Phosphate (P) is an essential macronutrient for plant growth. Roots employ adaptive mechanisms to forage for P in soil. Root hair elongation is particularly important since P is immobile. Here we report that auxin plays a critical role promoting root hair growth in Arabidopsis in response to low external P. Mutants disrupting auxin synthesis (taa1) and transport (aux1) attenuate the low P root hair response. Conversely, targeting AUX1 expression in lateral root cap and epidermal cells rescues this low P response in aux1. Hence auxin transport from the root apex to differentiation zone promotes auxin-dependent hair response to low P. Low external P results in induction of root hair expressed auxin-inducible transcription factors ARF19, RSL2, and RSL4. Mutants lacking these genes disrupt the low P root hair response. We conclude auxin synthesis, transport and response pathway components play critical roles regulating this low P root adaptive response.
Phosphorus is an essential element for plant growth and development, which is absorbed in the form of phosphate (P). This macro-nutrient is extremely reactive and readily precipitates with Al, Fe, or Ca making P one of the most limiting nutrients in soil. Plants have evolved a number of mechanisms to acquire soil P. Key adaptations under P deficient conditions include exudation of organic acids and phosphatases to solubilize soil P, induction of high affinity phosphate transporters and reprogramming of root system architecture (RSA).

Root architectural changes under P stress include altered root to shoot ratio, increased root surface area and shallower root angle to improve foraging in the P rich top soil. At the cellular level, plants adapt to low P by increasing root hair elongation and density. Root hairs are formed by specialized epidermal cells called trichoblasts. The distribution of trichoblasts is species dependent and in Arabidopsis, trichoblasts (or H cells) are arranged in cell files interspersed with files of non-root hair forming cells (atrichoblast or N cells). Molecular genetic studies have identified several key genes regulating root hair development. Among them bHLH transcription factors ROOT HAIR DEFECTIVE 6 (RHD6) and ROOT HAIR DEFECTIVE 6-LIKE 2 (RSL2), and ROOT HAIR DEFECTIVE 6-LIKE 4 (RSL4) appear to be crucial for hair morphogenesis. Whereas RHD6 is required for early stages of trichoblast development, RSL4 appears to be the key gene regulating hair length. RSL2 has been shown to be responsive to P deficiency. Despite these impressive recent advances, the molecular mechanisms regulating root hair elongation under P stress are much less well understood.

We report that the promotion of Arabidopsis root hair elongation under low external P requires the concerted activities of the auxin synthesis, transport, and response pathway components. Mutants disrupting these distinct auxin pathways attenuate the ability of root tissues to promote hair elongation in response to low external P conditions. Reporter lines revealed that under low P conditions auxin levels are up-regulated at the root tip, whilst targeted expression studies demonstrate that auxin is mobilized via AUX1 to the root hair differentiation zone. Elevated auxin levels in trichoblasts trigger a gene expression cascade, mediated by transcription factors ARF19, RSL2, and RSL4 that promotes hair elongation. In parallel papers, we demonstrate that this auxin-dependent root hair response to low external P is highly conserved in the monocot model rice and relies on TIR1 to promote hair elongation via elevated intracellular auxin and calcium signaling responses.

Results

IAA increases in Arabidopsis roots grown in low external P. Increased root hair elongation is a highly conserved adaptive response to low external P in many plant species including the eudicot and monocot models Arabidopsis (Fig. 1a) and rice. Earlier genetic evidence suggested that auxin may facilitate root hair elongation under low external P conditions. We initially tested whether auxin levels were up-regulated under low external P stress.

Fig. 1 Root hair elongation is a conserved response to low P in Arabidopsis and results in IAA accumulation in the root tip. a Representative images of 9 day old seedlings grown in 3 µM (LP) or 312 µM P (HP). Scale bar = 0.5 mm. b IAA abundance was measured in 9 days old root tips (tip) and root hair region (RH) in Arabidopsis seedlings grown under low (LP) and high P (HP). Error bars represent standard error, N = 4. * Indicates significant difference (q-value < 0.05) judged by Student’s t-test. c–h. Confocal images of 9 day old Arabidopsis roots expressing auxin response reporter DR5:VENUS (yellow) under low (LP) and high P (HP) showing median (c, d) and surface (e, f) views. g and h are maximum projections. Propidium iodide was used for counter staining (red). Scale bar = 50 µm.
P conditions. Arabidopsis seedling root tips and root hair zone samples were micro-dissected, then their IAA levels were quantified using gas chromatography-tandem mass spectrometry (GC-MS/MS). Our MS profiling results revealed a significant increase in IAA accumulation in the root hair zone of seedlings grown under low P (3 μM) versus high P (312 μM) (Fig. 1b). We next tested whether the low P induced IAA levels resulted in an increase in auxin response. The DR5:Venus auxin responsive reporter revealed an increased signal in the root apex (primarily in columella cells, quiescent center, stele, and lateral root cap cells) and root hair zone in seedlings grown under low P versus high P conditions (Fig. 1c–h). Our results reveal that Arabidopsis roots exposed to low external P conditions increase their IAA levels, triggering auxin responses in root tip and hair zone cells.

Root hair elongation under low P requires auxin synthesis. To probe the molecular events regulating root hair elongation under different external P conditions, we performed Agilent array-based gene expression profiling on seedling root tissues either grown under low or high P conditions (see Methods section and Gruber et al.5). Our array results revealed that transcript levels of the auxin biosynthesis gene TRYPTOPHAN AMINOTRANSFERASE OF ARABIDOPSIS1 (TAA1) increased in response to low external P conditions (Fig. 2a). This was validated using genomic TAA1::GUS translational reporter lines which revealed levels of the enzyme were up-regulated in root apices grown in low external P conditions (Supplementary Fig. 1). Array profiling also revealed that transcripts of the auxin inducible IAA degrading enzyme DIOXYGENASE FOR AUXIN OXIDATION1 (DAO1) are significantly upregulated under low P conditions (Fig. 2a). Our transcriptomic results suggest that the regulation of key auxin homeostasis genes may play an important regulatory role during root hair elongation under low P conditions.

To functionally characterize the role of auxin homeostasis in general, and the specific roles of IAA biosynthesis gene TAA1 and IAA degradation gene DAO1, the root hair responses to low P were characterized in loss of function taa1 and dao1 mutants. Our results revealed that, while there was a ~20 fold increase in root hair length under low P versus high P conditions in WT, this response was attenuated in taa1 mutant roots by more than 60% (Fig. 2b, c). We also performed P experiments using a buffered nutrient system (see Methods section). Irrespective, we observed low P responsive root hair elongation in WT, whereas aux1 and taa1 almost completely disrupted this response (Supplementary Fig. 2). We also performed split P experiments where anchor roots (see Methods section) from the same plants were transferred to LP and HP media. In these experiments, WT roots on LP (but not HP) showed root hair elongation (Supplementary Fig. 3). In contrast, taa1 roots grown on LP did not respond to low P conditions (Supplementary Fig. 3). These experiments revealed that root hair elongation under low P is a local adaptive response which is TAA1 dependent.

We also investigated the root hair phenotype of two different DAO1 alleles recently described by Porco et al.25. The dao1.2D allele (in which the T-DNA is inserted 3’ of the coding sequence, increases DAO1 expression)25 had shorter hairs under P deficient conditions (Fig. 2d, e). Our observation reveals that root auxin levels limit root hair growth. In contrast, in the null dao1-1 allele, root hairs were slightly longer compared to the wild type plants in both P sufficient and deficient conditions (Fig. 2d, e). Nevertheless, the absence of a strong phenotype in the loss of function allele dao1.1 (Fig. 2d, e), questioned the importance of the auxin degradation pathway and/or may indicate redundancy with other hormone homeostasis mechanisms such as auxin conjugation, as reported by Porco et al.25. Taken together, our results suggest that auxin synthesis, and possibly degradation, play an important role in promoting the local, adaptive root hair elongation response to low external P conditions.

AUX1 mediates shootward auxin transport in low P conditions. Auxin transport is also crucial for hair elongation in rice roots grown in low external P.23 Given that mutations in the rice auxin influx transporter OsAUX1 disrupt this root hair response under low P conditions,23, and Arabidopsis aux1 mutants have been reported to have a root hair defect,26 we tested whether mutations in the orthologous Arabidopsis AUX1 gene necropothesized this effect. Our results revealed that root hair elongation response under low P was severely disrupted in the loss of function Arabidopsis mutant aux1-22 compared to WT controls (Fig. 2b, c). Hence, AUX1-mediated auxin influx is required for the increased root hair elongation response to low external P in Arabidopsis as reported in rice.23

Next, we identified the root tissue(s) in which AUX1-mediated auxin transport is required to facilitate the root hair elongation response to low external P. AUX1 is predominantly expressed in several tissues at the apex of the Arabidopsis root, including proto-phloem, columella, lateral root cap (LRC), and epidermal cells.27-29 AUX1 expression in the LRC and epidermal cells has been reported to be important for root hair development under nutrient replete conditions.30 We tested whether expressing the auxin influx carrier in a sub-set of its expression domain is sufficient to restore the aux1 root hair elongation defect under low P conditions (Fig. 2b, c). We targeted expression of a HA-tagged AUX1 sequence in aux1-22 LRC and epidermal root cells using the tissue specific aux1-22 J0951 AUX1-HA line. Expression in the LRC and epidermal cells was confirmed using an anti-HA antibody to reveal AUX1 spatial expression in root tissues after performing immuno-localization studies in the aux1-22 J0951 AUX1-HA line (Fig. 3a, b). Next, we tested whether J0951 AUX1-HA expression rescued the defective aux1 root hair response under low P conditions. As shown in Fig. 3c, aux1-22 J0951 AUX1 rescued the aux1 root hair elongation defect under low P conditions. Hence, AUX1 expression in lateral root cap and epidermal cells is sufficient to rescue low P inducible root hair elongation. We reason that AUX1 is required to mobilize auxin from the root apex to hair differentiation zone (referred to as "shootward" auxin transport). Dindas et al.24 have demonstrated that AUX1 contributes more than 80% of the total root hair auxin uptake capacity. Hence, in the absence of auxin uptake carrier activity in aux1, the IAA-dependent root hair elongation response to low P conditions is severely perturbed.

ARF19 regulates root hair elongation in low P conditions. On reaching the root hair zone, auxin triggers a combination of short term and longer term signaling events. Dindas et al.24 demonstrated that auxin triggered AUX1-dependent membrane depolarization. TIR1-mediated auxin perception and Ca2+ signaling is closely coupled events in root hairs, occurring within seconds of each other. These rapid signaling events function to stabilize a calcium maxima close to the tip and promote integration of new cell wall materials. Longer term auxin responses like transcription are also required, to increase synthesis of new cell wall material, for example, to promote root hair elongation.

ARF7 and ARF19 are transcriptional activators32 that regulate the majority of auxin responsive gene expression in seedling root tissues33. To investigate the role ARF7 and ARF19 during auxin-dependent root hair elongation responses to low P, we characterized the phenotypes of loss of function arf7 and arf19 mutant alleles. We observed that arf7-1 seedlings exhibited a slight increase in root hair elongation compared to WT in low P
conditions (Supplementary Fig. 4). In contrast, the promotion of root hair elongation under low P versus high P conditions was attenuated in \textit{ARF19} knock out alleles (\textit{arf19-a} and \textit{arf19-b}; Fig. 4a–c) (Fig. 4b, c). Hence, our studies suggest that \textit{ARF19} (rather than \textit{ARF7}) represents a key transcription factor promoting auxin dependent root hair elongation in response to low external P.

How is \textit{ARF19} activity enhanced under low P conditions? \textit{ARF19} is reported to be strongly auxin-inducible\(^3\). Hence, elevated auxin levels under low P conditions may increase \textit{ARF19} transcript abundance. Consistent with this model, transcript profiling revealed that \textit{ARF19} mRNA abundance increases under low P conditions (Fig. 4d). We validated this result using a reporter composed of the \textit{ARF19} promoter driving the expression of 3 tandem copies of a nuclear localized VENUS fluorescent protein (\textit{pARF19::3×Venus-NLS}). \textit{pARF19}-driven reporter fluorescence was ~2 fold higher in low P versus high P conditions (Fig. 4e; Supplementary Fig. 5a). Expression of the \textit{pARF19}-

---

**Fig. 2** Auxin homeostasis is crucial for root hair elongation under low P. 

\textbf{a} Genes regulating auxin biosynthesis (\textit{TAA1}) and auxin degradation (\textit{DAO1}, \textit{DAO2}) are upregulated under low P (LP) conditions. *Indicate significant difference (q-value < 0.0001) judged by Student’s t-test and after Benjamini and Hochberg false discovery rate correction. 

\textbf{b, c} Representative images (\textbf{b}) and boxplot (\textbf{c}) showing root hair length in the root hair zone under low (LP) or high P (HP) in Columbia (Col), \textit{aux1} (\textit{aux1}-22) and \textit{taa1} mutant seedlings. \(N = 3\), scale bar = 0.5 mm. *, **, and *** indicate significant difference (q-value < 0.005, 0.0001 and 0.00001, respectively) judged by Student’s t-test.

\textbf{d, e} Representative images (\textbf{d}) and boxplot (\textbf{e}) showing root hair length under low (LP) or high (HP) P in Columbia (Col) and \textit{dao1} (\textit{dao1-2} & \textit{dao1-2D}) mutant seedlings. \(N = 3\), scale bar 0.5 mm. *, **, and *** indicate significant difference (q-value < 0.005, 0.0001, and 0.00001, respectively) judged by Student’s t-test.
driven reporter also extended throughout the root hair differentiation zone under low P conditions. We conclude from our mutant, transcriptomic, and reporter studies that ARF19 represents a key regulator for auxin-inducible root hair elongation under low P conditions.

Transcription factors RSL2 and RSL4 are induced in low P. ARF19 is likely to promote root hair elongation by regulating the expression of genes like RSL2 and RSL4 that encode basic Helix-Loop-Helix transcription factors that promote root hair initiation and elongation. We initially tested whether rsl4 mutant lines disrupt the low P root hair elongation response. Root hair lengths in rsl4-1 were shorter than WT (Fig. 5a, b). This mutant phenotype suggests that RSL4 functions as part of the low P inducible root hair elongation machinery. However, the low P root hair defect exhibited by the rsl4-1 allele was not as severe as observed with the arf19 mutant alleles (Fig. 4b, c). This implies that ARF19 regulates other genes, in addition to RSL4, to promote root hair elongation under low P conditions. Indeed, transcript profiling revealed that mRNAs for other closely related members of the bHLH family (such as RSL2) are up-regulated under low P conditions (Fig. 5c; Supplementary Fig. 6a). To test the role of RSL2 in regulating root hair elongation in low P, we characterized its loss of function mutant phenotype. This revealed a striking defect in the rsl2 mutant root hair response to low P, which was even more severe than rsl4 (Fig. 5c). Next, we monitored its expression using an RSL2::GFP reporter line. This also revealed an increase in the RSL2::GFP signal under low P conditions.

RSL4 has been shown to be induced by auxin. We searched for the presence of auxin response element(s) in the RSL4 (and RSL2) promoter sequence. Motif analysis revealed highly conserved auxin response elements in both RSL promoters (Supplementary Fig. 6b) that represent target(s) for ARFs such as ARF19. Consistent with the bHLH gene being a target for regulation by ARF19, we observed the sequential induction of ARF19 and then RSL4 mRNAs in our transcriptomic time course dataset (Fig. 4d and Supplementary Fig. 5b). RSL4 protein abundance is correlated with root hair elongation. This prompted us to monitor RSL2 and RSL4 levels under low P versus high conditions employing GFP fusion proteins driven by their native promoters. RSL2 and RSL4 levels are higher (Supplementary Fig. 5b) under low P than high P conditions (Fig. 5d and ref. 19). Spatially, the RSL2 and RSL4 GFP reporters overlap with the pARF19-driven GFP reporter in the root hair differentiation zone (Fig. 5c), consistent with these genes being targets for regulation by ARF19 under low P conditions.

Discussion

Developmental plasticity is crucial for plant roots to optimize foraging for resources in highly heterogeneous soil environments. Root hair elongation represents an important adaptive response to acquire immobile nutrients such as P. We report that Arabidopsis roots exposed to low external P conditions employ the hormone auxin to enhance uptake of this key nutrient by promoting root hair elongation.

Our study reveals that the promotion of root hair elongation under low external P requires the concerted activity of auxin synthesis, transport and transcription-related components, from which we have been able to construct a mechanistic framework for this important root adaptive response pathway. Using the DR5-based auxin response reporter and direct auxin measurements we show that IAA accumulates in the root hair zone when grown in low P conditions (Fig. 1b, c). Currently it is not clear where IAA is synthesized in response to low P conditions. Transcriptomic and genetic approaches reveal that TAA1 is required for this low P induced root hair response. TAA1 encodes a key enzyme in auxin biosynthesis, which is expressed in the root apex and induced under low P conditions (Fig. 2a). The taa1 mutant disrupts root hair elongation under low P (Fig. 2b, c). Taken together these results suggest that TAA facilitates auxin synthesis under low P conditions presumably in the root apex. Split root experiments suggest that root hair elongation is a local response and is independent of plant P status. As TAA1 is expressed in the root apex and IAA can be synthesized in the root apex, it is likely that there is local auxin synthesis in the root apex under low P. Consistent with this conclusion, targeted expression studies in Arabidopsis revealed that low P induced hair elongation required IAA to be mobilized via the AUX1 influx carrier from the root apex to the root differentiation zone (Fig. 3).

In a co-submitted paper, Giri et al. reported that this was also the case in rice roots. Our observations made in Arabidopsis and rice reveal the functional conservation of the AUX1-dependent shootward auxin transport pathway (from the root apex to hair differentiation zones) for this important root P adaptive response in these distantly related angiosperm species.

On reaching the root hair zone, AUX1-dependent IAA uptake triggers very rapid (i.e., seconds to minutes) and longer term (i.e., minutes to hours) signaling processes. Dindas et al. reported that the rapid auxin-dependent root hair response to low P relied

---

**Fig. 3** Expressing AUX1 in LRC and epidermal cells can restore auxin root hair defect under low P. a The schematic diagram of the Arabidopsis root apex. b Confocal image showing AUX1 expression in lateral root cap and epidermal cells in aux1-22J0951>>AUX1 line. Scale bar = 20 μm. c Representative images showing root hair growth in the driver line aux1-22J0951 and aux1-22J0951>>AUX1 lines under low (LP) or high P (HP). N = 3, scale bar = 0.5 mm.
**Fig. 4** Auxin Response Factor **ARF19** regulates root hair elongation under low P in Arabidopsis. 

a Cartoon showing positions of two different **arf19** mutant alleles used in this study (top) and Reverse Transcription-PCR results showing that **arf19-a** and **arf19-b** are null alleles. 

b Representative images showing root hair growth of Columbia (Col), **arf19-a** and **arf19-b** seedlings under low P (LP) or high P (HP) conditions. N = 3, scale bar = 0.5 mm. 

c Boxplot showing root hair growth of Columbia (Col), **arf19-a** and **arf19-b** seedlings under low P (LP) or high P (HP) conditions. *, **, and *** indicate significant difference (q-value < 0.005, 0.0001, and 0.00001, respectively) judged by Student’s t-test. 

d Expression profiling studies showing **ARF19** expression under low P (LP) or high P (HP) conditions. *Indicates significant difference (q-value < 0.05) judged by Student’s t-test and after Benjamini and Hochberg false discovery rate correction. 

e Fluorescence intensity of p**ARF19:3XVENUS-NLS** seedlings grown under low P (LP) and high P (HP). N = 3, scale bar = 100 µm.
on TIR1 to promote hair elongation via elevating intracellular auxin and calcium signaling. Maintenance of calcium maxima at the root tip is known to be essential for hair growth to coordinate secretion, endocytosis, and actin dynamics. Fascinatingly, Dindas et al. observed that the calcium maxima at the root hair tip was transient and dependent on auxin. Hence, to maintain this tip focused calcium gradient and promote hair elongation would require a sufficient duration and level of auxin, as experienced during root exposure to low external P. Increased auxin in root hair cells will also trigger longer term transcriptional responses. This increase in auxin response could help explain why the root hair defects in the auxin mutants axr1 and axr2 under control conditions, can be rescued by low external P treatment. Moreover, the existence of RSL4-dependent and independent auxin mediated root hair response pathways is consistent with the reduced severity of the rsl4-1 defect in low P compared to rsl2 and axr1 alleles (Figs. 4c and 5b).

In summary, we have constructed a mechanistic framework for the low P induced root hair elongation response pathway. We demonstrate that low P elevates IAA levels in the root apex facilitated by TAA1 mediated auxin synthesis and AUX1 dependent auxin transport. The resulting increase in IAA levels leads to the induction of ARF19 resulting in RSL2 and RSL4 induction in the elongation and differentiation zones respectively promoting root hair elongation (Fig. 6).

Methods

**Plant materials.** Arabidopsis ecotype Columbia was used as the wildtype in all experiments. arf19 mutant alleles arf19-a (SAIL_92_G09, N872822) and arf19-b (SK-24322, N1007173) were obtained from the Nottingham Arabidopsis Stock Centre and the insertion was confirmed by PCR using forward and reverse primers (obtained using T-DNA primer design tool) in combination with LB3 (for arf19-a)
and pSKTAIL-L1 (for arf19-b) border primers (Supplementary Table 1). ARF19 expression in these alleles was checked by Reverse Transcription-PCR using pri-

ermers ARF19_RT_For and ARF19_RT_Rev (Fig. 4a and Supplementary Table 1). Following mutant alleles: arf7-1, arf19 (arf19-a and arf19-b), rsl2, rsl3-1 (back-

crossed four times), aux1 (aux1-22), tol1, dao1.1, and dao1.2D and reporters: pARF19-3xVENUS-NLS; pRSL2::GFP-RSL2, and pRSL2-GFP-RSL4 were used in

phosphate treatment experiments.

Growth conditions. Seeds were surface sterilized with 70% ethanol for 10 min and

100% ethanol for 30 s followed by five washes with sterile water and stratified at 4 °

c for 48 h in dark. For phosphate treatment experiments, seeds were germinated

in 100% ethanol for 30 s followed by

Growth conditions

pARF19::3XVENUS-NLS; pRSL2::GFP-RSL2

and pSKTAIL-L1 (for arf19-b) border primers (Supplementary Table 1). ARF19 expression in these alleles was checked by Reverse Transcription-PCR using pri-

mers ARF19_RT_For and ARF19_RT_Rev (Fig. 4a and Supplementary Table 1). Following mutant alleles: arf7-1, arf19 (arf19-a and arf19-b), rsl2, rsl3-1 (back-

crossed four times), aux1 (aux1-22), tol1, dao1.1, and dao1.2D and reporters: pARF19-3xVENUS-NLS; pRSL2::GFP-RSL2, and pRSL2-GFP-RSL4 were used in

phosphate treatment experiments.

Growth conditions. Seeds were surface sterilized with 70% ethanol for 10 min and

100% ethanol for 30 s followed by five washes with sterile water and stratified at 4 °

c for 48 h in dark. For phosphate treatment experiments, seeds were germinated for 5 days on vertical 1/4th MS (Murashige and Skoog) Plates (supplemented with 1% agar, at pH 5.7) in growth room (22 °C continuous light; 100–125 µmol/m²/s). 5-day-old seedlings were transferred to MS salt (without phosphate) plates sup-

plemented with 1% agar and 3 µM (low) and 312 µM (high) P. pH of media was set to 5.7 and low P media was supplemented with equimolar concentration of KCl. We also performed P experiments using a buffered nutrient system. Plants were grown under sufficient (MB, 50 µM soluble P) or limiting (LR, 3 µM soluble P) conditions with a buffered phosphorus delivery regime. Plants were grown for 3 days (16 h day, 100–120 µmol min⁻¹) on 1/4 strength MS media without sucrose before being transferred to the respective treatment. After four days of growth, images were obtained on a dissecting scope and root hair lengths were measured. Ten plants per genotype, per treatment were measured. Error bars represent the standard error of the mean.

Transcriptome studies. For microarray experiments, plants were grown on 0.5

MS Difco agar as described previously using 625 and 100 µM phosphate for control and

P-deficient conditions, respectively. RNA was extracted from root material using an RNeasy plant mini kit (Qiagen) with on-column DNase treatment according to the manufacturer’s instructions. cDNA was prepared using random hexamer primers and SuperScript II reverse transcriptase (Life Technologies) according to the manufacturer’s protocol. RNA amplification, labeling and hybri-
dization to Agilent microarrays (Arabidopsis (V4, 021169) Gene Expression Microarray) were conducted following the manufacturer’s protocol (Agilent Technologies). Analysis of microarray data was performed with the R package

limma42. Raw feature intensities were background corrected using “normexp,” and “quantile” normalization method was used to normalize between arrays.

Differential expression was performed by fitting a linear model to log2-transformed
data by an empirical Bayes method42. The Benjamini and Hochberg method was used to correct for multiple testing. The transcriptome raw data is deposited in electronic data archive library (eDAL—https://edal.ipk-gatersleben.de) repository under the accession code https://doi.org/10.5447/IPK/2018/4.

Auxin estimation. Roots tip (~1 mm) and root hair zone (next 1.5 mm region) from

9-days-old seedlings grown on low and high P media were cut using a sharp

scissor and snap frozen in liquid nitrogen. Seventyfive roots were collected per

replicate with four replicates per experiment. Five-hundred picograms of 13C6-IAA

internal standard was added to each sample before purification. Auxin estimation was
done as described previously45, with minor modifications.

GUS quantification. GUS intensity was measured using imageJ software. Briefly,

Leica image in imageJ/fiji software was imported and “color tool” in image tab was used
to split channels and convert single image into three grayscale images. Red channel image was selected and “adjust tool” in image tab was used to assign values to pixels between 0 and 255. Intensity threshold was used to adjust the pixels in the selected range (corresponds to GUS staining) that are highlighted in red. Raw integrated density (i.e., the sum of the values of the pixels in the selection) was measured using the usual measurement option (Analyze/Measure menu option) with measurements limited to only the selected pixels.

DNA constructs and transgenic materials. pRSL2::GFP-RSL419 and pRSL2::GFP-

RSL419 transgenic lines were obtained from Liam Dolan, Department of Plant Sciences, University of Oxford, Oxford, OX1 3RB, UK.

The pARF19-3xVenUS-N7ARF19 was cloned using Gateway MultiSite cloning technology. The ARF19 fragment (−4906 to +452 bp) was PCR amplified using primers ARF19_For and ARF19_Rev (Supplementary Table 1) and cloned into pDONR P4-P1R to create pARF19 pDONR P4-P1R. The latter was recombined in a multbsite Gateway reaction with the following plasmids: 3 × mVenUS-N7 pDONR211 (containing triple mVenUS coding sequences and N7 nuclear localization signal), OCS terminator pDONR P2R-P3 (containing the stop codon followed by octopine synthase (OCS) terminator), and pK734GW (the destination vector containing selection kanamycin resistance gene for in planta selection). The resulting final plasmid was transformed into Agrobacterium tumefaciens

C58HMP90 strain by electroporation and then transformed into Col-0 plants by

floral dip method46.

Data availability. The transcriptome raw data is deposited in electronic data

archive library (eDAL—https://edal.ipk-gatersleben.de) repository under the accession code https://doi.org/10.5447/IPK/2018/4. Other data and materials supporting the findings of this study may be obtained from the corresponding authors upon reasonable request.

Received: 9 January 2017 Accepted: 16 March 2018
Published online: 12 April 2018

References

1. Stetter, M. G., Schmid, K. & Ludewig, U. Uncovering genes and ploidy

involved in the high diversity in root hair density, length and response to

local scarce phosphate in Arabidopsis thaliana. PLOS One 10, e012064 (2015).

2. Zhang, Z., Liao, H. & Lucas, W. Molecular mechanisms underlying phosphate

sensing, signaling and adaptation in plants. J. Int. Plant Biol. 56, 192–220 (2014).

3. Lambers, H. et al. Phosphorus nutrition of pastureceae in severely phosphorus-

impovery soils: are there lessons to be learned for future crops? Plant Physiol. 156, 1058–1066 (2011).

4. Lynch, J. P. Root phenes for enhanced soil exploration and phosphorus

acquisition: tools for future crops. Plant Physiol. 156, 1041–1049 (2011).

5. Gruber, B., Giehl, R., Friedel, S. & von Wirén, N. Plasticity of the Arabidopsis

root system under nutrient deficiencies. Plant Physiol. 163, 161–179 (2013).

6. Bates, T. R. & Lynch, J. P. Stimulation of root hair elongation in Arabidopsis

thaliana by low Pi availability. Plant Cell Environ. 19, 529–538 (1996).

7. Williamson, L. C., SPCP, Ribroux, Fitter, A. H. & Leyser, H. M. O. Phosphate

availability regulates root system architecture in. Arab. Plant Physiol. 126, 875–882 (2001).

8. Linkohr, B. L., Williamson, L. C., Fitter, A. H. & Leyser, H. M. O. Nitrate and

phosphate availability and distribution have different effects on root system architecture of Arabidopsis. Plant J. 29, 751–760 (2002).

9. López-Bucio, J. et al. Phosphate availability alters architecture and causes

changes in hormone sensitivity in the Arabidopsis root system. Plant Physiol. 129, 244–256 (2002).
10. López-Bucio, J. et al. An auxin transport independent pathway is involved in phosphate stress induced root architectural alterations in Arabidopsis. Identification of RIG as a mediator of auxin in pericycle cell activation. Plant Physiol. 137, 681–691 (2005).

11. Müller, M. & Schmidt, W. Environmentally induced plasticity of root hair development in Arabidopsis. Plant Physiol. 134, 409–419 (2004).

12. Nacey, P. et al. A role for auxin redistribution in the responses of the root system architecture to phosphate starvation in Arabidopsis. Plant Physiol. 138, 2061–2074 (2005).

13. Brown, L. K., George, T. S., Dupuy, L. X. & White, P. J. A conceptual model of root hair ideotypes for future agricultural environments: what combination of traits should be targeted to cope with limited P availability? Ann. Bot. 112, 317–330 (2013).

14. Zhou, J. et al. OsPHR2 is involved in phosphate-starvation signaling and excessive phosphate accumulation in shoots of plants. Plant Physiol. 146, 1637–1686 (2008).

15. Leavitt, R. G. Trichomes of the root in vascular cryptogams and angiosperms. Proc. Boston Soc. Nat. Hist. 31, 273–313 (1904).

16. Salazar-Henao, J. E., Vélez-Bermúdez, I. C. & Schmidt, W. The regulation and plasticity of root hair patterning and morphogenesis. Development 143, 1848–1858 (2016).

17. Lan, P., Li, W., Lin, W. D., Santi, S. & Schmidt, W. Mapping gene activity of Arabidopsis root hairs. Genome Biol. 14, R67 (2013).

18. Deal, R. B. & Henriksøff, S. A simple method for gene expression and chromatin preparation of individual cell types within a tissue. Dev. Cell 18, 1030–1040 (2010).

19. Yi, K., Menand, B., Bel, E. & Dolan, L. A basic helix-loop-helix transcription factor controls cell growth and size in root hairs. Nat. Genet. 42, 264–267 (2010).

20. Datta, S. P., Prescott, H. & Dolan, L. Intensity of a pulse of RSL4 transcription promotes root hair elongation by transcriptionally regulating the expression of genes required for cell growth. New Phytol. 212, 944–953 (2016).

21. Bruex, A. et al. A gene regulatory network for root epidermis cell differentiation in Arabidopsis. PLOS Genet. 8, e1002446 (2012).

22. Ritchie, M. E. et al. Lmiota powers differential expression analyses for RNA-sequencing and microarray studies. Nucleic Acid. Res. 43, e47 (2015).

23. Smyth, G. K. Linear models and empirical Bayes methods for assessing differential expression in microarray experiments. Stat. Appl. Genet. Mol. Biol. 3, Article 3 (2004).

24. Andersen, S. U. et al. Requirement of B2-type cyclin-dependent kinases for meristem integrity in Arabidopsis thaliana. Plant Cell 20, 88–100 (2008).

25. Clough, S. J. & Bent, A. F. Floral dip: a simplified method for Agrobacterium-mediated transformation of Arabidopsis thaliana. Plant J. 16, 735–743 (1998).

Acknowledgements
This work was supported by the awards from the Biotechnology and Biological Sciences Research Council [grant numbers BB/G023972/1, BB/R013748/1, BB/L026848/1, BB/M018431/1, BB/P016655/1, BB/M001086/1, BB/P010520/1]; the European Research Council FUTUREROOTS Advanced Investigator grant [grant number 294729]; Leverhulme Trust [grant number RPG-2016-499]; Royal Society [grant number WM130021, NA140381]; Newton International Fellowship [NF140287] and British Council Newton Bhabha (228144076). This work was also supported by funds from the University of Nottingham Future Food Beacon of Excellence Nottingham Research and PhD+ fellowship schemes; the Interuniversity Attraction Poles Program initiated by the Belgian Science Policy Office [P7/29]; the Swedish Governmental Agency for Innovation Systems (VINNOVA), and the Swedish Research Council (V.R.). NSF-MCB1158181 and a joint studentship between the University of Nottingham and Institut National de la Recherche Agronomique (INRA). We also thank Roger Granbom (Swedish University of Agricultural Sciences) for skilful technical assistance.

Author contributions
R.B., J.G., B.K.P., R.F.H.G., A.H., R.T., J.T., N.L., M.H., K.S., A.R., U.V., J.A., A.S., and J.Y. wrote the manuscript.

R.B., J.G, B.K.P., R.F.H.G., A.H., R.T., J.T., N.L., M.H., K.S., A.R., U.V., J.A., A.S., and J.Y. contributed to the experiments.

R.B., J.G, B.K.P., R.F.H.G., A.H., R.T., J.T., N.L., M.H., K.S., A.R., U.V., J.A., A.S., and J.Y. contributed to the data analysis.

R.B., J.G, B.K.P., R.F.H.G., A.H., R.T., J.T., N.L., M.H., K.S., A.R., U.V., J.A., A.S., and J.Y. wrote the manuscript.

R.B., J.G, B.K.P., R.F.H.G., A.H., R.T., J.T., N.L., M.H., K.S., A.R., U.V., J.A., A.S., and J.Y. contributed to the writing.

R.B., J.G, B.K.P., R.F.H.G., A.H., R.T., J.T., N.L., M.H., K.S., A.R., U.V., J.A., A.S., and J.Y. contributed to the discussion.

R.B., J.G, B.K.P., R.F.H.G., A.H., R.T., J.T., N.L., M.H., K.S., A.R., U.V., J.A., A.S., and J.Y. contributed to the figures.

R.B., J.G, B.K.P., R.F.H.G., A.H., R.T., J.T., N.L., M.H., K.S., A.R., U.V., J.A., A.S., and J.Y. contributed to the tables.

R.B., J.G, B.K.P., R.F.H.G., A.H., R.T., J.T., N.L., M.H., K.S., A.R., U.V., J.A., A.S., and J.Y. contributed to the references.

R.B., J.G, B.K.P., R.F.H.G., A.H., R.T., J.T., N.L., M.H., K.S., A.R., U.V., J.A., A.S., and J.Y. contributed to the ethical considerations.

R.B., J.G, B.K.P., R.F.H.G., A.H., R.T., J.T., N.L., M.H., K.S., A.R., U.V., J.A., A.S., and J.Y. contributed to the acknowledgments.

R.B., J.G, B.K.P., R.F.H.G., A.H., R.T., J.T., N.L., M.H., K.S., A.R., U.V., J.A., A.S., and J.Y. contributed to the author contributions.

R.B., J.G, B.K.P., R.F.H.G., A.H., R.T., J.T., N.L., M.H., K.S., A.R., U.V., J.A., A.S., and J.Y. contributed to the Competing interests.

Reprints and permission information is available online at http://npg.nature.com/reprintsandpermissions/.

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2018