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Data S2: Information on clustering analysis for the genes significantly regulated in the Mock roots of *phr2* vs WT grown at LP and *35S:PHR2* vs WT grown at HP samples in RNASeq.

Data S3: List of DEGs upregulated in wild type AM vs Mock LP and DEGs downregulated in *phr2* vs WT LP Mock roots.

Data S4: List of AM genes (AM genelists).

Data S5: Clustering analysis for the combined unique list of DEGs (14333 DEGs) significantly regulated in AM vs Mock samples of *phr2* under low phosphate (LP), *35S:PHR2* under high phosphate (HP) and WT under both phosphate conditions.

Data S6: Genes with reduced expression in *phr2* vs WT in AM and/or Mock samples at LP and required for or involved in AM symbiosis as determined by mutant analysis (red bold), RNAi (green) or overexpression (blue).

Data S7: Venn intersection of DEGs downregulated in *phr2* vs WT in AM and/or Mock samples at LP with DEGs upregulated in *smax1* or *d3/smax1* vs WT (Choi et al., 2020) and with AM genelists (Data S6).

Data S8: List of genes annotated to PHR2 binding sites (detected as MACS2 narrow peaks) and fasta sequence for PHR2 binding sites in ChIP-Seq of PHR2-FLAG.

Data S9: Occurrence analysis for P1BS/P1BS-like motifs in 3000 bp upstream sequence for genes with reduced expression in *phr2* and PHR2 targets in the two ChIP biological replicates.

Data S10: Direct PHR2 targets as identified by ChIP-Seq and RNASeq assays used in this work, therefore Set A + B genes in Fig. 3A and ChIP binding statistics for known targets of PHR2.

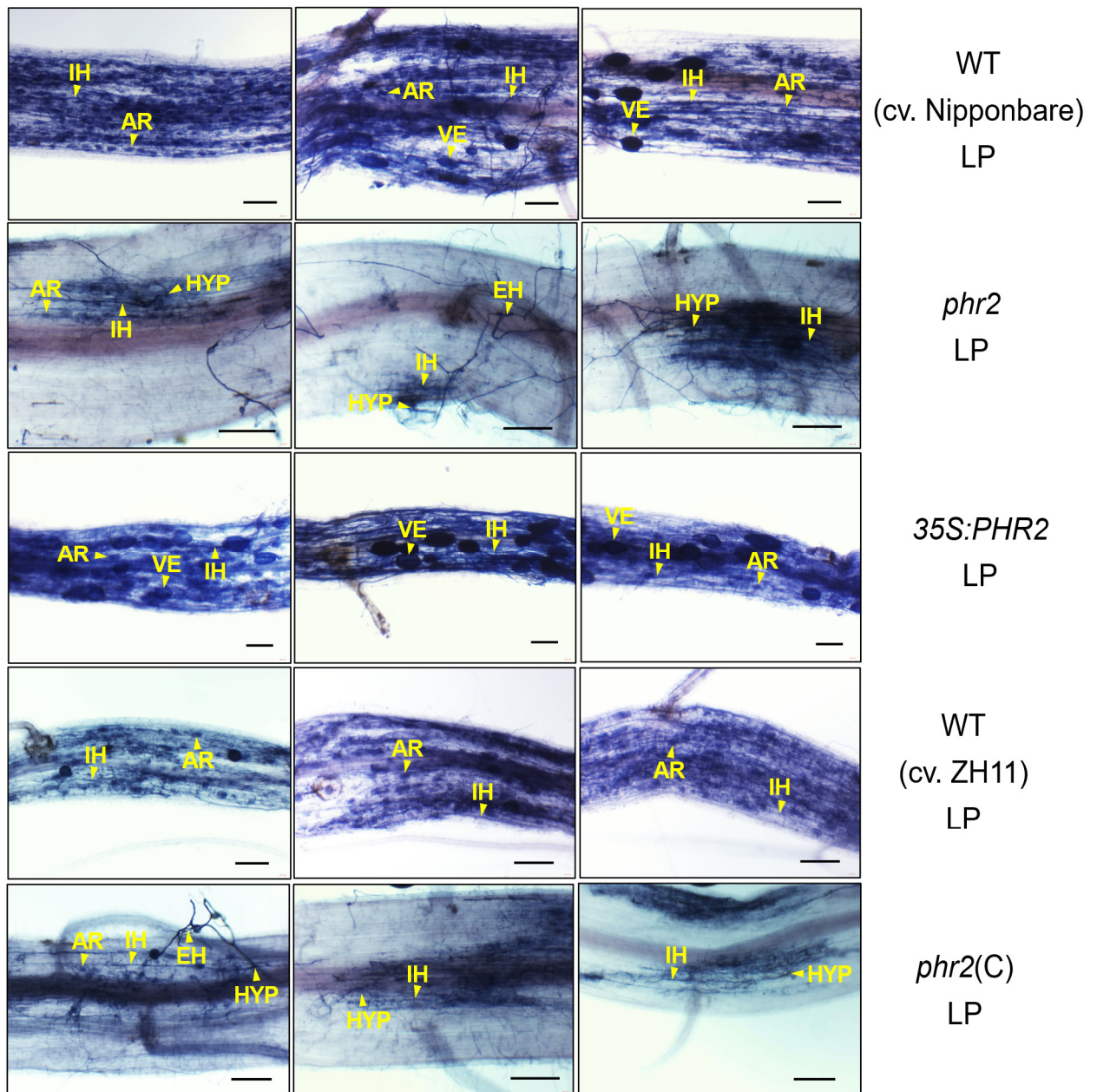


Fig. S1. Brightfield images of roots stained with acid-ink to visualize colonization of wild type (cv. Nipponbare), *phr2* and 35S:PHR2 and wild-type (cv. ZH11) and *phr2(C)* roots by *Rhizophagus irregularis* at 7 wpi when grown at LP (25 μ M P_i) in quartz sand. Scale bars, 200 μ m. Abbreviations: EH, extraradical hypha; HYP: hyphopodium; IH, intraradical hypha; AR, arbuscule, VE, vesicle.

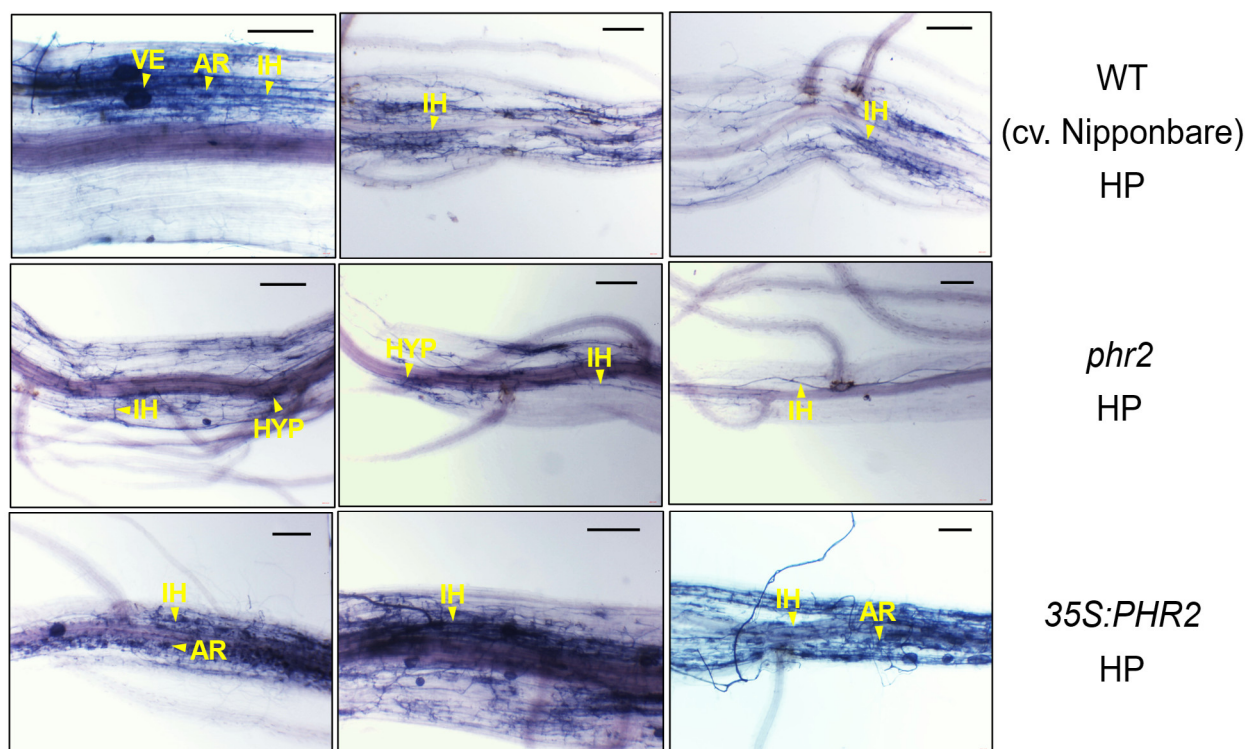


Fig. S2. Brightfield images of roots stained with acid-ink to visualize colonization of wild type (cv. Nipponbare), *phr2* and *35S:PHR2* roots by *Rhizophagus irregularis* at 7 wpi and grown at HP (500 $\mu\text{m P}_i$) in quartz sand. Scale bars, 200 μm . Abbreviations: EH, extraradical hypha; HYP: hyphopodium; IH, intraradical hypha; AR, arbuscule, VE, vesicle.

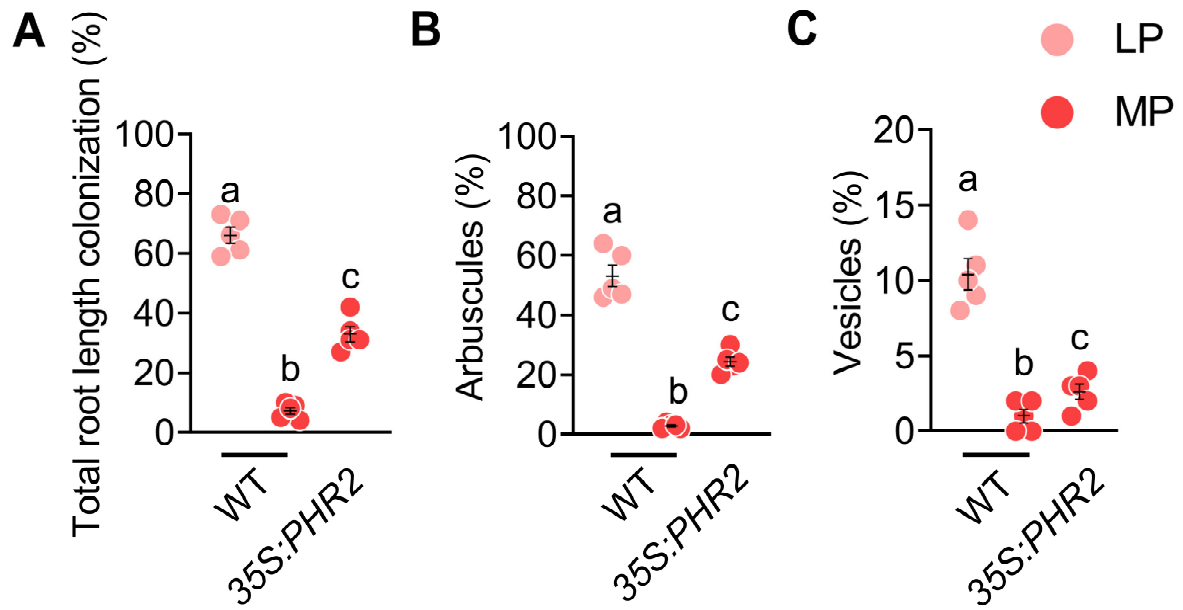


Fig. S3. Effect of low (LP, 25 μ M P_i) and medium (MP, 200 μ M P_i) phosphate conditions on total root colonization (**A**), arbuscules (**B**) and vesicles (**C**) at 7 wpi. Statistics: Individual data-points and mean \pm SE are shown. N=5; Brown-forsythe and Welch's One-Way ANOVA test with Games-Howell's multiple comparisons test. Different letters indicate statistical differences between samples.

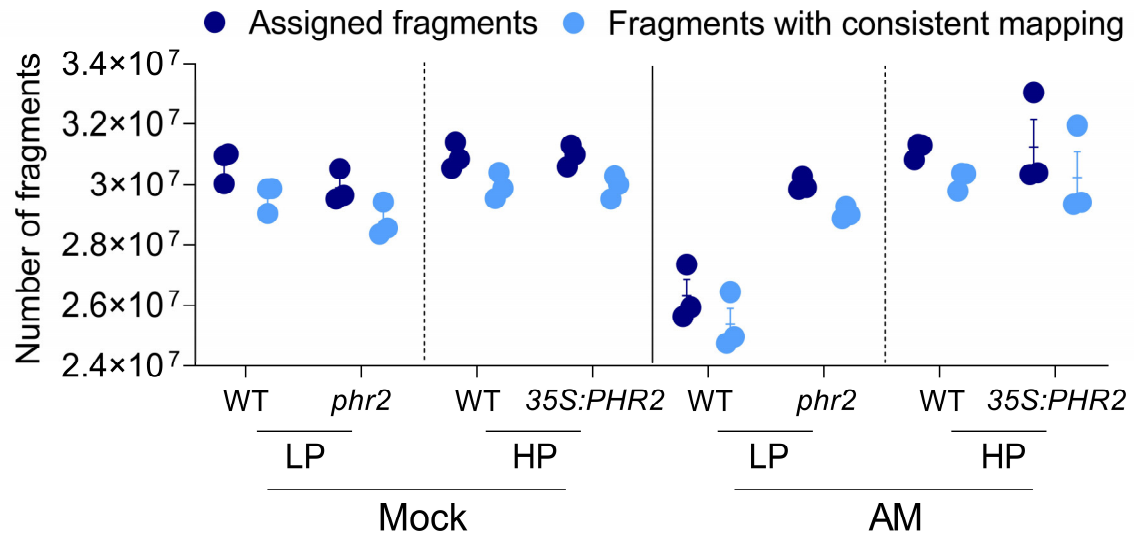


Fig. S4. Number of assigned fragments in RNA-Seq along with fragments consistently mapped to the transcriptome reference, $n=3$. Individual data-points and mean \pm SE are shown. Lower number of fragments in AM-inoculated wild-type roots grown at LP results from the fact that these sample also contain reads from *R. irregularis* ($\approx 10\%$ of total fragments).

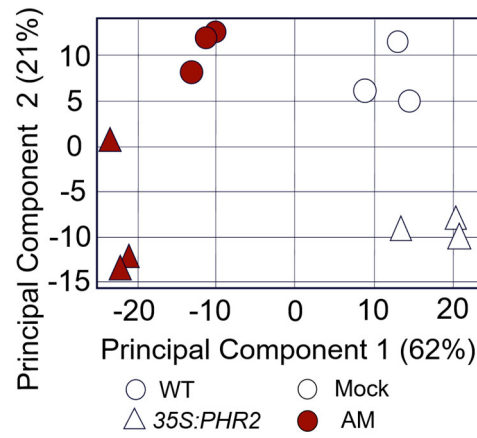


Fig. S5. PCA plot for the RNA-Seq based transcriptome of mock and AMF-inoculated wild-type and 35S:PHR2 roots grown at HP conditions.

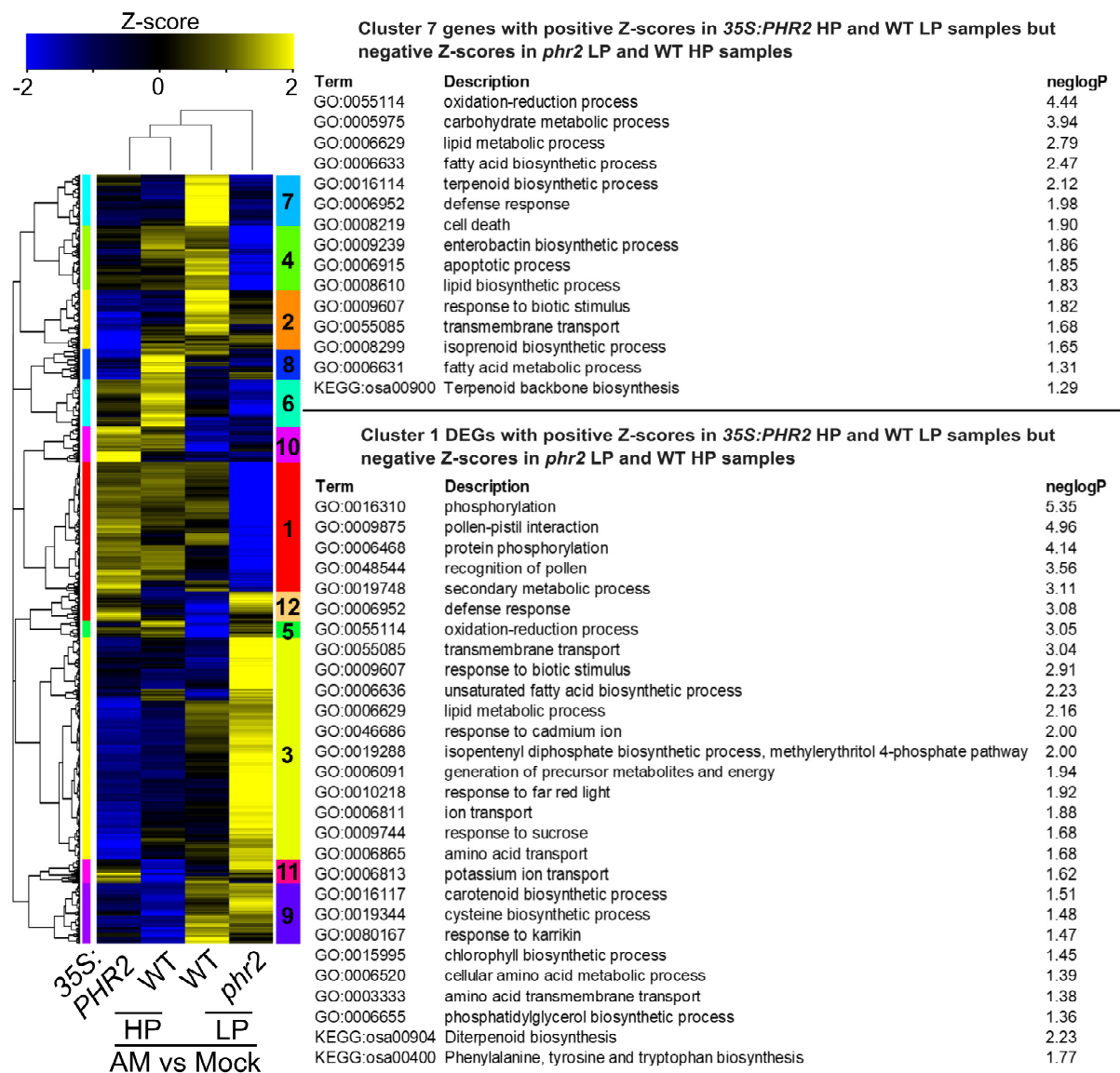
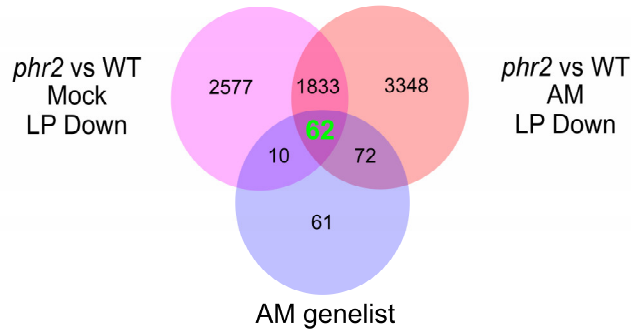


Fig. S6. Hierarchical clustering of combined DEGs (AM vs Mock samples, $\log_2(\text{Fold-change})$, $\log_2(\text{FC})$) from roots of *phr2* and 35S:PHR2 in LP and HP respectively and wild type at both P_i conditions. Z-scores represent scaled $\log_2(\text{FC})$. Colored bars on the left side of heatmap depict individual clusters (based on the dendrogram). Gene ontology (GO) enrichment analysis for selected DEGs in cluster-1 and -7, which showed positive Z-scores for 35S:PHR2 HP and WT LP samples and negative Z-scores for *phr2* LP and WT HP samples. GO term enrichment indicates functional categories important for AM symbiosis such as fatty acid, lipid and carotenoid metabolism, response to biotic stimulus and response to karrikin.

A



B

Locus ID (62 Genes)

Gene description

LOC_Os01g65000	AMT3;1, ammonium transporter protein
LOC_Os03g45290	Ankyrin repeat domain-containing protein (Medtr6g027840 VAPYRIN)
LOC_Os01g54270	D10, CCD8B
LOC_Os01g38580	D10-like, CCD8A
LOC_Os04g46470	D17, CCD7, carotenoid cleavage dioxygenase 7
LOC_Os11g01050	EXO70 exocyst complex subunit domain containing protein
LOC_Os03g29480	NSP1, GRAS transcription factor, nodulation-signaling pathway 1 protein
LOC_Os03g15680	NSP2, nodulation-signaling pathway 2 protein
LOC_Os05g41090	OsDMI3, OsCCaMK, calcium/calmodulin dependent protein kinases
LOC_Os06g02520	OsIPD3, OsCYCLOPS, interacting protein of DMI3
LOC_Os03g13080	OsLysM-RLK2, OsLYK5, MYR1/LYK2/RLK2/NFR5
LOC_Os06g44430	Protein kinase (Medtr4g129010 KIN2)
LOC_Os01g46860	PT11, inorganic phosphate transporter
LOC_Os04g10800	PT13, inorganic phosphate transporter
LOC_Os09g23640	STR1, ABC-2 type transporter domain containing protein
LOC_Os07g38070	SYMRRK, protein kinase, putative, expressed (Lj2g3v1467920 LjSymRK)
LOC_Os09g15240	Zaxinone Synthase (ZAS), carotenoid cleavage dioxygenase
LOC_Os08g05690	ABC transporter, ATP-binding protein, putative
LOC_Os05g50300	AMP1, AMP-binding enzyme, putative
LOC_Os04g39780	AMP-binding enzyme family protein
LOC_Os01g44950	AMP-binding enzyme, putative
LOC_Os07g38750	AP2 domain containing protein
LOC_Os12g39330	AP2 domain containing protein
LOC_Os02g48820	BCP, plastocyanin-like domain containing protein, putative, expressed
LOC_Os06g47130	C2domain containing protein
LOC_Os07g23450	C2H2 zinc finger protein
LOC_Os08g28240	carotenoid cleavage dioxygenase, putative
LOC_Os04g58680	CBF1/2, core histone H2A,H2B,H3,H4, putative
LOC_Os06g20120	CND41, chloroplast nucleoid DNA binding protein, putative, expressed
LOC_Os09g39530	Cupin-domain containing protein
LOC_Os07g33620	cytochrome P450 domain containing protein
LOC_Os01g50520	cytochrome P450 monooxygenase CYP711A12, putative, expressed
LOC_Os01g50590	cytochrome P450, putative, expressed
LOC_Os05g51240	D14L2a, Hydrolase, alpha/beta fold family domain containing protein
LOC_Os10g42400	DNA polymerase III, clamp loader complex, gamma/delta/delta subunit
LOC_Os01g46390	DUF538 domain containing protein, putative
LOC_Os05g49790	DUF538 domain containing protein, putative
LOC_Os08g42990	expressed protein
LOC_Os11g29630	expressed protein
LOC_Os02g18954	GDSL-like lipase,acylhydrolase, putative
LOC_Os02g44850	GDSL-like lipase,acylhydrolase, putative
LOC_Os04g47390	GDSL-like lipase,acylhydrolase, putative
LOC_Os07g19040	glycosyl hydrolase, putative
LOC_Os08g34258	inhibitor I family protein, putative
LOC_Os08g34249	inhibitor I family protein, putative, expressed
LOC_Os04g39180	KIN6, nodulation receptor kinase precursor, putative
LOC_Os02g20140	KinD, protein kinase domain containing protein
LOC_Os01g57400	lysM domain containing protein, putative
LOC_Os04g40570	MDR1, ABC transporter, ATP-binding protein, putative
LOC_Os05g47500	MDR-like ABC transporter
LOC_Os08g42590	mtN19, putative, expressed
LOC_Os01g52750	OsSub3 - Putative Subtilisin homologue
LOC_Os04g04750	peroxidase precursor, putative, expressed
LOC_Os03g57740	plastocyanin-like domain containing protein, putative
LOC_Os02g57700	protein kinase, putative, expressed
LOC_Os01g72710	putative RETICULATA-RELATED protein of unknown function
LOC_Os09g20970	receptor kinase, putative
LOC_Os03g38600	secretory carrier-associated membrane protein, putative, expressed
LOC_Os08g29510	Taurine catabolism dioxygenase TauD/TfdA domain containing protein
LOC_Os10g18510	UDP-glucuronosyl and UDP-glucosyl transferase domain containing protein
LOC_Os05g50680	WRKY83
LOC_Os01g53130	zinc finger, C3HC4 type domain containing protein, expressed

Fig. S7. Overlap of genes with decreased transcript levels in *phr2* in Mock or AM roots with AM genelist. (A) Venn diagram of DEGs downregulated in *phr2* vs WT Mock and AM roots at LP and AM genelist. (B) Genes common to all the three sets. Genes highlighted in red have been previously genetically shown to be required for AM development or function.

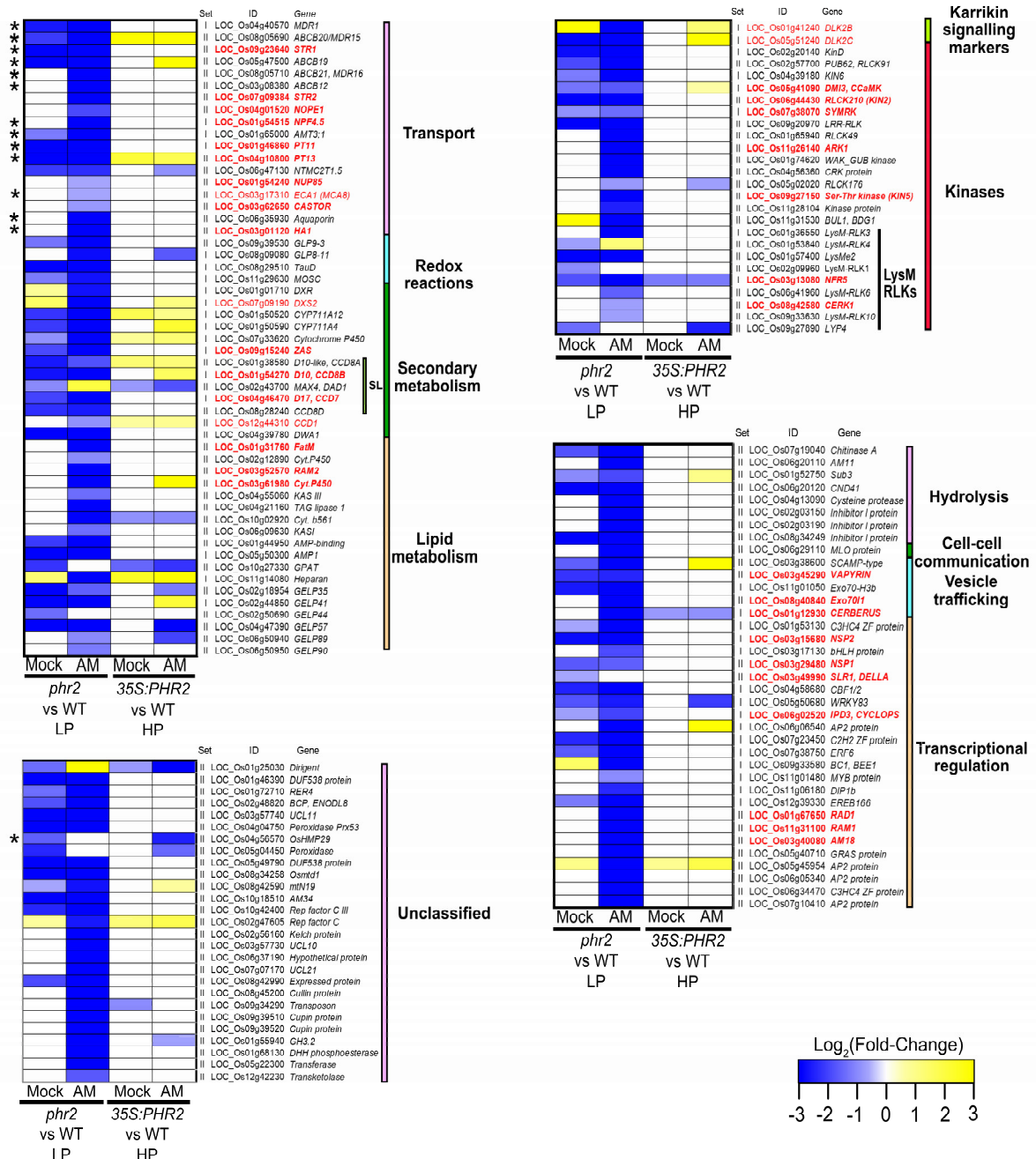


Fig. S8. Expression of genes required for or induced during AM depend on PHR2. Heatmaps for $\log_2(\text{Fold-change})$ of AM genes from set I and II (Figure 2G) for the comparisons *phr2* vs wild type at LP (25 μM) and 35S:PHR2 vs wild type at HP (500 μM). Colored bars on the right indicate functional categories to which the genes belong. Genes with genetically confirmed functions in AM symbiosis are indicated in red (bold for mutants, regular font for RNAi lines). 144 out of 205 genes (70%) in the AM genelist had reduced expression in *phr2* vs WT AM and/or Mock LP samples. Asterisks indicate orthologs of *Lotus japonicus* genes involved in transport and associated with P1BS elements in Supplementary table 2 of Lota et al., 2013. Out of 48 total genes in this table, only 31 could be retrieved from the *Lotus japonicus* genome assembly build 1.0 (<http://www.kazusa.or.jp/lotus/release1/>) and 14 out of these have reduced expression in rice *phr2* vs WT).

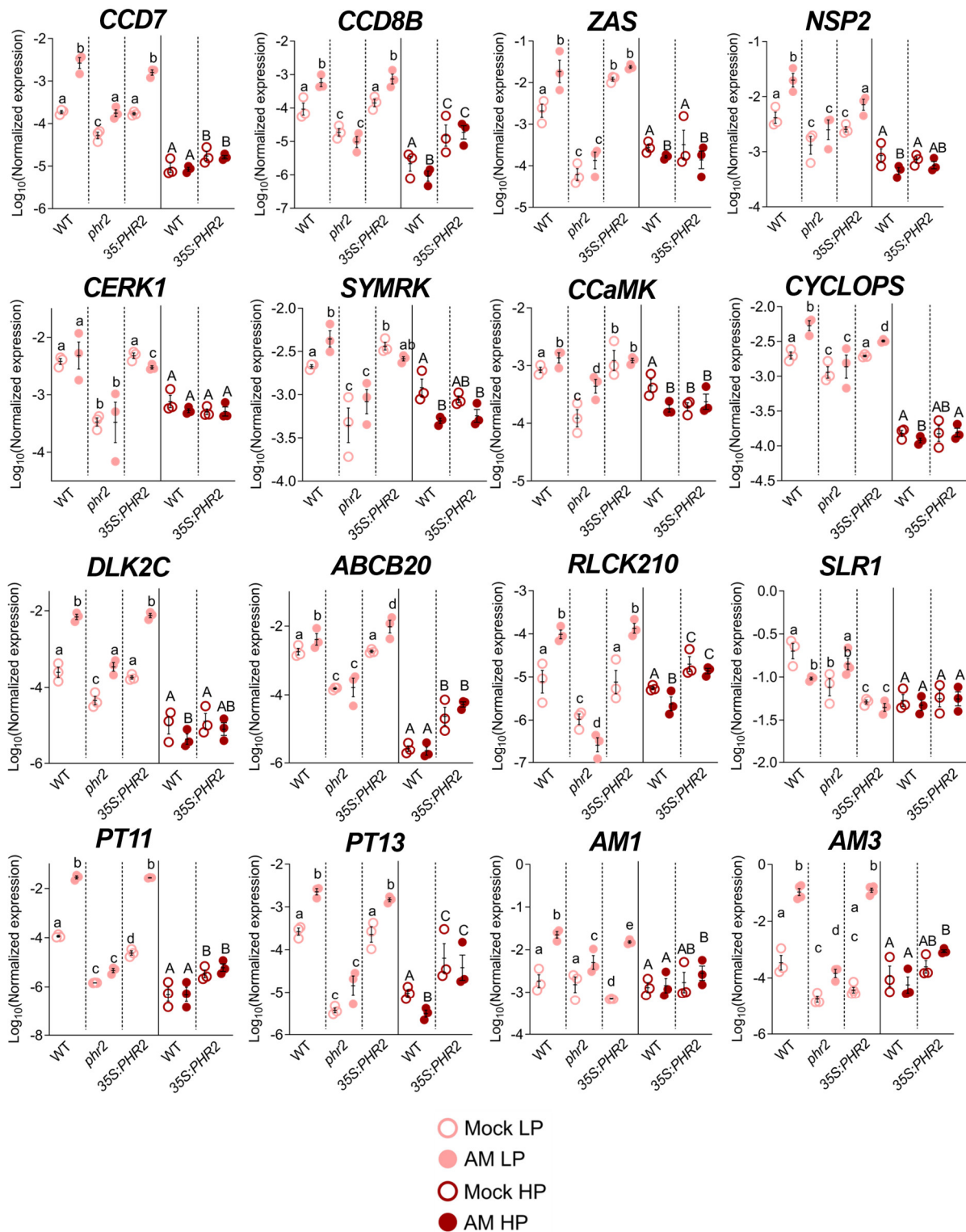


Fig. S9. RT-qPCR-based transcript accumulation of selected DEGs recapitulates the RNA-Seq results. Relative transcript accumulation in mock inoculated (Mock) and *R. irregularis* colonized (AM) roots of the indicated genotypes grown in quartz sand and fertilized with LP (25 μ M P_i) or HP (500 μ M P_i). Expression values of indicated genes were normalized to the geometric mean of the expression of two housekeeping genes, *UBIQUITIN* and *CYCLOPHILIN*. Letters indicate statistical differences between genotypes and treatments for

each phosphate level separately. Statistics: Individual data-points and mean \pm SE are shown. N=3. Brown-forsythe and Welch's One-Way ANOVA test with Games-Howell's multiple comparisons test was carried out for datapoints between samples at each phosphate level separately. Different letters indicate statistical differences between samples.

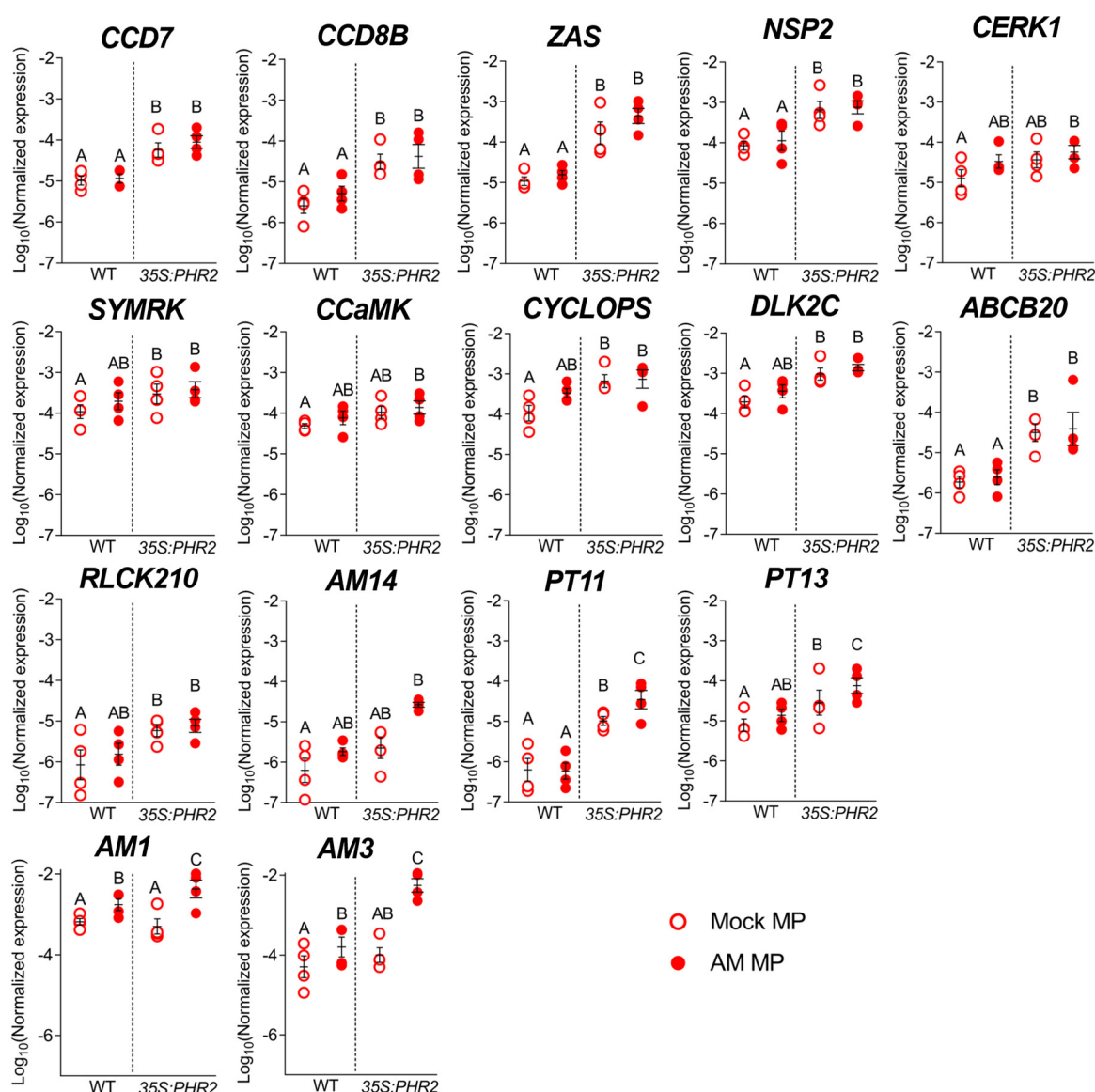


Fig. S10. RT-qPCR-based transcript accumulation of selected DEGs at medium phosphate. Relative transcript accumulation in mock inoculated (Mock) and *R. irregularis* colonized (AM) roots of the indicated genotypes grown in the same experiment as Fig. S3 in quartz sand and fertilized with MP is shown. Expression values of indicated genes were normalized to the geometric mean of the expression of two housekeeping genes, *UBIQUITIN* and *CYCLOPHILIN*. Letters indicate statistical differences between genotypes and treatments within each phosphate level. Statistics: Individual data-points and mean \pm SE are shown. N=3-4. Brown-forsythe and Welch's One-Way ANOVA test with Games-Howell's multiple comparisons test was carried out for datapoints between all samples. Different letters indicate statistical differences between samples.

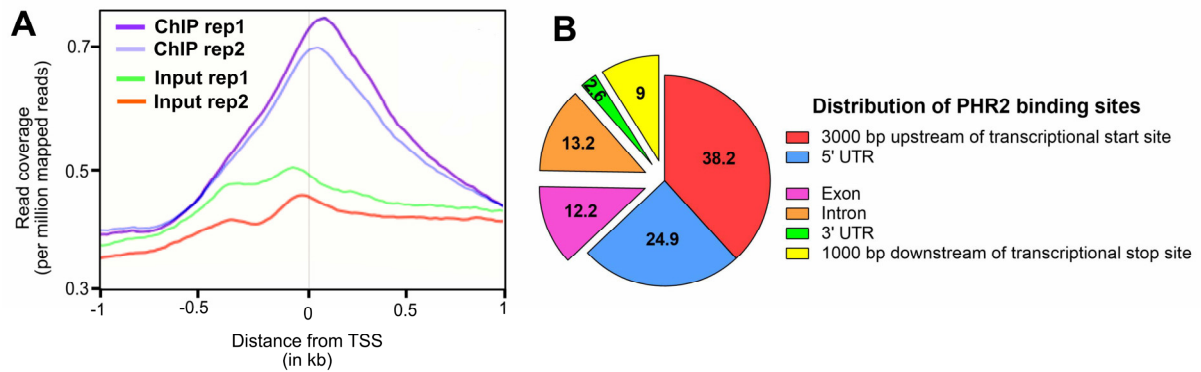
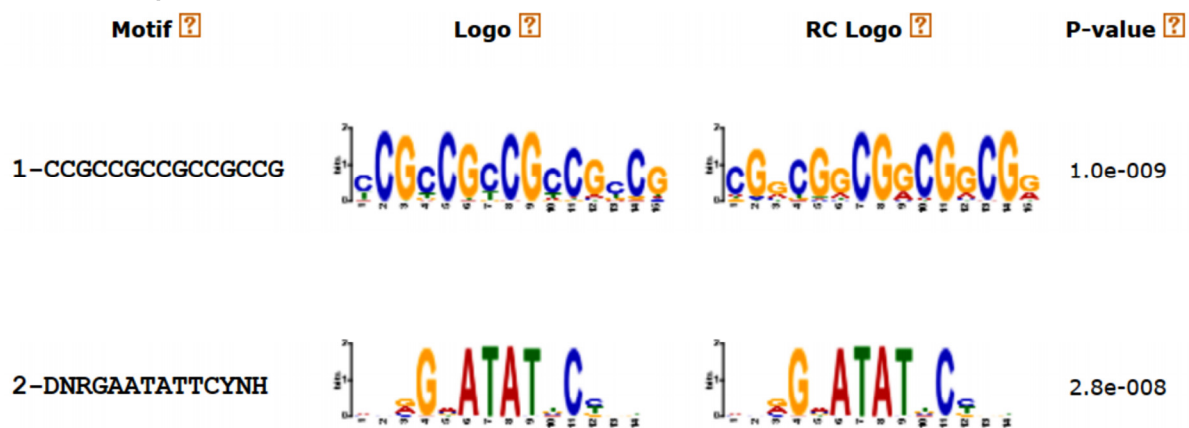
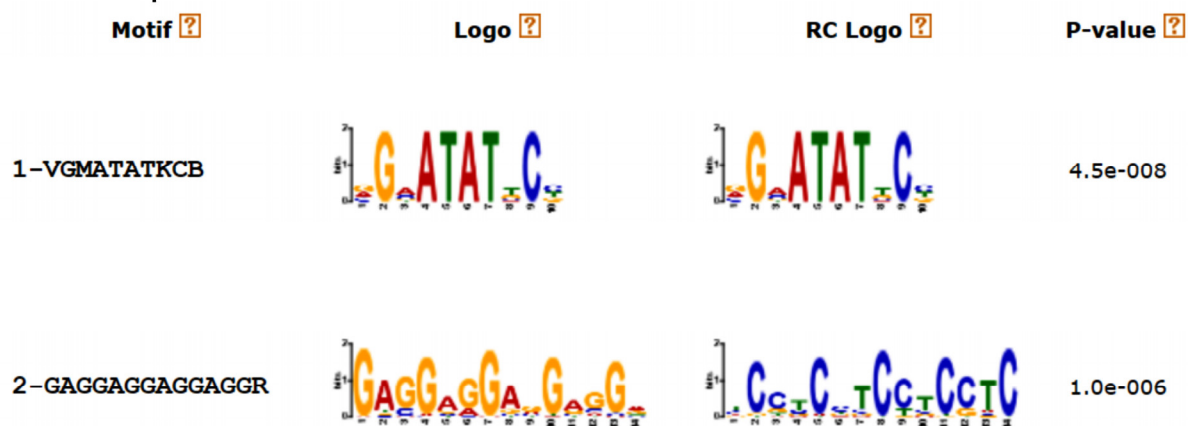


Fig. S11. ChIP-Seq binding peaks of PHR2-FLAG are enriched near the transcriptional start site. (A) Read coverage plot for the two biological replicates from ChIP-Seq with FLAG tagged PHR2 protein. TSS, transcriptional start site. (B) Distribution of PHR2-binding sites in the rice genome. ChIP-Seq read distribution in relation to transcriptional start site (TSS) suggested a slight skew in the distribution of PHR2 binding sites towards 1000 bp downstream of TSS. Correspondingly, PHR2 binding sites are enriched not only in 3000 bp region upstream of TSS (38.2%) but also in the regions downstream of TSS such as 5' UTR, exon and intron (24.9% + 12.2% + 13.2% = 50.3%).

ChIP replicate 1



ChIP replicate 2



5 **Fig. S12.** Motifs over-represented in DNA sequences with PHR2 binding sites. Analysis was carried out separately for the two biological ChIP-Seq replicates using STREME (<https://meme-suite.org/meme/tools/streme>).

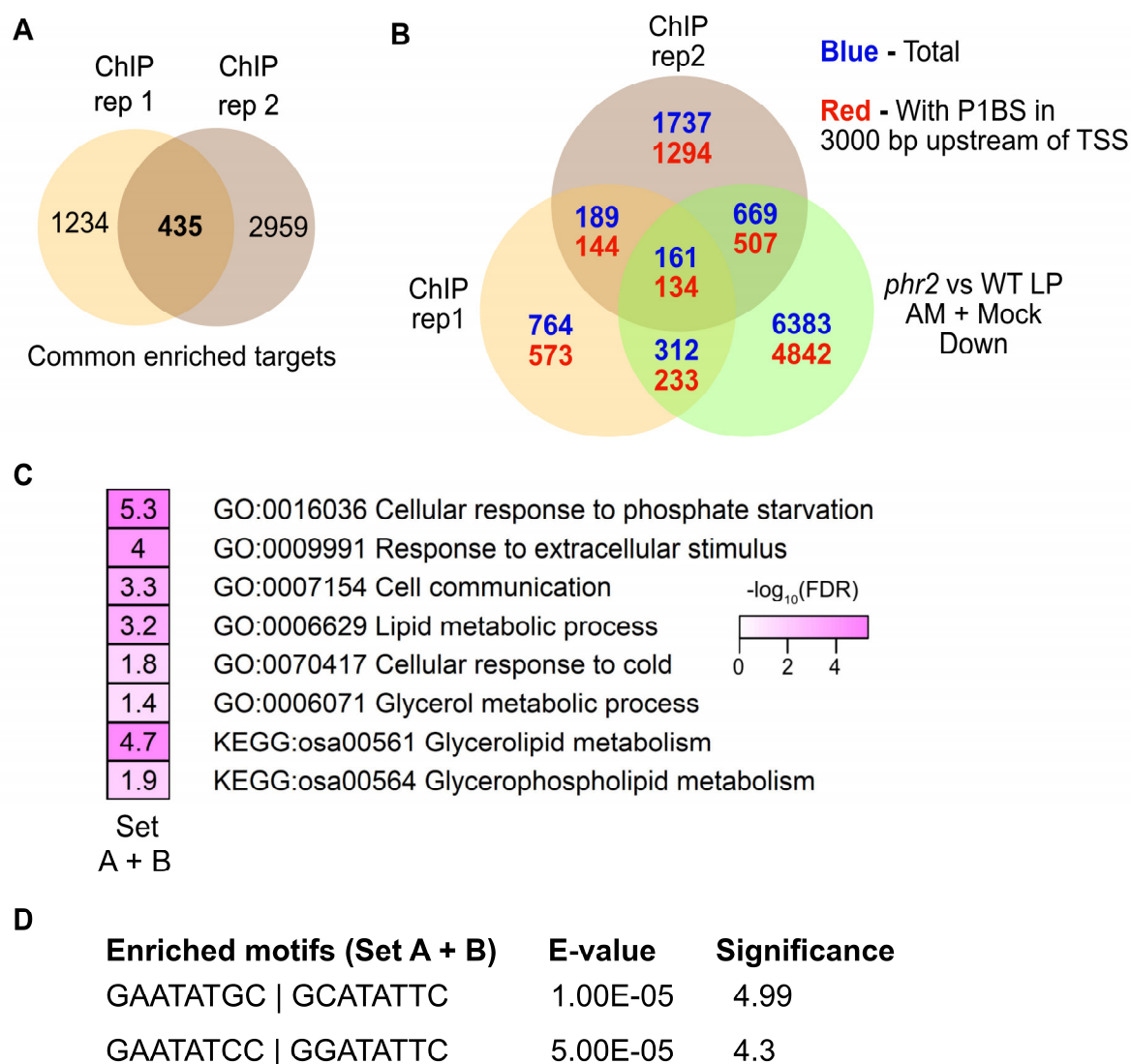


Fig. S13. Binding site analysis for rice PHR2. (A) Venn diagram showing overlap of PHR2 targets from two independent biological ChIP-Seq replicates. These 435 common PHR2 targets are genes annotated closest to PHR2-binding sites in both replicates. (B) Venn diagram showing overlap of PHR2 targets with DEGs with reduced expression in *phr2* vs WT AM + Mock samples at LP. Blue indicates the total number of genes and red with P1BS or P1BS-like motif in 3000 bp upstream region upstream of transcriptional start site (TSS). MSU IDs in the three individual gene sets were converted to RAPDB Locus IDs (to facilitate extraction of upstream sequence from RAPDB website, <https://rapdb.dna.affrc.go.jp/>). This resulted in a smaller number of genes than the original number of MSU ID DEGs. (C) GO-term enrichment in category “biological process” for Set A + Set B genes (167 genes out of 435 PHR2 targets which are repressed in AM or Mock root samples of *phr2* vs WT grown at LP as shown in Fig. 3A). Darker colors indicate stronger enrichment of GO-terms. GO-terms include categories involved in phosphate starvation signalling as well as AM. (D) Motifs enriched in 1000 bp sequence (upstream of TSS) for Set A + B genes in Fig. 3A.

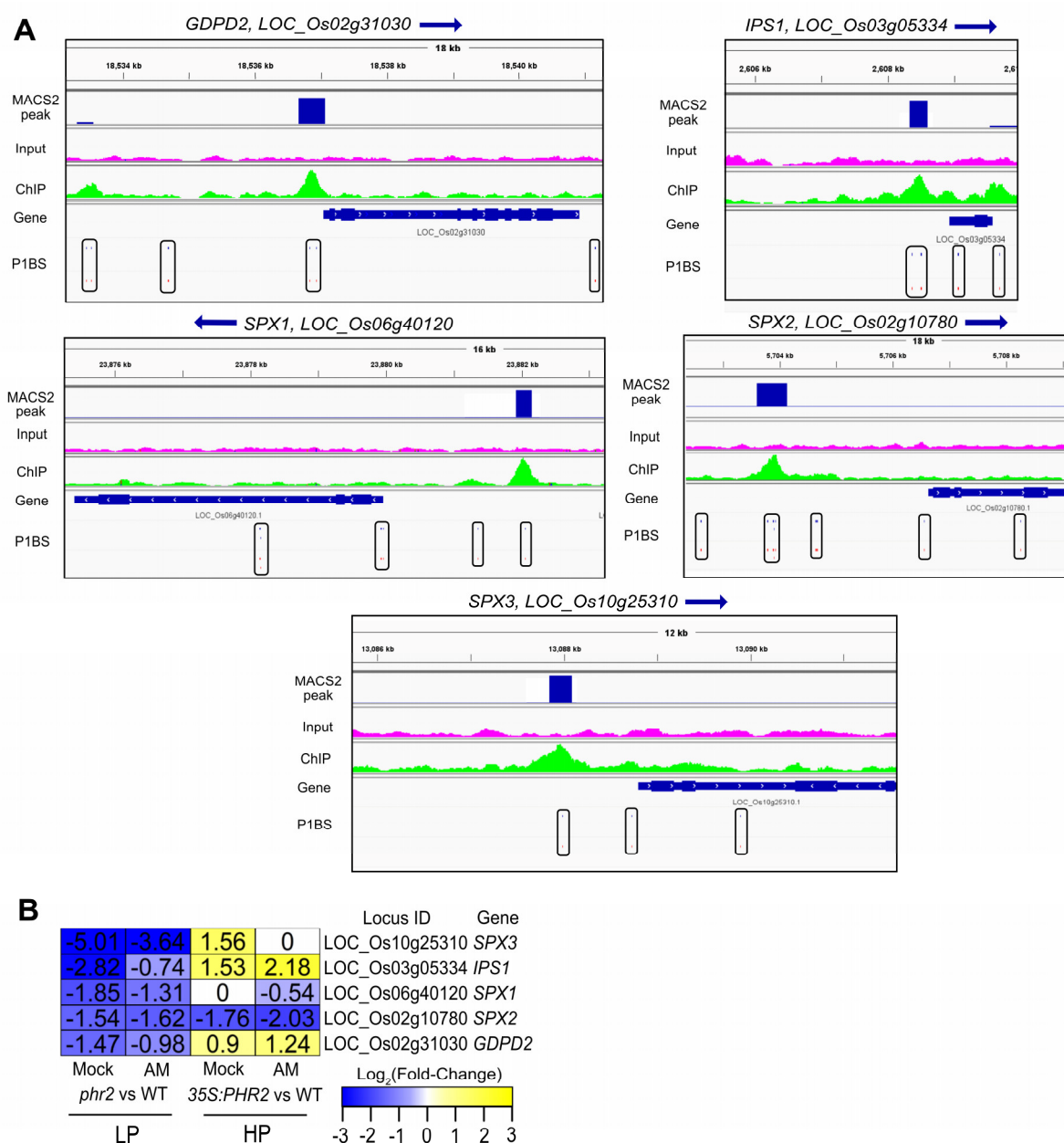


Fig. S14. IGV browser view of ChIP-Seq peaks adjacent to previously known PHR2 target genes. (A) ChIP-Seq peak profiles of genes which have been previously shown to be PHR2 targets. Gene orientation is indicated by the direction of the blue arrow close to the gene name. MACS2 peaks (blue bars) denote PHR2 binding sites corresponding to enrichment of PHR2-FLAG IP (green color) sequencing reads vs Input (pink color) sequencing reads. Positions of P1BS elements along the genomic coordinate are marked by enclosing the motifs in black rectangular boxes. **(B)** RNA-Seq based log₂(Fold-change) of these known PHR2 target genes in for *phr2* vs wild type and 35S:PHR2 vs wild type at LP (25 μM) and HP (500 μM), respectively. The phosphate level at HP (500 μM) maybe high enough to prevent the transcriptional induction of some of these phosphate starvation response genes such as *SPX1* and 2 in the 35S:PHR2 line.

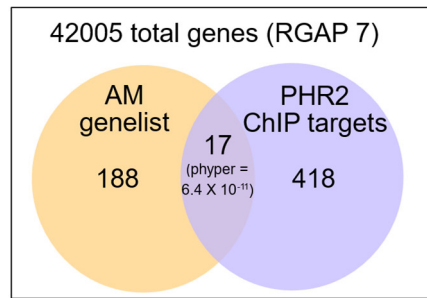


Fig. S15. Enrichment of PHR2 direct targets in AM genelists. Hypergeometric test was used to assess the statistical significance (phyper) of overlap of PHR2 ChIP targets with the AM genelists.

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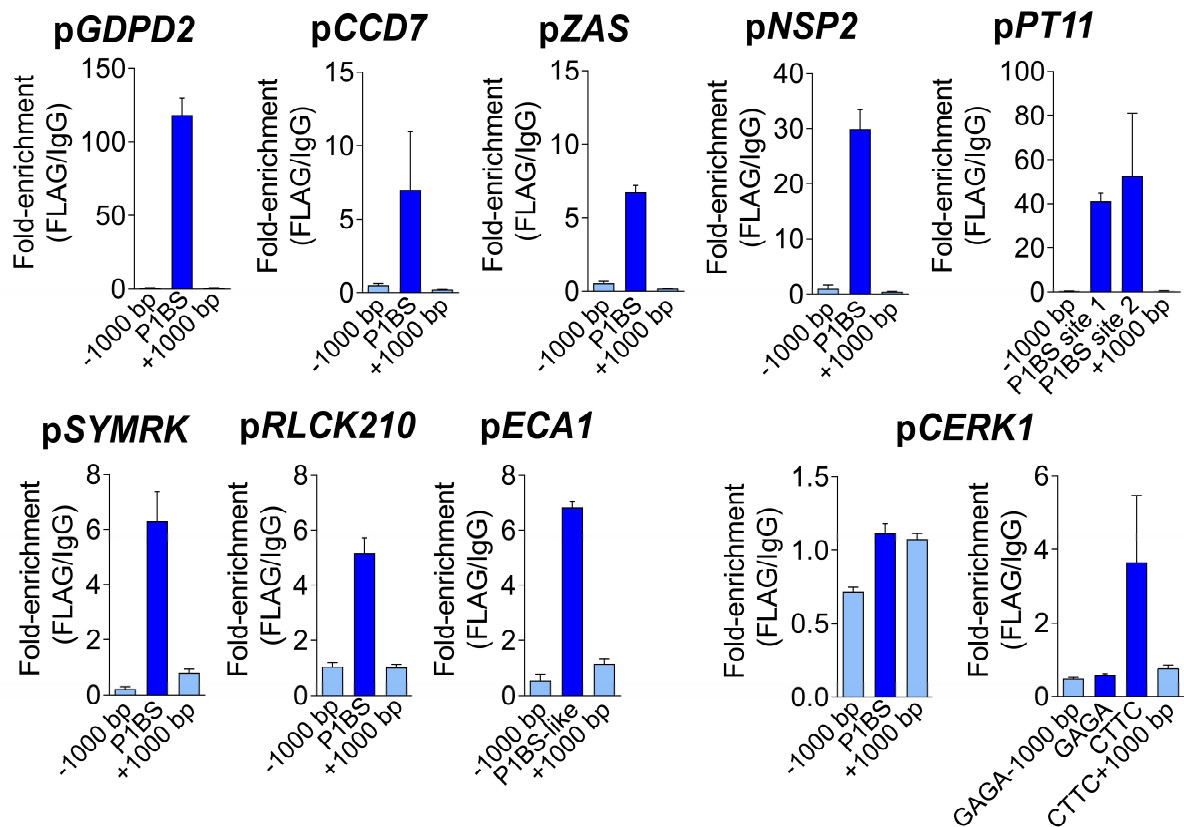


Fig. S17. Enrichment of PHR2 at P1BS promoter motifs detected by ChIP-qPCR. Primers were designed to amplify regions flanking motifs (P1BS, P1BS-like, GAGA, CTTC), and 1000 bp left (5') of these motifs (-1000 bp) and 1000 bp right (3') of these motifs (+1000 bp). Motifs: P1BS is GNATATNC; P1BS-like is AMATATYC; GAGA is GGAGAGGA; CTTC is TCCTCTTGTCTTC. Data: Individual data-points and mean \pm SE are shown. N=3.

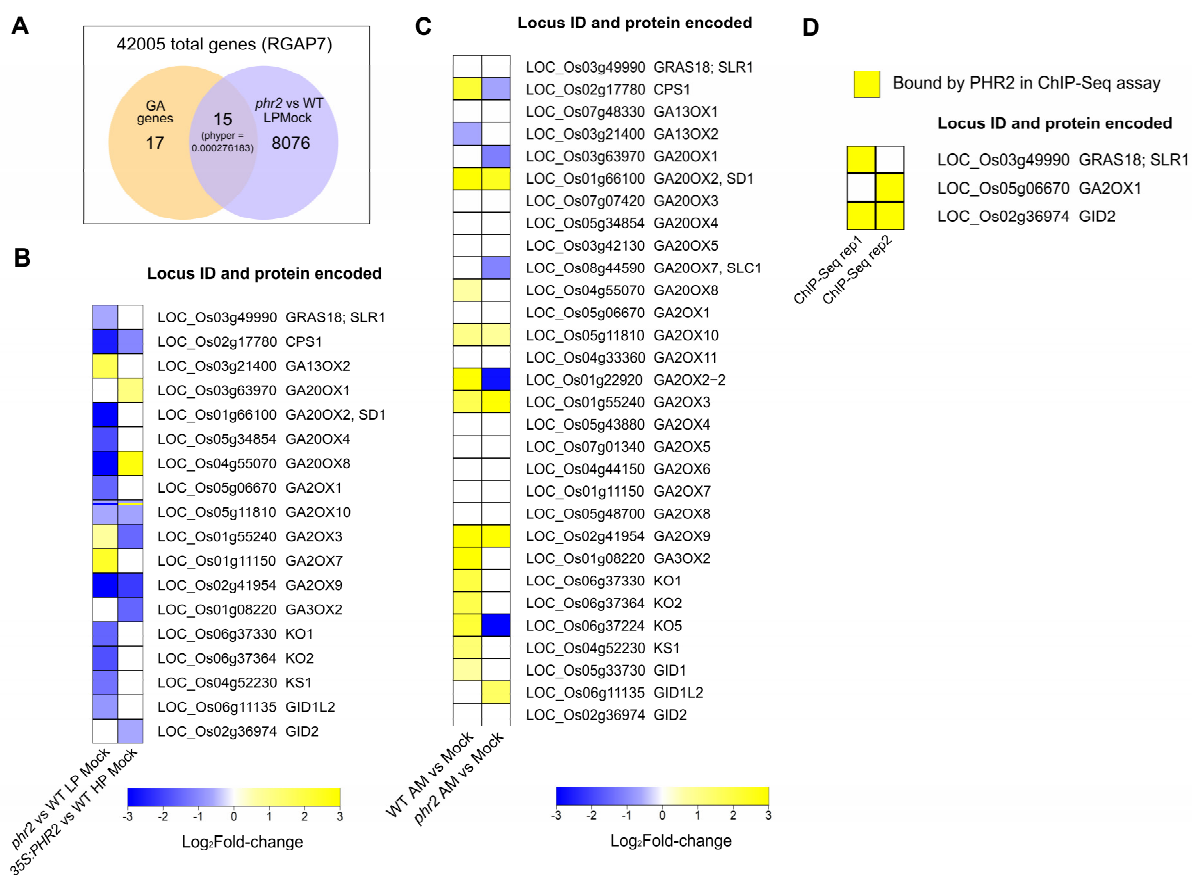


Fig. S18. Gibberellin-biosynthesis and -signaling related genes in RNASeq and ChIP-Seq. (A) Gibberellin (GA)-related genes are enriched in DEGs with reduced expression in non-inoculated *phr2* vs wild type. as shown by a hypergeometric test to assess the statistical significance (phyper). (B) Expression of GA-related genes in non-colonized *phr2* vs wild type roots at LP and 35S:PHR2 vs wild type roots at HP. (C) Comparison of gene expression for GA-related genes in *phr2* and wild type in AM vs Mock roots. (D) GA-related genes directly targeted by PHR2 as determined by ChIP-Seq.

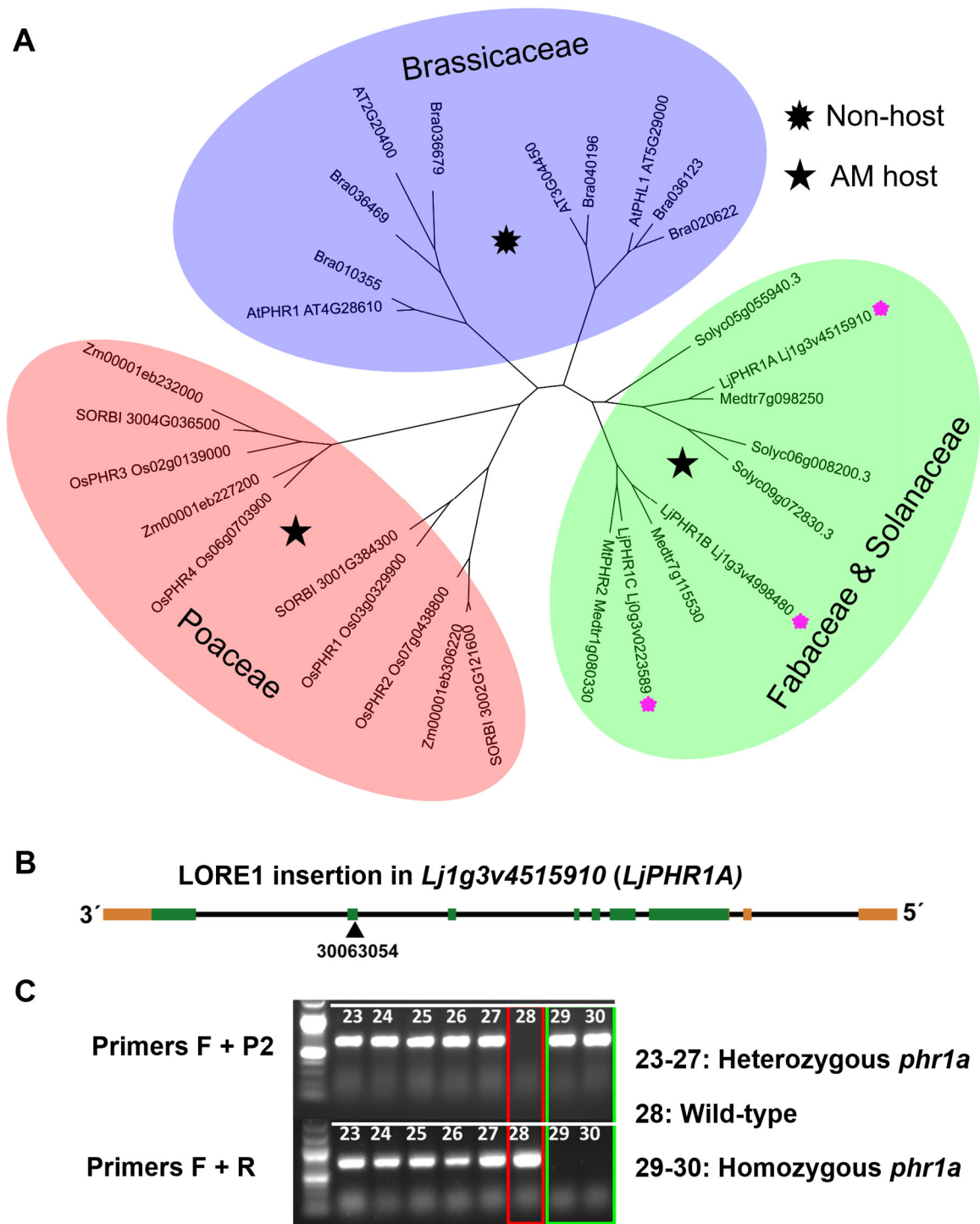


Fig. S19. *Lotus japonicus* PHR1A protein. (A) Phylogenetic tree of PHR proteins in representative *Brassicaceae*, *Poaceae*, *Fabaceae* and *Solanaceae*. The three *Lotus japonicus* PHR proteins are marked with pink stars. (B) Position of LORE1 insertion in *L. japonicus* PHR1A. The number indicates the Plant ID for LORE1 insertion. (C) Genotyping for LORE1 insertion in *phr1a*. The P2 primer sequence is located in the LORE1 insertion while F and R are PHR1A specific primers surrounding the insertion.

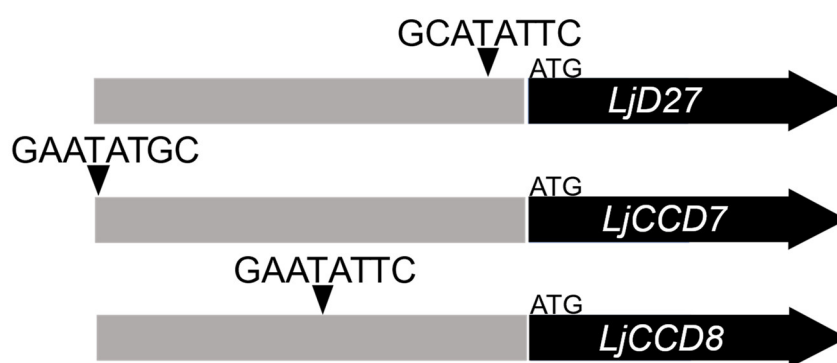


Fig. S20. Position of P1BS motifs in the promoters of strigolactone biosynthesis genes in *Lotus japonicus*. Promoter of length 1600 kb is represented in gray for each gene.

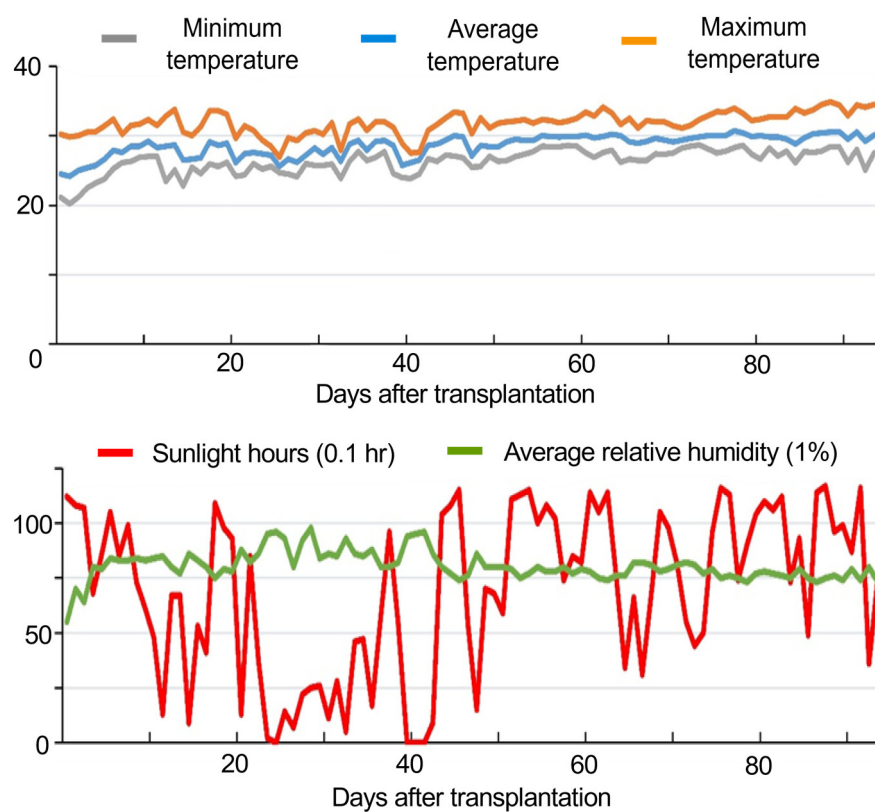


Fig. S21. Temperature, sunlight and relative humidity profiles during the greenhouse experiment in field soil.

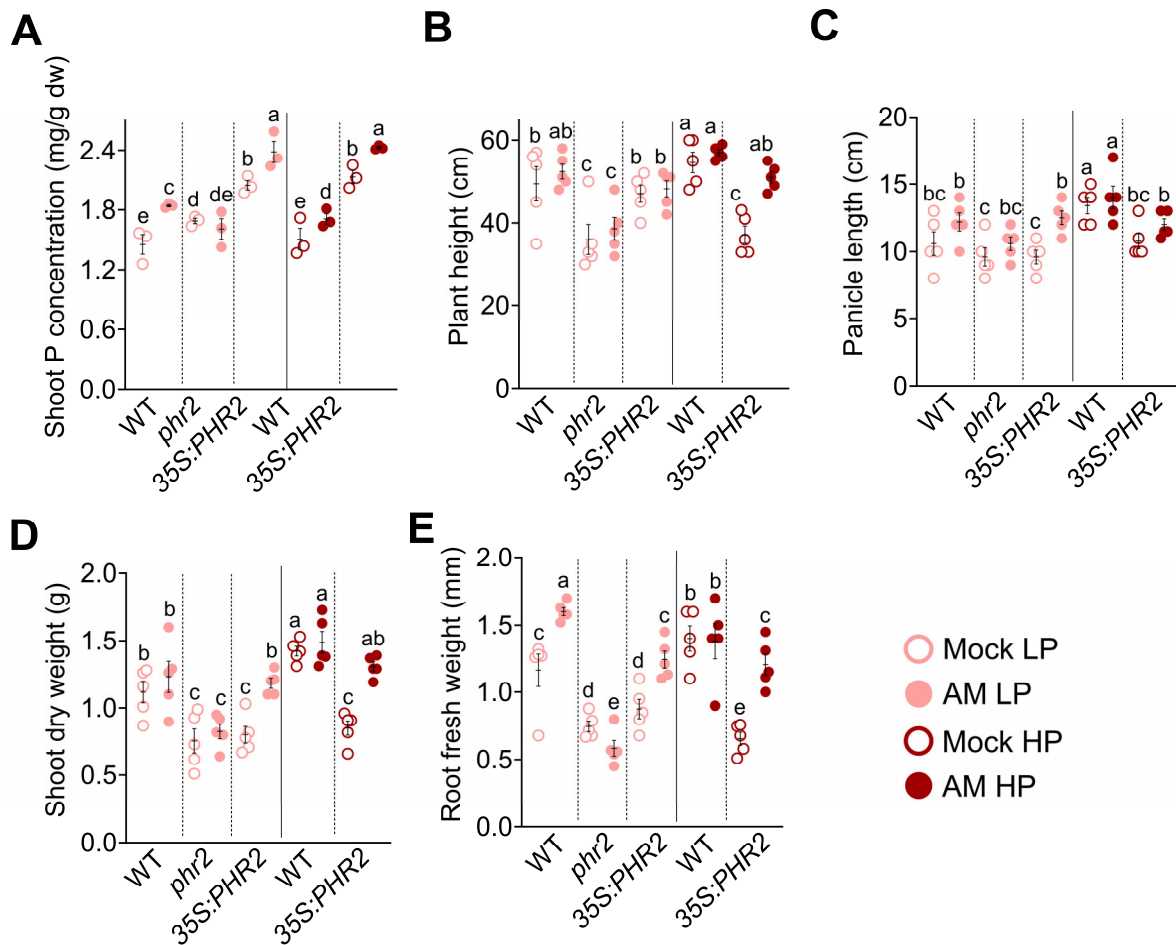


Fig. S22. PHR2 affects rice agronomic traits in a field soil. (A) Shoot phosphorus (P) concentration (mg/g dw), (B) Plant height (cm), (C) panicle length (cm), (D) shoot dry weight (g), (E) root fresh weight (g) in mock (Mock) and *R. irregularis* (AM) inoculated plants of wild type, *phr2* and 35S:PHR2 lines grown at LP (unfertilized) and of wild type and 35S:PHR2 lines grown at HP (fertilized with superphosphate fertilizer, P_2O_5). Traits were quantified for plants harvested at 110 days post transplanting into soil and inoculation. Statistics: Individual data-points and mean \pm SE are shown. N=3-5; Brown-forsythe and Welch's One-Way ANOVA test with Games-Howell's multiple comparison test was carried out for datapoints between all samples. Different letters indicate statistical differences between samples.

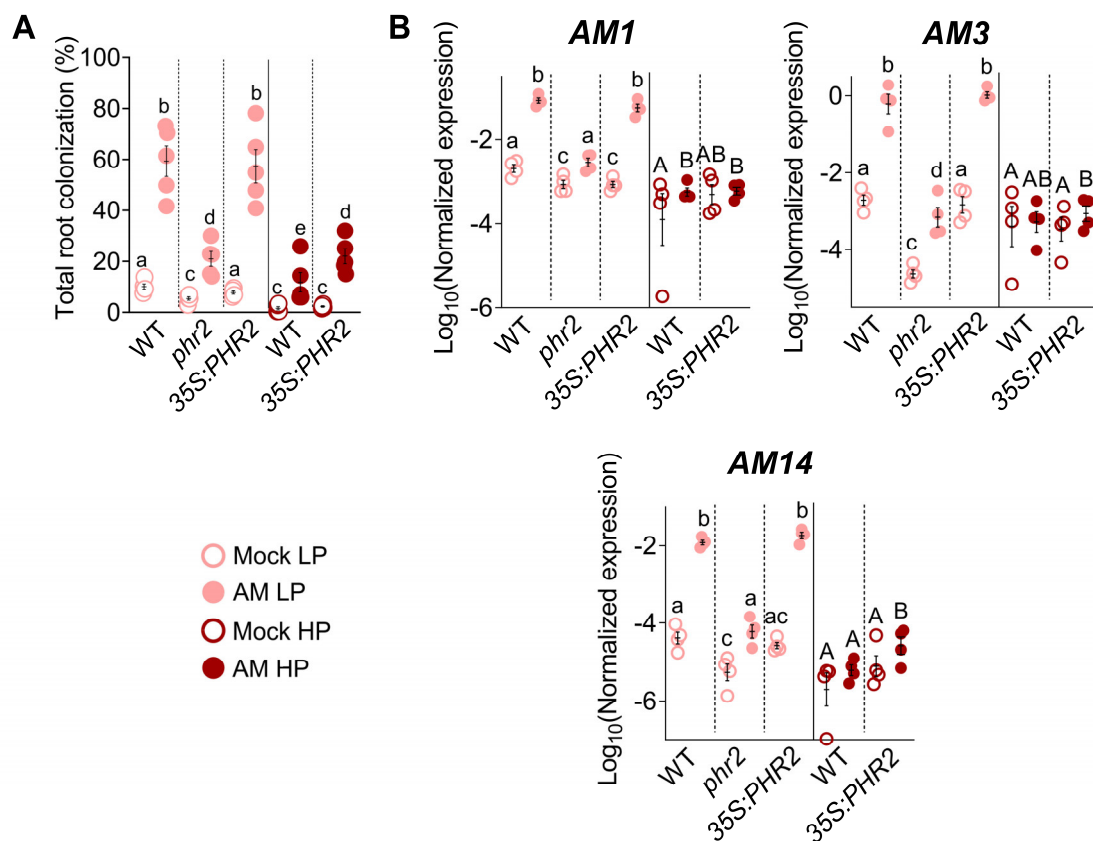


Fig. S23. Root colonization and RT-qPCR-based transcript accumulation of AM-marker genes in roots of plants grown in field soil. (A) Total root colonization (%) in *R. irregularis*-inoculated plants of wild type and *phr2* and 35S:PHR2 at LP (unfertilized) and in wild type and 35S:PHR2 lines at HP (fertilized with superphosphate fertilizer, P₂O₅). Roots were harvested at 110 days post transplantation into field soil and inoculation. Letters indicate statistical differences between genotypes, treatment and phosphate levels. Statistics: N=5; Kruskal-Wallis test with Dunn's posthoc comparison. **(B)** Relative transcript accumulation in mock inoculated (Mock) and *R. irregularis* colonized (AM) roots of the indicated genotypes grown in field soil and fertilized with LP or HP (as described in A). Expression values of indicated genes were normalized to the geometric mean of the expression of two housekeeping genes, *UBIQUITIN* and *CYCLOPHILIN*. Letters indicate statistical differences between genotypes and treatments within each phosphate level. Statistics: Individual data-points and mean ± SE are shown. (A) N=5; Brown-forsythe and Welch's One-Way ANOVA test with Games-Howell's multiple comparisons test between all samples. Different letters indicate statistical differences between samples. (B) N=3-4. Brown-forsythe and Welch's One-Way ANOVA test with Games-Howell's multiple comparisons test was carried out for datapoints between samples at each phosphate level separately. Different letters indicate statistical differences between samples.

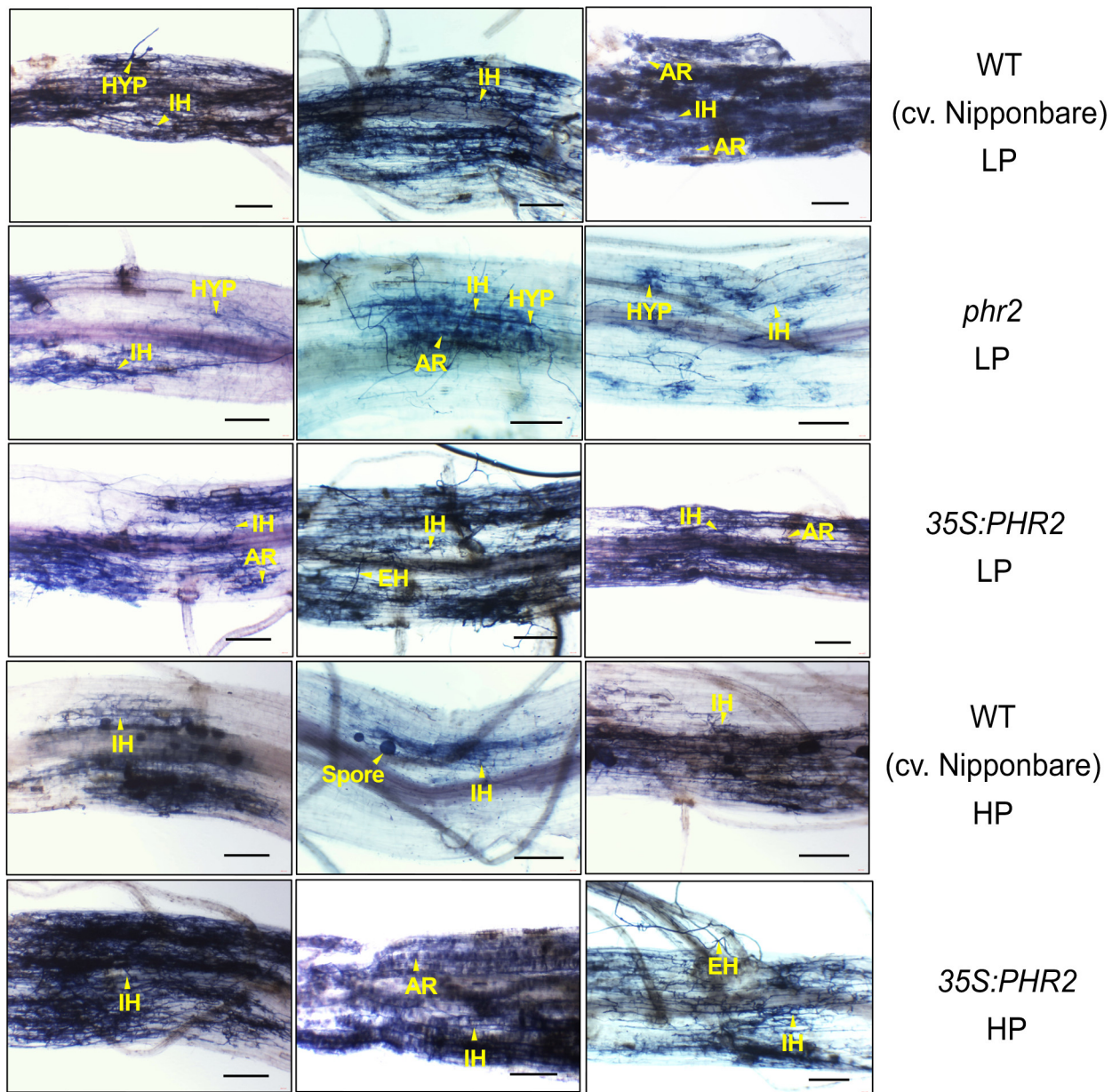


Fig. S24. Brightfield images of roots stained with acid-ink to visualize colonization of wild type (cv. Nipponbare), *phr2* and 35S:PHR2 roots by *R. irregularis* at 110 days post transplantation and grown at LP (unfertilized) or HP (fertilized with superphosphate fertilizer, P₂O₅) in field soil. Scale bars, 200 μm. Abbreviations: EH, extraradical hypha; HYP: hyphopodium; IH, intraradical hypha; AR, arbuscule, VE, vesicle.

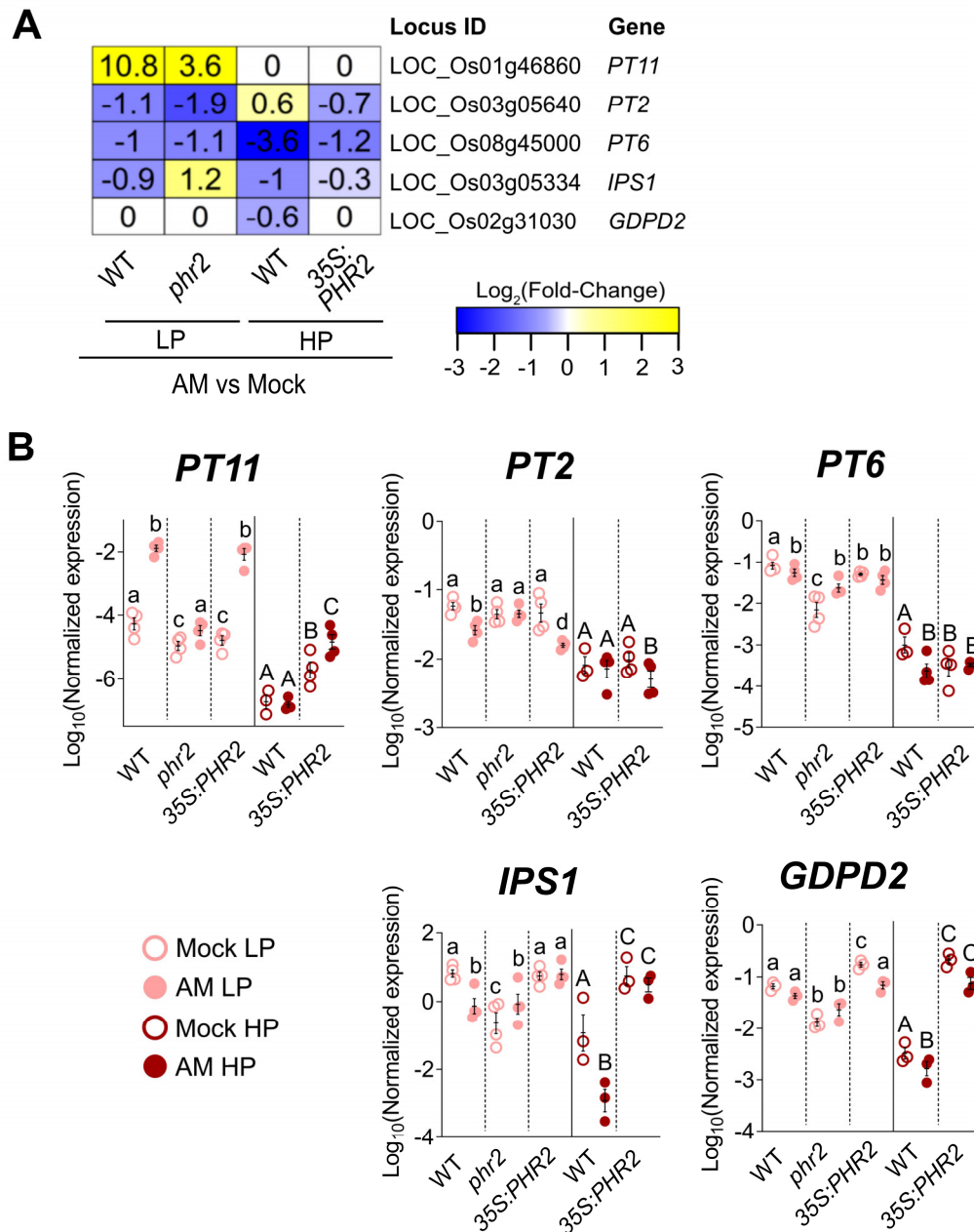


Fig. S25. RT-qPCR-based transcript accumulation of phosphate transporters and starvation marker genes in roots of plants grown in field soil. (A) RNA-seq-based fold-change of transcript accumulation (AM vs mock) of phosphate transporter genes involved in AM-mediated (*PT11*) or direct (*PT2*, *PT6*) P_i uptake, as well as the phosphate starvation marker genes (*IPS1*, *GDPD2*) at LP (25 μM P_i) or HP (500 μM P_i) in quartz sand. (B) RT-qPCR based relative transcript accumulation in mock inoculated (Mock) and *R. irregularis* colonized (AM) roots of the indicated genotypes grown in field soil and fertilized with LP or HP (as described in A) is shown. Roots were harvested at 110 days post transplanting and inoculation. Expression values of indicated genes were normalized to the geometric mean of the expression of two housekeeping genes, *UBIQUITIN* and *CYCLOPHILIN*. Letters indicate statistical differences between genotypes and treatments within each phosphate level. Statistics: Individual data-points and mean \pm SE are shown. N=3-4. Brown-forsythe and Welch's One-Way ANOVA test with Games-Howell's multiple comparisons test was carried out for datapoints between samples at each phosphate level separately. Different letters indicate statistical differences between samples.

Table S1. Primers used for RT-qPCR and genotyping.

Primer name	Forward Primer (5'→3')	Reverse Primer (5'→3')
CCD7-qRT	AATGCACTTGTGGCAAAACTAGAG	CATTGGAAAAGTGAGGTTCTTTGG
CCD8b-qRT	CAACTATGCCTTTTGGGTAAAG	AAAGTCTCGGCCAAATCCT
DLK2C-qRT	CGATGTTGCCATATAGTTGTGC	ACAAGGGAGCACACATGCAG
ZAS-qRT	TATGGAGGCCCTTGCAAACTTTGTC	CATTGTGTTTGCTAGTGATGATCTG
NSP2-qRT	TCAGCTGCTTCAACCACAGC	TGTTGGGACCCGTCCTCCTC
CYCLOPS-qRT	GGTTTGGCTTGGTACAGCATCT	GGGAGGCAGGTCATCACAA
CCaMK-qRT	AGGCCAACAGCAAGTGATCT	CGCAGATTATCCAGCTCCTC
ABCB20-qRT	GAAATGCTTGATAGGGACACAC	TGAAACTCAGTTCTTCCCATGA
SYMRK-qRT	CCTGGCATAAAAGGGCAATA	GTGCTTTCGATGGACCTCAT
CERK1-qRT	TGGAATCGTGATACATCCCCG	CAAGTTGTGTGGAATCTTCAG
SLR1-qRT	GACGTCAACGAACGCTCAATT	CGGAGTCCAGTCGTCGATCT
PT11-qRT	ATATCCAAGGCCTCGTTCCT	CCGATCAGCTGGATCATGT
PT13-qRT	CAGGACGAGTATGGCCTCTT	TCGAGGACGAACCAACAGA
RLCK210 (KIN2)-qRT	CCTCATGGAGATGGACAAGAG	GATACCATCTCCTCCTCCAAAC
AM1-qRT	ACCGTGTGGGAGATGGAGTT	CCTGCAGCTCTTCCTCATCT
AM3-qRT	CTGTTGTTACATCTACGAATAAGGAGAAG	CAACTCTGGCCGGCAAGT
AM14-qRT	CCAACACCGTTGCAAGTACAATAC	GCACTTTGAAATTGGACTGTAAGAAA
PT2-qRT	GACGAGACCGCCCAAGAAG	TTTTCAGTCACTCACGTCGAGAC
PT6-qRT	CCGCCCTGCAAACTGTA	CAACTGGCGGTTTCTTCGAT
OsUbiq-qRT	CATGGAGCTGCTGCTGTTCTAG	CAGACAACCATAGCTCCATTGG
CYCLOPHILIN2-qRT	AGCTCTCCTAGATCTGTGCTG	GCGATATCATAGAACGAGCGAC
IPS1-qRT	TTGGCAATTATTCGGTGGAT	ACCATTTACCATCCTCTTTATG
GDPD2-qRT	GCCCAGTCATCTTCCATGATA	CCAATTCATATCCGACCATCT
qPCR LjUbi	ATGCAGATCTTCGTCAAGACCTTG	ACCTCCCCTCAGACGAAG
qPCR LjPHR1A	CCGAATTGGAAGCATCCAAAGC	CTCGGAAGCTTGACTTTCGG
qPCR LjSYMRK	GAGGGTCAAAGGTGGATGA	GCGAACAATGGCGACCA
qPCR LjCCaMK	GGAGACAATGCAACTCTGTCTGA	CGGTGCTAGAGGGATCAATGA
qPCR LjCYCLOPS	GCTGGCAGATGAAAAAGAGC	GCGTGTTTGAGCACAAACATT
qPCR LjD27	GCCATCTCAATCGTTTATCAAG	GCTTCAGTGCTGGATCATC
qPCR LjCCD7	GTATGGAGTGTTTAAGATGCCC	TAAATGACTGCGTGGAAG
qPCR LjCCD8	GGACACGCTTAGGAAATTCG	TCTGTCACAATGGGATGTGC
LjPHR1A genotyping	TTGGTTATAAAGGACCGCAAG	TTCCTAACTAAGCTTGCCATAA
LORE1 P2		CCATGGCGGTTCCGTGAATCTTAGG

Table S2. Primers used for ChIP qPCR.

Primers with name appended with “motif”, “left” and “right” were used for amplifying sequences flanking P1BS (GNATATNC), or 1000 bp left and 1000 bp right of P1BS motif respectively (-1000 bp and +1000 bp in Fig. S17). In case of *ECA1*pro, primers with name appended with “motif” represents P1BS-like (AMATATYC). In case of *CERK1*pro, primers with names appended with “GAGAmotif” and “CTTC motif” were used for amplifying sequences amplifying GGAGAGGA and TCCTCTTGTTCCTTC elements respectively, while primer with names appended with “GAGAlleft” and “CTTCright” were used for amplifying sequences 1000 bp left of GAGA and 1000 bp right of CTTC respectively.

Primer name	Forward primer (5'→3')	Reverse primer (5'→3')
<i>CERK1</i> pro-ChIP-motif	TCGCAGTTTACAGTCGGAATC	TTGGATATACGGGCACACATTTA
<i>CERK1</i> pro-ChIP-left	GGCTGCTACATCACAAATTCAC	GGATGTGTTTCGGCTGGTATT
<i>CERK1</i> pro-ChIP-right	AAGAACACAGAGTGAGCTGTAA	GGGAAGAAAGGGAGAAGAAGAG
<i>CCD7</i> pro-ChIP-motif	GGGCCTATAACTGCATATTCTC C	GTGCCCACGTAATTTGAAAGAG
<i>CCD7</i> pro-ChIP-left	CCTTCACTTGGCGTTACAGA	CAGACACTAAACAGCACTACGA
<i>CCD7</i> pro-ChIP-right	CATGCAGGTTTCGTGGAGAC	TCACATTGCCCACCTTCTTC
<i>ZAS</i> pro-ChIP-motif	AGACACATGGATGCAGAGAAG	CGTGACGGATATTTCCAAGATGA
<i>ZAS</i> pro-ChIP-left	CATTGGTGTGCTGATGTTCTT	GCGGCCTACATTCTCAACTAT
<i>ZAS</i> pro-ChIP-right	GTCACATGGCATGCTACAAAC	AAATAACGGGTCCACCAATTTAAG
<i>SYMRK</i> pro-ChIP-motif	TGATTCCTCCCTTCCTCCTT	TTTCGTTCCGTGTCGTCATC
<i>SYMRK</i> pro-ChIP-left	ACAGTAACAAGGCTGAGTGTAT C	AAGCAGCAATCCATCTACTCC
<i>SYMRK</i> pro-ChIP-right	TCTGCAGGACAACAACCTTCA	GCAAATGGTAAAGAACAGCATCTA
<i>ECA1</i> pro-ChIP-motif	CGCACAGCAGAGCACAA	CCATTCTCCACTCTCCGTTTC
<i>ECA1</i> pro-ChIP-left	GAGATCGCATGGAACCGAAA	CTTCTCCTCTCTTCCGCATTG
<i>ECA1</i> pro-ChIP-right	TCGTGTAGGACAGACCTTGAT	AATCCTAGTCACCAGTCCTACC
<i>PT11</i> pro-ChIP-motif1	CGAGAGGAGAATGACGAAATCA	GCTCTTCTCCCATATCCATCAG
<i>PT11</i> pro-ChIP-motif2	TGATTGGCGATTCTACCATAC	GCGTAGCGGTAAATCGATGA
<i>PT11</i> pro-ChIP-left	ATGCGCCACACGTAGTC	CCATGATCGTCTCTAGCATCTTC
<i>PT11</i> pro-ChIP-right	AGCGGTGAAGCAGCAAA	CTCTAGATAAGTGGGACCGTACA
<i>GDPD2</i> pro-ChIP-motif	TGCCTTTGGACCGGAATATC	AAGGAAGGAAGCGGGAATG
<i>GDPD2</i> pro-ChIP-left	TGTGCTCTCGTGATGAATCTG	CTGTTCCACGACGGGTAA
<i>GDPD2</i> pro-ChIP-right	TGAGCTGCTGTTCCGATTC	TGTCGATCGATTGATTTCCC
<i>NSP2</i> pro-ChIP-motif	GCATTACGGGAAGCAACAAAG	GCTGAAGTCTGAAGACTGA
<i>NSP2</i> pro-ChIP-left	CCATAGGTGAGACTTGAGAG	CCCTTGGTACTTTAGAAATAGATGT
<i>NSP2</i> pro-ChIP-right	GCCTTGGCAACAAAGCTAAG	AGAAATGTGCCGAGAGAGATG
<i>CERK1</i> pro-ChIP-GAGAmotif	CAGTCCTGAACAGAGGACATAA G	GACTCCTCTCCAGACACTTCTA
<i>CERK1</i> pro-ChIP-CTTCmotif	GGAGTCAAGGTTAGTGGCTAAG	CTGTGTTCTTTGCTTACGGATG
<i>CERK1</i> pro-ChIP-GAGAlleft	GTCTACCTCCACATGTCTCAAC	CCTAGGAAGAGGCCTAGATACA
<i>CERK1</i> pro-ChIP-CTTCright	TACCCGGCCAACAACATC	TGAGGAACAGCCCGTAGT
<i>RLCK210</i> pro-ChIP-motif	TTTGTGAATGAATTAGGTGCGT	GTGTTGTATGAGTATTGTCAATGT
<i>RLCK210</i> pro-ChIP-left	TACTATCACACCACGCGTCTA	GAGATTAGTAGATGGTCCCTGTAATT T
<i>RLCK210</i> pro-ChIP-right	ACATGACCACCAGGCAAG	AATAAGACGGACGGTCAAACA

Table S3. Primers used for cloning.

Purpose	Name	Sequence
c <i>PHR2</i> cloning for p35S:c <i>PHR2</i>	MP503	ATGAAGACTTTACGGGTCTCACACCATGGAGAGAATAAGCACCAATCAGC
	MP508	ATGAAGACTTCAGAGGTCTCACCTTTCTGTCACCTGATTCTGAAACAAAAATTTAAGG
p <i>GDPD2</i> cloning for p <i>GDPD2</i> :GUS	MP595	TTTGGTCTCAGCGGTGTTTCATATATCTGATGTGACACGTC
	MP600	TTTGGTCTCACAGATATATTCGGAGGATGTCCTAGCTG
p <i>GDPD2m</i> cloning for p <i>GDPD2m</i> :GUS	MP595	TTTGGTCTCAGCGGTGTTTCATATATCTGATGTGACACGTC
	MP596	TTGGTCTCACGCGGACGGTCCAAAGGCACGC
	MP597	TTGGTCTCACGCGAATGGAGGATAAACCATCCGATCCGC
	MP598	TTGGTCTCAGGTCCGCGGAGGGGTGGGGATGCGTTC
	MP599	TTGGTCTCAGACCCATTCCCGCTTCCTTC
	MP600	TTTGGTCTCACAGATATATTCGGAGGATGTCCTAGCTG
p <i>PT11</i> cloning for p <i>PT11</i> :GUS	MP609	TTGGTCTCTGCGGGGAGCAATAGACGAGGGATGCC
	MP614	TTGGTCTCTCAGACTCCGATGATGCCGTCGATCG
p <i>PT11m</i> cloning for p <i>PT11m</i> :GUS	MP609	TTGGTCTCTGCGGGGAGCAATAGACGAGGGATGCC
	MP610	TTGGTCTCTTACGCGGAGGTAAATACATGAAAAATTAA AAGTTAGTTAGC
	MP611	TTGGTCTCTCGTACACTGAACTACCCATTACACACC
	MP612	TTGGTCTCTATTCCGCGGAGGCAGATAATCATGATTG
	MP613	TTGGTCTCTGAATACCAAAAACGACGCATTTCGCTCC
	MP614	TTGGTCTCTCAGACTCCGATGATGCCGTCGATCG
p <i>CCD7</i> cloning for p <i>CCD7</i> :GUS	MP549	TTTGGTCTCAGCGGGGGCGTGCACTGCAAGCATC
	MP550	TTTGGTCTCAATGATGTCTGCAAGGACCCAGAGCTCTAC
	MP551	TTTGGTCTCATATTCTCTGTTCTTTCCACC
	MP552	TTTGGTCTCACAGACTTTGGACTTGGCCTCCTTC
p <i>CCD7m</i> cloning for p <i>CCD7m</i> :GUS	MP549	TTTGGTCTCAGCGGGGGCGTGCACTGCAAGCATC
	MP591	TTTGGTCTCATCCGCGTAAGTTATAGCCCCGTTCTGTTTTGG ATTTTGATGGCACATTTTTC
	MP592	TTTGGTCTCAGGATCCGGGGAAAAATATTGAACTGGAATTAG
	MP552	TTTGGTCTCACAGACTTTGGACTTGGCCTCCTTC
pZAS cloning for pZAS:GUS	MP545	TTTGGTCTCAGCGGATATTTGGATGGTATGCAAAGCACATG
	MP546	TTTGGTCTCAAGTTACGTACTCCCTCTGTTTCAC
	MP547	TTTGGTCTCAAACCTACGTACATATACCTAACGTAAC
	MP548	TTTGGTCTCACAGATCTGCTAGTAAAAAAGCCTAAATCC
pZAS <i>m</i> cloning for pZAS <i>m</i> :GUS	MP545	TTTGGTCTCAGCGGATATTTGGATGGTATGCAAAGCACATG
	MP585	TTTGGTCTCAGTACTCAGAAAAAAATTTCCGTCCCTTGTC
	MP586	TTTGGTCTCAGTACGCGGACAACGGGTCGTAGTCTTTAGTTATC
	MP587	TTTGGTCTCATACGCGCAATAAAAAAGACGACAAAAAAAT ACATCATAAAAAATCGATG
	MP588	TTTGGTCTCACGTAGCATGGTTTTTTCTTTTCTTTCCAG
	MP589	TTTGGTCTCAGAAGTCACGGAACCATCTTGGTG
	MP590	TTTGGTCTCACTTCGCGGACAAGATGACAAATGGAATTTTCATCAC
	MP548	TTTGGTCTCACAGATCTGCTAGTAAAAAAGCCTAAATCC

Table S4. Plasmid construction by Golden Gate cloning (Level I, II and III). The Golden Gate toolbox was previously described⁶⁰.

Purpose	Name	Description
Golden Gate level I (LI) elements		
	LI <i>cPHR2</i>	PCR amplification of <i>OsPHR2</i> coding sequence from Nipponbare cDNA with MP503 + MP508 and assembly by Bpil cut ligation into LI pUC57 plasmid (BB03).
	LI C-D <i>GUS</i>	ref. 81
Golden Gate level II (LII) plasmids		
	LII F 3-4 p35S: <i>cPHR2</i>	Assembled by Bsal cut ligation from: LI A-C p35S (G009) + LI dy B-C (BB6) + LI <i>cPHR2</i> + LI D-E c-Myc (G070) + LI E-F 35S-T (G059) + LI dy F-G (BB09) + LII R 3-4 (BB24)
	LIIc F 1-2 p <i>Ubi</i> : <i>mCherry</i>	Assembled by Bsal cut ligation from: LI A-B p <i>Ubi</i> (G007) + LI B-C (BB06) dy + LI C-D <i>mCherry</i> (G023) + LI D-E (BB08) dy + LI E-F 35S-T (G059) + LI F-G dy (BB09) + LIIc F 1-2 (BB30)
	LIIc R 5-6 p35S: <i>mCherry</i>	Assembled by Bsal cut ligation from: LI A-B p35S (G009) + LI B-C (BB06) dy + LI C-D <i>mCherry</i> (G023) + LI D-E (BB08) dy + LI E-F 35S-T (G059) + LI F-G dy (BB09) + LIIc R 5-6 (BB30)
	LIIc F 3-4 p <i>Ol</i> : <i>GUS</i>	Assembled by Bsal cut ligation from: LI A-B Esp3I- <i>lacZ</i> dy (G082) + LI B-C dy (BB06) + LI C-D <i>GUS</i> + LI D-E dy (BB08) + LI nos-T (G006) + LI F-G dy (BB09) + LIIc F 3-4 (BB33)
Golden Gate level III (LIII) plasmids for plant transformation		
Overexpression of <i>OsPHR2</i> in <i>N. benthamiana</i> leaves	LIIIβ F A-B p35S: <i>cPHR2</i>	Assembled by Bpil cut ligation from: LII dy 1-2 (BB63) + LII dy 2-3 (BB39) + LII F 3-4 p35S: <i>cPHR2</i> + LII dy 4-5 ins (BB44) + LIIc R 5-6 p35S: <i>mCherry</i> + LIIIβ F A-B (BB53)
Esp3I compatible destination backbone for Localization of promoter activity	Esp3I cut ligation compatible backbone: LIIIβ fin p <i>Ubi</i> : <i>mCherry</i> _p <i>Ol</i> : <i>GUS</i> Esp3I	Assembled by Bpil cut ligation from: LIIc F 1-2 p <i>Ubi</i> : <i>mCherry</i> + LII 2-3 ins (BB43) + LIIc F 3-4 p <i>Ol</i> : <i>GUS</i> + LII dy 4-6 (BB41) + LIIIβ fin (BB52)
Bsal compatible destination backbone for Localization of promoter activity	Bsal cut ligation compatible backbone: LIIIβ fin p <i>Ubi</i> : <i>mCherry</i> _p <i>Ol</i> : <i>GUS</i> Bsal	Assembled by Esp3I cut ligation from: LIIIβ fin p <i>Ubi</i> : <i>mCherry</i> _p <i>Ol</i> : <i>GUS</i> Esp3I + LI A-B Esp3I-ccdB dy (G084)
Transactivation of p <i>GDPD2</i> : <i>GUS</i> in <i>N. benthamiana</i> leaves	LIIIβ fin p <i>Ubi</i> : <i>mCherry</i> _p <i>GDPD</i> : <i>GUS</i>	Assembled by Bsal cut ligation from: LIIIβ fin p <i>Ubi</i> : <i>mCherry</i> _p <i>Ol</i> : <i>GUS</i> Bsal + PCR amplicon MP595 + MP600 amplified from Nipponbare genomic DNA
Transactivation of p <i>GDPD2m</i> : <i>GUS</i> in <i>N. benthamiana</i> leaves	LIIIβ fin p <i>Ubi</i> : <i>mCherry</i> _p <i>GDPDm</i> : <i>GUS</i>	Assembled by Bsal cut ligation from: LIIIβ fin p <i>Ubi</i> : <i>mCherry</i> _p <i>Ol</i> : <i>GUS</i> Bsal + PCR amplicons MP595 + MP596, MP597 + MP598, MP599 + MP600

		amplified from LIIIβ fin <i>pUbi:mCherry_pGDPD:GUS</i>
Transactivation of <i>pPT11:GUS</i> in <i>N. benthamiana</i> leaves	LIIIβ fin <i>pUbi:mCherry_pPT11:GUS</i>	Assembled by BsaI cut ligation from: LIIIβ fin <i>pUbi:mCherry_pOI:GUS</i> BsaI + PCR amplicons MP609 + MP614 amplified from Nipponbare genomic DNA
Transactivation of <i>pPT11m:GUS</i> in <i>N. benthamiana</i> leaves	LIIIβ fin <i>pUbi:mCherry_pPT11m:GUS</i>	Assembled by BsaI cut ligation from: LIIIβ fin <i>pUbi:mCherry_pOI:GUS</i> BsaI + PCR amplicons MP609 + MP610, MP611 + MP612, MP613 + MP614 amplified from LIIIβ fin <i>pUbi:mCherry_pPT11:GUS</i>
Transactivation of <i>pCCD7:GUS</i> in <i>N. benthamiana</i> leaves	LIIIβ fin <i>pUbi:mCherry_pCCD7:GUS</i>	Assembled by BsaI cut ligation from: LIIIβ fin <i>pUbi:mCherry_pOI:GUS</i> BsaI + PCR amplicons MP549 + MP550, MP551 + MP552 amplified from Nipponbare genomic DNA
Transactivation of <i>pCCD7m:GUS</i> in <i>N. benthamiana</i> leaves	LIIIβ fin <i>pUbi:mCherry_pCCD7m:GUS</i>	Assembled by BsaI cut ligation from: LIIIβ fin <i>pUbi:mCherry_pOI:GUS</i> BsaI + PCR amplicons MP549 + MP591, MP592 + MP552 amplified from LIIIβ fin <i>pUbi:mCherry_pCCD7:GUS</i>
Transactivation of <i>pZAS:GUS</i> in <i>N. benthamiana</i> leaves	LIIIβ fin <i>pUbi:mCherry_pZAS:GUS</i>	Assembled by BsaI cut ligation from: LIIIβ fin <i>pUbi:mCherry_pOI:GUS</i> BsaI + PCR amplicons MP545 + MP546, MP547 + MP548 amplified from Nipponbare genomic DNA
Transactivation of <i>pZASm:GUS</i> in <i>N. benthamiana</i> leaves	LIIIβ fin <i>pUbi:mCherry_pZASm:GUS</i>	Assembled by BsaI cut ligation from: LIIIβ fin <i>pUbi:mCherry_pOI:GUS</i> BsaI + PCR amplicons MP545 + MP585, MP586 + MP587, MP588 + MP589, MP590 + MP548 amplified from LIIIβ fin <i>pUbi:mCherry_pZAS:GUS</i>

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