Ectopic Expression of a Self-Incompatibility Module Triggers Growth Arrest and Cell Death in Vegetative Cells

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Self-incompatibility (SI) is used by many angiosperms to reject self-pollen and avoid inbreeding. In field poppy (Papaver rhoeas), SI recognition and rejection of self-pollen is facilitated by a female S-determinant, PrsS, and a male S-determinant, PrpS. PrsS belongs to the cysteine-rich peptide family, whose members activate diverse signaling networks involved in plant growth, defense, and reproduction. PrsS and PrpS are tightly regulated and expressed solely in pistil and pollen cells, respectively. Interaction of cognate PrsS and PrpS triggers pollen tube growth inhibition and programmed cell death (PCD) of self-pollen. We previously demonstrated functional intergeneric transfer of PrpS and PrsS to Arabidopsis (Arabidopsis thaliana) pollen and pistil. Here, we show that PrpS and PrsS, when expressed ectopically, act as a bipartite module to trigger a self-recognition/self-destruct response in Arabidopsis independently of its reproductive context in vegetative cells. The addition of recombinant PrP to seedling roots expressing the cognate PrpS resulted in hallmark features of the P. rhoeas SI response, including S-specific growth inhibition and PCD of root cells. Moreover, inducible expression of PrsS in PrpS-expressing seedlings resulted in rapid death of the entire seedling. This demonstrates that, besides specifying SI, the bipartite PrpS-PrsS module can trigger growth arrest and cell death in vegetative cells. Heterologous, ectopic expression of a plant bipartite signaling module in plants has not been shown previously and, by extrapolation, our findings suggest that cysteine-rich peptides diversified for a variety of specialized functions, including the regulation of growth and PCD.

Pollen-pistil interactions are complex, crucial events in plant reproductive biology, involving bidirectional signaling between the pistil and the pollen landing on it. Many of the responses regulating pollination take place within the pollen grains, which constitute the highly reduced haploid male gametophyte. The pollen grain is composed of the specialized vegetative cell that contains within itself two sperm cells, complete with cell walls and plasma membranes. The pollen’s role is to deliver two sperm cells to the embryo sac so that double fertilization can take place. Thus, pollen represents a unique gametophytic structure; for example, serial analysis of gene expression studies have revealed that 83% of the pollen-expressed gene tags are pollen specific and thus thought to be critical for pollen function (da Costa-Nunes and Grossniklaus, 2003; Honys and Twell, 2004; Mergner et al., 2020).

Self-incompatibility (SI) is an important mechanism used by flowering plants to prevent selfing. It is controlled by a multiallelic S-locus allowing self/nonself recognition between pistil and pollen. In several SI systems, when male and female S-determinant allelic specificities match, self (incompatible) pollen is recognized and rejected before fertilization can occur. A key characteristic of SI determinants is that they are extremely tightly regulated, both in a developmental and a tissue-specific manner, being expressed solely in pistil
and pollen cells during a narrow developmental window, as the tissues approach maturity (Takayama and Isogai, 2005). SI in poppy (Papaver rhoeas) is controlled and specified by S-determinants expressed specifically in the stigma (PrsS; Foote et al., 1994) and pollen (PrpS; Wheeler et al., 2009). PrpS encodes a novel integral membrane protein with several predicted transmembrane domains; PrsS encodes a small, secreted protein and is the founding member of the large family of S-protein homologs (SPHs), which are found in most dicotyledonous plants, some fungi, and metazoa (Rajasekar et al., 2019). This family of small, secreted proteins has features similar to cysteine-rich peptides (CRPs), which include ligands known to be involved in diverse signaling pathways (Wheeler et al., 2010; Marshall et al., 2011; Bircheneder and Dresselhaus, 2018). PrpS and PrsS are secreted proteins, and their receptors (Sparks et al., 2013). Many signaling peptides are perceived by receptor-like kinases, and it is thought that much of the specificity of responses is due to the localized expression of ligands and their receptors (for review, see Breiden and Simon, 2016). For example, although CLAVATA3 CLE family peptides act in both roots and shoots (Fletcher et al., 1999), they nevertheless function in both organs specifically in apical meristematic tissues.

Heterologous expression of plant genes in other plant species has often been used to identify function phenotypically by dominant gene activity (Diener and Hirschi, 2000). Ectopic expression has also been used to demonstrate function; for example, Boutiller et al. (2002) showed that constitutive expression of the BABY BOOM transcription factor promotes cell proliferation and morphogenesis during embryogenesis. However, transfer of two genes encoding a receptor-ligand pair that are normally specifically expressed in certain tissues for a specific function to a completely different cellular context, to our knowledge, has not previously been explored. Thus far, examples of the ectopic expression of single genes in plant cells has typically been restricted to reiterate their function in other cell types to show functional relatedness or to recapitulate an evolutionarily divergent event using similar genes from different species. One of the best known examples is perhaps the expression of chimeric RNase genes in anthers of transformed tobacco (Nicotiana tabacum) and oilseed rape (Brassica napus) plants, which specifically destroyed the tapetal cells of developing pollen, resulting in male sterility (Mariani et al., 1990).

Here, we have examined the effect of the ectopic expression of PrpS and PrsS from P. rhoeas in vegetative cells of Arabidopsis, using characteristic markers of P. rhoeas SI-PCD to examine function. We show that the heterologous, ectopic expression of these genes, which specify a tightly controlled reproductive trait in the male gametophyte, can trigger a self-recognition: self-destruct response, resulting in growth arrest and PCD in vegetative sporophytic cells. Ectopic expression of PrpS and PrsS in Arabidopsis recapitulates major cellular aspects of the P. rhoeas SI response in vegetative cells, providing evidence that this heterologous, bipartite module can signal to similar cellular targets in different cell types.

RESULTS

PrpS Treatment Results in S-Specific Root Growth Inhibition of PrpS-Expressing Seedlings

In P. rhoeas, the interaction of cognate PrpS and PrsS triggers a Ca²⁺-dependent signaling network in pollen, resulting in a rapid growth arrest followed by PCD of incompatible pollen after SI induction (Franklin-Tong et al., 1993, 1995, 1997). To examine if the PrpS-PrsS module might also work outside the specific context of pollen-pistil interactions, we examined if growth
inhibition and PCD caused by the PrpS-PrsS module also could be triggered in other tissues. We therefore expressed PrpS1 under the control of a constitutive UBQ10 promoter in Arabidopsis plants and established five independent single T-DNA insertion lines (pUBQ10::PrpS1 lines 7, 11, 12, 13, and 16). Reverse transcription quantitative PCR (RT-qPCR) showed that PrpS1 mRNA was substantially expressed in these transgenic Arabidopsis seedlings (Fig. 1A; Supplemental Fig. S1). Focusing first on root growth, we did not observe differences in the root length between Arabidopsis Columbia-0 (Col-0) wild-type and pUBQ10::PrpS1 seedlings (Fig. 1B), demonstrating that PrpS1 expression alone did not alter seedling development. Next, we applied recombinant PrsS1 protein to 4-d-old root tips of wild-type and pUBQ10::PrpS1 seedlings. Exposure to PrsS1 protein did not show any inhibition effect of normal development and growth of Arabidopsis wild-type seedlings (Supplemental Fig. S2). However, PrsS1 treatment of pUBQ10::PrpS1 seedlings resulted in a rapid and complete inhibition of root growth (Fig. 1, B and C). The growth of pUBQ10::PrpS1 seedling roots was inhibited by recombinant PrsS1 protein in a dose-dependent manner (Fig. 1D). Treatment of pUBQ10::PrpS1 roots with 5 ng μL⁻¹ PrsS1 significantly inhibited their growth rate, while 10 ng μL⁻¹ or more completely blocked root elongation. This provides evidence that the PrpS-PrsS module, although its constitutive components are normally only expressed in pollen and pistil, respectively, and triggers a response in pollen specifically, can also act to trigger the inhibition of growth of vegetative, sporophytic cells.

The S-allele-specific inhibition is a key feature of the P. rhoeas SI response. To test this, we treated pUBQ10::PrpS1 and wild-type seedlings with either PrsS1 or PrsS3 recombinant protein. pUBQ10::PrpS1 seedling roots were strongly inhibited by the PrsS1 protein, while the PrsS3 protein had no effect; wild-type

![Figure 1. Expression of PrpS in transgenic Arabidopsis triggers root growth inhibition after cognate PrsS treatment.](image-url)

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**Figure 1.** Expression of PrpS in transgenic Arabidopsis triggers root growth inhibition after cognate PrsS treatment. A, RT-PCR shows the expression of PrpS1 mRNA in pUBQ10::PrpS1 transgenic seedlings. Actin2 was used as a housekeeping gene control. Quantification of the relative expression levels is shown in Supplemental Figure S1. B and C, S-specific inhibition of root growth of pUBQ10::PrpS1 seedlings after PrsS1 treatment. B, Images of 4-d-old seedlings 24 h after treatment with PrsS proteins (10 ng μL⁻¹). Black lines indicate the positions of root tips when treated. Only pUBQ10::PrpS1 seedlings (line 12) treated with PrsS1 (bottom, center) display inhibited root growth. This line was used for all the other experiments if not specified. Bar = 1 cm. C, Quantitation of increases in seedling root length from different transgenic lines (see A) treated with PrsS proteins (10 ng μL⁻¹) 24 h after treatment (means ± SD; n = 20–25 seedlings). All five lines had root growth significantly inhibited by PrsS1 when comparisons were made with either PrsS1 or mock treatment for each line (two-way ANOVA multiple comparison: NS, not significant; ***p < 0.001). D, Root growth of pUBQ10::PrpS1 seedlings was inhibited by PrsS1 in a dose-dependent manner. The x axis indicates time (days) after transfer of plates to the growth chamber. The arrow indicates when the treatment was added. Data shown are means ± SD (n = 20–25 seedlings). E, PrsS1 treatment induces rapid root growth inhibition of pUBQ10::PrpS1 seedlings in an S-specific manner. The arrow indicates the time point of PrsS1 addition (10 ng μL⁻¹). Two-way ANOVA shows that PrsS1 treatment significantly inhibited root growth (P < 0.001) while PrsS3 did not (P = 0.29), in comparison with wild-type (WT) seedlings treated with PrsS1. Data shown are means ± SD (n = 6).
seedling roots were not inhibited by any treatment (Fig. 1, B and C). As only a cognate PrpS-PrsS combination caused growth inhibition, this shows that the S-determinants maintain their S-specificity in Arabidopsis roots.

As the SI response in pollen triggers rapid inhibition of incompatible pollen tube growth, we examined the timing of inhibition of growth of the roots in more detail, using a perfusion chamber system in combination with confocal microscopy (Krebs and Schumacher, 2013). Under these conditions, wild-type seedling roots elongated at a rate of ~2.3 μm min⁻¹, and the addition of PrsS proteins did not affect this (Fig. 1E; Supplemental Fig. S3, A and D). However, the addition of PrsS₁ protein resulted in a rapid reduction of root growth of pUBQ10::PrpS₁ seedlings (P < 0.001, two-way ANOVA; Fig. 1E). Growth was completely inhibited within 5 to 20 min after the addition of PrsS₁ (Supplemental Fig. S3, B, E, and G). This inhibition was only observed with a cognate PrpS-PrsS interaction; in a compatible interaction, using the noncognate recombinant PrsS₁ protein, roots of pUBQ10::PrpS₁ seedlings grew at a similar rate to wild-type roots (P = 0.29, two-way ANOVA; Fig. 1E; Supplemental Fig. S3, C and F).

Taken together, these data demonstrate that PrpS and PrsS interaction in roots rapidly elicits the inhibition of growth. This response is strikingly similar to what was observed in P. rheas pollen tubes during the SI response (Thomas and Franklin-Tong, 2004). However, a key difference is that root is a multicellular organ that increases its length by diffuse growth, whereas the pollen tube is a single cell elongating by tip growth. Together, these data demonstrate that the PrpS-PrsS bipartite signaling module can operate ectopically to inhibit the growth of vegetative cells.

PrsS Triggers Cell Death and DEVDase Activation in PrpS-Expressing Seedlings

In P. rheas pollen, downstream of PrpS and PrsS interaction, after inhibition of growth, a distinctive PCD program is triggered. To investigate if this aspect of the SI response could be recapitulated in PrpS-expressing Arabidopsis roots, we examined root cells for evidence of death after PrsS treatment. We first examined plasma membrane permeability using propidium iodide (PI) staining and nuclear integrity using a nucleus-localized fluorescent protein marker line (pUBQ10::NLS-YC3.6 [Nagai et al., 2004] containing both nucleus-localized eCFP and cpVENUS). Twenty-four hours after the addition of PrsS₁ to pUBQ10::PrpS₁ seedling root tips, we found that many cells showed plasma membrane permeabilization to PI and loss of nuclear integrity, providing evidence of death (Fig. 2, A and B). This occurred in the whole root tip region, including the different cell types in the root cap, the root meristem, transition zone, and elongation zone (Fig. 2, A and B). Examining temporal changes to the root after PrsS₁ treatment, we observed a gradual increase in the number of dead cells (Supplemental Fig. S4). A significant increase in PI staining was initially observed in the lateral root cap 1 h after PrsS₁ treatment. At 2 h, cell death was observed in the columella root cap region. Cell death in the meristem was observed 4 h after PrsS₁ treatment, and the number of cells affected increased over time. In contrast, in the controls (mock treated and treated with PrsS₃), only a few PI-positive cells were observed in the root cap (Fig. 2B; Supplemental Fig. S4), which undergoes PCD as part of its regular developmental program (Fendrych et al., 2014). These results provide good evidence that cognate combinations of PrpS-PrsS, besides specifying SI, can operate to trigger cell death in vegetative cells.

To investigate if a similar pathway to that triggered in P. rheas pollen was utilized in the death of the root cells, as a DEVDase is implicated as a key PCD executor (Bosch and Franklin-Tong, 2007) in P. rheas pollen SI-PCD, we examined this protease activity in the pUBQ10::PrpS₁ seedling roots. The chemically synthesized probe CR(DEVD)₂ was employed to detect DEVDase activity in roots in vivo. In wild-type roots, consistent with the occurrence of normal, constitutive root cap PCD (Fendrych et al., 2014), DEVDase activity was detected in the outermost layer of the root in the root cap prior to treatment (Fig. 2C; Supplemental Fig. S5A). The addition of PrsS proteins to wild-type seedling roots did not affect DEVDase activity even after 4 h (Fig. 2C; Supplemental Fig. S5A). However, treatment of pUBQ10::PrpS₁ roots with PrsS₁ induced the activation of DEVDase activity in several different zones and cell types, including the root cap, meristem, and elongation zone (Fig. 2C; Supplemental Fig. S5B). When pUBQ10::PrpS₁ roots were treated with PrsS₁ protein, no major differences in DEVDase activity were observed compared with that in untreated roots (Supplemental Fig. S5, C and D). This demonstrates that DEVDase activation is induced by PrpS-PrsS interaction in these PrpS₁-expressing Arabidopsis seedling roots and that DEVDase activation is S-allele specific in these vegetative tissues. This suggests that a similar pathway is reconstituted in these vegetative cells by this bipartite module.

PrsS Treatment Triggers an S-Specific Ca²⁺ Signature in PrpS-Expressing Roots

We next investigated whether other hallmark downstream features of the P. rheas SI response were triggered in the PrpS-expressing roots after the addition of cognate PrsS proteins. To monitor the [Ca²⁺]cyt spatiotemporally, the genetically encoded calcium indicator YC3.6 (Nagai et al., 2004; Krebs et al., 2012) was coexpressed with PrpS₁ in Arabidopsis seedlings. We observed no obvious change in the [Ca²⁺]cyt when wild-type seedlings were treated with PrsS₁ protein (Fig. 3A; Supplemental Fig. S6A). However, when PrsS₁ protein was added to PrpS₁-expressing seedlings, we detected transient [Ca²⁺]cyt increases in their roots. The
increase was first observed in the elongation zone of the root, peaking ~10 min after PrsS protein addition, and subsequently gradually decreased back to the approximate resting level within ~25 min (Fig. 3A; Supplemental Fig. S6B). An increase in \([Ca^{2+}]_{\text{cyt}}\) in the meristem and columella regions was also observed (Supplemental Fig. S6B). These \([Ca^{2+}]_{\text{cyt}}\) dynamics were not observed in \(PrpS_1\)-expressing seedlings treated with \(PrsS_1\) protein (Fig. 3A; Supplemental Fig. S6C), demonstrating that this \([Ca^{2+}]_{\text{cyt}}\) response was \(S\)-specific. We also examined roots for increases in nuclear \(Ca^{2+}\) \([\text{Ca}^{2+}]_{\text{nuc}}\) after the addition of PrsS by introducing an NLS-YC3.6 construct into the \(pUBQ10::PrpS_1\) transgenic seedlings. We observed increases in \([Ca^{2+}]_{\text{nuc}}\) at the root tip, including columella, meristem, and elongation zone (Supplemental Fig. S7), which were spatiotemporally similar to the \([Ca^{2+}]_{\text{cyt}}\) response. Our observation of unsynchronized \(Ca^{2+}\) signatures in different parts of the root hints at a possible transmission of \(Ca^{2+}\) signaling between neighboring tissues in the \(pUBQ10::PrpS_1\) root triggered by \(PrsS_1\). As increases in \([Ca^{2+}]_{\text{cyt}}\) are a key feature of the SI response, our data suggest that we may be observing an SI-like response in vegetative tissues.

PrsS Induces \(S\)-Specific Cytoplasmic Acidification in \(PrpS\)-Expressing Roots

Another hallmark feature of the \(P. rhoeas\) SI is cytosolic acidification. We examined \(PrpS_1\)-expressing roots
treated with PrsS proteins for alterations in \([\text{pH}]_{\text{cyt}}\) using the genetically encoded pH-sensitive GFP variant, pHluorin (Moseyko and Feldman, 2001). After 30 min of PrsS treatment, PrpS1-expressing roots displayed a significant drop in \([\text{pH}]_{\text{cyt}}\) (\(P < 0.0001\), one-way ANOVA; Fig. 3, B and C). Further cytoplasmic acidification continued until \(1 \text{ h}\), and levels remained low, as the pHluorin 405:488 ratio at 3 h was not significantly different to that at 1 h (\(P = 0.7975\), one-way ANOVA; Fig. 3, B and C). This rapid drop in \([\text{pH}]_{\text{cyt}}\) was only observed in PrpS1-expressing roots treated with cognate PrsS1 proteins and not in wild-type seedlings treated with PrsS1/3 proteins or PrpS1-expressing roots treated with PrsS3 proteins (Supplemental Fig. S8). The temporal pH dynamics after PrpS-PrsS interaction in Arabidopsis root was similar to that observed in *P. rhoeas* pollen after SI induction. These data demonstrate that the cognate PrpS-PrsS interaction in Arabidopsis roots induces cytoplasmic acidification and further support the idea that a *P. rhoeas* SI-like signaling
pathway is triggered in Arabidopsis roots after the interaction of cognate PrpS and PrsS.

**PrsS Triggers Actin Cytoskeletal Remodeling in PrpS-Expressing Seedling Roots**

As highly characteristic alterations to the actin cytoskeleton are a key feature of *P. rheas* SI, we examined the dynamics of the actin cytoskeleton to see if this characteristic response also took place in roots. We added recombinant PrsS to *pUBQ10::PrpS1* transgenic seedling roots that also expressed the genetically encoded actin marker, LifeAct-mRuby2 (Dyachok et al., 2014; Bascom et al., 2018). Wild-type roots displayed typical actin filament bundles before and after PrsS1 application (Supplemental Fig. S9A). *PrpS1*-expressing roots showed a similar actin organization prior to the addition of recombinant PrsS1 (Fig. 3D). However, by 60 min after PrsS1 application, the mRuby2 signal in the *PrpS1*-expressing seedling roots was much reduced, fragmented actin filaments were detected, and small punctate actin foci had formed (Supplemental Fig. S9B). At 3 h, the actin foci were brighter and larger (Fig. 3D; Supplemental Fig. S9B). These distinctive actin alterations are very similar to what has been described for incompatible pollen in the *P. rheas* SI response (Snowman et al., 2002). In roots, we also observed abnormally thick actin bundles and actin aggregation around the nucleus at 3 h after cognate PrsS treatment (Supplemental Fig. S9B). *PrpS1*-expressing roots did not undergo any actin remodeling after treatment with recombinant PrsS3 protein (Fig. 3D), demonstrating that actin remodeling is an *S*-specific event. Together, these observations demonstrate that interaction of PrpS and PrsS in Arabidopsis roots triggers a signaling network involving hallmark features observed in incompatible pollen in the *P. rheas* SI response (Snowman et al., 2002), suggesting that they can recapitulate an SI-like response in vegetative tissues.

**PrpS Treatment Results in an *S*-Specific Cell Death of PrpS-Expressing Leaf Protoplasts**

As our data suggested that a *P. rheas* SI-PCD-like signaling pathway could be triggered in Arabidopsis root cells, we wondered whether this response might also be observed in other somatic cell types. We
therefore examined whether the viability of leaf protoplasts derived from PrpS1-expressing Arabidopsis plants might also be affected by PrsS1 protein treatment. We utilized a nucleus-localized eCFP (NLS-eCFP) signal as a cell viability marker for leaf protoplasts. After 8 h of incubation with PrsS1 protein, only PrpS1-expressing protoplasts showed a loss of the NLS-eCFP signal, together with abnormal cell shape and leakage of cellular contents (Fig. 4A). In contrast, treatment with PrsS1 protein or mock treatment with buffer had no effect; these control protoplasts appeared viable and intact and the same as untreated wild-type protoplasts (Fig. 4A). Quantitative, temporal analysis showed a gradual and significant decrease in the ratio of protoplasts displaying a positive NLS-eCFP signal. Prior to treatment, this was 95.7%, and it decreased to 71.2% at 1 h \((P < 0.05, \text{one-way ANOVA}; \text{Fig. 4B})\), progressively decreasing down to 15.6% after 8 h \((P < 0.001, \text{one-way ANOVA}; \text{Fig. 4B})\). This was not observed in the wild-type protoplasts or PrpS1-expressing protoplast incubated with PrsS2 proteins, which displayed NLS-eCFP signals not significantly different from the untreated controls \((P = 0.7532, \text{one-way ANOVA}; \text{Fig. 4B})\). These data demonstrate that the interaction between PrpS and PrsS in leaf protoplasts is sufficient to induce cell death in an S-specific manner. This provides further evidence that the S-determinants can operate ectopically in totipotent protoplasts.

Coexpression of PrsS and PrpS Triggers S-Specific Cell Death in Whole Plants

Finally, we investigated whether PrsS, when expressed in planta, was able to exert the same effect as treatment with recombinant PrsS protein in whole plants. We introduced PrsS into the pUBQ10::PrpS1 background line under the control of an estradiol-inducible promoter \((pH3:3XVE::PrsS1/pUBQ10::PrpS1, \text{referred to as XVE::PrsS1/PrpS1 hereafter})\). XVE::PrsS1/PrpS1 seeds completely failed to germinate on medium containing estradiol. In contrast, no significant difference in the germination rate \((95.5\%–97.5\%)\) of the background pUBQ10::PrpS1 line and the XVE::PrsS1/PrpS1 line was observed before and after estradiol induction (Table 1). This effect on seed germination demonstrated that simultaneous expression of cognate PrpS and PrsS in seeds induces cell death in planta.

To test this hypothesis and examine cell viability after estradiol induction further, we induced PrsS1/PrpS1 expression by transferring XVE::PrsS1/3/PrpS1 seedlings to medium containing estradiol. Root growth was rapidly inhibited after transfer to estradiol, whereas pUBQ10::PrpS1 and XVE::PrsS1/PrpS1 seedlings were not affected (Fig. 5, A and B; Supplemental Fig. S10). Strikingly, the XVE::PrpS1/PrpS1 seedlings were stunted and cotyledons were white after 48 h on estradiol (Fig. 5A). These data show that the estradiol-induced expression of PrsS1 (Supplemental Fig. S11) is sufficient to cause S-specific root growth inhibition and subsequent systemic PCD of the entire pUBQ10::PrpS1 seedling. Moreover, time-lapse examination of XVE::PrsS1/PrpS1 seedling roots expressing NLS-YC3.6 after estradiol treatment revealed localized increases in \([Ca^{2+}]_{\text{nuc}}\) 3 h after estradiol induction (Fig. 5C), providing evidence for estradiol-induced expression of PrsS and subsequent PrpS-PrsS interaction. At 5 h, a dramatic decrease in nuclear integrity was observed in root tips, and this continued for up to 11 h, when almost no cells with intact nuclei were observed in root tips (Fig. 5C). PI staining showed that besides the loss of nuclear integrity, plasma membrane permeability was also affected (Fig. 5D). Thus, cell death triggered by the coexpression of cognate PrpS and PrsS was observed in whole root tissues (Supplemental Fig. S12). Control plants that expressed noncognate PrsS3 and PrpS1 exhibited no major changes in nuclear integrity after estradiol induction (Supplemental Fig. S12). Together, these data demonstrate that the coexpression of cognate PrpS and PrsS induces the death of the whole plant in Arabidopsis. This suggests that this two-component system is capable of triggering cell death when they are expressed together, regardless of tissue or cell type.

DISCUSSION

The S-locus in P. rheasr encodes a pair of S-determinants, PrpS and PrsS. Their tissue- and development-specific expression, solely in pollen and pistil, respectively, is tightly regulated, and they interact in an allele-specific manner to specify and mediate the SI response within the male gametophyte pollen during early pollination. We previously demonstrated functional intergeneric transfer of the P. rheas

| Table 1. Coexpression of cognate PrsS and PrpS completely abolishes Arabidopsis seed germination |
|---------------------------------------------------------|-------------------------------------------------|------------------------------------------------------|
| Treatment                                               | pUBQ10::PrpS1                                  | XVE-PrsS1/PrpS1                                      | XVE-PrsS1/3/PrpS1                                    |
| Mock                                                    | 95.5% (128/134)                                | 96.1% (147/153)                                     | 97.5% (197/202)                                     |
| Estradiol                                               | 96.6% (113/117)                                | 0.0% (0/108)                                        | 97.0% (131/135)                                     |

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S-determinants to the reproductive system of Arabidopsis (de Graaf et al., 2012; Lin et al., 2015). Here, we show that PrpS and PrsS do not just function as S-determinants to specify SI but that they can operate beyond their usual reproductive context. We demonstrate that the effect of this self-recognition:self-destruct mechanism is not confined to the male gametophyte but that PrpS and PrsS can also act as a heterologous bipartite module to trigger a canonical SI-like response, resulting in growth inhibition and PCD independent of the reproductive context.

Figure 5. Ectopic expression of PrpS and PrsS in Arabidopsis triggers cell death in whole seedlings in an S-specific manner. A and B, Root growth of XVE-PrsS1/PrpS1 seedlings was inhibited after estradiol induction in an S-specific manner. A, Four-day-old seedlings were transferred to new medium containing 10 μM estradiol. Images were taken 48 h after treatment. White dashed lines indicate the positions of the root tips at the time of transfer. Estradiol-induced expression of PrsS1 resulted in the death of the whole seedling (top, center), whereas no obvious effect was observed when PrsS3 was expressed (top, right). B, Quantification of root length at 24 and 48 h after estradiol induction reveals the inhibition of root growth in XVE-PrsS1/PrpS1 lines upon transfer to estradiol plates, whereas the growth of roots of XVE-PrsS3/PrpS1 and pUBQ10::PrpS1 seedlings was not affected (means ± SD; n = 20 seedlings). C, Estradiol induction resulted in nuclear disintegration and cell death of XVE-PrsS1/PrpS1 seedlings. NLS-YC3.6 was used to monitor the nuclear integrity after estradiol induction over time. Confocal images of merged eCFP (green) and cpVENUS (red) channels are shown. The yellow signal (green-red overlap) shows intact nuclei; extensive nuclear disintegration (loss of yellow signal) was observed as early as 5 h after estradiol induction and was almost complete by 11 h. The fluorescence signal was so weak at 5 h that the confocal laser power was increased from 1.5% (0 and 3 h) to 3.5% (5, 7, 11, and 24 h) to allow visualization of the seedling. NLS-YC3.6 monitors [Ca2+]nuc and reveals that increases (red signal; indicated by white arrowheads) could be observed 3 h after induction. Bar = 100 μm. D, PI staining of a representative root at 24 h after estradiol induction reveals that virtually all the cells are dead (white signal). Bar = 100 μm.
when ectopically expressed in sporophytic tissues of Arabidopsis.

Pollen is a highly specialized gametophytic organism with very specific, precise functions related to reproduction. As such, pollen displays a distinct molecular profile that is distinct from all other plant tissues (da Costa-Nunes and Grossniklaus, 2003; Honys and Twell, 2004; Mergner et al., 2020). The finding here that PrpS and PrsS can act outside of this reproductive context to trigger an SI-like growth arrest and PCD response in vegetative cells of the sporophyte when expressed ectopically is surprising, exciting, and not predicted by our earlier studies. To our knowledge, ectopic transfer of a two-component module from the reproductive context into the vegetative sporophytic context to trigger an SI-like growth arrest and PCD reveals the existence of hundreds of predicted reproductive traits, which are normally responsible for controlling a reproductive milestone, as it demonstrates that these two genes, one has not previously been reported in plants. This is a breakthrough that PrpS and PrsS can act outside of this reproductive trait, are sufficient to trigger signaling to growth arrest and cell death in numerous cell types, independent of their particular tissue-specific and developmental context.

Our data showing that the PrpS-PrsS module can act ectopically provide potential new clues to the possible origin and evolution of bipartite genetic modules that act in cell-cell signaling networks. PrsS has homologs in a large family named after them, the SPHs (also known as plant self-incompatibility protein S1 homologs in the databases), comprising more than 1,800 homologous sequences in more than 70 plant species as well as in fungi and metazoa (Ride et al., 1999; Rajasekar et al., 2019). Over 90 SPHs have been identified in Arabidopsis (Rajasekar et al., 2019). Based on the large number of SPH family members, all encoding proteins with signal peptides, together with their wide distribution, it has previously been proposed that they may be ligands involved in a wide range of signaling pathways (Ride et al., 1999). It has been suggested that this family of proteins may have evolved to act as a versatile and stable scaffold to display a variety of peptides in the predicted extracellular loops, each interacting with a different receptor (Rajasekar et al., 2019). Our findings here, showing that PrsS can trigger responses in vegetative tissues, provide further hints that (depending on how they have evolved) perhaps other SPHs may be involved in signaling in different tissues.

PrsS and SPHs are members of the CRPs, which include the Brassica spp. pollen S-determinant SCR/SP11 (Schopfer et al., 1999; Takayama et al., 2000), defensins (Biurchedner and Dresselhaus, 2016), LUREs (Okuda et al., 2009; Takeuchi and Higashiyama, 2016), and rapid alkalinization factors (RALFs; Pearce et al., 2001; Li and Yang, 2018), which are known to interact with receptors to activate diverse signaling networks involved in plant growth, defense, and reproduction (Wheeler et al., 2010; Marshall et al., 2011; Takeuchi and Higashiyama, 2016; Liu et al., 2017). Although comparatively few secreted peptides have been shown to interact with receptors in plants, genome analysis has revealed the existence of hundreds of predicted secreted proteins that may act as ligands (Lease and Walker, 2006). It has been suggested that CRPs have diversified for a huge variety of specialized functions (Manners, 2007; Silverstein et al., 2007; Bircheneder and Dresselhaus, 2016); rapid evolution from an origin in plant defense to regulate plant reproduction has been proposed (Bircheneder and Dresselhaus, 2016). Analysis of Arabidopsis SPH genes in the available databases reveal that they are mainly, but not exclusively, expressed in reproductive tissues (Supplemental Figs. S13 and S14; Mergner et al., 2020). Notably, several SPHs are expressed in silique septum, silique valves, flower pedicels, and senescent leaves, which all undergo PCD in various cellular/developmental contexts (Beers, 1997; Gómez et al., 2014). This hints that this family may have evolved a general function in several diverse tissues to signal to growth and PCD, as we have found for PrsS in this study.

Examination of the literature and databases reveals that no functional data are currently available for any Arabidopsis SPHs. Nevertheless, association networks for one of the SPH genes, AT1G51250, using STRING analysis (Szklarczyk et al., 2019), for example, reveals associations/putative interactions with several proteins. These include APPB1 and AT4G02250, which are plant pectin methylesterase inhibitor proteins, implicated in mediating growth; RALFL8, RALFL15, and RALFL26 (RALF-like cell signaling peptides), implicated in regulating plant stress, growth, and development; and LCR72, a Cys-rich peptide, predicted to encode a PR protein, that itself interacts with other defensins. These interactions hint that this SPH may signal to regulate growth and stress response. As both RALFs and pectin methylesterase inhibitors are broadly expressed (Supplemental Figs. S15 and S16; Mergner et al., 2020), this suggests that some SPHs may also potentially interact with these proteins to mediate these responses in various tissues. However, to our knowledge, no studies to date have identified a function for any SPH in another tissue, and as no other partners for SPHs have been identified to date, we cannot speculate much further about the possible functions of putative homologs of SPHs or their putative interactors, which is currently a black box. Although PrpS, being a small transmembrane protein with no known homologs, is not a receptor in the classic sense, our findings here, showing that the PrpS-PrsS module can act as a receptor-ligand-like module outside its usual reproductive context in vegetative tissues, provide a rare example of a specialized bipartite gene module that can act in a cell-autonomous manner. Thus, our finding that PrpS-PrsS can function in vegetative tissues, together with information on SPH homologs and their possible interactors, may provide clues about how the SPHs might potentially have coevolved to function in different cell types, an interesting avenue to be explored in the future.

Although the downstream cellular responses observed here in Arabidopsis roots in response to PrsS are strikingly similar to what was observed in P. rhoeas pollen tubes during the SI response (Wilkins et al., 2014;
Our evidence that the PrpS-PrsS module can also inhibit diffuse growth and does not apparently distinguish between these two types of growth supports this concept. Moreover, it is of interest that the peptides of several other CRP members function to regulate different types of growth. For example, RALFs are involved in the arrest of root growth and development (Pearce et al., 2001; Haruta et al., 2014; Blackburn et al., 2020), LUREs (specifically expressed in synergid cells) act to control directional growth of pollen tubes to the embryo sac (Okuda et al., 2009), SCR/SP11 act as the male S-determinant in Brassica spp. to inhibit self pollen (Schopfer et al., 1999; Takayama et al., 2000), and ZmES4 induces pollen tube growth arrest and bursting to release sperm cells during fertilization (Amien et al., 2010; for review, see Kanaoka and Higashiyama, 2015; Higashiyama and Yang, 2017; Blackburn et al., 2020). Further studies are needed to determine if there is a common growth-arrest mechanism triggered by these different CRP-mediated signaling pathways. Moreover, it would be of considerable interest to investigate if PrsS interacts with RALFs as a putative candidate player in the SI signaling pathway in pollen, as they are involved in signaling via reactive oxygen species to inhibit primary root elongation (Haruta et al., 2014) and it has been established that reactive oxygen species are involved in the SI-PCD response in pollen (Wilkins et al., 2011).

We previously showed that PrpS and PrsS could function to mediate SI and PCD in Arabidopsis pollen, despite the fact that this species is self-compatible (de Graaf et al., 2012; Lin et al., 2015). We proposed that the P. rheaes SI system worked in Arabidopsis pollen because it could recruit existing proteins to form new signaling networks, by multitasking of endogenous components that can act in signaling networks that they do not normally operate in, to provide a specific, predictable physiological outcome. This successful transfer between species suggested that the signaling network and cellular targets downstream of the PrpS-PrsS interaction might be present in a wide range of angiosperm species (de Graaf et al., 2012), as this was the simplest explanation of why these genes work in such an evolutionarily diverged (more than 100 million years; Bell et al., 2010) species. However, we did not explore whether this might extend beyond the particular context of pollen involved in the SI response. Here, we have extended our studies to show that this pair of genes can also act in other cell types in Arabidopsis. Our findings here, showing that this module can trigger growth arrest and PCD in sporophytic vegetative cells, provide firm evidence for this idea of plug and play and extends it, by showing that PrpS-PrsS can also act in an ectopic situation to trigger a signaling network and response that appears to be common and ubiquitously expressed and not just restricted to pollen. As key components can be harnessed in different cell types to reconstitute key P. rheaes SI-PCD-like phenomena in vegetative cells, this provides hints about the functional diversification and recruitment of preexisting components and the plasticity of cell signaling downstream of the PrpS-PrsS interaction leading to growth arrest and PCD in plant cells.

Our study has substantially extended previous studies (de Graaf et al., 2012; Lin et al., 2015) and reveals that the events downstream of the P. rheaes PrpS-PrsS interaction can be triggered in different cell types of Arabidopsis. This lays the foundations for new opportunities to elucidate key mechanisms triggered by cognate PrpS-PrsS interactions. Although the P. rheaes SI system has provided an excellent model system to investigate cell-cell recognition, intracellular signaling, and PCD at the molecular level, the extremely limited genetic resources in this system have provided an obstacle to progress, as certain approaches were not possible. Our findings here suggest that Arabidopsis plants express an SI-like response in vegetative tissues with all the key features of P. rheaes SI, opening up new opportunities to genetically dissect the signaling networks involved. Expression of the bipartite PrpS-PrsS module in different tissues has the potential to be applied to devise biochemical or genetic approaches to search for downstream components. Using this system in vegetative tissue or whole plants has the advantage that it overcomes the bottleneck that many reproductive researchers are faced with (i.e. that of limited material), as collecting sufficient pollen at the correct developmental stage is laborious, time-consuming, and difficult to scale up. Being able to perform experiments on bulk plant tissue or on whole plants could allow us to identify new genes/proteins involved in the downstream pathway; these could then be examined and validated in pollen to establish if they authentically play a role in the SI response. For example, root growth assays could provide a simple assay for screening large sets of T-DNA mutants or chemical library screening. Biochemical approaches, such as purification of candidate proteins or profiling of PrpS-PrsS-induced metabolomic changes using pollen, are generally impossible due to the small amount of tissue available. Using a heterologous expression system to enable a bulk purification, from leaves or roots, for example, of putative proteases with caspase-like activities or actin-binding proteins implicated in the actin remodeling might be possible. In conclusion, this ectopic Arabidopsis self-recognition:self-destruct system will allow us to test new hypotheses about the cellular mechanisms and genetic components involved in the SI-PCD response and tip growth of plant cells in the future.

**MATERIALS AND METHODS**

**Plant Material and Growth Conditions**

Arabidopsis (Arabidopsis thaliana) Col-0 seeds were gas sterilized, sown out on LRC2 plates (2.15 g L\(^{-1}\) Murashige and Skoog medium basal salts [Duchefa
Cloning and Transgenic Lines

All the expression vectors were generated using either Gateway cloning (Invitrogen) or GreenGate cloning (Lampropoulos et al., 2013). High-fidelity Phusion DNA polymerase (New England Biolabs) was used for all the DNA fragment amplification.

The expression clones pUBQ10::PrpS1 were obtained using Gateway cloning (Invitrogen). PrpS1 gDNA was amplified using primers F-attB1-PrpS1/R-attB2-PrpS1 with gDNA of line BG16 (de Graaf et al., 2012) as template. The resulting PCR fragments were cloned into pDONR221 using BP clonase (Invitrogen) to obtain pEN-L4-pRPS5A::XVE-R1. The entry vector pEN-L4-pRPS5A::XVE-R1 was obtained from the PSB Gateway Vector collection (Fendrych et al., 2014). These entry clones were recombined into Gateway destination vector pB7m24GW (Karimi et al., 2002) using LR Clonase II plus enzyme (Invitrogen) to obtain the expression clone pUBQ10::PrpS1.

The expression clones pH3.3::XVE::PrpS1/3 were generated using Gateway cloning (Invitrogen). The DNA fragment of the H3.3 promoter was amplified using Kpn1 and XhoI restriction enzymes followed by DNA gel purification. Plasmid pEN-L4-pRPS5A::XVE-R1 (Huysmans et al., 2018) was digested using the same restriction enzymes, followed by DNA gel electrophoresis. The vector backbone without the RPS5A promoter was cut out and purified. The Kpn1-HF3.3-XhoI DNA fragment was ligated into the linearized vector backbone to generate pENT-L4-pH3.3::XVE-R1. PrpS1/3 gDNA was amplified using primer sets F-attB1-PrpS1/3-R-attB2-PrpS1 with plastid plR1::PrpS1/3 (Lin et al., 2013) as template. The resulting PCR fragments were cloned into pDONR221 using BP clonase (Invitrogen) to obtain pEN-L1-PrpS1-L2. The entry vector pEN-L4-pRPS5A::XVE-R1 was obtained from the PSB Gateway Vector collection (Fendrych et al., 2014). These entry clones were recombined into Gateway destination vector pB7m24GW using LR Clonase II plus enzyme (Invitrogen) to obtain the expression clone pH3.3::XVE::PrpS1/3.

The dual-expression clones pUBQ10::PrpS1_pUBQ10::PrpS1/3 were generated using GreenGate cloning. Promoter UBQ10 was amplified using primer sets F-attB1-PrpS1_pUBQ10/R-attB2-UBQ10 and F-UBQ10/R-UBQ10 with entry vector pEN-L4-pUBQ10_R1 as template. The resulting PCR fragments were cloned into pDONR221 using BP clonase (Invitrogen) to obtain the entry vector pEN-C-tRBCS-D, pEN-E-NLS-YC3.6-F, pEN-F-YC3.6 to obtain the dual-expression clone pH3.3::XVE::PrpS1_pUBQ10::PrpS1/3 using the Gateway cloning Kit (Thermo Fisher) to obtain the entry vectors pH3.3::XVE_YC3.6-F and pEN-D-pUBQ10::R-E. Similarly, pEN-B-PrpS1-pRPS5A::XVE::PrpS1/3 was generated by cloning of the PrpS1 DNA fragment amplified using primers F-B-PrpS1/R-R-C-PrpS1 into pET12 using the ClonetEC PCR Cloning Kit (Thermo Fisher) to obtain the entry vectors pEN-A-pUBQ10::B and pEN-D-pUBQ10::B. The expression vectors were transformed into E. coli Transetta DH5α (Novagen) and stored in −80°C. PrpS1 proteins were dialyzed in one-fifth strength LRC2 medium liquid medium overnight in 4°C before use. The concentration of PrpS1 proteins was determined using the Bradford assay (Bio-Rad), during which the standard curve was generated using BSA (Sigma-Aldrich). To examine the effect of PrpS1 proteins on seedling growth, 10 μL of PrpS1 proteins with the desired concentration (the PrpS1 protein concentration used in all the experiments was 10 ng μL−1, unless specified) needed for different experiments was added to the root tip of each seedling using a pipette on LRC2 plates. The plates were kept horizontally for 30 min to allow PrpS1 proteins to dry before being placed back in the growth chamber vertically. When the PrpS1 protein treatment was needed during live-cell imaging, a perfusion chamber system was adopted. These leaves were mounted and treated as described (Krebs and Schumacher, 2013) with minor modifications: instead of cotton, glass wool was used, and one-half-strength Murashige and Skoog medium was replaced with one-fifth-strength LRC2 solution. The leaves were imaged with a confocal laser scanning microscope (Bio-Rad). The laser power was adjusted to optimize signal-to-noise ratio, and the exposure time was set to 500 ms. Images were collected with a 40× objective. The images were analyzed using ImageJ (National Institutes of Health). The threshold was set at 0.2, and the intensity of the signal was normalized. The data were presented as mean ± SEM from three independent experiments. The Student’s t test was performed using GraphPad Prism 7.0. The significance level was set at P < 0.05.

Protoplast Preparation

Leaves from 4-week-old plants were harvested and the lower epidermis was removed using double-sided tape as described (Wu et al., 2009). These leaf samples were immediately transferred into a petri dish containing protoplast medium (1% [w/v] cellulose R10 Yakult and 0.25% [w/v] macerozyme R10 Yakult) in protoplast washing solution (0.4 M sorbitol, 10 mM CaCl2, 20 mM KCl, 0.1% [w/v] BSA, and 20 mM MES, pH 5.7 adjusted using KOH). Samples were incubated at room temperature with light on a orbital shaker set to 40 rpm for up to 2 h, followed by gentle filtration using a 70-μm cell strainer into a 50-mL tube. Protoplasts were washed three times with the protoplast washing solution by centrifuging at 100g for 5 min and aspirating off the supernatant, followed by suspension in protoplast washing solution, before being subjected to PrpS1 treatment. PrpS1 proteins used for protoplast treatment were dialyzed in protoplast washing solution without BSA overnight in 4°C. BSA was added back to the protoplast washing solution after PrpS1 protein concentration determination using the Bradford assay (Bio-Rad). PrpS1 protein treatment for protoplasts was carried out on a 12-well tissue culture plate. PrpS1 proteins were added to the protoplasts directly to a final concentration 10 ng mL−1. Protoplast washing solution was added as a mock control. During treatments, plates were placed in the Arabidopsis growth chamber with continuous light emitted by white fluorescent lamps (intensity of 120 μmol m−2 s−1) at 22°C. Fifty micro-liters of protoplast samples was taken from the plate at 0 h (before treatment) and at 1, 2, 4, and 8 h for viability examination and confocal imaging.
Estradiol Induction

β-Estradiol (Sigma-Aldrich) was dissolved in pure ethanol, and a 10 mM stock solution was prepared. The stock solution was stored in −20°C for up to 1 month. Four-day-old seedlings grown on LRC2 plates were transferred onto LRC2 plates containing estradiol (10 μM) or 0.1% (v/v) ethanol (mock control) for specified periods of time according to different experiments.

DEVDa ce Activity Assay

The CV-Caspase3&7 detection Kit (Enzo Life Science) was used for measuring the DEVDa ce activities of seedlings after PrpS protein treatment. DEVDa ce activity probe CR(DEVD)2 powder was reconstituted using 100 μL of dimethyl sulfoxide to obtain CR(DEVD)2 stock solution and kept in −20°C if not utilized immediately. Before use, the stock solution was diluted 1:5 in MilliQ water to make the staining solution. The working solution was made by further diluting the staining solution 1:20 in one-fifth-strength LRC2 solution. Samples were incubated in the working solution for 1 h at room temperature before imaging.

Imaging, Image Analysis, and Figure Preparation

Imaging of the root calcium signature was performed using a Zeiss LSM710 microscope using a PlanApochromat 20× objective (numerical aperture 0.8). YC3.6 or NLS-YC3.6 was excited with 405 nm, and fluorescence emissions of 460 to 515 nm and 515 to 570 nm were collected for eCFP and cpVenus, respectively. When PI staining was performed in conjunction with NLS-YC3.6 signal acquisition, seedling samples were mounted with one-fifth-strength LRC2 medium containing 5 μg mL⁻¹ PI. A new imaging track was set up for PI signal acquisition. PI was excited with 561 nm, and fluorescence emissions between 580 and 700 nm were collected.

Imaging of the root pHGFP signal was performed using a Zeiss LSM710 microscope using a PlanApochromat 20× objective (numerical aperture 0.8). pHGFP was excited with 405 and 488 nm, and fluorescence emissions between 495 and 545 nm were collected.

LifeAct visualization was acquired using a Leica SP8 confocal laser scanning system with HCPL APO CS2 40×/1.0 (water) objective and HyD detector. LifeAct-mRuby2 was excited with 559 nm, and fluorescence emissions between 570 and 700 nm were collected.

DEVDa ce activities were visualized using a Leica SP8 confocal laser scanning system with FluoStar VISIR 25×/0.95 (water) objective and HyD detector. Samples were excited with 592 nm, and fluorescence emissions between 610 and 690 nm were collected.

All the images were processed and analyzed using Fiji. To quantify the calcium signal from NLS-YC3.6 or YC3.6 images, selected regions of interest were extracted using Fiji for both the eCFP and cpVenus channels. Fractional ratio changes (ΔR/R) were calculated as (R – R₀)/R₀, where R₀ is the average ratio of the first 5 min (15 frames) of each measurement.

Statistical Analysis

Statistical analysis was performed using GraphPad Prism 8.0 for Windows.

Accession Numbers

Sequence data from this article can be found in the GenBank/EMBL data libraries under the following accession numbers: Actin2 (At3g18780), UBQ10 (At4g05320), and H3.3 (At4g40040).

Supplemental Data

The following supplemental materials are available.

Supplemental Figure S1. RT-qPCR shows that expression of PrpS is mRNA varies in different pUBQ10::PrpS lines.

Supplemental Figure S2. Treatment with PrpS proteins does not inhibit the growth of wild-type seedlings.

Supplemental Figure S3. Recombinant PrpS protein treatment triggers rapid root growth inhibition of PrpS-expressing seedlings in an S-specific manner.

Supplemental Figure S4. PrpS treatment results in S-specific cell death of PrpS-expressing seedling root cells.

Supplemental Figure S5. PrpS treatment results in S-specific activation of DEVDa ce in PrpS-expressing seedling roots.

Supplemental Figure S6. PrpS treatment triggers S-specific alterations in [Ca²⁺]cyt.

Supplemental Figure S7. PrpS treatment triggers nucleus-localized Ca²⁺ changes in an S-specific manner.

Supplemental Figure S8. PrpS treatment triggers S-specific cytosolic pH decreases in PrpS-expressing roots.

Supplemental Figure S9. PrpS treatment triggers S-specific formation of actin foci in PrpS-expressing roots.

Supplemental Figure S10. Coexpression of PrpS and Prps in Arabidopsis triggers root growth inhibition in an S-specific manner.

Supplemental Figure S11. Estradiol treatment induces the expression of Prps mRNA transcript in XVE-PrpS/PrpS lines.

Supplemental Figure S12. Coexpression of PrpS and Prps in whole Arabidopsis plants using estradiol triggers S-specific cell death in whole seedling roots.

Supplemental Figure S13. Transcript expression patterns of the SPH genes in Arabidopsis tissues.

Supplemental Figure S14. Protein expression patterns of the SPH genes in Arabidopsis tissues.

Supplemental Figure S15. Expression patterns of RALFs in Arabidopsis tissues.

Supplemental Figure S16. Expression patterns of pectin methylesterase inhibitors in Arabidopsis tissues.

Supplemental Table S1. Primers for vector construction and mRNA detection.

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Supplemental Figure S17. PrpS treatment triggers nucleus-localized Ca²⁺ changes in an S-specific manner.

Supplemental Figure S18. PrpS treatment triggers S-specific cytosolic pH decreases in PrpS-expressing roots.

Supplemental Figure S19. PrpS treatment triggers S-specific formation of actin foci in PrpS-expressing roots.

Supplemental Figure S20. Coexpression of PrpS and Prps in Arabidopsis triggers root growth inhibition in an S-specific manner.

Supplemental Figure S21. Expression patterns of RALFs in Arabidopsis tissues.

Supplemental Figure S22. Expression patterns of pectin methylesterase inhibitors in Arabidopsis tissues.

Supplemental Table S1. Primers for vector construction and mRNA detection.
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