Imaging of Dynamic Secretory Vesicles in Living Pollen Tubes of *Picea meyeri* Using Evanescent Wave Microscopy¹[W]

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Evanescent wave excitation was used to visualize individual, FM4-64-labeled secretory vesicles in an optical slice proximal to the plasma membrane of *Picea meyeri* pollen tubes. A standard upright microscope was modified to accommodate the optics used to direct a laser beam at a variable angle. Under evanescent wave microscopy or total internal reflection fluorescence microscopy, fluorophores localized near the surface were excited with evanescent waves, which decay exponentially with distance from the interface. Evanescent waves with penetration depths of 60 to 400 nm were generated by varying the angle of incidence of the laser beam. Kinetic analysis of vesicle trafficking was made through an approximately 300-nm optical section beneath the plasma membrane using time-lapse evanescent wave imaging of individual fluorescently labeled vesicles. Two-dimensional trajectories of individual vesicles were obtained from the resulting time-resolved image stacks and were used to characterize the vesicles in terms of their average fluorescence and mobility, expressed here as the two-dimensional diffusion coefficient $D^2$. The velocity and direction of vesicle motions, frame-to-frame displacement, and vesicle trajectories were also calculated. Analysis of individual vesicles revealed for the first time, to our knowledge, that two types of motion are present, and that vesicles in living pollen tubes exhibit complicated behaviors and oscillations that differ from the simple Brownian motion reported in previous investigations. Furthermore, disruption of the actin cytoskeleton had a much more pronounced effect on vesicle mobility than did disruption of the microtubules, suggesting that actin cytoskeleton plays a primary role in vesicle mobility.

Vesicle trafficking is fundamental to numerous activities in eukaryotic organisms and underlies many of the basic processes involved in cell growth and differentiation. The vesicle trafficking network is responsible for exocytosis and endocytosis (Rothman, 1994; Bretscher and Aguado-Velasco, 1998; Hepler et al., 2001; Zefirov et al., 2004; Šamaj et al., 2005). Exocytosis is the process by which secretory vesicles derived from the trans-Golgi network (TGN) attach and/or fuse with the plasma membrane, releasing their contents into the extracellular space. In contrast, endocytosis is responsible for the uptake of compounds from the extracellular milieu to be used in cellular metabolism, the uptake of signal molecules, and the uptake of membrane proteins and lipids to be recycled (Šamaj et al., 2005). Vesicle transport of cargo is carried out by coated vesicles from a donor compartment. These vesicles move on to the Golgi apparatus, where further posttranslational modifications are made. Uncoating and subsequent docking and fusion of the vesicles with a target compartment membrane then occur. Three major types of vesicle coating proteins, COPI, COPII, and clathrin, are involved in exocytotic and endocytotic vesicle trafficking. Plant cells grow when secretory vesicles derived from the TGN and endosomes fuse with the plasma membrane, into which they insert cellulose synthases. At the same time, they insert cellulose synthases. At the same time, they insert cellulose synthases. At the same time, they insert cellulose synthases. At the same time, they insert cellulose synthases. At the same time, they insert cellulose synthases. At the same time, they insert cellulose synthases. At the same time, they insert cellulose synthases. At the same time, they insert cellulose synthases.
animal cells (Rutter and Tsuboi, 2004; Sudhof, 2004; Vo et al., 2004; Zanner et al., 2004; Wang and Quick, 2005; Xu et al., 2005). Relatively few studies of vesicle trafficking in plant cell systems have been performed, largely due to the difficulties presented by the plant cell walls. However, in a study on vesicle transport during pollen tube growth, Derksen et al. (1995) used transmission electron microscopy (TEM) to quantitatively analyze the distribution and density of vesicles in cryo-fixed tobacco (Nicotiana tabacum) pollen tubes. Using video microscopy and TEM, Ketelaar (2002) subsequently estimated the number of exo- and endocytic vesicles in Arabidopsis (Arabidopsis thaliana) root hairs and proposed that the exocytosis and endocytosis rates were 4,202 and 6,901 vesicles per minute, respectively. Although indirect insight can be gained from electron microscopy or optical recording with confocal microscopy (Picton and Steer, 1983; Rutter and Knuiman, 1993; Parton et al., 2001, 2003), neither method can easily provide the high temporal resolution and information needed to determine the frequency or rate of fast exocytotic and endocytotic events occurring during pollen tube growth. By means of differential interference contrast microscopy, Foissner et al. (1996) recorded vesicle trafficking in the characean wound wall system. de Win et al. (1998) applied interactive computer-assisted video microscopy to analyze the movement patterns of the organelles in pollen tubes and found the movements were random in the tip regions. Sieberer and Emons (2000) measured the speed and orientation of organelar motion in root hairs of Medicago truncatula with diameters of 1 to 2 μm using video microscopy. Indeed, to our knowledge, no study to date has portrayed the dynamic motion of individual secretory vesicles in living pollen tubes (for studies of tip-growing root hairs, see Ovecka et al., 2005).

With the introduction of evanescent wave microscopy (EWM), direct monitoring of an individual vesicle during secretion has become possible (Tsuboi and Rutter, 2003; Kuhn and Pollard, 2005). Evanescent wave excitation of fluorescence is based on the decaying near-field generated near a dielectric interface upon the total internal reflection of light. Due to the light confinement of evanescent wave excitation, photobleaching and phototoxic reactions are generally of minor concern, as compared to conventional epifluorescence and confocal laser scanning microscopy. Therefore, EWM provides an ideal tool for long-term observations or when time-resolved imaging at high frame rates is desired.

In this study, we visualized secretory vesicle motions in living pollen tubes after labeling the vesicles with the endocytic/exocytic tracer FM4-64. This amphiphilic styryl dye has been used to investigate endocytosis and visualize the diverse membrane compartments in living fungal hyphae (Fischer-Parton et al., 2000). FM4-64 is a suitable probe for imaging the dynamic changes in size, morphology, and position of apical vesicles within growing pollen tubes because of its staining pattern, relatively high photostability, and low cytotoxicity (Bolte et al., 2004). Thus, it is an excellent dye for following dynamic processes in living cells over time without disturbing these processes (Šamaj et al., 2005; Dettmer et al., 2006; Šamaj, 2006). In this study, we used EWM for two-dimensional (2-D) secretory vesicle tracking to determine the role of the actin cytoskeleton and microtubules in vesicle trafficking and dynamics proximal to the plasma membrane.

RESULTS

Visualization of Secretory Vesicles Close to the Plasma Membrane in Living Pollen Tubes

Pollen tubes labeled with FM4-64 exhibited a distinct peripheral, bright staining pattern (Fig. 1A). Confocal microscopic analysis of labeled pollen tubes demonstrated that the peripheral staining was plasma membrane associated, not cell wall associated, as indicated by plasmolysis with 100 μM sorbitol in the presence of FM4-64 (Fig. 1C). Pretreatment of pollen tubes with 500 μM sodium azide impaired dye uptake; therefore, FM4-64 fluorescence could be observed only at the plasma membrane (Fig. 1E), but still allowed staining of the plasma membrane of the apical and subapical regions. Because FM4-64 is a membrane-selective fluorescent dye and does not label the cytoplasm or cell wall (Fig. 1C), it was used to detect vesicle trafficking in living pollen tubes.

Under EWM, apical region versus subapical region defined by de Win et al. (1996) and Wang et al. (2005) can be mostly distinguished because the apical region in Picea meyeri tubes with FM4-64 staining appeared crescent, which was different from inverted cone-shaped (V-shaped) clear zone in angiosperm pollen tubes. Fluorescent spots were distributed plentifully over the apical and subapical regions beneath the plasma membrane of the pollen tube proximal to the coverslip. The spot morphologies of the apical and subapical spots were similar. The density of spots, however, was much higher in the apical region than in the subapical region (Fig. 1G). The spots appeared dim because of the scattering effect induced by out-of-focus vesicles or organelles, such as the Golgi apparatus and endoplasmic reticulum deeper within the pollen tube cytoplasm (Fig. 1H). After processing with flattening and high-pass filters, the spots became clear and could be seen to have similar sizes (Fig. 1I). Spots were identified as vesicles when the average intensity in a 3×3-pixel region was 20% greater than the surrounding background gray value in three consecutive frames, the central intensity was a local maximum, and the spot was present in more than three consecutive images. Because the diameter of vesicles in P. meyeri ranged from 100 to 300 nm (Wang et al., 2005), twice the size as those in Lilium and Arabidopsis under the TEM, only those spots...
with diameter less than 400 nm were considered as secretory vesicles (TGN vesicles) for analysis; other larger fluorescent spots are considered not to be vesicles or organelles, which were excluded from analysis in this article.

Figure 1. Secretory vesicles in living pollen tubes of *P. meyeri*. A, A median focal plane optical section showing a typical FM4-64 staining pattern in a growing pollen tube. B, Corresponding bright-field image to A. C, Applying standard medium with 100 μM sorbitol and 2 μM FM4-64 showed dye fluorescence associated with retracted protoplast but not with cell wall. D, Corresponding bright-field image to C. E, Sodium azide impaired FM4-64 uptake. Note that dye remains at the plasma membrane. F, Corresponding bright-field image to E. G, Density of the vesicles in apical and subapical regions in pollen tubes (*n* = 8). H, First image in a stack of 500 taken at 200-ms intervals observed under evanescent wave microscope. I, Flattened and low-pass filtered image of H. For filtering, blurred versions of images on the left were first generated by flatten filtering and then enhanced with low-pass filter three times, at last yielding the distinct images by executing contrast enhancement. Bar = 25 μm for A to F, 5 μm for H and I.

Dynamics of FM4-64-Labeled Secretory Vesicles in Living Pollen Tubes

To explore vesicle motions, a series of images of growing pollen tubes labeled with FM4-64 was taken under EWM. Movies compiled from a large number
of images showed that the vesicles moved around a resting position in the apical and subapical regions of the pollen tubes (Supplemental Movies 1 and 2). These bright fluorescent spots showed short, nonlinear motions in various directions in living pollen tubes. Furthermore, two types of secretory vesicle mobility were observed along the pollen tubes in terms of running length and velocity: short-distance motion (Fig. 2A) and long-distance motion (Fig. 2B). Long-distance motions were defined as motions of >1 μm in distance and with a maximum velocity of >2 μm/s. All other motions were considered short-distance motions. Approximately 10% of vesicles within the evanescent field could be classified as undergoing long-distance motion, whereas the majority (>80%) of vesicles proximal to the plasma membrane underwent short-distance motion.

Short-distance motions often involved rapid changes or reversals in direction and velocity between consecutive runs, whereas the long-distance motions were directed to the apical region, as though these vesicles were guided to their targets. Moreover, the two types of motion differed in their velocities. The average velocity during short-distance motions was 1.09 ± 0.02 μm/s (n = 30 vesicles), with a maximum velocity of 3.5 μm/s. The run length averaged 75 ± 0.8 μm (n = 30 vesicles), with a duration of 100 s. In contrast, the average velocity during long-distance motions was 1.93 ± 0.05 μm/s (n = 30 vesicles), and the maximum velocity was 5.85 μm/s (Table I). In addition, vesicle motion could be estimated by the mean squared displacement (MSD) and the diffusion coefficient. The diffusion coefficient in two dimensions, $D^2$, was derived from the slope of a plot of the MSD versus time for $\Delta t \to 0$ (Fig. 2, C and D). For

Figure 2. Short-distance and long-distance motions of secretory vesicles in P. meyeri pollen tubes. A, The lateral mobility of a short-distance motion vesicle on a plot of x versus y coordinates. B, The lateral mobility of a long-distance motion vesicle on the plots of x versus y coordinates. C, MSD of short-distance motion plotted against time interval $\Delta t$. To calculate the MSD, we measured the distance of motion over an interval of fixed duration (0.2 s), with the interval starting first at 0 ms after the beginning of the recording, then at 200 ms, at 400 ms, and so on until at 9.6 s. All distances were squared. D, MSD of long-distance motion plotted against time interval $\Delta t$. The characteristic decay time of the MSD is given by a polynomial fit (black line).
Table 1. Two types of secretory vesicle mobility based on measurements of 30 vesicles observed under EWM in living pollen tubes

<table>
<thead>
<tr>
<th>Parameters of Vesicle Motions (Mean ± SD)</th>
<th>Long-Distance Motions</th>
<th>Short-Distance Motions</th>
</tr>
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<tbody>
<tr>
<td>Mean velocity (μm/s)</td>
<td>1.93 ± 0.05</td>
<td>1.09 ± 0.02</td>
</tr>
<tr>
<td>Maximum velocity (μm/s)</td>
<td>5.85</td>
<td>3.5</td>
</tr>
<tr>
<td>Minimal velocity (μm/s)</td>
<td>0.038</td>
<td>0.0081</td>
</tr>
<tr>
<td>Path length (μm/10 s)</td>
<td>11.5 ± 0.8</td>
<td>7.79 ± 1.2</td>
</tr>
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</table>

long-distance motions, $D^2$ ranged from $2.1 \times 10^{-2} \mu m^2/s$ to $0.62 \mu m^2/s$, with an average value of $0.11 \pm 0.07 \mu m^2/s$ ($n = 30$ vesicles). For short-distance motions, $D^2$ ranged from $8 \times 10^{-3} \mu m^2/s$ to $0.12 \mu m^2/s$, with an average value of $5.3 \times 10^{-2} \mu m^2/s$ ($n = 30$ vesicles).

Quantifying Individual Secretory Vesicle Motions in Living Pollen Tubes

Much variation in velocity was observed during the recording time. In the pollen tubes observed for long periods, single secretory vesicles occasionally traveled distances of 100 μm at a top speed of 6.87 μm/s (in one instance), but such cases were rare (less than one vesicle per cell per minute) and were therefore excluded from further analysis. During pauses between directed runs, complex, oscillatory behaviors were observed. As shown in Figure 3A, oscillations occurred preferentially during pauses and before direction reversals. The average oscillation frequency was $\omega_{osc} = 0.34 \pm 0.08$ per second ($n = 12$ vesicles).

Exocytosis, the release material from secretory vesicles, was identifiable as a sudden spread of fluorescence. Exocytosis of a vesicle was preceded by a transient brightening of the vesicle followed by a decay of its fluorescence intensity (Fig. 3B), with a characteristic time of $13.7 \pm 1.3 s$ ($n = 20$ pollen tubes). Fusion of vesicles appeared as a fluorescent spot spreading away from the site of fusion (Fig. 3C). The trace showed apparently random motion, superimposed with a slow drift. The trajectories were constrained in an irregular-shaped value of 1 μm in the $x$ to $y$ direction (Fig. 3D). For analysis, we measured the square of the distance traveled by the vesicles and plotted the MSD against the time interval (Fig. 3E). Table II summarizes the tracking parameters and other findings, as well as the ratios of vesicles with different motions in relation to all vesicles.

Effect of Actin Cytoskeleton Disruption on Secretory Vesicle Mobility

Disruption of the actin cytoskeleton by treatment with latrunculin B (LATB) or cytochalasin D (CD) caused significant alterations in secretory vesicle motions (Table III). The average vesicle path lengths in pollen tubes treated with LATB or CD was reduced to $7.34 \pm 1.51$ or $9.16 \pm 1.37 \mu m$ of displacement per 10 s, respectively. These values represent significantly shorter path lengths than those measured in control cells (Fig. 4A). LATB treatment dramatically reduced the vesicle velocity as well; the velocity ranged from 0.20 to 0.96 μm/s, with an average velocity of 0.62 ± 0.21 μm/s, which is about one-fourth that observed in the control pollen tubes. CD showed similar but less significant effect on the velocity, reducing it to 0.65 to 1.04 μm/s, with an average value of 0.81 ± 0.09 μm/s (Fig. 4B). The alterations in vesicle trajectories caused by LATB and CD treatment were characterized by plotting the $x$-$y$ coordinates of vesicles in the two experimental conditions. The results showed that long-distance motion occurred only in the untreated cells that had intact actin cytoskeletons (Supplemental Fig. 1A). Vesicles were much less mobile in the presence of both inhibitors (Supplemental Fig. 1, B and C), and their motions became random and were mostly confined to the restricted regions (Supplemental Fig. 1, G and H). Complex dynamics, such as the oscillations, regularly visualized in the control pollen tubes, were not observed in the LATB- and CD-treated pollen tubes (Fig. 4, C and D). Furthermore, the MSD curves (Fig. 4E) revealed that individual vesicle kinetics was clearly affected by these inhibitors, which reduced $D^2$ by 70% (Fig. 4F).

Effects of Microtubule Disruption on Secretory Vesicle Mobility

Microtubule disruption by treatment with the inhibitors oryzalin and colchicine led to more moderate alterations in secretory vesicle motion than did by actin cytoskeleton disruption. Incubation for 10 min with 100 μm oryzalin and 5 μM colchicine induced a 10% to 20% reduction in vesicle mobility (Supplemental Fig. 1, D and E). Vesicle tracking showed a clear reduction in vesicle mobility as detected in the $x$-$y$ plane; the average path lengths were 9.91 and 10.6 μm in 10 s for oryzalin and colchicines, respectively, which represent approximately 80% of the mobility found near the plasma membrane in control cells (Fig. 4A). When traces of individual vesicle motions in control pollen tubes were compared to those in oryzalin- and colchicine-treated cells, a marked difference in velocity was noted (Fig. 4B). In addition, both oryzalin and colchicine treatments had a significant impact on vesicle trajectories, which were much more confined in the lateral directions than those of the control cells (Supplemental Fig. 1, I and J). Plots of MSD versus time revealed that vesicle motions in treated pollen tubes had a number of different, distinctive features, which deviated from the regulated motions observed in control pollen tubes. The averaged MSD plots are shown in Figure 4E. When pollen tubes were exposed to colchicine and oryzalin for 10 min, $D^2$ of mobile vesicles was reduced by only 25% to 40% (Fig. 4F).
Tracking parameters of secretory vesicles in treated pollen tubes are summarized in Table IV.

Effects of Brefeldin A on Secretory Vesicle Trafficking

We also characterized the effect of brefeldin A (BFA) on the mobility of secretory vesicle in living pollen tubes. When the pollen tubes were observed after a 10-min incubation with BFA, the average path lengths in apical and subapical regions were found to be slightly shorter than the average path lengths measured in control cells. The average path lengths for the apical and subapical regions in BFA-treated pollen tubes were $11.76 \pm 2.35 \mu m$ and $9.42 \pm 1.17 \mu m$.

Figure 3. Trajectory of an individual secretory vesicle near the plasma membrane in living pollen tubes. A, Vesicle oscillation. Plot of the x-y velocity of an oscillating vesicle as a function of time (oscillation frequency $\omega_{osc} = 0.6$ per second) is shown. B, Time course of the fluorescence intensity of the vesicle shown in A. The characteristic decay time of the fluorescence is given by a polynomial fit (black line): $t = 13.7 \pm 1.3$ s. C, Individual exocytotic events of a FM4-64-labeled vesicle. Numbers indicate time (in seconds) relative to the moment of fusion. Lower sections show three-dimensional lumianance plots of four successive frames starting one frame before fusion. D, Lateral mobility of a vesicle on a plot of x versus y coordinates. E, MSD of an exocytotic vesicle plotted against time interval $\Delta t$. 
DISCUSSION

Imaging techniques are powerful tools for detecting vesicle trafficking in living cells, and they can provide information concerning the mechanisms of vesicle trafficking (Betz and Angleson, 1998; Fletcher et al., 2000; Baumann et al., 2005). Because confocal microscopy provides excellent optical sectioning, it has been widely used in the investigation of endocytosis and exocytosis in living cells (Ohara-Imaizumi et al., 2002; Bolte et al., 2004; Ma et al., 2004; Šamaj, 2006). However, vesicle diameters in pollen tubes range from 75 to 200 nm (Derkson et al., 1995), and, thus, vesicles are beyond the resolution limits of a conventional confocal microscope. In this study, we used a high numerical aperture objective and varied the angle of incidence of the laser beam to generate a penetration depth of the evanescent field of 70 to 400 nm proximal to the plasma membrane with cells adhered to a glass coverslip. EWM has provided valuable and complementary information about vesicle trafficking, especially in living cells, under lengthy observation periods (Watkins and Robertson, 1977; Oheim et al., 1999). In terms of depth discrimination, an evanescent wave microscope is up to 10-fold better than a confocal microscope (Steyer and Almers, 2001). Combined with a high-resolution charge-coupled device, which has extremely high temporal sensitivity, EWM is very useful in tracking 2-D motions of small particles near the coverslip under physiological conditions (Steyer et al., 1997; Sund et al., 1999; Sund and Axelrod, 2000; Wiegand et al., 2002; Mashanov et al., 2003; Ohara-Imaizumi et al., 2004; Xia et al., 2004; Zenisek et al., 2004). In this report, we examined the dynamics and motions of FM4-64-labeled vesicles proximal to the plasma membrane in living pollen tubes of *P. meyeri*. To our knowledge, this report is the first description of visualization of secretory vesicle tracking in living pollen tubes using EWM. This technique greatly facilitates the quantification of vesicle trafficking in plant cells.

Pollen tubes have been used as in vivo model systems for studying vesicles and organelle trafficking because they are highly polarized, tip growing, and easy to manipulate experimentally. Angiosperm pollen tubes have been chosen as the subjects of most previous investigations because they grow quickly. For example, the growth rate of *Lilium longiflorum* was reported to be about 5 to 25 μm/min (Parton et al., 2001). In our study, we used the slow-growing pollen tubes of *P. meyeri*, which have a growth rate of 20 μm/h (Anderhag et al., 2000), so that we could easily follow the secretory vesicle motion. Our current understanding of vesicle trafficking in pollen tubes is based primarily on ultrastructural studies and mathematical modeling of vesicle trafficking in relation to tip growth (Derkson et al., 1995; Ketelaar, 2002). Lancelle and Hepler (1992) defined an apical clear zone as a vesicular

### Table II. Overview of individual secretory vesicle tracking parameters on the basis of 200 vesicles observed under EWM in five living pollen tubes

<table>
<thead>
<tr>
<th>Vesicle Tracking Parameters (Mean ± sd)</th>
<th>Long-Distance Motions</th>
<th>Short-Distance Motions</th>
<th>Exocytotic Vesicles</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proportion (%)</td>
<td>Approximately 8%</td>
<td>Approximately 90%</td>
<td>&lt;1%</td>
</tr>
<tr>
<td>Area density (1/µm²)</td>
<td>0.011 ± 0.008</td>
<td>0.038 ± 0.007</td>
<td>N.D.</td>
</tr>
<tr>
<td>Total path length (µm/10 s)</td>
<td>16.5 ± 0.9</td>
<td>7.8 ± 1.2</td>
<td>10.4 ± 0.8</td>
</tr>
<tr>
<td>Mean velocity (µm/s)</td>
<td>1.5 ± 0.4</td>
<td>0.8 ± 0.3</td>
<td>0.9 ± 0.4</td>
</tr>
<tr>
<td>D² (µm²/s)</td>
<td>0.11 ± 0.07</td>
<td>0.053 ± 0.04</td>
<td>0.065 ± 0.03</td>
</tr>
</tbody>
</table>

### Table III. Disruption of the actin cytoskeleton reduced the secretory vesicle mobility (n = 30 pollen tubes)

<table>
<thead>
<tr>
<th>Parameter of Vesicle Motions (Mean ± sd)</th>
<th>10 µm LATB</th>
<th>10 µm CD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean velocity (µm/s)</td>
<td>0.62 ± 0.21</td>
<td>0.81 ± 0.09</td>
</tr>
<tr>
<td>Maximum velocity (µm/s)</td>
<td>0.65</td>
<td>1.04</td>
</tr>
<tr>
<td>Minimal velocity (µm/s)</td>
<td>0.20</td>
<td>0.96</td>
</tr>
<tr>
<td>Path length (µm/10 s)</td>
<td>7.34 ± 1.51</td>
<td>9.16 ± 1.37</td>
</tr>
<tr>
<td>D² (µm²/s)</td>
<td>0.042 ± 0.021</td>
<td>0.049 ± 0.013</td>
</tr>
</tbody>
</table>
Figure 4. Effects of cytoskeleton disruption on mobility of secretory vesicles. A, The average path lengths of vesicles of the nontreated (NT), LATB-treated (LATB), CD-treated (CD), oryzalin-treated (Ory), and colchicine-treated (Col) pollen tubes (n = 12). B, Bar graph depicting the mean and maximum velocities of vesicles in NT, LATB-, CD-, Ory-, and Col-treated pollen tubes (n = 12). C, Plot of the x-y velocity of an individual vesicle in a LATB-treated pollen tube. D, Plot of the x-y velocity of an individual vesicle in a CD-treated pollen tube. E, MSD of vesicle motions in LATB-, CD-, Ory-, and Col-treated pollen tubes plotted against time interval Δt. To calculate the MSD, we measured the distance moved over an interval of fixed duration (0.2 s), with the interval starting first at 0 ms after the beginning of the recording, then at 200 ms, at 400 ms, and so on until at 13 s. F, Calculated 2-D diffusion coefficient of vesicle motions, $D'$, of the NT, LATB-, CD-, Ory-, and Col-treated pollen tubes.
zone depleted of organelles that is located in the apical region of *L. longiflorum* pollen tubes using TEM, and Parton et al. (2001) used FM4-64 and confocal microcopy to reveal that the apical clear zone is a vesicle-rich, V-shaped region at the apex of *L. longiflorum* pollen tubes. In our study, however, we found that in gymnosperm the staining pattern at the tube apex did not correspond spatially to the previously described V-shaped, apical clear zone containing secretory vesicles, confirming that the cytoplasmic organization of gymnosperm pollen tubes was distinct from that of angiosperm pollen tubes (Justus et al., 2004; Wang et al., 2005; Chen et al., 2006).

In previous studies, some organelles moved linearly, in a mostly circulatory pattern (Cai et al., 2001), whereas the majority of organelles in *Pinus sylvestris* pollen tubes exhibited Brownian-like motions (Terasaka and Niitsu, 1994; de Win et al., 1996). The experiments described herein clearly show that vesicles were significantly held back when they moved, and that two classes of secretory vesicle motions are present. Most of the motions are short distance. Furthermore, we found that the tracks of vesicle motions were longer and straighter in the center than in the peripheral portions, and that vesicles moved more slowly at the apical regions than in the middle and subapical regions. Using a pharmacological approach with the actin-disrupting drugs LATB and CD, we confirmed that the intact actin cytoskeleton was essential for secretory vesicle transport in pollen tubes. Indeed, this notion is supported by recent data showing that actin-dependent motions of FYVE (Fab-1, YGL023, Vps27, and EEA1)-labeled endosomes within the clear zone are essential for the tip growth of root hairs (Voigt et al., 2005; Šamaj, 2006), and that active F-actin gels drive shape oscillations in animal cells and their cell fragments (Paluch et al., 2005).

In our investigations, we observed that exocytotic vesicles entered the evanescent field and then became brighter, losing their lateral mobility while they reached a stable maximum intensity, and then their intensity declined as the dye diffused away. From the spread of fluorescence of secretory vesicles, we deduced that exocytosis is a "full" fusion event involving collapse of the vesicles into the membrane as they release of their internal components. The fusion event in living pollen tubes was not consistent with the "kiss-and-run" model recently reported in synapses (Gandhi and Stevens, 2003; Wightman and Haynes, 2004), indicating that the mechanism of vesicle fusion events in plant cells differs from that of animal cells.
Figure 5. BFA-induced changes in dynamics of secretory vesicles in living pollen tubes. A, Histograms depicting the average path lengths in apical and subapical regions after 10 min of BFA treatment (n = 8). B, Bar graph depicting the average and maximum velocities of vesicles with or without BFA treatment. C, The plot of x-y velocity of an individual vesicle in a BFA-treated pollen tube. D, MSD of vesicle motion in a BFA-treated pollen tube. MSD was calculated as a function of the observation time. E, Data from vesicle motions in BFA-treated and control pollen tubes were pooled to calculate the 2-D diffusion coefficient \( D^* \). F, The average vesicle density changed in BFA-treated pollen tubes.
the actin cytoskeleton in an intact state, so that disruption of the microtubules indirectly alters vesicle mobility. Therefore, we can safely conclude that the actin cytoskeleton is the main underlying mechanism for vesicle mobility and that microtubules within the elongating pollen tube influence microfilament organization, thus affecting the mobility of vesicles.

The fungal metabolite BFA has been widely used as a useful tool in studies of the relationship between membrane trafficking and secretion. It has been reported as inhibiting vesicle coat formation, blocking endoplasmic reticulum-to-Golgi trafficking, and disrupting the Golgi apparatus (Donaldson et al., 1992; Yasuhara et al., 1995; Robineau et al., 2000; Wang et al., 2005). Interestingly, more recent evidence indicates that BFA affects both actin organization and actin-dependent endosomal mobility within the growing tips of root hairs (Samaj et al., 2002; Voigt et al., 2005). The experiments described herein indicate that BFA causes a significant decrease in vesicle density in both the apical and subapical regions, probably as a result of its inhibitory effect on the production of secretory vesicles. Furthermore, we observed a trend of decreasing vesicle velocity and mobility in pollen tubes after treatment with BFA. Although the effect was not significant, in light of the role of the actin cytoskeleton in regulating vesicle trafficking, the reduction in vesicle mobility may be interpreted as a direct consequence of the BFA-induced modification of the actin cytoskeleton. Taken together, our experiments indicate that in P. meyeri pollen tubes, BFA affects secretory vesicle formation more than it affects vesicle mobility.

In summary, a fluorescence excitation technique using variable penetration depths of evanescent waves was modified and applied to investigations of secretory vesicles trafficking in living P. meyeri pollen tubes. The secretory vesicles were labeled with the membrane-specific dye FM4-64. Using EWM, we observed exocytosis of secretory vesicles via a “full” fusion event, and we found that there were two types of vesicle motions showing different oscillation frequencies. Contrary to earlier investigations using confocal microscopy, our results indicate that secretory vesicle motions in living pollen tubes are not Brownian, and they are dependent on the actin cytoskeleton. In addition, BFA prominently inhibits vesicle formation, but not vesicle mobility, in living pollen tubes. EWM can be used not only as an independent assay for vesicle trafficking, but also as a new, powerful tool for studies on the dynamics of vesicles, organelles, and single molecules of plant cells, which may now be undertaken in much more detail than ever before.

MATERIALS AND METHODS

Plant Materials

Cones with mature pollen were collected from Picea meyeri Rehd. et Wils growing in the Botanical Garden of the Institute of Botany, Chinese Academy of Sciences, prior to the beginning of the pollination season in mid-April 2004. Cones were dried overnight at room temperature, and the dry pollen was stored at –20°C until further use.

Pollen Culture

Stored pollen was equilibrated at room temperature for 30 min and then carefully suspended in a culture medium (2 mg of pollen per ml medium) that contained 12% (w/v) Suc, 0.03% calcium chloride, and 0.01% boric acid in a shaker at 100 rpm. The pH of the medium was adjusted to 6.4 with phosphate buffered saline. Germination took place in the dark at 25°C.

Dye Labeling

FM4-64 dye was purchased from Sigma. In this experiment, loading of cells with FM4-64 dye was generally achieved by application at a concentration of 2 μM during the inhibition of pollen tubes by direct addition of dye solutions in the culture medium. After 5 min of loading with FM4-64, the medium containing dye was replaced with fresh medium lacking the dye.

Drug Treatment

Inhibitor stock solutions were made up in 100% ethanol (2 mM CD, 20 mM oryzalin, 50 mM colchicine, and 5 mg/mL BFA), except for LATB, which was a 1 mM stock in dimethyl sulfoxide. All of the inhibitors were purchased from Sigma. The final inhibitor concentrations in the germination media were up to 10 μM for CD, 10 mM for LATB, 100 μM for oryzalin, 5 mM for colchicine, and 5 μg/mL for BFA (Supplemental Fig. 2 shows the appropriate concentrations we selected). Actin cytoskeletons were disrupted using 10 mM LATB or 10 μM CD for 10 min after labeling with FM4-64 at 25°C, respectively. The microtubule was disrupted by incubating pollen tubes with 100 μM oryzalin or 5 mM colchicine for 10 min after labeling with FM4-64 at 25°C. BFA was applied together with FM dyes at a concentration of 5 μg/mL.

Confocal Microscope Imaging

Sorbitol (100 μM) and sodium azide (500 μM) were applied directly to the medium. The pollen tubes were examined and digital images were acquired using a laser scanning confocal system installed on an inverted microscope (IX81; Olympus). The samples were excited at 514 nm with a 30-mW argon ion laser operated at full power at an intensity of 3%, achieved by means of neutral-density filters, with a nearly closed pinhole and the gain adjusted to below 7.00. Emission was detected with a 530- to 600-nm band-path filter (Zeiss RS10; Carl Zeiss). Images were collected and processed using Adobe Photoshop 7.0 (Adobe Systems).

EWM

The total internal reflection system was constructed based on an inverted microscope (IX81; Olympus). Light from a multichannel argon laser (458, 488, 515 nm; 30 mW) was introduced to the microscope through a single-mode fiber and three illumination lenses. The light was focused at the back focal plane of a high aperture objective lens (Apo 100 × OHR; NA 1.65; Olympus). This allowed us to generate evanescent waves with penetration depths from approximately 60 to 400 nm in aqueous solution (n = 1.33) at an excitation wavelength of 515 nm laser. FM4-64-labeled cells were visualized by excitation with an argon laser set to 514 nm. Emission was detected with a 530- to 600-nm band-path filter (Zeiss RS10; Carl Zeiss).

Evanescent Wave Microscope Imaging Collection and Analysis

Fluorescence was gathered through a 100 × Apo OHR objective (NA 1.65; Olympus), and the evanescent wave microscope images were captured through the right lateral port of the inverted microscope onto a cooled charge-coupled device camera (Micromax, MMX-512-BFT; Princeton Instruments), which was operated with Metamorph 6.0 (Universal Imaging). Immersion oil with a high refractive index (n = 1.63) was used to bridge the optical contact between the objective and the cover glass, and the incident light for evanescent illumination was introduced from the objective lens. Time-lapse images were acquired every 200 ms and sampled into the computer through a frame grabber with genuine 16 Bit (216, 65,536 gray levels).
Image analysis was performed with Image-Pro Plus 5.1 (Media Cybernetics), Adobe Photoshop 7.0 (Adobe Systems), and Image 1.34 (Wayne Rasband, National Institutes of Health). The raw images were filtered to enhance the visibility of the vesicles. A high-pass fast Fourier transform filter, followed by a 3 × 3-pixel trimmed mean filter, was applied to remove non-uniform background noise, and a flattened filter was used to smooth the background. The pixel size was 0.125 µm with a 1.6× optical zoom, and the image size was typically 326 × 484 pixels. The fluorescence intensity was expressed in the 8-bit value of digitization as described previously (Tsuboi et al., 2000; Taraska and Almers, 2004).

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