

# The functional balance between the *WUSCHEL-RELATED HOMEBOX1* gene and the phytohormone auxin is a key factor for cell proliferation in *Arabidopsis* seedlings

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**Abstract** The *WUSCHEL-RELATED HOMEBOX1* (*WOX1*) transcription factor and its homolog *PRESSED FLOWER* (*PRS*) are multifunctional regulators of leaf development that act as transcriptional repressors. These genes promote cell proliferation under certain conditions, but the related molecular mechanisms are not well understood. Here, we present a new function for *WOX1* in cell proliferation. To identify the *WOX1* downstream genes, we performed a microarray analysis of shoot apices of transgenic *Arabidopsis thaliana* lines harboring [*35Sp::WOX1-glucocorticoid receptor (GR)*] in which the *WOX1* function was temporarily enhanced by dexamethasone. The downregulated genes were significantly enriched for the Gene Ontology term “response to auxin stimulus”, whereas the significantly upregulated genes contained auxin transport-associated *PINI* and *AUX1* and the auxin response factor *MP*, which are involved in formation of auxin response maxima. Simultaneous treatments of synthetic auxin and dexamethasone induced the formation of green compact calli and the unorganized proliferation of cells in the hypocotyl. A microarray analysis of *35Sp::WOX1-GR* plants treated with indole-3-acetic acid and dexamethasone revealed that *WOX1* and auxin additively influenced their common downstream genes. Furthermore, in the presence of an auxin-transport inhibitor, cell proliferation during leaf initiation was suppressed in the *prs* mutant but induced in a broad region of the peripheral zone of the shoot apical meristem in the ectopic *WOX1*-expressing line *FILP::WOX1*. Thus, our results clarify the additive effect of *WOX1/PRS* and auxin on their common downstream genes and highlight the importance of the balance between their functions in controlling cell proliferation.

**Key words:** auxin, cell proliferation, microarray, *PRESSED FLOWER (PRS)*, *WUSCHEL-RELATED HOMEBOX1 (WOX1)*.

## Introduction

Plant-specific *WUSCHEL RELATED-HOMEBOX* (*WOX*) family transcription factors play fundamental roles for plant development, such as body axis-dependent patterning and the maintenance of shoot, root and vascular stem cells (van der Graaff et al. 2009). The members of this family diverged during plant evolution and are conserved (Lian et al. 2014; Nardmann et al. 2009). Among *WOX* family members, *WOX1* and its paralog *PRESSED FLOWER (PRS)*; also called *WOX3*, which act as transcriptional repressors (Ikeda et al. 2009; Lin et al. 2013; Zhang et al. 2014), play crucial roles in the

leaf and floral organ development of seed plant species. *WOX1* and *PRS* promote cell proliferation during leaf initiation and leaf lamina growth (Alvarez et al. 2016; Cho et al. 2013; Ishiwata et al. 2013; Lin et al. 2013; McHale 1993; McHale and Marcotrigiano 1998; Nakata et al. 2012; Nakata and Okada, 2013; Nardmann et al. 2004; Scanlon and Freeling 1997; Scanlon et al. 1996; Tadege et al. 2011; Vandenbussche et al. 2009; Zhuang et al. 2012). *WOX1* and *PRS* are specifically expressed at the margins and the region between the adaxial and abaxial domains (hereafter, the middle domain) in leaf primordia and contribute to margin cell differentiation and the maintenance of adaxial-abaxial patterning (Honda et

Abbreviations: *WOX1*, *WUSCHEL-RELATED HOMEBOX1*; *PRS*, *PRESSED FLOWER*; *GR*, *glucocorticoid receptor*; DEX, dexamethasone; CHX, cycloheximide; GO, Gene Ontology; 2,4-D, 2,4-dichlorophenoxyacetic acid; IAA, indole-3-acetic acid; NPA, 1-*N*-naphthylphthalamic acid.

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al. 2018; Ishiwata et al. 2013; Matsumoto and Okada 2001; Nakata and Okada 2012; Nakata and Okada, 2013; Nakata et al. 2012; Nardmann et al. 2004; Tadege et al. 2011; Vandenbussche et al. 2009; Yoshikawa et al. 2016; Zhang et al. 2014; Zhuang et al. 2012).

In grasses, relatively high *WOX1* and *PR5* expression levels lead to the enlargement of leaves and stems, and increased biomass production (Ishiwata et al. 2013; Wang et al. 2017). In addition, overexpression in tobacco caused unorganized cell proliferation to form callus-like tumors (Tadege et al. 2011). However, extremely high expression levels also caused dwarfism in both grasses and eudicot species. In *Arabidopsis thaliana*, the overexpression of *WOX1* inhibits plant growth and negatively affects shoot meristem development (Zhang et al. 2011). The ectopic high expression of *WOX1* by the *FIL* promoter and the constitutive expression of *PR5* caused local abnormal outgrowths, which formed protrusions or ridges, but did not appear to globally increase the biomass (Matsumoto and Okada 2001; Nakata et al. 2012). Unorganized cell proliferation, such as calli formation in tobacco, did not occur when *WOX1* was overexpressed in *A. thaliana*. Such context-dependent phenotypes have made it difficult to determine the roles of *WOX1* and *PR5* in cell proliferation and have delayed the development of industrial applications.

In this study, we aimed to understand the function of *WOX1* in cell proliferation in more detail. To improve our knowledge of events occurring downstream of *WOX1*, we examined gene expression profiles of early responses to the transient activation of *WOX1* by glucocorticoid receptor (GR) and dexamethasone (DEX) in *A. thaliana*. The microarray analysis identified hundreds of *WOX1*-upregulated and downregulated genes. A Gene Ontology (GO) analysis revealed that auxin-responsive genes were significantly enriched among both the upregulated and downregulated genes. We then focused on the relationship between *WOX1* and the auxin pathway and performed further microarray and pharmacological analyses. The data allows a deeper understanding of the link between *WOX1* and the auxin pathway as related to cell proliferation.

## Materials and methods

### Plant materials and growth conditions

The *A. thaliana* accession Columbia-0 was used as wild type. Among the mutants and transgenic lines used in this study, *pr5-2*, *wox1-101*, *pr5-2 wox1-101*, *FILp::WOX1* (Nakata et al. 2012) and *DR5::GFP* (Ottenschläger et al. 2003) have been previously described. The generation of *35Sp::WOX1-GR* and *35Sp::GFP-GR* is described below.

Surface-sterilized seeds were sown and stored at 4°C in the dark for 2–3 day. Plants were grown under continuous white

fluorescent light at 22°C on solid or in liquid media. Solid and liquid media contained 0.5×Murashige and Skoog salts, 1% sucrose and 0.05% MES-KOH (w/v) pH 5.7 with or without 1.2% purified agar, respectively. For pharmacological analyses, the plants were incubated on solid or in liquid medium with or without DEX, 2,4-dichlorophenoxyacetic acid (2,4-D), indole-3-acetic acid (IAA) and 1-*N*-naphthylphthalamic acid (NPA).

### Microarray analyses

For microarray analyses, *35Sp::WOX1-GR* and *35Sp::GFP-GR* were generated using the GR–DEX system that was established in Aoyama and Chua 1997. For the construction of *35Sp::WOX1-GR* and *35Sp::GFP-GR*, a partial DNA fragment of *Rattus norvegicus* GR protein was amplified by PCR and cloned into the multiple cloning site of pBlueScript SKII(+) using the restriction enzymes *Bam*HI and *Kpn*I. Full-length sequences of *WOX1* and *G3GFP* (Kawakami and Watanabe 1997) were amplified by PCR and cloned into pBlueScript SKII(+) containing the GR fragment using the restriction enzymes *Sac*II and *Bam*HI, generating *WOX1-GR* and *GFP-GR*, respectively, DNA fragments. These were transferred separately to pDONR221 (Thermo Fisher Scientific, USA) and to pFAST-G02 by the GATEWAY reaction (Shimada et al. 2010). These constructs were introduced into Columbia-0 by vacuum infiltration using *Agrobacterium tumefaciens* strain ASE. The first generation of transgenic (T1) plants was screened for GFP fluorescent seeds (*35Sp::WOX1-GR* and *35Sp::GFP-GR*) and obtained homozygous T3 lines. Primer sequences are shown in Supplemental Table S1.

For the transient application of DEX and/or IAA before harvesting, plants were grown on solid medium for 6 day and transferred to liquid medium with or without 10 μM DEX, 10 μM cycloheximide (CHX) and 20 μM IAA. DEX, CHX and IAA treatments were started 6, 6 and 3 h, respectively, before sample collection.

For the microarray analysis, RNA was extracted from shoot apices, including leaf primordia, using a Plant RNeasy Mini Kit (QIAGEN, Germany), and their quality was checked using a Bioanalyzer 2100 (Agilent, USA). For both DEX and CHX treatments of *35Sp::WOX1-GR*, sampling was performed twice in two independent lines ( $n=4$ ). For the analysis of *35Sp::GFP-GR* and a combined application of DEX and IAA, sampling was performed from two independent lines ( $n=2$ ). Microarray analyses were performed using an Arabidopsis Whole Genome Microarray 4X44K Ver. 4 and the one-color method (Agilent). The labelled-cRNA preparation, hybridization and scanning were performed by Miltenyi Biotec Inc. (Germany). The signal intensities from the raw data obtained from individual experiments were normalized by dividing them by their median values. For the identification of *WOX1* downstream genes,  $p$ -values, as determined by Student's  $t$ -test, and  $q$ -values of the false discovery rate (FDR), as determined by the Benjamini-Hochberg method, were calculated with Microsoft Excel and the R stats package (<https://stat.ethz.ch/R-manual/R-devel/library/stats/html/00Index.html>). GO analyses were performed

with GOEAST (Zheng and Wang 2008). For the analysis of simultaneous applications of DEX and IAA, genes with a fold change of more than 1.5-fold in both of the independent *35Sp::WOX1-GR* lines were collected as differentially expressed genes.

The effects of interactions between two factors can be estimated using linear modeling (Brady et al. 2015). Therefore, to investigate the contribution of the interaction on individual gene expression levels, the following general linear model was fitted to  $\log_2$ -transformed signal values of the given probe:

$$S \sim \mu + D + I + D:I + \varepsilon \quad (1)$$

where  $S$ ,  $\mu$ ,  $\varepsilon$ ,  $D$ ,  $I$  and  $D:I$  are  $\log_2$ -transformed signal values, intercept, residuals, effects of DEX, IAA, and the interaction between DEX and IAA, respectively. The  $t$ - and  $p$ -values were obtained for DEX, IAA, and DEX:IAA coefficients, and the  $q$ -values were calculated using the Benjamini-Hochberg method. The probes showing signal values less than  $-3$  in all eight samples were eliminated to satisfy the normal distribution assumption required for the linear model.

The graphical presentations of the results were performed with Microsoft Excel, the R gplots package (<https://www.rdocumentation.org/packages/gplots/versions/3.0.1>), the R beeswarm package (<http://www.cbs.dtu.dk/~eklund/beeswarm/>) and the Python Matplotlib pyplot library ([https://matplotlib.org/api/pyplot\\_api.html](https://matplotlib.org/api/pyplot_api.html)). Raw data from the microarray analyses were deposited to the Gene Expression Omnibus (GSE79647: <http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?token=evihssyaddqdfmj&acc=GSE79647>).

### Reverse transcriptase-quantitative PCR (RT-qPCR)

For the RT-qPCR analysis, seedlings of two independent lines were grown on a solid medium for 6 day and were treated transiently with DEX and IAA under the same conditions as for the microarray analyses. Total RNA was extracted from the aerial plant parts using a Plant RNeasy Mini Kit. cDNA was synthesized using a QuantiTect reverse transcription kit (QIAGEN). Quantitative PCR was performed on a 7500 fast Real-Time PCR Systems (Thermo Fisher Scientific) using the QuantiTect SYBR Green PCR kit (QIAGEN). Relative expression levels were analyzed by the comparative CT ( $\Delta\Delta CT$ ) method using *ACT2* as an endogenous control. The experiment was performed with biological triplicates. Primer sequences are shown in Supplemental Table S1.

### Observation and staining

The stereoscopic observation was performed with a LEICA MZ FLIII (Leica, Germany). For scanning electron microscopy, samples frozen in liquid nitrogen for 1 min were observed with a XL30 scanning electron microscope (FEI, USA). For the observation of the DR5::GFP pattern, leaves were stained with  $50 \mu\text{g ml}^{-1}$  FM4-64 (Thermo Fisher Scientific) for 5 min and observed with a TCS SP8 (Leica) equipped with a white light laser and the time-gating detector HyD. For the analysis of

plants treated with DEX and/or 2,4-D, hypocotyls were stained by the modified pseudo-Schiff–propidium iodide staining method (Truernit et al. 2008) with some changes (Nakata et al. 2018), and observed with a LSM710 (Zeiss, Germany). For observation with a confocal laser scanning microscope, GFP, FM4-64 and propidium iodide fluorescence were detected with the excitation/emission values of 488/495–540 nm, 488/580–625 nm and 514/566–719 nm, respectively. Obtained images were processed using ImageJ (<https://imagej.nih.gov/ij/>).

## Results

### Enhancement of the WOX1 function by the GR–DEX system

To confirm the enhancement of the WOX1 function by the GR–DEX system, we investigated the phenotypes of *35Sp::WOX1-GR* transgenic lines treated with DEX. Of 24 independent T2 transformants, 13 lines showed growth defects when grown on the medium containing 125 nM DEX (example shown in Figure 1A). In these lines, the numbers of leaves decreased, and the leaves were narrow. We found adventitious protrusions on the abaxial sides of leaves in 5 of the 13 lines (Figure 1B). The leaf epidermises were covered with elongated oblong cells resembling leaf margin cells (Figure 1C). This phenotype occurs with the ectopic expression of WOX1, i.e. *FILP::WOX1* (Nakata et al. 2012). Using two of these five transgenic lines, we examined the effects of lower concentrations of DEX (5, 10, 25 nM DEX). The lower the DEX concentration, the milder the phenotype (Figure 1D). The relatively weak dwarf phenotype was consistent with the phenotype of *wox1-D* in a previous study (Zhang et al. 2011). We also found the growth inhibition of roots in addition to shoots. No positive effect of WOX1 on biomass was found in the tested concentration range. These phenotypes indicate that the WOX1 function was strongly enhanced by the DEX treatment in these two *35Sp::WOX1-GR* lines.

### Transient enhancement of WOX1 function alters auxin-related gene expression levels

To identify WOX1-downstream genes, we performed a microarray analysis of shoot apices that included leaf primordia harvested immediately after a 5–6-h DEX treatment. WOX1 expression increased more than 60 times in these two lines (Figure 2A). We found that 429 and 614 genes were upregulated and downregulated by the DEX treatment, respectively [Figure 2B; FDR  $< 0.05$ ,  $\log_2$  fold change ( $\log_2 FC$ )  $> 1$ ]. A GO analysis revealed the upregulated genes were enriched with GO terms, including “response to jasmonic acid” (17 genes,  $p$ -value =  $7.33\text{e-}6$ ), “response to wounding” (13 genes,  $p$ -value =  $2.77\text{e-}3$ ), “sterol biosynthetic process” (5 genes,  $p$ -value =  $6.84\text{e-}3$ ) and “response to auxin stimulus” (16 genes,  $p$ -value =  $4.62\text{e-}3$ ). The downregulated genes were

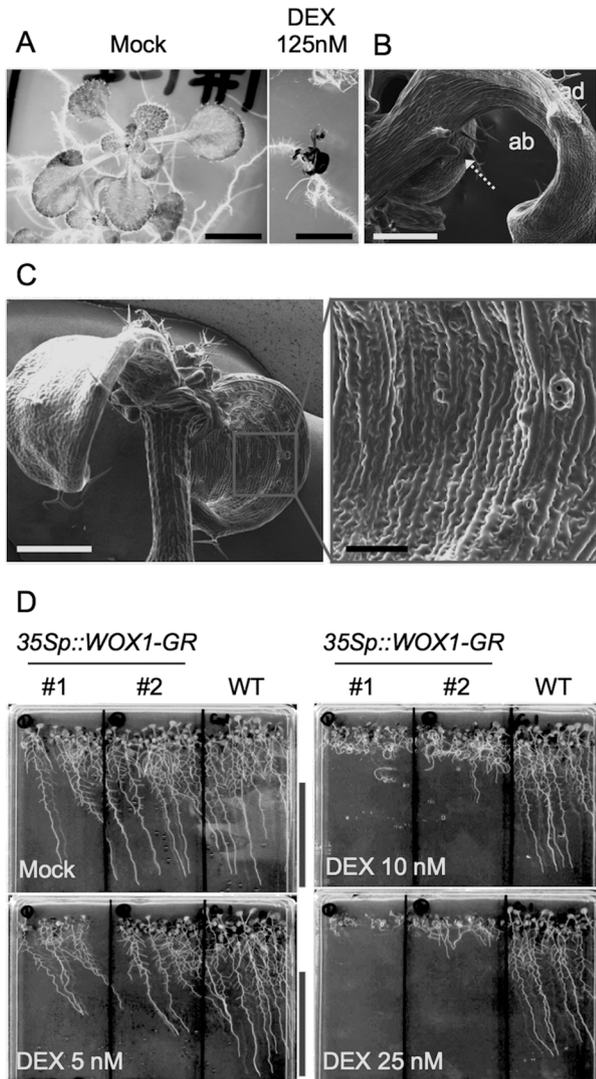


Figure 1. Phenotype of the DEX-treated *35Sp::WOX1-GR* Arabidopsis line. (A) Phenotype of *35Sp::WOX1-GR* plants treated with mock or 125 nM DEX as observed by stereomicroscopy. (B–C) Leaves of *35Sp::WOX1-GR* plants as observed by scanning electron microscopy. The arrow in (B) indicates an abaxial protrusion. “ad” and “ab” in (B) indicate the adaxial and abaxial sides, respectively. Almost all epidermis cells other than stomata have changed to an elongated oblong form. (D) The dose-dependent effects of DEX on the growth of *35Sp::WOX1-GR* plants 13 day after germination. Scale bars, 5 mm (A), 500  $\mu$ m (B, C left), 100  $\mu$ m (C right), 5 cm (D).

enriched with other GO terms, including “response to auxin stimulus” (37 genes,  $p$ -value=3.25e-14) and “regulation of transcription, DNA dependent” (72 genes,  $p$ -value=2.03e-6) (Figure 2B).

To clarify which genes have the potential to be directly affected by WOX1, we performed the same transcriptome analysis in the presence of CHX, which inhibits protein synthesis. Under these conditions, 117 and 380 genes were upregulated and downregulated by the DEX treatment, respectively (Figure 2B; FDR < 0.05,  $\log_2$  FC > 1). Because WOX1 acts as transcriptional repressor (Lin et al. 2013), it is plausible

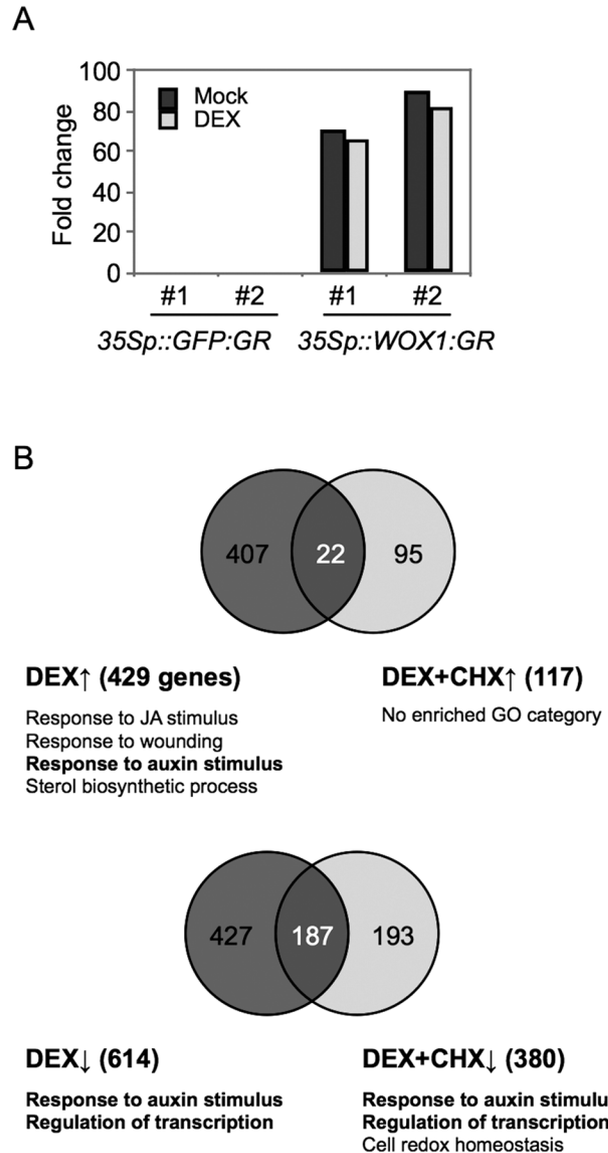


Figure 2. Genes altered in the DEX-treated *35Sp::WOX1-GR* plants with or without CHX. (A) Venn diagram illustrating the numbers of genes upregulated (upward-pointing arrow) and downregulated (downward-pointing arrow) by DEX and/or CHX. The enriched GO categories and the gene numbers are indicated beside the diagram. GO category numbers: GO:0009753 (response to jasmonic acid), GO:0009611 (response to wounding), GO:0009733 (response to auxin stimulus), GO:0016126 (sterol biosynthetic process), GO:0006355 (DNA-dependent regulation of transcription) and GO:0045454 (cell redox homeostasis). (B) A comparison between the expression level of WOX1 in the control (*35Sp::GFP-GR*) and *35Sp::WOX1-GR* plants.

that more genes are downregulated than upregulated. A GO analysis revealed the upregulated genes were not significantly enriched with any GO terms. In contrast, the downregulated genes were enriched with GO terms, including “response to auxin stimulus” (25 genes,  $p$ -value=7.93e-10), “regulation of transcription, DNA dependent” (48 genes,  $p$ -value=6.56e-5) and “cell redox homeostasis” (10 genes,  $p$ -value=3.84e-4). Among genes involved in “response to auxin stimulus”, 14 SAUR genes,

Table 1. “Response to auxin stimulus” genes downregulated by WOX1 with or without CHX.

Description	Gene name	AGI code	Fold change (D/M)* <sup>1</sup>		<i>p</i> -value* <sup>2</sup>	<i>q</i> -value* <sup>3</sup>	Fold change (D/M+C)* <sup>1</sup>		ProbeName
			Mean	SE			Mean	SE	
Aux/IAA	<i>IAA16</i>	AT3G04730	-1.47	0.06	0.000	0.019	-1.06	0.04	A_84_P23996
GH3	<i>DFL2</i>	AT4G03400	-1.10	0.04	0.000	0.018	-1.86	0.14	A_84_P20399
SAUR	<i>SAUR61</i>	AT1G29420	-4.11	0.19	0.000	0.020	-3.22	0.25	A_84_P537376
	<i>SAUR63</i>	AT1G29440	-4.10	0.12	0.000	0.017	-5.28	0.19	A_84_P279980
	<i>SAUR64</i>	AT1G29450	-3.17	0.20	0.001	0.024	-4.25	0.20	A_84_P19713
	<i>SAUR65</i>	AT1G29460	-3.16	0.17	0.001	0.021	-2.14	0.20	A_84_P15930
	<i>SAUR68</i>	AT1G29490	-3.06	0.15	0.000	0.020	-1.15	0.09	A_84_P843096
	<i>SAUR66</i>	AT1G29500	-1.58	0.11	0.001	0.025	-3.26	0.14	A_84_P11207
	<i>SAUR68</i>	AT1G29510	-1.69	0.09	0.000	0.021	-2.67	0.16	A_84_P22555
	<i>SAUR56</i>	AT1G76190	-1.63	0.11	0.001	0.025	-1.56	0.15	A_84_P19956
	<i>SAUR59</i>	AT3G60690	-1.13	0.11	0.003	0.035	-1.19	0.10	A_84_P11862
	<i>SAUR15</i>	AT4G38850	-3.68	0.33	0.002	0.031	-4.53	0.29	A_84_P16734
	<i>AT5G18030</i>		-1.28	0.13	0.004	0.037	-2.42	0.18	A_84_P152798
	<i>SAUR22</i>	AT5G18050	-1.28	0.16	0.006	0.046	-1.60	0.09	A_84_P141269
	<i>SAUR23</i>	AT5G18060	-1.39	0.10	0.001	0.026	-1.56	0.12	A_84_P94979
	<i>SAUR24</i>	AT5G18080	-1.48	0.09	0.001	0.024	-2.12	0.14	A_84_P272980
Others	<i>AIR3</i>	AT2G04160	-2.63	0.20	0.001	0.027	-2.51	0.12	A_84_P93389
	<i>ACS4</i>	AT2G22810	-3.11	0.31	0.003	0.035	-3.33	0.17	A_84_P17344

\*<sup>1</sup> Means of log<sub>2</sub> fold changes for four experiments; D/M indicates DEX/Mock; +C indicates “with CHX”. \*<sup>2</sup> *p*-values as assessed by a two-sided *t*-test. \*<sup>3</sup> False discovery rate as assessed by the Benjamini–Hochberg method.

*IAA16*, *DFL2*, *AIR3* and *ACS4* were downregulated regardless of the presence of CHX (Table 1). Genes involved in “regulation of transcription” contained 10 transcription factor families, including *bHLH*, *NAC*, *ERF/AP2* and *GRAS* (Table 2). Interestingly, five abscisic acid-receptor genes were downregulated by DEX regardless of the presence of CHX (Table 3). Among adaxial- and abaxial-specific genes, *ZPR1* and *ETT* were upregulated, while *AS2* and *YAB5* were significantly downregulated by the DEX treatment (Table 3). However, the changes in the expression levels of these genes, except for *YAB5*, were subtle in the presence of CHX (Table 3). Thus, transcriptional regulatory genes, auxin-responsive genes and abscisic acid-receptor genes were identified as potential direct targets of WOX1 repression.

Because significant numbers of genes categorized into “response to auxin stimulus” were upregulated and downregulated, we examined the expression profiles of genes involved in the auxin pathway (Table 4). Among the auxin biosynthetic genes, *TAA1*, *YUC1*, *YUC4*, *NIT1* and *NIT3* were upregulated, while *TAR2*, *YUC2*, *YUC5*, *NIT4* and *CYP83B1* were downregulated. Among transport-associated genes, *AUX1*, *LAX1* and *PIN1* were upregulated, while *PIN3* and *PIN7* were downregulated. The expression of the activator-type auxin response factor *MP* increased. The changes in all of these genes were statistically significant (Table 4). Regulation on *AUX1*, *LAX1*, *PIN1* and *MP* appeared to be mainly indirect because changes in the expression levels of these four genes are limited to the absence of CHX (Table 4).

To investigate the importance of WOX1 expression in the regulation of auxin transport-associated genes

and *MP* in developing leaves, auxin response patterns in the wild type, *prx wox1* and plants harboring the *FILp::WOX1* construct were examined by confocal laser scanning microscopy using the *DR5::GFP* marker. In the wild-type leaves, spots on the marginal tip showed the maximum fluorescence, but in the leaves of *prx wox1*, the spot pattern disappeared, and the auxin response was detected in a broad region of the leaf base (Figure 3A). In the leaves of the *FILp::WOX1* plants, there was no apparent change in the global pattern of auxin at the margins (Figure 3A), but the spots with the maximum fluorescence were also found at the tips of the ectopically developed abaxial protrusions (Figure 3B; an arrow head). The *DR5::GFP* pattern in *FILp::WOX1* leaves was consistent with the *PIN1p::PIN1-GFP* pattern in the same genotype (Alvarez et al. 2016). Thus, *WOX1* may affect auxin patterning at the leaf margins probably through the regulation of auxin transport.

In conclusion, the microarray data suggested that *WOX1* had bifurcated effects on the auxin pathway: the downregulation of auxin responsive genes and the upregulation of auxin transport-associated genes and *MP*.

### ***The combination of WOX1 and auxin induces unorganized cell proliferation***

To clarify whether the complicated relationship between auxin and *WOX1* is responsible for the phenotypic variation in cell proliferation, we analyzed the phenotypes of plants treated with a synthetic auxin analog, 2,4-D, together with enhanced *WOX1* activity. In the liquid medium, a relatively low 2,4-D concentration alone resulted in the formation of white

Table 2. “Regulation of transcription “ genes downregulated by WOX1 with or without CHX.

Description	Gene name	AGI code	Fold change (D/M) <sup>*1</sup>		<i>p</i> -value <sup>*2</sup>	<i>q</i> -value <sup>*3</sup>	Fold change (D/M+C) <sup>*1</sup>		ProbeName
			Mean	SE			Mean	SE	
bZIP	<i>bZIP3</i>	AT5G15830	-3.34	0.44	0.007	0.048	-4.42	0.29	A_84_P15068
	<i>bZIP44</i>	AT1G75390	-1.46	0.16	0.004	0.038	-1.30	0.12	A_84_P817973
			-1.48	0.13	0.002	0.030	-1.58	0.09	A_84_P817969
			-1.46	0.12	0.002	0.028	-1.52	0.08	A_84_P22038
C2H2	<i>GIS</i>	AT3G58070	-1.85	0.24	0.007	0.048	-2.10	0.10	A_84_P23199
ERF/AP2	<i>ERF106</i>	AT5G07580	-3.11	0.24	0.002	0.027	-2.22	0.16	A_84_P821249
			-3.13	0.25	0.002	0.028	-2.21	0.11	A_84_P12054
	<i>ERF107</i>	AT5G61590	-3.28	0.13	0.000	0.018	-5.80	0.09	A_84_P21661
	<i>RAP2.1</i>	AT1G46768	-1.67	0.06	0.000	0.018	-1.13	0.31	A_84_P108742
			-1.70	0.12	0.001	0.026	-1.56	0.07	A_84_P786987
			-1.80	0.10	0.001	0.021	-1.49	0.08	A_84_P844167
HB	<i>HAT1</i>	AT4G17460	-2.56	0.18	0.001	0.026	-2.51	0.17	A_84_P19589
	<i>HAT22</i>	AT4G37790	-1.51	0.20	0.008	0.049	-1.36	0.05	A_84_P18621
MYB	AT5G56840		-2.10	0.21	0.003	0.035	-1.49	0.09	A_84_P11239
	<i>RL3</i>	AT4G36570	-3.86	0.40	0.004	0.037	-3.01	0.19	A_84_P789861
NAC	<i>NAC020</i>	AT1G54330	-1.08	0.34	0.002	0.032	-1.23	0.25	A_84_P23710
	<i>NAC028</i>	AT1G65910	-1.23	0.10	0.000	0.021	-1.22	0.09	A_84_P11441
	<i>NAC003</i>	AT1G02220	-2.81	0.06	0.006	0.044	-2.17	0.05	A_84_P799508
WRKY	<i>WRKY7</i>	AT4G24240	-1.88	0.09	0.000	0.020	-1.06	0.03	A_84_P10055
GRAS	AT3G49950		-1.75	0.20	0.004	0.040	-1.57	0.03	A_84_P13705
	<i>SCL7</i>	AT3G50650	-2.40	0.13	0.001	0.021	-1.37	0.02	A_84_P17490
	<i>SCL15</i>	AT4G36710	-1.29	0.17	0.007	0.047	-1.16	0.01	A_84_P13917
			-1.22	0.07	0.001	0.023	-1.15	0.04	A_84_P104016
bHLH	<i>AKS2</i>	AT1G05805	-1.22	0.07	0.001	0.023	-1.15	0.04	A_84_P104016
	<i>AIF4</i>	AT1G09250	-1.38	0.13	0.003	0.034	-1.35	0.14	A_84_P820734
			-1.45	0.12	0.002	0.028	-1.32	0.10	A_84_P23695
	<i>bHLH030</i>	AT1G68810	-1.37	0.06	0.000	0.019	-2.10	0.12	A_84_P19990
	<i>bHLH129</i>	AT2G43140	-2.80	0.22	0.002	0.028	-1.82	0.09	A_84_P838362
			-3.21	0.25	0.002	0.028	-2.59	0.08	A_84_P150068
	<i>SPCH</i>	AT5G53210	-1.91	0.15	0.001	0.027	-1.11	0.10	A_84_P576326
	<i>PIF7</i>	AT5G61270	-2.65	0.23	0.002	0.030	-1.58	0.02	A_84_P827848
ZF			-2.85	0.10	0.000	0.018	-1.55	0.02	A_84_P553165
	<i>ZF3</i>	AT5G43170	-2.15	0.11	0.000	0.021	-2.25	0.10	A_84_P21591
			-2.20	0.11	0.000	0.021	-2.20	0.09	A_84_P826434

<sup>\*1</sup> Means of log<sub>2</sub> fold changes for four experiments; D/M indicates DEX/Mock; +C indicates “with CHX”. <sup>\*2</sup> *p*-values as assessed by a two-sided *t*-test. <sup>\*3</sup> False discovery rate as assessed by the Benjamini-Hochberg method.

Table 3. Other genes significantly upregulated or downregulated by WOX1 with DEX.

Description	Gene name	AGI code	Fold change (D/M) <sup>*1</sup>		<i>p</i> -value <sup>*2</sup>	<i>q</i> -value <sup>*3</sup>	Fold change (D/M+C) <sup>*1</sup>		ProbeName
			Mean	SE			Mean	SE	
ABA receptor	<i>PYR1</i>	AT4G17870	-2.44	0.10	0.000	0.019	-1.66	0.15	A_84_P120122
	<i>PYL4</i>	AT2G38310	-2.74	0.09	0.000	0.018	-2.62	0.09	A_84_P197484
	<i>PYL5</i>	AT5G05440	-1.94	0.15	0.001	0.027	-1.95	0.08	A_84_P146199
			-1.89	0.14	0.001	0.027	-1.97	0.11	A_84_P817892
			-1.85	0.08	0.000	0.019	-1.98	0.11	A_84_P852047
		<i>PYL6</i>	AT2G40330	-4.35	0.25	0.001	0.022	-3.69	0.11
Adaxial gene	<i>RCAR3</i>	AT5G53160	-1.36	0.06	0.000	0.020	-1.02	0.01	A_84_P94069
	<i>AS2</i>	AT1G65620	-1.14	0.08	0.001	0.025	-0.52	0.09	A_84_P15163
	<i>ZPRI</i>	AT2G45450	1.26	0.10	0.001	0.027	0.41	0.07	A_84_P62200
Abaxial gene	<i>YAB5</i>	AT2G26580	-1.15	0.06	0.000	0.020	-0.83	0.06	A_84_P516292
			-1.14	0.10	0.002	0.031	-0.94	0.06	A_84_P788325
	<i>ETT</i>	AT2G33860	1.31	0.06	0.000	0.019	0.43	0.13	A_84_P18296

<sup>\*1</sup> Means of log<sub>2</sub> fold changes for four experiments; D/M indicates DEX/Mock; +C indicates “with CHX”. <sup>\*2</sup> *p*-values as assessed by a two-sided *t*-test. <sup>\*3</sup> False discovery rate as assessed by the Benjamini-Hochberg method.

Table 4. Expression changes in auxin-pathway genes significantly upregulated or downregulated by WOX1 (FDR &lt;0.05).

Description	Gene name	AGI code	Fold change (D/M) <sup>*1</sup>		<i>p</i> -value <sup>*2</sup>	<i>q</i> -value <sup>*3</sup>	Fold change (D/M+C) <sup>*1</sup>		ProbeName
			Mean	SE			Mean	SE	
Biosynthesis	<i>TAA1</i>	AT1G70560	<b>1.03</b>	0.06	0.001	0.024	0.10	0.05	A_84_P19052
	<i>TAR2</i>	AT4G24670	<b>-1.41</b>	0.11	0.001	0.027	-0.70	0.12	A_84_P10057
	<i>YUC1</i>	AT4G32540	0.50	0.02	0.000	0.020	0.02	0.08	A_84_P22386
	<i>YUC2</i>	AT4G13260	<b>-2.01</b>	0.17	0.002	0.030	-0.91	0.37	A_84_P16644
	<i>YUC4</i>	AT5G11320	0.77	0.08	0.004	0.038	-0.10	0.47	A_84_P767924
			0.76	0.05	0.001	0.025	-0.59	0.07	A_84_P18684
	<i>YUC5</i>	AT5G43890	<b>-3.74</b>	0.15	0.000	0.019	<b>-3.03</b>	0.23	A_84_P23485
	<i>NIT1</i>	AT3G44310	0.71	0.05	0.001	0.025	0.22	0.13	A_84_P808866
			0.64	0.07	0.004	0.037	0.15	0.09	A_84_P21247
	<i>NIT3</i>	AT3G44320	<b>1.74</b>	0.11	0.001	0.025	1.23	0.06	A_84_P22193
	<i>NIT4</i>	AT5G22300	<b>-1.42</b>	0.09	0.001	0.024	<b>-1.21</b>	0.16	A_84_P10196
	<i>CYP83B1</i>	AT4G31500	-0.45	0.04	0.002	0.028	-0.13	0.06	A_84_P16706
	Transport	<i>AUX1</i>	AT2G38120	0.74	0.09	0.005	0.041	0.12	0.05
<i>LAX1</i>		AT5G01240	0.75	0.03	0.000	0.019	0.28	0.10	A_84_P859674
			0.94	0.10	0.003	0.036	0.45	0.05	A_84_P20541
<i>PIN1</i>		AT1G73590	<b>1.03</b>	0.08	0.001	0.027	0.10	0.05	A_84_P11175
<i>PIN3</i>		AT1G70940	-0.50	0.06	0.005	0.041	-0.19	0.06	A_84_P13326
<i>PIN7</i>		AT1G23080	<b>-1.08</b>	0.08	0.001	0.027	-0.69	0.25	A_84_P67534
Activator ARF		<i>MP</i>	AT1G19850	<b>1.42</b>	0.04	0.000	0.017	0.23	0.06
Repressor ARF	<i>ARF1</i>	AT1G59750	0.16	0.00	0.000	0.017	0.01	0.01	A_84_P18037
	<i>ARF6</i>	AT1G30330	-0.29	0.03	0.005	0.041	-0.37	0.06	A_84_P21873
	<i>ARF16</i>	AT4G30080	0.36	0.01	0.000	0.017	0.29	0.06	A_84_P101526

<sup>\*1</sup> Means of log<sub>2</sub> fold changes for four experiments; D/M indicates DEX/Mock; +C indicates "with CHX". <sup>\*2</sup> *p*-values as assessed by a two-sided *t*-test. <sup>\*3</sup> False discovery rate as assessed by the Benjamini-Hochberg method.

and friable tissues on hypocotyls (Figure 4A, a white arrow), while green compact calli were formed on hypocotyls by the simultaneous application of DEX with a relatively low 2,4-D concentration (Figure 4A). The higher the concentration of 2,4-D was, the slower the cell proliferation was, regardless of the presence of DEX (Figure 4A). Green compact calli on hypocotyls were also formed when plants were grown on the solid medium containing both DEX and 2,4-D (Figure 4B). In the green compact calli, unorganized cell proliferation, with random division planes, was observed (Figure 4B). This phenotype was clearly different from the root tissue-like layered structures formed on hypocotyls treated with 2,4-D alone (Figure 4B). Thus, the combination of WOX1 and auxin stimulated unorganized cell proliferation but the balance between the functions of the two factors is important in determining the rate of cell proliferation and the characteristics of the cells.

#### ***WOX1 and auxin additively influence the expression of their common downstream genes***

To investigate the effects of simultaneous auxin and DEX treatments on gene expression profiles, we performed microarray analyses (Figure 5). As shown in Figure 5A, 166 genes were upregulated commonly by DEX and by IAA (DEX-up-IAA-up; Supplemental Table S2), 87 genes were upregulated by DEX but downregulated by IAA (DEX-up-IAA-down; Supplemental Table S3), 108 genes were downregulated by DEX but upregulated by IAA

(DEX-down-IAA-up; Supplemental Table S4), and 272 genes were downregulated by both DEX and IAA (DEX-down-IAA-down; Supplemental Table S5). *MP*, *PIN1* and *AUX1* were found in DEX-up-IAA-up. DEX-up-IAA-down included *TAA1* and regulators of shoot apical meristem/leaf development, including *CUC2*, *CUC3* and *KLU*. The *Aux/IAA* genes, the *SAUR* genes, *ARGOS* and *CKX1*, which is a member of the *CKX* genes involved in cytokinin degradation, were included in DEX-down-IAA-up, while *CKX7* and *ARR7*, which is a negative regulator of cytokinin signaling, and the abscisic acid-receptor genes (*PYR1*, *PYL4-6* and *RCAR3*) were found in DEX-down-IAA-down (Supplemental Table S2).

To examine the global interaction between WOX1 and IAA, the expression levels of both genes under each condition were fitted to the statistical model. The density of genes potentially affected by a single treatment of DEX or IAA ( $p < 0.05$ ) was much higher than the average, while the density affected depending on the interaction between DEX and IAA was lower (Figure 5B). Based on these results, even though >600 genes were affected by both DEX and IAA, the effects of DEX and IAA are likely to be additive at the transcriptome level. In particular, in both the DEX-up-IAA-down and DEX-down-IAA-up categories, changes in gene expression by the simultaneous DEX plus IAA treatment (IAA+DEX) were intermediate between those of single DEX and IAA treatments (Figure 5C). As with the global trend, changes in the expression levels of regulators of shoot

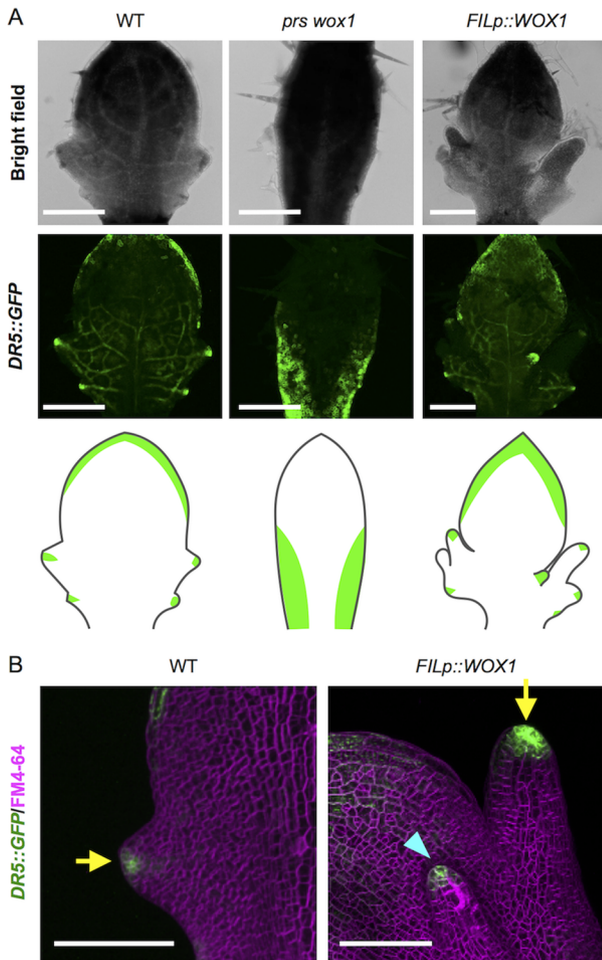


Figure 3. DR5::GFP fluorescent pattern in *prs wox1* and the *FILp::WOX1* plants. (A) Confocal laser scanning microscope images of developing leaves. The bottom columns show the leaf shape and the GFP pattern in the individual lines. (B) The GFP spots on a growing tip. The arrows indicate the marginal tip, and the arrow head indicates an abnormal abaxial tip. Scale bars, 200  $\mu\text{m}$  (A), 100  $\mu\text{m}$  (B).

apical meristem/leaf development that were categorized into DEX-up-IAA-down in the presence of IAA+DEX were offset between the changes after treatments with IAA and DEX alone (Figure 5D). *MP*, *PIN1* and *AUX1* were further upregulated by IAA+DEX (Figure 5E). The expression changes of *CUC2*, *CUC3*, *MP* and *PIN1* were confirmed by RT-qPCR (Figure 5F). These expression analyses indicate that WOX1 and auxin additively control the expression of several genes responsible for leaf development.

#### The WOX1/PRS is responsible for leaf initiation when normal auxin patterning is disrupted

The phenotype of *35Sp::WOX1-GR* simultaneously treated with DEX and 2,4-D suggested that the balance between the auxin and WOX1 functions is important for unorganized cell proliferation. To clarify whether the function of WOX1/PRS in leaf development is also closely related to that of the auxin pathway, we disrupted

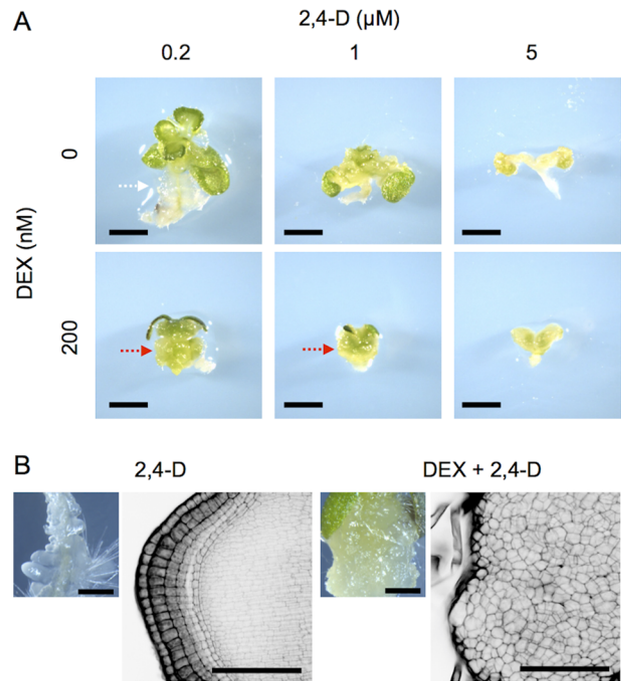


Figure 4. Effects of a simultaneous application of DEX and 2,4-D on Arabidopsis seedling phenotypes. (A) The *35Sp::WOX1-GR* plants grown on liquid media containing various concentrations of DEX and/or 2,4-D for three weeks. The relatively low concentration of 2,4-D led to white friable tissue formation (white arrow) in the absence of DEX but green compact calli in the presence of DEX (red arrows). (B) The *35Sp::WOX1-GR* plants grown on solid media containing 125 nM DEX and/or 1  $\mu\text{M}$  2,4-D for 12 day. The left panels are stereomicroscopic images, and the right ones are black-and-white reversal images observed with confocal laser scanning microscope after modified pseudo-Schiff–propidium iodide staining. Scale bars, 2 mm (A), 0.5 mm (B, left panels), 100  $\mu\text{m}$  (B, right panels).

the auxin pattern in seedlings of loss-of-function mutants by treatment with NPA, an inhibitor of polar auxin transport (Thomson et al. 1973) and analyzed the phenotype. In *A. thaliana*, leaf initiation phenotypes of all *prs*, *wox1* and *prs wox1* mutants had no, or only slight, phenotypic differences from the wild type in the absence of NPA (Nakata et al. 2012). NPA caused a severe inhibition of leaf initiation in the *prs* mutant background unlike the wild type (Figure 6A, B). The *wox1* mutation did not have any effect (Figure 6A). The phenotype of *prs* indicated that *PRS* plays an important role in leaf initiation when the normal auxin pattern is disturbed. For further characterization, we examined the phenotype of the *FILp::WOX1* plants treated with NPA. These seedlings frequently formed fused leaves or cup-shaped leaves in the presence of 10  $\mu\text{M}$  NPA (Figure 6C). Fused or cup-shaped leaves were not formed in the wild-type seedling treated with 10  $\mu\text{M}$  NPA (Figure 6C). The similar cup-shaped leaf phenotype was also observed in the 2,4-D-treated *FILp::WOX1* plants (Figure 6D). The leaf primordia are initiated by cell proliferation along periclinal division planes from the peripheral region of the shoot apical meristem. Thus, the cup-

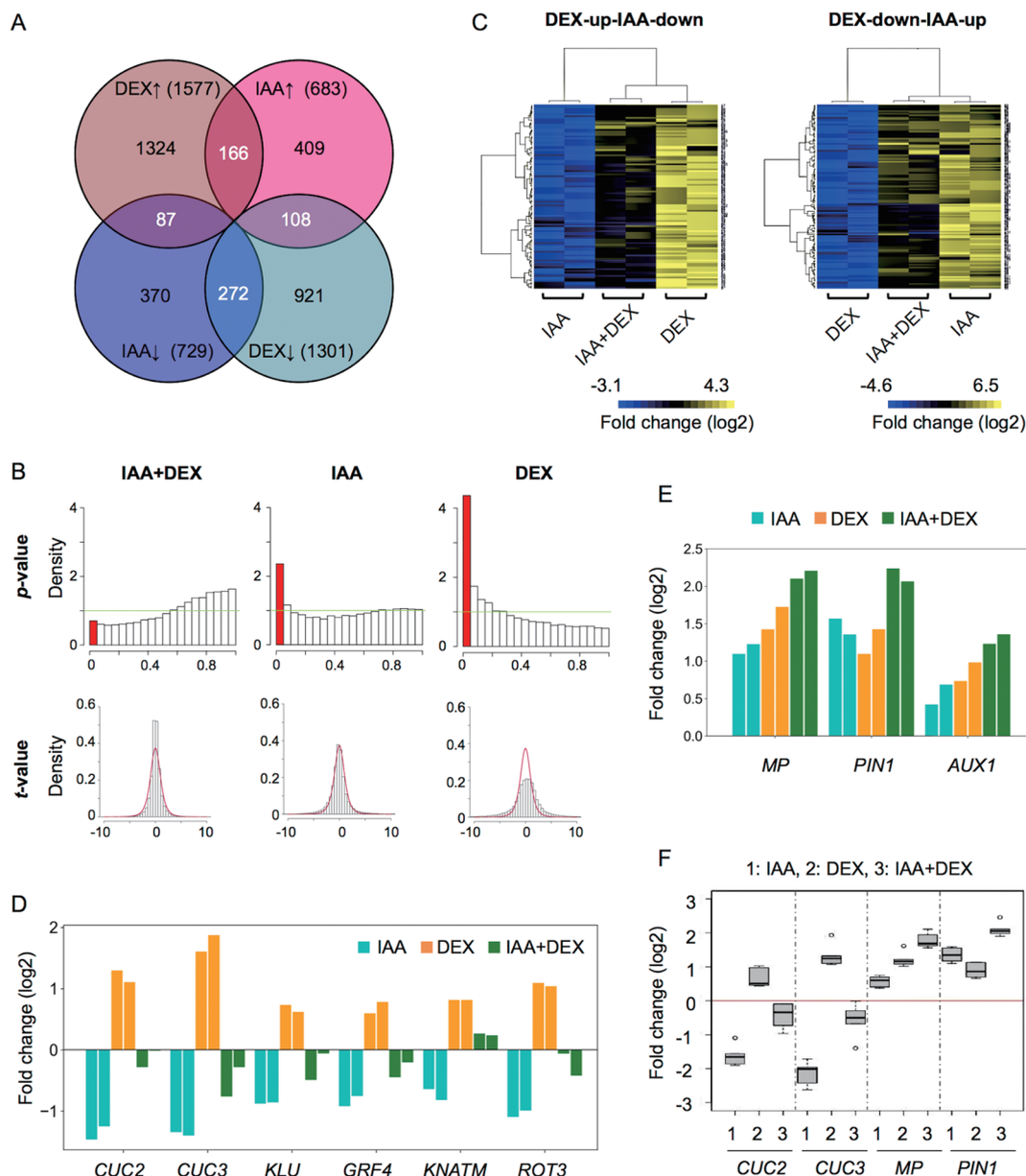


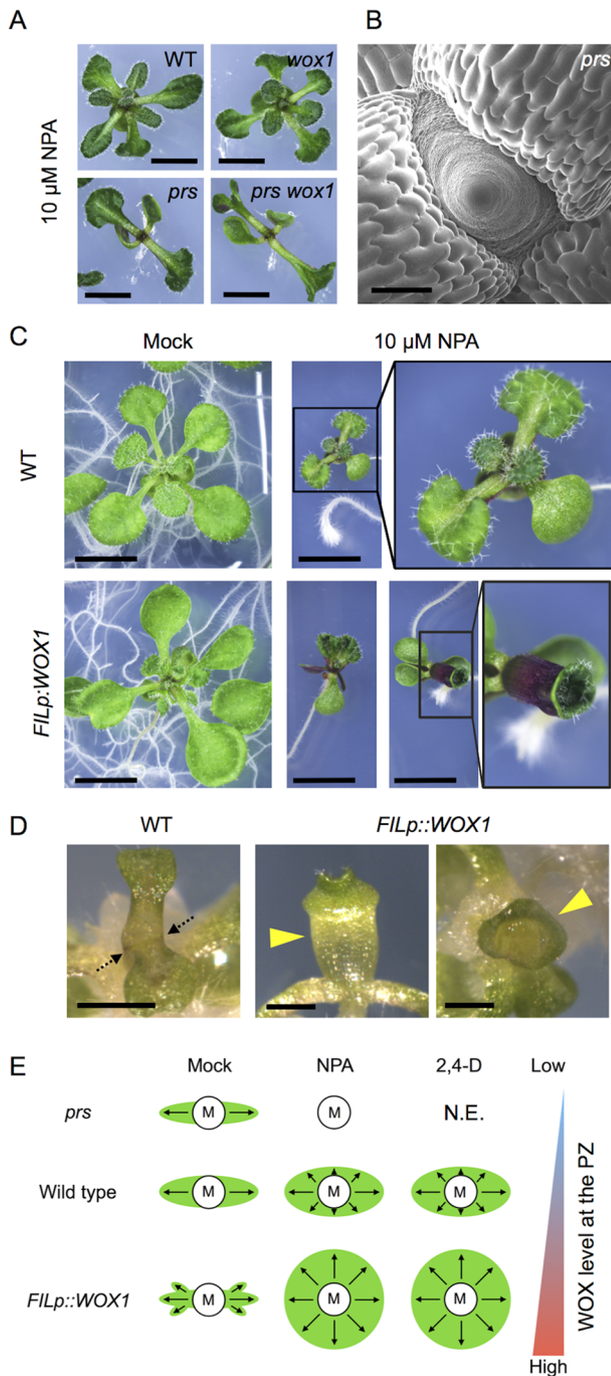
Figure 5. Genes altered in the *35Sp::WOX1-GR* treated with DEX and IAA. (A) Venn diagram illustrating the numbers of genes upregulated (upward-pointing arrow) and downregulated (downward-pointing arrow) with or without DEX and IAA. (B) Histograms showing the distributions of calculated  $t$ -values and  $p$ -values for the linear model's analysis of the interaction between DEX and IAA. See Materials and Methods for details. (C) Heat maps of fold changes in the expression levels of DEX-up-IAA-down and DEX-down-IAA-up genes compared with the mock treatment. Expression levels of DEX-up-IAA-down genes treated with IAA+DEX were greater than with IAA alone and lower than with DEX alone (left), while those of DEX-down-IAA-up genes treated with IAA+DEX were lower than with IAA alone and greater than with DEX alone (right). Data from different probes for a single gene are shown in different rows in the heat map. (D) Log<sub>2</sub>FCs in the expression levels of the DEX-up-IAA-down genes involved in shoot meristem and leaf development in comparison with the mock treatment. Two bars of the same color represent the results of two independent lines. (E) Log<sub>2</sub>FCs in the expression levels of *MP*, *PIN1* and *AUX1* in comparison with the mock treatment. Two bars of the same color represent the results of two independent lines. (F) Boxplots showing the results of the RT-qPCR analysis of *CUC2*, *CUC3*, *MP* and *PIN1*. Data were normalized by the *ACT2* expression level.

shaped leaf phenotype is considered to be a result from cell proliferation that occurred in the whole peripheral zone of the shoot apical meristem (Figure 6E). These findings indicate that *PRS* and *WOX1* are involved in cell proliferation in coordination with the auxin pathway during leaf initiation.

## Discussion

This study provides a new insight into the relationship between the function of auxin and *WOX1* at the gene expression level. In addition, it also suggested that the relationship is important for cell proliferation.

Here, we identified hundreds of genes affected by



**Figure 6.** Roles of WOX1/PRS and auxin in leaf development. (A) Phenotype of the aerial part of wild type, *prs*, *wox1* and *prs wox1* in the presence of 10  $\mu$ M NPA. Leaf initiation defects in 10  $\mu$ M NPA-treated *prs* and *prs wox1* plants. (B) Short pin-like phenotype of the 10  $\mu$ M NPA-treated *prs* shoot apex. (C) Phenotype of wild type and the *FILp::WOX1* plants treated with 10  $\mu$ M NPA. (D) Leaf phenotype of 1  $\mu$ M 2,4-D-treated *FILp::WOX1* plants. The arrows indicate the fusion of petioles between two leaves observed in wild type. The arrow heads indicate cup-shaped leaves observed in the *FILp::WOX1* plants. Plants shown in (A–D) were grown on a solid medium. (E) Summary of the leaf phenotypes of mock-, NPA- or 2,4-D-treated plants. Schematic diagram of the vicinity of the shoot apical meristem observed from above. “M” represents the center of the meristem, “PZ” represents the peripheral zone, and “N.E.” represents “not examined”. Arrows indicate the degree and direction of growth. Scale bars, 4 mm (A), 200  $\mu$ m (B), 1 cm (C), 1 mm (D).

the activation of *WOX1* using a microarray analysis of the shoot apices of *35Sp::WOX1-GR* plants after DEX treatments. Both the upregulated and downregulated genes were enriched for the GO term “response to auxin stimulus”, which suggests that the effects of *WOX1* on the auxin pathway are complex. In experiments with CHX, *WOX1* indirectly activated and directly repressed auxin responsive genes. The effects of DEX and IAA were globally additive, and changes in the expression levels of both the DEX-up-IAA-down and the DEX-down-IAA-up genes did not occur when plants were treated with IAA+DEX. In addition, *MP*, *PIN1*, *AUX1* and *LAX1* were upregulated by DEX. The expression sites of *MP*, *PIN1*, *AUX1* and *LAX1* overlap with those of *WOX1* and *PRS* at the leaf margins (Kasprzewska et al. 2015; Qi et al. 2014; Scarpella et al. 2006). A role for *WOX1* in auxin transport is indicated by the observation that the DR5::GFP signal is broadly distributed in the leaf base of *prs wox1* mutant and the auxin maxima were ectopically formed at the abaxial protrusions of the *FILp::WOX1* plants leaves. Based on these findings, we concluded that *WOX1* positively regulates auxin response and transport but suppresses the expression of some auxin downstream genes independent of auxin. Furthermore, the results inferred that *WOX1* stimulates only particular events downstream of auxin, while preventing others. The microarray analysis using a mutant of the *WOX1* homolog of *Medicago truncatula* (*stenofolia*) revealed the decreased expression of the *PIN1* homolog and the increased expression of auxin-responsive SAUR homologous genes (Tadege et al. 2011), which is consistent with our results. However, in *A. thaliana prs wox1*, the DR5::GFP signal remained in the basal region, which was different from the extremely depressed signal of the DR5 marker in the tobacco mutant of *WOX1* homolog (*lam1*) (Tadege et al. 2011). Therefore, the role of *WOX1* in the auxin pathway suggested by our study may be basically, but not fully, conserved among eudicot plants.

Recent studies have revealed that the expression levels of *WOX1* and *PRS* in leaf primordia are induced by auxin (Caggiano et al. 2017; Shi et al. 2017) and that the induction is directly mediated by MP (Guan et al. 2017). These findings, together with our analysis, indicate that there is a positive feedback between auxin and *WOX1/PRS* (Figure 7A). *WOX1* and *PRS* act as the middle-domain regulators (Nakata et al. 2012) and mutually regulate the adaxial and abaxial genes in *A. thaliana*, *Petunia x hybrida*, *M. truncatula* and rice (Guan et al. 2017; Honda et al. 2018; Nakata and Okada 2012; 2013; Nakata et al. 2012; Vandenbussche et al. 2009; Zhang et al. 2014; Table 3). However, the adaxial/abaxial pattern contributes to dynamic auxin pattern formation (Caggiano et al. 2017; Izhaki and Bowman 2007). The adaxial/abaxial genes affect auxin responsive genes (Huang et al. 2014; Merelo et al. 2013), while auxin is important for the adaxial/abaxial patterning

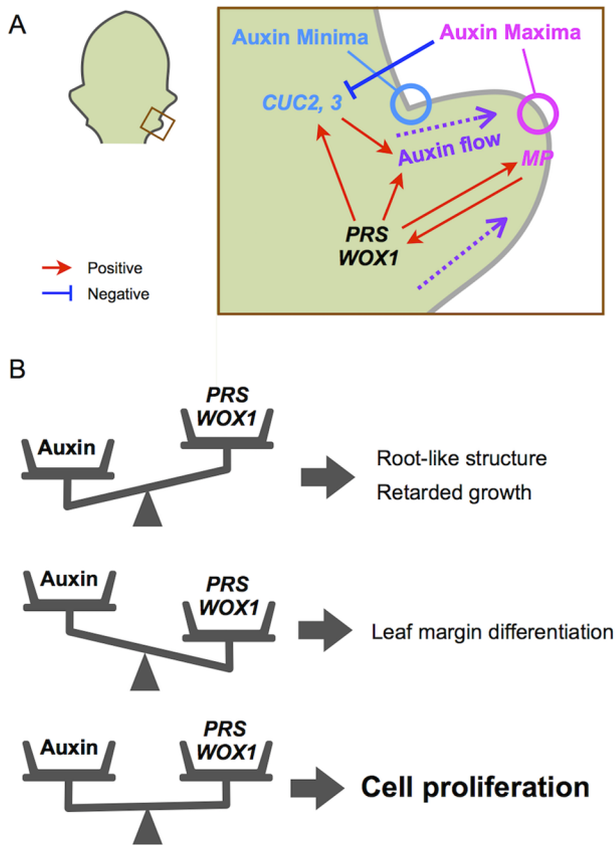


Figure 7. Models of the functions of *WOX1/PRS* and auxin. (A) The spatio-temporal model of the functions of *WOX1/PRS* and auxin in leaf development of wild type. (B) The model of the role of the balance between *WOX1/PRS* and auxin in controlling cell proliferation.

(Qi et al. 2014). Besides adaxial/abaxial associated genes, our data show that the expression levels of *CUC2* and *CUC3* were oppositely regulated by auxin and *WOX1* (Figure 7A). *CUC2*, which functions as a dimer with *CUC3* (Rubio-Somoza et al. 2014), modulates the growth patterning along the medial-lateral axis (Hasson et al. 2011; Kawamura et al. 2010; Nikovics et al. 2006) and cooperatively works with *PIN1*, *AUX1* and *LAX1* (Bilsborough et al. 2011; Kasprzewska et al. 2015). Thus, *WOX1/PRS*, MP-auxin, adaxial/abaxial genes and *CUC2/CUC3* establish a complicated regulatory network, and this network organizes leaf developmental events based on the adaxial–abaxial and the medial–lateral axes.

In addition to the role in leaf patterning, the combination of *WOX1* and auxin contributed to cell proliferation under certain conditions. The simultaneous application of DEX and 2,4-D to plants harboring *35Sp::WOX1-GR* construct induced unorganized cell proliferation. However, the application of DEX without 2,4-D caused dwarfism and the differentiation of the leaf-margin cells, while a high concentration of 2,4-D retarded cell proliferation, even in the presence of DEX. Based on these results, we propose a model in which the balance between the activity levels of auxin

and *WOX1* is a key factor in stimulating unorganized cell proliferation (Figure 7B). In cells where the level of auxin is low and *WOX1/PRS* is highly expressed, cell differentiation is triggered, resulting in the suppression of cell proliferation. Also, in a tissue where auxin is abundant but the expression level of *WOX1/PRS* is not high enough, root tissues are formed or cell proliferation is slowed or another tissue identities such as root are acquired. Cell proliferation occurs only when the amount of auxin and the expression level of *WOX1* are well balanced. In this model, the additive effect of *WOX1* and auxin at the gene expression level potentially explains why the balance between them is important for cell proliferation because only a subset of genes are strongly upregulated and downregulated whereas modulation of other *WOX1* and auxin downstream genes are offset.

This model can also explain the phenotypes of NPA-treated *prs* and NPA-treated the *FILp::WOX1* plants. The phenotype of the NPA-treated *prs* mutant indicated that *PRS* has the most important role in leaf initiation among the redundantly acting *WOX* family members. In vegetative shoots, *PRS* is expressed in the peripheral zone of the shoot apical meristem (Caggiano et al. 2017) as well as in early leaf primordia, in contrast with the faint or non-expression of *WOX1* (Caggiano et al. 2017; Matsumoto and Okada 2001; Nakata et al. 2012; Yu et al. 2017). Because NPA causes the broad distribution of the auxin response throughout the peripheral zone (Caggiano et al. 2017), the reduced total activity level of the *WOX* genes in the *prs* mutant might disrupt the balance between the activity levels of *WOX* and auxin, resulting in retarded cell proliferation during leaf initiation. However, because the *FIL* promoter induces relatively strong expression levels throughout the emerging primordia (Tameshige et al. 2013), in the *FILp::WOX1* plants, *WOX1* can be expressed in a broad region of the peripheral zone. In addition, the NPA treatment on the *FILp::WOX1* plants caused the retention of auxin, which may have generated a proper balance between *WOX1* and auxin throughout the peripheral zone, resulting in the formation of cup-shaped or fused leaves. 2,4-D is a substrate of *AUX1* but is not moved by an efflux carrier (Delbarre et al. 1996; Yang et al. 2006). Therefore, 2,4-D could trigger the cup-shaped phenotype similar to that induced by NPA. Although this model is consistent with the phenomenon found in this study, our observations were limited to mature leaves. Consequently, it is necessary to further verify this model by conducting detailed analyses of the leaf primordia at the cellular level.

In this study, unorganized cell proliferation was caused by the combined actions of *WOX1* and auxin but how these actions can be used to improve biomass is unclear. Because the balance between the actions of *WOX1* and auxin could be important, changes in the amount of

auxin that are dependent on leaf development may make it difficult to continue triggering cell proliferation using the combination of WOX1 and auxin. In addition, the constitutive expression of WOX1 strongly inhibited root growth in *A. thaliana*, which might be a negative factor that interferes with biomass improvement. Continuous cell proliferation in developing leaves can be caused by the prolonged expression of *PRS* and *WOX1* that results from the suppression of the TCP/NGA function (Alvarez et al. 2016). This report suggests that under particular cellular environments the increased expression levels of *PRS* and *WOX1* could result in biomass improvement, even in *A. thaliana*. Studies of grass species have suggested that the cytokinin pathway is responsible for WOX1-triggered biomass improvement (Wang et al. 2017). Here, both WOX1 and auxin downregulated *CKX7* and *ARR7*, which negatively affect the cytokinin pathway (Kölmer et al. 2014; To et al. 2007). Therefore, WOX1 may improve the biomass together with auxin through the activation of the cytokinin pathway. Interestingly, our microarray analysis also implied that both WOX1 and auxin suppress the abscisic acid pathway, which negatively affects leaf growth (Tardieu et al. 2010), through the downregulation of abscisic-acid receptors. To elucidate the regulatory mechanism of cell proliferation, it is necessary to understand the balance between auxin and WOX1, and the relationships with the cytokinin and with the abscisic-acid pathways. Fully understanding how the balance of WOX1/*PRS* and auxin controls cell proliferation could lead to their manipulation, which could improve biomass and allow for the modification of plant morphology in the future.

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### References

- Alvarez JP, Furumizu C, Efroni I, Eshed Y, Bowman JL (2016) Active suppression of a leaf meristem orchestrates determinate leaf growth. *eLife* 5: e15023
- Aoyama T, Chua NH (1997) A glucocorticoid-mediated transcriptional induction system in transgenic plants. *Plant J* 11: 605–612
- Bilborough GD, Runions A, Barkoulas M, Jenkins HW, Hasson A, Galinha C, Laufs P, Hay A, Prusinkiewicz P, Tsiantis M (2011) Model for the regulation of *Arabidopsis thaliana* leaf margin development. *Proc Natl Acad Sci USA* 108: 3424–3429
- Brady SM, Burow M, Busch W, Carlborg Ö, Denby KJ, Glazebrook J, Hamilton ES, Harmer SL, Haswell ES, Maloof JN, et al. (2015) Reassess the *t* Test: Interact with All Your Data via ANOVA. *Plant Cell* 27: 2088–2094
- Caggiano MP, Yu X, Bhatia N, Larsson A, Ram H, Ohno CK, Sappl P, Meyerowitz EM, Jönsson H, Heisler MG (2017) Cell type boundaries organize plant development. *eLife* 6: e27421
- Cho SH, Yoo SC, Zhang H, Pandeya D, Koh HJ, Hwang JY, Kim GT, Paek NC (2013) The rice *narrow leaf2* and *narrow leaf3* loci encode WUSCHEL-related homeobox 3A (OsWOX3A) and function in leaf, spikelet, tiller and lateral root development. *New Phytol* 198: 1071–1084
- Delbarre A, Muller P, Imhoff V, Guern J (1996) Comparison of mechanisms controlling uptake and accumulation of 2,4-dichlorophenoxy acetic acid, naphthalene-1-acetic acid, and indole-3-acetic acid in suspension-cultured tobacco cells. *Planta* 198: 532–541
- Guan C, Wu B, Yu T, Wang Q, Krogan NT, Liu X, Jiao Y (2017) Spatial Auxin Signaling Controls Leaf Flattening in Arabidopsis. *Curr Biol* 27: 2940–2950.e4
- Hasson A, Plessis A, Blein T, Adroher B, Grigg S, Tsiantis M, Boudaoud A, Damerval C, Laufs P (2011) Evolution and diverse roles of the *CUP-SHAPED COTYLEDON* genes in *Arabidopsis* leaf development. *Plant Cell* 23: 54–68
- Honda E, Yew C, Yoshikawa T, Sato Y, Hibara K, Itoh J (2018) *LEAF LATERAL SYMMETRY1*, a member of the *WUSCHEL-RELATED HOMEBOX3* gene family, regulates lateral organ development differentially from other paralogs, *NARROW LEAF2* and *NARROW LEAF3* in rice. *Plant Cell Physiol* 59: 376–391
- Huang T, Harrar Y, Lin C, Reinhart B, Newell NR, Talavera-Rauh F, Hokin SA, Barton MK, Kerstetter RA (2014) Arabidopsis KANADI1 acts as a transcriptional repressor by interacting with a specific *cis*-element and regulates auxin biosynthesis, transport, and signaling in opposition to HD-ZIPIII factors. *Plant Cell* 26: 246–262
- Ikeda M, Mitsuda N, Ohme-Takagi M (2009) *Arabidopsis* WUSCHEL is a bifunctional transcription factor that acts as a repressor in stem cell regulation and as an activator in floral patterning. *Plant Cell* 21: 3493–3505
- Ishiwata A, Ozawa M, Nagasaki H, Kato M, Noda Y, Yamaguchi T, Nosaka M, Shimizu-Sato S, Nagasaki A, Maekawa M, et al. (2013) Two *WUSCHEL*-related homeobox genes, *narrow leaf2* and *narrow leaf3*, control leaf width in rice. *Plant Cell Physiol* 54: 779–792
- Izhaki A, Bowman JL (2007) KANADI and class III HD-Zip gene families regulate embryo patterning and modulate auxin flow

- during embryogenesis in *Arabidopsis*. *Plant Cell* 19: 495–508
- Kasprzewska A, Carter R, Swarup R, Bennett M, Monk N, Hobbs JK, Fleming A (2015) Auxin influx importers modulate serration along the leaf margin. *Plant J* 83: 705–718
- Kawakami S, Watanabe Y (1997) Use of green fluorescent protein as a molecular marker tag of protein movement *in vivo*. *Plant Biotechnol* 14: 127–130
- Kawamura E, Horiguchi G, Tsukaya H (2010) Mechanisms of leaf tooth formation in *Arabidopsis*. *Plant J* 62: 429–441
- Köllmer I, Novák O, Strnad M, Schmölling T, Werner T (2014) Overexpression of the cytosolic cytokinin oxidase/dehydrogenase (CKX7) from *Arabidopsis* causes specific changes in root growth and xylem differentiation. *Plant J* 78: 359–371
- Lian G, Ding Z, Wang Q, Zhang D, Xu J (2014) Origins and evolution of WUSCHEL-related homeobox protein family in plant kingdom. *Sci World J* 2014: 534140
- Lin H, Niu L, McHale NA, Ohme-Takagi M, Mysore KS, Tadege M (2013) Evolutionarily conserved repressive activity of WOX proteins mediates leaf blade outgrowth and floral organ development in plants. *Proc Natl Acad Sci USA* 110: 366–371
- Matsumoto N, Okada K (2001) A homeobox gene, PRESSED FLOWER, regulates lateral axis-dependent development of *Arabidopsis* flowers. *Genes Dev* 15: 3355–3364
- McHale NA (1993) LAM-1 and FAT genes control development of the leaf blade in *Nicotiana sylvestris*. *Plant Cell* 5: 1029–1038
- McHale NA, Marcotrigiano M (1998) LAM1 is required for dorsoventrality and lateral growth of the leaf blade in *Nicotiana*. *Development* 125: 4235–4243
- Merelo P, Xie Y, Brand L, Ott F, Weigel D, Bowman JL, Heisler MG, Wenkel S (2013) Genome-wide identification of KANADII1 target genes. *PLoS One* 8: e77341
- Nakata M, Matsumoto N, Tsugeki R, Rikirsch E, Laux T, Okada K (2012) Roles of the middle domain-specific WUSCHEL-RELATED HOMEBOX genes in early development of leaves in *Arabidopsis*. *Plant Cell* 24: 519–535
- Nakata M, Okada K (2012) The three-domain model: A new model for the early development of leaves in *Arabidopsis thaliana*. *Plant Signal Behav* 7: 1423–1427
- Nakata M, Okada K (2013) The Leaf adaxial–abaxial boundary and lamina growth. *Plants* 2: 174–202
- Nakata MT, Sato M, Wakazaki M, Sato N, Kojima K, Sekine A, Nakamura S, Shikanai T, Toyooka K, Tsukaya H, et al. (2018) Plastid translation is essential for lateral root stem cell patterning in *Arabidopsis thaliana*. *Biol Open* 7: bio028175
- Nardmann J, Ji J, Werr W, Scanlon MJ (2004) The maize duplicate genes *narrow sheath1* and *narrow sheath2* encode a conserved homeobox gene function in a lateral domain of shoot apical meristems. *Development* 131: 2827–2839
- Nardmann J, Reisewitz P, Werr W (2009) Discrete shoot and root stem cell-promoting WUS/WOX5 functions are an evolutionary innovation of angiosperms. *Mol Biol Evol* 26: 1745–1755
- Nikovics K, Blein T, Peaucelle A, Ishida T, Morin H, Aida M, Laufs P (2006) The balance between the *MIR164A* and *CUC2* genes controls leaf margin serration in *Arabidopsis*. *Plant Cell* 18: 2929–2945
- Ottenschläger I, Wolff P, Wolverson C, Bhalerao RP, Sandberg G, Ishikawa H, Evans M, Palme K (2003) Gravity-regulated differential auxin transport from columella to lateral root cap cells. *Proc Natl Acad Sci USA* 100: 2987–2991
- Qi J, Wang Y, Yu T, Cunha A, Wu B, Vernoux T, Meyerowitz E, Jiao Y (2014) Auxin depletion from leaf primordia contributes to organ patterning. *Proc Natl Acad Sci USA* 111: 18769–18774
- Rubio-Somoza I, Zhou CM, Confraria A, Martinho C, von Born P, Baena-Gonzalez E, Wang JW, Weigel D (2014) Temporal control of leaf complexity by miRNA-regulated licensing of protein complexes. *Curr Biol* 24: 2714–2719
- Scanlon MJ, Freeling M (1997) Clonal sectors reveal that a specific meristematic domain is not utilized in the maize mutant *narrow sheath*. *Dev Biol* 182: 52–66
- Scanlon MJ, Schneeberger RG, Freeling M (1996) The maize mutant *narrow sheath* fails to establish leaf margin identity in a meristematic domain. *Development* 122: 1683–1691
- Scarpella E, Marcos D, Friml J, Berleth T (2006) Control of leaf vascular patterning by polar auxin transport. *Genes Dev* 20: 1015–1027
- Shi J, Dong J, Xue J, Wang H, Yang Z, Jiao Y, Xu L, Huang H (2017) Model for the role of auxin polar transport in patterning of the leaf adaxial–abaxial axis. *Plant J* 92: 469–480
- Shimada TL, Shimada T, Hara-Nishimura I (2010) A rapid and non-destructive screenable marker, FAST, for identifying transformed seeds of *Arabidopsis thaliana*. *Plant J* 61: 519–528
- Tadege M, Lin H, Bedair M, Berbel A, Wen J, Rojas CM, Niu L, Tang Y, Sumner L, Ratet P, et al. (2011) *STENOFOLIA* regulates blade outgrowth and leaf vascular patterning in *Medicago truncatula* and *Nicotiana sylvestris*. *Plant Cell* 23: 2125–2142
- Tameshige T, Fujita H, Watanabe K, Toyokura K, Kondo M, Tatematsu K, Matsumoto N, Tsugeki R, Kawaguchi M, Nishimura M, et al. (2013) Pattern dynamics in adaxial–abaxial specific gene expression are modulated by a plastid retrograde signal during *Arabidopsis thaliana* leaf development. *PLoS Genet* 9: e1003655
- Tardieu F, Parent B, Simonneau T (2010) Control of leaf growth by abscisic acid: Hydraulic or non-hydraulic processes? *Plant Cell Environ* 33: 636–647
- Thomson KS, Hertel R, Müller S, Tavares JE (1973) 1-N-naphthylphthalamic acid and 2,3,5-triiodobenzoic acid: In-vitro binding to particulate cell fractions and action on auxin transport in corn coleoptiles. *Planta* 109: 337–352
- To JP, Deruère J, Maxwell BB, Morris VF, Hutchison CE, Ferreira FJ, Schaller GE, Kieber JJ (2007) Cytokinin regulates type-A *Arabidopsis* Response Regulator activity and protein stability via two-component phosphorelay. *Plant Cell* 19: 3901–3914
- Truernit E, Bauby H, Dubreucq B, Grandjean O, Runions J, Barthélémy J, Palauqui JC (2008) High-resolution whole-mount imaging of three-dimensional tissue organization and gene expression enables the study of Phloem development and structure in *Arabidopsis*. *Plant Cell* 20: 1494–1503
- Vandenbussche M, Horstman A, Zethof J, Koes R, Rijpkema AS, Gerats T (2009) Differential recruitment of WOX transcription factors for lateral development and organ fusion in *Petunia* and *Arabidopsis*. *Plant Cell* 21: 2269–2283
- van der Graaff E, Laux T, Rensing SA (2009) The WUS homeobox-containing (WOX) protein family. *Genome Biol* 10: 248
- Wang H, Niu L, Fu C, Meng Y, Sang D, Yin P, Wu J, Tang Y, Lu T, Wang ZY, et al. (2017) Overexpression of the WOX gene *STENOFOLIA* improves biomass yield and sugar release in transgenic grasses and display altered cytokinin homeostasis. *PLoS Genet* 13: e1006649
- Yang Y, Hammes UZ, Taylor CG, Schachtman DP, Nielsen E (2006) High-affinity auxin transport by the AUX1 influx carrier protein. *Curr Biol* 16: 1123–1127
- Yoshikawa T, Tanaka SY, Masumoto Y, Nobori N, Ishii H, Hibara K, Itoh J, Tanisaka T, Taketa S (2016) Barley NARROW LEAFED DWARF1 encoding a WUSCHEL-RELATED HOMEBOX 3

- (WOX3) regulates the marginal development of lateral organs. *Breed Sci* 66: 416–424
- Yu T, Guan C, Wang J, Sajjad M, Ma L, Jiao Y (2017) Dynamic patterns of gene expression during leaf initiation. *J Genet Genomics* 44: 599–601
- Zhang F, Wang Y, Li G, Tang Y, Kramer EM, Tadege M (2014) STENOFOLIA recruits TOPLESS to repress ASYMMETRIC LEAVES2 at the leaf margin and promote leaf blade outgrowth in *Medicago truncatula*. *Plant Cell* 26: 650–664
- Zhang Y, Wu R, Qin G, Chen Z, Gu H, Qu LJ (2011) Over-expression of WOX1 leads to defects in meristem development and polyamine homeostasis in Arabidopsis. *J Integr Plant Biol* 53: 493–506
- Zheng Q, Wang XJ (2008) GOEAST: A web-based software toolkit for Gene Ontology enrichment analysis. *Nucleic Acids Res* 36(suppl\_2): W358–63
- Zhuang LL, Ambrose M, Rameau C, Weng L, Yang J, Hu XH, Luo D, Li X (2012) LATHYROIDES, encoding a WUSCHEL-related Homeobox1 transcription factor, controls organ lateral growth, and regulates tendril and dorsal petal identities in garden pea (*Pisum sativum* L.). *Mol Plant* 5: 1333–1345