct from Golgi stacks comes from examining the differential effects of the drug wortmannin. Whereas wortmannin at reversible concentrations (10 to 30 µM) primarily causes MVBs to vacuolate and lose their osmiophilic plaques, it leads to a reduction in the number of internal vesicles as well (Tse et al., 2004). These changes have also been recorded for multivesicular endosomes in mammalian cells (Bright et al., 2001; Sachse et al., 2002). Thus, wortmannin appears to be a useful MVB/PVC-specific drug, and is becoming a very useful tool for perturbing transport to the lytic vacuole in plant cells (daSilva et al., 2005).

Similar to wortmannin, the fungal macrocyclic lactone brefeldin A (BFA) has been widely used in studying protein trafficking in the secretory and endocytic pathways of eukaryotic cells. In mammalian cells, BFA targets a subclass of Sec7-type guanine nucleotide exchange factors (GEFs) required for converting the GDP-ribosylation factor 1 (Arf1p) to its GTP form (Jackson and Casanova, 2000), which then binds to the membrane to recruit coatomer and leads to the production of COPI-coated vesicles. Whereas, BFA disrupts the formation of COPI-coated
vesicles by inhibits the conversion of the Arf1p from its GDP form to a GTP form by locking the reaction intermediate in an inactive complex. Plant cells may have similar molecular target of BFA. For example, the *Arabidopsis* gene GNOM encoding an Arf GEF has been identified (Busch et al., 1996) and localized to the Golgi apparatus (Jackson and Casanova, 2000). In addition, the rapid release of COPI coats proteins from the Golgi stacks into the cytosol was also demonstrated in tobacco BY-2 cells using anti-Atγ-COPI antibodies in confocal immunofluorescence (Ritzenthaler et al., 2002).

The Golgi apparatus is likely the initial site in response to BFA in both plant and mammalian cells (Sciaky et al., 1997; Nebenfuhr et al., 2002). In addition to blocking protein transport from ER to the Golgi, BFA at low concentrations (2 to 10 µg mL⁻¹) also leads to rapid and dramatic changes in the morphology of the Golgi apparatus of mammalian cells where the Golgi apparatus gets extensively tubulated and fuses with the endoplasmic reticulum (Klausner et al., 1992). In addition, BFA may also act on endosomes and induce endosomes to become tubulated in mammalian cells (Lippincott-Schwartz et al., 1991; Hunziker et al., 1991; Wood et al., 1991). In plant cells, BFA also induced the Golgi apparatus to form ER-Golgi hybrid structures (Ritzenthaler et al., 2002), BFA compartments (Satiat-Jeunemaitre et al., 1996; Wee et al., 1998; Baldwin et al., 2001; Tse et al., 2004), or the loss of Golgi cis cisternae (Hess et al., 2000). In addition, BFA may induce trans-Golgi network (TGN) to incorporate into BFA compartments in *Arabidopsis* root cells (Dettmer et al., 2006). Moreover, some studies also demonstrated BFA (at 25 to 50 µM mL⁻¹) induces the endosomal compartments form aggregates in *Arabidopsis* root cells (Geldner et al., 2003; Grebe et al., 2003). However, little is known about the possible effects of BFA on PVCs/MVBs in the secretory pathway of plant cells.

We previously demonstrated that VSR proteins can be used as markers to define plant PVCs (Li et al., 2002). Recently, we developed two transgenic BY-2 cell lines expressing the Golgi marker GONST1-YFP reporter and the PVC marker YFP-BP-80 reporter and
demonstrated that the YFP-BP-80 reporter and VSR proteins defined PVCs with characteristics of multivesicular bodies (MVBs) in BY-2 cells (Tse et al., 2004). In addition, BFA at concentration less than 10 µg mL\(^{-1}\) induced the YFP-marked Golgi stacks but not the YFP-marked PVCs to form enlarged compartments in transgenic BY-2 cells. Therefore, these transgenic BY-2 cells are useful tools in studying the dynamic of Golgi and PVC in response to external stimulates in living cells.

In this study, we extended our understanding on BFA effects on PVCs/MVBs in plant cells. We used these two transgenic BY-2 cell lines expressing YFP/GFP reporters marking Golgi and PVC to test the hypothesis that BFA also affects other endomembrane organelles in addition to the ER and Golgi apparatus. Here we demonstrated that YFP-marked PVC and YFP-marked Golgi stacks showed different sensitivity in response to BFA treatment. BFA at low concentrations (5 to 10 µg mL\(^{-1}\)) induced YFP-marked Golgi stacks but not YFP-marked PVC to form aggregates. However, BFA at higher concentrations (50 or 100 µg mL\(^{-1}\)) induced both YFP-marked PVC and Golgi stacks to form typical enlarged structures in a dosage- and time-dependent manner. Confocal immunofluorescence studies demonstrated that BFA-induced aggregates derived from YFP-marked PVCs colocalized with VSR proteins but kept physically distinct from the Golgi-derived aggregates in BFA-treated cells. Further immunoEM and structural EM studies identified unique structures of Golgi- and PVC-derived aggregates. In addition, BFA also induced VSR-marked PVCs to form similar aggregates in root tip cells of pea, tobacco, mungbean and *Arabidopsis*. 
RESULTS

Development of the GFP-BP-80 Transgenic Cell Line

We have previously demonstrated that VSR proteins are markers for defining PVCs in plant cells (Li et al., 2002). More recently, we have developed two transgenic tobacco BY-2 cell lines expressing the Golgi reporter GONST1-YFP and the PVC reporter YFP-BP-80 and demonstrated that these two reporters colocalized with ManI and VSR antibodies, respectively (Tse et al., 2004). We further demonstrated that PVCs in tobacco BY-2 cells are multivesicular bodies (MVBs) that are marked by VSR proteins (Tse et al., 2004).

As a first step to compare reporter-marked Golgi to PVC directly in transgenic living tobacco BY-2 cells, we further developed a new transgenic tobacco BY-2 cell line expressing the GFP-BP-80 fusion in which the GFP was used to replace the YFP in the previous YFP-BP-80 construct (Tse et al., 2004). Similar to the transgenic cell line expressing the Golgi marker GONST1-YFP (Fig. 1A, panel 1), typical punctate fluorescent signals were detected from cells expressing the GFP-BP-80 (Fig. 1A, panel 2). Furthermore, organelles marked by either YFP-BP-80 or GFP-BP-80 (green) were mostly (more than 92% based on calculation from more than twenty labeled cells) colocalized with VSR antibodies (Fig. 1A, panels 3 & 4), indicating the PVC localization of both YFP-BP-80 and GFP-BP-80 in transgenic tobacco BY-2 cells.

The Golgi localization of the GONST1-YFP reporter and PVC localization of the GFP/YFP-BP-80 reporter in transgenic BY-2 cells were further confirmed by immunoEM studies where GFP antibodies specifically labeled Golgi stacks (Fig. 1B, left panel) or MVB (Fig. 1B, right panel) in transgenic BY-2 cells expressing either the Golgi- or PVC-reporters. Thus, similar to YFP-BP-80, the GFP-BP-80 reporter served as a tool for defining PVCs/MVBs in tobacco BY-2 cells. Therefore, from now on, both YFP-BP-80 and GFP-BP-80 cell lines were used as PVC markers in this study.
**PVC and Golgi have Different Sensitivity to Brefeldin A**

Brefeldin A (BFA) has been a useful tool in studying protein trafficking in the secretory pathway because this drug induced Golgi stacks to form enlarged compartments and prevented protein traffic from ER to Golgi (Hess et al., 2000; Jiang and Rogers, 1998; Ritzenthaler et al., 2002). Using transgenic tobacco BY-2 cell lines expressing Golgi and PVC reporters, we have recently demonstrated that Golgi and PVC had different sensitivity to wortmannin in which the drug induced the PVC but not Golgi to dilate (Tse et al., 2004). Since the Golgi and PVC are closely related organelles in the secretory pathway, using transgenic BY-2 cell lines expressing PVC reporters, we therefore wanted to study the possible effects of BFA on PVCs. As a control, a transgenic cell line expressing the Golgi reporter was used.

We first performed a dosage response experiment in which Day-3 transgenic cell lines expressing either the Golgi or PVC reporters were incubated with various concentrations of BFA (from 0 to 100 µg mL⁻¹) for one hour before the treated cells were collected and used in confocal imaging. Day-3 BY-2 cells have been used in our drug treatments studies because these cells are at their log phase (Matsuoka et al., 2004) and XFP-BP-80 fusions remain in PVCs as punctate patterns which will be targeted to vacuoles as diffusion patterns at Day-6 and Day-7 (Mitsuhashi et al., 2000; Lo and Jiang, 2006). As shown in Figure 2, in cells expressing the Golgi marker GONST1-YFP, typical enlarged compartments were observed in all cells treated with BFA at tested concentrations from 5 µg mL⁻¹ to 100 µg mL⁻¹ (Fig. 2A, arrows) and the average sizes of the visible BFA-induced aggregates were larger in cells treated with higher concentrations of BFA (Fig. 2C). To our surprise, in cells expressing the PVC marker YFP-BP-80, BFA at high concentrations (50 to 100 µg mL⁻¹) also induced the YFP-marked PVCs to form similar aggregates (Fig. 2B, arrows; Fig 2C), even though low BFA concentrations (5 to 10 µg mL⁻¹) did not have any visible effect on the size and number of YFP-marked PVCs (Fig. 2C and D). Similar results were obtained when the cells were treated with BFA with various concentrations for two hours before confocal imaging (Supplemental...
Fig. 1). In addition, similar results were also obtained when another transgenic cell line expressing the PVC marker GFP-BP-80 was used in a similar study (data not shown).

BFA at 5 to 10 µg mL\(^{-1}\) induced the GONST1-YFP-marked Golgi to form aggregates in transgenic BY-2 cells (Fig. 2A), such result is consistent with several previous studies showing the BFA-induced Golgi aggregation in different cell types (Wee et al., 1998; Baldwin et al., 2001; Tse et al., 2004; Miao et al., 2006). However, several other studies showed that BFA at such concentration also induced GFP-marked Golgi to form ER patterns in both BY-2 and tobacco leaf cells (Ritzenthaler et al., 2002; Saint-Jore et al., 2002). To find out the possible causes for such observed difference, we collected BY-2 cells at Day-3 and Day-6 for BFA treatment at 10 µg mL\(^{-1}\) for one hour before confocal imaging. Day-3 and Day-6 cells represent log phase and stationary phase, respectively (Matsuoka et al., 2004). As shown in Supplemental Figure 2, similar punctate patterns were observed in GONST1-YFP cells collected at either Day 3 or Day 6 (left panels 1 & 6). At the end of one-hour BFA treatment, major aggregation patterns were observed from Day-3 cells (left panels 2 & 3) or Day-2 cells (not shown), but typical ER patterns were observed from BFA-treated Day-6 cells (left panels 5 & 6) or Day-7 cells (not shown), indicating that the physiological status of GONST1-YFP BY-2 cells may affect its responsive patterns to BFA treatment. However, when the same BFA treatments were performed using transgenic Man1-GFP BY-2 cells, ER patterns were observed from both BFA-treated Day-3 (right panels 2 & 3) and Day-6 cells (right panels 5 & 6). We do not know if such observed difference is due to the difference between the trans-Golgi GONST1 and the cis-Golgi Man1 or due to the different cell line used, but such difference shall not affect this study on the effects of BFA on PVCs/MVBs where the GONST1-YFP cells were used as a control for XFP-BP-80-marked PVC/MVBs in BY-2 cells (see Discussion).

To study further the dynamics of the BFA-induced response of PVCs to high concentrations of BFA, a time-course study was carried out using these two transgenic cell lines. Day-3 individual transgenic cell lines were first incubated with BFA at 50 µg mL\(^{-1}\), a
minimum concentration that induced PVCs to form aggregates (Fig. 2), followed by sample collection at indicated time after BFA treatment for confocal imaging. As shown in Figure 3A, in cells expressing the Golgi marker GONST1-YFP, enlarged BFA-induced aggregates were already detected 15 minutes after the BFA treatment and these aggregated structures remained throughout the two-hour period of BFA treatment. In contrast, in cells expressing the PVC marker YFP-BP-80, no BFA-induced YFP-marked aggregates were detected (Fig. 3B and C) and the number of YFP-marked PVC remained unchanged (Fig. 3D) during the first 30 min of BFA treatment, but similar aggregates were observed from 45 minutes after BFA treatment and these aggregates remained detectable thereafter for two hours (Fig. 3B and C). Thus, when comparing to Golgi stacks, a 30-min delay was observed for PVCs to form BFA-induced aggregates in response to BFA at 50 µg mL⁻¹ under these conditions.

**Recovery of PVCs from BFA Treatment**

Low concentrations of BFA (typically between 5 and 10 µg mL⁻¹) have been commonly used in studying protein trafficking and organelle dynamics in tobacco BY-2 cells (Nebenfuhr et al., 2002). In cells treated with BFA at this low concentration, the normal Golgi apparatus can usually recover upon removing the drug from the culture media. It is thus believed that the formation of BFA-induced compartments and ER-Golgi hybrids is not caused by the toxic effect of BFA on BY-2 cells.

To find out if BFA at 50 or 100 µg mL⁻¹ is toxic to BY-2 cells and causes permanent damage, we performed a recovery study. Day-3 transgenic cells expressing the Golgi marker GONST1-YFP and PVC marker GFP-BP-80 were treated with BFA at 100 µg mL⁻¹ for one hour, followed by washing off the BFA with fresh media and then sample collections at indicated time points upon recovery for confocal imaging. As shown in Figure 4A, in cells expressing the Golgi marker GONST1-YFP, fluorescent signals in both aggregates and an ER pattern were detected 15 and 45 minutes after BFA washing, which was then followed by the
disappearance of aggregates, reappearance of punctate signals and weakening of the ER pattern after one hour, and a fully punctate pattern was eventually observed after two hours (Fig. 4A). These results indicate that the Golgi pattern can be recovered fully upon BFA removal. Similar patterns of changes were also detected in cells expressing the PVC marker GFP-BP-80 upon the removal of BFA and again normal punctate signals were recovered after two hours (Fig. 4B). Furthermore, the BFA-treated BY-2 cells during the recovery period still looked normal as judged by their appearance in the DIC images. Similar results were obtained when cells were treated with BFA at 50 µg mL⁻¹ for one hour, followed by washing off the BFA in recovery study in these two cell lines (Supplemental Fig. 3). These results demonstrate that BFA at 50 or 100 µg mL⁻¹ did not permanently damage the treated BY-2 cells and that the BFA-induced aggregates from either Golgi or PVC organelles can be fully recovered into its typical punctate patterns upon removal of BFA. Therefore, both Golgi apparatus and PVCs in transgenic BY-2 cells demonstrated the ability of recovery even in high dosage of BFA treatment at 50 or 100 µg mL⁻¹.

**BFA-induced Compartments Derived from Golgi and PVC Remain Distinct but are Closely Associated**

BFA at 5 to 10 µg mL⁻¹ induced the formation of aggregates that may represent the ER-Golgi hybrid structures in BY-2 cells (Ritzenthaler et al., 2002). Our results thus far indicated that BFA at 50 or 100 µg mL⁻¹ induced both Golgi apparatus and PVCs to form aggregates in transgenic BY-2 cells. Since both YFP-marked Golgi and PVC exhibit similar punctate patterns before BFA treatment and form similar aggregates in the presence of BFA, it is thus possible that the BFA-induced aggregates were derived from Golgi and PVC fusion.

To find out if the BFA-induced aggregates derived from Golgi and PVC remain distinct or fusion upon BFA treatment, we performed the following immunofluorescent labeling study. Day-3 transgenic BY-2 cells expressing GONST1-YFP and YFP-BP-80 reporters were treated
with BFA at 50 µg mL⁻¹ for one and two hours, followed by fixation and labeled with VSR antibodies for subsequent confocal imaging. As shown in Figure 5, the PVC/MVB marker VSR antibodies detected aggregates (red) in fixed transgenic cells expressing either the Golgi or PVC reporter (Fig. 5). Interestingly, in fixed transgenic BY-2 cells expressing the PVC marker YFP-BP-80, the BFA-induced aggregates derived from the YFP-marked PVCs (green) colocalized (more than 90%) with aggregates detected by VSR antibodies (red) (Fig. 5 panel 1, arrow heads in merged image). In contrast, in fixed BY-2 cells expressing the Golgi marker GONST1-YFP, the BFA-induced aggregates derived from the GONST1-YFP-marked Golgi (green) remained largely distinct (more than 90%) from the aggregates marked by the PVC marker VSR antibodies (red) at one or two hours after the BFA treatment (Fig. 5 panels 2 and 3, arrowhead vs. arrow in merged images). In addition, most of these aggregates derived from Golgi and PVC showed close association and tended to join together. These results demonstrate that BFA-induced aggregates derived from Golgi stacks are different from those derived from PVCs in BY-2 cells and that VSR proteins remain intact within the PVC-derived aggregates in BY-2 cells.

To further investigate the nature and relationship of the BFA-induced aggregates derived from Golgi and PVC in response to BFA treatment, we then performed a time-course experiment using transgenic cells expressing the Golgi marker GONST1-YFP. These transgenic cells were first incubated with BFA at 50 µg mL⁻¹, followed by sample collection and fixation at indicated time point before the fixed cells were used in labeling with VSR antibodies for subsequent confocal imaging analysis. As shown in Figure 6, BFA-induced aggregates marked by the Golgi marker GONST1-YFP (green) were already detected 10 min after BFA treatment and these aggregates remained similar throughout out the 2 hr treatment period (Fig. 6, panels 1 to 3, green). In contrast, the VSR-marked PVCs (red) remained unchanged during the first 10 min while VSR-marked aggregates became visible and gradually increased in two hours (Fig. 6, panels 1 to 3, red). In addition, aggregates derived from Golgi
and PVC remained distinct but closely associated (Fig. 6, merged images in panels 2 and 3). The differential response of Golgi and PVC to BFA treatment was consistent with results obtained from living cells (Fig. 2 and Fig. 3).

**The BFA Effects on the Golgi and PVC are Pharmacologically Distinguishable**

To find out if the effect of BFA on the formation of PVC-derived aggregates is a primary effect on a PVC target or a secondary effect of transport from the Golgi to PVC, we performed the following experiments using two phospholipase A₂ inhibitor, ONO-RS-082 and bromoenol lactone (BEL), that have been used to inhibit the effect of BFA in mammalian cells (de Figueiredo et al., 1998 and 1999). In addition, ONO in mammalian cells targets and disrupts the Golgi apparatus (de Figueiredo et al., 1999). Transgenic BY-2 cells expressing the Golgi marker GONST1-YFP were pretreated with ONO-RS-082 at 10 µM for 15 min, followed by addition of BFA at 10 or 100 µg mL⁻¹ for another one hour before confocal imaging. As shown in Figure 7, BFA at either 10 or 100 µg mL⁻¹ induced the GONST1-YFP-marked Golgi to form aggregates in cells without pretreatment with ONO-RS-082 (Fig. 7A, panels 1 and 2). Similar aggregates were observed in cells pretreated with ONO-RS-082 at 10 µM for 15 min followed by BFA treatment at either 10 or 100 µg mL⁻¹ (Fig. 7A, panels 3 and 4), indicating that this drug did not block the BFA effect on Golgi at either low or high concentrations. However, in transgenic BY-2 cell expressing the PVC marker GFP-BP-80, BFA at 100 µg mL⁻¹ induced the GFP-marked PVC to form aggregates in cells without pretreatment with ONO-RS-082 (Fig. 7A, panel 2), but no such aggregates were observed in cells pre-treated with ONO-RS-082 at 10 µM for 15 min (Fig. 7A, panel 4), indicating that the drug prevents the formation of BFA-induced PVC aggregates. When a similar drug BEL was used to pre-treat the cells before BFA treatment at 100 µg mL⁻¹, it did not block the formation of BFA-induced aggregates from Golgi or PVC (Supplemental Fig. 4), indicating that ONO-RS-082 was specific for PVC. These results clearly demonstrated that BFA acts on different targets on Golgi and PVC,
whereas ONO-RS-082 did not prevent the formation of BFA-induced aggregates from Golgi, the drug blocked the formation of BFA-induced PVC aggregates. Therefore, it is likely that BFA acts on distinct targets on Golgi and PVC respectively, even though its molecular mechanism of actions on Golgi and PVC remained to be illustrated in plant cells.

**Morphology of BFA-induced Golgi and PVC Aggregates**

Confocal immunofluorescent studies thus far demonstrate that BFA at high concentration (50 to 100 µg mL$^{-1}$) induced both Golgi and PVC to form aggregates that are closely adjacent to each other. To further identify and study the morphological structures of these BFA-induced aggregates and their relationship, immunoEM studies were then carried out where GFP and VSR antibodies were used to detect Golgi stacks and PVCs respectively in transgenic GONST1-YFP BY-2 cells. Day-3 GONST1-YFP transgenic BY-2 cells were treated with BFA at 100 µg mL$^{-1}$ for one hour before they were fixed by glutaraldehyde and embedded in Lowicryl – HM20. Ultra-thin sections were then prepared for immunoEM labeling. As shown in Figure 8, normal Golgi apparatus (panel A) was specifically labeled by GFP antibodies in the untreated control cells. In contrast, in BFA-treated cells, curved Golgi aggregates were labeled specifically by GFP antibodies (Fig. 8, panel B), which may likely represent the BFA-induced aggregates derived from Golgi apparatus.

Because no visible MVBs were detected in ultra-thin sections prepared from Lowicryl - HM20 blocks, we therefore prepared samples using a high-pressure freezing and frozen substitution protocol for immunoEM studies (Tse et al., 2004). As shown in Figure 8, in the control untreated cells, multivesicular PVCs were clearly labeled by VSR antibodies and the labeling was mainly on the peripheral regions (Fig. 8, panels C and D). Similarly, in BFA-treated cells, similar sizes (100 to 200 nm) of vesicles were also labeled specifically by VSR antibodies to their peripheral membranes (Fig. 8, panels E to H). Moreover, unlike untreated cells where PVC/MVBs were often identified individually, these VSR-labeled
vesicles were often detected as groups that tended to pack together and thus might be derived from PVC aggregation or vesiculation (Fig. 8, panels E to I). However, these aggregated vesicles containing VSR proteins seem to lose their MVB appearance because no visible internal vesicles were detected from these VSR-labeled PVCs under these conditions (see Fig. 8 E and G as examples).

To study further the details of these Golgi- or PVC-derived aggregates in response to BFA treatment, we then performed structural TEM studies using conventional chemical fixation and subsequent embedding in Spurr resin as previously described (Tse et al., 2004). First, Day-3 transgenic GONST1-YFP BY-2 cells were treated with BFA at 0, 10 (controls) and 100 µg mL\(^{-1}\) for one hour. Cells were then prepared for TEM analysis as described (Tse et al., 2004). As shown in Supplemental Figure 5, in untreated control cells, normal Golgi apparatus (panel A) and typical multivesicular PVCs (panel B) were detected. In addition, in cells treated with low concentration of BFA (at 10 µg mL\(^{-1}\)), typical BFA-induced Golgi aggregates with visible ER-Golgi hybrids (panel C) were observed while normal multivesicular PVCs remained unchanged in size and morphology (panel D). However, in cells treated with BFA at 100 µg mL\(^{-1}\), many aggregated vesicles with sizes of 100 to 200 nm but lacking internal vesicular structures were often observed in these BFA-treated cells (Fig. 9, panel A), a result consistent with the immunoEM study in which aggregated vesicles were found to be labeled by VSR antibodies in BFA-treated cells (Fig. 8, panels E to H). Such detection was due to BFA treatment because no such aggregated vesicles around the Golgi apparatus were observed in untreated cells (data not shown). Thus, these aggregated vesicles may represent BFA-induced aggregates derived from PVCs in BY-2 cells. Similar vesicle aggregates were also observed from Arabidopsis root tip cells treated with BFA at 100 µg mL\(^{-1}\) for one hour (Fig. 9B), indicating that such BEA effect on PVC is not limited to tobacco BY-2 cells.
BFA Induced VSR-marked PVCs to Form Aggregates in Root-Tip Cells of Other Plants

We have thus far demonstrated that BFA at high concentrations induced both Golgi and PVC to form aggregates in tobacco BY-2 cells, and that aggregates derived from Golgi and PVC remained distinct but closely associated. To find out if such a BFA response is unique to BY-2 cells or can also be observed in other cell types, we performed additional BFA induction experiments using root tip cells of various plants. Mature seeds of mungbean, pea, Arabidopsis and transgenic tobacco expressing the Golgi marker GONST1-YFP were first germinated before the root tips were treated with BFA at 0, 10 and 100 µg mL\(^{-1}\) for one hour, followed by fixation and immunolabeling with VSR antibodies. As shown in Figure 10, typical punctate patterns were observed in tobacco root tip cells expressing the Golgi marker GONST1-YFP, and these YFP-marked Golgi formed aggregates in the presence of BFA at either 10 or 100 µg mL\(^{-1}\) (Fig. 10, column 1), a result consistent with that observed in transgenic GONST1-YFP BY-2 cells. These results indicated that BFA at high concentration induced the Golgi to form aggregates in tobacco root tip cells. In contrast, BFA at low concentrations (0 or 10 µg mL\(^{-1}\)) did not induce any changes in the VSR-marked PVCs, but BFA at high concentrations (100 µg mL\(^{-1}\)) induced the VSR-marked PVCs to form aggregates in root tip cells of mungbean, pea and Arabidopsis (Fig. 10, columns 2 to 4). Moreover, the PVC-derived aggregates in Arabidopsis root tip cells (Fig. 9B) look similar to those in BY-2 cells (Fig. 9A). Therefore, BFA at high concentrations induced Golgi and PVC to form aggregates in root tip cells of various plants in a way that is similar to that of BY-2 cells.
DISCUSSION

Transgenic BY-2 Cells Expressing PVC Reporter are Useful Tools for Studying the Dynamics of PVCs

Transgenic tobacco BY-2 cells expressing GFP-tagged organelle markers have been useful tools for studying the dynamics of various endomembrane organelles including ER and Golgi apparatus (Satiat-Jeunemaitre et al., 1999). Prevacuolar/endosomal compartments are important organelles that mediate protein trafficking in the secretory and endocytic pathways of eukaryotic cells. Several proteins have been used as markers to define PVCs in plant cells, including syntaxin (Bassham et al., 1995; Conceição et al., 1997), small GTPase (Ueda et al., 2004) and VSR proteins (Li et al., 2002; Tse et al., 2004; Lam et al., 2005). More recently, we have generated transgenic tobacco BY-2 cell lines expressing the YFP-BP-80 reporter for PVCs. Using VSR antibodies and the YFP-BP-80 reporter expressing in transgenic tobacco BY-2 cells as markers, PVCs were identified as mobile MVBs in tobacco BY-2 cells (Tse et al., 2004). In this study, we used two transgenic BY-2 cells lines, expressing either the Golgi marker GONST1-YFP reporter or the PVC marker G/YFP-BP-80 reporter (Tse et al., 2004), to study the effects of BFA on PVCs in living cells by following changes of fluorescent signals upon drug treatment. We demonstrated that PVCs and Golgi apparatus showed different sensitivity towards BFA treatments where BFA at physiologically high concentrations induced PVCs to form aggregates that remained distinct from Golgi-derived BFA-induced aggregates.

PVCs and Golgi Have Different Sensitivity to BFA Treatment in BY-2 Cells

The Golgi apparatus provides a site for rapid action in both mammalian and plant cells where BFA induces Golgi to form ER-Golgi hybrid or a BFA compartment. However, mammalian cells and plant cells have different sensitivity to BFA treatment. For example, BFA at 2.8 μg mL\(^{-1}\) was sufficient and specific to induce the formation of BFA-compartments and block protein traffic from ER to Golgi in animal cells (Klausner et al., 1992), but a higher
concentration of BFA (i.e. at 10 \( \mu \text{g mL}^{-1} \)) was required to induce Golgi changes (Satiat-Jeunemaitre and Hawes, 1992; Driouich et al., 1993) and block ER to Golgi protein traffic (Jiang and Rogers, 1998) in plant cells. However, BFA at 10 \( \mu \text{g mL}^{-1} \) did not cause detectable morphological changes for PVCs in BY-2 cells (Tse et al., 2004). Therefore, the fate of Golgi apparatus in BFA-treated BY-2 cells is affected by BFA in a concentration dependent manner. These results are consistent with previous studies in maize root cells (Satiat-Jeunemaitre et al., 1996) and in tobacco BY-2 cells (Nebenfuhr et al., 2002; Ritzenthaler et al., 2002).

In addition to ER and Golgi apparatus, BFA may also act on endosomes in mammalian cells. For example, BFA treatment induced endosomes to become tubulated in mammalian cells (Lippincott-Schwartz et al., 1991; Hunziker et al., 1991; Wood et al., 1991), which may be due to the involvement of Arf1 in the formation of transport vesicles between early and late endosomes in mammalian cells (Gu and Gruenberg, 2000). Furthermore, BFA treatment caused changes in the membrane compositions of endosomes in mammalian cells and yeast (Donaldson and Jackson, 2000).

In this study, we extended our understating to BFA effects on the prevacuolar organelles using transgenic tobacco BY-2 cells expressing the PVC marker GFP-BP-80 reporter. We demonstrated that PVCs/MVBs in BY-2 cells formed aggregates in response to BFA treatment at high concentrations (i.e. at 50 to 100 \( \mu \text{g mL}^{-1} \)), but not at low concentrations (i.e. at 5 to 10 \( \mu \text{g mL}^{-1} \)). Such high BFA concentrations were within the reversible levels because these tested organelles recovered fully (based on the punctate fluorescent patterns) upon washing off the drug. Furthermore, the target of BFA effect on Golgi and PVC is distinct because the phospholipase A\(_2\) inhibitor ONO-RS-082 specifically prevented PVCs but not Golgi from forming aggregates in the presence of BFA. Furthermore, the delayed observation of PVC-derived aggregates compared to Golgi-derived aggregates indicated that Golgi apparatus might be more sensitive than PVC in response to BFA treatment, which might be due to the
different ARF GEFs (exchange factors for ARF GTPases) present in these two organelles (Geldner et al., 2003). However, it is not clear if BFA-induced PVC response is a consequence of the Golgi response or the effects of BFA on Golgi and PVC is independent to each other.

In addition, the PVC-derived aggregates as identified by VSR antibodies in confocal immunofluorescence remained distinct from the Golgi-derived aggregates in the same BFA-treated cells. ImmunoEM and structural TEM studies further identified the PVC-derived aggregates as clusters of VSR-labeled PVCs with similar size (about 100 to 200 nm in diameter) but lost the appearance of their internal vesicles. Such BFA effect on PVCs in BY-2 cells is different from the effect of wortmannin on PVCs in which wortmannin induced PVCs to form small vacuoles but the enlarged vacuoles still contained visible but reduced internal vesicles (Tse et al., 2004).

In addition, the BFA-induced PVC-derived aggregates were not limited to BY-2 cells, but also observed in several other cell types including root-tip cells of pea, tobacco, rice and Arabidopsis. Therefore, it seems that the BFA effect on PVCs is a general response in plant cells. It is thus interesting and important to find out the possible physiological consequences of PVC changes in response to BFA treatment and the molecular mechanism of BFA action on plant PVCs.

**PVC and Endosome might have Different Sensitivity to BFA in Plant Cells**

In this study, we have shown that Golgi and PVC have different sensitivity to BFA, where BFA at low concentrations (5 to 10 µg mL⁻¹) caused Golgi but not PVC to form aggregates, while high BFA concentrations (50 to 100 µg mL⁻¹) induced both Golgi and PVC to form aggregates. Several recent studies also demonstrated that BFA induced endosome to form aggregates in plant cells. The *Arabidopsis* GNOM protein (an exchange factor for ARF GTPases - ARF-GEFs), which plays important roles in mediating endosomal recycling, auxin transport and plant growth in *Arabidopsis*, was found to locate to endosome because this protein
colocalized with the internalized FM4-64 endosomal marker 30 minutes after uptake study (Geldner et al., 2003). When the *Arabidopsis* root tip cells were treated with BFA at 14 µg mL⁻¹ (50 µM), the GNOM-marked endosomal compartments formed aggregates (Geldner et al., 2003). However, these GNOM-marked endosomal compartments might be different from the PVCs marked by GFP-BP-80 in transgenic BY-2 cells in this study. First, the PVCs marked by GFP-BP-80 reporter did not form aggregates when transgenic BY-2 cells were treated with BFA at 10 µg/ml (Tse et al., 2004), a concentration induced GNOM-marked endosome to form aggregates in *Arabidopsis* cells (Geldner et al., 2003). Second, the VSR-marked PVCs were identified as MVBs in BY-2 cells, such multivesicular nature is consistent with the identity of late endosome (Tse et al., 2004). Third, 30 min after uptake study with the endosomal marker FM4-64, the PVCs marked by the GFP-BP-80 reporter were found colocalize with the internalized dye (Tse et al., 2004), a similar time required for internalization of FM4-64 to the GNOM-marked endosome in *Arabidopsis* root tip cells. However, the VSR-marked PVCs in BY-2 cells and the GNOM-marked endosomes might still represent two distinct endocytic compartments because the uptake rate of FM4-64 in BY-2 cells and *Arabidopsis* root cells might be different. Last, we have recently generated transgenic BY-2 cell lines expressing fusion proteins containing GFP and a rice secretory carrier associate membrane protein (SCAMP) (GFP-SCAMP) that mark both plasma membrane and internalized vesicles, where these GFP-marked vesicles are identified as early endosomes and BFA at low concentrations (5 to 10 µg mL⁻¹) induced them to form aggregates (Lam et al., 2006). In addition, the different sensitivity of Golgi, PVC and (early) endosome to BFA might lie in organelle-specific localization of different ARF GEFs, where endosome-specific GNOM determines the sensitivity of PIN1 recycling in *Arabidopsis* cells (Geldner et al., 2003). Taken together, the VSR-marked PVCs/MVBs might represent late endosomes merging from the endocytic pathway in tobacco BY-2 cells.
BFA-induced Golgi-derived Aggregates or ER Patterns

In this study, Day-3 or Day-2 BY-2 cells after subcultures were used to study the effects of BFA on PVC/MVB in tobacco BY-2 cells mainly for two reasons. First, cells at these two days are at their log phase, thus representing physiologically health stages, while cells at Day-6 or Day-7 represent stationary stage (Matsuoka et al., 2004). Second, in transgenic BY-2 cells expressing the PVC marker XFP-BP-80, typical punctate fluorescent patterns representing PVCs/MVBs (Tse et al. 2004) were observed in Day-2 or Day-3 cells, but such PVC patterns gradually disappear during later stages of cultures with fluorescent signals detected in vacuoles at Day-6 and Day-7 cells (Mitsuhashi et al., 2000; Lo and Jiang, 2006).

When Day-3 or Day-2 GONST1-YFP BY-2 cells were treated with BFA at 10 μg mL⁻¹ for one hour, typical Golgi aggregates were observed (e.g. Fig. 2A). In addition, similar BFA-induced aggregates were also observed in root cells of transgenic tobacco expressing the GONST1-YFP (Fig. 10). Such BFA-induced Golgi aggregations have been documented by many previous studies in various cell types including *Arabidopsis* root cells for the expressed the mammalian sialytransferase (ST) (Wee et al., 1998), tobacco BY-2 cells for GONST1-YFP (Tse et al., 2004; Miao et al., 2006), living onion epidermal cells for GONST1-YFP (Baldwin et al., 2001), *Arabidopsis* cotyledon epidermal cells for KAMΔC:mRFP (Kong et al., 2006), maize root cells for JIM84 (Satiat-Jeunemaitre et al., 1996; Satiat-Jeunemaitre and Hawes, 1992) and *Arabidopsis* root cells for JIM84/ADL6 (Jin et al., 2001). However, several other studies have also demonstrated that BFA induced the reporter-marked Golgi to form ER patterns in various cell types including tobacco epidermal cells for the STtmd-GFP (Boevink et al., 1998), tobacco BY-2 cells for Man1-GFP (Ritzenthaler et al., 2002), and tobacco leaf epidermal cells and BY-2 cells for ST-GFP (Saint-Jore et al., 2002). Therefore, it seems that there is a variety of response of tissues and Golgi markers to BFA. Furthermore, different BFA concentrations used and different lengths of BFA treatment time before imaging may also contribute to these observed variations.
In addition, the physiological status of the cells may also contribute such variation. Indeed, when Day-3 and Day-6 GONST1-YFP BY-2 cells were subjected to BFA treatment at 10 µg mL\(^{-1}\) for one hour, aggregation patterns and ER patterns were observed from these two stages of cells, respectively (Supplemental Figure 2). However, when another BY-2 cell line expressing Man1-GFP was used, ER patterns were observed from both Day-3 and Day-6 cells after BFA treatment (Supplemental Figure 2). However, we do not know if such difference between GONST1-YFP and Man1-GFP BY-2 cell lines is due to the difference between the \textit{trans}-Golgi localization of GONST1-YFP and the \textit{cis}-Golgi localization of Man1-GFP, or due to their different sensitivity to BFA treatment, or, even though unlikely, due to the different localization of GONST1-YFP and Man1-GFP to different populations of Golgi apparatus in BY-2 cells. Since the BFA-induced aggregates for GONST1-YFP have been demonstrated by several studies in living onion epidermal cells (Baldwin et al., 2001), tobacco BY-2 cells (Tse et al., 2004 and this study), and root cells of transgenic tobacco (this study), it would thus be reasonable to use GONST1-YFP BY-2 cells as a control for the XFP-BP-80 BY-2 cells to study the effect of BFA on PVC/MVB in BY-2 cells in this study.

**Concluding Remarks**

Several endomembrane organelles including ER and Golgi formed aggregates in response to BFA treatment, which also affect the transport of proteins from ER to Golgi in the secretory pathway. The development of transgenic tobacco BY-2 cell lines expressing GFP-marked PVCs and the identification of MVBs as PVCs have allowed us to study the dynamic response of PVCs to various drugs in living cells. For example, wortmannin treatment induced the GFP-marked PVCs to form small vacuoles and thus provided a quick tool for defining multivesicular PVCs in tobacco BY-2 cells (Tse et al., 2004). In this study, we demonstrated that BFA at high concentrations (50 or 100 µg mL\(^{-1}\)) induced both YFP-marked PVCs and Golgi to form typical enlarged aggregates but the PVC-derived aggregates remained physically
distinct from the Golgi-derived aggregates. In addition, the BFA-induced formation of aggregates derived from PVCs was specifically inhibited by a phospholipase A2 inhibitor ONO-RS-082, but this drug did not prevent the formation of BFA-induced Golgi-derived aggregates in BY-2 cells. The different changes of Golgi and PVC in response to wortmannin and BFA would provide quick tools to distinguish PVC from Golgi and identify PVC that will serve as a first step to study the molecular mechanism of BFA effect on PVC-mediated protein traffic in the secretory and endocytic pathways of plant cells. Several questions can be addressed in future research. For example, what is the molecular target of BFA on PVC? Does BFA also affect PVC-mediated protein trafficking between Golgi and PVC or between PVC and plasma membrane? What are the molecular mechanisms of PVC-mediated protein trafficking in the plant secretory pathway? What is the physiological significance of BFA-induced PVC-derived aggregates in plant cells? Our current research is addressing some of these questions to further understand the roles of PVCs in plant cells.
MATERIALS AND METHODS

General methods for construction of recombinant plasmids, characterization of cloned inserts, transformation of tobacco (N. tabacum) BY-2 cells, maintenance of transgenic tobacco (N. tabacum) BY-2 cells, and preparation and characterization of antibodies have been described previously (Jiang and Rogers, 1998; Jiang et al., 2000; Cao et al., 2000; Jiang and Rogers, 2001; Tse et al., 2004). BY-2 cells were maintained by either subculture twice a month on agar plates or every seven days in liquid cultures at room temperature (21 to 22 °C). Liquid cultures were kept in shakers at 125 rpm.

Generation of Transgenic BY-2 Cells Expressing the GFP-BP-80 Reporter

GFP with a signal peptide sequence from proaleurain (spGFP) was amplified by PCR and subcloned into the pYFP-BP-80 construct via HindIII / EcoRI sites to replace the pYFP and resulted in pGFP-BP-80 (Tse et al., 2004). The resulting pGFP-BP-80 construct was then transformed into the Agrobacterium LBA4404, followed by transforming BY-2 cells as previously described (Tse et al., 2004). In addition, transgenic tobacco plants expressing the Golgi marker GONST1-YFP and the PVC reporter pYFP-BP-80 were generated via Agrobacterium-mediated transformation as previously described (Jiang et al., 2000).

Brefeldin A Treatment and Recovery Studies

For BFA treatment experiments, aliquots of the Brefeldin A (stock at 2.5 mg mL⁻¹ in DMSO) solution were added to Day-2 and Day-3 old suspension cultures (log phase cultures) to give the proper final concentrations. BFA-treated cells were then removed from the cultures at indicated time for direct confocal imaging, or fixed for confocal immunofluorescence and electron microscopy. For recovery experiments, BY-2 cells were treated with BFA at indicated concentrations, followed by centrifugations at low-speed and washing three times with fresh MS medium before samples were used for confocal imaging or fixation. On average,
more than 100 cells were observed to obtain similar results for each experiment. All drug
treatment experiments had been repeated at least 2 to 3 times with similar results to make sure
that the drugs were functional and the cells used were at the same or similar physiological
stages.

**ONO-RS-082 and Bromoenol Lactone (BEL) Treatment**

For drug treatment using OBO-RS-082 (BIOMOL, USA) and bromoenol lactone (BEL)
(Sigma, USA), aliquots of the ONO-RS-082 and BEL (stock at 1mM in DMSO) solution were
added to 2-to-3 day-old suspension cultures to give the proper final concentrations and
incubate for 15 min, followed by addition of BFA at indicated final concentrations and incubate
for one hour. The treated cells were then collected and subjected to direct confocal imaging.

**Antibodies**

The production and characterization of VSR antibodies were described previously (Tse et al.,
2004). Secondary lissamine rhodamine-conjugated affinity-purified anti-rabbit antibodies
were purchased from Jackson ImmunoResearch Laboratories (West Grove, PA, U.S.A.).

**Confocal Immunofluorescence Studies**

Fixation and preparation of Tobacco BY-2 cells, labeling and analysis by epifluorescence and
confocal immunofluorescence have been described previously (Jiang and Rogers, 1998; Jiang
et al., 2000; Li et al., 2002). The parameters for collecting confocal images within the linear
range were settled as previously described (Jiang and Rogers, 1998). For single labeling, 4 µg
mL\(^{-1}\) of polyclonal rabbit VSR antibodies were incubated at 4°C overnight. All confocal
fluorescence images were collected using a Bio-Rad Radiance 2100 system (Hemel
Hempstead, UK). Images were processed using Adobe Photoshop software (San Jose, CA) as
previously described (Jiang and Rogers, 1998).
Electron Microscopy of Resin-Embedded Cells

For samples embedded in LR White and Lowicryl (HM20) for immunoEM studies, cells were fixed in 1 ml of a primary fixative solution containing 0.25% (v/v) glutaraldehyde and 1.5% (v/w) paraformaldehyde in 50 mM phosphate buffer, pH 7.4, for 15 minutes at room temperature and then transferred to 4°C for additional 16 hours. After washing with phosphate buffer at room temperature, cells were dehydrated in an ethanol series and then embedded in LR White and Lowicryl (HM20) resin.

For samples embedded in Spurr’s resin for structural TEM studies, cells were fixed in 1 ml of a primary fixative containing 2% (v/v) glutaraldehyde and 0.1 ml of saturated picric acid in 25 mM CaCo buffer, pH 7.2, for 15 minutes at room temperature and then transferred to 4°C for additional 16 hours. After washing with 25 mM CaCo buffer, pH 7.2, cells were further subjected to a secondary fixative solution containing 2% (w/v) osmium tetroxide and 0.5 % (w/v) potassium ferrocyanide in 25 mM CaCo buffer, pH 7.2, for 2 hours at room temperature. The cells were then washed with 25 mM CaCo buffer, pH 7.2, followed by contrasting at 2% aqueous uranyl acetate for 2 hours at room temperature. After washing twice in water, cells were dehydrated in an acetone series and finally embedded in Spurr’s resin.

For sample preparation using high-pressure freezing/frozen substitution, the procedures were performed essentially as described previously (Tse et al., 2004). Ultra-thin sections were then prepared from these blocks and used in either immunoEM or structural TEM studies as described (Tse et al., 2004). VSR and GFP antibodies (at 40 µg mL⁻¹ diluted with PBS containing 1% BSA) were used in immunoEM studies.

ACKNOWLEDGMENTS

This work was supported by grants from the Research Grants Council of Hong Kong
(CUHK4156/01M, CUHK4260/02M, CUHK4307/03M and CUHK4580/05M), NSF of China (30529001) and CUHK Scheme C to L. Jiang. We are grateful to Prof. David G. Robinson (University of Heidelberg, Germany) and John C. Rogers (National Science Foundation, USA) for their continuous support in our studies. A portion of this work has been presented in abstract form for a poster for the American Society of Plant Biologists Annual Meeting 2005 (http://abstracts.aspb.org/pb2005/public/P51/7276.html).
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Figure Legends

Figure 1. Subcellular Localization of Two Reporters in Transgenic BY-2 Cells.

(A). Confocal immunofluorescent study of reporters. Similar punctate signals were detected from transgenic BY-2 cells expressing either the Golgi marker GONST1-YFP (panel 1) or the PVC marker GFP-BP-80 (panel 2), where both YFP-BP-80 and GFP-BP-80 reporter (green) largely colocalized with VSR antibodies (red). DIC = Differential Interference Contrast images showing the morphology of the tested cells. n = nucleus. Bars = 50 µm.

(B). Immunogold EM localization of GONST1-YFP and GFP-BP-80. Ultra-thin sections prepared from high-pressure frozen/freeze-substituted transgenic GONST1-YFP BY-2 cells were labeled with either GFP antibodies (left panel) to detect the Golgi-localized GONST1-YFP reporter or VSR antibodies (right panel) to detect the multivesicular PVCs. Bars = 200 nm.

Figure 2. Brefeldin A Induces YFP-marked PVCs to Form Aggregates in a Dose-dependent Manner.

(A) and (B). Transgenic BY-2 cells expressing either the Golgi marker GONST1-YFP (panel A) or the PVC marker YFP-BP-80 (panel B) were incubated with BFA at various concentrations as indicated for one hr before they were collected for confocal imaging of YFP signals. Arrows indicate examples of BFA-induced aggregates in cells expressing these two reporters. DIC = Differential Interference Contrast images showing the morphology of the tested cells. n = nucleus. Bars = 50 µm.

(C). The average sizes of the fluorescence signals of GONST1-YFP and YFP-BP-80 in transgenic BY-2 cells treated with BFA at 50 µg mL⁻¹ at indicated times (calculated using image J software on images of 500 x 500 pixels with 200 dpi).

(D). The average numbers of YFP-BP-80-marked PVCs in transgenic BY-2 cells treated with
BFA at indicated concentrations (0, 5 and 10 µg/ml) for one hour.

**Figure 3. Time Course Formation of BFA-induced Golgi or PVC Aggregates.**

(A) and (B). Transgenic BY-2 cells expressing the Golgi marker GONST1-YFP (panel A) and the PVC marker YFP-BP-80 (panel B) were first incubated with BFA at 50 µg mL⁻¹, followed by sample collection at indicated times for confocal imaging of YFP signals. Arrows indicate examples of BFA-induced aggregates derived from either the YFP-marked Golgi (A) or the YFP-marked PVCs (B). DIC = Differential Interference Contrast images showing the morphology of the tested cells. n = nucleus. Bars = 50 µm.

(C). The average sizes of the fluorescence signals of GONST1-YFP and YFP-BP-80 in transgenic BY-2 cells treated with BFA at 50 µg mL⁻¹ at indicated times (calculated using imageJ software on images of 500 x 500 pixels with 200 dpi).

(D). The average numbers of YFP-BP-80-marked PVCs in a single transgenic BY-2 cell at indicated times upon BFA treatment at 50 µg/ml.

**Figure 4. Recovery of BFA-induced Aggregates.**

Transgenic BY-2 cells expressing the Golgi marker GONST1-YFP (panel A) and the PVC marker GFP-BP-80 (panel B) were treated with BFA at 100 µg mL⁻¹ for 1 hr before BFA was washed off with fresh MS medium, followed by sample collection at indicated times for confocal imaging. DIC = Differential Interference Contrast images showing the morphology of the tested cells. n = nucleus. Bars = 50 µm.

**Figure 5. BFA-induced PVC Aggregates are Distinct from Golgi Aggregates.** Transgenic BY-2 cells expressing the Golgi marker GONST1-YFP and the PVC marker YFP-BP-80 were treated with BFA at 50 µg mL⁻¹ for 1 hr (panels 1 and 2) or 2 hr (panel 3) prior to fixation. The fixed cells were then labeled with VSR antibodies to detect PVCs (red), while the YFP
reporters (green) were ready for detection. Arrowheads in the merged image of panel 1 indicated examples of colocalization between the PVC reporter YFP-BP-80 and VSR antibodies. Arrowheads and arrows in panels 2 and 3 indicated examples of distinct but closely associated BFA-induced aggregates derived from Golgi (green) and PVC (red) (termed Golgi-PVC hybrid). DIC = Differential Interference Contrast images showing the morphology of the tested cells. Bars = 50 µm.

**Figure 6. The Formation of Golgi-PVC Hybrids was Time-dependent.**
Transgenic BY-2 cells expressing the Golgi reporter GONST1-YFP were treated with BFA at 50 µg mL⁻¹, followed by sample collection at indicated time points and used for fixation. The fixed cells were then labeled with VSR antibodies to detect PVCs (red), while the YFP-marked Golgi (green) was ready for detection. DIC = Differential Interference Contrast images showing the morphology of the tested cells. Bar = 50 µm.

**Figure 7. The target of BFA effect on Golgi and PVC is Distinct**
Transgenic BY-2 cells expressing the Golgi reporter GONST1-YFP (panel A) and PVC reporter GFP-BP-80 (panel B) were treated with ONO-RS-082 at 10 µM for 15 min, followed by BFA treatment at 10 or 100 µg mL⁻¹ for 1 hr as indicated (panels 3 and 4) before confocal imaging. Controls included BFA treatment at 10 or 100 µg mL⁻¹ for 1 hr before confocal imaging (panels 1 and 2). DIC, Differential Interference Contrast images showing the morphology of the tested cells. Bars = 50 µm.

**Figure 8. Immunogold EM Identification of BFA-induced Aggregates Derived from Golgi and PVC.**
(A). GFP antibodies labeled the Golgi stacks in untreated transgenic GONST1-YFP cells using ultra thin sections of Lowicryl HM20.
(B). GFP antibodies labeled the curved Golgi structures in transgenic GONST1-YFP cells treated with BFA at 100 µg mL\(^{-1}\), using ultra-thin sections of Lowicryl HM20.

(C) & (D). Multivesicular PVCs as detected by VSR antibodies in untreated cells using ultra-thin sections from high pressure freezing and freeze substituted material.

(E) to (H). PVC or PVC aggregates as detected by VSR antibodies in cells treated with BFA at 100 µg mL\(^{-1}\), using ultra-thin samples of high pressure freezing and frozen substitution. Bars = 200 nm.

(I). Overview of BFA-induced PVC aggregates in BY-2 cells treated with BFA at 100 µg mL\(^{-1}\) for 1 hr, followed by high pressure freezing and frozen substitution. Bars = 500 nm.

**Figure 9. Ultrastructural Analysis of BFA-treated Transgenic GONST1-YFP Cells and Arabidopsis root tip cells.**

(A). Overview of BFA-induced Golgi and PVCs aggregates in BY-2 cells treated with BFA at 100 µg mL\(^{-1}\) for 1 hr.

(B). Overview of BFA-induced Golgi and PVCs aggregates in Arabidopsis root tips treated with at BFA 100 µg mL\(^{-1}\) for 1 hr. Bars = 200 nm.

**Figure 10. BFA also Induced VSR-marked PVCs to Form Aggregates in Other Plant Cell Types.**

Root tip cells from germinating seeds of transgenic GONST1-YFP tobacco, mungbean, pea and Arabidopsis were treated with BFA at 0, 10 and 100 µg mL\(^{-1}\) for 1 hr as indicated before they were either used for direct confocal imaging (for tobacco cells) to ready detect Golgi apparatus, or fixed and labeled with VSR antibodies (for pea, mung bean and Arabidopsis root tip cells) to detect PVCs. Bar = 50 µm.
Supplemental Figure 1. Brefeldin A Induces YFP-marked PVCs to Form Aggregates in a Dose-dependent Manner.

Transgenic BY-2 cells expressing either the Golgi marker GONST1-YFP (panel A) or the PVC marker YFP-BP-80 (panel B) were incubated with BFA at various concentrations as indicated for one hr before they were collected for confocal imaging of YFP signals. Arrows indicate examples of BFA-induced aggregates in cells expressing these two reporters. DIC = Differential Interference Contrast images showing the morphology of the tested cells. n = nucleus. Bars = 50 µm.

Supplemental Figure 2. The formation of BFA-Induced Golgi Aggregates in GONST1-YFP Cells is Stages Dependent.

Day-3 and Day-6 transgenic GONST1-YFP (left panel) or Man1-GFP (right panel) BY-2 cells were treated with BFA at 10 µg mL⁻¹ for 1 hr before image collection. DIC = Differential Interference Contrast images showing the morphology of the tested cells. Bars = 50 µm.

Supplemental Figure 3. Recovery of BFA-induced Aggregates.

Transgenic BY-2 cells expressing the Golgi marker GONST1-YFP (panel A) and the PVC marker GFP-BP-80 (panel B) were treated with BFA at 100 µg mL⁻¹ for 1 hr before BFA was washed off with fresh MS medium, followed by sample collection at indicated times for confocal imaging. DIC = Differential Interference Contrast images showing the morphology of the tested cells. n = nucleus. Bars = 50 µm.

Supplemental Figure 4. The target of BFA effect on Golgi and PVC is Distinct

Transgenic BY-2 cells expressing the Golgi reporter GONST1-YFP (panel A) and PVC reporter GFP-BP-80 (panel B) were treated with BEL at 10 µM for 15 min, followed by BFA treatment at 10 or 100 µg mL⁻¹ for 1 hr as indicated (panels 3 and 4) before confocal imaging.
Controls included BFA treatment at 10 or 100 µg mL⁻¹ for 1 hr before confocal imaging (panels 1 and 2). DIC, Differential Interference Contrast images showing the morphology of the tested cells. Bars = 50 µm.

Supplemental Figure 5. Ultrastructural Analysis of BFA-treated Transgenic GONST1-YFP Cells.

(A) Golgi stacks from an untreated cell.

(B) Multivesicular PVC in an untreated cell.

(C) Golgi aggregates and ER-Golgi hybrids from cells treated with BFA at 10 µg mL⁻¹.

(D) Multivesicular PVC from cells treated with BFA at 10 µg mL⁻¹.
