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Inositol pyrophosphates promote the interaction of SPX domains with the coiled-coil motif of PHR transcription factors to regulate plant phosphate homeostasis

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Phosphorus is an essential nutrient taken up by organisms in the form of inorganic phosphate (Pi). Eukaryotes have evolved sophisticated Pi sensing and signaling cascades, enabling them to stably maintain cellular Pi concentrations. Pi homeostasis is regulated by inositol pyrophosphate signaling molecules (PP-InsPs), which are sensed by SPX domain-containing proteins. In plants, PP-InsP-bound SPX receptors inactivate Myb coiled-coil (MYB-CC) Pi starvation response transcription factors (PHRs) by an unknown mechanism. Here we report that a InsP₈-SPX complex targets the plant-unique CC domain of PHRs. Crystal structures of the CC domain reveal an unusual four-stranded anti-parallel arrangement. Interface mutations in the CC domain yield monomeric PHR1, which is no longer able to bind DNA with high affinity. Mutation of conserved basic residues located at the surface of the CC domain disrupt interaction with the SPX receptor in vitro and in planta, resulting in constitutive Pi starvation responses. Together, our findings suggest that InsP₈ regulates plant Pi homeostasis by controlling the oligomeric state and hence the promoter binding capability of PHRs via their SPX receptors.

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hosphorus is an essential building block for many cellular components such as nucleic acids and membranes. It is essential for energy transfer and storage, and can act as a signaling molecule. Pro- and eukaryotes have evolved intricate systems to acquire phosphorus in the form of inorganic phosphate (Pi), to maintain cytosolic Pi concentrations and to transport and store Pi as needed. In green algae and plants, transcription factors have been previously identified as master regulators of Pi homeostasis and Pi starvation responses (PSR)^{1,2}. Phosphorus starvation response 1 (CrPsr1) from *Chlamydomonas* and PHOSPHATE STARVATION RESPONSE 1 (AtPHR1) from Arabidopsis were founding members of plant-unique MYB-type coiled-coil (MYB-CC) transcription factors³. PHR transcription factors were subsequently characterized as regulators of PSRs in diverse plant species⁴⁻⁶. In *Arabidopsis*, there are 15 MYB-CCs with PHR1 and PHL1 controlling the majority of the transcriptional PSRs⁷. Knockout mutations in Arabidopsis thaliana PHR1 (AtPHR1) result in impaired responsiveness of Pi starvation induced (PSI) genes and perturbed anthocyanin accumulation, carbohydrate metabolism, and lipid composition^{2,8,9}. Overexpression of AtPHR1 causes elevated cellular Pi concentrations and impacts the transcript levels of AtPHO2, which codes for an E2 ubiquitin conjugase involved in PSR, via increased production of its micro RNA miR399d^{8,10}. PHR binds to a GNATATNC motif (P1BS), found highly enriched in the promoters of PSI genes and in other cis-regulatory motifs, activating gene expression^{2,7}. AtPHR1 is not only implicated in Pi homeostasis but also in sulfate, iron, and zinc homeostasis, as well as in the adaption to high-light stress 11-14. Moreover, AtPHR1 shapes the plant root microbiome by negatively regulating plant immunity¹⁵.

AtPHR1 and OsPHR2 have been previously reported to physically interact with stand-alone SPX proteins ^{16–19}, additional components of PSR in plants ^{17,20–22}. SPX proteins may regulate PHR function by binding to PHRs under Pi-sufficient condition, keeping the transcription factor from entering the nucleus ^{23–25}. Alternatively, binding of SPX proteins to PHRs may reduce the ability of the transcription factors to interact with their promoter core sequences ^{18,19,23,24,26}. Two mechanisms were put forward regarding the regulation of the SPX–PHR interaction in response to changes in nutrient availability: SPX domains were proposed to act as direct Pi sensors, with the SPX–PHR interaction occurring in the presence of millimolar concentrations of Pi^{18,19}. Alternatively, the integrity of the SPX–PHR complex could be regulated by protein degradation. Indeed, SPX degradation via the 26S proteasome is increased under Pi starvation ^{23,24,27}.

Fungal, plant, and human SPX domains²⁸ have been independently characterized as cellular receptors for inositol pyrophosphates (PP-InsPs), which bind SPX domains with high affinity and selectivity^{29,30}. PP-InsPs consist of a fully phosphorylated myo-inositol ring, carrying one or two pyrophosphate groups at the C1 and/or C5 position, respectively³¹. In plants, inositol 1,3,4-trisphosphate 5/6-kinase catalyzes the phosphorylation of phytic acid (InsP₆) to 5PP-InsP₅ (InsP₇ hereafter)³². The diphosphoinositol pentakisphosphate kinases VIH1 and VIH2 then generate 1,5(PP)₂-InsP₄ (InsP₈ hereafter) from InsP₇^{29,33–35}. Plant diphosphoinositol pentakisphosphate kinases have been genetically characterized to play a role in jasmonate perception and plant defense responses³⁴ and, importantly, in nutrient sensing in Chlamydomonas36 and Arabidopsis29,35. vih1 vih2 double mutants lack the PP-InsP messenger InsP8, over accumulate Pi, and show constitutive PSI gene expression 29,35. A vih1 vih2 phr1 phl1 quadruple mutant rescues the vih1 vih2 seedling phenotypes and displays wild-type-like Pi levels, suggesting that VIH1, VIH2, PP-InsPs, and PHRs are part of a common signaling pathway³⁵. In line with, the AtSPX1-AtPHR1 interaction is reduced in vih1 vih2 mutant plants when compared

to wild type²⁹. Thus, biochemical and genetic evidence implicates $InsP_8$ in the formation of a SPX-PHR complex^{29,35}.

Cellular InsP₈ pools are regulated by nutrient availability at the level of the VIH enzymes themselves. Plant VIH1 and VIH2, and diphosphoinositol pentakisphosphate kinases from other organisms are bifunctional enzymes, with an N-terminal kinase domain that generates InsP₈ from InsP₇ and a C-terminal phosphatase domain that hydrolyzes InsP₈ to InsP₇ and InsP₆^{35,37,38}. The relative enzymatic activities of the two domains are regulated in the context of the full-length enzyme: under Pi starvation, cellular ATP levels are reduced, leading to a reduction of the VIH kinase activity and a reduction of InsP₈^{29,35}. Pi itself acts as an allosteric regulator of the phosphatase activity^{35,37}. Thus, under Pisufficient growth conditions, InsP₈ accumulates and triggers the formation of a SPX–InsP₈–PHR complex. Under Pi starvation, InsP₈ levels drop and the complex dissociates²⁹.

How the $InsP_8$ -bound SPX receptor inactivates PHR function remains to be understood at the mechanistic level. It has been previously reported that AtPHR1 binds P1BS as a dimer². Addition of SPX domains reduces the DNA-binding capacity of PHRs as concluded from electrophoretic mobility shift assays (EMSAs) 18,19,23 . Qi et al. 26 reported that AtPHR1 recombinantly expressed as a maltose-binding protein (MBP) fusion protein forms monomers in solution and binds DNA. This process can be inhibited by preincubating the recombinant transcription factor with AtSPX1 in the presence of high concentrations of $InsP_6^{26}$. A recent crystal structure of the AtPHR1 MYB domain in complex with a promoter core fragment supports a dimeric binding mode of MYB-CC transcription factors 39 . Here we investigate the oligomeric state of PHRs, their DNA-binding kinetics, and the targeting mechanism of the interacting SPX receptors.

Results

PP-InsPs trigger AtSPX1-AtPHR1 complex formation in yeast. The interaction of AtSPX1 with AtPHR1 has been previously characterized in yeast two-hybrid assays¹⁹. We reproduced the interaction of full-length AtPHR1 and AtSPX1 (Fig. 1a), and verified that all four stand-alone AtSPX proteins (AtSPX1-4) interact with a AtPHR1 fragment (AtPHR1²²⁶⁻³⁶⁰) that contains the MYB domain and the CC domain in yeast (Supplementary Fig. 1a). This is in line with previous findings, reporting interaction of SPX domains with larger PHR fragments also containing the MYB and CC domains (AtSPX1-AtPHR1²⁰⁸⁻³⁶² and OsSPX1/2-OsPHR2²³¹⁻⁴²⁶)^{18,19}.

We next tested whether the SPX-PHR interactions observed in yeast are mediated by endogenous PP-InsPs. The putative PP-InsPbinding surface in AtSPX1 was mapped by homology modeling, using the previously determined Chaetomium thermophilum Gde1-InsP₆ complex structure (PDB-ID 5IJJ) as template³⁰. We replaced putative PP-InsP-binding residues from the previously identified phosphate binding cluster (PBC: AtSPX1Y25, K29, K139) and Lysine (K) surface cluster (KSC: AtSPX1K136, K140, K143)30 with alanines (Supplementary Fig. 1b). The resulting AtSPX1PBC and AtSPX1^{KSC} mutant proteins failed to interact with AtPHR1^{226–360} in yeast two-hybrid assays, whereas mutation of a conserved lysine residue outside the putative PP-InsP-binding site (structural control, AtSPX1K81) to alanine had no effect (Supplementary Fig. 1b). We next deleted the known yeast PP-InsP kinase Vip1, which converts InsP₆ to 1PP-InsP₅ and InsP₇ to InsP₈, or the PP-InsP kinase Kcs1, which converts InsP₆ to InsP₇ and 1PP-InsP₅ to InsP₈^{40,41} (Supplementary Fig. 1c, d). We found that deletion of either kinase reduced the interaction between wild-type AtSPX1 and AtPHR1²²⁶⁻³⁶⁰ (Supplementary Fig. 1c). The interaction between the plant brassinosteroid receptor kinase BRI1 and the inhibitor protein BKI1, known to occur independently of

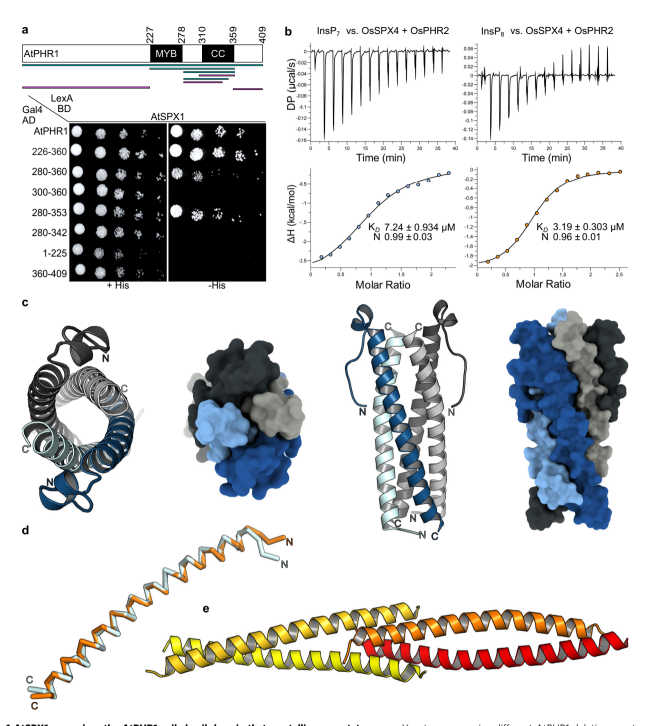


Fig. 1 AtSPX1 recognizes the AtPHR1 coiled-coil domain that crystallizes as a tetramer. a Yeast co-expressing different AtPHR1 deletion constructs fused to the Gal4-activation domain (AD; prey) and full-length wild-type AtSPX1 fused to the LexA-binding domain (BD; bait) were grown on selective SD medium supplemented with histidine (+His; co-transformation control) or lacking histidine (-His; interaction assay) to map a minimal fragment of AtPHR1 sufficient for interaction with AtSPX1. Shown are serial dilutions from left to right. A schematic overview of the tested interacting (in cyan) and non-interacting (in magenta) AtPHR1 fragments is shown alongside (MYB, MYB-DNA-binding domain; CC, coiled-coil domain). **b** Isothermal titration calorimetry assays of InsP₇ (400 μM 5PP-InsP₅; left panel) and InsP₈ (500 μM 1,5(PP)₂-InsP₄; right panel) binding to OsSPX4-OsPHR2 (30 μM), respectively. Raw heats per injection are shown in the top panel and the bottom panel represents the integrated heats of each injection, fitted to a one-site binding model (solid line). The insets show the dissociation constant (K_D) and binding stoichiometry (N) (±fitting error). **c** Ribbon and surface diagrams of the AtPHR1 CC four-stranded anti-parallel tetramer. Helices contributing to the dimer interface are shown in light and dark blue, respectively. Corresponding, symmetry-related helices completing the tetramer are shown in light and dark gray. **d** Structural superposition of two AtPHR1 core CC helices (C_a trace, in light blue) and ScCtp1 (PDB-ID 4 × 01, in orange)⁴⁴. R.m.s.d. is ~1 Å comparing 45 corresponding C_a atoms. **e** Ribbon diagram of the ScCtp1 dimer-of-dimers CC domain, with contributing helices colored from yellow to red.

PP-InsPs⁴², was not affected in either Δ vip1 or Δ kcs1 mutants (Supplementary Fig. 1c).

Using quantitative isothermal titration calorimetry (ITC)-binding assays, we have previously determined dissociation constants (K_D) for InsP₆ and InsP₇ binding to a OsSPX4–OsPHR2 complex to be ~50 and ~7 μ M, respectively³⁰. A side-by-side comparison of InsP₇ and InsP₈ binding to OsSPX4–OsPHR2 by ITC revealed dissociation constants of ~7 and ~3 μ M, respectively (Fig. 1b). Taken together, the SPX–PHR interaction is mediated by PP-InsPs, with the bona fide Pi signaling molecule InsP₈ being the preferred ligand in vitro.

AtSPX1 interacts with a unique four-stranded coiled-coil domain in AtPHR1. We next mapped the SPX-PP-InsP-binding site in AtPHR1 to a fragment (AtPHR1²⁸⁰⁻³⁵³), which comprises the CC domain and a 30-amino acid spanning N-terminal extension, in yeast two-hybrid experiments (Fig. 1a). We sought to crystallize an AtSPX1-PP-InsP-AtPHR1 complex either in the pre- or absence of P1BS fragments. We obtained crystals of a putative AtSPX1-InsP₈-AtPHR1²⁸⁰⁻³⁶⁰ complex diffracting to 2.4 Å resolution and solved the structure by molecular replacement, using isolated SPX domain structures as search models³⁰. Iterative cycles of model building and crystallographic refinement yielded, to our surprise, a well-refined model of AtPHR1^{280–360} only (see "Methods"). Analysis with the program PISA revealed the presence of a crystallographic tetramer in which four long αhelices fold into an unusual anti-parallel four-stranded coiled-coil (Fig. 1c). AtPHR1^{280–360} residues 292–356 and 310–357 are visible in the electron density maps from chain A and B, respectively. Residues 292–311 in chain A fold into a protruding loop region that harbors a small α-helix, and appear disordered in chain B (Fig. 1c). The anti-parallel a-helices in AtPHR1 closely align with a root mean square deviation (r.m.s.d.) of ~0.5 Å comparing 45 corresponding C_α atoms. Structural homology searches with the program DALI⁴³ returned different coiled-coil structures, with a monomer of the tetrameric coiled-coil domain of the yeast transcription factor Ctp1 representing the closest hit (DALI Z-score 5.9, r.m.s.d. is ~1 Å comparing 45 corresponding C_a atoms) (Fig. 1d)⁴⁴. However, no anti-parallel four-stranded coiled-coil domain with structural similarity to AtPHR1 was recovered, with, for example, the Ctp1 dimer-of-dimers domain having a very different configuration (Fig. 1e)⁴⁴.

We next assessed the oligomeric state of AtPHR^{280–360} using size-exclusion chromatography coupled to right-angle light scattering (SEC-RALS) and determined an apparent molecular weight of ~37.5 kDa, thus confirming that the isolated AtPHR1 CC forms tetramers in solution (theoretical molecular weight of the monomer is ~9.5 kDa) (Fig. 2a). Two additional crystal structures of AtPHR1^{280–360} obtained in different crystal lattices all revealed highly similar tetrameric arrangements (Supplementary Fig. 2 and Supplementary Table 5). Although we found purified full-length AtPHR1 to be too unstable for SEC-RALS analysis, full-length OsPHR2 formed a tetramer in solution (Supplementary Fig. 3). In contrast, untagged AtPHR1^{222–358}, which comprises the CC and the MYB domains only, runs as a dimer (Fig. 2a, b black traces), in agreement with earlier reports².

Mutations in the CC domain abolish AtPHR1 oligomerization and DNA binding in vitro. The observed oligomeric state differences between full-length OsPHR2, AtPHR1^{280–360} (CC), and AtPHR1^{222–358} (MYB-CC) prompted us to investigate the putative dimerand tetramerization interfaces in our AtPHR1 structures with the program PISA⁴⁵. We found the dimerization (~1400 Å² buried surface area) and the tetramerization (~1900 Å² buried surface area) interfaces to be mainly

formed by hydrophobic interactions (Supplementary Fig. 4a, b). Both interfaces are further stabilized by hydrogen bond interactions and several salt bridges (Supplementary Fig. 4a, b). Importantly, all contributing amino-acids represent sequence fingerprints of the plant-unique MYB-CC transcription factor subfamily and are highly conserved among different plant species (Supplementary Fig. 4c). We identified residues specifically contributing to the formation of a CC dimer (Olig1: AtPHR1^{L319}, AtPHR1^{L337}, shown in cyan in Fig. 2 and Supplementary Fig. 4) or tetramer (Olig2: AtPHR1^{L317}, AtPHR1^{L327}, AtPHR1^{L341}, shown in dark orange in Fig. 2 and Supplementary Fig. 4) in our different CC structures (Supplementary Table 5). We replaced these residues by asparagine to generate two triple mutants in AtPHR1^{222–358} and AtPHR1^{280–360}, respectively. We found in SEC-RALS assays that both mutant combinations dissolved AtPHR1^{280–360} tetramers and AtPHR1^{222–358} dimers into monomers, respectively (Fig. 2a, b).

Analysis of the secondary structure content of wild-type AtPHR^{280–360} using circular dichroism (CD) spectroscopy revealed a 100% α -helical protein (Supplementary Fig. 5a), in agreement with our structural model of the AtPHR1 CC domain (Fig. 1c). In contrast, we estimated the secondary structure content of the Olig1 and Olig2 mutant proteins to be ~50% α -helical and ~50 random coil (Supplementary Fig. 5a). The CD melting spectrum for wild-type AtPHR^{280–360} indicated the presence of a well-folded protein with a melting temperature ($T_{\rm m}$) of ~50 °C, while we could not reliably determine $T_{\rm m}$'s for the Olig1 and Olig2 mutant proteins (Supplementary Fig. 5b). We conclude that mutation of either AtPHR1^{L319}, AtPHR1^{I333}, AtPHR1^{I337} or AtPHR1^{L317}, AtPHR1^{L327}, AtPHR1^{I341} to asparagine disrupts the tetrameric coiled-coil domain of AtPHR1 and affects the structural integrity of the contributing α -helix.

It has been recently reported that the AtPHR1 MYB domain associates with its target DNA as a dimer³⁹. We thus studied the capacity of AtPHR1^{222–358} oligmerization mutants to interact with the P1BS in qualitative EMSA and quantitative grating-coupled interferometry (GCI) assays. AtPHR1^{222 – 358} Olig1 could still interact with the P1BS in EMSAs indistinguishable from wild type (Fig. 2c). However, AtPHR1^{222 – 358} Olig1 and AtPHR1^{222 – 358} Olig2 bound a biotinylated P1BS immobilized on the GCI chip with ~20-fold reduced affinity when compared to the wild-type control (Fig. 2d-f). Together, our experiments suggest that PHR1 may exist as a tetramer or dimer in solution, and that disruption of its plant-unique CC domain interface reduces the capacity of the transcription factor to bind its DNA recognition site.

CC surface mutations abolish PHR-SPX interactions but do not interfere with DNA binding in vitro. We next sought to identify the binding site for SPX in the PHR CC domain. In our structures, a conserved set of basic residues maps to the surface of the four CC helices (shown in magenta in Fig. 3a and Supplementary Fig. 4c). A similar set of surface-exposed basic residues has been previously found to form the binding site for PP-InsPs in various SPX receptors³⁰. Mutation of AtPHR1^{K325}, AtPHR1^{H328}, and AtPHR1^{R335}, but not of AtPHR1^{K308}, AtPHR1R318, and AtPHR1R340 to alanine, disrupted the interaction of AtPHR1 with AtSPX1 in yeast (Fig. 3b). We simultaneously mutated the residues corresponding to AtPHR1K325, AtPHR1H328, and AtPHR1R335 to alanine in OsPHR2 (OsPHR2KHR/A). The mutant transcription factor showed no detectable binding to OsSPX4-InsP7 in quantitative ITC assays, but maintained the ability to bind the P1BS (Fig. 3c, d). In line with this, mutation of the KHR motif does not alter the oligomeric state of AtPHR1²⁸⁰⁻³⁶⁰ as concluded from SEC-RALS experiments (Fig. 2a magenta traces).

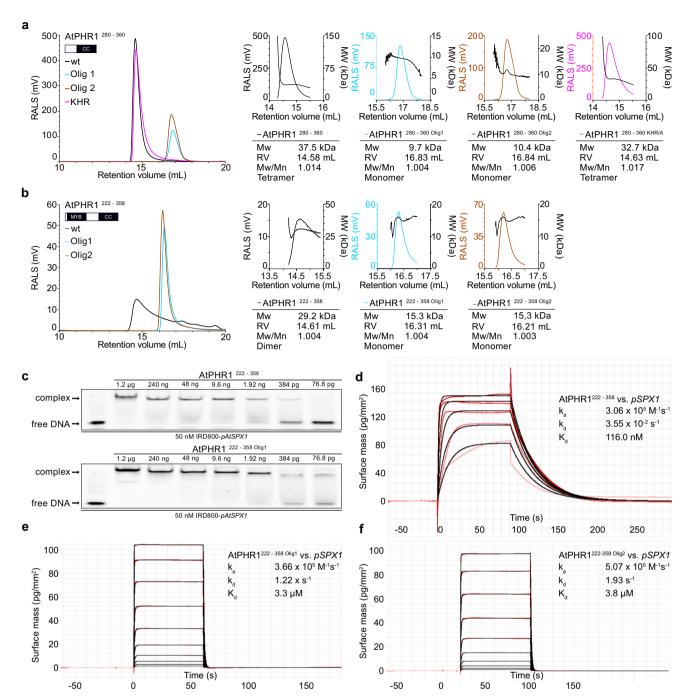


Fig. 2 Mutations in the AtPHR1 coiled-coil domain impair oligomerisation and DNA binding. a Analytical size-exclusion chromatography traces of wild-type AtPHR1 CC (wt, black line), AtPHR1²⁸⁰⁻³⁶⁰ Olig1 (Olig1, cyan), AtPHR1²⁸⁰⁻³⁶⁰ Olig2 (Olig2, orange), and of AtPHR1²⁸⁰⁻³⁶⁰ KHR/A (KHR, magenta). The corresponding right-angle light scattering (RALS) traces are shown alongside, the molecular masses are depicted by a black line. Table summaries provide the molecular weight (Mw), retention volume (RV), dispersity (Mw/Mn), and the derived oligomeric state of the respective sample. **b** Analysis of AtPHR1 MYB-CC (AtPHR1²²²⁻³⁵⁸) as described in **a. c** Qualitative comparison of the interaction of AtPHR1²²²⁻³⁵⁸ (upper panel) or AtPHR1²²²⁻³⁵⁸ Olig1 (lower panel) binding to IRD800-*pAtSPX1* in electrophoretic mobility shift assays. Experiment was performed twice with similar results. **d-f** Quantitative comparison of the interaction of AtPHR1²²²⁻³⁵⁸, AtPHR1²²²⁻³⁵⁸ Olig1, or AtPHR12²²⁻³⁵⁸ Olig2 with *pSPX1* by grating-coupled interferometry (GCI). Sensorgrams show raw data (red lines) and their respective fits (black lines). Table summaries provide the derived association rate (k_a), the dissociation constant (K_d), and the dissociation constant (K_d).

Using nuclear magnetic resonance (NMR) spectroscopy, we next tested whether the KHR motif in AtPHR1 is directly involved in PP-InsP ligand recognition. We titrated InsP₈ into ¹⁵N, ²H-labeled AtPHR1^{280–360}, and recorded TROSY spectra using potassium phosphate (KPi) as control. Five backbone amide moieties exhibited chemical shift perturbations in the presence of

 $InsP_8$ but not in the presence of the KPi control. We acquired titration spectra using increasing concentrations of $InsP_8$ and estimated dissociation constants for $InsP_8$ based on three representative peaks (Supplementary Fig. 6). The derived dissociation constant is in the millimolar range and saturation could not be reached in the available concentration range

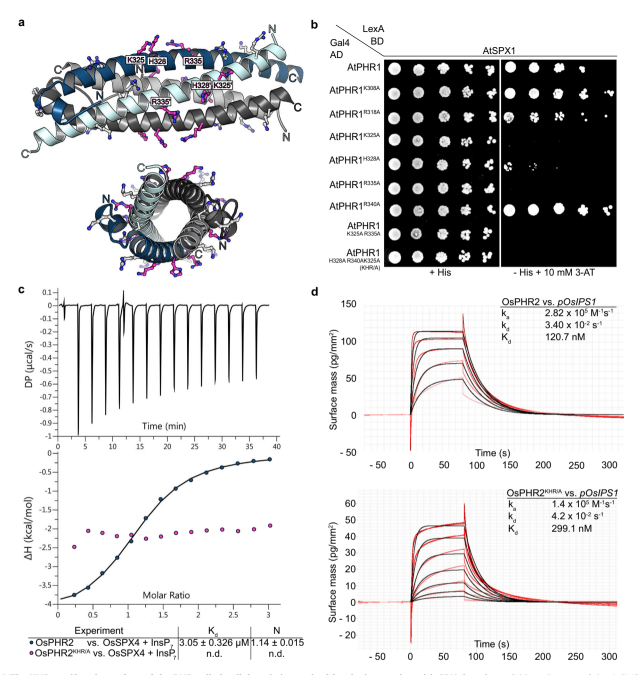


Fig. 3 The KHR motif at the surface of the PHR coiled-coil domain is required for the interaction with SPX domains. a Ribbon diagram of the AtPHR1 CC domain with conserved basic residues located at the surface of the domain shown in bonds representation. The KHR motif (AtPHR1^{K325}, AtPHR1^{H328}, AtPHR1^{R335}) is highlighted in magenta. **b** Mutational analysis of the basic residues in AtPHR1 CC. Yeast co-expressing AtPHR1²²⁶⁻³⁶⁰ variants in which surface-exposed basic residues have been replaced with alanine fused to the Gal4-AD (prey) and AtSPX1 fused to the LexA-BD (bait) were grown on selective SD medium supplemented with histidine (+His; co-transformation control) or lacking histidine and supplemented with 10 mM 3-amino-1,2,4-triazole (3-AT) (—His + 10 mM 3-AT; interaction assay) to identify residues required for interaction with AtSPX1 in yeast two-hybrid assays. Shown are serial dilutions from lift to right. **c** Isothermal titration calorimetry (ITC) assay of wild-type OsPHR2 and OsPHR2^{KHR/A} (300 μM) vs. OsSPX4 (20 μM)-5PP-InsP₅ (100 μM). Raw heats per injection are shown in the top panel, the bottom panel represents the integrated heats of each injection, fitted to a one-site binding model (solid line). The insets show the dissociation constant (K_D) and binding stoichiometry (N) (± fitting error, n.d. no detectable binding). **d** Quantitative comparison of the interaction of OsPHR2 (top panel) or OsPHR2KHR/A (bottom panel) with *pOsIPS1* by GCI. Sensorgrams show raw data (red lines) and their respective fits (black lines). The insets show summarize association rates (K_A), dissociation rates (K_A), and the dissociation constant (K_A) of the respective sample.

(Supplementary Fig. 6). We next repeated the same set of experiments using the AtPHR1^{KHR/A} mutant protein, located the same peaks in the transverse relaxation-optimized spectroscopy (TROSY) spectra and estimated similar dissociation constants (Supplementary Fig. 6).

Taken together, three highly conserved basic residues located at the surface of the PHR coiled-coil domain are critical for the interaction with the PP-InsP-bound SPX receptor (Supplementary Fig. 4c). The NMR titration experiments suggest that AtPHR1 does not contribute to the specific binding of InsP $_8$ and

that the low affinity interaction between the CC domain and $InsP_8$ does not involve the KHR motif (Supplementary Fig. 6).

Mutation of the AtPHR1 KHR motif impairs AtSPX1 binding and Pi homeostasis in planta. We next tested whether mutation of the SPX-binding site in AtPHR1 can modulate its function in Pi homeostasis in *Arabidopsis*. We expressed wild-type and pointmutant versions of AtPHR1 carrying an N-terminal FLAG tag under the control of its native promoter in a phr1-3 loss-offunction mutant⁸. At seedling stage, we found that AtPHR1 single, double, and triple point mutations complemented the previously characterized Pi deficiency phenotype of phr1-3^{2,8}. (Fig. 4a and Supplementary Fig. 7a). After transferring the seedlings to soil, variable growth phenotypes became apparent 21 days after germination (DAG) (Supplementary Fig 7b). From three independent lines per genotype, we selected one line each showing similar AtPHR1 transcript levels for all experiments shown in Fig. 4 (Fig. 4 and Supplementary Fig. 8a). Comparing these lines, we found that AtPHR1^{K325A,R335A} double and AtPHR1^{K325A},H328A,R335A triple mutants, but not the single mutants, displayed severe growth phenotypes, with the triple mutant showing the strongest defects (Fig. 4a). We next determined cellular Pi levels in all independent lines and found that (i) Pi levels are positively correlated with AtPHR1 expression levels (Supplementary Fig. 8a), that (ii) all AtPHR1 mutant proteins tested accumulate Pi to significantly higher levels when compared to wild type and *phr1-3*, and that (iii) the AtPHR1^{K325A,H328A,R335A} triple mutant displayed the highest Pi levels (Fig. 4b and Supplementary Fig. 8b-d). In line with this, PSI gene expression is misregulated in AtPHR1^{K325A,R335A} double and AtPHR1^{K325A,H328A,R335A} (AtPHR1^{KHR/A}) triple mutants (Fig. 4c). In co-immunoprecipitation assays in Nicotiana benthamiana and in Arabidopsis, we found the interaction of AtPHR1^{K325A,H328A,R335A} with AtSPX1 to be reduced when compared to wild-type AtPHR1 (Fig. 4d and Supplementary Fig. 8).

We next studied the genetic interaction between PHR1 and VIH1/2. As previously reported, the severe phenotypes of vih1-2 vih2-4 seedlings are partially rescued in the phr1 phl1 vih1-2 vih2-4 quadruple mutant, suggesting that VIH1/2-generated InsP8 regulates the activity of PHR1 and PHL1 by promoting the binding of SPX receptors^{29,35}. We performed the orthogonal genetic experiment, by complementing the phr1 phl1 mutant with AtPHR1KHR/A expressed under the control of the AtPHR1 promoter and carrying a N-terminal enhanced green fluorescent protein (eGFP) tag (Fig. 4e, see "Methods"). The complemented lines displayed intermediate growth phenotypes and constitutive PSI gene expression (Fig. 4e, f). Thus, SPX-InsP₈ mediated regulation of PHR1 and PHL1 has to be considered one of several PP-InsP regulated processes affected in the *vih1-2 vih2-4* mutant. Together, our in vivo experiments reveal that SPX receptors interact with the CC domain of AtPHR1 via the surface-exposed Lys325, His328, and Arg335, and that this interaction negatively regulates PHR activity and PSRs.

Discussion

PHR transcription factors have been early on recognized as central components of the PSR in green algae and in plants, directly regulating the expression of PSI genes^{1,2,7}. In *Arabidopsis* and in rice *spx* mutants of then unknown function also showed altered PSI gene expression^{20,22}. This genetic interaction was later substantiated by demonstrating that stand-alone plant SPX proteins can interact with PHR orthologs from *Arabidopsis* and rice^{18,19,23}. The biochemical characterization of SPX domains as cellular receptors for PP-InsPs and the genetic identification of VIH kinases

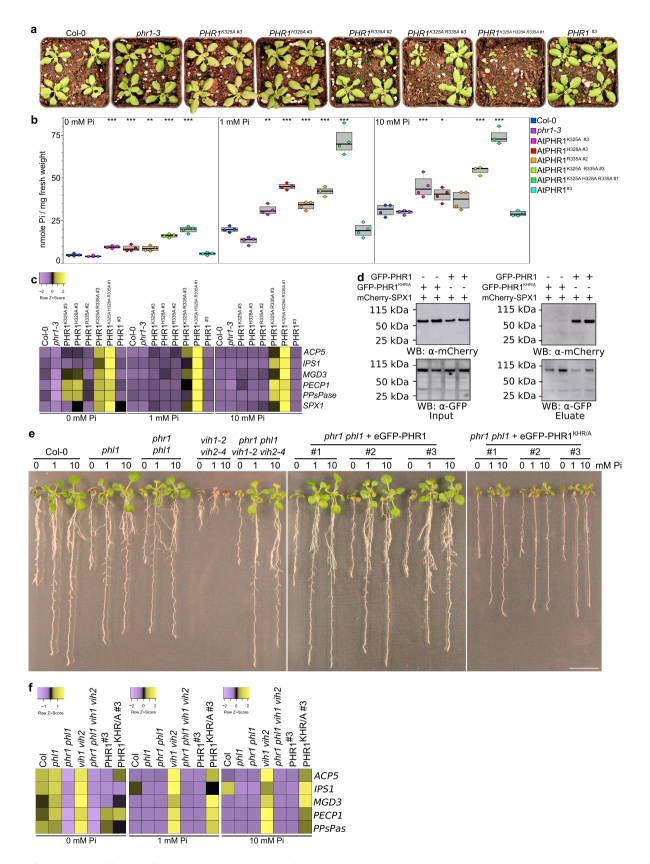
as master regulators of PSR in plants suggested that PP-InsPs, and specifically InsP₈ mediates the interaction of SPX proteins with PHRs in response to changing nutrient conditions^{29,30,35}.

Our quantitative DNA-binding assays demonstrate that AtPHR1 MYB-CC binds the P1BS from the AtSPX1 promoter with high affinity ($K_{\rm d} \sim 0.2~\mu{\rm M}$), in agreement with previously reported binding constants for different MYB-CC constructs ($K_{\rm d} \sim 0.01-0.1~\mu{\rm M}$)^{26,39}. Different oligomeric states have been reported for various PHR MYB-CC constructs^{2,26}. Our AtPHR1 MYB-CC construct behaves as a dimer in solution, consistent with the recently reported crystal structure of the AtPHR1 MYB-DNA complex and with earlier reports^{2,39}. Purified full-length OsPHR2, however, appears to be a homotetramer in solution (Supplementary Fig. 3).

In yeast two-hybrid assays, we found that AtSPX1-4 all are able to interact with AtPHR1 (Fig. 1a and Supplementary Fig. 1a). We mapped their conserved interaction surface to the plantunique CC motif of PHRs (Fig. 1a). Crystal structures of this fragment reveal an unusual, four-stranded anti-parallel coiled-coil domain (Fig. 1c). Given the fact, that AtPHR1 MYB-CC forms dimers in solution (Fig. 2b), we cannot exclude the possibility that the CC tetramers represent crystal packing artifacts. However, we did observe identical CC tetramers in three independent crystal lattices (Supplementary Fig. 2) and in solution (Fig. 2a). The residues contributing to the dimer and to the tetramer interfaces are highly conserved among all plant MYB-CC transcription factors (Supplementary Fig. 4). Mutation of either interface blocks AtPHR1 oligomerization in vitro (Fig. 2a, b) and reduces DNA binding (Fig. 2d-f). An attractive hypothesis would thus be that AtPHR1 binds its target promoter as a dimer, but can potentially form homo-tetramers, or hetero-tetramers with other MYB-CC type transcription factors sharing the conserved, plant-unique CC structure and sequence (Supplementary Fig. 4). Notably, PHR1 PHL1 heteromers have been previously described⁷.

We found that SPX-PHR complex formation is mediated by endogenous PP-InsPs in yeast cells, as deletion of the yeast PP-InsP kinases Vip1 and Kcs1 abolished the interaction, and mutation of the PP-InsP-binding surface in AtSPX1 interfered with AtPHR1 binding (Supplementary Fig. 1b, c). In line with this, SPX-PHR complexes are found dissociated in vih1 vih2 mutant plants²⁹. It is of note that the observed differences in binding affinity for InsP₇ and InsP₈ to SPX-PHR in vitro (Fig. 1b)²⁹ cannot fully rationalize the apparent preference for InsP₈ in vivo^{29,35}. We identified the binding surface for SPX-InsP₈ by locating a set of highly conserved basic residues exposed at the surface of the CC domain (Fig. 3a and Supplementary Fig. 4). Mutation of this KHR motif did not strongly impact the ability of isolated OsPHR2 to bind pOsIPS1 in vitro (Fig. 3d), but disrupted the interaction of AtPHR1 with AtSPX1 in yeast (Fig. 3b). The corresponding mutations in OsPHR2 had a similar effect on the interaction with OsSPX4 in quantitative ITC assays (Fig. 3c). Expression of AtPHR1KHR/A in the phr1-3 mutant resulted in Pi hyperaccumlation phenotypes and constitutive PSI gene expression in Arabidopsis (Fig. 4a-c). The intermediate growth phenotypes of vih1 vih2 phr1 phl1 mutants complemented with AtPHR1KHR/A clearly suggests that PP-InsPs do not only regulate the activity of PHR1 and PHL1 in plants, but likely the function of other (SPX domain-containing) proteins³⁰ (Fig. 4e). Notably, binding of AtPHR1KHR/A to AtSPX1 was reduced in co-immunoprecipitation assays when compared to wild-type AtSPX1 (Fig. 4d and Supplementary Fig. 9). Thus, our and previous finding suggest that InsP₈ can promote the association of SPX receptors and PHR transcription factors. The newly identified basic surface area in PHR CC, harboring the conserved KHR motif, likely forms part of the SPX-PHR complex interface (Fig. 3a).

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It is of note that addition of AtSPX1 to AtPHR1 has been previously demonstrated to reduce AtPHR1's ability to bind to P1BS in the presence of InsP₆²⁶. We could not quantify these interactions in ITC or GCI binding assays, as PHR CC formation is much preferred over SPX–InsP₈ binding at the protein concentrations required in these assays. We speculate that InsP₈

bound SPX proteins can bind to the basic residues we identified in the PHR CC domain to control the oligomeric state and hence the promoter binding capacity of PHRs. As these residues are conserved among all plant MYB-CC proteins this may suggest that transcription factors outside the PHR subfamily may be regulated by SPX domains and PP-InsPs, possibly rationalizing

Fig. 4 Mutation of the AtPHR1 KHR motif impairs interaction with AtSPX1 and Pi homeostasis in planta. a Growth phenotypes of Col-0 wild type, *phr1-3* of *phr1-3* complementation lines expressing FLAG-AtPHR1, FLAG-AtPHR1^{K325A}, FLAG-AtPHR1^{K325A}, FLAG-AtPHR1^{K325A}, FLAG-AtPHR1^{K325A}, FLAG-AtPHR1^{K325A}, FLAG-AtPHR1^{K325A}, and FLAG-AtPHR1^{K325A} under the control of the *AtPHR1* promoter at 21 DAG grown in Pi-sufficient conditions. One representative line per complementation construct is shown (specified by a #), additional lines are shown in Supplementary Fig. 4. **b** Plot representing the cellular Pi content of the lines shown in **a**. Bold black line, median; box, interquartile range (IQR); whiskers, lowest/highest data point within 1.5 IQR of the lower/upper quartile. Seedlings were germinated and grown on vertical ^{1/2}MS plates for 8 days, transferred to ^{1/2}MS plates supplemented with either 0, 1, or 10 mM Pi and grown for additional 7 days. For each line, four plants were measured in technical duplicates. Pi contents of all lines can be found in Supplementary Fig. 4 (*p < 0.05; **p < 0.01; ***p < 0.001). **c** Heat maps of PSI marker gene (*ACP5, IPS1, MGD3, PECP1, PPsPase, SPX1*) expression analyses of the lines shown in **a**, represented as *Z*-scores. For each line, three biological replicates were analyzed in technical triplicates by qRT-PCR. **d** Co-immunoprecipitation (Co-IP) experiment assessing the ability for immobilized GFP-AtPHR1 and GFP-AtPHR1^{KHR/A} to interact with mCherry-AtSPX1 in *N. benthamiana*. Input western blottings are shown alongside. Experiment was performed twice with similar results. **e** Genetic interactions in the VIH-PHR signaling pathway. Col-0 wild type and the indicated mutant seedlings were grown on ^{1/2}MS plates for 7 DAG, transferred to ^{1/2}MS plates supplemented with either 0, 1, or 10 mM Pi and grown for additional 7 days. For complementation analyses, wild-type AtPHR1 or AtPHR1^{KHR/A} was expressed as an N-terminal eGFP fusion protein under the control of the

the severe phenotypes of vih1 vih2 mutant plants (Fig. 4e and Supplementary Fig. 4). The recent findings that VIH kinases and PHRs act together in plant PSR and that SPX-PHR complexes are dissociated in vih1 vih2 mutants further suggest that InsP₈ is the bona fide signaling molecule promoting the association between SPX domains and MYB-CCs^{29,35}. Repressive SPX-PHR complexes consequently form only under Pi-sufficient conditions, where InsP₈ levels are high^{29,35}. Under Pi starvation, when InsP₈ levels are reduced, SPX-PHR complexes dissociate, enabling the transcription factors to acquire the oligomeric state required for high affinity promoter binding. The physiological and mechanistic investigation of this central process may, in the long term, contribute to the development of Pi starvation resilient crops. This could in turn sustain the use of the essential and nonrenewable resource rock phosphate, which is currently consumed at an alarming scale.

Methods

Molecular cloning, constructs, and primers. For a detailed description of the cloning strategies, constructs and primers used in this study, please refer to Supplementary Tables 2 and 3.

Generation of stable transgenic A. thaliana lines. All stable transgenic A. thaliana lines are listed in Supplementary Table 1. Constructs were introduced into Agrobacterium tumefaciens strain pGV2260 and A. thaliana plants were transformed via floral dipping⁴⁶. Transformants were identified by mCherry fluorescence with a Zeiss Axio Zoom.V16 stereo microscope (mRFP filter) and a HXP200C illuminator. Homozygous T3 lines have been identified for complementation lines expressing FLAG-AtPHR1, FLAG-AtPHR1K325A, FLAG-AtPHR1H328A, and FLAG-AtPHR1R335A under the control of the native AtPHR1 promoter. For complementation lines expressing FLAG-AtPHR1K325A R335A and FLAG-AtPHR1^{K325A} H^{328A} R^{335A} under the control of the native *AtPHR1* promoter, T2 lines were used throughout, and homozygous and heterozygous transformants were selected for all experiments by mCherry fluorescence as described above. T3 homozygous lines expressing eGFP-PHR1 under the control of the AtPHR1 promoter were identified by their Hygromycin resistance. PHR1 was amplified from Arabidopsis cDNA and introduced into pH7m34GW binary vector. Point mutations were introduced by site-directed mutagenesis (primers are listed in Supplementary Table 3).

Yeast two-hybrid experiments. (Screen) AtSPX1 $^{1-252}$ was used as a bait and screened against an *A. thaliana* seedling cDNA library by Hybrigenics Services. AtSPX1 $^{1-252}$ was cloned into the pB29 vector providing a C-terminal LexA-DNA-binding domain and transformed into yeast strain L40 α Gal4 (MATa). Prey genes were cloned into the pP6 vector providing a N-terminal Gal4-activation domain and transformed into yeast strain YHGX13 (MAT α). After mating haploid bait and prey strains, positive interactions were detected by growth on histidine deficient medium.

Yeast strains and media. For all experiments, either the diploid TATA strain (Hybrigenics Services) or the haploid L40 strain was used (Supplementary Table 2). Cells were routinely maintained on yeast extract-peptone-adenine-dextrose (YPAD) plates (20 g/L glucose, 20 g/L bacto-peptone, 10 g/L yeast extract, 0.04 g/L adenine hemisulfate, and 20 g/L agar). Experiments were performed on synthetic dropout (SD) plates (6.7 g/L yeast nitrogen base with adenine hemisulfate

and without leucine, tryptophan, and histidine, 20 g/L glucose, and 20 g/L agar) supplemented with $0.076\,\mathrm{g/L}$ histidine or $10\,\mathrm{mM}$ 3-amino,1,2,4-triazole (3-AT).

Yeast transformation. One yeast colony was resuspended in 500 μL sterile $\rm H_2O$, plated on YPAD plates, and grown for 2 days until the whole plate was covered with yeast. Yeast cells were then resuspended in 50 mL YPAD liquid medium and the OD600nm was determined (2 × 10^6 cells/mL were used for one transformations). Cells were centrifuged at 3000 × g and 4 °C for 5 min, resuspended in 25 mL Tris-EDTA (TE) buffer, centrifuged again at 3000 × g and 4 °C for 5 min, resuspended in 2 mL LiAc/TE buffer, centrifuged at 16,000 × g and room temperature (RT) for 15 s, and finally resuspended in 50 μL/transformation LiAc/TE buffer. The transformation mix (0.5 μg bait plasmid, 0.5 μg prey plasmid, 10 μL ssDNA (10 mg/mL), 50 μL yeast cells, 345 μL 40% (w/v) PEG3350 in LiAC/TE) was prepared and incubated at 30 °C for 45 min, followed by incubation at 42 °C for 30 min. Finally, yeast cells were centrifuged at 6500 × g and RT for 15 s, resuspended in TE buffer, plated on SD plates lacking leucine and tryptophan, and incubated at 30 °C for 3 days.

Yeast spotting dilution assay. Positive transformants were selected on SD plates without tryptophan and leucine, and incubated at 30 °C for three days. Cells were counted, washed in sterile water and spotted in 5 times dilution (5000, 1000, 200, 40, and 8 cells) on SD plates without either tryptophan and leucine, or tryptophan, leucine, and histidine supplemented with 10 mM 3-AT. Plates were incubated at 30 °C for 3 days.

Protein expression and purification. For in vitro biochemistry, AtPHR1²⁸⁰⁻³⁶⁰ wt/ KHR/Olig1/Olig2 and AtPHR1222-358 wt/Olig1/Olig2 were cloned into the pMH-HT protein expression vector, providing a N-terminal 6 × His affinity tag with a tobacco etch virus (TEV) protease recognition site. OsPHR2¹⁻⁴²⁶ wt/KHR/A was cloned into the pMH-HSgb1T protein expression vector, providing a N-terminal 8×His-Strep-GB1 affinity tag with a TEV cleavage site. OsSPX2^{1–321} was cloned into the pMH-HSsumo protein expression vector, providing a N-terminal 8xHis-Strep-Sumo affinity tag. All constructs were transformed into Escherichia coli BL21 (DE3) (argU, ileY, leuW) RIL cells. For recombinant protein expression, cells were grown at 37 °C in terrific broth (TB) medium to an OD_{600nm} of ~0.6. After reducing the temperature to 18 °C, protein expression was induced with 0.3 mM isopropyl β-D-galactoside (IPTG) for 16 h. Cells were centrifuged at 4500 \times g and 4 °C for 1 h, resuspended in lysis buffer (50 mM Tris-HCl pH 7.8, 500 mM NaCl, 0.1% (v/v) IGEPAL, 1 mM MgCl, 2 mM β-mercaptoethanol), snap-frozen in liquid nitrogen, and stored at -80 °C. For protein preparation, cells were thawed, supplemented with cOmpleteTM EDTA-free protease inhibitor cocktail (Roch), DNaseI, and lysozyme, and disrupted using a sonicator. Cell lysates were centrifuged at $7000 \times g$ and 4 °C for 1 h, sterile filtered, supplemented with 20 mM imidazole, and loaded onto a 5 mL HisTrap HP Ni²⁺ affinity column (GE Healthcare). After washing with several column volumes (CVs) of lysis buffer supplemented with 20 mM imidazole, high-salt buffer (50 mM Tris-HCl pH 7.8, 1 M NaCl, 2 mM β-mercaptoethanol), and high-phosphate buffer (200 mM K₂HPO₄/KH₂PO₄ pH 7.8, 2 mM β-mercaptoethanol), proteins were eluted in a gradient from 20 to 500 mM imidazole in lysis buffer. The purified proteins were cleaved by TEV or Sumo protease overnight at 4 °C (1:100 ratio) and dialyzed against lysis buffer for PHR1²²²⁻³⁵⁸ wt/Olig1/Olig2 and PHR1²⁸⁰⁻³⁶⁰ wt/KHR/Olig1/Olig2 fragments, against modified lysis buffer (25 mM Tris-HCl pH 7.8, 300 mM NaCl, 0.1% (v/v) IGEPAL, 1 mM MgCl, 2 mM β-mercaptoethanol) for OSPHR2¹⁻⁴²⁶ wt/KHR/A, and against modified anion exchange buffer (20 mM Tris-HCl pH 6.5, 500 mM NaCl) for OsSPX4¹⁻³²¹. PHR1²⁸⁰⁻³⁶⁰ wt/KHR/Olig1/Olig2 and OsPHR2¹⁻⁴²⁶ wt/KHR/A were subjected to a second Ni²⁺ affinity purification in either lysis buffer or modified lysis buffer, respectively, and the flow-throughs were collected and concentrated. PHR1²²²⁻³⁵⁸ wt/Olig1/Olig² were subjected to cation exchange (50 mM HEPES pH 7.5, 50-1000 mM NaCl) and OsSPX41-321 was subjected to anion exchange (20 mM Tris-HCl, 50-1000 mM NaCl). Fractions corresponding to

the respective proteins were pooled and concentrated. All proteins were loaded onto a HiLoad Superdex 75 pg HR26/60 column (GE Healthcare), pre-equilibrated in gel filtration buffer A (20 mM Tris/HCl pH 7.5, 300 mM NaCl, 0.5 mM tris(2-carbox-yethyl)phosphine (TCEP)) for PHR1 $^{222-358}$ wt/Olig1/Olig2, or in gel filtration buffer B (20 mM Tris/HCl pH 7, 200 mM NaCl, 0.5 mM TCEP) for the remaining proteins. Fractions containing the respective proteins were pooled and concentrated. Purified and concentrated protein was immediately used for further experiments or snapfrozen in liquid nitrogen and stored at $-80\,^{\circ}\mathrm{C}$. The AtPHR1 $^{280-360}$ fragment used for crystallization was cloned into the pMH-

HS-Sumo protein expression vector, providing a N-terminal 8 × His-StrepII tandem affinity tag and a Sumo fusion protein. The construct was transformed into E. coli BL21 (DE3) RIL cells. For recombinant protein expression, cells were grown at 37 °C in TB medium to an OD_{600nm} of ~0.6. After reducing the temperature to 16 °C, protein expression was induced with 0.3 mM IPTG for 16 h. Cells were centrifuged for 20 min at 4000 × g and 4 °C, then the cell pellet was washed with PBS, snap-frozen in liquid nitrogen and stored at -80 °C. For the purification of a putative AtSPX1-InsP₈-AtPHR1²⁸⁰⁻³⁶⁰ complex, the AtPHR1 cell pellet was thawed and mixed with twice the amount of cells expressing a His-Strep-MBP-AtSPX1^{1–251} fusion protein, which provides a N-terminal, TEV-cleavable MBP. Lysis buffer (200 mM KP_i pH 7.8, 2 mM β-mercaptoethanol) supplemented with 0.1% (v/v) IGEPAL, 1 mM MgCl₂, 10 mM imidazole, 500 units TurboNuclease (BioVision), 2 tablets Protease Inhibitor Cocktail (Roche), and 0.1 mM InsP₈ was added and cells were disrupted using an EmulsiFlex-C3 (Avestin). Cell lysates were centrifuged at 7000 × g and 4 °C for 1 h. The cleared supernatant was sterile filtered and loaded onto a 5 mL HisTrap HP Ni²⁺ affinity column (GE Healthcare). After washing with several CVs of lysis buffer, the protein was eluted with 250 mM imidazole in lysis buffer. The purified His-Strep-Sumo-AtPHR1/His-Strep-MBP-AtSPX1 fusion proteins were cleaved by TEV and Sumo protease treatment overnight at 4 °C, while dialyzing in a buffer containing 200 mM KP, pH 7.8, 100 mM NaCl, 2 mM β -mercaptoethanol. Imidazole (10 mM) was added to the cleaved protein sample and a second Ni²⁺ affinity step was performed in order to remove the cleaved-off His-Strep-Sumo/MBP fusion tags as well as the 6 × His-tagged Sumo and TEV proteases. The flow-through was concentrated and loaded onto a HiLoad Superdex 75 pg HR16/60 column (GE Healthcare), pre-equilibrated in gel filtration buffer (20 mM Tris/HCl pH 7.8, 250 mM NaCl, 2.5 mM InsP₆, 0.5 mM TCEP). Fractions containing the co-eluting AtSPX1 $^{1-251}$ and AtPHR1 $^{280-360}$ proteins were pooled and concentrated. A second SEC step was performed using a HiLoad Superdex 200 pg HR26/60 column (GE Healthcare) and the same gel filtration buffer as above. It was later found that $AtSPX1^{1-251}$ expressed in *E. coli* is largely unfolded and unable to bind InsP8 and hence no complex with AtPHR1280-360 was observed in our crystals. Purified and concentrated protein was immediately used for further experiments or snap-frozen in liquid nitrogen and

For production of [U-15N,2H]-labeled AtPHR1280-360 variants for NMR spectroscopy, AtPHR1^{280–360} and AtPHR1^{280–360} KHR/A in pMH-HT were transformed and expressed in E. coli BL21 (DE3) RIL cells. Luria-Beltani liquid medium (2 mL) was inoculated with freshly transformed cells. After overnight growth at 37 °C, cells were transferred to M9H2O medium and grown overnight at 37 °C. From this culture, 150 μL were transferred into 1 mL M9_{H2O} medium and grown at 37 °C overnight. Cells were transferred into 100 mL M9_{D2O} medium and grown overnight at 37 °C. The culture was added to 0.9 L of $M9_{\rm D2O}$ medium and grown at 37 °C to an OD_{600nm} of ~0.4, then the temperature was shifted to 16 °C. After 1 h, protein expression was induced with 0.3 mM IPTG and cells were grown overnight, reaching a final OD_{600nm} of 1.2-1.4. The cell culture was harvested at $7800 \times g$ and the cell pellet was stored at -20 °C. The cell pellet was resuspended in 50 mM Tris pH 7.8, 500 mM NaCl, 0.1% IGEPAL, 1 mM MgCl₂, 2 mM βmercaptoethanol, 20 mM imidazole, and DNAseI (AppliChem). The resuspension was homogenized by magnetic stirring and after 10 min lysozyme was added. The cells were lysed by sonication and subsequently centrifuged at $42,500 \times g$ for 45 min. The collected supernatant was applied on 5 mL Histrap HP (GE Healthcare) equilibrated in a buffer A (50 mM Tris pH 7.8, 500 mM NaCl and 2 mM βmercaptoethanol) and washed with a wash buffer 1 (50 mM Tris pH 7.8, 200 mM KPi, and 2 mM β-mercaptoethanol) and with wash buffer 2 (50 mM Tris pH 7.8, 1 M NaCl, 2 mM β-mercaptoethanol), followed by a final wash step with buffer A. His-tagged protein was eluted with a buffer B (50 mM Tris pH 7.8, 500 mM NaCl, 1 M imidazole, and 1 mM β-mercaptoethanol). The fractions containing the eluted protein were incubated with TEV protease overnight at 4 °C. The sample was dialyzed against buffer A at $4\,^{\circ}$ C and purified by a second Ni²⁺ affinity step. Cleaved AtPHR1^{280–360} from the flow-through of the column was concentrated to 3 mL and purified further by SEC on a Superdex 75 16/600 (GE Healthcare) equilibrated in 25 mM HEPES pH 7.0, 200 mM NaCl and 0.5 mM TCEP at a flow rate of 1 mL/min. Fractions containing AtPHR1^{280–360} were pooled, concentrated to 0.5 mL, and the protein concentration was determined via Bradford assay (AppliChem). Samples were flash-frozen in liquid nitrogen and subsequently used for NMR spectroscopy experiments.

Isothermal titration calorimetry. All ITC experiments were performed at 25 °C using a MicroCal PEAQ-ITC system (Malvern Panalytical) equipped with a 200 μ l sample cell and a 40 μ l injection syringe. InsP₇ and InsP₈ were produced as described⁴⁷. All proteins were dialyzed against ITC buffer (20 mM HEPES pH 7.0,

200 mM NaCl) and PP-InsP ligands were diluted in ITC buffer prior to all measurements. A typical titration consisted of 15 injections, the protein concentrations in the syringe and in the cell are provided in the respective figure legend. Data were analyzed using the MicroCal PEAQ-ITC analysis software (v1.21).

Crystallization and crystallographic data collection. Two hexagonal crystal forms containing AtPHR1 $^{280-360}$ only developed in sitting drops consisting of 0.2 µL protein at a concentration of 12 mg/mL and 0.2 µL reservoir solution (0.1 M phosphate citrate pH 4.2, 0.2 M NaCl, 20% (w/v) PEG 8000). Crystals were cryoprotected by adding reservoir solution containing 10% (v/v) ethylene glycol directly to the drop and subsequently snap-frozen in liquid nitrogen. A third, tetragonal crystal form developed in 0.1 M Bis-Tri pH 6.5, 0.1 M NaCl, 1.5 M (NH₄)₂SO₄. Crystals were cryo-protected by serial transfer into reservoir solution supplemented with 15% (v/v) glycerol and snap-frozen in liquid nitrogen. Crystal forms 1, 2, and 3 diffracted to ~2.4, ~2.5, and ~1.9 Å resolution, respectively. Data were collected at beam-line PXIII of the Swiss Light Source, Villigen, Switzerland. Data processing and scaling were done in XDS⁴⁸.

Crystallographic structure solution and refinement. The AtPHR1 $^{280-360}$ structure was solved by molecular replacement using the previously described SPXCtGde1 (PDB-ID:5IJJ) core helices as search model in calculations with the program PHASER 49 . The structure was completed in iterative cycles of manual model building in COOT 50 and restrained refinement in phenix.refine 51 or Refmac5 52 . Residues 280–294 and 278–280 appear disordered in the final model. Quality of the structural model was assessed by using MolProbity 53 , refinement statistics are shown in Supplementary Table 5. Structure visualization was done with PyMOL (Molecular Graphics System, Version 1.8, Schrödinger, LLC) and ChimeraX 54 . The structure of AtSPX1 was modeled using the program SWISS-MODEL 55 and the SPXHsXPR1 domain structure of the human phosphate exporter as template (PDB-ID:5IJH, GMQE score \sim 0.49, QMEAN4 score \sim –2.27, 29.5% sequence identity 30). Conserved PP-InsP-binding residues in AtSPX1 were determined by aligning sequences with previously described SPX domains 30 using the program T-coffee 36 .

Right-angle light scattering. The oligomeric state of AtPHR1 variants was analyzed by SEC paired with a refractive index detector using an OMNISEC RESOLVE/REVEAL combined system (Malvern). Instrument calibration was performed with a bovine serum albumin (BSA) standard (Thermo Scientific Albumin Standard). Samples of 50 µL containing 2–10 mg/mL AtPHR1 (wild-type AtPHR1^{280–360}, AtPHR1^{280–360} Olig¹, AtPHR1^{280–360} Olig², AtPHR1^{280–360} KHR/A, wild-type AtPHR1^{222–358}, AtPHR1^{222–358} Olig¹, and AtPHR1^{222–358} Olig²) in OMNISEC buffer (20 mM HEPES pH 7.5, 150 mM NaCl) were separated on a Superdex 200 increase 10/300 GL column (GE Healthcare) at a column temperature of 25 °C and a flow rate of 0.7 ml min⁻¹. Data were analyzed using the OMNISEC software (v10.41).

DNA oligonucleotide annealing. DNA oligonucleotides were dissolved in annealing buffer (10 mM HEPES-NaOH pH 8.0, 50 mM NaCl, 0.1 mM EDTA). Equal volumes of the equimolar DNA oligonucleotides were mixed and incubated in a heat block for 5 min at 95 °C. Subsequently, DNA oligonucleotides were cooled down to room temperature for 90 min. Double-stranded DNA oligonucleotides were aliquoted and stored at -20 °C.

Electrophoretic mobility shift assay. Mini-PROTEAN TBE precast gels (5%; Bio-Rad) have been pre-electrophoresed in 0.5× TBE buffer for 60 min at 70 V. Reactions mixes have been prepared following the Odyssey* Infrared EMSA kit manual (LI-COR) without the use of optional components, including 50 nM of IRDye800 end-labeled oligos (refer to Supplementary Table 4a; Metabion), and a 1:5 dilution series of wild-type AtPHR1^{222–358} or AtPHR1^{222–358} Olig1 (1.2 μg to 76.8 pg). Reaction mixes have been incubated for 30 min at room temperature in the dark and 2 μL of 10× Orange Loading Dye (LI-COR) have been added to each sample prior to loading on a 5% TBE gel. Gels have been electrophoresed until orange dye migrated to the bottom of the gel (~1 h) at 70 V in the dark. Gels have been scanned with the 800 nm channel of an Odyssey imaging system (LI-COR).

Grating-coupled interferometry. All GCI experiments were performed at 4 °C using a Creoptix WAVE system (Creoptix sensors) with 4PCP WAVE chips (Creoptix sensors). Chips were conditioned with borate buffer (100 mM sodium borate pH 9.0, 1 M NaCl) and subsequently neutravidin was immobilized on the chip surface via standard amine-coupling: activation (1:1 mix of 400 mM N-(3-dimethylaminopropyl)-N'-ethylcarbodiimide hydrochloride and 100 mM N-hydroxysuccinimide), immobilization (30 μg ml⁻¹ of neutravidin in 10 mM sodium acetate, pH 5.0), passivation (5% BSA in 10 mM sodium acetate pH 5.0), and quenching (1 M ethanolamine, pH 8.0). Biotinylated oligos (Supplementary Table 4b; Metabion) were captured on the chip. Analytes were injected in a 1:2 dilution series starting from 4 μM (AtPHR1²²³⁻³⁵⁸), 20 μM (AtPHR1²²³⁻³⁵⁸ Olig1, AtPHR1²²³⁻³⁵⁸ Olig2), or 10 μM (OsPHR2, OsPHR2KHR/A) in GCI buffer (for OsPHR2: 20 mM HEPES pH 7.9, 200 mM NaCl; for AtPHR1: 20 mM HEPES pH

7.5, 300 mM NaCl). Blank injections every fourth cycle were used for double referencing and a dimethylsulfoxide (DMSO) calibration curve (0%, 0.5%, 1%, 1.5%, 2%) for bulk correction. Data were corrected and analyzed using the Creoptix WAVE control software (corrections applied: X and Y offset, DMSO calibration, double referencing, refractive index correction), and a one-to-one binding model was used to fit all experiments.

CD spectroscopy. Far-UV CD spectra and melting curves were acquired with a Chirascan V100 CD spectrometer holding the protein solutions in quartz cuvettes of 1 mm optical path, scanning from 260 to 190 nm at 1 nm/s with a slit width of 1 nm against an air background obtained with the same settings without any cuvette. Protein concentration was 0.08 mg/mL (8.5 μ M) in all experiments, the buffer was 10 mM KPi pH 7.5. A reference spectrum of the buffer showed only noise within ± 0.5 mdeg. All the final spectra reported were obtained with a single scan, as we observed very good signal-to-noise and no saturation of the detector voltage even at low wavelengths. Melting curves were acquired with 2 °C stepwise increments and 30 s intervals from 24 to 98 °C. Data were fitted in the range from 195 nm to 250 nm using the three-component (a-helix, β -strand, and random coil) fitting (http://lucianoabriata.altervista.org/jsinscience/cd/cd3.html) as previously described 57 .

NMR spectroscopy. TROSY spectra of $[U^{-15}N,^2H]$ -AtPHR1^{280–360} and $[U^{-15}N,^2H]$ -AtPHR1^{280–360} KHR/A were acquired on a 600 MHz spectrometer with a cryo-probe at 30 °C. The NMR buffer contained 25 mM HEPES pH 7.0, 200 mM NaCl and 0.5 mM TCEP. NMR data were analyzed by CCPNMR 2.4.2⁵⁸. The dissociation constant K_D was obtained by calculating chemical shift perturbations of each point by and plotted against $InsP_8$ concentration. The data points were fitted in MATLAB.

Plant material, seed sterilization, and plant growth conditions. All A. thaliana plants used in this study were of the Columbia (Col-0) ecotype. Seeds of the T-DNA insertion lines phr1-3 (SALK_067629) and phl1 (SAIL_731_B09) were obtained from the European Arabidopsis Stock Center. Homozygous phr1-3 and phl1 lines were identified by PCR using T-DNA left and right border primers paired with gene-specific sense and antisense primers (Supplementary Table 3d). The phr1 phl1 double mutant was kindly provided by Yves Poirier (University of Lausanne, Switzerland); vih1-2 vih2-4 double and phr1 phl1 vih1-2 vih2-4 quadruple mutants have been reported previously³⁵. Seeds were surface sterilized by incubation in 70% (v/v) CH₃-CH₂-OH for 10 min, followed by incubation in 0.5% (v/v) sodium hypochlorite for 10 min, and subsequently washed four times in sterile H₂O. Seeds were placed on full half-strength Murashige-Skoog plates⁵⁹ containing 1 (w/v) % sucrose and 0.8 (w/v) % agar (1/2MS plates), and stratified for 2-3 days at 4 °C in the dark prior to transfer into a growth cabinet. Plants were grown on vertical ^{1/2}MS plates at 22 °C under long day conditions (16 h light-8 h dark) for 8 to 11 days.

Western blotting. Proteins were transferred to nitrocellulose membrane (GE Healthcare, AmershamTM HighbondTM-ECL) via wet western blotting at 4 °C and 30 V overnight. Membranes were blocked in TBS-Tween (0.1%)-Milk (5%) for 1 h at room temperature. For mCherry detection, membranes were incubated overnight with an anti-mCherry antibody (ab167453, dilution 1:2000; Abcam) followed by 1 h incubation with an anti-rabbit-horseradish peroxidase (HRP) antibody (dilution 1:10,000, Calbiochem). For GFP detection, membranes were incubated overnight with an anti-GFP-HRP antibody (130-091-833, dilution 1: 1000, Miltenyi Biotec). For FLAG detection, membranes were incubated overnight with an anti-FLAG-HRP antibody (A8692, dilution 1:1000, Sigma). For SPX1 detection, membranes were incubated overnight with an anti-SPX1 antibody (dilution 1:1000, kind gift of Professor Mingguang Lei) followed by one hour incubation with an anti-rabbit-HRP antibody (dilution 1:10,000, Calbiochem). Antibodies were diluted in TBS-Tween (0.05%)-Milk (2.5%). Membranes were detected with SuperSignal™ West Femto Maximum Sensitivity Substrate (34095, Thermo Scientific™).

Determination of cellular Pi concentrations. To determine cellular Pi concentration at seedling stage, plants were transferred from $^{1/2}MS$ plates to -Pi $^{1/2}MS$ plates containing 1 (w/v)% sucrose and 0.8 (w/v)% agarose supplemented with either 0, 1, or 10 mM Pi (KH₂PO₄/K₂HPO₄ pH 5.7) at 7 DAG, and grown at 22 °C under long day conditions. At 14 DAG, seedlings were weighted and harvested into 1.5 mL tubes containing 500 μL nanopure H₂O. Samples were frozen at -80 °C overnight, thawed at 80 °C for 10 min, refrozen at -80 °C, incubated at 80 °C in a thermomixer shaking at 1400 r.p.m. for 1 h, and briefly centrifuged to sediment plant tissue. Pi content was measured by the colorimetric molybdate assay 60 . In brief, 600 μL ammonium molybdate solution (0.44 g of ammonium molybdate tetra hydrate in 97.3 mL nanopure water; add 2.66 mL concentrated (18 M) H₂SO₄ to a final volume of 100 mL), 100 μL 10% ascorbic acid, and 300 μL sample (250 μL nanopure water + 50 μL extracts) or 300 μL NaPi standard solution were mixed. Samples were incubated at 37 °C for 1 h and absorbance at 820 nm was measured.

RNA analyses. At 14 DAG, 50–150 mg seedlings were harvested in 2 ml Eppendorf tubes containing two metal beads each, shock-frozen in liquid nitrogen and ground in a tissue lyzer (MM400, Retsch). RNA extraction was performed using the ReliaPrep RNA Tissue Miniprep System (Promega) including in column DNaseI treatment to remove genomic DNA. First strand cDNA synthesis was performed from 1 to 2.5 μ g of total RNA using Superscript II RT (Invitrogen) with oligo(dT) primers. Quantitative reverse transcriptase PCR (qRT-PCR) was performed in 10 μ L reactions containing 1× SYBR-Green fluorescent stain (Applied Biosystems) and measured using a 7900HT Fast Real Time PCR-System (Applied Biosystems). qRT-PCR program: 2'–95 °C; 40× (30"–95 °C; 30"–60 °C; 20"–72 °C); melting curve 95 °C–60 °C–95 °C. A primer list can be found in Supplementary Table 3e. Expression levels of target genes were normalized against the housekeeping gene *Actin2*. For every genotype, three biological replicates were analyzed in technical triplicates.

Transient transformation of N. benthamiana. For each construct, 4 ml of A. *tumefaciens* strain pGV2260 suspension culture were grown overnight at 28 °C. Cells were collected by centrifugation at $700 \times g$ for 15 min and resuspended in transformation buffer (10 mM MgCl₂, 10 mM MES pH 5.6, 150 μ M acetosyringone). Cell density was measured and set to a final OD_{600nm} of 0.5 for SPX1 and PHR1, and to 0.1 for the silencing suppressor P19. Suspension cultures were incubated for 2 h in the dark at room temperature and subsequently mixed at a volume ratio of 1:1:1 (SPX1:PHR1:P19). *N. benthamiana* leaves were infiltrated using a 0.5 ml syringe and 3 leaf disks (d=1 cm) per sample were harvested after 3 days, snap-frozen in liquid nitrogen, and stored at -80 °C.

Co-immunoprecipitation. For co-immunoprecipitation experiments with proteins transiently expressed in N. benthamiana, samples were ground in liquid nitrogen with plastic mortars and proteins were extracted with 600 μL of homogenization buffer (50 mM Tris-HCl pH 7.5, 150 mM NaCl, 0.25% Triton X-100, 5% (v/v) glycerol, 1 mM phenylmethylsulfonyl fluoride, cOmpleteTM EDTA-free protease inhibitor cocktail (Roche). Samples were incubated at 4 °C for 10 min with gently rotation and subsequently centrifuged at 16,000 × g and 4 °C for 15 min. Supernatants were transferred to fresh tubes and further centrifuged at $16,000 \times g$ and 4 °C for 15 min. Supernatants were transferred to fresh tubes, while 50 μL of each supernatant were taken and mixed with 10 μL 6× SDS sample buffer (input), the remaining supernatants were mixed with 50 μL magnetic μMACS anti-GFP beads (Miltenyi Biotec) and incubated at 4 °C for 2 h with gently rotation. MACS columns (Miltenyi Biotec) were used with a µMACS Separator (Milteyi Biotec). MACS columns were washed with 200 µL of homogenization buffer and samples were loaded. Columns were washed either four times with 200 µL of homogenization buffer and once with wash buffer 2 (Miltenyi Biotec), or three times with 200 μL of homogenization buffer, three times with wash buffer 1 (Miltenyi Biotec) and once with wash buffer 2. Columns were incubated with 20 μL preheated elution buffer (Miltenyi Biotec) for 5 min at room temperature. Elution buffer (50 μL) was added and eluates were recovered. Inputs and eluates were boiled for 5 min at 95 °C prior and separated on 9% SDS-polyacrylamide electrophoresis gels. Coimmunoprecipitation experiments for proteins stably or natively expressed in A. thaliana were performed as previously described²⁹.

Statistics. All statistical analyses and data plots have been performed and generated with R version $3.5.2^{61}$ and the packages "Hmisc⁶²," "agricolae⁶³," "car⁶⁴," "multcompView⁶⁵," and "multcomp⁶⁶." qRT-PCR data were power transformed with the Box–Cox transformation and a significant one-way analysis of variance (ANOVA) followed by a Dunnett's post hoc test was performed for multiple comparisons of several genotypes vs. wild type (Col-0) shown in Supplementary Fig. 8a. Pi content was analyzed with a one-way ANOVA followed by a Dunnett's post hoc test for multiple comparisons of several genotypes vs. wild type (Col-0) shown in Fig. 4b and Supplementary Fig. 8b–d. (*p<0.5; **p<0.01; ***p<0.001)

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

Data availability

Data supporting the findings of this manuscript are available from the corresponding authors upon reasonable request. A reporting summary for this article is available as a Supplementary Information file. Coordinates and structure factors have been deposited in the Protein Data Bank (PDB) with accession codes 6TO5 (form1), 6TO9 (form2), and 6TOC (form 3). The associated X-ray diffraction images and data processing files have been deposited at https://doi.org/10.5281/zenodo.3570977 (form2), and https://doi.org/10.5281/zenodo.3570977 (form2), and https://doi.org/10.5281/zenodo.3571040 (form 3). Source data are provided with this paper.

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Competing interests

The authors declare no competing interests.

Additional information

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