POD1 Regulates Pollen Tube Guidance in Response to Micropylar Female Signaling and Acts in Early Embryo Patterning in Arabidopsis

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The pollen tube germinates from pollen and, during its migration, it perceives and responds to guidance cues from maternal tissue and from the female gametophyte. The putative female cues have recently been identified, but how the pollen tube responds to these signals remains to be unveiled. In a genetic screen for male determinants of the pollen tube response, we identified the pollen defective in guidance1 (pod1) mutant, in which the pollen tubes fail to target the female gametophyte. POD1 encodes a conserved protein of unknown function and is essential for positioning and orienting the cell division plane during early embryo development. Here, we demonstrate that POD1 is an endoplasmic reticulum (ER) luminal protein involved in ER protein retention. Further analysis shows that POD1 interacts with the Ca2+ binding ER chaperone CALRETICULIN3 (CRT3), a protein in charge of folding of membrane receptors. We propose that POD1 modulates the activity of CRT3 or other ER resident factors to control the folding of proteins, such as membrane proteins in the ER. By this mechanism, POD1 may regulate the pollen tube response to signals from the female tissues during pollen tube guidance and early embryo patterning in Arabidopsis thaliana.

INTRODUCTION

Fertilization completes the cycle from the haploid gametophyte generation to the diploid sporophyte generation in the life history of animals and plants. In single-celled organisms and animals, the egg cells may emit attractant molecules, which form a concentration gradient to guide the sperm cells by chemotaxis. The sperm cells perceive and respond to these attractant signals to target the egg cells and achieve fertilization (Higashiyama and Hamamura, 2008). However, the sperm cells in flowering plants have lost their mobility during evolution. To compensate for this, a new mechanism called siphonogamy evolved in which a pollen tube is produced by the male gametophyte to deliver the two sperm cells to the female gametophyte. Pollen tubes grow unidirectionally and are guided by multiple signals from the maternal tissues and the female gametophytes (embryo sacs). Indeed, pollen tube guidance is a precisely regulated process analogous to axon guidance in animals (Palanivelu and Preuss, 2000). Pollen tube guidance can be divided into sporophytic and gametophytic guidance (Shimizu and Okada, 2000; Higashiyama et al., 2003). Sporophytic guidance refers to pollen tube growth that is guided by female sporophytic signals within the transmitting tract. Gametophytic guidance is directed by signals from the embryo sac, and it is often divided into funicular guidance and micropylar guidance. Funicular guidance directs the tubes from the septum (placenta) surface to the funicular surface, and micropylar guidance directs the tubes from the funiculus into the microprolar opening of the ovule.

Genetic studies revealed that the female gametophyte plays an important role in pollen tube guidance (Hülskamp et al., 1995; Ray et al., 1997; Shimizu and Okada, 2000). For example, cell ablation experiments in Torenia fournieri showed that the synergid cells of the embryo sac are key to attracting pollen tubes (Higashiyama et al., 2001). Several proteins produced in the embryo sac, such as MYB98 in the synergid cells (Kasahara et al., 2005; Márton et al., 2005), CENTRAL CELL GUIDANCE in the central cell (Chen et al., 2007), and GAMETE-EXPRESSED3 in the egg cell (Alandete-Saez et al., 2008), have been shown to be involved in micropylar pollen tube guidance. Recently, the secreted defensin-like peptides LUREs have been shown to be able to guide pollen tube growth in T. fournieri (Okuda et al., 2009). LUREs are Cys-rich proteins that contain a motif conserved among antimicrobial peptides.
addition, maize (Zea mays) EA1, a small peptide secreted from the egg apparatus, is also essential for micropylar pollen tube guidance (Mártion et al., 2005). These results suggest that the attracting cues from the embryo sac are small secreted peptides. Such peptides are encoded by a large family of genes in Arabidopsis thaliana, and more investigation is necessary to elucidate which are the attracting or repelling cues (Jones-Rhoades et al., 2007).

Although the pollen tube has long been considered a model system to study polar cell growth in plants (Yang, 1998; Hepler et al., 2001; Cheung and Wu, 2008; Kost, 2008; Cai and Cresti, 2009), very little is known about how it perceives and responds to the attractants to achieve directed growth. It was recently shown that Glu receptor-like proteins function as Ca^{2+} channels to regulate the cytosolic [Ca^{2+}]{$_{\text{cyt}}$} in pollen tubes, and, interestingly, the activities of these Glu receptor-like proteins are modulated by α-Ser in pistils (Michael et al., 2011). This is a conserved mechanism also used in neurotransmission in animal systems. Recently, it was reported that two endoplasmic reticulum (ER)–localized K+ transporters in pollen are involved in funicular guidance in vivo and micropylar guidance in a semi-in vivo assay (Lu et al., 2011). These findings suggest a role for cation and pH dynamics in pollen tube guidance. Mutation of the Rab GTPase RABA4D severely reduced the growth rate and efficiency of micropylar pollen tube targeting (Szumlanski and Nielsen, 2009). In addition, mutations of HAPLESS/GENERATIVE CELL–SPECIFIC1, a conserved, sperm cell–specific gene mediating cell fusion in flowering plants (Mori et al., 2006; Liu et al., 2010), also impairs pollen tube growth (von Besser et al., 2006).

Guided cell growth is a common phenomenon in the plant and animal kingdoms, and much has been learned in yeast and animal systems (Madden and Snyder, 1998; Huber et al., 2003; O’Donnell et al., 2009), but very little is known about the signaling processes in plants (Chae and Lord, 2011; Cheung and Wu, 2008). Pollen tube guidance is a species-specific process (Swanson et al., 2004; Higashiyama et al., 2006), which serves as an ideal model system to investigate mechanisms involved in signaling and guided cell growth in plants. To identify male factors controlling pollen tube guidance, we performed a forward genetic screen and identified the mutant pollen defective in guidance1 (pod1), which was found to be specifically defective in the pollen tube response to female attractants during micropylar guidance. Molecular analysis shows that POD1 is a novel ER luminal protein involved in ER protein retention and interacts with CALRETICULIN3 (CRT3), a luminal chaperone involved in Ca^{2+} homeostasis and ER quality control. This work demonstrates that POD1 plays a specific role in the micropylar response and is also essential for cell patterning during early embryogenesis in Arabidopsis.

RESULTS

Genetic Screen to Identify Male Determinants in Pollen Tube Guidance

To investigate how pollen tubes perceive and respond to the female guidance signals during pollen tube growth, we used a series of tests to identify mutants that were defective in pollen tube entry into the gametophyte. As a first step, we screened Arabidopsis Ds and T-DNA insertion lines for reduced transmission efficiency of the mutation through the male gametophyte. In this broad screen, we selected mutations that affect many processes, including pollen development, pollen function, and pollen tube guidance. Second, we tested the candidate mutants to determine whether their pollen could target ovules in a limited pollination assay. A limited number of pollen grains (<40) from these candidate mutants were pollinated manually onto a wild-type pistil, which harbors ∼50 to 60 ovules. This eliminates competition between pollen tubes and ensures that each pollen tube has the opportunity to target one ovule. To observe the entry of the pollen tubes into the ovules, 12 h after pollination the pistil was stained with aniline blue, which specifically labels the callose wall of the pollen tube. Mutants that displayed normal pollen tube growth but failed to enter the micropylar opening of the ovule were chosen for further investigation and designated as pod1.

The pod1 mutant was isolated from our Arabidopsis Ds mutant pool (Sundaresan et al., 1995). The Ds element used for mutagenesis contains a kanamycin resistance gene (Kanr), so the transmission of the mutation can be tracked by the Kanr segregation of its progeny. Progeny from a self-pollinated pod1 plant showed a Kanr/Kanr (kanamycin-sensitive) segregation ratio of 1:1 (550:554, n = 1104) (Table 1), and this ratio is stable over three consecutive generations, indicating that the mutant is heterozygous for the Ds insertion and its fertility is compromised. In addition, reciprocal crosses between the wild type and pod1 mutants were performed. When pod1/POD1 pistils were pollinated with wild-type pollen, the Kanr/Kanr segregation ratio of the F1 progeny was 1:1 (500:498). This ratio was maintained in three independent crosses, indicating that the transmission of the Ds through the female gametophyte is not affected and the pod1 ovule is completely fertile. However, when wild-type pistils were pollinated with pollen from a pod1/POD1 plant, the Kanr/Kanr segregation ratio of the F1 progeny was 0.04:1 (51:1215) with a transmission efficiency of 4.1%. This indicates that pollen development or/and function is severely affected in the pod1 mutant.

Pollen Germination and Tube Growth Are Normal in pod1

To investigate whether the reduced male transmission of pod1 is caused by a pollen developmental defect, we first checked the

Table 1. Segregation Analysis of pod1/POD1 Mutants

<table>
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<tr>
<th>Parental Genotype</th>
<th>Male</th>
<th>Female</th>
<th>Kanr</th>
<th>Kanr</th>
<th>Kanr/Kanr</th>
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<tr>
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<td>550</td>
<td>554</td>
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<tr>
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<td>1:1</td>
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<tr>
<td>pod1-1/POD1-1</td>
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<td>1215</td>
<td>0.04:1</td>
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<td></td>
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<tr>
<td>pod1-2/POD1-2</td>
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<td>500</td>
<td>1:1</td>
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<td></td>
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<tr>
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<td>342</td>
<td>0.53:1</td>
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<td></td>
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<td>pod1-1/POD1-1</td>
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<td>254</td>
<td>1.06:1</td>
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pod1-1, 1-2, and 1-3 refer to different Ds or T-DNA insertion alleles. LP: experiment carried out under limited pollination conditions. a, more than 10 replicates; b, three replicates.
morphology of mature pod1 pollen grains by 4’,6-diamidino-2-phenylindole staining and Alexander staining for cell viability. The results showed that the pollen grains from pod1/POD1 plants are morphologically normal and contain two generative nuclei and one vegetative nucleus at maturity (n = 1000) (see Supplemental Figure 1 online); no difference in morphology or cell viability was observed between mutant and wild-type pollen. This indicates that pod1 pollen develop normally. We next used an in vitro pollen germination assay to test whether the reduced male transmission of pod1 is caused by a pollen germination defect. A mean value of 81% germination (n = 857, from six independent Kanr plants) is obtained for pollen grains from pod1/POD1 plants, which is comparable to that of the wild-type pollen grains (81%, n = 211, χ² = 0.0057, P > 0.05) (Figure 1). In addition, no abnormality in pollen tube morphology or growth in vitro was observed (Figures 1A and 1B). To test pollen tube growth in vivo, 6 to 12 pollen grains from pod1/POD1 were pollinated on each wild-type pistil (24 h after emasculation). The pollinated pistils were collected 2 h later and stained with aniline blue. We found that 93.3% of the pollen grains (n = 453, χ² = 0.06, P = 3.841) germinated on the stigma, their pollen tubes entered the style, and the tubes grew within the transmitting tract at the same growth rate (Figures 2A and 2B). These data indicate that the germination rate and tube growth of pod1 pollen in the sporophytic tissues are the same as that of POD1 pollen. Taken together, these data demonstrate that the pod1 mutation does not affect pollen morphology, germination, or pollen tube growth in vitro and in vivo.

The Micropylar Response of the pod1 Pollen Tube Is Compromised

To further investigate the effect of the pod1 mutation on pollen function, we pollinated pollen grains from pod1/POD1 plants on wild-type or pod1/POD1 stigma under limited pollination conditions and then analyzed the Kanr/Kanr ratio of the F1 progeny. When wild-type stigmas were pollinated with pollen grains from pod1/POD1 plants, the Kanr/Kanr ratio of the F1 progeny was 0.53:1 (n = 523). The ratio in this limited-pollination test is higher than the 0.04:1 seen in the reciprocal crosses described above, but lower than the expected 1:1 ratio expected if pollen function is normal. This indicates that pod1 pollen transmission increases when there is no competition for ovules, suggesting that the competence of pod1 pollen is compromised. To determine whether pollen tube guidance was compromised, we tracked the path of pollen tube growth under limited pollination conditions. Some tubes approach the micropyle but fail to enter the micropylar opening of the ovule, and some pollen tubes entered the micropyle directly (Figure 2). Because the pollen used here comes from a heterozygous pod1/POD1 plant, 50% of the pollen grains should be pod1 and 50% should be POD1; if pod1 is gametophyte specific, then the POD1 pollen tubes should migrate normally, but the pod1 tubes may show defective migration. When we examined pollen tube migration, we found that the normal pollen tubes entered the micropyle directly (50%) and no twisting growth around the micropyle was observed (Figures 2C and 3A). The defective pollen tube phenotypes can be divided.

Figure 1. Germination Ratio and Pollen Tube Length in Vitro.
(A) DIC image of pollen grains from pod1/POD1 germinated in vitro.
(B) Pollen grains from the wild type germinated in vitro. Bars = 50 μm.
(C) Percentage of germination of pollen from pod1/POD1 and the wild type (WT).
(D) Pollen tube length of pod1/POD1 and the wild type 1 h after germination in vitro.
into two categories: (1) pollen tubes growing on the ovule surface or twisting around the micropyle or just turning back on the funicular surface (26.5%, \(n = 684\)) (Figures 2D, 3B, and 3C) or (2) a portion of pollen tubes twisting initially around the micropyle of the ovule and finally entering the ovule (23.5%, \(n = 684\)) to complete fertilization (Figures 2E to 2H). Furthermore, scanning electron microscopy confirmed that the 50% of pollen tubes defective in micropylar entry also displayed abnormal growth behavior (\(n = 391\)) (Figure 3). For example, we often observed two pollen tubes targeting the same ovule (Figures 3D and 3E), with the first pollen tube failing to enter the micropyle and the second pollen tube entering the embryo sac. We speculate that the first one is a \(pod1\) pollen tube and the second one is a \(POD1\) tube. Pollen tubes defective in funicular guidance in the \(pod1/POD1\) mutant were not observed in our assay. We also investigated the behavior of wild-type pollen tubes under limited pollination condition.

Figure 2. Phenotype of Pollen Tubes Germinated in Planta.

Pollen tubes are stained with aniline blue and visualized by fluorescence microscopy. (A) Pollen grains of a \(pod1/POD1\) plant germinated on the stigma and grown for 2 h. (B) Pollen grains of a wild-type plant germinated on the stigma and grown for 2 h. (C) Wild-type pollen tubes enter the micropyle directly. (D) to (I) Pollen tubes of \(pod1/POD1\) plant growing in wild-type pistils. (E) to (H) Images of the same ovule in different focal planes showing the configuration of the \(pod1\) pollen tube outside the micropyle. The pollen tube twisting outside the micropyle and growing on the integument (D) or twisting outside the micropyle and finally entering the ovule ([E] to [H]). Bar graph of the percentage of the pollen tubes showing each phenotype (\(n = 684\)). The nontargeting phenotype is shown in (D), and the targeting phenotype is shown in (E) to (H). Arrows indicate the pollen tubes. Arrowhead indicates the micropyle. WT, wild type. Bars = 400 μm in (A) and (B) and 100 μm in (C) to (H).
(pollen grains from the wild-type plants were pollinated on the wild-type stigma). Occasionally, wild-type tubes that did not target to the microlype (0.26%, \( n = 1155 \)) were observed. Together, these experiments under limited pollination indicate that \( pod1 \) pollen tubes are defective specifically in micropylar pollen tube guidance.

### POD1 Is Essential for Early Embryo Development

The low transmission rate of the \( pod1 \) mutation indicated that there could be additional defects in the \( pod1 \) mutants. For example, under limited pollination conditions, we found that 23.5% of the \( pod1 \) pollen tubes can enter the micropyle, but the Kan\(^{+}\)/Kan\(^{-}\) ratio rises from 1:1 to 1.06:1 (\( n = 508 \)), when the pollen of \( pod1/POD1 \) were pollinated on the stigma of \( pod1/POD1 \). However, under normal pollination conditions, where pollen is in excess, we seldom observe aborted seed due to the very low transmission efficiency of the \( pod1 \) mutation. The low transmission rate indicates that the \( pod1/POD1 \) embryos may be inviable. To confirm this, we self-pollinated \( pod1/POD1 \) pistils using limited pollen, then opened the silques and examined the ovules (Ding et al., 2006). When the sibling embryos reached the globular embryo stage (88.5%) within the same silique (Figure 4A), 11.5% of the embryos were arrested at several early stages and also showed aberrant division planes (\( n = 578 \)) (Figures 4B to 4I). The putative mutant zygotes underwent cell division with the aberrant cell plate usually at the wrong position or in the wrong orientation, yielding abnormal apical and basal cells (Figures 4B and 4C). The subsequent division plane of these cells was also aberrant. In the wild type, the division plane of apical cell was parallel to the apical-basal axis, but the apical cell of the putative \( pod1/pod1 \) often divided perpendicular to the apical-basal axis (Figures 4D to 4F). Rarely, the pre-embryo was able to divide further and formed an abnormal embryo, which gradually collapsed, and a suspenosor with a likely reversed apical-basal axis (Figures 4G to 4I). These observations indicate that the cell plate does not form at the right position or orientation in the putative \( pod1/pod1 \) embryos. The percentage of embryo abortion is \( \sim 11.5\% \), which is half of the percentage of \( pod1 \) pollen tube targeting under limited pollination conditions, so the aborted embryos are most likely the \( pod1/pod1 \) homozygotes. Consistent with this, \( pod1/pod1 \) homozygous seedlings were never recovered in the progeny of plants fertilized by limited pollination. These results indicate that \( POD1 \) plays an essential role in cell plate orientation/positioning in early embryo patterning.

### POD1 Encodes a Novel Conserved Protein with Unknown Function

To identify the gene disrupted in \( pod1 \) mutant, the flanking sequence of the \( Ds \) element was isolated by thermal asymmetrical interlaced PCR and sequenced. Examination of the resulting sequence showed that the \( Ds \) was inserted into the first exon of \( At1G67980 \), 31 bp downstream of the ATG start codon, resulting in an 8-bp duplication (5‘-GCCAAAAC-3’) typical for \( Ds \) insertions (Figure 5A). This \( Ds \) insertion allele is designated \( pod1-1 \). We also obtained two additional alleles, \( pod1-2 \) (SALK_049247) with a T-DNA insertion in the fourth intron and \( pod1-3 \) with a \( Ds \) insertion at 54 bp downstream of the ATG start codon in the first exon (Figure 5A). For both \( pod1-2/POD1 \) and \( pod1-3/POD1 \), when pollinated by wild-type pollen, the progeny display a Kan\(^{+}\)/Kan\(^{-}\) ratio of 1:1 (500:500 for \( pod1-2 \) and 580:582 for \( pod1-3 \); Table 1), indicating 100% transmission through the female gametophyte. The efficiency of transmission through the male gametophyte of \( pod1-2 \) is 4% (\( n = 850 \)) and the transmission of \( pod1-3 \) is 7% (\( n = 520 \)). Moreover, pollen mutant for \( pod1-2 \) or \( pod1-3 \) showed defective micropylar pollen tube guidance (see Supplemental Figure 2 online). The analysis of these mutant alleles indicates that \( At1G67980 \) is the \( POD1 \) gene, and the disruption of this gene results in a defective pollen tube response to guidance cues from the female gametophyte.

To further confirm that the phenotype is caused by mutation in \( At1G67980 \), a genetic complementation experiment was performed. A genomic fragment including a 1.628-kb promoter, the coding region including the exons and introns, and a 482-bp fragment downstream the stop codon was cloned and transformed into \( pod1/POD1 \) heterozygous plants. After double antibiotic selection for the \( Ds \) and the transgene, 30 T1 transgenic...
lines were obtained. The T2 generation seeds from six randomly chosen independent T1 lines were plated on Murashige and Skoog media supplemented with kanamycin for Kan segregation analysis. The Kan/Kan^r ratio rose to 1.8–3:1, and plants homozygous for the Ds insertion were obtained in the T3 generation of these lines (Table 2). In addition, a construct containing the POD1 coding sequence driven by the pollen-specific promoter LAT52 was transformed into wild-type plants. Six randomly selected lines out of 18 T1 transgenic plants were analyzed for pollen tube guidance, and all of them showed defective micropylar pollen tube guidance (see Supplemental Figure 3 online). Together, these data demonstrate that At1G67960 is indeed the POD1 gene. The CDS of POD1 driven by the LAT52 promoter can rescue the male defect of pod1 mutant and increase the Kan^r/Kan^s of progeny from pod1-1/POD1 from 1:1 up to 2:1 (n = 2000). The LAT52 promoter is expressed in the vegetative cell but not the sperm cell (Twell, 1992; Eady et al., 1994). This demonstrates that POD1 functions in the vegetative cell of pollen tube to control the pollen tube response.

POD1 encodes a predicted protein of 624 amino acids belonging to the eukaryotic membrane protein superfamily pfam05346, the distinguishing feature of which is the presence of a domain of unknown function 747 (DUF747). POD1 homologs exist in many eukaryotic organisms, with the amino acid sequence similarity highest in the DUF747 domain. In addition to the DUF747 domain, POD1 contains a Lys-rich motif (LR; KRKRSKKKKKK) at the N terminus, followed by an Arabidopsis-specific Gly-rich motif (GGGGSGSSGGG) and five predicted

Figure 4. Early Embryo Development in pod1/POD1.
(A) to (I) When the sibling embryos in the same silique reach the globular embryo stage (A), the putative homozygous mutant embryos are blocked at different stages (B) to (I).
(B) and (C) Mutant embryos are blocked after the first zygotic division.
(D) to (F) Mutant embryos development are blocked when the apical cell divides once with a abnormal horizontal or oblique division plane.
(G) to (I) Mutant embryos with abnormal division patterns in the apical cell and basal cell lineage. Arrows indicate the presence of an aberrant cell plate. Bars = 20 μm.

Figure 5. The Structure and Subcellular Localization of POD1.
(A) Three insertion alleles of pod1 designated as pod1-1 (Ds insertion line), pod1-2 (T-DNA insertion line), and pod1-3 (Ds insertion line).
(B) Domain structure of POD1 protein. The DUF747 domain is underlined. GR, Gly-rich motif; aa, amino acids.
(C) A confocal image showing POD1-GFP localization in an Arabidopsis protoplast. Bar = 50 μm.
(D) A confocal image of the same cell as (C) showing the localization of the ER marker mCherry-HDEL.
(E) Merged image of (C) and (D) shows the colocalization of POD1-GFP and mCherry-HDEL.
(F) A confocal image showing POD1-GFP localization in tobacco leaf cells. Bar = 50 μm.
(G) Immunoblot showing that POD1 protein is detected in total proteins (S12) and the soluble fraction (S100), but not the microsomal fraction (P100). CNX1 and PDIL1-1 are the controls as the membrane protein and soluble protein, respectively. Markers of molecular weight and POD1 protein are indicated on the left with arrows and arrowheads, respectively.
proteins are and animal counterparts, respectively. Furthermore, plant POD1 motif shows that there is high similarity within the plant homologs ER retention signals (Boulaflous et al., 2009). Sequence alignment reveals that the first (NRR1) is conserved in plants, but the second di-arginine (NRR2) is unique in the plant clade showed that the first (NRR1) is conserved in plants, monocots, moss, ferns, and yeast indicates that, in addition to pod1 mutation. Two fusion constructs, POD1-GFP and GFP-POD1, driven by the native POD1 promoter, were introduced into the pod1 mutant and were able to rescue the phenotype completely. To visualize the cellular localization of POD1, the 10-d-old seedlings were stained with the ER-specific dye ER-Tracker Blue-White DPX and then observed by confocal microscopy (Yi et al., 2009). The signal of POD1-GFP or GFP-POD1 colocalized with the ER-Tracker dye staining (see Supplemental Figure 6 online), confirming POD1’s ER localization.

To further clarify whether POD1 is a membrane protein or a soluble protein in the ER, we performed cellular fractionation by ultracentrifugation of total proteins of wild-type inflorescences. POD1 protein was detected in the soluble fraction (S100) by a POD1-specific antibody but was not detected in the microsomal fraction (P100). As a control, the ER membrane protein CXN1 was detected in the P100, and the ER luminal protein Protein Disulfide Isomerase 1-1 (PDI1-1) was only detected in the S100 fraction (Figure 5G). Together, these data confirm that POD1 is an ER luminal protein.

### POD1 Is Involved in ER Protein Retention

To dissect the functional domains of POD1, deletion and mutation experiments were performed. Sequence comparison of POD1 with its homologs in vertebrates, invertebrates, dicots, monocots, moss, ferns, and yeast indicates that, in addition to the DUF747 domain, the N- and C-terminal domains are conserved in plants (see Supplemental Figures 4 and 7 online). To determine whether the plant-specific domains are essential for POD1 function, we made two truncation constructs driven by the native POD1 promoter; the first construct lacked the N-terminal 60–amino acid residues, including the di-arginines (NRR1 and NRR2) and the LR motif (POD1<sup>CDS-Exon6</sup>), and the second lacked the last C-terminal 22 amino acids, which is hydrophilic and basic, including the tri-arginine motif (CRRR) (POD1<sup>CDS-2330</sup>) (Figure 6A). The constructs were cotransformed with mCherry-HDEL into Arabidopsis POD1 belongs to the plant clade, and the animal homologs form another clade (see Supplemental Figure 5 online).

### POD1 Is an ER Luminal Protein

To determine the subcellular localization of POD1, we transformed a POD1-GFP (green fluorescent protein) fusion construct driven by the constitutive 35S promoter of Cauliflower mosaic virus into Arabidopsis mesophyll protoplasts and tobacco (Nicotiana tabacum) leaf cells. To mark the ER, we cotransformed the cells with an ER marker, mCherry-HDEL (Batoko et al., 2000). POD1-GFP colocalized with mCherry-HDEL in Arabidopsis protoplasts (Figures 5C to 5E) and tobacco leaf cells (Figure 5F), showing a reticulate ER network. Surprisingly, no membrane localization of POD1-GFP was observed although five putative transmembrane domains were predicted by hydrophobicity analysis. To make sure that these constructs are functional, we tested whether they could complement the pod1 mutation. Two fusion constructs, POD1-GFP and GFP-POD1, driven by the native POD1 promoter, were introduced into the pod1 mutant and were able to rescue the phenotype completely. To visualize the cellular localization of POD1, the 10-d-old seedlings were stained with the ER-specific dye ER-Tracker Blue-White DPX and then observed by confocal microscopy (Yi et al., 2009). The signal of POD1-GFP or GFP-POD1 colocalized with the ER-Tracker dye staining (see Supplemental Figure 6 online), confirming POD1’s ER localization.

Table 2. Complementation Analysis of the pod1/POD1 Mutant

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<th>T2 Transgenic Plants</th>
<th>Kan&lt;sup&gt;a&lt;/sup&gt;</th>
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</tbody>
</table>

Constructs as indicated were transformed into pod1-1/POD1-1 background, and kanamycin selection was performed in T2 progeny. Genomic POD1 construct contains 1.628-kb promoter, coding region (including exons and introns), and 482-bp sequence downstream stop codon. The suffix numbers 6, 10, 18, 20, 21, and 26 indicate different transgenic lines. pLAT52:POD1 refers to POD1 CDS driven by LAT52 promoter, pPOD1:POD1<sup>RXRRR>RXAAA</sup> and pPOD1:POD1<sup>LR</sup> refer to mutated POD1 CDS driven by the POD1 promoter, respectively. The Kan<sup>a</sup>/Kan<sup>b</sup> ratio was obtained from at least five independent transgenic plants.

hydrophobic transmembrane domains (Figure 5B). In general, the Lys/Arg-rich motif functions as a putative nuclear localization signal. However, it can also serve as a recognition site for molecular chaperones or in some cases it can serve as an interaction site for protein–protein or protein–RNA interaction (Wang et al., 2000; Boeddrich et al., 2006; Teft et al., 2009). The LR may also facilitate the interaction with diacylglycerol and/or acidic phospholipids for the full function of the enzyme activity (Rodríguez-Alfaro et al., 2004). In addition, two N-terminal di-arginine motifs (NRR1 and NRR2) and one C-terminal tri-arginine motif (CRRR) are present in POD1; these may serve as potential ER retention signals. To test these possibilities, sequence comparison of POD1 with its homologs in vertebrates, invertebrates, dicots, monocots, moss, ferns, and yeast indicates that, in addition to pod1 mutation. Two fusion constructs, POD1-GFP and GFP-POD1, driven by the native POD1 promoter, were introduced into the pod1 mutant and were able to rescue the phenotype completely. To visualize the cellular localization of POD1, the 10-d-old seedlings were stained with the ER-specific dye ER-Tracker Blue-White DPX and then observed by confocal microscopy (Yi et al., 2009). The signal of POD1-GFP or GFP-POD1 colocalized with the ER-Tracker dye staining (see Supplemental Figure 6 online), confirming POD1’s ER localization.

To further clarify whether POD1 is a membrane protein or a soluble protein in the ER, we performed cellular fractionation by ultracentrifugation of total proteins of wild-type inflorescences. POD1 protein was detected in the soluble fraction (S100) by a POD1-specific antibody but was not detected in the microsomal fraction (P100). As a control, the ER membrane protein CXN1 was detected in the P100, and the ER luminal protein Protein Disulfide Isomerase 1-1 (PDI1-1) was only detected in the S100 fraction (Figure 5G). Together, these data confirm that POD1 is an ER luminal protein.
N-terminal deletion (POD1DN-ter-GFP) disrupted the ER localization of POD1 (Figure 6A), respectively. Mutated POD1-GFP constructs (POD1m-GFP) were transiently expressed in Arabidopsis protoplasts. The result shows that mutations in neither NRR1 nor NRR2 disrupted the ER localization of POD1 (see Supplemental Figure 8 online). In the case of CRRR, the POD1RXRRR>RXAAA mutation, but not POD1RXRRR>RXAA or POD1RXRRR>RXAA, impaired the ER localization of mCherry-HDEL but not POD1 itself (Figures 6H to 6J). These data indicate that the tri-arginine motif of POD1 is essential for ER retention of HDEL proteins and this process requires at least one Arg in the tri-arginine motif.

To investigate whether ER retention is essential for pollen tube guidance, POD1RXRRR>RXAAA driven by its native promoter was introduced into the wild-type and pod1/POD1 plants. Indeed, micropylar pollen tube targeting was affected in the transgenic plants under limited pollination conditions (Figure 7A). However, after twisting, ~80% of the pollen tubes (n = 800) finally enter the micropyle and finish fertilization. The POD1RXRRR>RXAAA construct can recover the Kan'/Kan+ ratio of pod1/POD1 from 1:1 to 2:1 in the T2 generation (n = 1786). In addition, pollen tubes carrying the POD1RXRRR>RXAAA construct displayed a zigzag morphology (Figure 7A), and this implies that the POD1RXRRR>RXAAA mutation also impairs pollen tube morphology.

As shown in Figures 6B to 6D, the deletion of the N-terminal 60 amino acids (POD1DN-ter-GFP) impairs the ER localization of POD1. This raises the possibility that an ER localization signal may exist in the N-terminal region. To our surprise, POD1DN-ter-GFP was mainly targeted to the nucleus (Figures 6K to 6M). This indicated that the N-terminal sequence may be targeted into the nucleus by the LR motif, although we did not observe nuclear localization of the full-length POD1 protein. Usually the Lys-rich motif is predicted to be a nuclear localization signal. To analyze the function of this potential nuclear localization signal, POD11LR with the KKRKSKKK sequence deletion was fused with GFP reporter gene. Transient assay in protoplasts showed that the localization pattern of POD11LR-GFP did not change compared with that of POD1, but a portion of mCherry-HDEL escaped from the ER (Figures 6N to 6P). This is nearly the same as that of the POD1RXRRR>RXAAA mutation. Interestingly, transformation of this mutated POD1 into wild-type plants also reduced the ratio of microplar pollen tube targeting (78%, n = 496) and resulted in abnormal tube morphology in vivo, similar to that of POD1RXRRR>RXAAA (Figure 7B). Consistent with this, the POD11LR mutation can also restore the Kan'/Kan+ ratio of pod1/POD1 plants to 2:1 (n = 1291). These results confirm the indispensable role of the conserved Lys/Arg-rich motifs for POD1. These studies suggest that the N-terminal Lys/Arg-rich motif and the C-terminal tri-arginine of POD1 are essential for HDEL protein retention, which is required for pollen tube response in Arabidopsis. An HDEL retrieval signal generally exists in some ER luminal chaperones. Therefore, POD1 may be involved in the ER retention of these chaperones.

**Figure 6.** Mutation Analysis of POD1.

(A) A diagram showing different mutations and deletions of POD1 protein for POD1-GFP fusion constructs. aa, amino acids. 
(B) to (P) Confocal images showing the localization of POD1m-GFP ([B], [E], [H], [K], and [N]), ER marker mCherry-HDEL ([C], [F], [J], and [O]) in respective channels, and the merged images ([D], [G], [I], [M], and [P]). 
(B) to (D) Confocal images of the same protoplast showing the N-terminal deletion (POD1DN-ter-GFP) disrupted the ER localization of POD1. 
(E) to (G) Confocal images of the same protoplast showing C-22aa deletion (POD1DC-22aa-GFP) abolished the ER localization of POD1. 
(H) to (J) Confocal images of the same protoplast showing that the CRRR mutation (POD1RXRRR>RXAAA-GFP) disrupted the ER localization of mCherry-HDEL. Arrows indicate the mislocalization of mCherry-HDEL. 
(K) to (M) Confocal images of the same protoplast showing the nuclear localization of POD11LR-GFP. 
(N) to (P) Confocal images of the same protoplast showing POD11LR-GFP disrupted the ER localization of mCherry-HDEL. Arrows indicate the mislocalization of mCherry-HDEL.

DN-ter, POD1 with the 60-amino acid N-terminal sequence deleted; DC-22aa, POD1 with the C-terminal 22 amino acids deleted; Nter, The N-terminal sequence including the LR; ΔLR, POD1 with the LR deleted. Bar = 50 μm.

**POD1 Interacts with CRT3**

To investigate whether POD1 acts as an ER chaperone as it plays a role in ER protein retention, a yeast two-hybrid screen was performed. One candidate protein isolated was CRT3. To
confirm the interaction, the full-length POD1 CDS and CRT3 CDS with the signal peptide for ER targeting deleted were cloned into pGBK7 and pGADT7, respectively. The yeast cells cotransformed with these two plasmids grew well on the Trp-, Leu-, His-, Ade-dropout media (Figure 8A), indicating an interaction between these two proteins. To further test if POD1 also interacts with the other two closely related ER chaperones, CRT1 and CALNEXIN (CNX) with both the signal peptide and the transmembrane domain deletion were cloned into pGADT7. Yeast cells cotransformed with pGBK7-POD1 and pGADT7-CRT1 or pGADT7-CNX did not grow on the Trp-, Leu-, His-, Ade-dropout media. This suggests that POD1 interacts specifically with CRT3 but not with its homologs CRT1 and CNX. The interaction between POD1 and CRT3 was further confirmed by bimolecular fluorescence complementation (BiFC). Protoplasts cotransformed with NYFP-CRT3 and CYFP-POD1 showed a yellow fluorescent protein (YFP) signal (Figure 8B), and no YFP signal was detected in the control experiment.

Consistent with the interaction of CRT3 with POD1, CRT3 is expressed in pollen grains and pollen tubes, as revealed by Arabidopsis microarray analysis (Qin et al., 2009). Similarly, the transcripts of CRTs are detected in pollen grains, pollen tubes, and pistils in Haemanthus (Lenartowska et al., 2009). Together, these data suggest that POD1 most likely interacts with CRT3 and plays an important role in ER quality control.

Expression Pattern of POD1 in Planta

To characterize the temporal and spatial regulation of POD1, its expression pattern was analyzed. First quantitative RT-PCR (qRT-PCR) was performed with RNAs from root, stem, leaf, inflorescence, silique, and shoot. The data showed that POD1 mRNA is present in multiple tissues, with a relatively high level in inflorescence, silique, root, and shoot (Figure 9A). To provide a more precise expression pattern, a genomic sequence fusion with ß-glucuronidase (GUS) or GFP driven by the POD1 native promoter was transformed into Arabidopsis, and four of five transgenic lines showed positive GUS staining. The GFP line is the same as in Supplemental Figure 6 online. GUS staining of transgenic lines and confocal microscopy of GFP-expressing transgenic lines showed that POD1 is expressed in mature pollen and pollen tubes (Figures 9B, 9C, 9H, and 9I). Although pod1 shows no female gametophytic defect, POD1 is expressed abundantly in synergids and weakly in the antipodal cells (Figures 9D, 9J, and 9K). When pollen from the GUS line was pollinated onto the wild-type stigma, the GUS signal was observed at the tip of the pollen tube (Figure 9E). When the tubes burst in the embryo sac, the POD1-GUS protein was released into the synergid cell (Figure 9F). Consistent with the phenotype of embryo abortion, POD1 is also expressed in the early embryo...
and endosperm (Figure 9G). The expression pattern of *POD1* suggests that it functions in multiple developmental processes.

**DISCUSSION**

We identified a male gametophytic mutant specifically defective in micropylar pollen tube guidance and cell plate patterning during early embryogenesis. Confocal microscopy and cellular fractionation showed that *POD1* is an ER luminal protein. Mutagenesis indicates that the C-terminal tri-arginine motif and the N-terminal Lys/Arg-rich motif are essential for the function of *POD1* in the ER retention of proteins with an HDEL motif. Interaction of *POD1* with CRT3 indicates that *POD1* might act as a cochaperone in the ER lumen. *POD1* likely acts as a component of the CRT3 complex, which is involved in ER retention and perhaps also in ER-mediated protein secretory pathways, to regulate the pollen tube response to female gametophyte signals and cell plate patterning during early embryogenesis in plants.

**POD1 Functions in ER Protein Retention**

ER resident proteins are necessary for normal ER function. They usually contain specialized targeting signals (the signal peptide) consisting of specific amino acids, which is recognized by the signal recognition particle and translocated into the ER. The
signal peptide is often located at the N or C terminus or sometimes internal to the proteins. In general, the signal peptides are cleaved by signal peptidase, but in some cases, they are retained in the mature protein sequence (Lingappa et al., 1979; Guo et al., 2011). Our data suggest that the N-terminal 60 amino acids and the C-terminal 22 amino acids are essential for the localization of POD1 in the ER. Therefore, the ER targeting signal of POD1 may reside in these two regions, but no typical ER retention or targeting signal peptide is predicted in these regions. The fact that both the N- and C-terminal GFP fusion proteins of POD1 showed ER localization suggests that neither the N nor C terminus is cleaved by signal peptidase. So, we propose two possible molecular mechanisms of the ER localization of POD1. First, POD1 does not possess the typical ER retention or targeting sequence, and its translocation into the ER may be a signal recognition particle independent. Second, the N- or C-terminal domains required for POD1 ER retention may interact with ER resident proteins that possess the HDEL retrieval signals. Both scenarios require more studies on POD1 sequence and its interacting proteins. Further detailed mapping of the N- and C-terminal domains essential for the ER localization of POD1 will be helpful to identify new mechanism of the ER retention signal.

POD1 functions in the retention of ER proteins with HDEL retrieval signals by an unknown mechanism. It is known that ER luminal resident proteins are constitutively secreted to the Golgi. To be retained in the ER, soluble ER-resident proteins have to escape from the bulk flow of secreted proteins. Many ER luminal proteins, such as CRT, PDIL, CNX, and BiP, have a C-terminal H/KDEL motif that acts as an ER retrieval signal (Munro and Pelham, 1987). H/KDEL is recognized and retrieved by the integral membrane H/KDEL receptors in the Golgi complex (Pagny et al., 1999; Capitani and Sallese, 2009). However, the amount of secretion of resident proteins, such as CRT3 and BiP, is very low, and one possible reason for this is that these chaperones may associate with newly folding proteins (Pfeffer, 2007). So H/KDEL acts by retrieval rather than retention and other mechanisms yet to be unravelled (Capitani and Sallese, 2009). This view is also supported by the fact that some ER chaperones, like SDF2 and ERdj3b, lack an H/KDEL motif at the C terminus. Our data show that POD1 acts as an ER luminal protein without the K/HDEL motif and plays a role in the retrieval of HDEL proteins. The basic motifs in the N and C terminus are likely involved in this process. Two dominant-negative mutations, POD1R239R-R240A and POD1L273L, cause migration of mCherry-HDEL protein out of the ER and impair pollen tube guidance and morphology. Therefore, ER retention of specific ER resident proteins by POD1 plays an important role in pollen tube guidance. Similarly, mutations in YER1140W, the yeast homolog of POD1, result in the escape of a well-characterized ER chaperone (BiP) from the ER (Copic et al., 2009) and consequently inhibit filamentous growth (Jin et al., 2008) and induce unfolded protein response (Jonikas et al., 2009). So HDEL protein retention may be a general function of POD1 and its homologs in eukaryotic organisms.

The Role of POD1 in Pollen Tube Guidance and Early Embryogenesis

Pollen tube growth in the pistil is a continuous process of guidance and reception. Previous genetic and molecular analy-

sis suggests that different mechanisms function during sporophytic and gametophytic pollen tube guidance (Ray et al., 1997; Higashiyama et al., 1998; Shimizu and Okada, 2000; Chae and Lord, 2011). The ER luminal POD1 acts specifically in the micropylar response to female cues, but obviously through an indirect mechanism. The interaction with CRT3 suggests that POD1 may be involved ER quality control, in which ER chaperones assist and monitor the proper maturation and formation of the nascent proteins in the secretory pathway. In Arabidopsis, ~30% of total proteins enter the secretory pathway through the ER, and among them 33% are transmembrane proteins.

ER-dependent maturation of membrane receptors during host–pathogen interactions in plants has been reported. CRTs are multifunctional ER chaperones involved in protein folding and Ca²⁺ homeostasis. Three CRTs are found in the Arabidopsis genome, and CRT1 and CRT2 function as general ER chaperones, while CRT3 is specifically associated with the formation of some membrane receptors, such as elongation factor Tu receptor (EFR) (Christensen et al., 2010). The maturation of EFR is dependent on the ER quality control components STT3A, SDF2, BiP, and CRT3. These proteins are essential for the N-glycosylation, accumulation, folding, and maturation of EFR. The processing of EFR is abolished and the immunity response to the pathogen is lost when the genes encoding the chaperones are knocked out (Nekrasov et al., 2009; Häweker et al., 2010). In rice (Oryza sativa), the ER chaperone BiP3 interacts with the pattern recognition receptor XA21, which is involved specifically in resistance to pathogen Xoo (Park et al., 2010). It has also been shown that SHEPHERD, an ER-resident HSP90-like protein in Arabidopsis, controls the formation and/or folding of CLAVATA, a receptor controlling meristem activity (Ishiguro et al., 2002). Similarly, the folding of the bradykinin receptor in ES cells and the Toll receptor in mice is also dependent on ER chaperones (Nakamura et al., 2001; Randow and Seed, 2001). All these findings suggest that the ER quality control system is essential for formation and accumulation of receptors. In this context, it is plausible to speculate that POD1, via modulating CRT3 function, plays a key role in the ER quality control of putative membrane receptors perceiving the female signal molecules, such as LUREs, during pollen tube guidance in plants.

In addition to protein folding and vesicle trafficking, ion homeostasis, which is fundamental to the axis of polarity in yeast, animals, and plants, requires the proper function of the ER. So another possibility is that POD1 affects the folding of ER-localized cation transporters, such as the putative K⁺ transporters CHX21 and 23 (Lu et al., 2011) and cyclic nucleotide-gated channels (Friesch et al., 2007), and subsequently disrupts the Ca²⁺ and K⁺ homeostasis in the ER and Ca²⁺ signaling. Another possibility is that POD1 may regulate cytosolic Ca²⁺ homeostasis and its regional concentration by tethering CRT3 activity to mediate the precise micropylar guidance. Finally, POD1 may have a general effect on proteins that are folded or modified in the ER. To clarify these possibilities, more factors involved in the pollen tube response need to be identified. It would be interesting to compare the proteome of pod1 with that of the wild-type pollen tubes, which may facilitate the identification of POD1-interacting proteins that may be the male factors directly involved in pollen tube guidance.
In addition to its role in pollen tube guidance, POD1 is also important for early embryogenesis. Occasionally, the pod1 egg cells can be fertilized by the pod1 sperm cells under limited pollination to form arrested zygotes or embryos that are defective in positioning and orientation of the cell plate. In mouse, mutation of TAP71, the putative homolog of POD1, results in a reversed posterior-anterior axis polarity of the axial skeleton, suggesting a role for TAP71 in body axis formation (Howell et al., 2007). Similar reversion of apical-basal polarity likely takes place in pod1 embryos in Arabidopsis, as shown earlier. The knockdown of POD1 homolog NP507972 in Caenorhabditis elegans causes embryo lethality, but no detailed analysis of embryo polarity has been reported (http://www.wormbase.org/).

The mechanism of the pod1 embryo defect is not clear, but it is plausible to speculate that the ER quality control and subsequently the secretory systems that require the function of POD1 are needed for positioning and orientation of the cell plate in zygotes and embryos. In plant cells, studies on cell plate orientation have revealed roles of microtubule and associated proteins on preprophase band and phragmoplast formation (Müller et al., 2009) but have not yet provided clues on the role of the ER in these processes. It is believed that cell plate formation requires the deposition of newly synthesized materials from Golgi apparatus–derived secretory vesicles. Recently, it was also shown that protein secretion and endocytic pathways are essential for cell plate formation (Dhonoukhé et al., 2006; Reichardt et al., 2007; Boutté et al., 2010). The inhibition of ER-Golgi trafficking causes the retention of newly synthesized proteins, such as the cytokinesis-specific syntaxin KNOLLE in the ER, therefore preventing cell plate formation (Reichardt et al., 2007). In conclusion, POD1 might function in ER quality control or the protein secretory system that is required for proper positioning and orientation of the cell plate during embryogenesis.

METHODS

Plant Materials and Growth Conditions

Seeds of Arabidopsis thaliana were surface-sterilized with 20% bleach for 10 min, washed four times with sterile water, and plated onto Murashige and Skoog medium (Sigma-Aldrich) supplemented with 50 mg/L kanamycin for Ds and T-DNA insertion lines or 40 mg/L hygromycin for pCAMBIA-based constructs. After 7 d of growth on a Murashige and Skoog plate in the greenhouse (16 h light/8 h dark, 22°C), seedlings are transferred to soil for further growth. SALK_049247 was obtained from The Arabidopsis Information Resource (TAIR) seed stock center. pod1-3 is a Ds insertion line obtained from De Ye’s lab at China Agricultural University.

Phenotypic Analysis

To investigate pollen development, flowers were dipped several times in the 4',6-diamidino-2-phenylindole or Alexander solution droplets directly on microscope slides and observed under a fluorescence microscope. For the pollen in vitro germination assay, mature pollen grains were spread onto the pollen germination medium containing 10 mM CaCl2, 10 mM Ca(NO3)2, 10 mM MgSO4, 10 mM H3BO3, 1.8% Suc, and 0.1% agarose and cultured for 5 to 12 h at 25°C. The aniline blue staining, GUS staining, and scanning electron microscopy were performed as described previously (Chen et al., 2007). For embryo observation, the ovules were dissected out with a fine needle, cleared according to Li et al. (2010), and observed using a Zeiss microscope equipped with differential interference contrast (DIC) optics.

Protein Extraction and Immunoblot Analysis

Flowers from Arabidopsis were collected and ground in liquid nitrogen to a fine powder, and then total protein was extracted with ice-cold extraction buffer (50 mM Tris, pH 7.5, 1 mM EDTA, and 0.25 M Suc supplemented with PMSF and protease inhibitor cocktail [Roche]). The mixture was centrifuged at 1,000 g for 25 min at 4°C. Supernatant (S1) was filtered through four layers of miracloth (Calbiochem). The filtered supernatant was centrifuged at 12,000 g for 30 min. Then the supernatant (S2) was transferred to a new ultracentrifuge tube (Beckman) and subsequently centrifuged at 100,000 g for 2 h at 4°C. The supernatant (S100) was collected and the pellet (P100) was rinsed several times with cold extraction buffer. The pellet was then incubated with detergent buffer (5 mM HEPES-KOH, pH 7.0, 20% Triton X-100, and 1% Suc) on ice for 30 min. All the samples were mixed with SDS-PAGE loading buffer and boiled before gel fractionation. Immunoblots were performed using specific antibody, which was generated with the POD1 peptide INRCRRRSSHLHND as antigen to immune rabbit and affinity purified. Antibody specificity of POD1, CNX1, and PDIL1-1 was verified using protein extracts from the wild type and corresponding mutant or transgenic plants carrying 3SS:FLAG-POD1.

Constructs and Plant Transformation

All primers are listed in Supplemental Table 1 online. For the complementation construct, a 5479-bp genomic fragment containing 1.628-kb promoter region and the transcribed region (including intron and exon) plus a 482-bp fragment downstream stop codon was amplified with primers POD1-F and POD1-R from Arabidopsis genomic DNA and ligated to pCAMBIA1300 to generate p1300-gPOD1-Ter, which was transformed by floral dip method into pod1 mutants (Clough and Bent, 1998). Similarly, the pPOD1-gPOD1-GUS fusion with GUS CDS inserted before the stop codon was made and transformed into Arabidopsis plants for POD1 gene expression analysis. The POD1 CDS (1975 bp in length) amplified with POD1-F and POD1-R was cloned into p1300 flanked by the LAT52 promoter and the 498 bp POD1 3’ untranslated region amplified with primers T-F and T-R to generate LAT52-POD1. The LAT52-POD1AS construct was cloned in the same way as the LAT52-POD1 but with the POD1 CDS in a reverse direction.

For subcellular localization of POD1, pBSK-35S:POD1-EGFP-polyA was made by subcloning POD1 CDS into pBSK-35S:EGFP-polyA at the SalI-BamHI site. All the deletion and mutation constructs were cloned into pBSK-35S-EGFP-polyA to produce a POD1-GFP fusion. For the BIFC constructs, the full-length CDS of POD1 and CRT3 was cloned into the Xbal and KpnI sites of pSPYNE and pSPYCE plasmid (Walter et al., 2004). The protoplast transformation was performed by the method described by Asai et al. (2002). The protoplasts were transformed with NYFP-CRT3 and CYFP-POD1, and protoplasts were transformed with NYFP-CRT3 plus CYFP and NYFP plus CYFP-POD1 as the negative control. The protoplasts were then viewed with a Zeiss LSM510 META laser scanning microscope with a 488- and 543-mm laser. For ER staining, seedlings were stained in 2 μM ER Tracker Blue-White DPX (Invitrogen) solution diluted in W5 buffer (154 mM NaCl, 125 mM KCl, 5 mM Glc, and 0.03% MES, pH 5.8) for 15 min. The signal was excited with a 405-nm laser and detected using the band-pass 420- to 480-nm filter.

Yeast Two-Hybrid Assay

The POD1 CDS and CRT3 with its ER targeting signal deleted were amplified from floral cDNAs with primers POD1y-F/POD1y-R and CRT3y-
The total RNA was extracted from different Arabidopsis tissues using TRIzol reagent (Invitrogen). First-strand cDNA synthesis was performed in a 20-μL reaction mixture with 1 μg of DNase-digested (Roche) total RNA using the superscript III reverse transcriptase (Invitrogen) with random primers (TaKaRa) according to the manufacturer’s instructions. The cDNA product was diluted and 2 ng was used in each qRT-PCR. Real-time PCR reactions containing SYBR Green PCR master mix were performed on the ABI PRISM 7900 HT Real-Time PCR system (Applied Biosystems). The following thermal reaction was used: 50°C for 2 min, 95°C for 5 min, and 40 cycles of 95°C for 15 s and 60°C for 1 min. Three technical replicates were performed with each cDNA sample. Primers for 18s rRNA were used to normalize the qRT-PCR data. The analysis of the qRT-PCR data was performed according to the geNORM manual (http://medgen.ugent.be/~jvdesomp/genorm/). The primers used were POD1q-F and POD1q-R. Primer sequences are provided in Supplemental Table 1 online.

Supplemental Figure 1. The Development of pod1 Pollen Is Normal.

Supplemental Figure 2. pod1-2 and pod1-3 Pollen Tubes Showing the Micropylar Targeting Defect.

Supplemental Figure 3. Defective Pollen Tube Guidance in pod1 Antisense Transgenic Plants.

Supplemental Figure 4. Sequence Alignment of POD1 and Its Homologs.

Supplemental Figure 5. Phylogenetic Tree of Arabidopsis POD1 and Its Homologs.

Supplemental Figure 6. POD1-GFP Is Localized in the ER.

Supplemental Figure 7. Sequence Alignment of the N Terminus and C Terminus Showing NRR1, NRR2, CRRR, LR, and C-22aa Sequences.

Supplemental Figure 8. POD1RXXXR→AXXXA and POD1RR→AA Localization in the ER.

Supplemental Table 1. List of Primers Used in This Study.

Supplemental Data Set 1. List of Primers Used in This Study.

Bioinformatics and Phylogenetic Analysis

We used the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/) and TAIR (http://www.Arabidopsis.org) to analyze the cDNA and genomic sequence. The phylogenetic tree is processed with MEGA5 software. The alignment in Supplemental Figure 4 was generated using the alignment program ClustalW (network protein sequence analysis). The alignment used for phylogeny is produced with ClustalW2 program at the European Bioinformatics Institute (http://www.ebi.ac.uk/clustalw/index.html). The weight matrix used was Gonnet with the penalty of the gap opening 10 and gap extension 0.1 for slow pairwise alignment and gap opening 10, gap extension 0.2, and gap distance 5 for multiple sequence alignment. The alignment was manually fine-tuned with DNA-MAN software (http://www.lynnnon.com/pf/framepc.html) and has been made available in FASTA format (see Supplemental Data Set 1 online). The phylogeny of the aligned sequences was generated with neighbor-joining algorithm (Saitou and Nei, 1987) by the MEGA5 software (Tamura et al., 2011), and bootstrap analysis was conducted with 1000 replications.

Accession Numbers

Sequence data from this article can be found in the Arabidopsis Genome Initiative or GenBank/EMBL databases under the following accession numbers: POD1, AT1g67960; CNX1, AT5G61790; CRT3, AT1G08450; and PDIL1-1, AT1G21750.

Supplemental Data

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

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Author Contributions

H.-J.L. and W.-C.Y. designed the research and wrote the article. H.-J.L., Y.X., and T.W. performed the research and analyzed the data. F.C. and Q.X. provided technical assistance. H.-J.L. and W.-C.Y. provided the plant material of Arabidopsis thaliana. J.L. and W.-C.Y. provided the antibody of CNX and PDIL2-1. D.-J.J. and D.Y. provided the antibody of CNX and PDIL2-1. D.-J.J. and D.Y. provided the plant material of pod1-3. D.-Q.S. and J.L. provided technical assistance.

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References

F/CRT3y-R (primer sequences in Supplemental Table 1 online) and subcloned into pGBK7T and pGADT7 (Clontech), respectively. The CRT1 cDNA was amplified with CRT1-F and CRT1-R, and CALNEXIN was amplified with CNX-F and CNX-R from plasmid, which was a kind gift from Jianming Li (Michigan State University).

qRT-PCR

The total RNA was extracted from different Arabidopsis tissues using TRIzol reagent (Invitrogen). First-strand cDNA synthesis was performed in a 20-μL reaction mixture with 1 μg of DNase-digested (Roche) total RNA using the superscript III reverse transcriptase (Invitrogen) with random primers (TaKaRa) according to the manufacturer’s instructions. The cDNA product was diluted and 2 ng was used in each qRT-PCR. Real-time PCR reactions containing SYBR Green PCR master mix were performed on the ABI PRISM 7900 HT Real-Time PCR system (Applied Biosystems). The following thermal reaction was used: 50°C for 2 min, 95°C for 5 min, and 40 cycles of 95°C for 15 s and 60°C for 1 min. Three technical replicates were performed with each cDNA sample. Primers for 18s rRNA were used to normalize the qRT-PCR data. The analysis of the qRT-PCR data was performed according to the geNORM manual (http://medgen.ugent.be/~jvdesomp/genorm/). The primers used were POD1q-F and POD1q-R. Primer sequences are provided in Supplemental Table 1 online.

Sequence data from this article can be found in the Arabidopsis Genome Initiative or GenBank/EMBL databases under the following accession numbers: POD1, AT1g67960; CNX1, AT5G61790; CRT3, AT1G08450; and PDIL1-1, AT1G21750.

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Author Contributions

H.-J.L. and W.-C.Y. designed the research and wrote the article. H.-J.L., Y.X., and T.W. performed the research and analyzed the data. F.C. and Q.X. provided the antibody of CNX and PDIL2-1. D.-J.J. and D.Y. provided the plant material of pod1-3. D.-Q.S. and J.L. provided technical assistance.

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