The epidermis coordinates thermoresponsive growth through the phyB-PIF4-auxin pathway

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In plants, an elevation in ambient temperature induces adaptive morphological changes including elongated hypocotyls, which is predominantly regulated by a bHLH transcription factor, PIF4. Although PIF4 is expressed in all aerial tissues including the epidermis, mesophyll, and vascular bundle, its tissue-specific functions in thermomorphogenesis are not known. Here, we show that epidermis-specific expression of PIF4 induces constitutive long hypocotyls, while vasculature-specific expression of PIF4 has no effect on hypocotyl growth. RNA-Seq and qRT-PCR analyses reveal that auxin-responsive genes and growth-related genes are highly activated by epidermal, but not by vascular, PIF4. Additionally, inactivation of epidermal PIF4 or auxin signaling, and overexpression of epidermal phyB suppresses thermoresponsive growth, indicating that epidermal PIF4-auxin pathways are essential for the temperature responses. Further, we show that high temperatures increase both epidermal PIF4 transcription and the epidermal PIF4 DNA-binding ability. Taken together, our study demonstrates that the epidermis regulates thermoresponsive growth through the phyB-PIF4-auxin pathway.
thermomorphogenesis refers to the morphological changes that occur in plants in response to a rise in ambient temperature\(^1\). It is characterized by elongated stems and hypocotyls, hyponastic leaf growth, and thinned leaves\(^1\). These morphological changes are considered to enhance leaf cooling capacity by increasing the leaf transpiration rate, thereby helping plants survive under heat stress\(^3,\text{4}\). A bHLH transcription factor, phytochrome interacting factor 4 (PIF4), is a key regulator of thermomorphogenesis\(^5\). The expression of PIF4 mRNA is induced when plants are exposed to high temperatures\(^5\). The resulting PIF4 protein directly activates the expression of the auxin biosynthetic genes (YUCCA8 (YUC8), TAA1, and CYP79B2), growth-promoting factors (ATHB2, PEFs, and LNGs), and brassinosteroid biosynthetic genes that induce hypocotyl growth\(^6\)–\(^10\). The high-temperature activation of PIF4 mRNA expression is mediated by the evening complex (EC)\(^11\)–\(^12\). The EC, consisting of ELF3, ELF4, and LUX, binds directly to the promoter of PIF4 and represses its expression, and it has been reported that high temperatures release this repression by preventing the binding of EC to the PIF4 promoter\(^11\)–\(^13\).

In addition to thermomorphogenesis, PIF4 is also involved in a wide range of developmental and environmental responses including photomorphogenesis, shade avoidance responses, hypocotyl gravitropism, phototropism, leaf senescence, and stomatal development\(^14\)–\(^20\). PIF4 was first identified as a component of the phytochrome signaling pathway\(^21\). Light-activated phyB interacts directly with, and phosphorylates, PIF4, resulting in the degradation of the PIF4 protein via the 26S proteasome pathway\(^20\),\(^22\). Recently, it was revealed that phyB functions as a thermosensor for thermomorphogenesis and, therefore, phyB may regulate PIF4 activity in response to changes in ambient temperature as well as light conditions\(^23\),\(^24\). Blue light receptors CRY1 and CRY2 also directly interact with PIF4 in a blue light-dependent manner\(^25\),\(^26\). Interaction with CRY1 represses PIF4 activity by interfering with its transcriptional activation activity\(^26\). PIF4 also mediates the thermosensory activation of flowering by direct regulation of the expression of FLOWERING LOCUS T (FT) in a temperature-dependent manner\(^27\), and it suppresses immunity through the negative regulation of defense-related gene expression\(^28\).

Growth regulation by various hormones is dependent on PIF4 activity\(^15\). In addition to the direct regulation of auxin biosynthetic gene expression, PIF4 also regulates auxin signaling by directly increasing the activities of auxin response factors (ARFs), at least partly by enhancing ARF binding to the promoters of target genes\(^29\),\(^30\). Growth regulation by two other hormones, brassinosteroids (BR) and gibberellic acids (GA), is also mediated by PIF4. BR promotes hypocotyl growth by repressing BIN2 kinase activity and activating the BZR1 transcription factor. BIN2 is a GSK3-like kinase that was reported to directly phosphorylate PIF4 and reduce its stability in the absence of BR signal\(^31\). The BR-regulated transcription factor, BZR1, directly interacts with PIF4 and this interaction regulates the expression of thousands of genes, promoting hypocotyl growth\(^8\). It has also been reported that BZR1 activates the expression of PIF4\(^32\). DELLA, negative regulators of GA signaling pathways, directly interact with both BZR1 and PIF4, and prevent them from binding to target gene promoters\(^33\),\(^35\). GA signals induce the degradation of DELLA repressor proteins, resulting in the de-repression of BZR1 and PIF4 activity, and thereby promoting growth. Consistent with the BR and GA signals being required for PIF4 activity, these phytohormones have been shown to be necessary for thermomorphogenic growth\(^36\).

Although the functional roles of PIF4 in the regulation of growth and development of aerial tissues have been extensively studied, the tissue-specific roles of PIF4 are still poorly understood. It was previously shown that PIF4 overexpression in vasculature promotes flowering, and endodermal PIF1, a PIF4 homolog, regulates hypocotyl negative gravitropism\(^37\)–\(^39\). Surprisingly, it was reported that epidermal, but not endodermal, phyB activity is required for light regulation of hypocotyl negative gravitropism\(^40\). Consistent with this fact, epidermal phyB is able to induce the phosphorylation and degradation of endodermal PIFs in response to light, implying that phyB generates an unidentified mobile signal that moves between the epidermis and endodermis\(^40\). Epidermal phyB was also shown to regulate many light responses, including hypocotyl growth, seed germination, and shade avoidance responses\(^40\). However, whether epidermal phyB also regulates these responses in a non-cell autonomous manner has not been determined yet.

In this study, we analyzed the tissue-specific roles of PIF4 by generating and characterizing transgenic plants expressing PIF4 or a dominant negative version of PIF4 under the control of either an epidermis- or vasculature-specific promoter. We found that epidermal PIF4, but not vascular PIF4, regulates thermoresponsive hypocotyl growth. It was also observed that the block of epidermal auxin signaling and epidermal overexpression of phyB inhibit thermoresponsive growth. We also found that the expression of PIF4 mRNA in the epidermis was induced at high temperatures. Our results demonstrate that high temperatures induce thermomorphogenesis through the activation of the PIF4-auxin module in the epidermis.

**Results**

**Tissue-specific expression of PIF4-YFP in a pif mutant.** PIF4 regulates a wide range of growth and developmental processes such as environmental or hormonal regulation of hypocotyl elongation, hypocotyl phototropism and negative gravitropism, chlorophyll biosynthesis, stomatal development and flowering\(^4\),\(^14\)–\(^15\),\(^19\),\(^20\),\(^22\),\(^41\),\(^42\). According to the Cell-Type Specific Arabidopsis eFP Browser\(^43\), PIF4 mRNA is expressed in all aerial tissues, with the highest expression in the vasculature (Supplementary Fig. 1a). To confirm the tissue-specific expression patterns of PIF4 mRNA, we analyzed β-glucuronidase (GUS) activity in transgenic plants expressing the GUS gene controlled by the PIF4 promoter (PIF4::GUS). Consistent with the data from the Arabidopsis eFP Browser, PIF4::GUS expression was strongly detected in the vasculature of cotyledons (Fig. 1a). A weak GUS signal was also detected in other tissues, including the mesophylls and epidermis, but no GUS signal was detected in the roots (Fig. 1a). To ascertain whether PIF4 in the vasculature is involved in the regulation of various developmental processes including the hypocotyl growth, we generated transgenic plants expressing PIF4-YFP under the control of vascular-specific promoter SUCROSE TRANSPORTER 2 (SUC2) in a pif1:pif3:pif4:pif5 quadruple mutant (pifq) background. We also generated transgenic plants expressing PIF4-YFP driven by MERISTEM LAYER1 (ML1) and CHLOROPHYLL A/B BINDING PROTEIN 3 (CAB3) promoter to examine the physiological functions of epidermal and mesophyll PIF4, respectively. These promoters have been successfully used in previous studies to drive the tissue-specific expression of target genes\(^44\)–\(^46\). We confirmed that the expression of PIF4-YFP is mainly restricted to the vascular bundles when under the control of the SUC2 promoter (SUC2p::PIF4:pifq) (Fig. 1b). In ML1p::PIF4:pifq transgenic plants, PIF4-YFP expression was detected specifically in the epidermis (Fig. 1c). In contrast, the PIF4-YFP signal driven by the CAB3 promoter was observed in the epidermis in addition to the mesophyll (Supplementary Fig. 1b), probably due to the leaky expression of the CAB3 promoter. Therefore, we only analyzed SUC2p::PIF4; pifq and ML1p::PIF4:pifq transgenic plants to determine the tissue-specific functions of PIF4.
Epidermal PIF4 promotes hypocotyl growth. PIF4 promotes hypocotyl growth in response to various environmental and hormonal signals. To determine the tissues in which PIF4 mediates hypocotyl growth, we measured the hypocotyl length of wildtype, pifq, ML1p::PIF4;pifq, and SUC2p::PIF4;pifq seedlings grown under white light. Consistent with previous reports, pifq mutants had short hypocotyls compared to those of the wild-type seedlings (Fig. 2a)\(^{41,49}\). The hypocotyl lengths of ML1p::PIF4;pifq seedlings were substantially longer than those of wild type and pifq mutant (Fig. 2a), indicating that epidermal PIF4 is capable of promoting hypocotyl growth. In contrast, vascular expression of PIF4 driven by the SUC2 promoter was not able to restore the short hypocotyls of pifq mutant seedlings (Fig. 2a), although the level of PIF4 in SUC2p::PIF4;pifq plants was higher than those in wild type and ML1p::PIF4;pifq plants (Fig. 2b).

A recent study showed that the circadian clock activity in the epidermal cells is important in the regulation of hypocotyl and petiole elongation\(^{48}\). We confirmed that while perturbation of the epidermal clock system by overexpression of CCA1 specifically in the epidermis (CER6p::CCA1) promotes the hypocotyl growth, neither vascular (SUC2p::CCA1) nor mesophyll (CAB3p::CCA1) clock perturbation had a significant effect on hypocotyl growth\(^{48}\) (Fig. 2c). A previous study showed that PIF4 expression is constitutively high in CCA1-overexpressing plants\(^{49}\). Intriguingly, PIF4 expression was highly elevated in the CER6p::CCA1, but showed only a marginal increase in SUC2p::CCA1 seedlings compared to those in wild-type seedlings (Fig. 2d), suggesting the hypocotyl elongation seen in CER6p::CCA1 plants might be attributable to the increased PIF4 levels in the epidermis. In support of this, the down-regulation of PIF4 expression by artificial microRNA against PIF4 (PIF4mi) suppressed the long hypocotyls of CER6p::CCA1 (Supplementary Fig. 2). Therefore, these results provide further support that epidermal PIF4 promotes hypocotyl growth.

To obtain clues for molecular mechanisms underlying the epidermal PIF4 regulation of hypocotyl growth, we performed RNA-Seq analysis of wild type, pifq, ML1p::PIF4;pifq, and SUC2p::PIF4;pifq seedlings grown under white light for 5 days. The RNA-Seq analysis identified 1281 differentially expressed genes (DEGs) between wild type and pifq seedlings (Supplementary Data 1). We refer to these DEGs as the global PIF quartet (PIF1, PIF3, PIF4, and PIF5)-regulated genes. We also compared the transcriptome of pifq with either ML1p::PIF4;pifq or SUC2p::PIF4;pifq and identified 575 epidermal PIF4 (ML1p::PIF4)-regulated genes and 1765 vascular PIF4 (SUC2p::PIF4)-regulated genes (Supplementary Data 1).

Thereafter, using Gene Set Enrichment Analysis (GSEA), we examined how the global PIF quartet-regulated genes are regulated by each tissue-specific PIF4 (Fig. 2e, f, Supplementary Fig. 3). The global PIF quartet-activated gene set shows a more significant positive enrichment in ML1p::PIF4;pifq than that in SUC2p::PIF4; pifq when compared to pifq (Supplementary Fig. 3a, b), suggesting that the PIF quartet-activated genes are more robustly activated by epidermal PIF4 than by vascular PIF4. In contrast, the global PIF4 quartet-repressed gene set shows a more significant negative enrichment in SUC2p::PIF4;pifq than that in ML1p::PIF4;pifq when compared to pifq (Supplementary Fig. 3c, d). This suggests that vascular PIF4 may play a dominant role in the regulation of PIF quartet-repressed genes. On the other hand, we tested the statistical significance of overlaps between each of the gene sets shown in Fig. 2e, f (Supplementary Data 1). The global PIF quartet-regulated genes showed a highly significant overlap with either of the epidermal PIF4- or vascular PIF4-regulated genes (Supplementary Data 1). However, the overlap between epidermal PIF4-regulated genes and vascular PIF-regulated genes were far less significant in both up- and down-regulated DEGs, showing an overlap enriched only 2.5-fold than that by random chance in both cases (Supplementary Data 1). The distinct DEGs of epidermal PIF4 and vascular PIF4 indicate high tissue-dependency in PIF4 target gene regulation, and also validate robust tissue-specific expression of PIF4 by ML1 and SUC2 promoters (Fig. 1b, c).

Gene ontology (GO) analyses revealed that GO terms including “response to auxin stimulus,” “cell wall organization,” and “cell growth” were highly enriched only in epidermal PIF4-activated genes (Fig. 2g). The results are very consistent with the positive role of epidermal PIF4 in hypocotyl growth (Fig. 2a), suggesting that epidermal PIF4 regulates hypocotyl growth through the auxin pathway. Interestingly, “iron ion transporter” and “response to nitrate” GO terms were highly enriched in the vascular PIF4-repressed genes (Fig. 2g). Thus, it is possible that the vascular PIF4 may mediate these iron and nitrogen responses. It was previously shown that PIF4 negatively regulates defense responses\(^{28}\). In GO analyses, the GO term of defense responses was enriched in both epidermal PIF4-repressed and vascular PIF4-repressed genes (Fig. 2g). Thus, it appears that both epidermal and vascular PIF4 participate in the regulation of defense responses. To confirm that epidermal PIF4 modulates auxin responses, we compared epidermal or vascular PIF4-regulated genes with the auxin-regulated genes in hypocotyls\(^{50}\). Of 266 epidermal PIF4-activated genes, 75 (61.6 expected randomly) were auxin-activated genes (Fig. 2h, Supplementary...
Regulated PIF4 direct target genes, 35 were auxin-activated genes (Fig. 2i, Supplementary Data 3). Of 75 epidermal PIF4-activated/repressed genes, were PIF4 direct target expected randomly), but not epidermal PIF4-repressed and addition, auxin-repressed genes were enriched in epidermal PIF4-vascular PIF4-activated/repressed genes, were PIF4 direct target genes51. A large portion (75 of 266) of epidermal PIF4-activated genes (11.2 expected randomly) were auxin-activated genes (Fig. 2h). In contrast, only 14 of 492 vascular PIF4-activated genes (266) of epidermal PIF4 were normalized to those of PP2A. Error bars indicate s.d. (n = 3). *P < 0.05 and **P < 0.01 (two-tailed Student’s t-test). The expression levels of PIF4 were normalized to those of PP2A. Error bars indicate s.d. (n = 3). *P < 0.05 and **P < 0.01 (two-tailed Student’s t-test). The Venn diagram shows the overlap between genes that are up-regulated by global PIF quartet (G.PIFQ), ML1 promoter-driven PIF4 (ML1:PIF4 UP), and SUC2 promoter-driven PIF4 (SUC2:PIF4 UP). The Venn diagram shows the overlap between genes that are down-regulated by the global PIF quartet, ML1:PIF4, and SUC2:PIF4. GO analyses of ML1:PIF4-regulated and SUC2:PIF4-regulated genes. Total: Arabidopsis total genes. Percentage of auxin (picloram)-activated or repressed genes in global PIF quartet-regulated or ML1:PIF4-regulated or SUC2:PIF4-regulated genes. Source data are provided as a Source Data file.

Epidermal PIF4 regulates thermoresponsive hypocotyl growth. Next, we examined whether epidermal or vascular PIF4 activity regulates thermoresponsive hypocotyl growth. The hypocotyl lengths of wild-type plants exposed to 28 °C for 3 days were approximately five times longer than those grown constitutively at 20 °C (Fig. 3a). In contrast, the hypocotyl growth in the pifq mutant was almost insensitive to the increase in ambient temperature (Fig. 3a). The hypocotyls of ML1p::PIF4:pifq plants grown at 20 °C were longer than those of wild-type plants also grown at 20 °C, and the lengths were further increased by exposure to high temperatures (Fig. 3a). In contrast, the hypocotyl growth of SUC2p::PIF4:pifq plants was only marginally increased by high temperature (Fig. 3a), like that of pifq. These results show that epidermal PIF4 activity is essential for the hypocotyl growth response to high temperatures.

**Fig. 2 Epidermal PIF4 promotes hypocotyl growth.** a Hypocotyl lengths of seedlings grown under the continuous white light at 20 °C for 7 days. Error bars indicate s.d. (n = 15 plants). **P < 0.01 (two-tailed Student’s t-test); NS, not significant (two-tailed Student’s t-test P ≥ 0.05). b qRT-PCR analysis of PIF4. Seedlings were grown in the same conditions as a. The expression levels of PIF4 were normalized to those of PP2A. Error bars indicate s.d. (n = 3). *P < 0.05 and **P < 0.01 (two-tailed Student’s t-test). c Hypocotyl lengths of seedlings grown under continuous white light at 20 °C for 7 days. Error bars indicate s.d. (n = 15 plants). **P < 0.01 (two-tailed Student’s t-test). d PIF4 expression levels in the seedlings grown in the same growth conditions as c. The expression levels of PIF4 were normalized to those of PP2A. Error bars indicate s.d. (n = 3). *P < 0.05 and **P < 0.01 (two-tailed Student’s t-test). e The Venn diagram shows the overlap between genes that are up-regulated by global PIF quartet (G.PIFQ), ML1 promoter-driven PIF4 (ML1:PIF4 UP), and SUC2 promoter-driven PIF4 (SUC2:PIF4 UP). f The Venn diagram shows the overlap between genes that are down-regulated by the global PIF quartet, ML1:PIF4, and SUC2:PIF4. g GO analyses of ML1:PIF4-regulated and SUC2:PIF4-regulated genes. Total: Arabidopsis total genes. h Percentage of auxin (picloram)-activated or repressed genes in global PIF quartet-regulated or ML1:PIF4-regulated or SUC2:PIF4-regulated genes. i Percentage of PIF4 direct target genes in ML1:PIF4-regulated or SUC2:PIF4-regulated genes. Source data are provided as a Source Data file.

In contrast, only 14 of 492 vascular PIF4-activated genes (11.2 expected randomly) were auxin-activated genes (Fig. 2h). In addition, auxin-repressed genes were enriched in epidermal PIF4-repressed genes (38 of 309, 4.3 expected randomly) (Fig. 2h, Supplementary Data 2). These results provide further support that epidermal PIF4 activates auxin responses.

We also compared epidermal or vascular PIF4-regulated genes with previously identified PIF4 direct target genes31. A large portion (75 of 266) of epidermal PIF4-activated genes (11.2 expected randomly), but not epidermal PIF4-repressed and vascular PIF4-activated/repressed genes, were PIF4 direct target genes (Fig. 2i, Supplementary Data 3). Of 75 epidermal PIF4-regulated PIF4 direct target genes, 35 were auxin-activated genes (Supplementary Data 3), suggesting that epidermal PIF4 directly regulates auxin responses.
PIF4 has previously been shown to directly activate auxin biosynthesis and signaling to induce hypocotyl growth at high temperatures\cite{8,51}. Consistent with this fact, RNA-Seq analyses found that epidermal PIF4-regulated genes include many auxin-responsive genes in hypocotyls (Fig. 2h). To further examine whether epidermal PIF4 promotes hypocotyl growth through the auxin pathway, we measured the expression levels of the auxin biosynthetic gene YUCCA8 (YUC8) and auxin-responsive genes (IAA19 and ATHB2) in seedlings that were grown continuously at 20 °C or were shifted to 28 °C for 4 h. These genes are also PIF4 direct target genes\cite{5,31}. Consistent with the previous studies\cite{5,31}, the expression of YUC8, IAA19, and ATHB2 was highly induced by exposure to high temperatures in wild-type plants, but not in pifq mutant plants (Fig. 3b, c, Supplementary Fig. 4). The expression

**Fig. 3 Epidermal PIF4 activity is required for thermoresponsive hypocotyl growth.** a Hypocotyl lengths of seedlings grown under continuous white light at 20 °C for 7 days or 20 °C for 4 days followed by 28 °C for 3 days. Error bars indicate s.d. (n = 15 plants). The P-value for the interaction term (genotype x temperature) calculated by two-way ANOVA is shown at the top. Black asterisks above the bars (in a–h) indicate significant differences (**P < 0.01 and *P < 0.05, two-tailed Student’s t-test). NS, not significant (P ≥ 0.05). Numbers above the bars (in a, d, f) indicate the ratio of hypocotyl lengths of seedlings grown at two different temperatures (28 °C/20 °C). b, c qRT-PCR analyses of YUC8 and ATHB2 expression. Seedlings were grown in 12 h light/12 h dark cycles (12 L:12D) at 20 °C for 4 days and transferred under the continuous white light on 5th day. The growth temperature was increased to 28 °C or kept at 20 °C for 4 h at Zeitgeber Time (ZT) 20-24 before harvesting for total RNA extraction. Gene expression levels were normalized to APX3 and presented as values relative to that of wild type (WT) kept at 20 °C. Error bars indicate s.d. (n = 3). The P-value for the interaction (genotype x temperature) is shown at the top. d Hypocotyl lengths of WT, pifq, and ML1p::PIF4mi seedlings grown under continuous white light at 20 °C for 7 days or 20 °C for 4 days followed by 28 °C for 3 days. Error bars indicate s.d. (n = 15 plants). The P-value for the interaction (genotype x temperature) is shown at the top. e, f Hypocotyl lengths of WT, ML1p::PIF4ΔN-SRDX, and SUC2p::PIF4ΔN-SRDX seedlings grown under continuous white light at 20 °C for 7 days or 20 °C for 4 days followed by 28 °C for 3 days. Representative seedlings are shown in e and error bars in f indicate s.d. (n = 15 plants). The P-value for the interaction (genotype x temperature) is shown at the top. g, h qRT-PCR analyses of YUC8 and IAA19. WT and transgenic seedlings were grown in 12 h light/12 h dark cycles (12 L:12D) at 20 °C for 4 days and transferred under the continuous white light on 5th day. The growth temperature was increased to 28 °C or kept at 20 °C for 4 h at ZT20-24 before harvesting for total RNA extraction. Gene expression levels were normalized to APX3 and presented as values relative to that of wild type (WT) kept at 20 °C. Error bars indicate s.d. (n = 3). The P-values for the interaction (genotype x temperature) are shown at the top. Source data are provided as a Source Data file.
levels of these same genes in ML1p::PIF4pifq plants grown at 20 °C showed a marked increase, to similar levels observed in the wild-type plants exposed to 28 °C (Fig. 3b, c, Supplementary Fig. 4), indicating that epidermal PIF4 activates auxin responses, even at 20 °C. While ATHB2 expression in ML1p::PIF4pifq plants was further increased by exposure to high temperature, the expression of YUC8 and IAA19 was not significantly affected (Fig. 3b, c, Supplementary Fig. 4). The additional increase in ATHB2 expression by high temperature may account for the longer hypocotyls of ML1p::PIF4pifq seedlings grown at 28 °C compared to those grown at 20 °C (Fig. 3a). In contrast to ML1p::PIF4pifq plants, YUC8, IAA19, and ATHB2 expression was not induced by exposure to high temperature in SUC2p::PIF4pifq plants, similar to the pifq mutant (Fig. 3b, c, Supplementary Fig. 4) and consistent with their thermo-insensitive hypocotyl growth phenotype (Fig. 3a). Together, these results show that epidermal, but not the vascular, PIF4 activates auxin responses, thereby triggering thermomorphogenesis.

To further confirm that thermomorphogenesis is dependent on epidermal PIF4 activity, we generated transgenic plants expressing an epidermis-specific PIF4 artificial microRNA (ML1p::PIF4mi) and examined thermoresponsive hypocotyl growth in the transgenic plants. We found that PIF4 mRNA expression was reduced by approximately half compared to that in wild-type plants (Supplementary Fig. 5), likely due to the PIF4mi-induced reduction in epidermal PIF4 mRNA levels. The hypocotyl growth of two independent ML1p::PIF4mi plants was insensitive to the high-temperature exposure (Fig. 3d), similar to the pifq mutant, supporting that epidermal PIF4 activity is required for high temperature-induced hypocotyl growth. However, it is possible that the small microRNAs may have moved to neighboring cells from the cells where the microRNAs were initially biosynthesized; therefore, the thermo-insensitive hypocotyl growth seen in ML1p::PIF4mi plants could be attributable to reduced PIF4 levels in neighboring tissues other than the epidermis.

To exclude this possibility, we utilized PIF4, with the N-terminal deleted, fused to the EAR repression domain (PIF4AN-SRDX) to inhibit PIF4 activity in a tissue-specific manner. PIF4AN fused to the strong transcriptional repressor SRDX is expected to inhibit endogenous PIF4 activity by competitively binding to the promoters of PIF4 target genes and repressing their expression. Deletion of the PIF4 N-terminus, that includes the active phytochrome binding motif (APB), was shown to increase the PIF4 stability under light-enhancing the negative effects of PIF4-SRDX on endogenous PIF4 activity. The PIF4AN-SRDX chimera driven by the UBOQ10 promoter (UBOQ10::PIF4AN-SRDX), which is strongly active in all aerial tissues, greatly suppressed hypocotyl growth at high temperatures (Supplementary Fig. 6a, b), indicating that the PIF4AN-SRDX chimera represses endogenous PIF4 activity, as expected. To repress PIF4 activity only in the epidermis and vasculature, we expressed the PIF4AN-SRDX protein under the control of the ML1 and SUC2 promoters, respectively (ML1p::PIF4AN-SRDX and SUC2p::PIF4AN-SRDX). In agreement with the results obtained from the analyses of ML1p::PIF4mi, hypocotyl growth in ML1p::PIF4AN-SRDX transgenic plants was not significantly enhanced by high temperatures (Fig. 3e, f). In contrast, SUC2 promoter-driven PIF4AN-SRDX had no significant effect on high temperature-induced hypocotyl growth (Fig. 3e, f). Consistent with thermo-insensitive hypocotyl growth, the expression of the PIF4 target genes YUC8 and IAA19 was not induced by exposure to high temperature in ML1p::PIF4AN-SRDX transgenic plants, in contrast to that observed in wild-type and SUC2p::PIF4AN-SRDX plants (Fig. 3g, h). Together, these results demonstrate that epidermal, but not vascular, PIF4 regulates thermoresponsive hypocotyl growth.

Epidermal auxin regulates thermoresponsive hypocotyl growth. We next tested whether epidermal PIF4-activated auxin responses are essential for high temperature-induced hypocotyl growth. We generated transgenic plants expressing a stabilized mutant form of IAA3/SHY2 (shy2–2) under the control of the ML1 promoter, in which the epidermal expression of shy2–2 blocks the transcriptional responses to auxin in the epidermis and analyzed thermoresponsive hypocotyl growth. As shown in Fig. 4a, blocking auxin responses in the epidermis significantly reduced the hypocotyl growth response to an elevation in ambient temperature, indicating that epidermal auxin responses are required for high temperature-induced hypocotyl elongation. Consistent with our observation, a previous study showed that epidermal expression of a stabilized mutant version of IAA17 protein suppressed the hypocotyl growth response to both shade and high temperatures. These results suggest that the environmental regulation of hypocotyl growth may be determined by epidermal auxin responses.

Epidermal phyB inhibits temperature-induced hypocotyl growth. The light-activated phyB represses PIF4 activity through both the proteasome-mediated degradation of PIF4 and the prevention of PIF4 from binding to target genes. It was previously shown that epidermal phyB represses hypocotyl growth. Given our results indicating that epidermal PIF4 promotes hypocotyl growth, it is highly likely that epidermal phyB impacts on hypocotyl growth by repressing the epidermal PIF4 activity. If this is the case, the overexpression of epidermal phyB suppresses thermomorphogenesis. To test this hypothesis, we generated transgenic plants expressing PHYB under control of the ML1 promoter in phyB mutant background (phyb-9). Total PHYB levels were higher in ML1p::PHYB;phyb-9 than those in the wild type (Supplementary Fig. 7), implicating that epidermal phyB is overexpressed in ML1p::PHYB;phyb-9. As expected, while the hypocotyl lengths in wild-type seedlings were increased more than two times by high temperature, those in ML1p::PHYB;phyb-9 transgenic plants were only marginally increased (Fig. 4b). Consistent with the defect in thermoresponsive hypocotyl growth, the expression of PIF4 target gene (YUC8) was largely repressed and insensitive to exposure to high temperature in ML1p::PHYB;phyb-9 transgenic plants (Fig. 4c). In contrast, the expression of PIF4 was similarly induced by high-temperature exposure in wild type and ML1p::PHYB;phyb-9 transgenic plants (Fig. 4d). Taken together, these results suggest that the overexpressed epidermal phyB suppresses thermomorphogenesis through the inactivation of epidermal PIF4 post-transcriptionally, probably in a cell-autonomous manner.

Role of epidermal and vascular PIF4 in skotomorphogenesis. In the dark, PIF4, together with three additional PIFs (PIF1, PIF3, and PIF5), promotes skotomorphogenesis including hypocotyl growth, a closed cotyledon, and hypocotyl negative gravitropism. To examine whether epidermal and vascular PIF4 are also involved in these developmental processes, we examined hypocotyl length, cotyledon opening, and hypocotyl negative gravitropism in ML1p::PIF4pifq and SUC2p::PIF4pifq transgenic plants in the dark. Surprisingly, and in contrast to the light conditions, the hypocotyl lengths of both ML1p::PIF4pifq and SUC2p::PIF4pifq transgenic plants were shorter than those of wild-type plants, similar to the pifq mutant (Fig. 5a). This indicates that epidermal or vascular expression of PIF4 is not sufficient to promote hypocotyl growth in etiolated seedlings. This also indicates that the mechanisms underlying PIF-mediated hypocotyl elongation are different under light and dark conditions. While both wild type and ML1p::PIF4pifq plants had closed cotyledons,
SUC2p::PIF4:pifq and pifq plants had opened cotyledons (Fig. 5b), showing that epidermal PIF4 is involved in inhibiting the cotyledon opening of etiolated seedlings. Similarly to hypocotyl elongation, both epidermal and vascular PIF4 failed to rescue the hypocotyl negative gravitropism of pifq mutants (Fig. 5c), consistent with previous studies showing that endodermal PIFs mediate negative hypocotyl gravitropism37,38.

**Vascular PIF4 plays a dominant role in flowering.** PIF4 is also involved in the thermosensory flowering pathway by directly activating the expression of FLOWERING LOCUS T (FT) at high temperatures. It has previously been reported that PIF4 in the vasculature is capable of promoting flowering38,39. To ascertain whether epidermal PIF4 is also able to induce flowering, we measured the flowering time in ML1p::PIF4:pifq and SUC2p::PIF4:pifq plants under a non-inductive photoperiod (8-h light: 16-h dark) and high temperature (27 °C) condition, where PIF4 is known to promote flowering27. Consistent with the previous reports38,39, SUC2 promoter-driven PIF4 expression resulted in very early flowering, whereas epidermal PIF4 expression only slightly accelerated flowering compared to that in wild-type plants (Supplementary Fig. 8a), indicating that vascular PIF4 plays a dominant role in the promotion of flowering. To further verify this result, we determined the flowering time in ML1p::PIF4ΔN-SRDX and SUC2p::PIF4ΔN-SRDX transgenic plants under an inductive photoperiod (16 h-light: 8 h-dark). As shown in Supplementary Fig. 8b, while inactivation of PIF4 specifically in the vascular tissues (SUC2p::PIF4ΔN-SRDX) greatly delayed flowering, inactivation of PIF4 in the epidermis (ML1p::PIF4ΔN-SRDX) did not significantly affect flowering time. Consistent with the late-flowering phenotypes, FT expression levels showed a dramatic reduction in SUC2p::PIF4ΔN-SRDX plants (Supplementary Fig. 8c). Although the flowering time was not significantly affected by the epidermal inactivation of PIF4 under long days, the FT expression levels in ML1p::PIF4ΔN-SRDX plants were approximately half that seen in wild-type plants (Supplementary Fig. 8c). This indicates that epidermal PIF4 may also affect flowering time under certain
conditions, although to a much lesser extent than that by vascular PIF4. Taken together, these results show that vascular PIF4 has a greater effect on the regulation of flowering than that by epidermal PIF4.

Epidermal PIF4 DNA-binding is enhanced by high temperatures. The stability of the PIF4 protein is known to be regulated by light. Upon irradiation, light-activated phytochromes interact directly with PIF4, and the interaction induces PIF4 phosphorylation. The phosphorylated form of PIF4 is then ubiquitinated and degraded via the 26S proteasome pathway. To examine whether the stability of PIF4 in the epidermis is degraded by light-activated phytochrome signals, we determined PIF4 protein levels in ML1p::PIF4;pifq transgenic plants grown in the dark for 5 days or irradiated by red light for 2 h after the 5-day incubation in the dark. Figure 6a shows that the levels of PIF4-YFP were significantly reduced after 2 h of red-light irradiation, indicating that light induces the degradation of PIF4 in the epidermis. Consistent with this result, the levels of PIF4-YFP were increased when the light-grown ML1p::PIF4;pifq plants were transferred to darkness (Supplementary Fig. 9). It was recently reported that epidermal phyB is sufficient for many light-mediated responses, including photomorphogenesis. Thus, it is highly likely that epidermal phyB mediates light-induced PIF4 degradation in a cell-autonomous manner.

Our genetic analyses showed that epidermal PIF4 controls thermoresponsive hypocotyl growth, suggesting that the epidermal PIF4 level or activity increases in response to a rise in ambient temperature to promote thermomorphogenesis. To test whether this is indeed the case, we first determined whether the level of epidermal PIF4 driven by the ML1 promoter (ML1p::PIF4) is affected by ambient temperature changes. The PIF4-YFP protein levels in ML1p::PIF4-YFP;pifq plants were not significantly altered either by 6 h or 24 h of high-temperature exposure under light (Fig. 6b). Similarly, the dark stabilization of epidermal PIF4 was not accelerated at high temperatures (Supplementary Fig. 9). These results suggest that epidermal PIF4 protein stability may not be affected by high temperatures.

Next, we determined whether the DNA-binding of epidermal PIF4 is altered by changes in ambient temperatures. A temperature shift from 16 °C to 28 °C was used because the difference in YUC8 expression was significant between these two temperature conditions (Supplementary Fig. 10). Chromatin immunoprecipitation (ChIP) assays confirmed that epidermal PIF4 directly binds to the promoters of YUC8, IAA19, and ATHB2 (Fig. 6c), which is consistent with the activation of these genes by epidermal PIF4 (Fig. 3b, c). The binding of epidermal
PIF4 to the promoters of these genes was significantly increased by a temperature shift from 16 °C to 28 °C (Fig. 6c). Given that the levels of epidermal PIF4 were not significantly different between the two conditions (Supplementary Fig. 11), the ChIP results indicate that the DNA-binding ability of epidermal PIF4 is enhanced by high temperatures.

**Epidermal PIF4 expression is induced by high temperatures.**
Since PIF4 mRNA is transcriptionally induced at high temperatures\(^1,5\), it is probable that epidermal PIF4 expression is also increased by high temperatures. To test this hypothesis, we first compared the tissue-specific GUS signals in PIF4p::GUS plants grown either at 16 °C for 5 days or grown at 16 °C and transferred to 28 °C for 24 h. Consistent with previous studies, the GUS signal in PIF4p::GUS plants showed a significant overall increase after a 24-h exposure to high temperatures (Fig. 6d). A cotyledon cross-section showed that PIF4p::GUS expression was limited mainly to the vasculature and was weak in the epidermis and mesophyll of the plants grown at 16 °C, but PIF4p::GUS expression showed an increase both in the epidermis and mesophyll with high-temperature exposure (Fig. 6e). To further confirm that epidermal PIF4 expression is increased at high temperatures, we isolated the epidermal tissues from the cotyledons of plants grown either at 20 °C or transferred to 28 °C for 24 h. The PIF4 mRNA levels in the epidermal tissues showed a five-fold increase with high-temperature exposure (Fig. 6f). In addition, the PIF4 target genes YUC8 and IAA29 were also activated by high temperatures in the epidermis (Fig. 6g, h). We also determined the level of epidermal PIF4 protein regulated by its own promoter (PIF4p::PIF4-Myc) in the isolated epidermis. In accordance with the induction of epidermal PIF4 mRNA, the

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**Fig. 6** Both epidermal PIF4 DNA-binding and epidermal PIF4 mRNA expression is increased at high temperatures. **a** Red light induces the degradation of epidermal PIF4 protein. ML1p::PIF4-YFP;pifq seedlings were grown in the dark and transferred under red light for 2 h before harvesting. Immunoblotting was probed using an anti-GFP antibody. Ponceau S staining is shown for equal loading. **b** High temperature effects on the stability of epidermal PIF4 protein. ML1p::PIF4-YFP;pifq seedlings were grown under white light constitutively at 20 °C or grown at 20 °C and shifted to 28 °C for 6 or 24 h. Immunoblotting was probed using an anti-GFP antibody. Ponceau S staining is shown for equal loading. **c** ChIP assays showing epidermal PIF4 DNA-binding is increased by high temperatures. WT and ML1p::PIF4;pifq seedlings kept at 16 °C or shifted to 28 °C for 24 h were used for ChIP assays. Enrichment of DNA was calculated as the ratio between ML1p::PIF4;pifq and WT control, normalized to that of the rDNA gene as an internal reference. Error bars indicate s.d. (n = 3). **P < 0.01 and *P < 0.05** (two-tailed Student’s t-test). **d** The effects of high temperatures on the expression of PIF4p::GUS. PIF4p::GUS seedlings were grown under white light constitutively at 16 °C or grown at 16 °C and shifted to 28 °C for 24 h. Cross-sections of cotyledons of seedlings are shown in **e**. Scale bars = 50 μm. **f**–**h** The expression of PIF4 and its target genes is increased by high temperatures in the epidermis. Wild-type seedlings were grown 20 °C or were shifted to 28 °C for 24 h. Total RNA was extracted from the isolated epidermal tissues. Gene expression levels were normalized to APX3 and presented as values relative to that of wild type at 20 °C. Error bars indicate s.d. (n = 3). **P < 0.05** and **P < 0.01** (two-tailed Student’s t-test). **i** The level of PIF4-Myc under the control of PIF4 promoter is increased by high temperatures in the epidermis. Epidermal tissues were isolated from PIF4p::PIF4-Myc;pifq seedlings grown at 20 °C or exposed to 28 °C for 24 h. PIF4-Myc protein in the isolated epidermis was determined by immunoblotting using an anti-Myc antibody. Ponceau S staining is shown for equal loading. The experiments (**a**, **b**, and **i**) were repeated at least two times with similar results. Source data are provided as a Source Data file.
level of the epidermal PIF4 protein regulated by the PIF4 promoter was also increased in response to high-temperature exposure (Fig. 6i). These results indicate that epidermal PIF4 protein levels are increased at high temperatures through the transcriptional activation of PIF4 expression.

Discussion

Plant organs are composed of different tissues that are functionally specialized. Therefore, to understand how plants respond to the environmental changes, it is essential to identify the specific functions of each tissue in each environmental response. PIF4 acts as a central signaling hub that integrates multiple environmental signals for growth regulation, and it is expressed in all aerial tissues15. Although vascular PIF4 has previously been shown to regulate flowering38,59, the function of epidermal and mesophyll PIF4 has not yet been determined. In this study, we show that epidermal, but not vascular, PIF4, mediates thermo-responsive hypocotyl growth by characterizing transgenic plants expressing epidermal-specific PIF4 or a dominant negative form of PIF4 (PIF4-SRDX). We also show that high temperatures transcriptionally activate epidermal PIF4 and enhance epidermal PIF4 DNA-binding ability, which results in the activation of auxin-related genes in the epidermis. Taken together, our findings demonstrate that epidermal PIF4 levels and activity are increased at high temperatures, which then activates epidermal auxin responses in a cell-autonomous manner, thereby facilitating thermomorphogenesis in Arabidopsis.

Several studies have demonstrated that the epidermis drives and restricts shoot growth and morphology in plants. Growth-promoting hormones such as BR and auxin have been shown to act in the epidermis to determine hypocotyl growth35,58. Epidermal phyB was recently shown to be necessary for the light regulation of hypocotyl growth40. BR, auxin, and light regulate hypocotyl growth through the modulation of PIF4 activity15. Therefore, our finding that epidermal PIF4 mediates thermo-responsive hypocotyl growth is in accordance with the epidermal growth control model. Gibberelic acids (GA) also regulate hypocotyl growth, but the tissue-specific actions of GA on the regulation of hypocotyl growth have not been defined. Since the interaction between DELLAs and PIF4 mediates the GA regulation of hypocotyl growth34, it is plausible that GA also acts in the epidermis to control hypocotyl growth.

It was previously shown that PIF4 promotes hypocotyl growth partly through the auxin pathway6,7. In agreement with this, our transcriptome analyses show that auxin-responsive genes are highly enriched in epidermal PIF4-activated genes (Fig. 2). qRT-PCR analyses with ML1p::PIF4::pifq and ML1p::PIF4AN-SRDX confirm that epidermal PIF4 activity is essential for the high-temperature activation of auxin biosynthetic and responsive genes (Fig. 3). In addition, the block of the epidermal auxin responses suppressed high temperature-induced hypocotyl growth35 (Fig. 4a). Thus, high temperature appears to activate epidermal auxin biosynthesis and response to trigger thermomorphogenesis including hypocotyl elongation. The high-temperature activation of hypocotyl growth and gene expression was impaired by the overexpression of epidermal phyB (Fig. 4b, c), which could inhibit this process at least in part by directly inhibiting the action of epidermal PIF4. Taken together, these results demonstrate that the epidermal phyB-PIF4-auxin signaling module determines thermomorphogenetic growth.

The epidermal auxin response was recently shown to be necessary for the hypocotyl growth response to shade as well as high temperatures55. In the epidermis, the auxin response was suggested to promote hypocotyl growth partly by activating the brassinosteroid signaling pathway55. However, how shade activates the epidermal auxin and BR pathways remains poorly understood. Shade-induced hypocotyl growth is also mediated by many PIF factors including PIF4, PIF3, and PIF720,59. These PIFs promote hypocotyl growth through auxin and BR pathways. Epidermal phyB was previously shown to be required for the hypocotyl response to shade40. Therefore, it is likely that shade enhances epidermal PIF activities through the inactivation of epidermal phyB, thereby activating epidermal auxin and BR signaling. Given that phyB directly interacts with PIFs and the interaction is required for the inhibition of PIFs20,59, epidermal phyB may regulate the activities of epidermal PIF factors probably in a cell-autonomous manner. Our results show that epidermal phyB inhibits thermorepressive hypocotyl (Fig. 4b). Given that the activity of phyB is regulated by ambient temperatures as well as light, both light and temperature signals are likely to be integrated through epidermal phyB to coordinate plant growth with changes in the environment.

We observed that the epidermal PIF4 protein level was increased by high temperatures when it is regulated by its own promoter, but not by the epidermis-specific ML1 promoter (Fig. 6). These results suggest that the transcriptional regulation of PIF4 is a major mechanism underlying high-temperature induction of the epidermal PIF4 protein. In support of this notion, both the PIF4 promoter-driven GUS signals and the expression of endogenous PIF4 mRNA showed increases in the epidermis in response to high temperatures. Although the ML1 promoter-driven PIF4 protein level was not significantly increased, the DNA-binding ability of epidermal PIF4 in ML1p::PIF4::pifq plants was significantly increased by high temperature (Fig. 6c), suggesting that high temperatures induce thermomorphogenesis by enhancing epidermal PIF4 DNA-binding as well as epidermal PIF4 mRNA expression. Consistent with the enhanced epidermal PIF4 binding to its target promoters, the expression of AThB2 and YUC8 was increased by high temperatures in ML1p::PIF4::pifq (Fig. 3c, Supplementary Fig. 10). The high-temperature inactivation of phyB may be partly responsible for the enhanced epidermal PIF4 DNA-binding because phyB-PIF4 interaction was previously shown to prevent PIF4 from binding to DNA56.

Transcriptional regulation of PIF4 is mediated by the circadian clock. The Evening Complex (EC), consisting of ELF3, ELF4, and LUX, has been reported to mediate the high-temperature activation of PIF4 expression12,13. EC binds directly to the PIF4 promoter and inhibits its expression, but high temperatures reverse this inhibition by preventing EC from binding to the PIF4 promoter through unknown mechanisms11,13,60. In addition to EC, TOC1 and PRR5 also directly repress PIF4 expression in the evening3,61. Whether these factors also mediate the high-temperature activation of PIF4 expression remains to be determined. Our results show that epidermal PIF4 mRNA expression is induced by high temperatures (Fig. 6). It is likely that epidermal ELF3 is directly involved in this process in a cell-autonomous manner. Given that CCA1 directly represses ELF3 expression62, the increased PIF4 expression by epidermal CCA1 overexpression (CER6p::CCA1) (Fig. 2d) suggests that epidermal ELF3 could regulate PIF4 mRNA expression. In support of our hypothesis, epidermal ELF3 driven by the ML1 promoter significantly repressed hypocotyl growth and partially restored the thermosensitivity of PIF4 mRNA expression in the elf3 mutant (Supplementary Fig. 12). Further experiments are required to identify the molecular mechanisms by which epidermal ELF3 activity is regulated by ambient temperatures.

A recent study revealed that PIF7, in addition to PIF4, is required for high temperature-induced hypocotyl growth and gene expression53. The PIF7 protein increases in response to elevated ambient temperatures and directly regulates auxin
biosynthetic and signaling genes\textsuperscript{63}. Interestingly, PIF7 interacts with PIF4 (ref. 63). The combination of this interaction and the phenotypes of the pif4 and the pif7 single mutants suggests that the PIF4/PIF7 heterodimer may regulate thermomorphogenesis. Epidermal PIF4 is required for thermomorphogenesis (Fig. 3), so there is a high likelihood that PIF7 also acts in the epidermis to control the high-temperature responses through heterodimerization with epidermal PIF4.

In conclusion, this study demonstrates the crucial role of epidermal PIF4 in the regulation of thermomorphogenic responses. Collectively, our results suggest that an increase in ambient temperature is perceived in the epidermis probably through phyB and EC, which facilitates the activation of epidermal PIF4. The epidermal PIF4 then induces thermomorphogenesis through the epidermal auxin pathway. A recent study showed that sensing of temperature is perceived in the epidermis probably through phyB, which means that PIF4, possibly the mobile auxin signal, is activated by the epidermal PIF4 in cotyledons. Future investigations are required to identify the mechanisms underlying the high-temperature regulation of epidermal PIF4 protein level and activity.

**Methods**

**Plant materials and growth conditions.** Arabidopsis thaliana plants were grown in a greenhouse with a 16 h/8 h light/dark cycle at 22 °C for general growth and seed harvesting. All the Arabidopsis thaliana plants used in this study were in a Col-0 ecotype background. To generate transgenic plants with tissue-specific PIF4 expression, the MLI (epidermal) and SUC2 (vascular) promoters were amplified using the primer pairs indicated in Supplementary Table 1 and fused to the coding sequence of PIF4. MLI::P::PIF4 and SUC2::P::PIF4 were cloned into a modified 35S::EarleyGate vector (promoterless version of 35S::EarleyGate 101), and transformed into quadruple pif7 mutants. To generate the tissue-specific PIF4AN-SRDX transgenic plants, the MLI and SUC2 promoters were fused to a fragment of PIF4 lacking its N terminus (amino acid 1–96) and fused to SRDX. MLI::P::PIF4AN-SRDX and SUC2::P::PIF4AN-SRDX were cloned into the modified 35S::EarleyGate vector and transformed into Col-0. To generate MLI::P::PIF4mi transgenic plants, we used Web MicroRNA Designer \textsuperscript{35} to design an artificial microRNA sequence (5’-TTAATGAGGATACGTGGC3’-3) targeting PIF4 and transformed the MLI promoter-driven artificial microRNA into Col-0. CEB6::CCA1, SUC2::CCA1, and CAY1::CCA1 mutant seeds were kindly provided by Motomu Endo\textsuperscript{66}, as were PIF4c::GUS seeds\textsuperscript{85}.

**Hypocotyl length measurement.** Seeds sterilized with 70% (v/v) ethanol and 0.1% (v/v) Triton X-100 were plated on MS medium (PhytoTechnology Laboratories) supplemented with 0.7% phytoagar. After three days of incubation at 4 °C, seedlings were irrigated with white light for 6 h to promote germination and then incubated in specific light (light intensity: 40 μmol m\(^{-2}\) s\(^{-1}\) and temperature conditions. Seedlings were photocopied and hypocotyl lengths were measured using ImageJ software (http://rsb.info.nih.gov/ij/).

**Measurement of hypocotyl negative gravitropism.** To measure hypocotyl negative gravitropism, seeds imbibed with 15190 (2016). Total RNA was extracted from seedlings using the MiniBEST Plant RNA Extraction Kit (Takara) according to the manufacturer’s instructions. 1 μg of total RNA was processed for preparing mRNA sequencing library using MGIeasy RNA Directional Library Prep Kit (MGI) according to the manufacturer’s instruction. Sequencing was conducted on the MGSeq system (MGI) with 100 bp paired-end reads. The quantification of transcript expression was performed using Salmon\textsuperscript{67}. Differential expression analysis was performed using DESeq2 software\textsuperscript{86}. Differentially expressed genes between samples were defined based on the criteria (expression fold change > 1.5 and adjusted P-value < 0.05). Gene ontology enrichment analysis was performed using the Plant GeneSet Enrichment Analysis Toolkit (PlantGSEA)\textsuperscript{88}.

**Confocal imaging.** Confocal microscopy was performed using Zeiss LSM 710 confocal microscope with 40× water-immersion objective lens. For imaging, seedlings were grown under continuous white light (11w/m\(^2\), 23 °C) for 4 days on 1/2 Murashige-Skoog medium. After staining the cell walls and nuclei with propidium iodide (PI, diluted to a final concentration of 50 μM with distilled water) for 10 min, seedlings were mounted on a slide glass with distilled water. The emission range of GFP fluorescence was 500 to 540 nm whereas the PI emission range was restricted to 630 to 660 nm in order to minimize the signal overlap between two fluorescence dyes.

**Isolation of epidermal cells.** Epidermal cells were isolated from the seedlings grown in the specific growth conditions as described\textsuperscript{66}. Briefly, Arabidopsis cotyledons were cut with microscissors and dropped in the 1.5 ml EPP tube containing 1 ml of enzyme solution (0.75% (wt/vol) cellulase ‘onozuka’ R-10, 0.25% (wt/vol) macerozyme R-10, 0.4 M mannitol, 8 mM CaCl\(_2\) and 5 mM MES-KOH). The tubes were sonicated gently until the cotyledons become transparent and dissolved. The dissolved cotyledons were spread on a petri dish. The epidermis was carefully separated from the vasculature using a micropipette under a stereomicroscope. Isolated epidermal tissues were frozen in liquid nitrogen and used for total RNA or protein extraction. The isolation processes were carried out at the same temperature as the growth temperature of the sample plants.

**qRT-PCR gene expression analysis.** Total RNA was extracted from seedlings using the TaKaRa MiniBEST Plant RNA Extraction Kit (Takara) according to the manufacturer’s instructions. M-MLV reverse transcriptase (Thermo Fisher Scientific) was used for cDNA synthesis. Quantitative real-time PCR (qRT-PCR) was performed using CFX96 (Bio-Rad) and the EvaGreen master mix (Solgent). Gene expression levels were normalized to that of PP2A and are shown relative to the wild-type expression levels. Gene-specific primers are listed in Supplementary Table 1.

**Chromatin immunoprecipitation (ChIP) assays.** Wild type and MLI::P::PIF4-pik7 seedlings kept at 16 °C or shifted to 28 °C for 24 h were cross-linked in 1% formaldehyde solution for 20 min under vacuum. After cross-linking, seedlings were ground in liquid nitrogen. The cross-linked chromatin complex was resuspended by adding lysis buffer (50 mM Tris-HCl at pH 8.0, 2 mM EDTA, 150 mM NaCl and 0.5% Triton X-100, 0.1% Na deoxycholate and 0.1% SDS) and sheared by sonication to reduce the average DNA fragment size to around 0.3 to 0.5 kb. The fragmented chromatin complex was then immunoprecipitated by anti-GFP antibody (GFP-Tag (A-11122) rabbit polyclonal antibody; Thermo Fisher Scientific). The beads were washed with a low-salt buffer (50 mM Tris-HCl at pH 8.0, 2 mM EDTA, 150 mM NaCl and 0.5% Triton X-100), high-salt buffer (50 mM Tris-HCl at pH 8.0, 2 mM EDTA, 0.5 M LiCl, 0.5% NP-40 and 0.5% deoxycholate) and TE buffer (10 mM Tris-HCl at pH 8.0 and 1 mM EDTA), and eluted with an elution buffer (1% SDS and 0.1 M NaHCO\(_3\)). After reverse cross-linking, the PIF4–YFP-bound DNA fragments were purified using a PCR purification kit (Macherey-Nagel). Co-immunoprecipitated DNA fragments were quantified by real-time PCR using specific primers (Supplementary Table 1).

**RNA-Seq analysis.** Seedlings were grown under white light for 5 days at 20 °C. Total RNA was extracted from the seedlings using the MiniBEST Plant RNA Extraction Kit (Takara) according to the manufacturer’s instructions. 1 μg of total RNA was processed for preparing mRNA sequencing library using MGIeasy RNA Directional Library Prep Kit (MGI) according to the manufacturer’s instruction. Sequencing was conducted on the MGSeq system (MGI) with 100 bp paired-end reads. The quantification of transcript expression was performed using Salmon\textsuperscript{67}. Differential expression analysis was performed using DESeq2 software\textsuperscript{86}. Differentially expressed genes between samples were defined based on the criteria (expression fold change > 1.5 and adjusted P-value < 0.05). Gene ontology enrichment analysis was performed using the Plant GeneSet Enrichment Analysis Toolkit (PlantGSEA)\textsuperscript{88}.

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

**Data availability**

RNA sequencing data were deposited into the Gene Expression Omnibus database under accession number GSE133564 and the NCBI Sequence Read Archive under accession number SRP212662. Arabidopsis mutants and transgenic lines used in this study are available upon request from the corresponding author. The source data for Figs. 2a-d, 3a-d, 3f-h, 4a-d, 5a, 6a-c and 6d-i and Supplementary Figs. 2 and 9-11 are provided as a Source Data file.

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Author contributions

Sa.K., G.H., Se.K., E.O., and G.C. conceived the study and designed the experiments. Sa.K., G.H., T.N.T., H.K., J.J., Ja.K., and So.K. carried out the experiments. E.O., G.C., and Ju.K. supervised the work. Sa.K., G.H., S.K., T.N.T., H.K., G.C., and E.O. wrote the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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