

The Role of LORELEI in Pollen Tube Reception at the Interface of the Synergid Cell and Pollen Tube Requires the Modified Eight-Cysteine Motif and the Receptor-Like Kinase FERONIA

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In angiosperms, pollen tube reception by the female gametophyte is required for sperm release and double fertilization. In *Arabidopsis thaliana lorelei (lre)* mutants, pollen tube reception fails in most female gametophytes, which thus remain unfertilized. *LRE* encodes a putative glycosylphosphatidylinositol (GPI)-anchored surface protein with a modified eight-cysteine motif (M8CM). *LRE* fused to citrine yellow fluorescent protein (*LRE*-cYFP) remains functional and localizes to the synergid plasma membrane-rich filiform apparatus, the first point of contact between the pollen tube and the female gametophyte. Structure-function analysis using *LRE*-cYFP showed that the role of *LRE* in pollen tube reception requires the M8CM, but not the domains required for GPI anchor addition. Consistently, *LRE*-cYFP-TM, where GPI anchor addition domains were replaced with a single-pass transmembrane domain, fully complemented the pollen tube reception defect in *lre-7* female gametophytes. Ectopically expressed and delivered *LRE*-cYFP from pollen tubes could non-cell-autonomously complement the pollen tube reception defect in *lre* female gametophytes, only if they expressed *FERONIA*. Additionally, pollen tube-expressing *LRE* variants lacking domains critical for GPI anchor addition also rescued *lre* female gametophyte function. Therefore, *LRE* and *FERONIA* jointly function in pollen tube reception at the interface of the synergid cell and pollen tube.

INTRODUCTION

The success of plant reproduction depends on a series of cell-cell interactions between the male and female gametophytes (Kessler and Grossniklaus, 2011; Palanivelu and Tsukamoto, 2012; Beale and Johnson, 2013; Qu et al., 2015). In *Arabidopsis thaliana*, the mature male gametophyte (pollen) consists of two sperm cells and a vegetative cell (Twell, 2011), whereas the female gametophyte consists of five accessory cells (three antipodal cells and two synergids) and two gametic cells (one egg cell and one central cell) (Yadegari and Drews, 2004; Yang et al., 2010; Song et al., 2014). Synergids facilitate the fusion of one sperm cell to the egg cell and one sperm cell to the central cell to complete double fertilization, forming an embryo and endosperm, respectively (Russell, 1992). Synergids are specialized secretory cells with a unique structure called the filiform apparatus (FA) at the micropylar pole. In the FA, the plasma membrane surface area is vastly increased by many finger-like projections, which extend into the synergid cytoplasm (Mansfield et al., 1991; Huang and Russell, 1992; Kasahara et al., 2005). Thus, the unique structure of the FA increases the area of contact between the pollen tube and the synergid cell and

facilitates secretion of small peptides to influence pollen tube attraction and pollen tube reception, which is defined as pollen tube burst and release of sperm cells for double fertilization (Russell, 1992; Huck et al., 2003; Okuda et al., 2009; Amien et al., 2010; Takeuchi and Higashiyama, 2012).

In *Arabidopsis*, genes expressed in female and male gametophytes regulate pollen tube reception. For example, the female gametophyte-expressed gene *FERONIA* (*FER*) encodes a receptor-like kinase. The *fer* female gametophyte fails to induce pollen tube reception and consequently remains unfertilized (Huck et al., 2003; Escobar-Restrepo et al., 2007; Haruta et al., 2014). Many mutants cause a similar phenotype due to defects in the female gametophyte: these include *scylla* (*sy*), *lorelei* (*lre*), *nortia* (*nta*), *evan*, and *turan* (Capron et al., 2008; Rotman et al., 2008; Kessler et al., 2010; Tsukamoto et al., 2010; Lindner et al., 2015). Of these, the identity of *SYL* remains unknown. *LRE* encodes a putative glycosylphosphatidylinositol (GPI)-anchored surface protein (Capron et al., 2008; Tsukamoto et al., 2010), and *NTA* encodes a transmembrane protein, a member of the Mildew Resistance Locus O family (Kessler et al., 2010). *LRE* and *NTA* function with *FER* in pollen tube reception (Kessler et al., 2010; Duan et al., 2014; Li et al., 2015; also see below). *TURAN* and *EVAN* may *N*-glycosylate synergid-expressed proteins that are involved in pollen tube reception (Lindner et al., 2015). Also, maize (*Zea mays*) *EMBRYO SAC4* induces pollen tube burst via opening of the potassium channel *KZM1* in the pollen tube (Amien et al., 2010).

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In the male gametophyte, transcription factor genes *MYB97*, *MYB101*, and *MYB120* are preferentially induced in pollen tubes growing through pistils (Qin et al., 2009; Leydon et al., 2013). In the triple *myb* mutant, the pollen tube coils in the female gametophyte and fails to discharge sperm cells (Leydon et al., 2013; Liang et al., 2013), demonstrating an active role for the pollen tube in pollen tube reception. Loss of *AUTO-INHIBITED Ca²⁺ ATPASE9 (ACA9)*, which encodes a calcium efflux pump, results in some pollen tubes failing to discharge the sperm, even though they reach the synergid and arrest growth in the synergid cell; thus, *aca9* tubes differ from triple *myb* mutant tubes in that they fail to coil and burst (Schiøtt et al., 2004). Interestingly, *abstinence by mutual consent (amc)* mutants show pollen tube reception defects only when an *amc* pollen tube encounters an *amc* female gametophyte, indicating that pollen tube reception requires interactions between the male and the female gametophyte and that both gametophytes share common signaling components (Boisson-Dernier et al., 2008).

Among the proteins involved in pollen tube reception, FER and LRE function together in the FER-RAC/ROP signaling complex to regulate reactive oxygen species (ROS) production in the micro-pylar region of female gametophytes (Duan et al., 2014). Additionally, ROS produced in the female gametophyte are important for pollen tube reception, and ROS-mediated pollen tube rupture requires calcium (Duan et al., 2014). Indeed, changes in cytoplasmic calcium ($[Ca^{2+}]_{cyto}$) observed during pollen tube-synergid interactions (Iwano et al., 2012; Denninger et al., 2014; Hamamura et al., 2014) showed that the initiation of calcium oscillations and the increase in $[Ca^{2+}]_{cyto}$ are affected in *fer* and *lre* mutant synergids (Ngo et al., 2014). In addition to affecting ROS and calcium production in ovules, LRE also has an intracellular role in chaperoning FER from the endoplasmic reticulum (ER) to the FA of the synergids (Li et al., 2015). FER and NTA also function together in pollen tube reception: The preferential relocation of NTA from puncta in synergids to the FA upon pollen tube arrival requires FER (Kessler et al., 2010), indicating that NTA functions downstream of FER. Calcium may also play an important role in mediating the function of NTA. Unlike FER and LRE, NTA is required to modulate the magnitude of the calcium signatures in the synergids and therefore likely functions downstream of FER and LRE (Ngo et al., 2014).

LRE is expressed in ovules but not in the pollen or pollen tubes (Tsukamoto et al., 2010). Consistent with this, *lre* mutants only showed defects in the female gametophyte (Capron et al., 2008; Tsukamoto et al., 2010). *LRE* encodes a putative GPI-anchored surface protein. In eukaryotes, GPI-anchored pre-pro-proteins (nascent full-length protein) usually contain an N-terminal signal peptide (SP) for ER translocation, a proline-rich unstructured ω -1 region, and two domains that are critical for addition of the GPI anchor: (1) the ω -site region containing four tiny residues, one of which will serve as the ω site, and (2) a hydrophobic tail at the C terminus of the protein (Eisenhaber et al., 1998, 2003). In the ER lumen, the transamidase cleaves the pro-protein (from which the SP has been removed) at the ω site, removes the hydrophobic tail, and covalently links the remainder of the pro-protein to a GPI anchor, which is synthesized in the ER. Subsequently, the mature protein with the GPI-anchor traffics through the endomembrane system to the cell surface, where it associates with the plasma membrane using the GPI anchor (Varma and Mayor, 1998; Mayor and Riezman, 2004).

Consistent with the prediction that the GPI anchors LRE to the membrane, transiently expressed GFP-LRE localized to the surface of Arabidopsis mesophyll protoplasts (Capron et al., 2008) and citrine yellow fluorescent protein (cYFP)-LRE localized to the FA (Lindner et al., 2015). However, whether the predicted GPI anchor addition domains in LRE are important for its function in pollen tube reception has not been experimentally tested. LRE interacts with an extracellular domain of FER in yeast two-hybrid and pull-down assays, and FER-GFP localization in the FA is affected in *lre* mutants (Li et al., 2015). These results and characterization of LLG1, a putative paralog of LRE, indicate that LRE/LLG1 interacts with FER in the ER lumen and functions as a chaperone to bring FER to the FA (Li et al., 2015). Once localized to the FA, LRE could function as a coreceptor with FER to perceive signals from the pollen tube to trigger calcium profile changes and ROS production (Denninger et al., 2014; Duan et al., 2014; Ngo et al., 2014). However, the amino acid residues in LRE that are important for it to function in pollen tube reception are not yet clear. Additionally, whether LRE has a separate function at the synergid cell surface, which is independent of its intracellular function in the synergids, remains to be established.

We hypothesized that LRE localizes to the synergid cell surface using a GPI anchor and regulates pollen tube reception. Our analysis showed that the predicted GPI anchor addition domains in LRE are critical for its localization in the FA but not its function. We discovered that a unique 12-amino acid domain between the 5th and the 6th cysteines, including the highly conserved Asn-Asp dipeptide, is critical for LRE function in pollen tube reception. Finally, non-cell-autonomous and extracellular complementation of *lre* synergids with pollen tube-expressed *LRE* demonstrated that LRE and FER have a synergid cell surface-specific function in pollen tube reception.

RESULTS

LRE Encodes a Cysteine-Rich Protein with Domains Implicated in GPI Anchor Addition

LRE encodes a putative GPI-anchored surface protein (Capron et al., 2008; Tsukamoto et al., 2010), as it has the domains predicted to be required for GPI anchor addition (ω -site region and GAS domain; Figure 1; Supplemental Figure 1). Additional protein sequence analysis revealed that *LRE* encodes a cysteine-rich protein (CRP) with eight cysteines in the ectodomain, similar to proteins containing an eight-cysteine motif (8CM; José-Estanyol et al., 2004). The cysteines in an 8CM protein form disulfide bridges to maintain the tertiary structure of a scaffold containing conserved helical regions connected by variable loops, which confer functional specificity to the protein (José-Estanyol et al., 2004). In a typical 8CM, one amino acid separates the 5th and the 6th cysteines; by contrast, in *LRE*, a unique 12-amino acid domain separates them (Figure 1; Supplemental Figure 1). Sequence alignment showed conservation of spacing of the eight cysteines and the 12-amino acid domain between the 5th and the 6th cysteines (Supplemental Figure 1). Interestingly, this 12-amino acid domain is relatively variable with a highly conserved Asn-Asp dipeptide (Figure 1; Supplemental Figure 1), suggesting that the

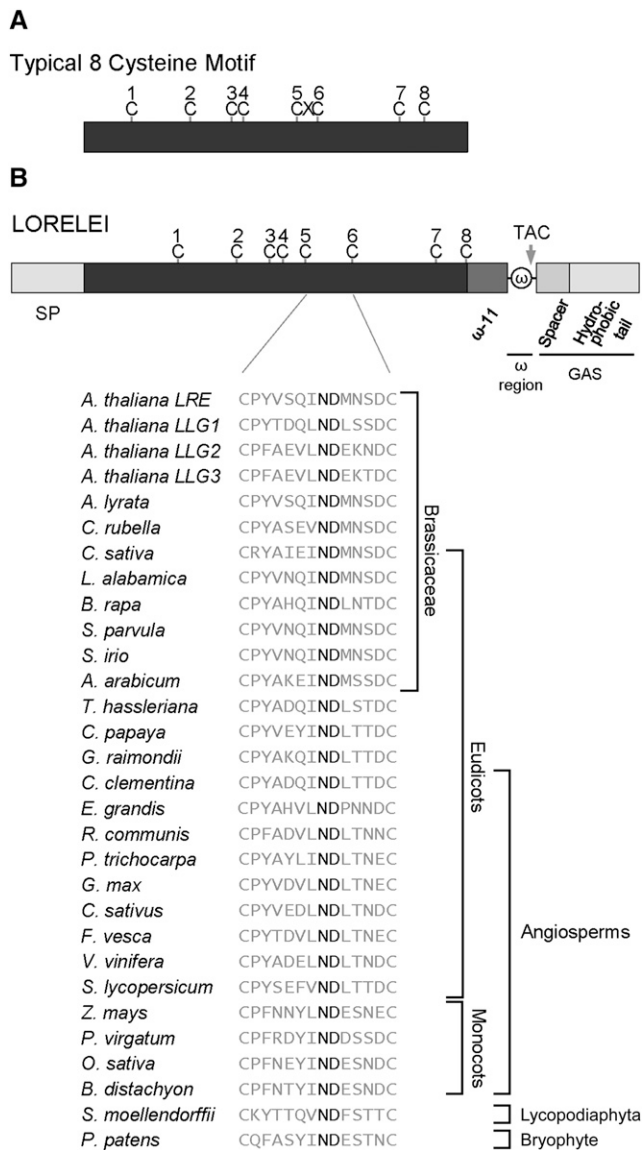


Figure 1. LRE Encodes a Putative GPI-Anchored Surface Protein with a M8CM.

(A) Diagram of a typical 8CM, in which the 3rd and 4th cysteines are adjacent to each other and the 5th and 6th cysteines are separated by one amino acid. C, cysteine; X, any other amino acid.

(B) Diagram of the LRE pre-pro protein. Predicted domains critical for GPI anchor addition to the pro-protein and positions of eight cysteines in LRE are indicated. A unique domain of 12-amino acids between the 5th and the 6th cysteines in LRE from Arabidopsis and in proteins from indicated plants that share the highest sequence similarity with full-length Arabidopsis LRE are shown. The highly conserved Asn-Asp dipeptide (ND) in this 12-amino acid domain is shown in bold. Multiple sequence alignment of full-length protein sequences shown here is presented in Supplemental Figure 1. ω , omega site; GAS, GPI attachment signal with a spacer region (darker gray) and a hydrophobic tail (lighter gray). Arrow points to the omega site cleaved by the transamidase complex (TAC).

Asn-Asp dipeptide is critical for LRE function. Thus, LRE encodes a putative GPI-anchored surface protein with a modified 8CM (M8CM).

Polar Localization of LRE-cYFP in the Filiform Apparatus of Synergids

To determine if the predicted GPI anchor addition domains and the M8CM in LRE are necessary for its function, we generated LRE-cYFP fusions for structure-function analysis. We used cYFP as the reporter because it is pH insensitive (Griesbeck et al., 2001) and can fluoresce even in the acidic apoplast (Gjetting et al., 2012), where the ectodomain of GPI-anchored surface proteins is expected to localize (Schultz et al., 1998). Additionally, cYFP was used to determine plasma membrane localization of other Arabidopsis GPI-anchored surface proteins (Simpson et al., 2009).

We generated N-terminal (*ProLRE:cYFP-LRE*) and C-terminal (*ProLRE:LRE-cYFP*) constructs (Figure 2A; Supplemental Figure 2A), determined the subcellular localization of LRE-reporter fusion proteins in the synergids, and verified that they retained LRE function. Although *ProLRE:cYFP-LRE* (N-terminal fusion construct) complemented the reduced seed set defect in *lre-7/lre-7* plants (Supplemental Figure 2D), cYFP expression was not detected in unfertilized ovules of all 14 primary transformants (T1) analyzed, and these were not characterized further. However, T1 plants carrying *ProLRE:LRE-cYFP* (C-terminal fusion construct) expressed cYFP in synergids (Supplemental Figures 3A, 3B, and 3I; also see below) and complemented the reduced seed set defect in *lre-7/lre-7* plants (Supplemental Figure 2D). We isolated three single-insertion lines in the T2 generation (Supplemental Table 1), identified plants that are homozygous for the transgene, and used these to analyze the subcellular localization and function of LRE-cYFP in greater detail; this approach was also undertaken for other constructs reported here (see below).

Confocal laser scanning microscopy showed that LRE-cYFP localized at the micropylar end of synergids (Figures 2B to 2F), where the FA is located (Huang and Russell, 1992; Kasahara et al., 2005). Indeed, the FA region contained nearly 56% of the total cYFP signal in synergids (Figures 2E and 2F). This is in contrast to EGFP-ROP6C, a membrane protein that localizes both in the FA and outside the FA, throughout the synergid plasma membrane (Escobar-Restrepo et al., 2007). Polar localization of LRE-cYFP in the FA is consistent with LRE being a putative GPI-anchored surface protein that functions in pollen tube reception (Capron et al., 2008; Tsukamoto et al., 2010). Besides its polarized localization in the FA, we consistently detected LRE-cYFP fluorescence in puncta in the synergid cell cytoplasm (Figures 2B to 2F). To explore the intracellular localization of LRE-cYFP, we performed colocalization experiments with an organelle marker (Golgi, ER, or peroxisome; Nelson et al., 2007) expressed under the control of the LRE promoter (Supplemental Figure 4A). None of these organelle markers colocalized with LRE-cYFP puncta in the synergids (Supplemental Figures 4B to 4M), and the identity of intracellular LRE-cYFP puncta remains unknown.

Upon pollen tube arrival in the female gametophyte, NTA-GFP relocates from the fluorescent puncta in the cytoplasm to the FA of synergids (Kessler et al., 2010). To test if LRE-cYFP localization also changes upon pollen tube arrival, *ProLRE:LRE-cYFP* pistils

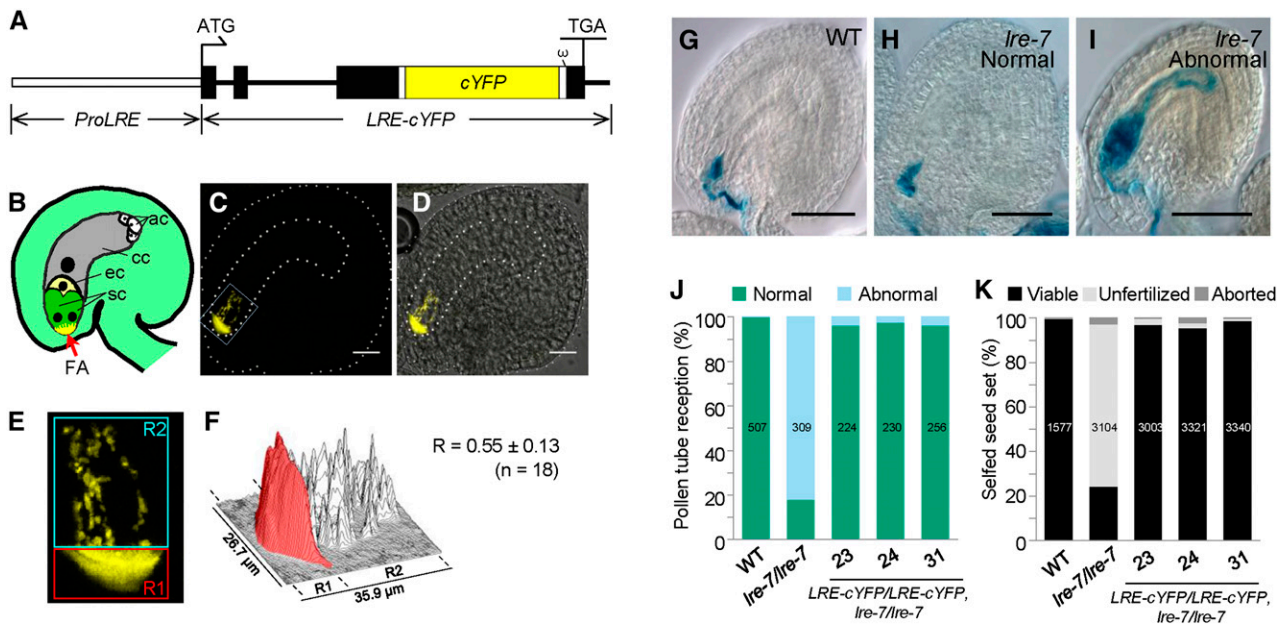


Figure 2. LRE-cYFP Shows a Polarized Localization in the Filiform Apparatus and Is Fully Functional in Pollen Tube Reception.

(A) Diagram of *ProLRE:LRE-cYFP* in which the cYFP is placed toward the C-terminal portion of the LRE protein and the *LRE-cYFP* is expressed from the *LRE* promoter (*ProLRE*, open rectangle). Filled black rectangles and black lines refer to exons and introns, respectively, in the *LRE* gene. ω , omega site; ATG and TGA, the start and stop codons, respectively, in *LRE*. The small white rectangle that appears before and after cYFP points to 9-amino acid and 11-amino acid linkers, respectively.

(B) Diagram of an unfertilized Arabidopsis ovule containing a mature female gametophyte. sc, synergid cell; ec, egg cell; cc, central cell; ac, antipodal cell.

(C) A representative fluorescent image showing LRE-cYFP localization in an ovule. The female gametophyte and the ovule are outlined with thick and thin white dashed lines, respectively. Bar = 20 μ m.

(D) A merged image of fluorescent image shown in (C) and a bright-field image (data not shown) of the same ovule. Bar = 20 μ m.

(E) A close-up view of the portion of the ovule within the white rectangle in (C). cYFP signal in the region of the filiform apparatus (red box) and the remainder of the synergids (cyan box) were quantified. R1 and R2, regions of interest 1 and 2, respectively.

(F) Surface plot of quantification of cYFP signal intensity within the boxed areas in (E). cYFP signal intensity in the filiform apparatus region is highlighted in red. R (mean \pm sd) indicates the proportion of cYFP signal intensity in the filiform apparatus relative to the total cYFP signal intensity in the synergids [$R = R1/(R1 + R2)$]. n, number of images, from three independent transformants (J) and (K), used to calculate R.

(G) to (I) Representative images showing pollen tube reception in ovules of indicated genotype when crossed to *pLAT52:GUS* pollen. This GUS staining-based assay was used to score pollen tube reception (Figures 2J, 3F, 4F, and 5D). Bar = 50 μ m.

(G) Normal pollen tube reception in a wild-type ovule.

(H) Normal pollen tube reception in an *Ire-7* ovule.

(I) Abnormal pollen tube reception (pollen tube coiling due to overgrowth) in an *Ire-7* ovule.

(J) and (K) The *ProLRE:LRE-cYFP* construct fully complements the pollen tube reception defect (J) and the reduced seed set defect (K) in *Ire-7/Ire-7* plants. Total number of ovules (J) or seeds (K) analyzed are in the center of each column. Number below each column, three lines used that are single insertion and homozygous for the *ProLRE:LRE-cYFP* transgene in the *Ire-7/Ire-7* background.

were pollinated with pollen carrying *ProLAT52:DsRed*. The localization of LRE-cYFP did not change in the FA after pollen tube reception (Supplemental Figures 3C to 3H and 3J).

LRE-cYFP Functions in Pollen Tube Reception

To test if LRE-cYFP is functional, we performed four experiments. Using a GUS staining-based assay (Tsukamoto et al., 2010), we showed that LRE-cYFP fully complemented the pollen tube reception defect in the *Ire-7* female gametophytes (Figures 2G to 2J). Additionally, the seed set in *Ire-7/Ire-7* was also fully restored to wild-type levels (Figure 2K), indicating that LRE-cYFP is functional.

If LRE-cYFP is functional, it should also rescue reduced transmission of the *Ire* mutation through the female gametophyte (Capron et al., 2008; Tsukamoto et al., 2010). Indeed, in lines containing *ProLRE:LRE-cYFP*, the transmission efficiency of the *Ire-7* mutation increased from 0.11 (11%) to nearly 1 (~100% transmission; Table 1). Additionally, when the transgenic plant was the female parent, the progeny of the cross revealed an increased transmission of *ProLRE:LRE-cYFP*, showing that LRE-cYFP complemented the defects in the *Ire-7* female gametophyte (Supplemental Table 1); no such increase in transmission was observed when the transgenic plant was the male parent (Supplemental Table 1). Taken together, our results demonstrated that LRE-cYFP is functional.

Table 1. *ProLRE:LRE-cYFP* Restored the Reduction in Transmission of the *lre-7* Mutation through the Female Gametophyte

Female Parent ^a	Male Parent ^a	Observed No. of Progeny		TE (R/S) ^b	χ^2 ^c	P Value
		Basta Rd	Basta ^{Sd}			
Wild type	<i>lre-7/+</i>	180	164	1.10	0.740	0.388
<i>lre-7/+</i>	Wild type	22	194	0.11	137.000	<0.001
Line 23	Wild type	141	140	1.01 ^e	0.004	0.952
Wild type	Line 23	138	131	1.05	0.180	0.670

^aLine numbers refer to a *ProLRE:LRE-cYFP* transformant in the *lre-7/lre-7* background containing a single insertion of the *ProLRE:LRE-cYFP* transgene. Genotype of the transgenic line used is homozygous for the transgene (*ProLRE:LRE-cYFP/ProLRE:LRE-cYFP*) and heterozygous for the *lre-7* mutation (*lre-7/+*).

^bTransmission efficiency (TE) was calculated as the ratio of Basta resistance (R) to susceptibility (S) in the progeny of the indicated cross.

^c χ^2 was calculated based on the expectation of a 1:1 segregation of Basta resistance to susceptibility in the progeny of a cross between the wild type and *lre-7/+*.

^dBasta-resistant (Basta^R) and Basta-susceptible (Basta^S) progeny. Basta resistance gene is linked with the T-DNA that is inserted into the *LRE* gene in *lre-7* mutant.

^eWhen segregation ratio in row three was compared to that in row two as the expected segregation ratio, the $\chi^2 = 491.31$ and P value < 0.001.

The Predicted Signal Peptide Is Essential for LRE-cYFP Expression in the Synergids

To test the importance of the predicted SP in LRE, we generated 12 T1 transformants with *ProLRE:LRE Δ SP-cYFP* (Supplemental Figure 2B); none showed cYFP signal in unfertilized ovules. RT-PCR on five randomly chosen lines showed that *ProLRE:LRE Δ SP-cYFP* was transcribed in unpollinated pistils (Supplemental Figure 2C). The lack of an LRE Δ SP-cYFP signal may be due to degradation of mislocalized LRE Δ SP-cYFP pro-protein, containing the C-terminal hydrophobic tail, in the cytoplasm (Hessa et al., 2011). Consistent with the lack of expression, *ProLRE:LRE Δ SP-cYFP* did not complement the seed set defect in *lre-7/lre-7* plants (Supplemental Figure 2D), showing that the predicted SP in LRE is essential for its expression in the synergids.

Loss of the GPI Anchor Addition Domains in LRE Disrupts LRE-cYFP Localization in the Filiform Apparatus

A prediction algorithm for plant GPI-anchored surface proteins, Big-PI plant predictor (Eisenhaber et al., 2003), identified Ser-139 as the ω site in LRE. Loss of the ω site and the resulting failure to receive a GPI anchor could disrupt LRE-cYFP localization in the plasma membrane-rich FA. However, synergid cell localization of LRE-cYFP $\Delta\omega$ (Figure 3A), in which Ser-139 was deleted, was indistinguishable from LRE-cYFP localization (compare Figure 3B with Figures 2C to 2F). One possible interpretation of these results is that a cryptic ω site in LRE may function in the absence of Ser-139. Indeed, in an LRE sequence lacking Ser-139, Big-PI plant predictor identified Ala-141 as the ω site, and it identified no other ω site in an LRE sequence that lacked both Ser-139 and Ala-141. When both Ser-139 and Ala-141 were deleted (Figure 3A), LRE-cYFP $\Delta 2\omega$ localization in the FA was dramatically lower compared with LRE-cYFP (cYFP signal intensity decreased from ~56 to ~17%; Figure 3C). Additionally, there was a notable increase in the diffuse cYFP signal in the synergid cell cytoplasm. These results raise the possibility that removal of the predicted ω sites led to defects in GPI anchor addition and disrupted the trafficking of LRE-cYFP $\Delta 2\omega$ to the FA, consistent with studies showing that GPI

attachment is required for transport of GPI-anchored proteins from the ER to the cell surface (Doering and Schekman, 1996; Mao et al., 2003).

If loss of ω sites affects LRE-cYFP localization in the FA, then deletion of the GAS domain (Ser-143–Thr-165) should also produce LRE-cYFP without a GPI anchor (Mao et al., 2003) that fails to accumulate in the FA. Indeed, we observed a dramatic reduction in FA localization of LRE-cYFP Δ GAS (cYFP signal intensity decreased from ~56 to ~16%; Figure 3D). Additionally, the absence of the GPI anchor addition domains was associated with an enhanced diffuse signal in the cytoplasm (Figure 3D), which may be due to inefficient protein sorting and/or enhanced retention of these proteins in the ER (Doering and Schekman, 1996). These results were similar to LRE-cYFP $\Delta 2\omega$ localization and consistent with the model that LRE localization in the FA requires a GPI anchor. Unlike in LRE-cYFP and LRE-cYFP $\Delta 2\omega$, in 4/4 lines analyzed, LRE-cYFP Δ GAS was seen in the intercellular spaces of the integument cells surrounding the micropyle, and the number of ovules showing this extracellular accumulation increased over time (Supplemental Figure 5), indicating that LRE-cYFP Δ GAS is released outside the female gametophyte. These results are consistent with other GPI-anchored surface proteins lacking the GAS domain, which were released from the cell into the extracellular space at enhanced levels (Mao et al., 2003).

LRE-cYFP Localization in the Filiform Apparatus Is Disrupted in a Mutant Defective in GPI8, a Putative Subunit of the Transamidase Involved in GPI Anchor Addition

To further test if the GPI anchor addition domains in LRE are important for its localization in the FA, we examined LRE-cYFP localization in a mutant defective in GPI8, a critical subunit of the transamidase, which catalyzes the endoproteolysis reaction that removes the GAS from the pro-protein before covalently attaching the GPI anchor to the ω site. We reasoned that if GPI anchor addition domains were important for localization of LRE-cYFP in the FA, then the transamidase that uses these domains to catalyze GPI anchor addition would also be critical for the FA localization of LRE-cYFP. Based on this, it could be expected that the FA

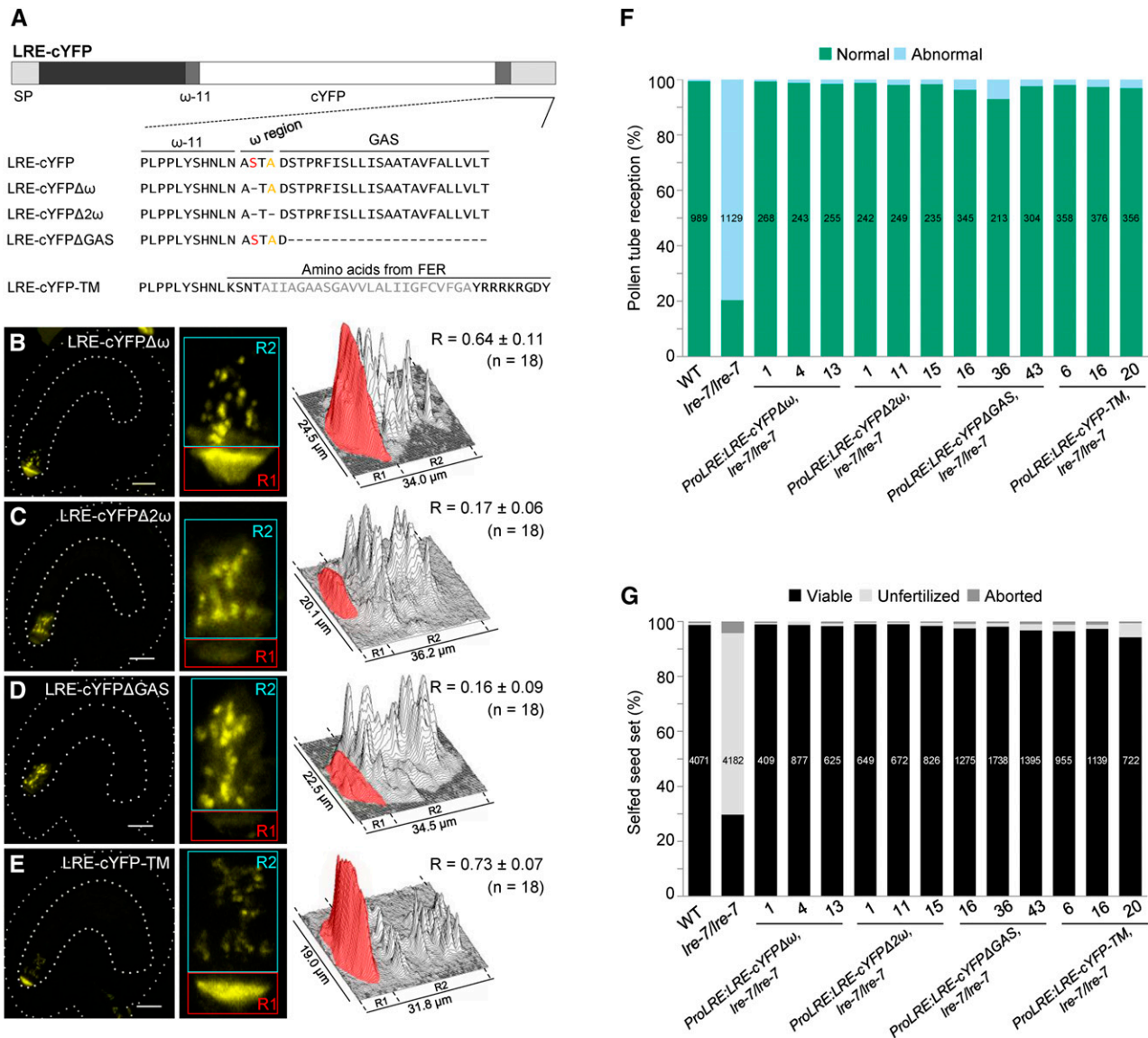


Figure 3. Predicted GPI Anchor Addition Domains Are Necessary for LRE Localization in the Filiform Apparatus but Not Its Function in Pollen Tube Reception.

(A) Diagram of the LRE-cYFP protein. Wild-type or altered amino acid sequence on either side of the predicted ω amino acid in each construct is indicated below the diagram. In LRE, the best predicted ω and a cryptic ω site are labeled in red and orange, respectively. Each dash in the protein sequence represents a deletion of the corresponding amino acid in wild-type LRE protein sequence. GAS, GPI attachment signal (lighter gray rectangle). In LRE-cYFP-TM, predicted transmembrane region of FER is in gray.

(B) to (E) Localization of LRE-cYFP $\Delta\omega$ **(B)**, LRE-cYFP $\Delta 2\omega$ **(C)**, LRE-cYFP Δ GAS **(D)**, and LRE-cYFP-TM **(E)** in the synergids of the female gametophyte. Left panels, representative image showing the localization of each fusion protein; middle panels, close-up view of the micropylar region in corresponding left panel images; right panels, surface plots showing quantification of cYFP signal in the region of filiform apparatus (red box) and the remainder of the synergids (cyan box) as in Figure 2F. Bar = 20 μ m.

(F) and **(G)** Constructs with altered GPI addition domains fully complement pollen tube reception defect **(F)** and the reduced seed set defect **(G)** in *lre-7/lre-7* plants. Total number of ovules **(F)** and seeds **(G)** analyzed are in the center of each column. Number below each column, the three lines used that are single insertion and homozygous for the indicated transgenes in the *lre-7/lre-7* background.

localization of LRE-cYFP would be affected in a *gpi8* mutant female gametophyte, as the GPI anchor will not be added to LRE-cYFP.

In yeast (*Schizosaccharomyces pombe*) and human cells, *gpi8* mutants are defective in GPI attachment and cause a decrease in cell surface display of GPI-anchored proteins (Benghezal et al., 1996; Chen et al., 1996; Yu et al., 1997). The Arabidopsis genome contains a single gene (*GPI8*) that encodes a protein with high sequence similarity to yeast (43.8% identity) and human (45.7% identity) GPI8 and contains the postulated protease catalytic site that is conserved in C13 clade of cysteine proteases (Supplemental Figures 6A and 6B) (Zacks and Garg, 2006). The expression of SKU5, a known GPI-anchored surface protein (Sedbrook et al., 2002), is affected in *gpi8-1*, a weaker point mutant allele of *GPI8* that was identified as an enhancer of stomatal clustering in an *erecta-like1 erecta-like2* double mutant; this decrease in expression is likely due to the failure in removing the GAS from SKU5 (Bundy et al., 2016). Many GPI-anchored proteins play essential roles in the male gametophyte (Lalanne et al., 2004). Consistent with this, transmission of *gpi8-2*, a stronger T-DNA allele of *GPI8* (Supplemental Figure 6C), through the male gametophyte is abolished (Supplemental Table 2); however, *gpi8-2* is transmitted through the female gametophyte (Supplemental Table 2). This result, combined with the finding that *GPI8* is expressed in the synergids (Wuest et al., 2010), makes it feasible to test the effect of *gpi8-2* on LRE-cYFP localization in the FA.

We crossed *ProLRE:LRE-cYFP* homozygous plants with the *gpi8-2/+* mutants and analyzed LRE-cYFP localization in the F1 progeny, which were either *+/+* or *gpi8-2/+* (Supplemental Table 3 and Supplemental Figure 6D). In all the progeny that were wild type for *GPI8*, cYFP-positive ovules showed only one pattern of localization: in fluorescent puncta and a polarized localization in the FA, as expected for LRE-cYFP. However, in every *gpi8-2/+* progeny analyzed, the cYFP-positive ovules showed two localization patterns: about half of the cYFP-positive ovules had polarized LRE-cYFP localization and in the other half, the cYFP signal was diffuse and dramatically reduced in the FA (from ~60 to ~13%; Figures 4A to 4E; Supplemental Table 3), a localization pattern similar to LRE-cYFP Δ 2 ω and LRE-cYFP Δ GAS (Figures 3C and 3D).

To test if the effect of *gpi8-2* is specific to proteins with GPI anchor addition domains, such as LRE-cYFP, we generated *gpi8-2/+* plants expressing a variant of LRE-cYFP, in which a single-pass transmembrane (TM) domain of FER, an integral membrane protein, replaced the GPI anchor addition domains (Figure 3A). LRE-cYFP-TM is expected to be unaffected in *gpi8-2* mutants, as TM-containing proteins do not require the transamidase for their localization to the cell surface. Indeed, all cYFP-positive ovules in *gpi8-2/+* pistils carrying *ProLRE:LRE-cYFP-TM* showed only one localization pattern: a preferential accumulation of cYFP in the FA (Supplemental Figure 7 and Supplemental Table 4), compared with the two localization patterns of *ProLRE:LRE-cYFP* in *gpi8-2/+* pistils (Figures 4A to 4E; Supplemental Table 3). These results are consistent with our finding that GPI anchor addition domains in LRE play a role in localizing LRE-cYFP to the FA (Figure 3).

The GPI Anchor Addition Domains in LRE Are Not Necessary for Its Function in Pollen Tube Reception

If LRE localization in the FA is affected by the loss of transamidase, *gpi8-2* and *lre* female gametophytes may show the same defects in pollen tube reception and seed set. Contrary to this expectation, *gpi8-2* mutants showed wild-type pollen tube reception and seed set (Figures 4F and 4G). Transmission of the *gpi8-2* mutation through female gametophytes was also not affected (Supplemental Table 2), suggesting that LRE function does not require the GPI anchor addition domains.

The lack of pollen tube reception and seed set phenotypes in *gpi8-2/+* plants prompted us to perform complementation tests with LRE-cYFP Δ 2 ω and LRE-cYFP Δ GAS to investigate if LRE function requires the GPI anchor addition domains. Despite dramatically reduced localization in the FA (Figures 3C and 3D), LRE-cYFP Δ 2 ω and LRE-cYFP Δ GAS almost fully complemented the pollen tube reception and seed set defects in *lre-7/lre-7* plants (Figures 3F and 3G). Transmission of either *ProLRE:LRE-cYFP Δ 2 ω* or *ProLRE:LRE-cYFP Δ GAS* through the *lre-7* female gametophyte (Supplemental Table 5) was also comparable to the transmission of *ProLRE:LRE-cYFP* through the *lre-7* female gametophyte (Supplemental Table 1), indicating that LRE can induce pollen tube reception despite lacking the potential ω sites or GAS domain and suggesting that LRE function does not require GPI anchor addition domains.

An LRE Variant Lacking the GPI Anchor Addition Domains but Attached to a Transmembrane Domain Can Function in Pollen Tube Reception

To further test if the GPI anchor addition domains in LRE are necessary for its function in pollen tube reception, we tested if LRE-cYFP-TM is functional in pollen tube reception. We chose LRE-cYFP-TM because in this transgenic protein, both the ω site and the GAS domain of LRE have been replaced with a single-pass TM domain (Figure 3A). Additionally, it localized to the FA of the female gametophyte (Figure 3E) and is expected to produce an LRE variant that remains tethered to the plasma membrane via the TM.

LRE-cYFP-TM almost fully rescued the pollen tube reception and seed set defects in *lre-7/lre-7* plants (Figures 3F and 3G; Supplemental Figure 8). Transmission of *ProLRE:LRE-cYFP-TM* through the *lre-7* female gametophyte was also enhanced, confirming that LRE-cYFP-TM is functional (Supplemental Table 5). These results indicate that even if it lacks GPI anchor addition domains but is attached to a TM domain, LRE can still function in pollen tube reception. Complementation by LRE-cYFP-TM and by LRE variants that lack the GPI anchor addition domains (LRE-cYFP Δ 2 ω and LRE-cYFP Δ GAS) indicates that a domain shared by these three constructs, such as the M8CM in the ectodomain of LRE, might be important for its function in pollen tube reception.

LRE Function in Pollen Tube Reception Requires a Unique 12-Amino Acid Domain in the M8CM

To identify which domain within the LRE ectodomain is critical for its function in pollen tube reception, we tested if loss of the

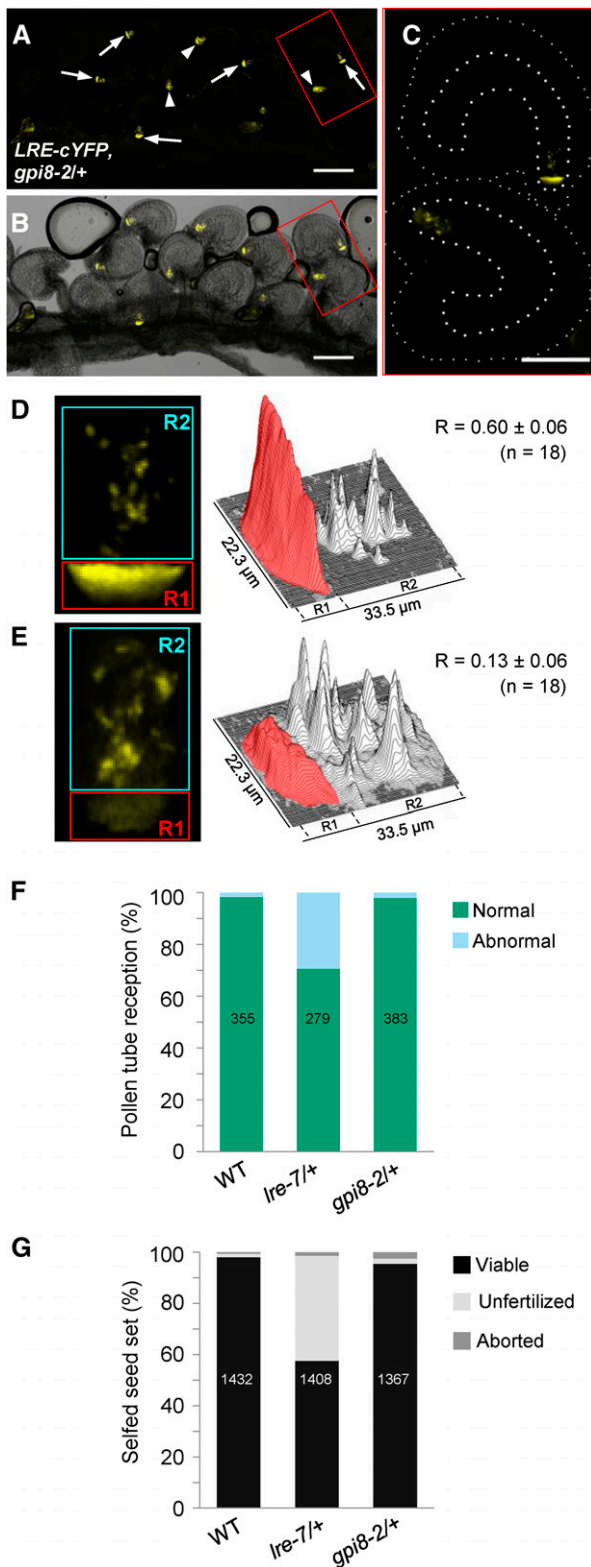


Figure 4. LRE-cYFP Localization in the Filiform Apparatus Is Affected by a Mutation in *GPI8*, a Critical Component of the Transamidase.

12-amino acid domain in the M8CM of the ectodomain, uniquely found in LRE family proteins (Figure 1; Supplemental Figure 1), affects LRE function. We produced an LRE with a typical 8CM by replacing the 12-amino acid domain between the 5th and the 6th cysteines in LRE with a single Leu (Figure 5A), the most prevalent amino acid in Arabidopsis proteins containing a typical 8CM (61 out of 98 proteins; José-Estanyol et al., 2004). LRE_(CLC)-cYFP signal intensity in the FA was ~ 4.6 times lower than that of LRE-cYFP (from ~ 56 to $\sim 12\%$; compare Figure 5B with Figures 2C to 2F). LRE_(CLC)-cYFP did not rescue pollen tube reception, seed set, or transmission defects in *ire-7/ire-7* plants (Figures 5D and 5E; Supplemental Figure 8 and Supplemental Table 6). These results demonstrated that the 12-amino acid domain between the 5th and the 6th cysteines in LRE is necessary for its localization in the FA and its function in pollen tube reception.

To investigate if the highly conserved Asn-Asp dipeptide in the 12-amino acid domain (Figure 1; Supplemental Figure 1) is necessary for LRE localization and function, we replaced the Asn-Asp with two alanines in LRE-cYFP (Figure 5A). Unlike LRE_(CLC)-cYFP, LRE_(ND88AA)-cYFP localized to the FA normally in unfertilized ovules (Figure 5C), indicating that loss of Asn-Asp does not affect sub-cellular localization of LRE. However, LRE_(ND88AA)-cYFP is not functional, as evident from pollen tube reception and seed count assays and transmission analysis (Figures 5D and 5E; Supplemental Figure 8 and Supplemental Table 6). These results showed that loss of the Asn-Asp dipeptide abolished LRE function in pollen tube reception. Taken together, these results show that the unique 12-amino acid domain in the M8CM of LRE is important for LRE function in pollen tube reception.

Pollen Tube-Expressed LRE Complements the Pollen Tube Reception Defect in the *ire* Female Gametophyte

Intracellular and synergid cell surface (FA) localization of LRE-cYFP raises the question of where LRE functions in pollen tube reception. As the first point of contact between the pollen tube and the synergid cell, the FA may be the critical location for LRE function. Alternatively, intracellular LRE may be critical for pollen tube reception for the following reasons: First, in our study, the levels of the two fusion proteins that successfully complemented *ire-7* (LRE-cYFP $\Delta 2\omega$ and LRE-cYFP Δ GAS) were dramatically reduced in the FA, but they did localize intracellularly (Figures 3D

(A) and **(B)** Localization of the LRE-cYFP fusion protein in a *gpi8-2/+* pistil. White arrows, female gametophytes with a polarized cYFP localization in the filiform apparatus; white arrowheads, sibling female gametophytes with a diffuse cYFP localization throughout the synergids. Bar = 100 μ m. **(C)** Enlarged view of two ovules within the red box in **(A)** and **(B)**. Ovules and female gametophytes are outlined in thin and thick white dashed lines, respectively. Bar = 50 μ m.

(D) and **(E)** cYFP signal intensity quantification in ovules within a *gpi8-2/+* pistil. Left panels, representative images of the micropylar region of ovules with polarized **(D)** and diffuse **(E)** cYFP localization in the synergids of ovules within a *gpi8-2/+* pistil. cYFP signal in the region of filiform apparatus (red) and the remainder of the synergids (cyan) were quantified as in Figure 2F.

(F) and **(G)** *gpi8-2* mutation does not cause pollen tube reception **(F)** or seed set defect **(G)**.

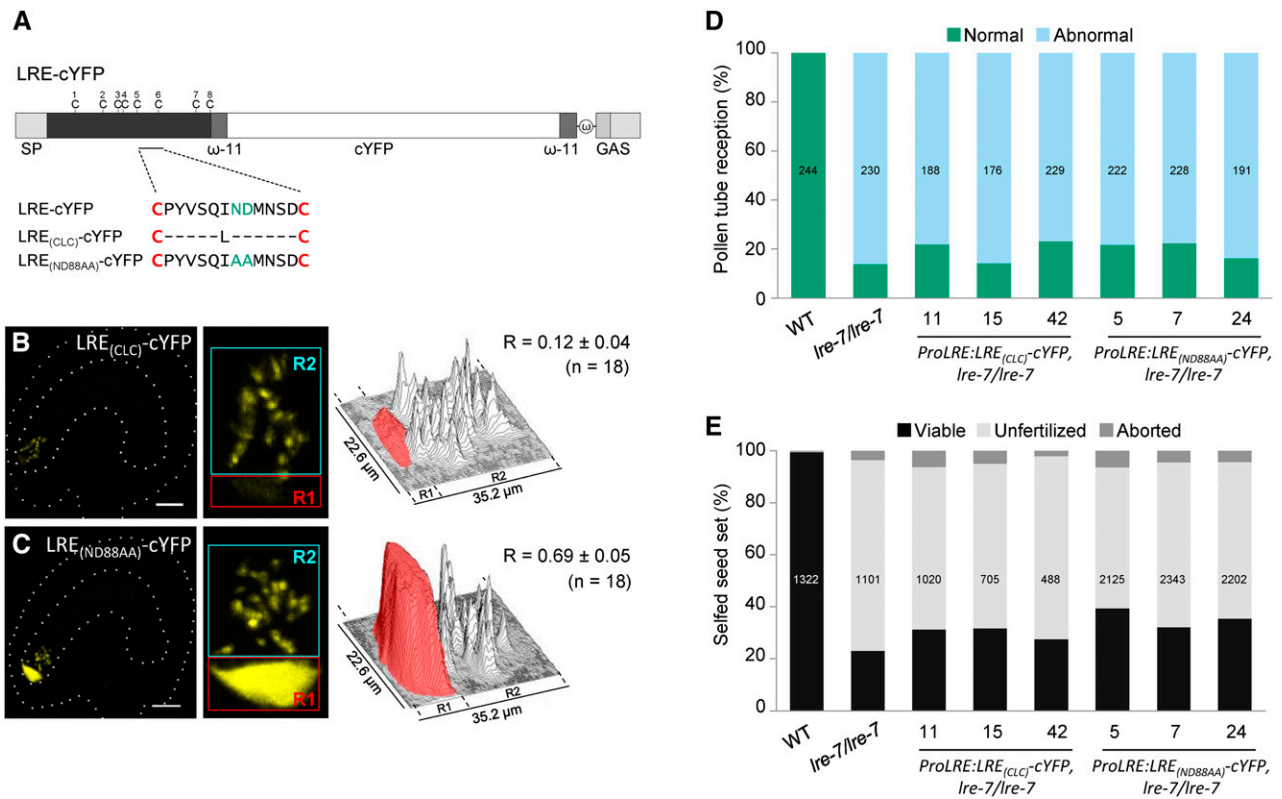


Figure 5. The 12-Amino Acid Residues between the 5th and 6th Cysteines of LRE and the Conserved Asn-Asp Dipeptide Are Necessary for LRE Function in Pollen Tube Reception.

(A) Diagram of the LRE-cYFP fusion protein. Amino acid sequence between the 5th and the 6th cysteines of LRE in each construct are indicated below the diagram. Each dash in the protein sequence represents a deletion of the corresponding amino acid in the wild-type LRE protein sequence. ω , omega site; GAS, GPI attachment signal containing a spacer region (darker gray rectangle) and a hydrophobic tail (lighter gray rectangle).

(B) and **(C)** LRE_(CLC)-cYFP **(B)** and LRE_(ND88AA)-cYFP **(C)** fusion proteins in the synergids of the female gametophyte. Left panels, representative image showing the localization of indicated fusion protein; middle panels, close-up view of the micropylar region in corresponding left panel images; right panels, surface plots showing quantification of cYFP signal in the region of filiform apparatus (red box) and the remainder of the synergids (cyan box) as in Figure 2F. Bar = 20 μ m.

(D) and **(E)** None of these three constructs complemented the pollen tube reception defect **(D)** or the reduced seed set **(E)** defect in *lre-7/lre-7* plants. Total number of ovules **(D)** and seeds **(E)** analyzed are in the center of each column. Number below each column, three lines used that are that are single insertion and homozygous for the indicated transgenes in the *lre-7/lre-7* background.

to 3F and 3G; Supplemental Table 5); second, a recent report showed that LLG1/LRE functions intracellularly as a chaperone to facilitate cell surface localization of the FER receptor kinase (Li et al., 2015). However, since the intracellular function of LLG1/LRE precedes its function in the cell surface, it remains unclear whether LRE has a separate function in the synergid cell surface.

To test these possibilities, we supplied LRE-cYFP extracellularly from the pollen tube, a cell where LRE is not normally expressed, and determined if it can complement the pollen tube reception defect in *lre* synergids. To avoid potential complications from ectopically expressed LRE-cYFP interfering with pollen development and germination, we expressed LRE-cYFP from the PECTIN LYASE-LIKE SUPERFAMILY PROTEIN3 (PLL3) promoter (Sun and van Nocker, 2010), which is preferentially induced in pollen tubes after their growth in the pistil (Qin et al., 2009) (Supplemental Figures 9A and 9B).

LRE may inhibit pollen tube growth by initiating a signal cascade in the pollen tube when it reaches the synergid cell, probably after

being released from the GPI anchor (Capron et al., 2008) upon cleavage by enzymes such as phospholipase C or D (Wang, 2001; Sharom and Lehto, 2002). Therefore, ectopically expressed LRE-cYFP may affect pollen tube growth. However, we observed normal transmission of *ProPLL3:LRE-cYFP* through the wild-type male gametophyte, when in competition with nontransgenic pollen, indicating that LRE-cYFP expression did not compromise in vivo pollen tube functions (Supplemental Table 7). We therefore used *ProPLL3:LRE-cYFP* to test if pollen tube-expressed LRE can induce pollen tube reception in *lre* synergids.

Indeed, we observed increased seed set in *lre-5* pistils pollinated with pollen from *ProPLL3:LRE-cYFP* plants (Supplemental Figure 9C), suggesting that pollen tube-expressed LRE-cYFP can function in pollen tube reception. To further characterize this non-cell-autonomous complementation, we isolated three single insertion lines (Supplemental Table 7) that were homozygous for *ProPLL3:LRE-cYFP*. An aniline blue staining-based

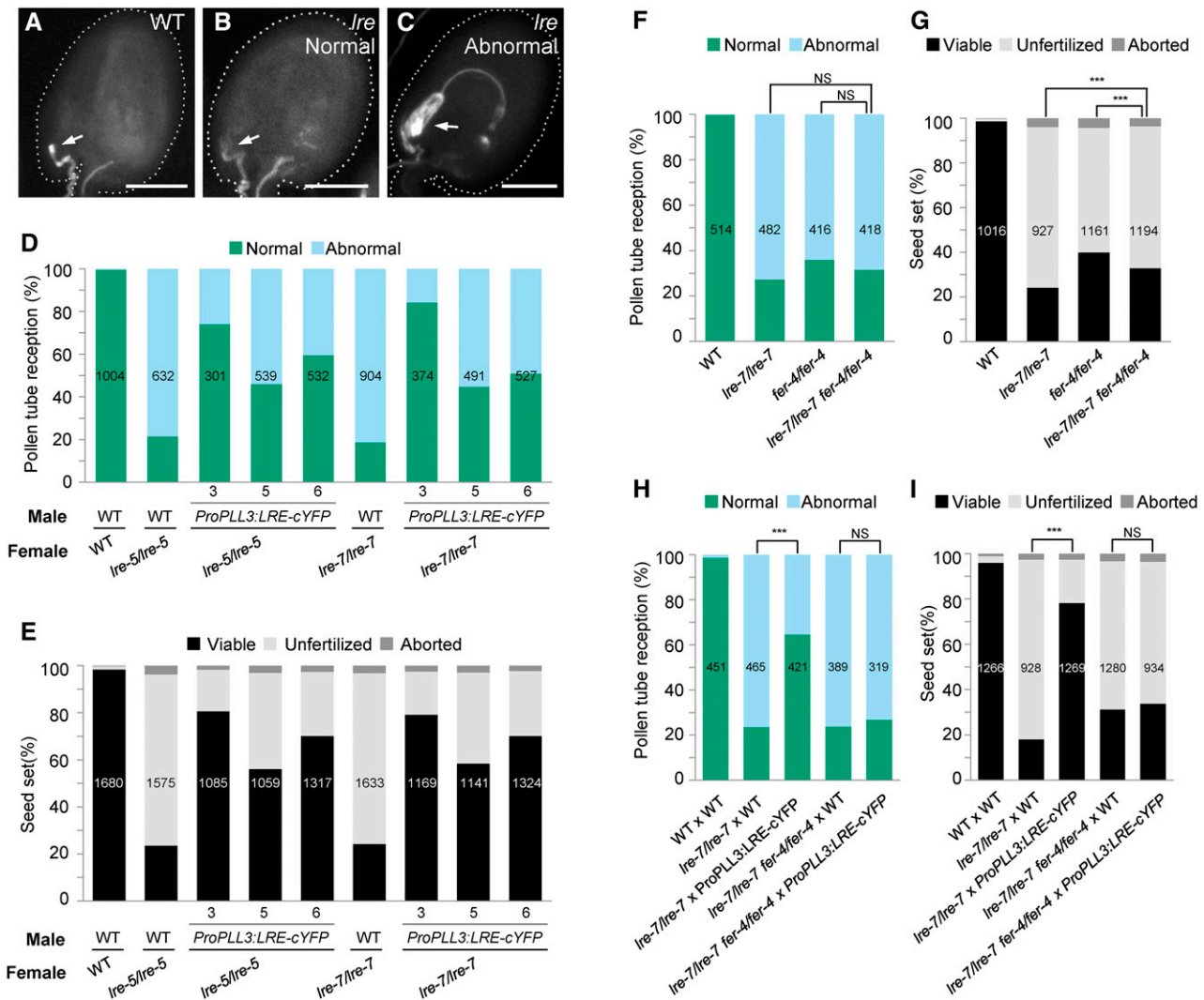


Figure 6. LRE-cYFP from Pollen Tubes Rescues Female Gametophyte Defects Caused by the *Ire-7* Mutation.

(A) to (C) Representative images from pollen tube reception assays using aniline blue staining. Ovules are outlined in white dashed line. Normal pollen tube reception in wild-type (A) and *ire-7/ire-7* ovules (B); abnormal pollen tube reception in *ire-7/ire-7* ovule (C). Bar = 50 μ m.

(D) and (E) Pollen tube-expressed LRE-cYFP rescued pollen tube reception (D) and seed set defects (E) of the *ire-7* female gametophyte, respectively. Total number of ovules (D) and seeds (E) analyzed, respectively, is in the center of each column.

(F) and (G) *ire-7/ire-7 fer-4/fer-4* double mutant showed similar level of pollen tube reception as *ire-7/ire-7* and *fer-4/fer-4* single mutant and a small but significant difference (χ^2 test) in seed set defect when compared with either *ire-7/ire-7* or *fer-4/fer-4* single mutant (G). Total number of ovules (F) and seeds (G) analyzed is in the center of each column. NS, not significant. P values = 0.192 (column 2 versus column 4 in [F]) and 0.1957 (column 3 versus column 4 in [F]), respectively. ***P value < 0.001 (P values = 6.34×10^{-5} for column 2 versus column 4 and 0.0007 for column 3 versus column 4 in [G]).

(H) and (I) *FER* is required in synergids to rescue the pollen tube reception (H) and seed set (I) defects in the *ire-7* female gametophyte by the pollen tube-expressed LRE. Total number of ovules (H) and seeds (I) analyzed is in the center of each column. NS, not significant. P value = 0.3994 (column 4 versus column 5 in [H]) and 0.4094 (column 4 versus column 5 in [I]) respectively. ***P value < 0.001; P value < 2.2×10^{-16} for column 2 versus column 3 in both (H) and (I).

pollen tube reception assay showed that substantially more *Ire* female gametophytes undergo pollen tube reception with *ProPLL3:LRE-cYFP* pollen compared with wild-type pollen (Figures 6A to 6D). Consistent with this, *Ire* pistils also showed improved seed set when pollen tubes expressing LRE-cYFP were used (Figure 6E). To further test whether pollen tubes expressing LRE-cYFP can rescue the *Ire* female gametophyte

defect, we examined the transmission of the *ire-5* and *ire-7* mutations through the female gametophyte, using homozygous *ProPLL3:LRE-cYFP* plants as the male parent. As expected, *ProPLL3:LRE-cYFP* pollen enhanced the transmission of *ire-5* and *ire-7* through the female gametophyte from 18% to 62 to 72% and 12% to 60 to 72%, respectively (Table 2). Consistent with these results, *ProPLL3:LRE-cYFP* is also transmitted at an enhanced

Table 2. *ProPLL3:LRE-cYFP* in the Male Gametophyte Restored the Reduction in Transmission of the *lre-7* Mutation through the Female Gametophyte

Female Parent	Male Parent ^a	Observed No. of Progeny		TE (R/S) ^b	χ^2 ^c	P Value
		Resistant ^d	Susceptible ^d			
Wild type	<i>lre-7/+</i>	405	397	1.02	–	–
<i>lre-7/+</i>	Wild type	100	542	0.18	–	–
<i>lre-7/+</i>	Line 3	150	215	0.70	180.77	<0.001
<i>lre-7/+</i>	Line 5	176	285	0.62	179.09	<0.001
<i>lre-7/+</i>	Line 6	226	316	0.72	281.24	<0.001
Wild type	<i>lre-5/+</i>	435	414	1.05	–	–
<i>lre-5/+</i>	Wild type	53	448	0.12	–	–
<i>lre-5/+</i>	Line 3	94	130	0.72	233.25	<0.001
<i>lre-5/+</i>	Line 5	203	328	0.62	429.17	<0.001
<i>lre-5/+</i>	Line 6	158	264	0.60	321.88	<0.001

^aLine numbers refer to three independent transformants in the wild-type background containing a single insertion of the *ProPLL3:LRE-cYFP* transgene; genotype of each transgenic line used is homozygous for the transgene (*ProPLL3:LRE-cYFP / ProPLL3:LRE-cYFP*).

^bTransmission efficiency (TE) was calculated as the ratio of Basta or kanamycin resistance (R) to susceptibility (S) in the progeny of the indicated cross.

^c χ^2 was calculated based on Basta or kanamycin resistance to susceptibility segregation ratio (R:S) in the progeny of the crosses *lre-7/+*(♀) × wild type(♂) or *lre-5/+*(♀) × wild type(♂), respectively, as the expected segregation ratio.

^dResistant and susceptible progeny to Basta or kanamycin. Basta and kanamycin resistance genes are linked to the T-DNA inserted in the *LRE* gene in *lre-7* and *lre-5*, respectively.

rate to the progeny if crossed to *lre-7* females compared with wild-type females (Supplemental Table 7). These results suggested that at the interface of the pollen tube and the synergid cell, LRE has a function that is independent of its intracellular function in the synergids.

In the synergid cells, complementing *lre-7* female gametophyte with LRE variants LRE-cYFP Δ 2 ω or LRE-cYFP Δ GAS revealed that the GPI anchor addition domains in LRE are not necessary for its function in pollen tube reception. To investigate if this is the case with pollen tube-expressed LRE, we ectopically expressed LRE-cYFP Δ 2 ω or LRE-cYFP Δ GAS in the pollen tube and examined if they are functional in pollen tube reception. Our results showed that pollen carrying *ProPLL3:LRE-cYFP Δ 2 ω* or *ProPLL3:LRE-cYFP Δ GAS* transgenes rescued the pollen tube reception defect in the *lre* female gametophyte to a similar extent as did *ProPLL3:LRE-cYFP* (Supplemental Figure 9C). These results showed that GPI anchor addition domains in LRE are not necessary for pollen tube-expressed LRE at the interface of the pollen tube and the synergid cell to induce pollen tube reception.

Alternatively, the pollen tube-expressed LRE-cYFP or LRE-cYFP Δ 2 ω or LRE-cYFP Δ GAS might be released from the pollen tubes and endocytosed into synergids and then carry out an intracellular function. To test this possibility, we used the *PLL3* promoter to express LRE-cYFP-TM (Supplemental Figures 9A and 9B), as it was almost fully functional when expressed in synergids (Figures 3A, 3F, and 3G) and is expected to remain tethered to the pollen tube plasma membrane. Although the YFP signal was primarily in the cytoplasm and there was only a weak plasma membrane localization in the pollen tubes, in all seven T1 transformants, seed set was restored when pollen tube-expressed LRE-cYFP-TM was pollinated onto *lre/lre* pistils (Supplemental Figure 9C). *ProPLL3:LRE-cYFP-TM* also was transmitted at an enhanced rate to the progeny if crossed to an *lre-7* female compared with the wild type (Supplemental Table 8),

confirming that LRE-cYFP-TM is functional when expressed in pollen tubes. These results are consistent with the model that LRE has a function at the interface of the pollen tube and the synergid cell, independent of its intracellular function in the synergids.

LRE and FER Require Each Other at the Interface of the Pollen Tube and the Synergid Cell to Induce Pollen Tube Reception

LRE acts as a chaperone for FER in the synergid cell and promotes its localization in the FA (Li et al., 2015). If FER localization in the FA is entirely dependent on LRE, then complementation of pollen tube reception defect in the *lre* female gametophyte by pollen tube-expressed LRE must depend on other protein(s) in the FA, besides FER, to induce pollen tube reception. We tested this possibility by examining if pollen tube-expressed LRE can complement the pollen tube reception defect in a female gametophyte that lacks both LRE and FER.

For this, we first established an *lre-7/lre-7 fer-4/fer-4* double homozygous mutant and assessed the seed set and pollen tube reception phenotypes in the double mutant pistils. There was no statistically significant difference in the pollen tube reception defect in the double mutant compared with either the *lre-7/lre-7* or *fer-4/fer-4* single mutant (Figure 6F). However, there was a small but statistically significant difference in seed set between each of the single mutants when compared with the *lre/lre fer/fer* double mutant (Figure 6G). Nevertheless, similar levels of pollen tube reception defect in single and double mutants provide genetic evidence in support of the proposal that, similar to its paralog LLG1, LRE functions with FER to induce pollen tube reception in the synergid cell (Li et al., 2015).

We next crossed *ProPLL3:LRE-cYFP* pollen onto the *lre-7/lre-7 fer-4/fer-4* double mutant pistil and found that pollen tube-expressed LRE-cYFP did not rescue the pollen tube reception

defect in the *lre-7 fer-4* double mutant female gametophyte (Figure 6H) and consequently also did not rescue the seed set defect in the *lre-7/lre-7 fer-4/fer-4* double mutant pistil (Figure 6I). Taken together, these results indicated that LRE-cYFP delivered by the pollen tube requires FER in the synergid cell to rescue the pollen tube reception defect in the *lre* female gametophyte and that LRE and FER require each other at the interface of the pollen tube and the synergid cell to induce pollen tube reception.

DISCUSSION

Non-Cell-Autonomous and Extracellular Complementation Analysis with Pollen Tube-Expressed LRE Clarified the Role of LRE in Pollen Tube Reception

Based on the excessive pollen tube growth phenotype in *lre* female gametophytes, we initially hypothesized that LRE participates in pollen tube reception by inhibiting/reducing pollen tube growth after it interacts with the synergids. This pause in pollen tube growth may then activate additional signaling between the pollen tube and the synergids to complete pollen tube reception. Near normal transmission efficiency of *ProPLL3:LRE-cYFP* in the wild-type background to the progeny through the male gametophyte, even in direct competition with nontransgenic pollen, showed that LRE-cYFP in pollen tubes did not affect any male gametophyte function (Supplemental Table 7; *ProPLL3:LRE-cYFP/+* crossed to the wild type). These results argue against the possibility that LRE by itself functions as a growth arrest signal in the pollen tube. Consistent with this, pollen tubes did not show premature growth arrest when interacting with ovules showing extracellular accumulation LRE-cYFP Δ GAS in the micropyle (Supplemental Figure 5).

Still, pollen tube-expressed LRE-cYFP complemented pollen tube reception defects in the *lre-7* female gametophyte (Figures 6D and 6E), showing that a pollen tube interacting with the synergid cell is necessary for LRE function. One way synergids may regulate LRE function is by possessing a mechanism to release the LRE ectodomain from the GPI anchor and activate it, akin to what is known in other organisms (Müller and Bandlow, 1993; Wang, 2001; Sharom and Lehto, 2002; Capron et al., 2008; Haruta et al., 2014). However, non-cell-autonomous and extracellular complementation of the *lre-7* female gametophyte defects by pollen tube-expressed LRE-cYFP-TM, which is expected to remain tethered to the pollen tube plasma membrane, suggests that pollen tube reception does not require release of the LRE ectodomain. Additionally, synergid-expressed LRE-cYFP-TM fully complemented pollen tube reception defects in the *lre-7* female gametophytes, arguing against release of the LRE ectodomain as a mode of LRE function in pollen tube reception.

Another proposed function for LRE is in synergid maturation, when they acquire the competency to induce pollen tube reception (Capron et al., 2008; Rotman et al., 2008). In this model, LRE does not mediate interactions between the pollen tube and the synergids; instead, it is part of a signaling pathway important for synergid development prior to the arrival of the pollen tube. Non-cell-autonomous and extracellular complementation of the *lre-7* female gametophyte defects by pollen tube-expressed

LRE-cYFP argues against such a function for LRE, as it is highly unlikely that *lre* mutant synergids use pollen tube-expressed LRE-cYFP to rapidly mature and induce pollen tube reception. Indeed, a pollen tube reception assay did not show any overt abnormalities in the interactions between LRE-cYFP-expressing pollen tubes and *lre* female gametophytes (Figure 6D).

Separate, FER-Dependent Function for LRE at the Interface of the Pollen Tube and the Synergid Cell in Pollen Tube Reception

LRE is proposed to have two functions in the synergid cell: chaperoning FER in the ER en route to the FA and functioning as a coreceptor with FER in the FA (Li et al., 2015). However, since FER localization in the FA depends on intracellular LRE, it has not been experimentally demonstrated if LRE has any synergid surface-specific function, such as its proposed role as a coreceptor with FER (Li et al., 2015) to signal ROS production (Duan et al., 2014). That is, the *lre* phenotype related to FER signaling at the plasma membrane can also be interpreted as an indirect consequence of the failure of FER to reach the plasma membrane in the absence of LRE. To directly investigate if there is an independent function for LRE in the interface of the pollen tube and the synergids, we ectopically expressed LRE in the pollen tube and used it to rescue the pollen tube reception function of *lre* synergids. Non-cell-autonomous and extracellular complementation of *lre* revealed that indeed LRE has a function at the interface of the pollen tube and the synergid cell and that the two functions of LRE can be uncoupled.

Based on the current model of LRE function as a coreceptor with FER (Li et al., 2015), it would be expected that pollen tube-expressed LRE functioned with FER in the FA, when it rescued pollen tube reception defects in *lre* female gametophytes (Figures 6D and 6E; Supplemental Figure 9C). However, since optimal FER localization in the FA is dependent on LRE (Li et al., 2015), it is not clear what protein functioned with pollen tube-expressed LRE to rescue the pollen tube reception phenotype in the *lre* female gametophyte. We speculated that in *lre* female gametophytes, perhaps sufficient amounts of FER reached the FA and functioned with pollen tube-expressed LRE. We considered this possibility based on the finding that all *lre* ovules showed some FER-GFP in the FA region despite increased intracellular retention of FER-GFP in the *lre* synergids (Li et al., 2015). Consistent with this, in this study, we showed that pollen tube-expressed LRE-cYFP requires FER in the *lre* female gametophyte to rescue the pollen tube reception defect associated with that *lre* female gametophyte (Figures 6H and 6I), indicating that some amount of FER is present in the FA of *lre* synergid cell, at least sufficient enough to rescue the pollen tube reception defect.

Although pollen tube-expressed LRE-cYFP, LRE-cYFP Δ 2 ω , LRE-cYFP Δ GAS, or LRE-cYFP-TM complemented the *lre-7* phenotypes, restoration was not to wild-type levels (Figures 6D and 6E; Supplemental Figure 9C), pointing to additional roles for LRE in the synergid cell, such as intracellular LRE chaperoning FER to the synergid cell surface (Li et al., 2015). Alternatively, the incomplete complementation could be due to the possibility that pollen tube-expressed LRE variants, compared with their synergid cell-expressed counterparts, are not optimal for functioning in

pollen tube reception. Nevertheless, non-cell-autonomous, extracellular complementation of the pollen tube reception defect in the *lre* female gametophyte by pollen tube-expressed LRE-cYFP, LRE-cYFP $\Delta 2\omega$, LRE-cYFP Δ GAS, or LRE-cYFP-TM establishes that LRE performs a synergid cell surface-specific function during interactions between the pollen tube and the synergid cell.

Both *lre* and *fer* are partially transmitted through the female gametophyte, and normal pollen tube reception is detected in a limited number of *lre* or *fer* female gametophytes (Figures 6F and 6G). Such an outcome is perhaps possible due to at least two scenarios: Either synergid-expressed proteins that function redundantly with LRE-FER complex to induce pollen tube reception or additional players, besides LRE-FER complex, might mediate pollen tube reception. Our observation that the *lre-7 fer-4* double mutant had similar levels of pollen tube reception and seed set defects compared with either *lre-7* or *fer-4* single mutants argues against the first possibility. Instead, our results demonstrate that LRE and FER require each other to function in pollen tube reception and predict the existence of another minor signaling pathway to induce pollen tube reception.

The GPI Anchor Addition Domains Are Not Necessary for LRE Function in Pollen Tube Reception

Prediction programs identified GPI anchor addition domains in the LRE sequence with high confidence, and these domains are also highly conserved in LRE family members. Deleting these domains in LRE (LRE-cYFP $\Delta 2\omega$ and LRE-cYFP Δ GAS) led to a dramatic decrease in protein localization in the FA. Localization of LRE-cYFP in the FA was also affected in *gpi8-2* female gametophytes, which are expected to lack a core component of the transamidase. These results point to the need for a GPI anchor in localizing LRE in the FA. Surprisingly, mutant proteins lacking GPI anchor addition domains fully complemented pollen tube reception and seed set defects. Our complementation results indicate that the GPI anchor addition domains are not necessary for LRE function in pollen tube reception.

An alternative interpretation of the complementation results with LRE-cYFP $\Delta 2\omega$ and LRE-cYFP Δ GAS is that sufficient amounts of these proteins may be available in the FA or pollen tube surface for function in the transgenic lines. In both synergid cells and pollen tubes, for example, sufficient amounts of LRE-cYFP $\Delta 2\omega$ may have reached the cell surface and associated with the plasma membrane using the lingering C-terminal hydrophobic tail (Galian et al., 2012), which is expected to be removed by the transamidase from the wild-type LRE pro-protein. Or, in LRE-cYFP Δ GAS, although much of the mutant protein is retained in the cytoplasm (Figure 3D; Supplemental Figure 9B), detectable levels of the protein were released from the female gametophyte to the surrounding integument cells lining the micropyle (Supplemental Figure 5). During this release, perhaps sufficient amounts of mutant proteins were in the FA or in the pollen tube surface and induced pollen tube reception.

LRE functions as a chaperone of FER from the ER to the synergid cell periphery and is required for optimal localization of FER in the FA (Li et al., 2015). It remains to be experimentally demonstrated if this intracellular function was also rescued by LRE-cYFP $\Delta 2\omega$ and LRE-cYFP Δ GAS expressed in the synergids. Our

results showing that pollen tube-expressed LRE complemented *lre* female gametophytes in a FER-dependent manner indicated that some amount of FER is present in the FA of *lre* female gametophytes. Even if LRE-cYFP $\Delta 2\omega$ or LRE-cYFP Δ GAS were unable to rescue the chaperone function of LRE in the synergid cells, they most likely functioned with the available FER in FA of *lre* female gametophytes to induce pollen tube reception.

Based on the full complementation of pollen tube reception and seed set defects in the *lre-7* female gametophyte by synergid-expressed LRE-cYFP-TM, which lacks GPI anchor addition domains, we propose that the GPI anchor is not necessary for LRE function in pollen tube reception. Consistent with this proposal, LRE homologs in some plants do not contain the GAS domain or the ω site region, even though they all contain the portion of the protein that shares sequence similarity to the LRE ectodomain (Supplemental Figure 1). Still, most putative LRE homologs have highly conserved GPI anchor addition domains. One possibility that may reconcile these observations is that the GPI anchor may enhance the efficiency of LRE function in pollen tube reception and hence remain fixed in the population. For example, the GPI anchor may afford flexibility to LRE in setting up signaling microdomains, similar to mammalian GPI-anchored surface proteins (Varma and Mayor, 1998), at the site of the pollen tube-synergid cell interaction, as the exact site of this interaction is determined stochastically.

A Novel 12-Amino Acid Domain in the M8CM of LRE Is Essential for LRE Function in Pollen Tube Reception

Results with synergid-expressed LRE-cYFP-TM also showed that the LRE ectodomain without the GPI anchor addition domains is sufficient to induce pollen tube reception. For some GPI-anchored proteins, proper structure of the ectodomain requires a GPI anchor (Bütikofer et al., 2001). However, LRE does not appear to require the GPI anchor, as LRE lacking the GPI anchor addition domains fully complements the pollen tube reception defects in *lre* female gametophytes. Lack of dominant-negative phenotypes in *lre* synergids and pollen tubes expressing LRE-cYFP-TM indicates it did not acquire novel functions, as in some GPI-anchored surface proteins in which a TM was swapped for a GPI anchor (Shenoy-Scaria et al., 1992, 1993).

We identified LRE as a CRP, as it contains eight cysteines that are arranged in a pattern reminiscent of cysteines in an 8CM protein. This adds LRE to the list of CRPs that play diverse roles in plant reproduction (Marshall et al., 2011; Beale and Johnson, 2013). However, LRE is notable in that it contains an M8CM, with a unique 12-amino acid domain separating the 5th and the 6th cysteines. Additionally, unlike other CRPs, most of which are released from the cell, LRE belongs to a group of CRPs that contain GPI-anchor addition domains (Fliegmann et al., 2011). The novel 12-amino acid domain in LRE is ancient, as it is present even in the putative LRE homolog in the moss (*Physcomitrella patens*) (Figure 1; Supplemental Figure 1). Identification of the M8CM in LRE supports the hypothesis that the 8CM serves as a sequence scaffold to evolve new proteins with different functions (José-Estanyol et al., 2004).

Our study showed that a 12-amino acid domain between the 5th and the 6th cysteines, especially the highly conserved Asn-Asp

dipeptide domain, in LRE is critical for its function and thus provided insights into the mechanism of LRE function in the female gametophyte. M8CM in LRE awaits additional characterization, including biochemical approaches to determine its disulfide bond linkages (Tang and Speicher, 2004) and structural studies to solve its three-dimensional structure. Preliminary three-dimensional structural prediction of LRE in the Phyre2 structure prediction website (<http://www.sbg.bio.ic.ac.uk/phyre2/html/page.cgi?id=index>) showed that this 12-amino acid domain may form a loop between two alpha helices and may mediate interactions between LRE and other proteins.

Although LRE_(CLO)-cYFP accumulated to normal levels in the unknown organelle within synergid cells, it was barely detectable in the FA, indicating that this 12-amino acid domain in the M8CM may mediate interactions between LRE and protein sorting machinery of the endomembrane system that is critical for its trafficking to the FA or between LRE and other proteins that are necessary for retention of LRE in the FA. It is also possible that the 12-amino acid domain in M8CM is critical for chaperoning FER from the ER to the FA (Duan et al., 2014; Li et al., 2015). Alternatively, this domain may mediate functional interactions between LRE and FER during pollen tube reception. For example, the interaction may facilitate binding of the LRE/FER complex with their putative ligand from the pollen tube or synergids or activate FER either by aiding it in reaching the correct confirmation or attaining maturation to initiate signaling in the synergid cell. Failure to mediate functional interactions with FER might be the basis for LRE_(ND88AA)-cYFP not complementing the defects in the *lre* female gametophyte, despite localizing normally in the synergid cell.

An Integrated Approach to Study Subcellular Localization and Function of LRE in Pollen Tube Reception in the Synergids

We analyzed the subcellular localization of LRE and its function in pollen tube reception in the synergids. For localization studies, the FA, an easily identifiable structure of a highly secretory cell, allowed quick monitoring of localization of LRE. Whether it is due to preferential secretion of LRE into the FA or concentration of LRE in the plasma membrane-rich structure of the FA, polarized localization of LRE in the FA allowed us to rapidly test the role of the GPI-anchor addition domains in subcellular localization of LRE in the synergids.

For functional analysis, the synergids also presented an important opportunity to perform genetic studies. Synergids are haploid and essential for sperm release from the pollen tube for fertilization. Loss of LRE function affects the function of the synergids in pollen tube reception, reducing *lre* transmission through the female gametophyte (Capron et al., 2008; Tsukamoto et al., 2010). Conversely, complementation of the pollen tube reception defect by an *LRE* transgene will increase the transmission of both the *lre* mutation and the transgene. With LRE-cYFP, both types of transmission analysis produced the same conclusions (Table 1; Supplemental Table 1). Therefore, for functional analysis of all the variants of LRE, we used enhanced transmission of the transgene through the *lre* female gametophyte to assay complementation, as it was easier to establish the plants of the required genotype (*LRE transgene/+ lre-7/lre-7*) for this analysis.

Utilizing these two important opportunities, in this study, we fused the cYFP reporter to the LRE variants and used stable transformants to perform subcellular localization, pollen tube reception and seed set assays, transmission analysis, and non-cell-autonomous complementation studies. This integrated analysis allowed us to test the relevance of subcellular localization of LRE to its function, which led to the surprising finding that even when lacking domains critical for GPI anchor addition, LRE can still function in pollen tube reception by the synergid cell.

METHODS

LRE Sequence Analysis

Proteins with highest sequence similarity to *Arabidopsis thaliana* LRE in the indicated plants were identified, assembled, aligned, and box-shaded to identify the M8CM. Genes encoding LRE-like proteins in the indicated plants were identified by comparing the *LRE* coding sequence against genomic sequence of corresponding species using tBLASTx on the Accelerating Comparative Genomics (CoGe) Platform (<https://genomevolution.org/CoGe/CoGeBlast.pl>), with default settings (E-value cutoff, 0.001; word size, 3; gap existence penalty, 11; gap extending penalty, 1). Top 100 hits were retrieved from BLAST and assembled in Geneious software (Geneious R6.1.8) using default settings; ~3 kb of the genomic sequence on each side of the assembly was retrieved from the CoGe Organism View tool (<https://genomevolution.org/CoGe/OrganismView.pl>), and the coding region was identified by aligning it with the *LRE* coding sequence. For *Physcomitrella patens*, the coding sequences of *LRE*-like genes were identified using RNaseq reads (Chang et al., 2014) in Integrated Genome Browser (<http://bioviz.org/igb/index.html>). In plants where multiple *LRE*-like genes were found, the one that shared highest identity with *Arabidopsis LRE* was chosen for further analysis. Protein alignment was performed on ClustalW Web server (<http://www.ch.embnet.org/software/ClustalW.html>), with the default settings (scoring matrix, Blossum; opening gap penalty, 10; extending gap penalty, 0.05; end gap penalty, 10; separation gap penalty, 0.05), and the box shade was generated by Mobyle box shade tool (<http://mobyle.pasteur.fr/cgi-bin/portal.py#forms:boxshade>). In each protein sequence, the signal peptide was predicted by SignalP 4.1 (<http://www.cbs.dtu.dk/services/SignalP/>) (Petersen et al., 2011), and the ω site was predicted by BIG-PI plant predictor (http://mendel.imp.ac.at/gpi/plant_server.html) (Eisenhaber et al., 2003).

For GPI8 sequence analysis, *Homo sapiens* GPI8 (hGPI8) and *Schizosaccharomyces pombe* GPI8 (yGPI8) protein sequence were retrieved from NCBI (<http://www.ncbi.nlm.nih.gov/>) and aligned with *Arabidopsis* GPI8 using ClustalW. Pairwise protein identity and similarity were calculated using the SIAS tool (<http://imed.med.ucm.es/Tools/sias.html>).

Plant Materials and Growth Conditions

Arabidopsis seeds were surface sterilized and plated on 0.5× Murashige and Skoog plates containing the corresponding antibiotics. Columbia (Col-0) is the ecotype of all *Arabidopsis* seeds used in this study. Seeds on plates were stratified at 4°C for 2 to 3 d before moving them to the growth chamber maintained at 20°C and continuous light (Philips F17T8/TL741 fluorescent tube light bulb, 100 to 150 $\mu\text{mol} \cdot \text{m}^{-2} \cdot \text{s}^{-1}$). The 7- to 10-d-old seedlings were transferred to the soil and were grown as described (Kessler et al., 2010). *lre-5*, *lre-7* (CS66104), and *fer-4* were reported previously (Tsukamoto et al., 2010; Li et al., 2015). The *gpi8-2/+* (CS853564) seeds were from ABRC.

Cloning Transgenic Constructs

The cYFP (E1403; Tian et al., 2004) reporter gene was fused to wild-type and mutated *LRE* and expressed from the *LRE* promoter. To fuse the cYFP

(E1403; Tian et al., 2004) reporter gene either toward the N terminus or C terminus of *LRE*, fragments were PCR amplified using primers and templates listed in Supplemental Table 9. The fragments were then fused by overlap PCR and the *LRE-cYFP* fusion under the control of *LRE* promoter was cloned into *pENTR/D-TOPO* (Invitrogen; K2400-20). For the N-terminal fusion construct (*ProLRE:cYFP-LRE*), the *cYFP* coding sequence (Tian et al., 2004) was cloned immediately downstream of the predicted secretion signal sequence (Supplemental Figure 2A). The C-terminal fusion construct (*ProLRE:LRE-cYFP*) was generated by placing the *cYFP* coding sequence slightly upstream of the predicted ω site (Figure 2A). Cloned transgenes in *pENTR/D-TOPO* were swapped into the Gateway destination vector *pH7WG* using Clonase II enzyme mix (Life Technologies; 11791020). The resultant plasmids containing the N-terminal and C-terminal fusions of *cYFP* to *LRE* were designated *ProLRE:cYFP-LRE* and *ProLRE:LRE-cYFP*, respectively. Swapping also resulted in these plasmids inheriting unique *Ascl* restriction site from the entry vector, which facilitated cloning of modified *LRE-cYFP* constructs (see below).

For all other mutated, truncated, or ectopically expressed *LRE-YFP* constructs used in this study, fragments with desired changes in *LRE* coding regions were introduced by PCR using primers and templates listed in Supplemental Table 9. Full-length constructs were assembled by overlap PCR and then cloned into the *ProLRE:LRE-cYFP* plasmid that was linearized with *SpeI/Ascl* and using In-Fusion HD Cloning Plus (Clontech; 638909). Linearizing the plasmid followed by in-fusion cloning essentially resulted in swapping wild-type *LRE-cYFP* with the modified *LRE-cYFP* transgene.

For constructs with organelle markers expressed from the *LRE* promoter, vectors carrying the organelle marker (*G-rb*, *Px-rb*, and *ER-rb* plasmids; Nelson et al., 2007) were used to PCR amplify the organelle marker. The *LRE* promoter and the *mCherry* coding sequence along with *NOS* or *35S* terminator were PCR amplified using the primers and templates listed in Supplemental Table 9. These fragments were then assembled by overlap PCR to obtain the desired transgene that was cloned into *EcoRI* linearized *pFGC19* using In-Fusion HD Cloning Plus.

Plant Transformation

Transformation solution containing *Agrobacterium tumefaciens* (GV3101 strain) harboring the desired plasmid were sprayed onto Arabidopsis inflorescences (Chung et al., 2000). Hygromycin-resistant transformants were selected as described (Harrison et al., 2006).

Isolation of Single Insertion Lines

For each construct, multiple T1 hygromycin-resistant transformants were selected and moved to soil. T1 plants are expected to be heterozygous for the transgene at the insertion locus. Upon bolting, stage 12c buds (Smyth et al., 1990) were emasculated and 24 h after emasculation *cYFP* expression in synergid cells was scored. For each construct, the number of T1 lines scored for *cYFP* expression and seed set are provided in Supplemental Figures 2 and 8. Among these lines, candidate single insertion lines were identified based on ~50% ovules containing the *cYFP* signal. Pollen from candidate single insertion lines were then crossed to wild-type female and progeny were analyzed for segregation of hygromycin resistance. Those lines that showed a 1:1 segregation of hygromycin resistance:susceptibility were considered single insertion lines.

The above approach was used for all constructs except *ProLRE:LRE-cYFP Δ GAS*. When the progeny of crosses were checked for hygromycin resistance, we obtained slightly higher than expected transmission of the *ProLRE:LRE-cYFP Δ GAS* transgene through the male gametophyte (we do not yet understand why), for the construct carrying this transgene (Supplemental Table 5). Hence, for this construct, single insertion lines were merely identified using 1:1 segregation of *cYFP*-positive ovules in T1 hygromycin-resistant transformants.

RNA Isolation and RT-PCR

RNA was isolated according to Qin et al. (2009). Briefly, ~20 pistils were collected 24 h after emasculation for each sample and were frozen in -80°C until RNA extraction. RNA was isolated using RNeasy Plant Mini Kit (Qiagen; catalog #74904) according to manufacturer's instruction. RNA samples were then subjected to RNase-free DNase I (Life Technologies; catalog #AM2222) treatment to remove potential DNA contamination, cleaned up using the RNeasy MinElute Cleanup Kit (Qiagen; catalog #74204), and tested for RNA integrity on Agilent Bioanalyzer 2100 (Agilent Technologies).

For RT-PCR, cDNA was synthesized from 5 μg total RNA using the ThermoScript RT-PCR System (Life Technologies; catalog #11146-024), and genes of interest (*LRE-cYFP* transgene and *ACTIN11*) were amplified using TaKaRa Ex Taq DNA polymerase (Fisher Scientific; catalog #TAK_RR01BM), with primers listed in Supplemental Table 9.

Confocal Imaging

Fluorescent images were taken using a Leica SP5 confocal laser scanning microscope system. For *cYFP* imaging, samples were excited with a 488-nm laser line, and emission spectra between 510 and 550 nm were collected. For colocalization analysis, *cYFP* and *mCherry* signals were detected in sequential scan mode, with the excitation at 488 and 543 nm, and emission spectra at 510 to 550 nm and 580 to 660 nm, respectively, were collected. *YFP* images were processed and quantified with ImageJ software (<http://imagej.nih.gov/ij/>). Single confocal sections were used for colocalization analyzed on Leica Application Suite (LAS 2.6.0.7200).

Pollen Tube Reception Assays

In vivo pollen tube reception assays were done as previously described (Tsukamoto et al., 2010). Briefly, *ProLAT52:GUS* pollen was crossed to emasculated stage 14 pistils of the indicated plants. Crossed pistils were then collected 24 h after pollination, stained for GUS activity, and imaged using differential interference contrast optics in a Zeiss Axiovert 100 microscope. Aniline blue staining was performed according to Mori et al. (2006). Stained pistils were mounted with 15% glycerol, and pollen tube behavior was scored under a Zeiss Axiovert 100 fluorescent microscope.

Accession Numbers

Sequence data from this article can be found in the GenBank/EMBL libraries under accession numbers *GPI8* (*AT1G08750*), human *GPI8* (NP_005473), yeast *GPI8* (CAC13970), *LRE* (*At4g26466*), and *FER* (*At3g51550*).

Supplemental Data

Supplemental Figure 1. Multiple sequence alignment of LRE-like proteins.

Supplemental Figure 2. *cYFP-LRE* and *LRE-cYFP*, but not *LRE Δ SP-cYFP*, complemented the *lre-7* seed set defect.

Supplemental Figure 3. FA localization of *LRE-cYFP* remains unchanged upon pollen tube arrival or burst.

Supplemental Figure 4. Puncta of *LRE-cYFP* in synergids does not colocalize with the ER, Golgi, or peroxisome markers.

Supplemental Figure 5. *LRE-cYFP Δ GAS* is released from the synergids.

Supplemental Figure 6. Genotyping of F1 progeny from a cross between *gpi8-2/+*(\varnothing) and *ProLRE:LRE-cYFP* (σ).

Supplemental Figure 7. The *gpi8-2* mutation does not affect LRE-cYFP-TM localization in the FA.

Supplemental Figure 8. Seed set of T1 plants carrying synergid cell-expressed constructs.

Supplemental Figure 9. Pollen tube-expressed LRE-cYFP, LRE-cYFP Δ 2 ω , LRE-cYFP Δ GAS, and LRE-cYFP-TM complemented *lre* seed set defect.

Supplemental Table 1. Enhanced transmission of the *ProLRE:LRE-cYFP* transgene through the *lre-7* female gametophyte.

Supplemental Table 2. Transmission of the *gpi8-2* mutation through the male gametophyte, but not through the female gametophyte, is abolished.

Supplemental Table 3. Abnormal LRE-cYFP localization cosegregates with the *gpi8-2* mutation.

Supplemental Table 4. LRE-cYFP-TM localization in the FA of synergids is not affected by the *gpi8-2* mutation.

Supplemental Table 5. Enhanced transmission of transgenes with mutated GPI anchor addition domains through the *lre-7* female gametophyte.

Supplemental Table 6. Normal transmission of mutated M8CM transgenes through the *lre-7* female gametophyte.

Supplemental Table 7. *ProPLL3:LRE-cYFP* is transmitted at an enhanced rate to the progeny when crossed to a *lre-5* female but not wild-type female.

Supplemental Table 8. *ProPLL3:LRE-cYFP-TM* is transmitted at an enhanced rate to the progeny when crossed to a *lre-7* female but not wild-type female.

Supplemental Table 9. Primers used in this study.

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AUTHOR CONTRIBUTIONS

X.L., E.S., and R.P. designed the experiments. X.L., C.C., N.P., Y.W., J.N., M.B., and C.H. performed the experiments. R.P. and X.L. wrote the article.

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