



A species-specific functional module controls formation of pollen apertures

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Pollen apertures are an interesting model for the formation of specialized plasma-membrane domains. The plant-specific protein INP1 serves as a key aperture factor in such distantly related species as *Arabidopsis*, rice and maize. Although INP1 orthologues probably play similar roles throughout flowering plants, they show substantial sequence divergence and often cannot substitute for each other, suggesting that INP1 might require species-specific partners. Here, we present a new aperture factor, INP2, which satisfies the criteria for being a species-specific partner for INP1. Both INP proteins display similar structural features, including the plant-specific DOG1 domain, similar patterns of expression and mutant phenotypes, as well as signs of co-evolution. These proteins interact with each other in a species-specific manner and can restore apertures in a heterologous system when both are expressed but not when expressed individually. Our findings suggest that the INP proteins form a species-specific functional module that underlies formation of pollen apertures.

Pollen grains of flowering plants are surrounded by a robust wall, called exine. In most species, exine is deposited on the pollen surface non-uniformly, with certain regions of the surface receiving little to no exine material¹. These regions develop into pollen apertures that help pollen to hydrate, change volume and germinate²⁻⁶. Across species, apertures vary greatly in their number, positions and morphology, contributing to diverse, species-specific patterns on the pollen surface^{1,2,7,8}. Recently, we and others have demonstrated that before forming apertures, developing pollen forms distinct aperture domains in their plasma membrane, which accumulate specific combinations of proteins and lipids^{6,9-11}. Apertures can thus be used to study how cells develop polarity and form membrane domains, as well as to understand how these mechanisms evolved to create the tremendous diversity of aperture patterns found in nature.

Aperture domains of the plasma membrane appear at the tetrad stage of pollen development, during which the four products of male meiosis (microspores) are transiently kept together under the common callose wall^{6,10}. The positions, number and morphology of the aperture membrane domains in microspores correspond to the aperture pattern of mature pollen. For example, in *Arabidopsis* pollen, apertures are shaped like three long and narrow meridional furrows (Fig. 1a). Accordingly, in *Arabidopsis* tetrads, each microspore develops three linear meridional domains of the plasma membrane, which attract the proteins D6 PROTEIN KINASE-LIKE3 (D6PKL3) and INAPERTURATE POLLEN1 (INP1)^{10,11}. In contrast, in rice and other grasses, pollen has only one small round aperture, positioned at the distal pole. Correspondingly, the tetrad-stage microspores in rice develop at their distal poles a single aperture domain shaped like a tiny ring, which attracts the rice orthologue of INP1 (OsINP1)⁶.

In both *Arabidopsis* and rice, as well as in maize, the INP1 protein was shown to be a major aperture factor whose loss causes a

complete loss of apertures (Fig. 1b)^{6,9,12}. INP1 is a plant-specific protein with a single recognizable domain, the DELAYED IN GERMINATION1 (DOG1) domain, whose function is unknown⁹. Although the biochemical function of INP1 remains to be identified, in both *Arabidopsis* and rice these proteins appear to play a role in keeping the aperture domains of the plasma membrane in close contact with the overlying callose wall^{6,10}, which possibly protects these domains from the deposition of exine materials.

Since the role of INP1 as an essential aperture factor is conserved in such distantly related species as *Arabidopsis*, rice and maize, it is reasonable to assume that INP1 orthologues across angiosperms are probably all involved in aperture formation. Intriguingly, though, many INP1 proteins show substantial sequence divergence and cannot substitute for the loss of *Arabidopsis* INP1 (refs. 9,12). This suggests that, despite their conserved involvement in aperture formation, INP1 proteins are probably functionally species-specific. We have previously proposed that such species specificity might be due to the presence of unknown aperture factors that have co-evolved with INP1 and help it to perform its function¹².

Here, we present an aperture factor, INP2, that fulfills the role of a species-specific partner for INP1. INP2 resembles INP1 in its protein structure, patterns of expression, trends of evolutionary divergence, mutant phenotype and genetic interactions. We provide evidence that INP2 is also functionally species-specific and that it physically interacts with INP1. Furthermore, we demonstrate that tomato orthologues of INP1 and INP2, which are unable to restore apertures in *Arabidopsis* mutants when only one of them is expressed, gain the ability to function in *Arabidopsis* when expressed together. The two INP proteins, therefore, behave as co-evolved species-specific partners that form a functional module required for the formation of pollen apertures.

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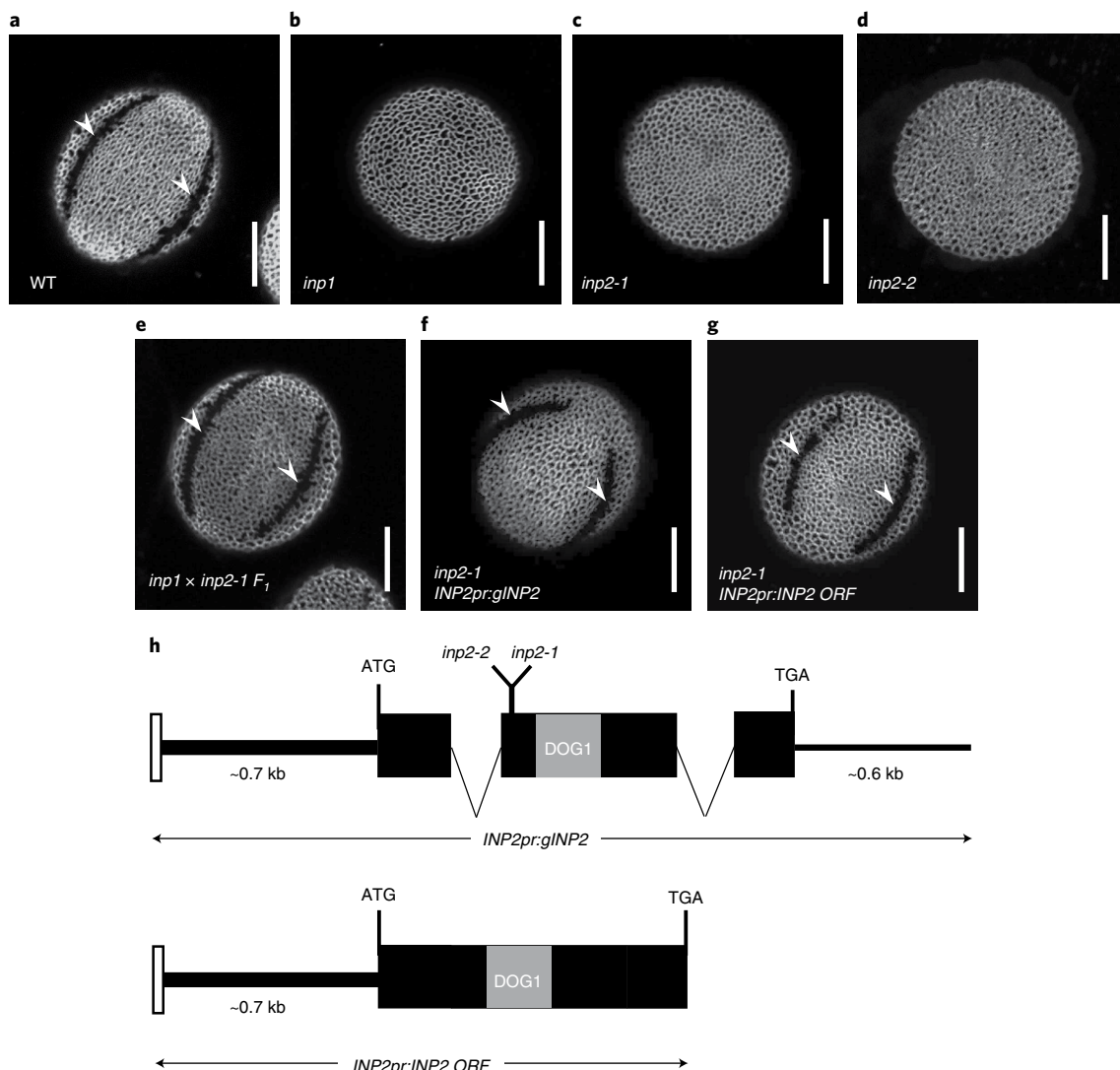


Fig. 1 | INP2 is a new factor essential for the formation of pollen apertures. **a–g**, Confocal images of pollen grains stained with auramine O. Scale bars, 10 μ m. **a**, Wild-type (WT) *Arabidopsis* pollen has three equidistant furrow-like apertures (two are visible here, arrowheads). **b**, *inp1* pollen completely lacks apertures. **c,d**, Similar to *inp1*, pollen of *inp2-1* (**c**) and *inp2-2* (**d**) mutants has normal exine but completely lacks apertures (>100 pollen grains were imaged, with similar results; for *inp2-2*, two independent CRISPR plants were obtained, producing similar phenotypes). **e**, Pollen of the F_1 progeny of the cross between *inp1* and *inp2* develops normal apertures (arrowheads), indicating that mutations disrupt different genes (eight plants (\geq 50 pollen grains per plant) were imaged, with similar results). **f,g**, *INP2pr:gINP2* (**f**) and *INP2pr:INP2 ORF* (**g**) transgenes restore apertures (arrowheads) in *inp2* (7/8 and 15/15 independent T_1 lines, respectively; \geq 50 pollen grains per line were imaged, with similar results). **h**, *INP2* gene model and structure of the *INP2pr:gINP2* and *INP2pr:INP2 ORF* complementation constructs. Black boxes indicate the protein-coding sequence of At1g15320. The region encoding the DOG1 domain is indicated by the grey box. The white box denotes a short region from the preceding gene, At1g15330, which was included in the constructs. Both the ~0.7-kb upstream region and the ~0.6-kb downstream region were included in the genomic construct. Positions of the *inp2-1* and *inp2-2* mutations are indicated on the gene model.

Results

A new *Arabidopsis* mutant has the inaperturate pollen phenotype identical to the phenotype of the *inp1* mutant. To discover genes involved in the formation of pollen apertures, we performed a forward genetic screen on an M_2 population of *Arabidopsis* plants mutagenized with ethyl methanesulfonate. Since changes in pollen shape can serve as a proxy for aperture formation defects^{13,14}, we screened these plants for unusual pollen shapes under dissecting microscopes. One mutant produced pollen that looked much rounder than the wild-type pollen, strongly resembling the phenotype of the *inp1* mutants. An examination by confocal microscopy showed that, like *inp1*, pollen of this mutant completely lacks apertures (inaperturate phenotype) but had otherwise normal exine (Fig. 1c).

To test whether the mutation represented an allele of *INP1* or disrupted another gene, we crossed the new mutant with the *inp1-1* null mutant. In the F_1 progeny of this cross, all pollen had normal apertures (Fig. 1e), demonstrating that the defect affected a gene other than *INP1*. This result also showed that, similar to *inp1* and other previously discovered aperture mutants, the new mutation affected a gene with the sporophytic function. Because of the similarities with the *inp1* mutant, we named the new gene *INAPERTURATE POLLEN2* (*INP2*) and its mutant allele *inp2-1*.

The *inp2-1* mutation disrupts the At1g15320 gene. Using positional cloning, we mapped the *inp2-1* defect to a 146-kilobase (kb) region at the top of chromosome 1, containing 51 genes. To

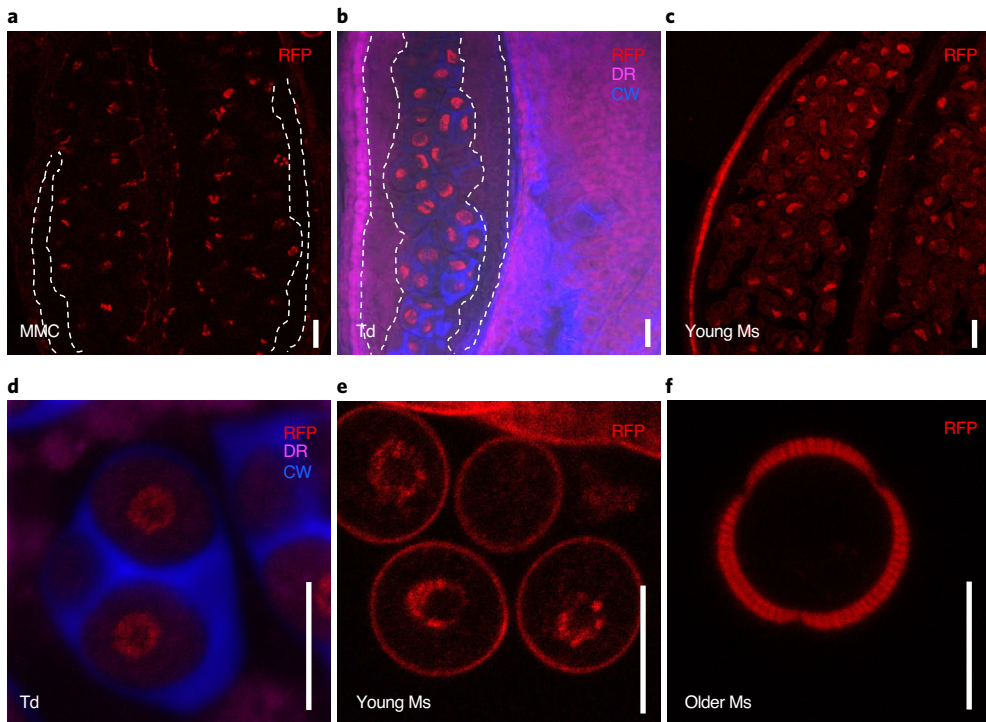


Fig. 2 | *INP2* is expressed in the male reproductive lineage at the time of aperture formation. Images of anthers at different developmental stages (a–c) and magnified images of the cells from the male reproductive lineage at different developmental stages (d–f) expressing the transcriptional fusion construct *INP2pr:H2B-RFP*. Nuclear signal of H2B-RFP (red) is found in dividing microspore mother cells (MMC; a), tetrads of microspores (Td; b,d) and young free microspores (Ms; c,e). Older microspores (f) do not show nuclear H2B-RFP signal (peripheral red signal is due to the autofluorescence of the developing exine). No signal was observed in the tapetal layer of the anther (outlined by the white dashed lines in a,b). Besides RFP, the images in b,d show staining for callose wall (blue, CW, calcofluor white) and membranous structures (magenta, DR, CellMask Deep Red). Five independent T_1 lines were imaged, with similar results. Scale bars, 10 μ m.

narrow down the list of gene candidates, we inspected their predicted identities as well as patterns of their messenger RNA expression reported in the TRAVA RNA-seq database¹⁵. We focused on the genes expressed in young flower buds (flowers 12–18 in the TRAVA nomenclature), as these buds include the tetrad stage of development associated with aperture formation. One gene, At1g15320, was prioritized as a particularly strong candidate as it is predicted to be expressed nearly exclusively in young buds (Extended Data Fig. 1) and encodes a protein with structural similarities to INP1 (below). Sequencing of this gene from *inp2-1* revealed a G-to-A substitution which created an early stop codon (Trp84Stop) (Fig. 1h). To independently confirm that *INP2* is At1g15320, we targeted At1g15320 in the wild-type Col-0 background with CRISPR-Cas9 and generated an allele (*inp2-2*) with a two-nucleotide deletion that caused a frame shift after the amino acid 83 (Fig. 1h). The CRISPR mutant displayed the same inaperturate pollen phenotype as the original *inp2-1* allele (Fig. 1d).

To further verify the identity of At1g15320 as *INP2* and to define its regulatory regions, we created transgenic constructs containing either the genomic region of At1g15320 (including introns and the ~0.6-kb region downstream of the stop codon) or its open reading frame (ORF) (Fig. 1h). These constructs were placed under the control of the putative native promoter (a region of ~0.7-kb between the start codon of At1g15320 and the preceding gene At1g15330) and transformed into *inp2-1*. Both constructs successfully restored apertures in transgenic plants—15/15 T_1 plants with the ORF construct and 7/8 T_1 plants with the genomic construct (Fig. 1f–g). Taken together, our results demonstrate that (1) At1g15320 encodes INP2 (a new factor essential for aperture formation) and (2) that the 0.7-kb upstream region is sufficient to drive functional expression

of *INP2*. This promoter region was then used for all subsequent *INP2* constructs transformed into *Arabidopsis*.

INP2 shares structural similarity with INP1. INP2 is a plant-specific protein of unknown biochemical function which shares certain similarities with INP1. Both proteins have similar size (273 amino acids for INP1 versus 307 amino acids for INP2), are usually encoded in angiosperm genomes by single-copy genes and contain the same domain—the plant-specific DELAYED IN GERMINATION1 (DOG1) domain (PFam14144) (Fig. 1 and Extended Data Fig. 2a). This domain, typically associated with seed dormancy proteins and TGA bZIP transcription factors¹⁶, is the only recognizable domain in both INP proteins. Interestingly, although INP1 and INP2 share only limited homology with each other (23% sequence identity; Extended Data Fig. 2a), the protein-fold recognition software Phyre2 (ref. ¹⁷) selected the same template for homology modeling of both proteins and predicted similar structures, with three alpha-helices, for their C-terminal regions (Extended Data Fig. 2b,c).

Protein alignments of INP1 and INP2 with their respective orthologues from other plants also revealed that, in eudicots, these proteins typically contain a region enriched in Asp and Glu residues. However, these acidic regions are positioned differently between INP1 and INP2. In the INP1 proteins, the acidic region follows the DOG1 domain^{9,12}, whereas in the INP2 proteins it is located ahead of the DOG1 domain (Extended Data Fig. 3).

For *Arabidopsis thaliana* INP2 (AtINP2), multiple algorithms also predicted the existence of a transmembrane (TM) domain at its N terminus (Extended Data Fig. 4), with most of the protein expected to be outside the cell, facing the extracellular space. Yet,

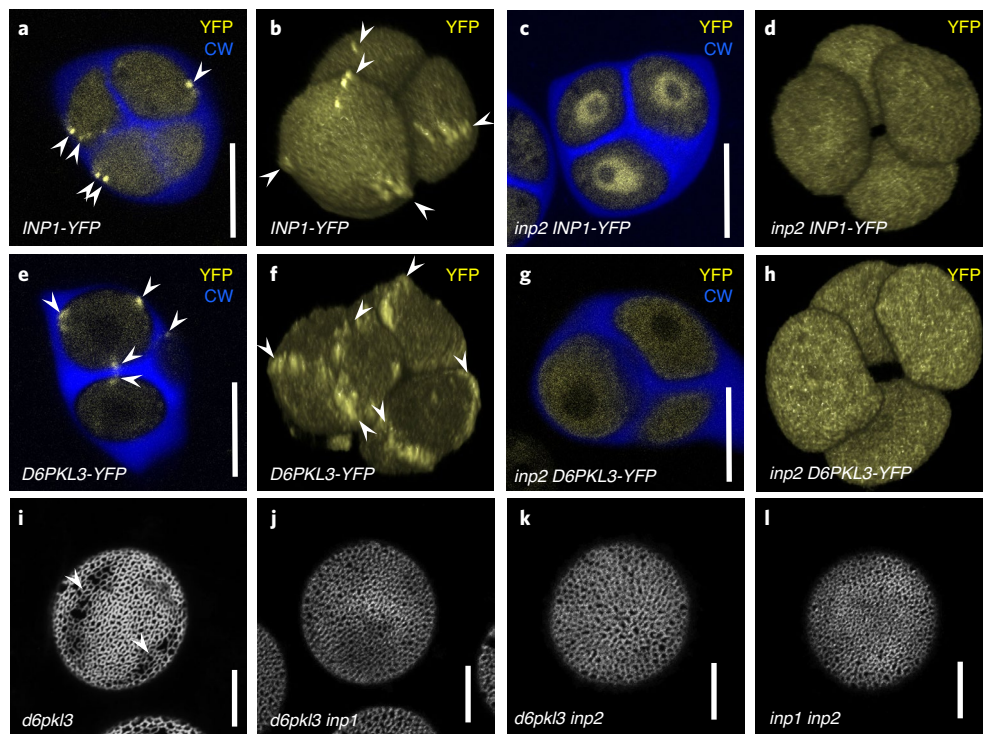


Fig. 3 | INP2 is required for INP1 and D6PKL3 accumulation at the aperture domains and both *inp1* and *inp2* are epistatic to *d6pk3*. **a–h**, INP1-YFP and D6PKL3-YFP localization in tetrads of microspores in the presence and absence of INP2. Confocal optical sections (**a, c, e** and **g**) and three-dimensional reconstructions of tetrads of microspores (**b, d, f** and **h**). YFP signal is shown in yellow and callose wall (CW, stained by calcofluor white) is shown in blue. Arrowheads point to the YFP signal at the aperture domains. INP1-YFP localizes to the aperture PM domains in the wild type (**a, b**) but loses this localization in the *inp2* mutant (**c, d**), instead becoming enriched in the nucleoplasm. Experiments in (**c, d**) were repeated three times, with similar results. Likewise, D6PKL3-YFP localizes to the aperture domains in the wild type (**e, f**) but loses this localization in the *inp2* mutant (**g, h**). Experiments in **g, h** were repeated two times, with similar results. **i–l**, *inp1* and *inp2* mutations are epistatic to *d6pk3* and do not cause additional phenotypic changes when combined. Confocal images of pollen grains stained with auramine O. The *d6pk3* mutant pollen often develops apertures partially covered with exine (arrowheads) (**i**), whereas double mutants *d6pk3 inp1* (**j**), *d6pk3 inp2* (**k**) and *inp1 inp2* (**l**) completely lack apertures. Three or more plants (≥ 50 pollen grains per plant) were imaged in **i–l**, with similar results. Scale bars, 10 μ m.

the algorithms failed to identify a TM domain in many orthologues of AtINP2, including the highly related proteins from *A. lyrata* and other members of the Brassicaceae family, suggesting that this is not a common feature of INP2 proteins. No lipid modifications are predicted for INP2.

INP2 is expressed in the developmental lineage of pollen at the time of aperture formation. Publicly available RNA-seq data show that, like INP1, INP2 is expressed nearly exclusively in young buds containing pollen at or around the tetrad stage during which apertures form (Extended Data Fig. 1). To test whether

in these buds INP2 is expressed in the male reproductive lineage, we expressed the nuclear marker histone H2B tagged with red fluorescent protein (RFP) under the control of the INP2 promoter (*INP2pr:H2B-RFP*) in the wild-type Col-0 plants. This reporter, with its concentrated localization in the nucleus, was specifically chosen to help visualize the expression from the INP2 promoter, since, like *INP1*, INP2 is predicted to be expressed at low levels (Extended Data Fig. 1). The nuclear RFP signal was found in the dividing microspore mother cells, tetrad-stage microspores and young free microspores (Fig. 2). The signal was absent in older microspores, the surrounding somatic tapetal cell layer and other

Fig. 4 | INP1 and INP2 physically interact. **a**, Yeast two-hybrid assay of interaction between INP1 and INP2^{ΔN} (lacking the N-terminal region). BD, DNA-binding domain; AD, activating domain; SD, synthetic defined medium. To test for the presence of both BD and AD constructs, leucine (L) and tryptophan (W) were excluded from the medium. To test for protein interaction, yeast were grown on media lacking L, W and histidine (H) and containing 20 mM 3-aminotriazole (3-AT). **b**, BiFC experiments. INP1 and INP2 proteins fused, respectively, to the N- and C-terminal parts of YFP (YN and YC) were cotransformed into tobacco leaves to test for interaction. Cotransformation of INP1-YN with only YC and cotransformation of INP2-YC with only YN were used as negative controls. Top panels show YFP signal in leaf epidermis. Bottom panels show merged YFP and bright-field images. Scale bars, 50 μ m. **c**, Co-immunoprecipitation experiments. INP1-HA₃/INP2-GFP and INP1-GFP/INP2-HA₃ pairs (or just single tagged proteins as negative controls) were co-expressed in tobacco leaves, precipitated with anti-GFP and visualized with anti-GFP or anti-HA. IP, immunoprecipitation; IB, immunoblot. ‘Mock’ indicates protein extract from leaves infiltrated only with buffer. **d**, Split-luciferase assay. Tobacco leaves were divided into sectors co-expressing indicated proteins containing the N-terminal (NLuc) and C-terminal (CLuc) parts of the firefly luciferase. Panels on the left show the bright-field images and panels on the right show the corresponding luminescence images. **e**, Y2H assay in which the DOG1 domains of INP1 (INP1^{DOG1}) and INP2 (INP2^{DOG1}) were tested for interaction with each other, self-interaction and interaction with the full-length INP1 and with INP2^{ΔN}. The description is the same as for **a**, except that 3 mM 3-AT was used here. Experiments in **a–c** and **e** were repeated three times and experiments in **d** were repeated two times (each time using multiple leaves from multiple plants), with similar results.

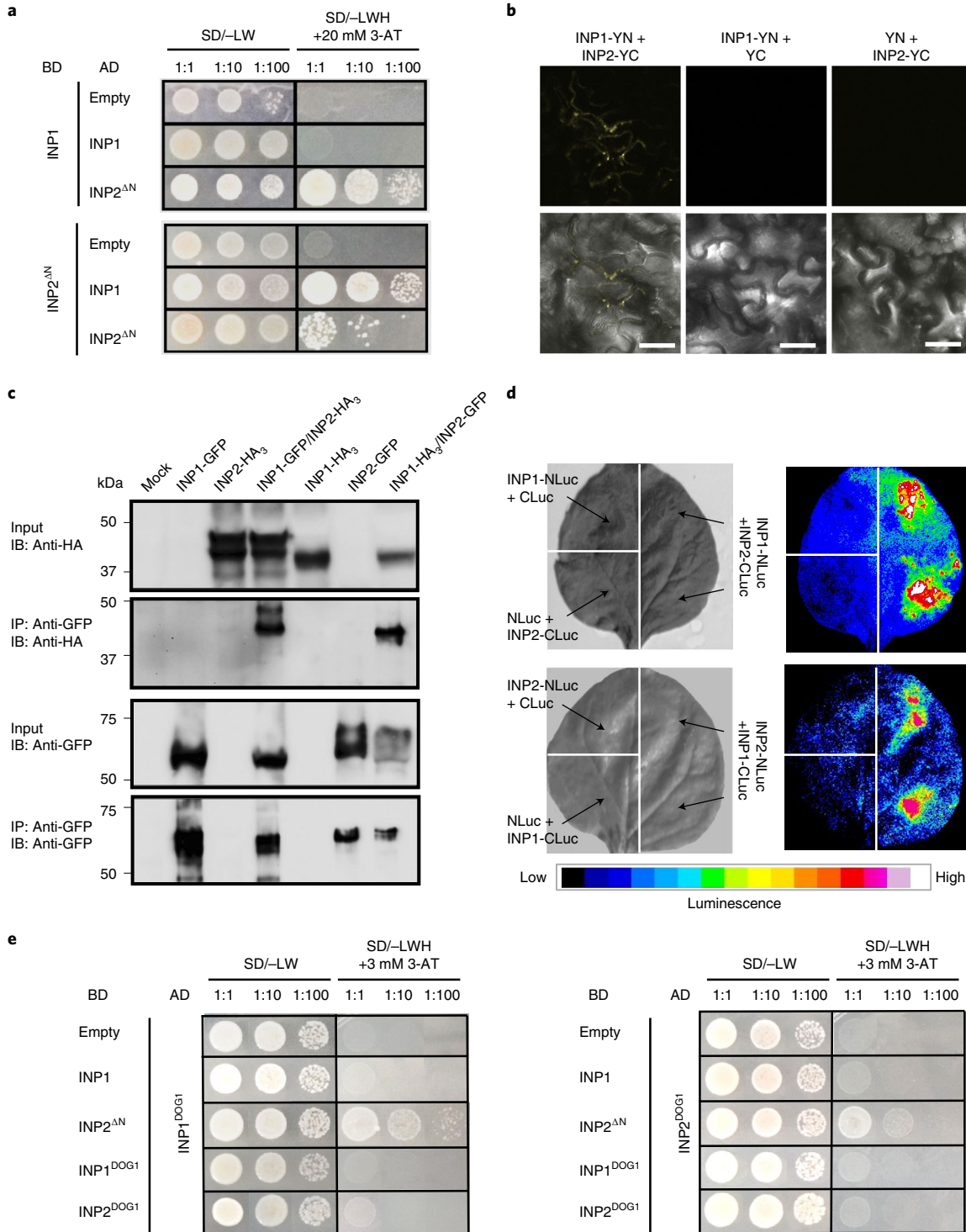
anther layers (Fig. 2). This expression pattern matches that of INP1 (refs. ^{9,10}).

To visualize the subcellular localization of the INP2 protein, we first created five constructs, in which INP2 was tagged with yellow fluorescent protein (YFP) at four positions: at the N terminus, at the C terminus (either directly or following an 18-amino acid linker), after the predicted TM region and internally—within the low-conservation region (below). However, none of the YFP-tagged constructs rescued the *inp2* mutant, suggesting that INP2 does not tolerate addition of sizable tags. This notion was supported by further experiments in which partial rescue of the mutant phenotype was

achieved with constructs expressing INP2 tagged at the C terminus with one or three copies of the small HA tag. Of these two types of constructs, the shorter HA₁ construct produced better rescue (Methods), yet no protein signal was detected in these lines with anti-HA in anther sections or whole-mount preparations, possibly owing to the low levels of the INP2 expression. This prevented us from determining whether INP2, like INP1, specifically localizes to the aperture domains in the plasma membrane of microspores.

Localization of INP1 and D6PKL3 to plasma-membrane aperture domains depends on the presence of INP2.

To test whether INP2



Tree scale: 1

Plant taxa

- Basal angiosperms
- Monocots—Arecales
- Monocots—Poales
- Basal eudicots
- Basal core eudicots
- Asterids—Asterales
- Rosids—Vitales
- Asterids—Gentianales
- Asterids—Ericales
- Asterids—Ariales
- Asterids—Solanales
- Asterids—Lamiaceae
- Rosids—Malvales
- Rosids—Fagales
- Rosids—Rosales
- Rosids—Malpighiales
- Rosids—Sapindales
- Rosids—Brassicaceae

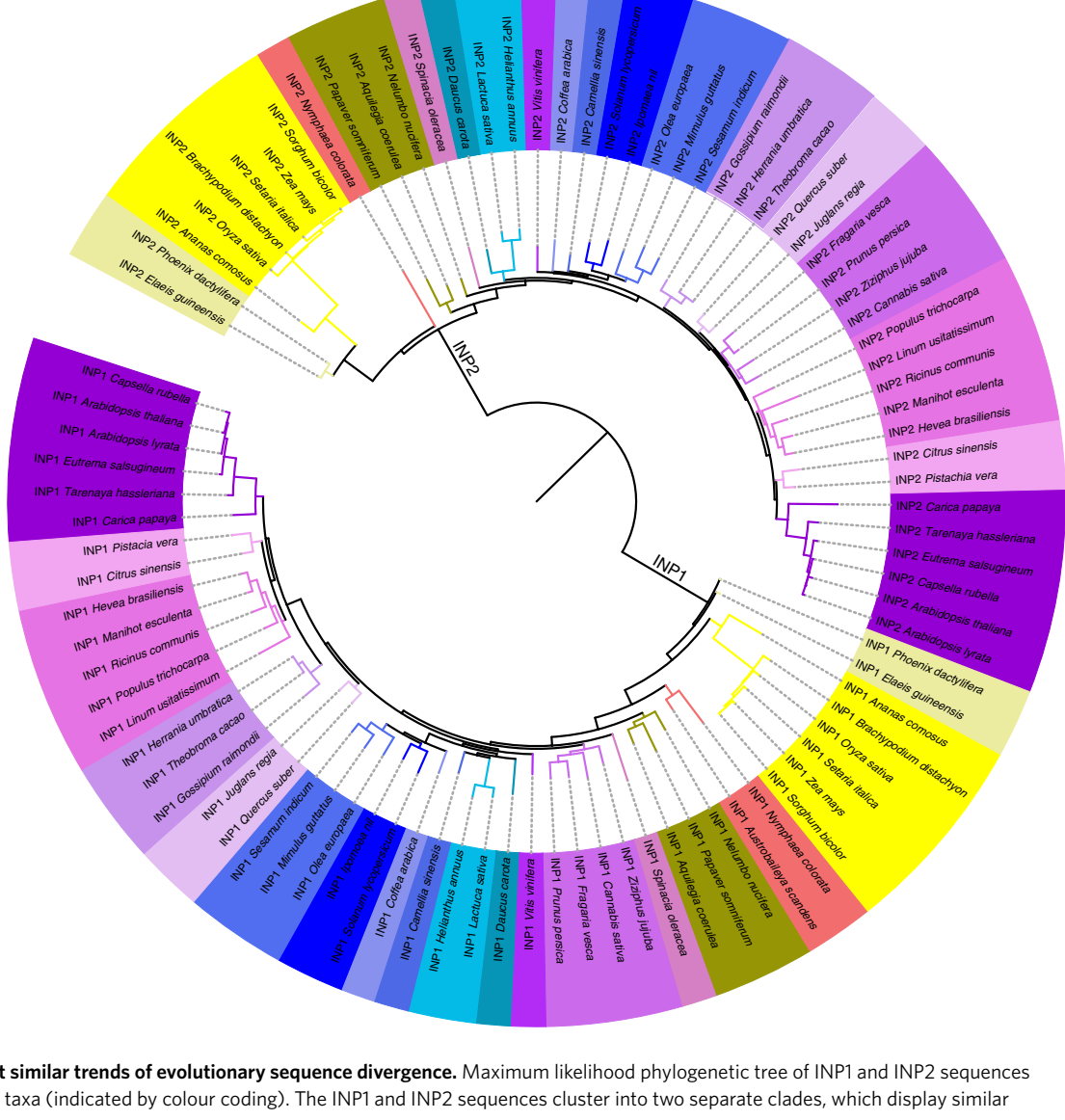


Fig. 5 | INP1 and INP2 exhibit similar trends of evolutionary sequence divergence. Maximum likelihood phylogenetic tree of INP1 and INP2 sequences from a variety of angiosperm taxa (indicated by colour coding). The INP1 and INP2 sequences cluster into two separate clades, which display similar topology.

contributes to the distinct positioning of INP1 and another recently identified aperture factor, *D6PKL3*, both of which accumulate at the microspore aperture domains (Fig. 3a,b,e,f), we introgressed the previously characterized reporter constructs *DMC1pr:INP1-YFP* (ref. ¹⁰) and *D6PKL3pr:D6PKL3-YFP* (ref. ¹¹) into the *inp2* mutant background. In the absence of INP2, INP1-YFP failed to localize to the aperture domains of the plasma membrane, instead showing notable enrichment in the nucleoplasm (Fig. 3c,d). This result suggests that INP2 is involved either in targeting INP1 to the aperture domains or in keeping it there. Likewise, in the absence of INP2, the membrane-associated kinase *D6PKL3-YFP* lost its association with the aperture domains, instead displaying diffuse cytoplasmic localization (Fig. 3g,h). As *D6PKL3* reacts the same way to the absence of INP1 (ref. ¹¹), both INP1 and INP2 are thus required to keep it at the aperture domains.

To test for epistatic relationships between these aperture factors, we created double mutants of *inp1 d6pk3* and *inp2 d6pk3*. Single mutations in *D6PKL3* do not completely abolish aperture formation, instead producing 'shadows of apertures' that are partially

covered with exine (Fig. 3i)¹¹. Yet both double mutants produced completely inaperturate pollen (Fig. 3j,k), indicating that *inp1* and *inp2* are both epistatic to *d6pk3*. To investigate the possibility of synergistic interactions between INP1 and INP2, we also created the *inp1 inp2* double mutant. Its phenotype, however, was identical to those of single mutants (Fig. 3l), showing that the simultaneous loss of INP1 and INP2 does not cause any additional observable effects (for example, in the exine deposition) and suggesting that these proteins behave as bona fide aperture factors. Taken together, the results presented so far are consistent with the notion that INP1 and INP2 occupy very similar positions in the aperture formation pathway and might coordinate their activities.

INP1 and INP2 are interacting proteins. Since INP1 and INP2 exhibit similarities in their protein structures, patterns of expression, mutant phenotypes and genetic interactions, we suggested that they might physically interact. To explore this possibility, we used several approaches. An initial yeast two-hybrid (Y2H) assay with the full-length INP1 and INP2 did not result in yeast growth

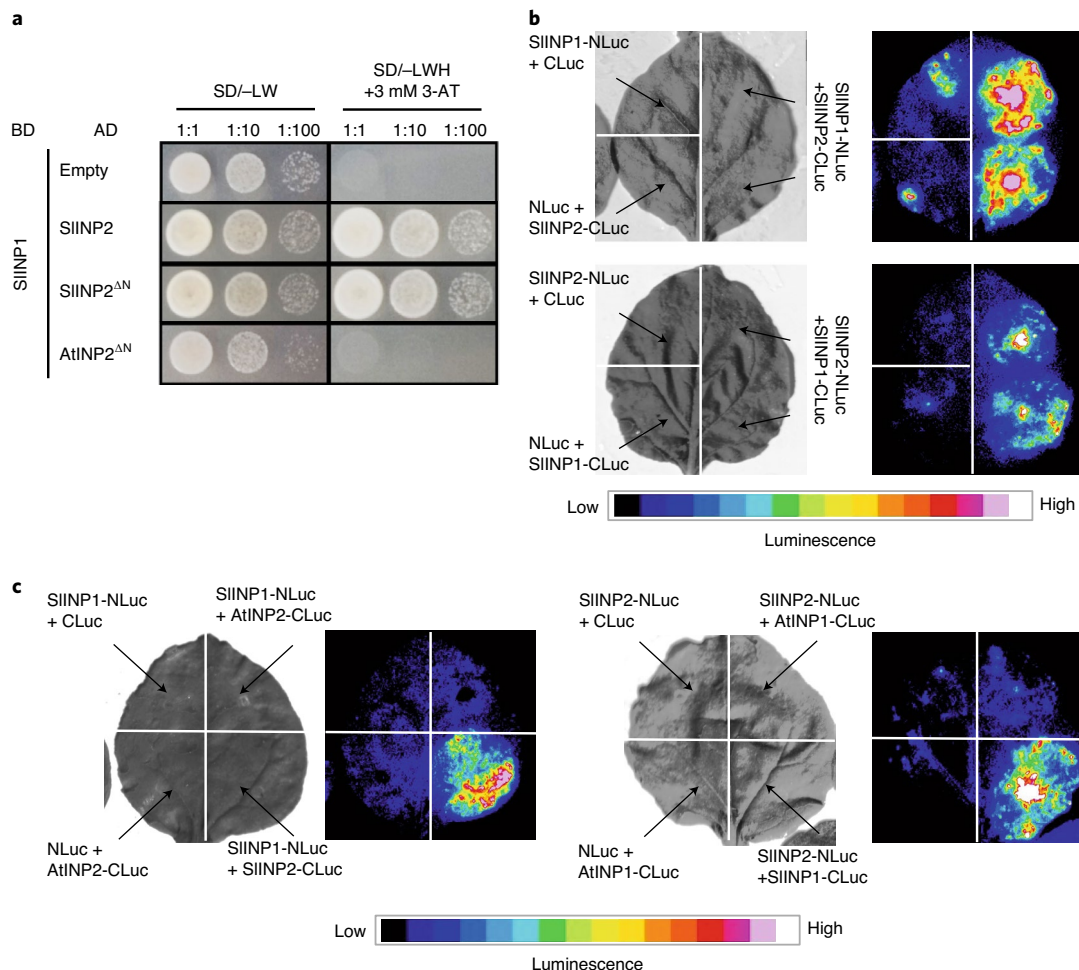


Fig. 6 | INP1 and INP2 interact in a species-specific manner. **a**, Y2H assay testing SIINP1 interactions with SIINP2 (or SIINP2^{ΔN} lacking the N-terminal region) and AtINP2^{ΔN}. To test for the presence of both BD and AD constructs, leucine (L) and tryptophan (W) were excluded from the medium. To test for protein interaction, yeast were grown on media lacking L, W and histidine (H) and containing 3 mM 3-aminotriazole (3-AT). **b**, Split-luciferase assay testing the ability of SIINP1 and SIINP2 to interact. Tobacco leaves were divided into sectors co-expressing indicated proteins containing the N-terminal (NLuc) and C-terminal (CLuc) parts of the firefly luciferase. Panels on the left show the bright-field images and panels on the right show the corresponding luminescence images. **c**, Split-luciferase assay testing the ability of INP1 and INP2 from *Arabidopsis* and tomato to interact with a protein from another species. Only the same-species interactions were observed. The description is the same as for **b**. All experiments were repeated at least twice, with similar results.

indicative of protein interaction. We reasoned, however, that lack of yeast growth would be expected if INP2 indeed had a TM domain at its N terminus and most of the protein was extracellular.

We, therefore, expressed INP2 in yeast without its first 24 amino acids, which contained the predicted TM domain. This truncated INP2 (INP2^{ΔN}) showed strong interaction with INP1 in the Y2H system (Fig. 4a). In addition, this assay revealed that INP2 may be able to self-interact (Fig. 4a). We further verified the ability of INP1 and INP2 to interact in planta by expressing them in tobacco leaf cells and performing co-immunoprecipitation, bifluorescent molecular complementation (BiFC) and a split-luciferase assay (Fig. 4b–d).

The DOG1 domain is the only recognizable protein domain present in these proteins. Although its function is unknown, it has been proposed that this domain might participate in protein–protein interactions¹⁸. We therefore used the Y2H assay to test the ability of the DOG1 domains from INP1 and INP2 to interact with each other and with the full-length (or nearly full-length in the case of INP2^{ΔN}) proteins (Fig. 4e). INP1^{DOG1} was able to interact with INP2^{ΔN}. In contrast, INP2^{DOG1} failed to interact with INP1 but showed some ability to interact with INP2^{ΔN}, consistent with the finding that INP2

may self-interact. However, no interactions occurred when only the DOG1 domains were present (Fig. 4e), suggesting that these regions probably interact with other portions of INP2.

INP1 and INP2 exhibit similar trends of evolutionary sequence divergence. We previously reported that INP1 greatly diversified in angiosperm lineages^{9,12}. Still, in several species these divergent orthologues were found to be involved in the formation of pollen apertures and able to localize to specific plasma-membrane aperture domains^{9,12}, suggesting that, despite the substantial difference in primary sequences, all INP1 proteins in angiosperms probably function as aperture factors. However, INP1 proteins appear to exhibit a notable degree of functional species specificity, since the divergent INP1 orthologues were not able to complement the aperture defects of the *Arabidopsis* *inp1* mutant¹². A possible interpretation of this result is that INP1 proteins might require the presence of co-evolved partners to perform their function.

To see whether INP2 shows signs of co-evolution with INP1, we performed BLAST searches for INP2 homologues followed by phylogenetic analysis, revealing notable parallels between INP1 and INP2. Although proteins with the DOG1 domain appeared as

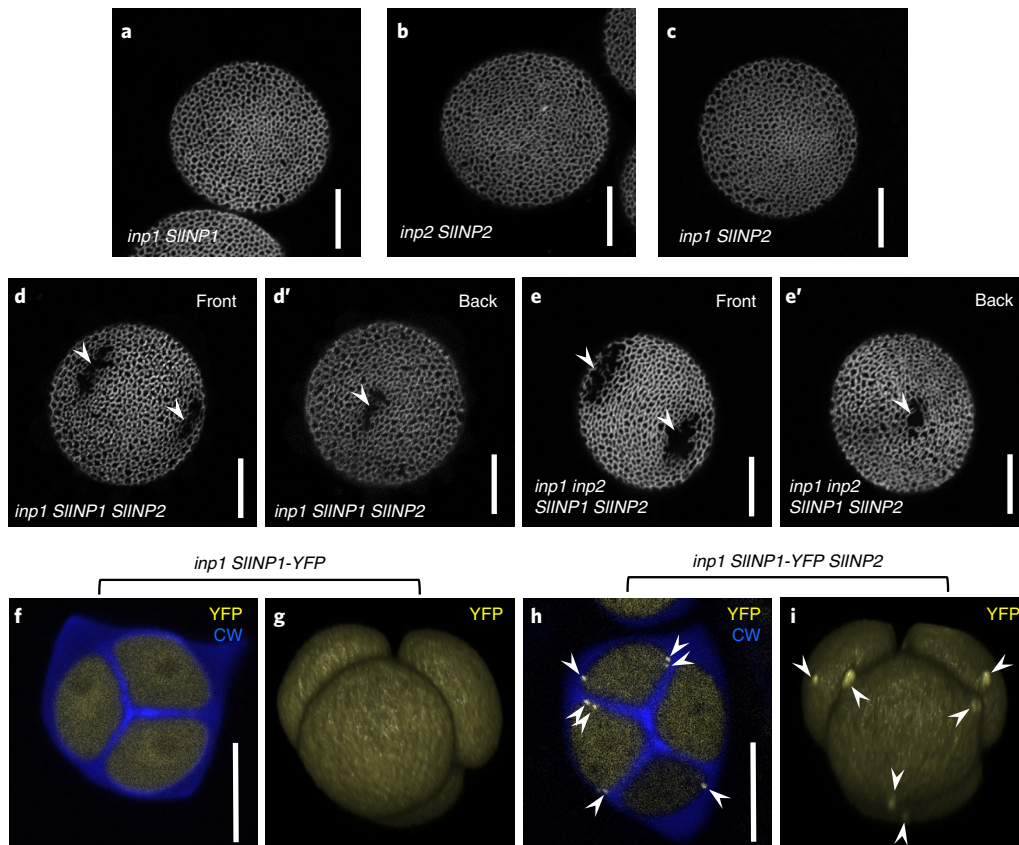


Fig. 7 | Tomato orthologues of INP1 and INP2 fail to function in *Arabidopsis* when expressed individually but gain this ability when co-expressed.

a–c, Neither SIINP1 (**a**) nor SIINP2 (**b,c**) are able to restore apertures in *Arabidopsis* pollen when expressed on their own. More than ten T_1 plants (≥ 50 pollen grains per plant) were analysed, with similar results. **d–e'**, When both SIINP1 and SIINP2 are expressed in *Arabidopsis*, they restore short to medium apertures (arrowheads) in the *inp1* (**d,d'**) and *inp1 inp2* (**e,e'**) *Arabidopsis* mutants. Confocal images of pollen grains stained with auramine O. Both front and back views are shown for the same pollen grains in (**d,d'**) and (**e,e'**) to demonstrate positions of apertures. Experiments were repeated twice, with similar results (~90% of plants had short- to medium-size apertures and the rest had no apertures). **f–i**, SIINP1-YFP localizes to the aperture domains in the presence of SIINP2 (**h,i**) but not when expressed on its own (**f,g**). Confocal optical sections (**f,h**) and three-dimensional reconstructions of tetrads of microspores (**g,i**). YFP signal is shown in yellow and callose wall (CW, stained by calcofluor white) is shown in blue. Arrowheads point to the YFP signal at the aperture domains. Experiments were repeated twice, with similar results. Scale bars, 10 μ m.

early as green algae, we found distinct, well-supported INP1 and INP2 protein lineages only in gymnosperms and angiosperms. In angiosperms, they have greatly diversified and display similar trends of evolutionary divergence, resulting in phylogenetic trees of similar topology (Fig. 5). Orthologues of both INP1 and INP2 exist in various families of rosids, asterids, basal eudicots, monocots and magnoliids. The INP1 sequence is also present among the transcripts from two ANA-grade basal angiosperms, *Austrobaileya* and *Nymphaea*. Failure to find an INP2 homologue in *Austrobaileya*, despite finding one in *Nymphaea*, could be due to the incompleteness of the database. Interestingly, both INP1 and INP2 are absent from the genome of *Amborella*, another basal angiosperm whose genome was published several years ago¹⁹.

Degrees of sequence divergence within the INP1 and INP2 angiosperm lineages are generally consistent with the phylogenetic relationships between species (Supplementary Table 1 and Fig. 5). Both *Arabidopsis* INP1 and INP2 (AtINP1 and AtINP2) share between ~95 and ~70% protein sequence identity with their respective orthologues from closely related species in the Brassicaceae and Cleomaceae families. Sequence identity with orthologues from more distantly related eudicots drops to ~40–50%. In monocots, the similarities to AtINP1 and AtINP2 are further reduced: proteins from Arecales and Bromeliaceae families (for example, palms

and pineapple) exhibit ~45 to ~30% sequence identity with AtINP1 and AtINP2. In both INP1 and INP2 lineages, particularly distinct clades are formed by proteins from grasses (Poaceae) (Fig. 5 and Supplementary Table 1): within each INP group, these proteins diverged greatly from the rest of their lineages (showing ~35 and ~20–25% identity, respectively, to AtINP1 and AtINP2) but retained >80% identity to their orthologues from other species of Poaceae despite the long evolutionary history of this monocot family²⁰.

INP1 and INP2 are functionally species-specific. The similar evolutionary trends displayed by INP1 and INP2, as well as the ability of these proteins to interact, led us to suggest that INP2 might serve as a species-specific partner for INP1. We tested this idea using the orthologues of INP1 and INP2 from tomato *Solanum lycopersicum* (SIINP1 and SIINP2) which both share ~45% amino acid identity with their *Arabidopsis* counterparts. Using the Y2H and split-luciferase assays, we confirmed the ability of SIINP1 and SIINP2 to interact (Fig. 6a,b). Furthermore, in both assays, the tomato INP proteins specifically interacted with each other and not with the *Arabidopsis* proteins (Fig. 6a,c).

We demonstrated previously that SIINP1 was unable to localize to aperture domains and restore apertures when expressed in the *Arabidopsis* *inp1* mutant¹² (Fig. 7a). Here, we placed SIINP2

under the control of the *AtINP2* promoter and transformed the *AtINP2pr:SlINP2* construct into the *Arabidopsis inp2-1* mutant. Similar to *SlINP1*, *SlINP2* failed to restore apertures in the *Arabidopsis inp2* mutant (12/12 T_1 plants; Fig. 7b), suggesting that *INP2* probably also exhibits functional species specificity.

Co-expression of *SlINP1* and *SlINP2* restores aperture formation in *Arabidopsis* mutants. Since individually expressed *SlINP1* and *SlINP2* did not restore apertures in *Arabidopsis* (Fig. 7a,b), we tested if together they could gain functionality in that species. To this end, we transformed *INP2pr:SlINP2* into *inp1* (as expected, apertures were not restored in all 14 T_1 plants; Fig. 7c) and crossed these plants with the previously characterized *DMC1pr:SlINP1-YFP* *inp1* transgenic lines¹². In the resulting *inp1* progeny that inherited just a single transgene, no apertures were restored. But, remarkably, among the *inp1* progeny that inherited both transgenes, 91% of plants (64/70) produced pollen with short- to medium-size apertures (Fig. 7d,d').

Because *SlINP1* was tagged with YFP, we assessed its localization in *Arabidopsis* tetrad-stage microspores. As shown previously¹², when expressed on its own, *SlINP1-YFP* fails to accumulate at the plasma-membrane aperture domains, instead localizing diffusely in the cytoplasm (Fig. 7f,g). However, in the presence of *SlINP2*, *SlINP1-YFP* gained the ability to successfully assemble into distinct puncta at the plasma-membrane aperture domains (Fig. 7h,i).

We further crossed the *SlINP1 SlINP2 inp1* plants with *inp2* plants to generate the double *inp1 inp2* mutants carrying both transgenes. In the F_2 and F_3 generations, those double mutants that inherited both tomato transgenes also produced pollen with apertures (Fig. 7e,e'). Taken together, these results demonstrate that *SlINP1* and *SlINP2* act as species-specific partners in pollen aperture formation.

Certain regions of *INP2* mediate its species specificity in *Arabidopsis*. To identify sequences in *INP2* responsible for the species specificity of this protein, we divided it into seven regions, on the basis of position relative to the transmembrane and DOG1 domains and evolutionary conservation (Fig. 8a and Extended Data Fig. 3). Regions were chosen as follows: (1) the N terminus (N), which in *AtINP2* encompasses the predicted TM region; (2) the acidic region, which, besides its enrichment in Asp and Glu residues, is fairly divergent across species; (3) the middle region; (4) the DOG1 domain; (5) the low-conservation region (LCR), which shows high sequence divergence across species; (6) the conserved C-terminal region (CTR); and (7) the highly divergent C-tail. To test which of the *AtINP2* regions were necessary for its function, we created seven chimaeric transgenic constructs in which most of the sequence came from *AtINP2*, while one region at a time was replaced with the corresponding sequence from *SlINP2* (Fig. 8a). In addition, to test if any of the *AtINP2* regions were sufficient for its function in *Arabidopsis*, we created a complementary set of seven constructs; in those, most of the protein was from *SlINP2* and a single region came from *AtINP2* (Fig. 8a). These 14 constructs were transformed into the *Arabidopsis inp2* mutant and their ability to complement the mutant phenotype was assessed.

The *AtINP2* constructs in which the N terminus, the middle region, the LCR or the CTR were replaced with the *SlINP2* sequences all restored apertures in *Arabidopsis* (19/19, 10/12, 10/12, 19/20 T_1 plants, respectively) (Fig. 8a,b,d,f,g), indicating that *AtINP2* can

tolerate the presence of tomato sequences in these four regions. The ability of the construct with the Sl N terminus to function in *Arabidopsis* was surprising, as this region in *SlINP2* is not predicted to contain a TM domain.

In contrast, the *AtINP2* constructs with the *SlINP2* acidic region or the DOG1 domain failed to rescue apertures in *Arabidopsis* (Fig. 8a,c,e) (11/11 T_1 plants for both constructs), demonstrating that these regions are critical for species-specific interactions. In addition, although five out of 12 T_1 plants expressing *AtINP2* with the tomato C-tail had some ability to produce short apertures (Fig. 8h), overall, the *AtINP2^{SlC-tail}* protein performed poorly: in ten out of 12 T_1 plants all or some pollen grains lacked apertures (Fig. 8h'). The divergent C-tail, therefore, probably also contributes to the protein's species specificity.

The complementary set of the *SlINP2* constructs with single *AtINP2* regions showed that none of the *AtINP2* regions was sufficient on its own to convert *SlINP2* into a protein able to function in *Arabidopsis* (Fig. 8a and Extended Data Fig. 5; ten or more T_1 plants were analysed for each construct). This suggests that sequences from more than one region contribute to the *INP2* species specificity.

We then explored the extent to which *AtINP2* can tolerate the simultaneous replacement of the regions which, individually, did not impact its functionality. The chimaeric *INP2s* in which either the N terminus and the middle region or the LCR and the CTR came from *SlINP2* were still functional in *Arabidopsis* (Fig. 8a,i,j); 16/18 T_1 plants for *AtINP2^{SlN+mid}* and 18/18 T_1 plants for *AtINP2^{SlLCR+CTR}*). In contrast, the simultaneous replacement of the middle region and the CTR (*AtINP2^{Slmid+CTR}*) resulted in a pronounced loss of protein activity: pollen developed either only very short apertures and 'shadows of apertures' (Fig. 8a,k; 10/17 T_1 plants) or no apertures at all (7/17 T_1 plants). The replacement of all four of these regions (*AtINP2^{SlN+mid+LCR+CTR}*) resulted in a completely non-functional protein (Fig. 8a,l; 13/13 T_1 plants), indicating that while each of these four regions plays a less prominent role in the species-specific functionality of *INP2* compared to the acidic region, the DOG1 domain and the C-tail region, together they still provide important contributions.

Discussion

The diversity of pollen aperture patterns in nature probably reflects the diversity of mechanisms controlling formation of these structures. In this study, we identified and characterized the new pollen aperture factor *INP2*, which is essential for this process and acts as a species-specific partner for the previously discovered aperture factor *INP1*. While not closely related, *INP1* and *INP2* share multiple similarities, including their matching patterns of expression, identical mutant phenotypes and the presence of the DOG1 domain (Figs. 1–3). Phylogenetic analysis suggests that the two *INP* genes are the result of an ancient gene duplication that occurred in the common ancestor of gymnosperms and angiosperms, with the diverging genes evolving non-redundant functions important for the formation of pollen apertures. *INP1* and *INP2* proteins interact with each other, display similar evolutionary trends and show functional species specificity (Figs. 4–7), indicating that they have co-evolved to form a species-specific functional module that promotes aperture formation. The notion of species specificity of the components of this module is strongly supported by the ability of the tomato *SlINP1* and *SlINP2* proteins to restore apertures in *Arabidopsis* only when

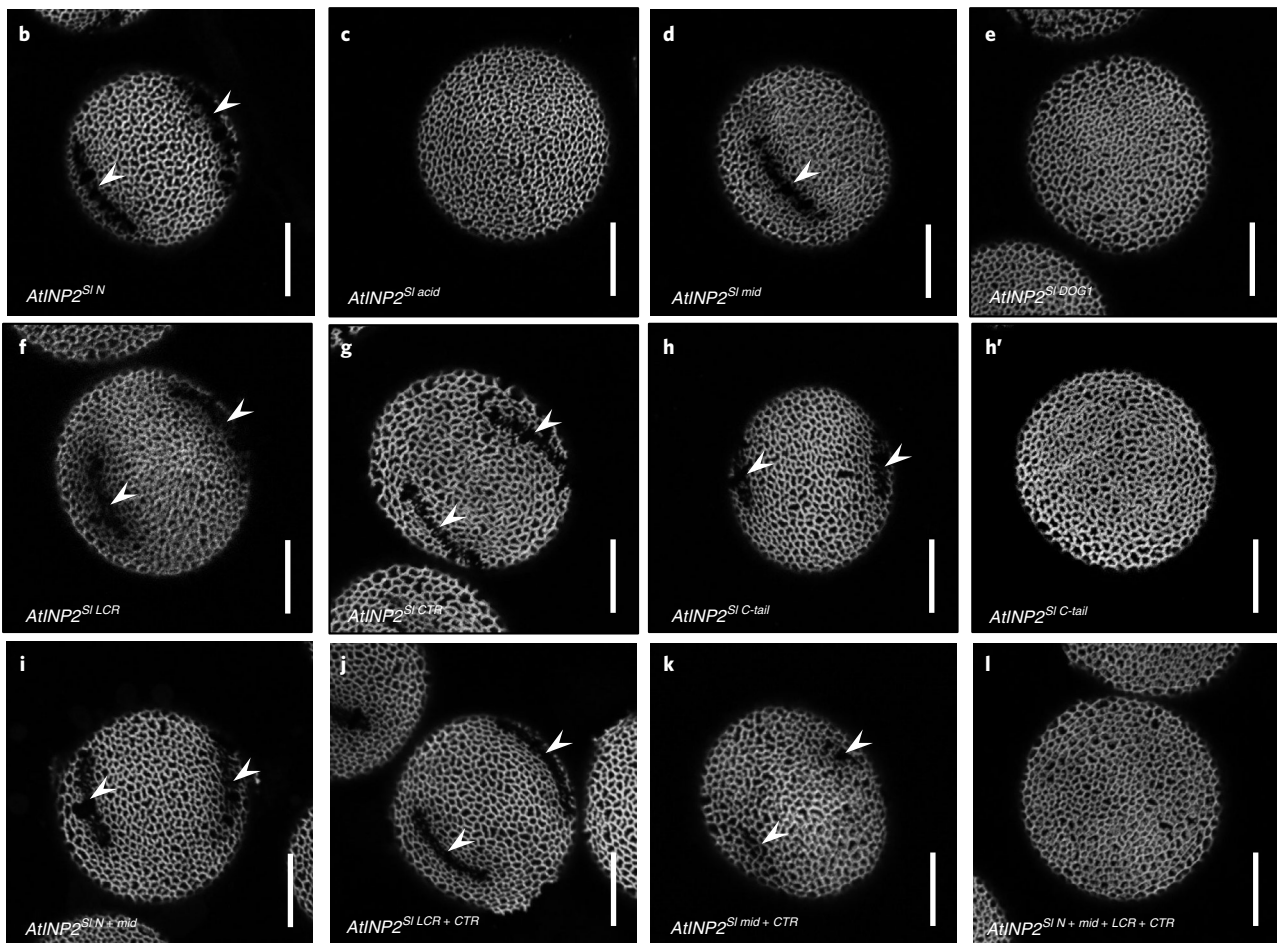
Fig. 8 | Certain regions of *INP2* mediate its species specificity. **a**, A diagram of 18 *INP2* chimaeric constructs containing regions from *Arabidopsis* (At, green) and tomato (Sl, magenta). The protein was divided into seven regions. The ability of a construct to restore apertures in the *Arabidopsis inp2* mutant is indicated by '+', the failure to restore apertures is indicated by '-' and the ability to restore apertures in some but not all transgenic lines is indicated by '+/-'. **b–m**, Representative images of pollen grains produced by transgenic *inp2* plants expressing different chimaeric *INP2* constructs. Over ten independent T_1 lines were tested for each construct (≥ 50 pollen grains per line), with all or nearly all lines producing similar results, except in **h**, **h'** and **k** where, as described, some plants produced short apertures and others no apertures. Apertures are indicated by arrowheads. Scale bars, 10 μm .

they are co-expressed but not when expressed individually (Fig. 7). We have also demonstrated that SIIINP1, which, on its own, does not assemble at the aperture domains in *Arabidopsis* tetrads¹², gains

the ability to do this in the presence of SIIINP2 (Fig. 7h,i). Our data show that several regions of INP2, including the DOG1 domain, contribute to its species specificity (Fig. 8).

a

Construct	N	Acidic	Middle	DOG1	LCR	CTR	C-tail	Rescue
<i>AtINP2^{SI} N</i>	SI				At			+
<i>AtINP2^{SI acid}</i>	At	SI			At			-
<i>AtINP2^{SI mid}</i>		At	SI			At		+
<i>AtINP2^{SI DOG1}</i>			At	SI		At		-
<i>AtINP2^{SI LCR}</i>			At		SI		At	+
<i>AtINP2^{SI CTR}</i>			At			SI	At	+
<i>AtINP2^{SI C-tail}</i>			At				SI	+-
<i>SIIINP2^{At} N</i>	At			SI				-
<i>SIIINP2^{At acid}</i>	SI	At			SI			-
<i>SIIINP2^{At mid}</i>		SI	At		SI			-
<i>SIIINP2^{At DOG1}</i>			SI	At		SI		-
<i>SIIINP2^{At LCR}</i>			SI		At		SI	-
<i>SIIINP2^{At CTR}</i>			SI			At	SI	-
<i>SIIINP2^{At C-tail}</i>			SI				At	-
<i>AtINP2^{SI N + mid}</i>	SI	At	SI		At			+
<i>AtINP2^{SI LCR + CTR}</i>			At		SI		At	+
<i>AtINP2^{SI mid + CTR}</i>		At	SI	At		SI	At	+-
<i>AtINP2^{SI N + mid + LCR + CTR}</i>	SI	At	SI	At	SI		At	-



INP1 and INP2 are both proteins of unknown function. Accumulating evidence for the *Arabidopsis* INP1 indicates that it is a late-acting aperture factor that becomes attracted to the plasma-membrane domains in microspores that are already pre-specified as aperture sites^{12,14,21}. As such, even though INP1 and its partner INP2 are critical for aperture formation, they are unlikely to directly define positions and morphology of apertures and to contribute to the aperture diversity in that way. This idea is supported by the phenotype of the apertures restored in the *Arabidopsis* *inp1* mutant by the expression of the tomato SLINP1/SLINP2 complex: the restored apertures did not resemble the corporate tomato apertures (PalDat: www.palddat.org)^{8,12} but were more like the colpate apertures of *Arabidopsis* (Fig. 7d,e'). It is, therefore, intriguing why these essential, yet late-acting aperture factors have diversified so much among angiosperms. This could indicate that different species have variations in the upstream mechanisms or differences in other interactors of the INP proteins. Relevant to this point, the apertures restored by the tomato complex in *Arabidopsis* were shorter than the normal *Arabidopsis* (or tomato) apertures (Fig. 7d,e'). While this could be due to reduced expression of the transgenes compared to that of endogenous genes, an alternative possibility is that, for optimal function, the tomato complex requires some additional species-specific component(s).

After aggregating at the aperture domains, the INP1 proteins in both *Arabidopsis* and rice participate in keeping the plasma membrane at these domains near the overlying callose wall and preventing the deposition of the exine precursor, primexine, at these sites^{6,10}. How can the newly discovered INP2 protein contribute to aperture formation? There are several possibilities and further investigations will be needed to distinguish between them. For instance, along with INP1, INP2 might directly be a part of the protein complex that assembles at the aperture domains and mediates their interaction with the callose wall. Alternatively, INP2 might be involved in the delivery of INP1 to its positions at the aperture domains. AtINP1 appears to aggregate on the extracellular side of the aperture domains, yet lacks a clear signal peptide or any obvious means to become anchored at the plasma membrane¹⁰. The proposed topology of AtINP2, with the TM domain at its N terminus and most of the protein outside the cell, is consistent with the idea that the INP1/INP2 complex in *Arabidopsis* is extracellular. It is tempting to speculate that the interaction of AtINP1 with the extracellular portion of AtINP2 could provide a way to anchor AtINP1 at the aperture domains. Yet, further experimentation will be needed to validate the topologies of AtINP2 and its orthologues from other species, many of which, like SLINP2, lack predicted TM domains; to find a way to visualize the INP2 subcellular localization; and to establish whether the INP proteins from species with very different aperture patterns function in the same way. Intriguingly, OsINP1 from rice was recently shown to interact with the cytoplasmic portion of a lectin receptor-like kinase⁶, suggesting a possibility that in grasses, whose single pore-like apertures differ greatly from the three furrows in *Arabidopsis* and tomato, INP1 might have a role at the cytoplasmic side of the aperture domains. Since both INP lineages in grasses have greatly diverged from their counterparts in eudicots and some other monocots, it would be very interesting to determine whether in that plant family INP2 proteins are also involved in aperture formation and function in a complex with INP1 or whether they have evolved other functions.

In conclusion, our study uncovered a new essential player in the poorly understood mechanism underlying the formation of important patterning elements on the pollen surface and demonstrated that the two DOG1 domain-containing aperture factors form a protein complex whose components contribute to the species specificity of this molecular mechanism.

Methods

Plant materials and growth conditions. Plants were grown at 20–22 °C with the 16-h light/8-h dark cycle in growth chambers or in a greenhouse at the Biotechnology facility at Ohio State University (OSU). Besides the genotypes generated in this study, the following genotypes were used: Columbia (Col-0), Landsberg *erecta* (Ler), *inp1-1* (ref. ⁹), *DMC1pr:INP1-YFP inp1-1* (refs. ^{10,14}); *DMC1pr:SLINP1-YFP inp1-1* (ref. ¹²); and *D6PKL3pr:D6PKL3-YFP d6pkl3-2* (ref. ¹¹).

Forward genetic screen. The genetic screen which led to the isolation of the *inp2-1* mutant was recently described²¹. In brief, M₂ plants (~10,000) from eight pools of ethylmethane sulfonate-treated lines of Ler background were screened for the presence of morphological abnormalities in their pollen (for example, in size, shape, light reflection, ease of pollen release from anthers) identifiable with standard dissecting stereomicroscopes (Zeiss Stemi-2000C and Nikon SMZ745) at 75–80× magnification. Particular attention was paid to changes in pollen shape, known to be associated with aperture defects^{9,13}. For primary screening, dry pollen did not undergo any treatment. At this level of magnification, pollen of the *inp2-1* mutant looked rounder than the wild-type pollen. Pollen was then stained with auramine O as described in ref. ¹⁴ and aperture defects were observed with confocal microscopy. Mutant *inp2-1* was then backcrossed with Ler once. To test for complementation, *inp2-1* mutant was crossed with *inp1-1* and the pollen of their F₁ progeny was observed with dissecting and confocal microscopes.

Mapping the *inp2* defect. Mutant *inp2-1* with Ler background was crossed with Col-0 and the resulting F₂ population was screened under a dissecting microscope for the presence of the round-pollen mutant phenotype. DNA was isolated from 189 mutants. First, the bulked segregant analysis²² placed the mutation to the top of chromosome 1. This was followed by map-based positional cloning using individual F₂ mutants²². The INDEL-based polymerase chain reaction (PCR) markers for this analysis were generated as previously described¹¹, using combined information from the 1,001 Genomes Project database (<http://signal.salk.edu/atg1001/index.php>)²³ and the *Arabidopsis* Mapping Platform²⁴. The mutation was mapped to a 146-kb region between 5,151,424 and 5,297,411 base pairs (bp).

To determine which of the 51 genes in this interval was responsible for the aperture defect in *inp2*, we used information from the *Arabidopsis* RNA-seq database TRAVA (travadb.org)¹⁵ to identify genes expressed in the young buds at or near the tetrad stage of pollen development (flowers 12–18). Although 37 of the 51 genes are expressed in buds at these stages, expression of only one of them, At1g15320, is specifically restricted to these tissues and stages. The finding that this gene, like *INP1*, encodes a protein with the DOG1 domain led us to further prioritize it as a strong candidate for *INP2*. Sequencing of this gene from the *inp2-1* mutant revealed the presence of the point mutation which leads to a premature stop codon (Trp84Stop).

Sequence retrieval and phylogenetic analysis. Sequences of INP1 and INP2 homologues were retrieved by TBLASTN from NCBI (<https://blast.ncbi.nlm.nih.gov/Blast.cgi>), Phytozome v.12.1 (<https://phytozome.jgi.doe.gov/pz/portal.html>), the 1000 Plants project (OneKP-China National Gene Bank (<https://db.cngb.org/onekp/>)²⁵ and PLAZA (<https://bioinformatics.psug.ugent.be/plaza/>)²⁶. Accession numbers are provided in Supplementary Table 1. For multiple sequences alignment, MAFFT v.7.017 (L-INS-i algorithm) was used. The alignment positions with more than 20% gaps were removed with trimAl²⁷. ModelFinder²⁸ accessed through IQ-TREE²⁹ tested 546 protein models to find the best-fit model of evolution (INP1 + INP2; JTT + R5). Lastly, the IQ-TREE program³⁰ was used to construct phylogenetic trees, with the maximum likelihood method and 1,000 bootstrap replicates. The trees were visualized in iTOL v.5 (ref. ³¹).

Confocal microscopy. Preparation and imaging of mature pollen and tetrads were performed as previously described¹⁴. Three-dimensional reconstruction of tetrads was done using NIS Elements v.4.20 software (Nikon).

Generation of the CRISPR allele of INP2. The guide RNA for the At1g15320 gene was selected with the help of the CRISPR-PLANT platform (<https://www.genome.arizona.edu/crispr/>)³² and its sequence was cloned into the pHE401E vector³³ as described³⁴ using primers Oligo-01-F-INP2-T1/Oligo-R-INP2-T1 (Supplementary Table 2). The resulting construct was transformed into the *Agrobacterium tumefaciens* strain GV3101 and *Arabidopsis* Col-0 plants were transformed using the floral dip method³⁵. The T₁ transformants were selected on hygromycin plates and 18 seedlings were transferred to soil. Two of the 18 T₁ plants displayed the inaperturate phenotype. Sequencing of the At1g15320 gene from these two plants revealed that both had the same homozygous single-nucleotide deletion which occurred one nucleotide before the PAM sequence and caused a shift in the ORF after the codon 83.

INP2 complementation and expression constructs. Primers used in this study are listed in Supplementary Table 2. All fragments for cloning were amplified with high-fidelity Phusion DNA polymerase (NEB, M0530 or Thermo Fisher, F530). To create the *INP2pr:INP2* ORF construct, the *INP2* promoter (a fragment of 701 bp

from the end of the preceding gene to the start codon of *INP2*) and ORF were amplified, respectively, with primer pairs *INP2pr*-IF-F/ *INP2pr*-IF-R and *INP2* ORF-IF-F/ *INP2*-Stop-IF-R from the Col-0 genomic DNA and from the *INP2* complementary DNA clone **DQ446252** obtained from the Arabidopsis Biological Resource Center. The resulting two fragments were cloned into the *SacI* and *SpeI* sites in the pGR111 binary vector³⁶ using in-fusion cloning (Takara, no. 638950). A *BamHI* site was introduced in front of the *INP2* start codon to facilitate subsequent cloning. To create the *INP2prgINP2* construct, the genomic fragment, which included the coding sequence, introns and the 567-bp region downstream of the *INP2* stop codon, was amplified from Col-0 genomic DNA with primers *INP2* ORF-IF-F/g*INP2*-*SpeI*-R and cloned downstream of the *INP2* promoter between the *BamHI* and *SpeI* sites. Constructs were verified by sequencing and transformed into the *A. tumefaciens* strain GV3101. The *inp2* plants were then transformed by floral dip. Transgenic *T₁* plants were selected with BASTA and the presence of transgenes was confirmed with specific primers.

To generate the reporter construct *INP2pr:H2B-RFP*, the *H2B-RFP* fusion gene was cloned into the *BamHI*/*SpeI* sites downstream of the *INP2* promoter in pGR111 and the construct was transformed into the Col-0 plants.

To express *INP2* tagged at the C terminus with one or three copies of HA (HA₁ and HA₃), we first used the *in vivo* assembly (IVA) cloning in *E. coli*³⁷ to combine the *INP2* ORF (without the stop codon), an eight-amino acid Gly-rich linker (G₈PGS) and the corresponding HA tag in the pGEM-T-Easy vector (Promega). After confirming the sequences, the tagged *INP2* sequences were placed into pGR111 under the control of the *INP2* promoter and the resulting constructs were transformed into *inp2*. The shorter construct with HA₁ produced better rescue. Out of the 14 *INP2-HA₁* *T₁* lines, 11 had long to medium-size apertures, two had short apertures and one had no apertures. Out of the 13 *INP2-HA₃* *T₁* lines, four had medium-size apertures, four had short apertures and five had no apertures. Note that the construct with HA₃ inserted directly after *INP2* (without the Gly-rich linker) essentially failed to rescue the mutant phenotype (2/9 *T₁* lines produced short apertures and the rest produced no apertures).

SIINP1, SIINP2 and Arabidopsis/tomato chimaeric INP2 constructs. The tomato paralogues of *INP1* and *INP2* were identified in the tomato genome, respectively, as Solyc08g079050 and Solyc03g116770. The *DMC1pr:SIINP1-YFP* construct was previously described¹² and the previously characterized transgenic lines of *DMC1pr:SIINP1-YFP* *inp1* were used in this study. To create the *INP2pr:SIINP2* construct, the tomato genomic DNA was amplified with primers *BamHI-SIINP2-BF*/*SpeI-SIINP2-AR*. The resulting fragment was digested with *BamHI*/*SpeI* and used to replace the *AtINP2* gene in the *INP2prgINP2* construct. The construct was transformed into the *inp2* and *inp1* mutants, transgenic *T₁* plants were selected with BASTA and the presence of the transgene was confirmed with specific primers.

To genotype the *F₂* populations which segregated both the *DMC1pr:SIINP1-YFP* and *INP2pr:SIINP2* transgenes in combination with the *inp1* or *inp1* *inp2* mutations, the following primers and conditions were used: for *SIINP1*, 2-SI-F/ Sly *INP1*-R-NcoI primers; for *SIINP2*, AD23/AD8 primers; for *inp1*, a cleaved amplified polymorphism (CAPS) marker was used (22600-DF/22600-DR primers, SacI cuts the mutant allele, digestion products resolved on 1.5% agarose gels); for *inp2*, a derived CAPS (dCAPS) marker was used (At1g15320-BF/AD402 primers, AccI cuts the mutant band, digestion products resolved on 4% agarose gels). PCR with 40 cycles (98 °C for 15 s, 55 °C for 15 s, 72 °C for 30 s) were performed in all cases.

To quickly generate multiple chimaeras of *AtINP2* and *SIINP2*, we used the ORF versions of the constructs and the IVA cloning method³⁷. The *AtINP2* ORF was first amplified using primers pGEM-*INP2*-IF-F/*INP2*-pGEM-IF-R and cloned with the help of the in-fusion procedure into the pGEM-T-Easy vector digested with *SacI* and *NcoI*. A *BamHI* site and a *SpeI* site were introduced, respectively, at the beginning and at the end of the *AtINP2* sequence for ease of subsequent subcloning into the binary pGR111 vector. To obtain the *SIINP2* cDNA, total mRNA was isolated from young tomato buds with Trizol (Thermo Fisher, no. 15596026) and converted into cDNA as previously described¹⁴. The *SIINP2* ORF was then amplified with primers AD19/AD16 and used to replace *AtINP2* between the *BamHI* and *SpeI* sites with the help of the IVA method in pGEM-T-Easy vector, which itself was amplified with AD17/AD14 primers. The resulting pGEM-T-Easy-based *AtINP2* and *SIINP2* constructs were used as backbone templates for chimaeras, while pGR111-based constructs were used as the *INP2*-region templates in IVA cloning. Thirty primers (AD1 through AD28, pGEM-*INP2*-IF-F and *INP2*-pGEM-IF-R) were used in different combinations (as indicated in Supplementary Table 2) to obtain all 18 chimaeras. IVA reactions were performed as described in ref. ³⁷: products of single PCR amplified with sets of four primers were treated with *DpnI* (15 min at 37 °C) to degrade templates and directly transformed into *E. coli*. Single-region chimaeras were used as backbone templates to generate multiregion chimaeras. All chimaeric sequences were verified by sequencing, cut out with the *BamHI*/*SpeI* digestion, subcloned downstream of the *INP2* promoter in the pGR111 vector and resequenced again. The final constructs were transformed into the *inp2* mutant; transgenic *T₁* plants were selected with BASTA and the presence of the transgenes was confirmed with specific primers.

Yeast two-hybrid assay. The Y2H assays were performed as previously described¹⁴. The DNA-binding domain construct *pB29-INP1* and the activation-domain construct *pP6-INP1* were described previously¹¹. Other constructs were created by cloning the coding sequences of *AtINP2*, *SIINP1*, *SIINP2* or their truncated forms into the same vectors. Constructs were cotransformed in indicated combinations into the NMY51 yeast strain. Positive bait-prey cotransformants were selected on the synthetic dropout medium lacking Leu and Trp (-LW). To test for interaction, cotransformed yeast cells were grown on the medium that lacked Leu, Trp and His and contained either 3 mM or 20 mM 3-amino-1,2,4-triazole (-LWH + 3-AT).

Bifluorescent molecular complementation assay. To create the *35SPr:nYFP-AtINP1* and *35SPr:cYFP-AtINP2* constructs, *AtINP1* and *AtINP2* ORFs were, respectively, amplified with AD17-BHL/AD18-BHL and AD469/470 primers, digested with *PacI*/*XbaI* and inserted into *35SPr:nYFP*₍₁₋₁₅₈₎ and *35SPr:cYFP*₍₁₅₉₋₂₃₈₎ binary vectors³⁸. Constructs were transformed into *A. tumefaciens* strain GV3101. Bacterial cultures containing these constructs were grown to optical density $OD_{600}=0.4$ and co-infiltrated into tobacco (*Nicotiana benthamiana*) leaves along with agrobacteria expressing the RNA silencing suppressor P19 (ref. ³⁹) at the 1:1:2 ratio. Infiltrated leaves were grown for 5 d, after which samples were collected and imaged on a Nikon A1 + confocal microscope using identical settings for all imaging. Samples were excited with a 514-nm laser and YFP emission was collected at 522–555 nm.

Co-immunoprecipitation. *AtINP1* and *AtINP2* ORFs were amplified through two sequential PCR reactions (first PCR with primers AD326/488 for *AtINP1* and primers AD370/489 for *AtINP2*; second PCR with primers AD122/123 for both genes) and inserted into the pDONR207 vector through the Gateway BP recombination reaction (Invitrogen, no. 11789020). Gateway LR recombination reaction was then used to transfer these sequences into the pCsVMV:GFP-C-999 or pCsVMV:HA-C-1300 vectors⁴⁰ (a gift from D. Somers, OSU), producing four constructs: CsVMV-*AtINP1*-GFP, CsVMV-*AtINP1*-HA3, CsVMV-*AtINP2*-GFP and CsVMV-*AtINP2*-HA3. All constructs were verified by sequencing and transformed into the *A. tumefaciens* strain GV3101 by electroporation. Tobacco leaves were infiltrated with different combinations of constructs (as described for BiFC above), harvested 5 d after infiltration and stored at -80 °C for at least 1 d. A total 1 μ l of monoclonal anti-GFP (A-11120, Molecular Probes by Thermo Fisher Scientific) was incubated with 1 μ l of protein A/G PLUS-agarose (SC-2003, Santa Cruz Biotechnology) and 8 μ l of IP buffer (50 mM Tris-HCl pH 7.5, 150 mM NaCl, 0.5% NP-40, 1 mM EDTA, 3 mM DTT, 1 mM PMSF, 5 μ g ml^{-1} leupeptin, 1 μ g ml^{-1} aprotinin, 1 μ g ml^{-1} pepstatin) overnight at 4 °C with gentle agitation. Frozen tobacco leaves were ground in liquid nitrogen and 500 μ l of ground tissue were mixed with 500 μ l of IP buffer. These samples were vortexed for 2–3 min and centrifuged at 18,400g for 10 min at 4 °C. Supernatant was added to the anti-GFP matrix agarose and the mixture was incubated for 2 h at 4 °C with gentle agitation, followed by centrifugation at 845g at 4 °C for 3 min to collect immune complexes. Pellets were washed three times with ice-cold IP buffer, once with 1× PBS buffer and eluted with 50 μ l of 2× SDS-PAGE sample buffer. Immunoblotting was performed as previously described¹¹. The following antibodies were used to detect the fusion proteins: rabbit anti-GFP (Abcam; ab6556), rat anti-HA (Sigma, 11867423001), anti-rabbit IgG peroxidase-conjugated (SeraCare/KPL; 5220-0283/04-15-06) and anti-rat IgG peroxidase-conjugated (SeraCare/KPL; 5220-0364/04-16-06). All antibodies were diluted 1:2,000 in TBS-T buffer (1× TBS, 0.1% Tween 20) supplemented with 4% non-fat milk. After the final washes, the membranes were processed with SuperSignal West Pico PLUS Chemiluminescent Substrate (Thermo Scientific, 34577) and imaged with a MyECL imager (Thermo Scientific).

Split-luciferase assay. The coding sequences of *INP1* and *INP2* from *Arabidopsis* and tomato were PCR amplified, cloned into pDONR207 (Invitrogen) and transferred into pCAMBIA1300-^{GW}Luc and pCAMBIA1300-^{GW}CLuc (ref. ⁴¹) (a gift from D. Mackey, OSU) using the Gateway technology. Resulting constructs were transformed individually into *A. tumefaciens* strain GV3101. Agrobacteria were collected and resuspended in infiltration buffer (10 mM MgCl₂, 10 mM MES, 150 μ M acetosyringone) at a final concentration of $OD_{600}=0.8$. Pairwise combinations of suspensions were infiltrated into young tobacco leaves, which were then allowed to grow for 3 d in light. A total 12–16 leaves were collected from five to ten plants, the abaxial side of leaves was sprayed with 1 mM luciferin (Biosynth, L-8220) and kept in the dark at 4 °C for 30 min. The bioluminescence images were captured using Azure Sapphire Biomolecule Imager (Azure Biosystems) and converted to heatmaps using the 16-colour look-up table from ImageJ v.1.53a.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

All data supporting the findings of this study are available within the article, Supplementary Information files or from the corresponding author on reasonable request. Source data are provided with this paper.

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Author contributions

B.H.L., R.W. and A.A.D. conceived and designed the experiments. B.H.L., R.W., I.M.M., S.H.R., P.A., M.H.T., K.A., P.C., A.H. and A.A.D. performed the experiments. E.P.A., A.A.D. and I.B.Z. performed phylogenetic analysis. B.H.L., R.W., I.M.M., K.A., P.C., E.P.A. and A.A.D. analysed the data. A.A.D. wrote the article and all authors revised and approved the final manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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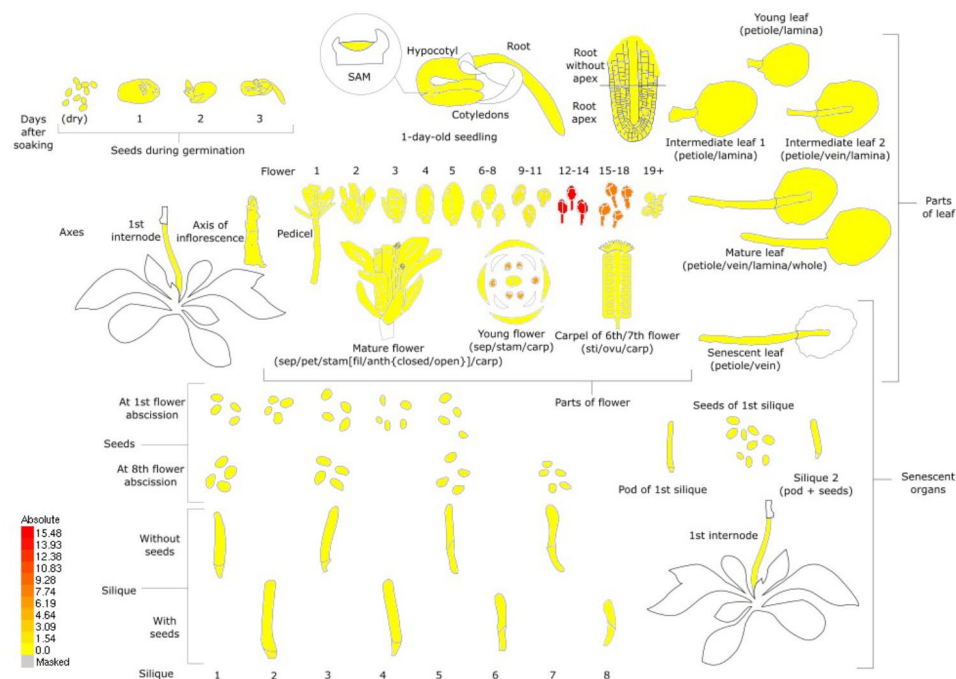
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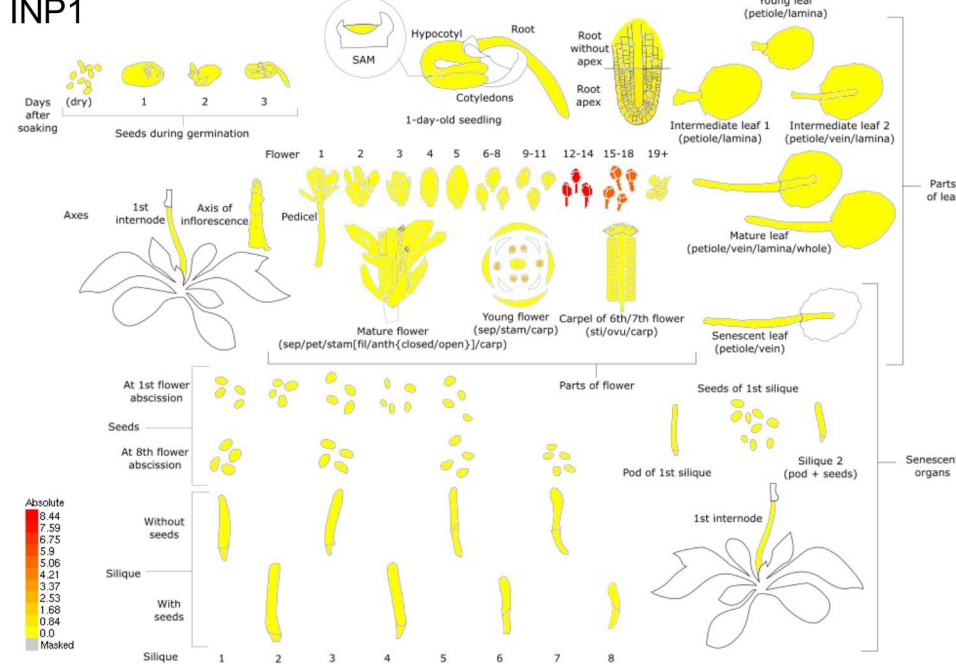
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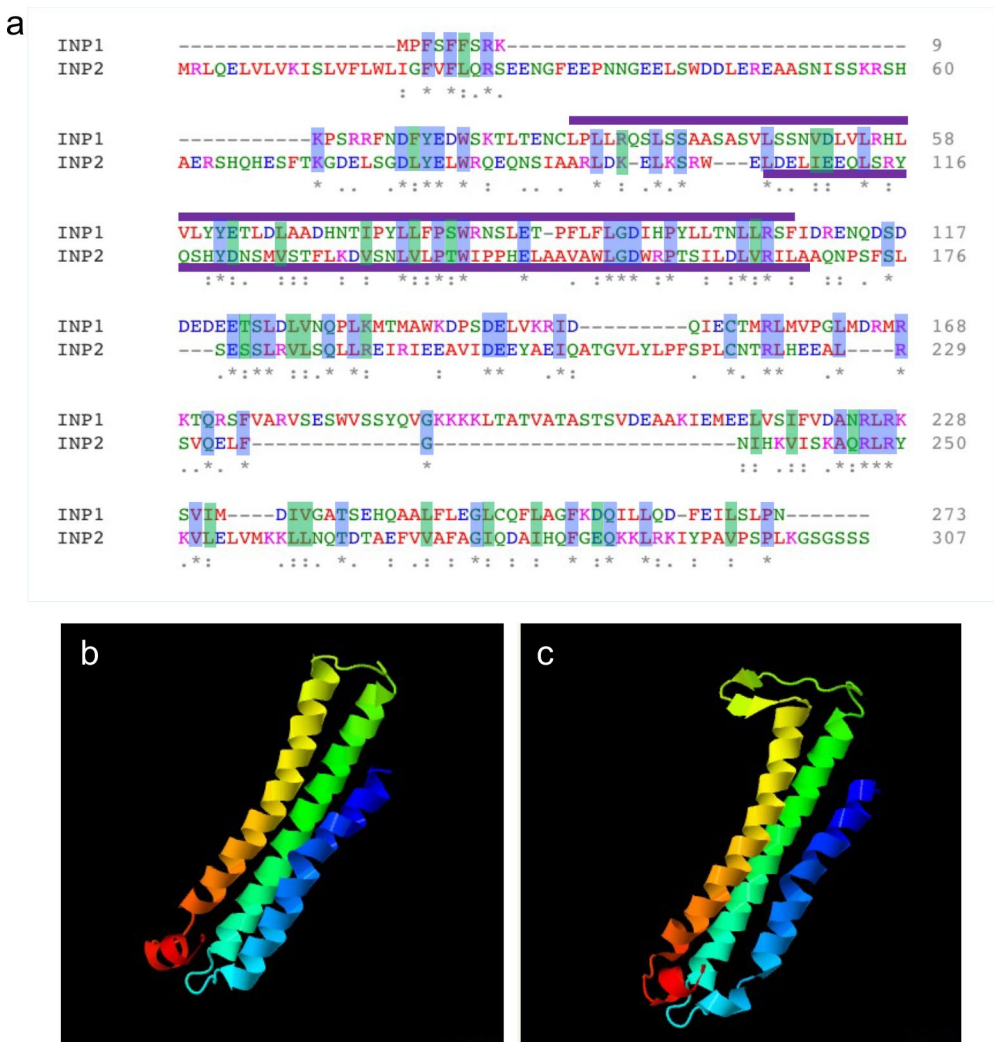
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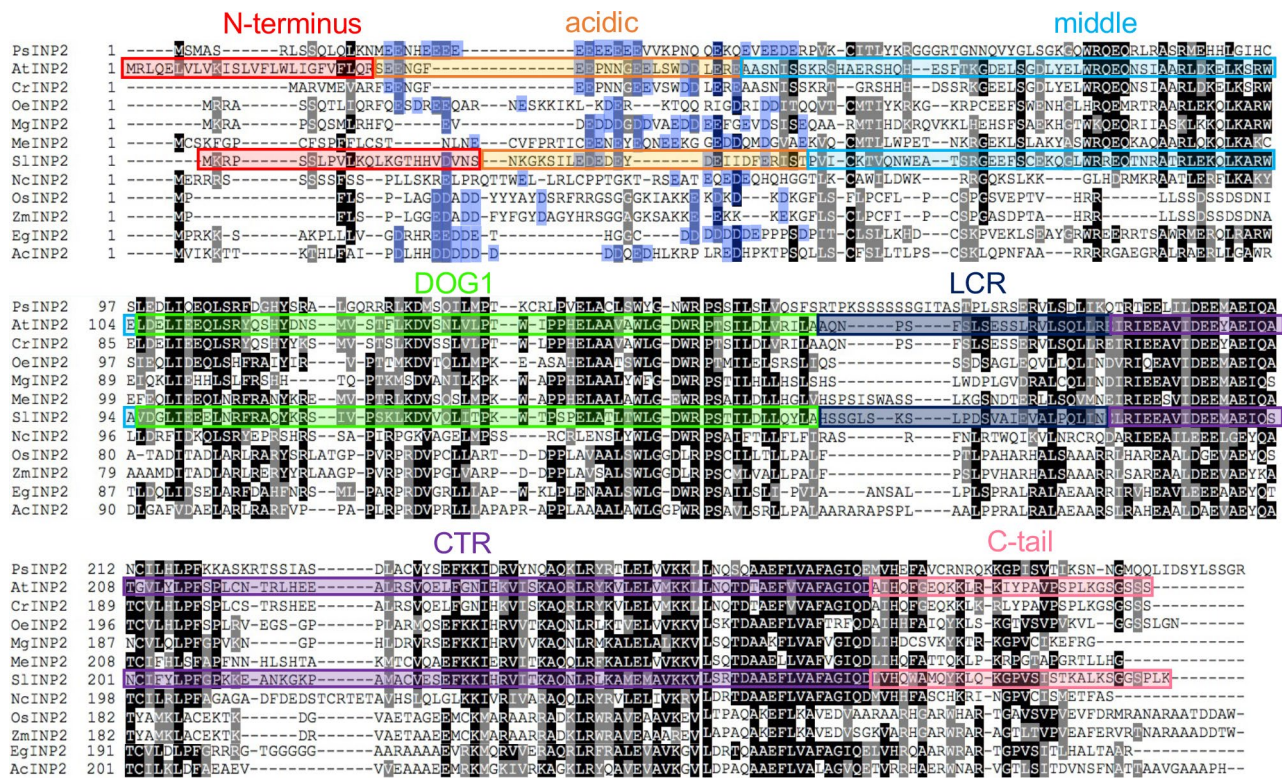
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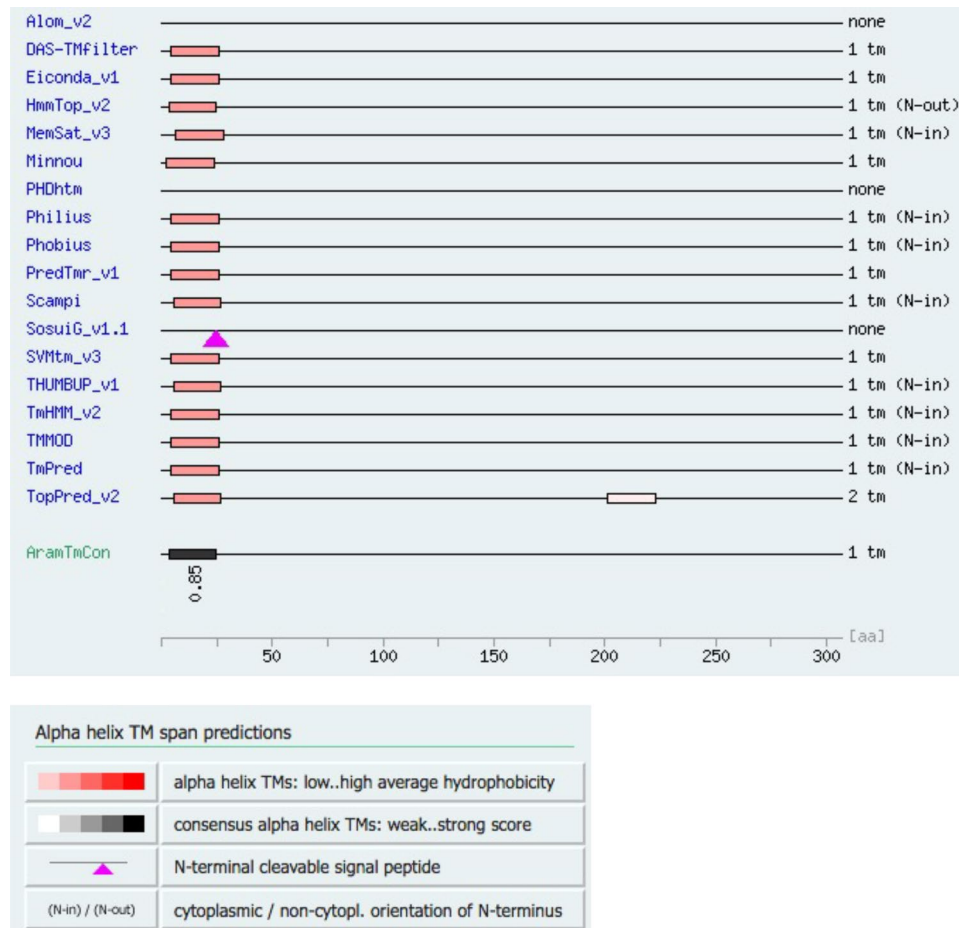
Extended Data Fig. 1 | INP2 and INP1 display similar expression patterns, with both genes showing highest expression in young developing buds. The RNA-seq data for INP2 (a) and INP1 (b) are from the dataset of Klepikova et al.¹⁵ and visualized with the BAR eFP Browser.



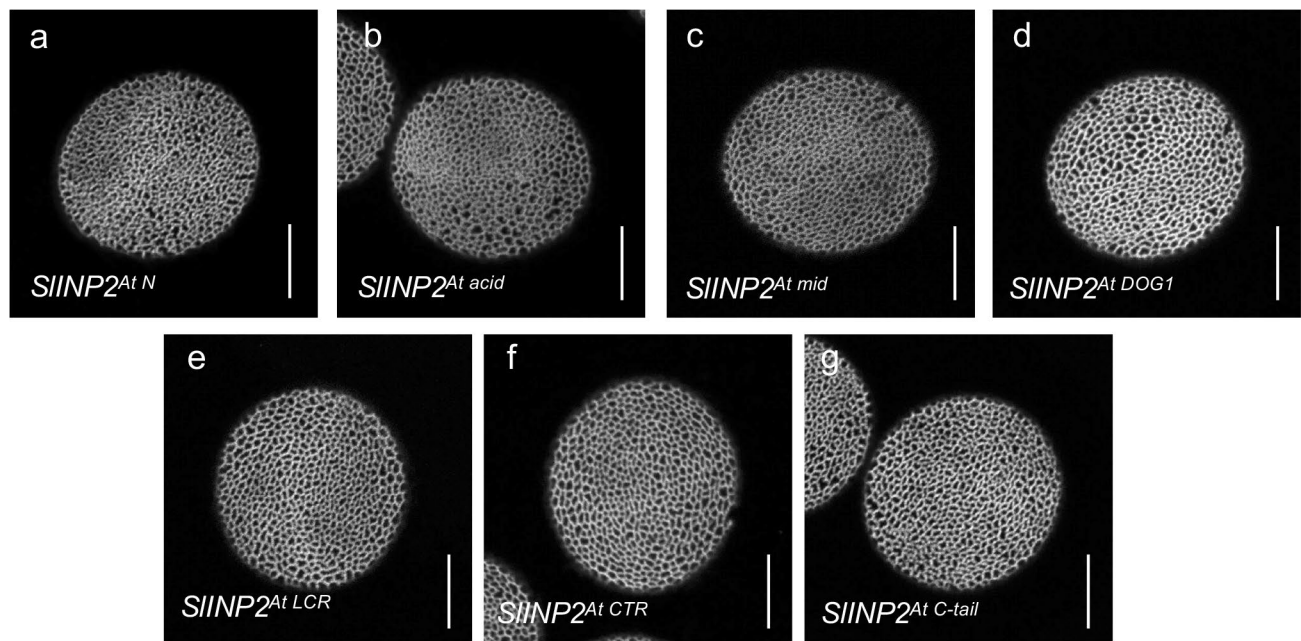
Extended Data Fig. 2 | INP1 and INP2 proteins both contain the DOG1 domain and have similar structural organization predicted for their C-terminal parts. **a**, Protein alignment between INP1 and INP2 proteins. Identical and similar (V/I/L, D/E, K/R, N/Q and S/T) residues are shaded, respectively, in blue and green. The positions of the DOG1 domains predicted by Pfam are indicated by purple lines. **b–c**, Protein structures predicted by Phyre2 for C-terminal parts of INP1 (**b**) and INP2 (**c**) (confidence: >97% for both proteins). In both cases, the modelled regions cover 114 amino acids, which constitute, respectively, 42% of INP1 and 37% of INP2. The same template (c4clvB, nickel-cobalt-cadmium resistance protein NccX from *Cupriavidus metallidurans* 31a) was selected by the program in both cases.



Extended Data Fig. 3 | Alignment of INP2 proteins from representatives of different angiosperm taxa. The following species were used (from top to bottom): *Papaver somniferum* (basal eudicots, Papaveraceae), *Arabidopsis thaliana* (rosids, Brassicaceae), *Capsella rubella* (rosids, Brassicaceae), *Olea europaea* (asterids, Oleaceae), *Mimulus guttatus* (asterids, Phrymaceae), *Manihot esculenta* (rosids, Euphorbiaceae), *Solanum lycopersicum* (asterids, Solanaceae), *Nymphaea colorata* (basal angiosperms, ANA, Nymphaeaceae), *Oryza sativa* (monocots, Poaceae), *Zea mays* (monocots, Poaceae), *Elaeis guineensis* (monocots, Arecaceae), *Ananas comosus* (monocots, Bromeliaceae). The seven regions selected for creating AtINP2/SlINP2 chimeras are indicated by differently coloured rectangles. Aspartate (D) and glutamate (E) residues in the acidic region are shaded in blue. Black shading indicates identical amino acids and grey shading indicates similar amino acids present at the same position in at least half of the aligned proteins.



Extended Data Fig. 4 | *Arabidopsis* INP2 likely contains a transmembrane domain at its N terminus. Multiple TM discovery algorithms predict existence of the transmembrane domain at the N terminus of INP2 from *Arabidopsis thaliana* (AtINP2), with the consensus score of 0.85 generated by the plant membrane protein database Aramemnon (AramTMCon).

**Extended Data Fig. 5 | None of the seven AtINP2 regions is sufficient on its own to convert SII NP2 into a protein able to function in Arabidopsis.**

Confocal images of pollen grains produced by the transgenic *inp2* plants expressing seven versions of chimeric *SII NP2* constructs in which one region at a time was replaced with the corresponding regions from *AtINP2*. At least 10 independent T_1 lines were tested for each construct (≥ 50 pollen grains per line), with similar results. Scale bars = 10 μm .

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Software and code

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Data collection For bioinformatics analysis, BLAST search was conducted using NCBI website (<http://www.ncbi.nlm.nih.gov/>), Phytozome v. 12.1 (<https://phytozome.jgi.doe.gov/pz/portal.html>), the 1,000 Plants project (OneKP-China National Gene Bank (<https://db.cngb.org/onekp/>)) and PLAZA (<https://bioinformatics.psb.ugent.be/plaza/>). For confocal imaging, NIS Elements v.4.20 software (Nikon) was used.

Data analysis 3D reconstruction of tetrads was done using NIS Elements v.4.20 software (Nikon). For multiple sequences alignment, MAFFT v7.017 (L-INS-i algorithm) was used (<https://www.ebi.ac.uk/Tools/msa/mafft/>). The IQ-TREE program v. 1.6.12, (<http://www.iqtree.org/>) was used to construct phylogenetic trees. The phylogenetic trees were visualized in iTOL v. 5 (<https://itol.embl.de/>). Split-luciferase assay analysis was done with ImageJ v. 1.53a.

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Public RNA-seq data for gene expression analysis were obtained from the TRAVA database (travadb.org).

Protein sequences were obtained from TAIR (<https://www.arabidopsis.org/>), NCBI (<http://www.ncbi.nlm.nih.gov/>), Phytozome v. 12.1 (<https://phytozome.jgi.doe.gov/pz/portal.html>), the 1,000 Plants project (OneKP-China National Gene Bank (<https://db.cngb.org/onekp/>)) and PLAZA (<https://bioinformatics.psb.ugent.be/plaza/>).

bioinformatics.psb.ugent.be/plaza/.

Markers for mapping were generated using the 1,001 Genomes Project database (<http://signal.salk.edu/atg1001/index.php>)²³ and the Arabidopsis Mapping Platform (<http://amp.genomics.org.cn/>).

Fig. 4C contains associated Source data.

All data supporting the findings of this study are available within the article, Supplementary Information files or from the corresponding author upon reasonable request.

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Sample size	Sample sizes were consistent with those in our and others' previously published similar studies (e.g. Lee et al. (2018) <i>Plant Cell</i> 30:2038-2056; Zhang et al. (2020) <i>Nat. Plants</i> 6, 394–403). All experiments were performed using at least three independent biological replicates. For transgenic lines, a minimum of eight independent lines were created and analyzed. Sample sizes and the number of times an experiment was performed are described in figure legends and main text.
Data exclusions	No data were excluded.
Replication	At least three independent biological replicates were used for each experiment, with similar results. For transgenic lines, a minimum of eight independent lines were created and analyzed, with similar results.
Randomization	Plants of different genotypes were used as study groups. When more plants were available than the required sample size, plants were chosen randomly for analysis.
Blinding	Experiments were not blinded. Data were collected according to the genotypes of samples. Since comparisons were generally qualitative, blinding was not relevant.

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Methods

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<input checked="" type="checkbox"/>	<input type="checkbox"/> ChIP-seq
<input checked="" type="checkbox"/>	<input type="checkbox"/> Flow cytometry
<input checked="" type="checkbox"/>	<input type="checkbox"/> MRI-based neuroimaging

Antibodies

Antibodies used

For immunoprecipitation: anti-GFP antibody (monoclonal (clone 3E6, lot 1711553), mouse, Molecular Probes by ThermoFisher Scientific, #A-11120).
To detect proteins on Westerns (all antibodies were used at 1:2000 dilution):
anti-GFP antibody (polyclonal, rabbit, Abcam; ab6556)
anti-HA antibody (monoclonal (clone 3F10), rat, Sigma, 11867423001)
anti-rabbit IgG peroxidase-conjugated antibodies (SeraCare/KPL; 5220-0283/04-15-06)
anti-rat IgG peroxidase-conjugated antibodies (SeraCare/KPL; 5220-0364/04-16-06)

Validation

Anti-GFP monoclonal: <https://www.thermofisher.com/antibody/product/GFP-Antibody-clone-3E6-Monoclonal/A-11120> and publications listed thereof.
Anti-GFP polyclonal: <https://www.abcam.com/gfp-antibody-ab6556.html> and publications listed thereof.
Anti-HA: <https://www.sigmaaldrich.com/content/dam/sigma-aldrich/docs/Roche/Bulletin/1/roahahabul.pdf>

