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# WAG2 represses apical hook opening downstream from gibberellin and PHYTOCHROME INTERACTING FACTOR 5

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#### **SUMMARY**

When penetrating the soil during germination, dicotyledonous plants protect their shoot apical meristem through the formation of an apical hook. Apical hook formation is a dynamic process that can be subdivided into hook formation, maintenance and opening. It has previously been established that these processes require the transport and signaling of the phytohormone auxin, as well as the biosynthesis and signaling of the phytohormones ethylene and gibberellin (GA). Here, we identify a molecular mechanism for an auxin-GA crosstalk by demonstrating that the auxin transport-regulatory protein kinase WAG2 is a crucial transcription target during apical hook opening downstream from GA signaling. We further show that WAG2 is directly activated by PHYTOCHROME INTERACTING FACTOR 5 (PIF5), a light-labile interactor of the DELLA repressors of the GA pathway. We find that wag2 mutants are impaired in the repression of apical hook opening in dark-grown seedlings and that this phenotype correlates with GA-regulated WAG2 expression in the concave (inner) side of the apical hook. Furthermore, wag2 mutants are also impaired in the maintenance or formation of a local auxin maximum at the site of WAG2 expression in the hook. WAG2 is a regulator of PIN auxin efflux facilitators and, in line with previous data, we show that this kinase can phosphorylate the central intracellular loop of all PIN-FORMED (PIN) proteins regulating apical hook opening. We therefore propose that apical hook opening is controlled by the differential GA-regulated accumulation of WAG2 and subsequent local changes in PIN-mediated auxin transport.

KEY WORDS: Arabidopsis, WAG2 protein kinase, Apical hook, Auxin transport, Gibberellin

#### **INTRODUCTION**

Gibberellins (GA) are phytohormones that regulate a number of physiological responses in plants such as germination, elongation growth, greening, flowering time as well as apical hook formation (Achard et al., 2003; Alabadí et al., 2004; Gallego-Bartolomé et al., 2011; Ueguchi-Tanaka et al., 2007; Vriezen et al., 2004). All GA responses known to date are mediated by DELLA proteins, which have emerged as key repressors of the GA pathway that regulate different classes of transcriptional regulators such as the light-labile PHYTOCHROME INTERACTING FACTORs (PIFs; namely PIF1, PIF3, PIF4 and PIF5) and the related but light-stable bHLH transcription factors ALCATRAZ and SPATULA (Arnaud et al., 2010; de Lucas et al., 2008; Feng et al., 2008; Gallego-Bartolomé et al., 2011; Gallego-Bartolomé et al., 2010).

GA controls DELLA protein abundance by binding to the GIBBERELLIN INSENSITIVE DWARF 1 (GID1) receptors and then promotes DELLA protein degradation via E3 ubiquitin ligases such as *Arabidopsis* SCF<sup>SLEEPY1(SLY1)</sup> (Ueguchi-Tanaka et al., 2007). Therefore, GA and DELLAs antagonistically control the activity of their downstream regulators, e.g. the DNA-binding activity of PIFs is repressed by DELLA interactions and GA relieves this DELLA-imposed restraint by promoting DELLA degradation (de Lucas et al., 2008; Feng et al., 2008). At the same time, PIF abundance is also downregulated by light as light-induced phytochrome-interactions promote PIF protein degradation (Bauer et al., 2004; Castillon et al., 2009; Lorrain et al., 2008;

AUX/IAA proteins (Chapman and Estelle, 2009). At the organismal level, the polar transport of auxin within the plant plays a pivotal role in the control of plant development and differentiation. Cell-to-cell polar auxin transport is mediated by the AUXIN-RESISTANT 1/LIKE-AUX 1 (AUX1/LAX) auxin import carriers and PIN-FORMED (PIN) auxin efflux facilitators that may function together with MULTIDRUG RESISTANCE/PHOSPHOGLYCOPROTEIN (MDR/PGP) transporters (Kleine-Vehn and Friml, 2008). Particularly instructive for the understanding of polar auxin transport is the localization of PIN auxin efflux facilitators that are polarly distributed in the plasma membrane of many cells in the growing plant. In *Arabidopsis* 

thaliana, at least five members of the PIN protein family with in

part specific and in part overlapping biological functions have been implicated in PIN-dependent polar auxin transport, PIN1, PIN2,

PIN3 and PIN4, as well as PIN7 (Blilou et al., 2005). Although auxin transport itself cannot be directly visualized, models for

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Monte et al., 2004; Park et al., 2004; Shen et al., 2005). As such, the repression of PIFs by DELLAs is particularly important when PIFs are abundant, notably in the dark, whereas phytochrome-dependent PIF degradation may be the predominant pathway controlling PIF abundance and activity in the light (Leivar and Quail, 2011). Although two recent reports reveal the identity of PIF transcription factor target genes using chromatin immunoprecipitation at the genome-wide level (Hornitschek et al., 2012; Oh et al., 2009), only a few studies have examined direct PIF targets in a biological context (Cheminant et al., 2011; Franklin et al., 2011; Gallego-Bartolomé et al., 2011; Li et al., 2012; Richter et al., 2010; Sun et al., 2012).

The phytohormone auxin regulates a large number of growth

processes in plants such as tissue differentiation, organ formation

and tropic responses (Teale et al., 2006). At the cellular level, auxin

responses are controlled by the auxin response factor

transcriptional regulators and by the inhibitory auxin-labile

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auxin transport by far and large support the notion that the knowledge on PIN polarity is sufficient to predict cell-to-cell auxin transport and to explain auxin-dependent growth processes (Grieneisen and Scheres, 2009; Wisniewska et al., 2006).

AGC kinases are a family of serine/threonine kinases that share homology with the well-characterized protein kinase A, protein kinase G and cAMP-dependent kinase from animals (Galván-Ampudia and Offringa, 2007). In *Arabidopsis*, a subset of the AGC kinases, PINOID (PID), PID2, WAG1 and WAG2 have been directly implicated in the control of auxin transport. PIDs and WAGs form a subgroup within the AGCVIII family and at least PID and the WAGs regulate PIN polarity by phosphorylating PINs at conserved phosphorylation sites (Cheng et al., 2008; Dhonukshe et al., 2010; Friml et al., 2004; Huang et al., 2010; Santner and Watson, 2006). The four members of the D6 PROTEIN KINASE (D6PK) subgroup of AGCVIII also regulate auxin transport and phosphorylate PINs; they have, however, been proposed to regulate PIN auxin transport activity rather than PIN polarity (Dhonukshe et al., 2010; Zourelidou et al., 2009). Interestingly, the blue light receptors PHOTOTROPIN1 (PHOT1) and PHOT2, which promote the auxin transport-dependent hypocotyl bending in response to lateral light, are also AGCVIII kinases (Christie et al., 2011; Galván-Ampudia and Offringa, 2007).

A number of growth and developmental processes in plants are regulated by auxin as well as by GA. To date, only a few studies have elucidated crosstalk mechanisms that can explain aspects of the interdependency of the two hormone pathways (Björklund et al., 2007; Frigerio et al., 2006; Fu and Harberd, 2003; Jacobs and Case, 1965; Scott et al., 1967; Willige et al., 2011). Auxin stimulates GA biosynthesis and it can be proposed that auxin promotes growth at least in part by increasing GA hormone synthesis and thereby degradation of the GA-labile DELLA growth repressors (Frigerio et al., 2006), e.g. the cessation of root growth in the absence of auxin transport from the shoot and the accumulation of DELLA proteins in the root may be explained through this mechanism (Fu and Harberd, 2003). Inversely, evidence for a control of auxin transport and signaling by GA has also been provided. Auxin transport in the stem is less efficient in GA signaling mutants and GA deficiency reduces PIN protein abundance, possibly by targeting PINs for degradation in the vacuole (Willige et al., 2011).

The development of the apical hook is a specific example for a process that is controlled by auxin transport as well as by GA signaling (Gallego-Bartolomé et al., 2011; Lehman et al., 1996; Li et al., 2004; Zádníková et al., 2010). The apical hook is formed during skotomorphogenesis of dicotyledonous seedlings and it protects the sensitive shoot apical meristem while the hypocotyl is growing through the soil to reach the light. Apical hook development can be divided into the hook formation, hook maintenance and hook opening phase (Raz and Ecker, 1999; Zádníková et al., 2010). The local accumulation of auxin is necessary for the formation and maintenance of the apical hook to restrict the elongation of cells in the concave (inner) side of the hook, which induces asymmetric growth in the apical part of the hypocotyl (Kuhn and Galston, 1992; Raz and Ecker, 1999). Recently, the auxin influx facilitators AUX1 and LAX3, as well as the auxin efflux facilitators PIN1, PIN3, PIN4 and PIN7, were identified as being necessary for normal hook development (Vandenbussche et al., 2010; Zádníková et al., 2010). In particular, the loss of PIN3 activity impairs the establishment of the asymmetric auxin maximum in the apical hook (Friml et al., 2002; Zádníková et al., 2010).

Like auxin transport, GA biosynthesis and signaling are prerequisite for the formation and maintenance of the apical hook (Achard et al., 2003; Alabadí et al., 2004; An et al., 2012; Gallego-Bartolomé et al., 2011; Vriezen et al., 2004): although the loss of GA biosynthesis in the gal mutant results in the loss of hook formation, the weaker DELLA gain-of-function mutant gai-1 is able to form a weak apical hook that opens prematurely. By contrast, DELLA loss-of-function mutants form an exaggerated hook, indicating that DELLA protein function is required to control the degree of hook formation in the wild type. From two recent studies, it has emerged that GA controls the transcription of the ethylene biosynthesis genes AMINOCYCLOPROPANE CARBOXYLIC ACID SYNTHASE 5 (ACS5) and ACS8, of the ethylene response regulator HOOKLESS 1 (HLS1), as well as that of the auxin efflux facilitators PIN3 and PIN7 (An et al., 2012; Gallego-Bartolomé et al., 2011). The mode of transcriptional GA control could be elucidated for two of these GA target genes by chromatin immunoprecipitation (ChIP) where it was shown that ACS8 and HLS1 expression is directly controlled by the DELLA interactors PIF5 and ETHYLENE INSENSITIVE 3/EIN3-LIKE1 (EIN3/EIL1), respectively.

We have examined genes of the auxin transport machinery with regard to their transcriptional regulation by GA. In this study, we identified the AGC kinase WAG2 as a GA-regulated auxin transport gene. We further found that WAG2 represses apical hook opening in dark-grown Arabidopsis seedlings and that WAG2 is expressed in a GA-regulated manner in the concave side of the apical hook. We further identified the DELLA interactor PIF5 as a transcriptional regulator that activates WAG2 expression by binding to a G-box motif in the WAG2 promoter. Finally, in line with previous data on the role of WAG2 as regulator of auxin transport polarity (Dhonukshe et al., 2010), our data suggest that WAG2 may control apical hook opening by phosphorylating PIN proteins and by influencing the formation or maintenance of a specific local auxin maximum in the apical hook.

# MATERIALS AND METHODS

# **Biological material**

Unless otherwise stated in the text, mutants and transgenic lines used in this study have been previously described: ga1 (Willige et al., 2007); gai-1 (Peng et al., 1997); rga-24 gai-t6 (King et al., 2001); sly1-10 (McGinnis et al., 2003); DR5:GUS (Ulmasov et al., 1997); DR5rev:GFP (Friml et al., 2003); pif1, pif3, pif4, pif5 and PIF5ox (Fujimori et al., 2004; Huq et al., 2004; Khanna et al., 2007; Leivar et al., 2008; Lorrain et al., 2008; Monte et al., 2004); PIN1:GFP, pin3 and PIN3:GFP (Benková et al., 2003; Zádníková et al., 2010); wag1 and wag2 (Santner and Watson, 2006).

#### Growth conditions and physiological experiments

Seeds were stratified for 3 days at 4°C, exposed to white light for 8 hours and grown in the dark at 21°C. *ga1* mutant seeds were incubated for 5 days at 4°C in 100 µM GA3 to induce germination. The seeds were then thoroughly washed five times in water to remove excess GA before exposure to light for 20 hours. To measure growth parameters, all seedlings were grown on vertically oriented plates containing half-strength MS medium without sucrose. At the indicated time points, plates were scanned to determine apical hook angles or hypocotyl lengths using the ImageJ software (NIH). The angle of a completely closed apical hook was defined as 180°, whereas the angle of a fully opened hook was defined as 0°.

#### Cloning and transgenic material

To generate *WAG1pro:GUS* and *WAG2pro:GUS*, 2.5 kb and 2.9 kb promoter fragments were amplified from Col-0 genomic DNA using the primers WAG1pro-LP/RP and WAG2pro-LP/RP, respectively. These sequences were inserted as *EcoRI-NcoI* fragments into pCAMBIA1391Z. At least six transgenic lines were generated in the Col-0 ecotype and

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analyzed. Plant transformations were performed as previously described (Clough and Bent, 1998). The gene fragments of the cytoplasmic loops of PIN1, PIN2, PIN3, PIN4 and PIN7 were amplified by RT-PCR with specific PINLOOP-FW and PINLOOP-RV primers from *Arabidopsis thaliana* (Columbia) mRNA. The *WAG2*-coding sequence was amplified by PCR from cDNA clone U84295 (ABRC). The kinase-dead WAG2 (*WAG2in*) coding sequence was created using overlap extension PCR with the primers (WAG2mut5a and WAG2mut3a). The Gateway system-compatible fragments were then cloned into the expression vector pDEST15 (Invitrogen) to generate *GST:PIN* and *GST:WAG2*. See supplementary material Table S1 for a list of primers.

#### Quantitative real-time PCR

Extraction of total RNA and cDNA synthesis was conducted as described previously (Richter et al., 2010). The cDNA equivalent of 42 ng of total RNA was used in a 10 µl PCR reaction on a CFX96 Real-Time System Cycler with iQ SYBR Green Supermix (Bio-Rad). A 40-cycle two-step amplification protocol (10 seconds at 95°C, 30 seconds at 60°C) or a 40-cycle three-step amplification protocol (20 seconds at 95°C, 20 seconds at 58.5°C, 25 seconds at 72°C) was used for all measurements. Primer sequences are listed in supplementary material Table S1. Unless otherwise stated, the average and the standard error of at least three technical replicates pooled from three biological replicates are shown.

# **Biochemical analyses**

Protein extraction and immunoblotting were performed as previously described (Willige et al., 2011). Anti-GFP for detection of PIN1:GFP (1:3000; Invitrogen) or for detection of DR5:GFP (1:4000; Roche), anti-PIN3 (1:3000; NASC) and anti-RGA [1:1000 (Willige et al., 2011)] were used for immunoblotting. For ChIP, seedlings were grown in the dark for 3 days. ChIP quantitative RT-PCR was performed and analyzed as described previously (Oh et al., 2007; Fode and Gatz, 2009). Anti-HA agarose and anti-HA antibody (Roche) were used for precipitation and detection of PIF5:HA. The in vitro phosphorylation assays were conducted as described previously (Zourelidou et al., 2009).

#### Cell biological and histological analyses

For GUS staining, etiolated seedlings were fixed in heptane for 15 minutes and incubated in GUS staining solution [100 mM Na-phosphate buffer (pH 7.0), 2 mM K<sub>4</sub>Fe(CN)<sub>6</sub>, 2 mM K<sub>3</sub>Fe(CN)<sub>6</sub>, 0.1% Triton X-100, 1 mg/ml X-Gluc]. GUS-stained seedlings were photographed using a Leica MZ16 stereo-microscope with a PLAN-APOX1 objective (Leica). Microscopy of PIN1:GFP, PIN3:GFP and DR5rev:GFP was performed on 2- to 4-day-old etiolated seedlings using an Olympus BX61 fluorescence microscope with a digital camera XM10 (Olympus) or a FV1000/IX81 laser scanning confocal microscope.

#### **RESULTS**

# WAG2 is a GA-regulated AGC protein kinase

We have previously shown that *Arabidopsis* GA biosynthesis and pathway mutants are deficient in polar auxin transport (Willige et al., 2011). In search for possible molecular mechanisms that underlie the control of auxin transport by GA, we also examined the GA-dependent transcriptional regulation of genes encoding AGCVIII kinase family members that have been or may be implicated in the regulation of polar auxin transport as well as auxin transport proteins (Fig. 1A). To this end, we tested seedlings of the GA biosynthesis mutant gal for GA-dependent changes in transcript abundance by quantitative real time (qRT)-PCR. This experiment identified WAG2 as the gene whose transcript abundance is most strongly increased in response to GA3 treatment (Fig. 1A). As light-regulated signaling by the PIF transcription regulators may interfere with GA signaling, we were next interested in examining GA-regulation of WAG2 in dark-grown seedlings to eliminate a possible co-regulation of WAG2 expression by these two stimuli. In addition, this experiment revealed a strong

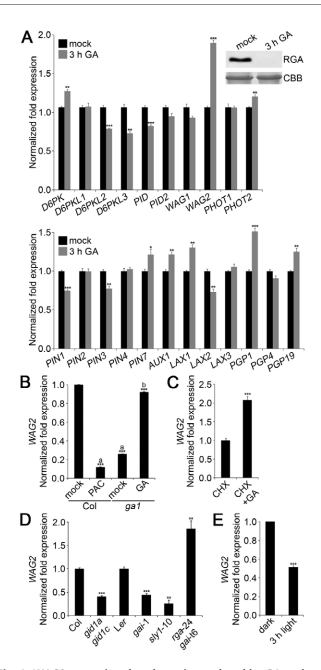


Fig. 1. WAG2 transcript abundance is regulated by GA and light. (A) qRT-PCR of biologically characterized AGC kinase and auxin transporter genes of 7-day-old light-grown seedlings following a 3 hour GA3 (100  $\mu$ M) treatment. The inset with an immunoblot with anti-RGA antibody of 45 µg total protein extracts indicates the efficiency of the GA3 treatment at the analyzed time point. (B) qRT-PCR analysis of WAG2 after GA3 (1 µM) treatment of 4-day-old dark-grown ga1 seedlings or wild-type (Col) seedlings grown in the presence of the GA biosynthesis inhibitor paclobutrazol (PAC). (C) qRT-PCR analysis of short-term GA3 (100  $\mu$ M, 1 hour)-treated dark-grown seedlings in the presence of the protein biosynthesis inhibitor cycloheximide (CHX, 50 μM). (**D**) Basal expression levels of WAG2 in different GA signaling mutants as detected by qRT-PCR. (E) qRT-PCR analysis of dark-grown seedlings and of dark-grown seedlings that were transferred to white light (150  $\mu$ M m<sup>-2</sup> s<sup>-1</sup>). Shown are the average and the standard deviation of two biological replicates. For all experiments, *UBC21* was used for normalization. Asterisks indicate the significances (Student's t-test: \*0.01<P<0.05; \*\*0.001 < P < 0.01; \*\*\*P < 0.001). In B, a and b indicate the comparisons to Col (mock) and ga1 (mock), respectively.

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GA-dependent increase in WAG2 transcript abundance in darkgrown gal seedlings, and inversely a decrease in WAG2 abundance in wild-type seedlings that had been depleted from GA by treatment with the GA biosynthesis inhibitor paclobutrazol (PAC) (Fig. 1B). As we observed that the GA-mediated regulation of WAG2 transcription can also take place in the presence of the protein synthesis inhibitor cycloheximide (CHX), we concluded that the GA response is dependent on a protein synthesisindependent mechanism such as GA-induced DELLA protein degradation (Fig. 1C). We then found that WAG2 abundance is reduced in other GA pathway mutants such as gidla gidlc, gai-1 and sly1-10, which are known to accumulate DELLA repressors and to be increased in the DELLA loss-of-function mutant rga-24 gai-t6 (Fig. 1D) (Peng et al., 1997). Finally, we found that WAG2 transcript abundance is reduced after the transfer of dark-grown seedlings to light, suggesting that WAG2 transcription is induced by light-labile regulators such as the PIFs (Fig. 1E). In summary, we concluded that WAG2 expression is regulated by DELLAs and may be downstream from PIFs, their light-labile interaction partners.

# WAG2 is a repressor of apical hook opening

To gain an insight into the possible role of GA-regulated WAG2 transcription, we phenotyped previously characterized wag2 mutants (Santner and Watson, 2006). As WAG2 expression is elevated in dark-grown seedlings, we paid particular attention to the effects of the loss of WAG2 function during skotomorphogenic growth and noticed with interest a decreased apical hook angle in 4-day-old wag2 mutants (Fig. 2A). At the same time, we did not observe any other phenotypes in dark-grown seedlings, such as changes in hypocotyl elongation (supplementary material Fig. S1). Interestingly, we measured an increased opening of the apical hook only in 3- and 4-day-old wag2 mutant seedlings (during the hook opening phase), and not in 2-day-old seedlings during the hook formation phase, suggesting that WAG2 is required for the repression of hook opening (Fig. 2B). As WAG2 had previously been found to genetically interact with its closest paralog, WAG1, in the control of root growth, we also included wag1 single and wag1 wag2 double mutants in our analysis (Santner and Watson, 2006). However, we found no evidence for a contribution of WAG1 to the apical hook opening phenotype of wag2, neither in the wag1 single mutant nor in the wag1 wag2 double mutant (Fig. 2A,B). We thus conclude that WAG2 represses apical hook opening in darkgrown seedlings and that WAG2 functions independently from WAG1 in this process.

We also tested whether GA is required and sufficient for the normalization of apical hook development in wag2 and wag2 ga1 seedlings. The GA-deficient ga1 mutant fails to form an apical hook and therefore apical hook formation requires GA treatments. In a comparison of GA-treated ga1 and wag2 ga1 mutants, we noted with interest that the normalization of the ga1 phenotype by GA was impaired in the presence of the wag2 mutation, indicating that WAG2 represses apical hook opening in dark-grown seedlings downstream from GA (Fig. 2C).

In order to gain insight into the tissue-specific expression pattern of WAG2 and to understand the divergent role of WAG1 in apical hook opening, we generated transgenic Arabidopsis lines that express the GUS reporter under control of WAG1 and WAG2 promoter fragments (2.5 kb and 2.9 kb, respectively), WAG1pro:GUS and WAG2pro:GUS. Our analysis of the WAG1 and WAG2 expression patterns indicated a specific expression of WAG2pro:GUS but not of WAG1pro:GUS in the concave side of

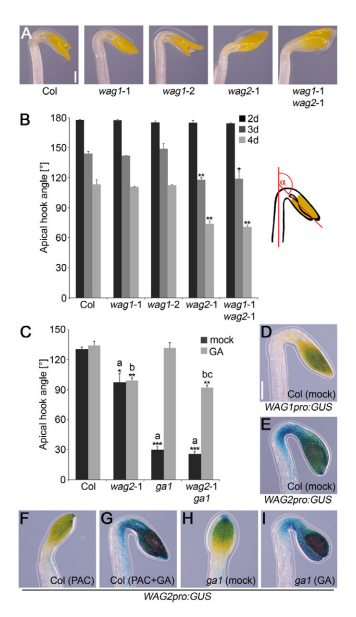


Fig. 2. Apical hook opening is accelerated in wag2 mutants. (A) Representative photographs of the apical hook of 4-day-old darkgrown seedlings of the genotypes indicated in the figure. Scale bar: 200 µm. (B) Quantification of the apical hook angle of 2-, 3- and 4-day (d)-old dark-grown seedlings of the wag1 and wag2 genotypes indicated in the figure reveals an accelerated hook opening in wag2 mutants when compared with the wild type or wag1 mutants. Shown are the averages and the standard deviations of two biological replicates using minimum 20 seedlings per genotype per experiment. Asterisks indicate the significance (Student's t-test: \*0.01<P<0.05; \*\*0.001<P<0.01) in comparison with the wild type at the specific time point. (C) Quantification of the apical hook angle in 3.5-day-old seedlings grown in the absence and presence of 1 µM GA3. Shown are the average and the standard deviation of two biological replicates (n>20). Asterisks indicate the significances (Student's t-test: \*0.01<P<0.05; \*\*0.001<P<0.01; \*\*\*P<0.001). a, b and c indicate the comparison with Col (mock), Col (GA) or ga1 (GA), respectively. (**D.E**) Photographs of representative 3-day-old dark-grown transgenic WAG1pro:GUS and WAG2pro:GUS seedlings. (F,H) WAG2pro:GUS activity following PAC treatment of the wild type (F) or in the GAdeficient ga1 background (H). (G,I) WAG2pro:GUS expression following GA treatment of PAC-treated seedlings (G) or ga1 mutants (I). Scale

bar: 200 µm.

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the apical hook (Fig. 2D,E). The expression of WAG2pro:GUS in the apical hook was GA regulated as shown after crossing WAG2pro:GUS into the GA-deficient gal background or treatment of WAG2pro:GUS lines with the GA biosynthesis inhibitor paclobutrazol (PAC) (Fig. 2F-I). WAG2pro:GUS expression thus mirrors the expression of the WAG2 gene as observed in our qRT-PCR experiments (Fig. 1) and reveals the specific expression of WAG2 in the concave side of the apical hook, which is of particular interest with regard to the phenotype of the wag2 mutant. At the same time, the apparent absence of WAG1pro:GUS expression in the apical hook, even after extended staining for GUS reporter activity, is consistent with our finding that WAG1 does not contribute to the wag2 hook phenotype.

## WAG2 is regulated by PIF5

As WAG2 expression is controlled by light and GA, we reasoned that WAG2 expression may be controlled by PIF transcription factors. This hypothesis was particularly intriguing as PIF5 had previously been reported to regulate apical hook maintenance (Gallego-Bartolomé et al., 2011; Khanna et al., 2007). We used qRT-PCR to examine the expression of WAG2 in previously characterized pif1, pif3, pif4 and pif5 single, double and quadruple mutants (Fig. 3A). This analysis revealed a strongly decreased WAG2 transcript abundance in *pif5* mutants. At the same time, *WAG2* transcript levels were increased in pif4 mutants but slightly decreased in pif4 pif5 (pif45) double mutants. These findings are consistent with the notion that PIF5 is an activator and PIF4 is a repressor of WAG2 expression. Although WAG2 expression was unaltered in pif1 and pif3 single mutants, WAG2 expression was decreased in pif1 pif3 (pif13), suggesting that PIF1 and PIF3 may be functionally redundant activators of WAG2 expression. Importantly, our analysis of an overexpressor line of PIF5 (PIF5ox) supported the concept of PIF5 being an activator of WAG2 expression (Fig. 3B). The results from the qRT-PCR analysis were confirmed when we examined the expression of WAG2pro:GUS transgenes in the pif5 and PIF5ox backgrounds, respectively (Fig. 3C). Owing to an apparent linkage between the WAG2pro:GUS transgene and the PIF5 locus, we had to use a different WAG2pro: GUS parental line (line #2) for the cross with pif5, which exhibited a lower (line 2) basal WAG2pro:GUS expression level than our WAG2pro:GUS reference line (line 1). Regardless of this fact and consistent with our qRT-PCR data, we found that WAG2pro:GUS expression in the apical hook is reduced in the pif5 mutant background and increased in PIF5ox when compared with their respective wild-type segregants (Fig. 3C).

We next examined the contribution of PIF5 to apical hook opening. In agreement with the previously published study, we detected also in our experimental conditions decreased and increased apical hook angles in *pif5* and *PIF5ox* lines, respectively (Fig. 3D; supplementary material Fig. S2). These findings are thus consistent with the notion that the reduced and increased *WAG2* transcript abundance in *PIF5* loss-of-function mutants and overexpression lines may be causative for their differential apical hook phenotype.

To examine whether *WAG2* is a direct transcription target of PIF5, we performed ChIP experiments followed by qRT-PCR with dark-grown seedlings using *PIF5ox*, which expresses an HA-tagged PIF5. Our binding site searches identified one G-box as putative PIF-binding site in the promoter of *WAG2*. Our ChIP experiment then revealed that this site is specifically bound by PIF5:HA, thus PIF5 may be a direct regulator of *WAG2* (Fig. 3E). Consistent with the differential transcriptional regulation of *WAG2* and *WAG1*, we did not find G-boxes in the promoter of *WAG1*.

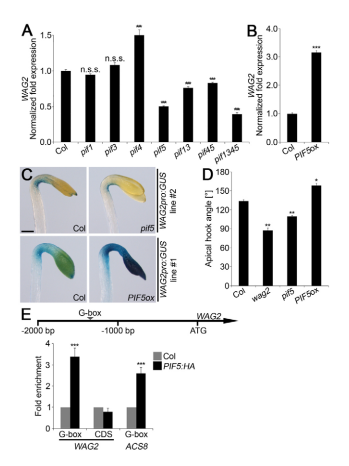


Fig. 3. WAG2 expression is PIF dependent. (A,B) qRT-PCR analysis of 3-day-old dark-grown pif mutant and PIF5 overexpression (PIF5ox) seedlings. UBC21 (A) and AT3G50685 (B) were used for normalization. (C) WAG2pro:GUS expression in wild-type and the pif5 or PIF5ox background. Owing to the apparent linkage of WAG2pro:GUS (line #1) to the PIF5 locus, a different WAG2pro:GUS transgenic line (line #2) had to be used for the cross with pif5. To account for the variability in the basal WAG2pro:GUS expression between these lines, wild-type segregants of either cross were analyzed and used for comparisons with the respective PIF5 mutant or overexpressor. (D) Apical hook phenotype of 3.5-day-old seedlings of the genotypes indicated in the figure. Shown are the average and the standard deviation of two biological replicates (n>20). (**E**) Scheme of the WAG2 promoter and ChIP of PIF5:HA from dark-grown PIF5ox Arabidopsis seedlings. A region in the WAG2-coding sequence (CDS) was used as negative control. The previously reported binding site in the ACS8 promotor was used as a positive control. Asterisks in all experiments indicate the significances in comparison with Col (Student's t-test: n.s.s., not statistically significant; \*0.01<P<0.05; \*\*0.001<P<0.01; \*\*\*P<0.001).

# WAG2 is required for a local auxin maximum in the apical hook

The analysis of mutants of the PIN auxin efflux facilitators implicated PIN1, PIN3, PIN4 and PIN7 in the maintenance of the apical hook (Zádníková et al., 2010). These analyses also identified *pin3* as the single *PIN* gene mutant with the strongest apical hook phenotype. As WAG2 is an established regulator of PIN polarity (Dhonukshe et al., 2010), a link between *WAG2* function and auxin transport regulation appeared reasonable. We therefore tested the genetic interaction between *WAG2* and *PIN3* in *wag2 pin3* double mutants and compared their apical hook phenotype with that of the single mutants. This analysis revealed an enhancement of the apical

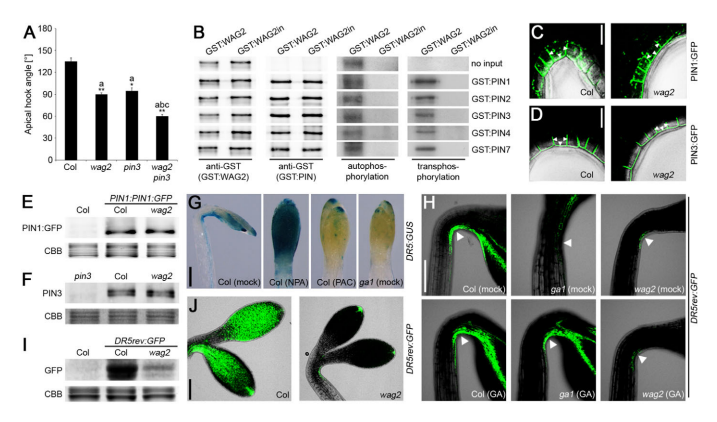


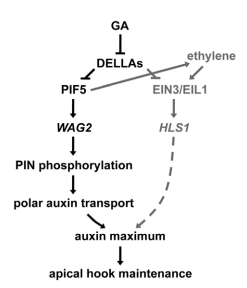
Fig. 4. WAG2 phosphorylates PIN proteins and regulates the apical hook auxin maximum. (A) Apical hook opening phenotype of 3.5-dayold seedlings. Shown is the average and the standard deviation of two biological replicates (n>20). Asterisks indicate the significances (Student's ttest: \*0.01<P<0.05; \*\*0.001<P<0.01). a, b and c indicate the comparison with Col, to wag2 or to pin3, respectively. (B) Phosphorylation experiment of recombinant GST-tagged PIN fragments by recombinant GST-tagged WAG2 or kinase-dead WAG2in. The two panels on the left represent immunoblots to demonstrate equal loading of the purified proteins. The two panels on the right are autoradiographs that reveal autophosphorylation of the active GST:WAG2 but not of GST:WAG2in and transphosphorylation of GST-tagged cytoplasmatic loops of PINs by GST:WAG2. (C,D) Confocal images of the concave apical hook region of (C) PIN1:PIN1:GFP- and (D) PIN3:PIN3:GFP-expressing, 3.5-day-old skotomorphogenic wild-type and wag2 mutants. Arrowheads indicate the polar distribution of PIN:GFP proteins. Scale bars: 25 μm. (E) Immunoblot with anti-GFP antibody of 15 µg total protein extracts prepared from the cotyledons and 1-2 mm of the apical hypocotyl of 3.5-day-old dark-grown PIN1:PIN1:GFP-expressing seedlings. (F) Immunoblot with anti-PIN3 antibody of 15 μg membrane protein extracts from the cotyledons and 1 to 2 mm of the apical hypocotyl of 3.5-day-old dark-grown seedlings. (G) Photographs of shoot apices of DR5:GUS-expressing wild-type and ga1 3.5day-old dark-grown seedlings grown, where indicated, on 1 µM NPA- or 1 µM PAC-containing media. Scale bar: 200 µm. (H) Confocal images of the apical hook of DR5rev:GFP-expressing 3.5-day-old dark-grown wild-type, ga1 and wag2 mutants. Scale bar: 200 μm. (I) Immunoblot with anti-GFP antibody of 15 µg total protein extracts prepared from the cotyledons and 1 to 2 mm of the apical hypocotyl of 3.5-day-old skotomorphogenic DR5rev:GFP-expressing seedlings. Arrowheads indicate the position of the apical hook auxin maximum. (J) Confocal images of the cotyledons of DR5rev:GFP-expressing 3.5-day-old skotomorphogenic wild-type and wag2 seedlings. Scale bar: 200 µm.

hook opening defect in the double mutant when compared to the single mutants suggesting that *WAG2* and *PIN3* act together to repress apical hook opening (Fig. 4A). As it is known that also other PINs participate in apical hook opening, it can of course be envisioned that other PINs are involved in WAG2-regulated hook development.

It has previously been reported that WAG2 phosphorylates PIN2 and thereby controls PIN2 polarity in cortical root cells (Dhonukshe et al., 2010). As PIN2 is not expressed in the apical hook (Zádníková et al., 2010) and as we had observed an enhancement of the apical hook phenotype in the *wag2 pin3* double mutant, we questioned whether WAG2 may phosphorylate PIN3 and other members of the PIN protein family besides PIN2. Using recombinant PIN fragments that correspond to their intracellular cytoplasmic loop, we could demonstrate that recombinant WAG2 phosphorylates – in addition to PIN2 – PIN1, PIN3, PIN4 and PIN7, at least in vitro, thus all PINs with an established role in cell-to-cell auxin transport and apical hook development (Fig. 4B). We

subsequently examined wild-type and wag2 apical hooks to find out whether the loss of WAG2 had a consequence on the polarity or distribution of PIN1:GFP and PIN3:GFP. This analysis did not reveal any striking differences in PIN distribution or polarity in the epidermis and cortex of the apical hook, suggesting that changes in PIN polarity may not be responsible for the wag2 apical hook phenotype (Fig. 4C,D). At the same time, our observations by epifluorescence microscopy indicated that PIN1:GFP and PIN3:GFP are strongly expressed in the inner tissue of the apical hook (supplementary material Fig. S3). Unfortunately, specifically in the apical hook region and in contrast to the remaining hypocotyl, these endodermis and stele cell layers are not amenable to confocal microscopy analysis. As we cannot image PIN proteins in this region, we are also unable to draw any conclusions on the effect of WAG2 on PIN protein behavior, e.g. polarity. To test whether changes in PIN protein abundance are the cause for the changes in apical hook formation in wag2, we also performed immunoblots to detect PIN1 and PIN3 in wild-type and wag2

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**Fig. 5. Model for the proposed role of GA in apical hook opening.** GA promotes the degradation of DELLA proteins. The DELLA repressors impair PIF5 activity, which is a transcriptional regulator of the AGCVIII kinase gene *WAG2* and of the ethylene biosynthesis gene *ACS8*. WAG2 phosphorylates PIN proteins and by this means the kinase regulates polar auxin efflux and the auxin (signaling) maximum at the concave side of the apical hook. This maximum prevents premature hook opening by impairing the elongation of cells localized at this position. Additionally, the DELLA proteins impair and ethylene promotes the transcriptional activity of EIN3/EIL1. EIN3 and EIL1 induce the expression of *HLS1*, which regulates the auxin signaling maximum in the apical hook by an unknown mechanism. Findings presented by Gallego-Bartolomé et al. (Gallego-Bartolomé et al., 2011) and An et al. (An et al., 2012) are depicted in gray.

seedlings (Fig. 4E,F). As PIN protein levels were identical in both genotypes, we concluded that changes in PIN protein abundance are not the cause for the apical hook phenotype of *wag2*.

We next examined auxin distribution in the apical hook region using the well-established DR5 auxin (response) reporters. DR5:GUS, as well as DR5rev:GFP expression analysis revealed a strong signal of the reporters in the concave side of the apical hook in dark-grown Arabidopsis seedlings (Fig. 4G,H). In agreement with previous reports, the lateral apical hook expression pattern of DR5:GUS, as well as apical hook formation were absent in seedlings that had been treated with the auxin transport inhibitor NPA or the GA biosynthesis inhibitor PAC (Fig. 4G,H) (Gallego-Bartolomé et al., 2011; Lehman et al., 1996; Zádníková et al., 2010). In addition, when we introduced DR5:GUS and DR5rev:GFP into the gal mutant, the DR5 signal for the apical hook was not visible, unless the gal mutant had been treated with GA (Fig. 4G,H). These findings confirmed the recently established observations from other laboratories showing that auxin transport and GA biosynthesis are not only required for apical hook formation but also for the formation of a lateral auxin maximum that correlates with apical hook formation (Gallego-Bartolomé et al., 2011; Zádníková et al., 2010). Importantly, our analysis of DR5rev:GFP in the wag2 mutant by confocal microscopy, as well as by immunoblotting revealed a strong reduction of the DR5:GFP signal in the concave side of the apical hook of wag2 seedlings and, as we noted with interest, also in their cotyledons (Fig. 4H-J). Interestingly, the difference in DR5rev:GFP expression was more prominent in 4-day-old seedlings than in 2-day-old seedlings

(supplementary material Fig. S4). As GA treatments did not restore or influence the DR5rev:GFP signal in the *wag2* mutant, this analysis suggests that WAG2 positively regulates the lateral auxin maximum in the apical hook (Fig. 4H). In view of the previously established function of WAG2 as an auxin transport regulatory kinase, we are tempted to speculate that this is the consequence of altered or reduced auxin transport in *wag2* mutants.

#### **DISCUSSION**

The present study reveals the role of WAG2 as a repressor of apical hook opening in dark-grown *Arabidopsis* seedlings. We show that WAG2 is specifically expressed in the concave side of the apical hook and that wag2 mutants open their apical hook faster than the wild type. At least two pieces of evidence point to a role for WAG2 in controlling auxin transport. First, we show that wag2 mutants fail to establish the strong lateral auxin maximum that is characteristic for the apical hook. Second, and as already shown in a previous study for the auxin efflux facilitator PIN2, we show that WAG2 phosphorylates all PINs – at least in vitro. As WAG2 had previously been implicated in the regulation of PIN polarity by others (Dhonukshe et al., 2010; Ding et al., 2011; Sorefan et al., 2009), we examined whether changes in PIN polarity are causative for the wag2 phenotype and the apparent changes in auxin accumulation in the apical hook. However, we could not find any evidence for a hook maintenance phase-specific or WAG2dependent change in the polarity of PIN1 and PIN3 when examining GFP-tagged variants of these two PINs. At the same time, our microscopic analysis also indicated that PIN1 and PIN3 are expressed in the inner tissue of the hypocotyl and the apical hook but, unfortunately, the inner tissues of the apical hook are not amenable to confocal studies. In addition, this has been reported in different biological contexts by others; based on our experiments, we cannot draw any conclusions on the role of WAG2 in controlling PIN protein polarity (Dhonukshe et al., 2010; Ding et al., 2011; Sorefan et al., 2009). Previously published studies have shown that auxin transport from the endodermis to the outer tissues is dynamically controlled during phototropism and gravitropism. Similar auxin tranport mechanisms may govern apical hook formation and therefore an analysis of PIN polarity in the inner tissues of the apical hook would be highly desirable but can at present not be realized. It also did not escape our attention that DR5 auxin responses are strongly reduced in wag2 cotyledons. As auxin transported from the cotyledons may be required for the establishment of the auxin maximum in the apical hook, it may well be that the loss of the apical hook auxin maximum is a consequence of the reduced auxin levels in the cotyledons that we infer from the reduced DR5rev:GFP expression in this tissue. As several reports point at a role of PIF transcription factors in the control of auxin biosynthesis and response (Franklin et al., 2011; Hornitschek et al., 2012; Li et al., 2012; Nozue et al., 2007; Sun et al., 2012), the correlation of WAG2 expression and the apparent auxin maximum in the apical hook could also be the consequence of a direct regulation of auxin distribution via the PIFs. As the local auxin maximum is strongly reduced in wag2 mutants, the maintenance of this auxin maximum should, however, be WAG2 dependent, either through the postulated role of WAG2 in controlling auxin transport or another role of WAG2 on auxin biosynthesis and signaling.

Our interest in *WAG2* had arisen from our observation that *WAG2* abundance is increased after GA treatment. In recent years, different families of transcription factors have been reported that are controlled by DELLA repressors of the GA pathway (Schwechheimer, 2011).

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We inferred from our observation that WAG2 expression is controlled by light that WAG2 is downstream from the light-labile PIF transcription factors. Comparing WAG2 expression patterns in different pif mutant backgrounds and subsequent ChIP analyses identified PIF5 as one PIF transcription factor that activates WAG2 expression downstream from GA, DELLAs and light. The role of PIF5 as a regulator of WAG2 is also supported by the apical hook phenotype of pif5 mutants and PIF5 overexpressors.

Based on the close relationship between WAG2 and WAG1, and the fact that both kinases had previously been shown to act in a functionally redundant manner in the control of root growth, we also examined the functional redundancy between the two kinases as repressors of apical hook opening. Interestingly, we found no evidence for a role of *WAG1* in apical hook formation. Most striking is our observation that *WAG1* is not expressed in the apical hook region and that the *WAG1* promoter, in contrast to the WAG2 promoter, does not contain any G-boxes, which are known binding sites for PIF transcription factors.

In summary, our results propose a model whereby PIF5, the activity of which is regulated by GA, DELLAs, and light, controls *WAG2* expression in dark-grown seedlings. We further propose that the tissue-specific expression and activity of *WAG2* in the concave side of the dark-grown seedling alters PIN polarity or PIN activity by direct PIN phosphorylation and that thereby WAG2 contributes to the regulation of a lateral auxin maximum that is required for apical hook maintenance in dark-grown seedlings (Fig. 5). Interestingly, we also observed that *WAG2* expression in the concave side of the dark-grown seedling can be detected throughout dark-grown seedling development whereas the phenotype of *wag2* mutants is restricted to the apical hook opening phase. Therefore, our results also suggest that other regulatory mechanisms must exist that control hook formation.

Importantly, DELLA proteins have most recently also been shown to repress the ethylene signaling proteins EIN3 and EIL1 (An et al., 2012). Ethylene promotes apical hook formation and *ein3 eil1* mutants are insensitive to the ethylene effect on apical hook formation. Ethylene and GA control the expression of the apical hook regulatory protein HLS1 and this control is achieved through inhibitory interactions of the GA-labile DELLAs with EIN3 and EIL1, direct regulators of *HLS1* expression. Importantly, this study also suggests that, besides the ethylene/GA-dependent regulation of *HLS1*, a GA-controlled auxin transport-dependent process must exist that modulates hook curvature through asymmetric auxin accumulation. In the context of this very recent study, the GA-dependent control of *WAG2* expression is a good candidate for this elusive auxin transport regulatory mechanism (Fig. 5).

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# **Competing interests statement**

The authors declare no competing financial interests.

### Supplementary material

Supplementary material available online at http://dev.biologists.org/lookup/suppl/doi:10.1242/dev.081240/-/DC1

# References

Achard, P., Vriezen, W. H., Van Der Straeten, D. and Harberd, N. P. (2003). Ethylene regulates arabidopsis development via the modulation of DELLA protein growth repressor function. *Plant Cell* 15, 2816-2825. Alabadí, D., Gil, J., Blázquez, M. A. and García-Martínez, J. L. (2004). Gibberellins repress photomorphogenesis in darkness. *Plant Physiol.* 134, 1050-1057

- An, F., Zhang, X., Zhu, Z., Ji, Y., He, W., Jiang, Z., Li, M. and Guo, H. (2012). Coordinated regulation of apical hook development by gibberellins and ethylene in etiolated Arabidopsis seedlings. Cell Res. 22, 915-927.
- Arnaud, N., Girin, T., Sorefan, K., Fuentes, S., Wood, T. A., Lawrenson, T., Sablowski, R. and Østergaard, L. (2010). Gibberellins control fruit patterning in Arabidopsis thaliana. *Genes Dev.* 24, 2127-2132.
- Bauer, D., Viczián, A., Kircher, S., Nobis, T., Nitschke, R., Kunkel, T., Panigrahi, K. C., Adám, E., Fejes, E., Schäfer, E. et al. (2004). Constitutive photomorphogenesis 1 and multiple photoreceptors control degradation of phytochrome interacting factor 3, a transcription factor required for light signaling in Arabidopsis. *Plant Cell* 16, 1433-1445.
- Benková, E., Michniewicz, M., Sauer, M., Teichmann, T., Seifertová, D., Jürgens, G. and Friml, J. (2003). Local, efflux-dependent auxin gradients as a common module for plant organ formation. *Cell* **115**, 591-602.
- Björklund, S., Antti, H., Uddestrand, I., Moritz, T. and Sundberg, B. (2007). Cross-talk between gibberellin and auxin in development of Populus wood: gibberellin stimulates polar auxin transport and has a common transcriptome with auxin. *Plant J.* 52, 499-511.
- Blilou, I., Xu, J., Wildwater, M., Willemsen, V., Paponov, I., Friml, J., Heidstra, R., Aida, M., Palme, K. and Scheres, B. (2005). The PIN auxin efflux facilitator network controls growth and patterning in Arabidopsis roots. *Nature* 433, 39-44
- Castillon, A., Shen, H. and Huq, E. (2009). Blue light induces degradation of the negative regulator phytochrome interacting factor 1 to promote photomorphogenic development of Arabidopsis seedlings. Genetics 182, 161-171
- Chapman, E. J. and Estelle, M. (2009). Mechanism of auxin-regulated gene expression in plants. Annu. Rev. Genet. 43, 265-285.
- Cheminant, S., Wild, M., Bouvier, F., Pelletier, S., Renou, J. P., Erhardt, M., Hayes, S., Terry, M. J., Genschik, P. and Achard, P. (2011). DELLAs regulate chlorophyll and carotenoid biosynthesis to prevent photooxidative damage during seedling deetiolation in Arabidopsis. *Plant Cell* 23, 1849-1860.
- Cheng, Y., Qin, G., Dai, X. and Zhao, Y. (2008). NPY genes and AGC kinases define two key steps in auxin-mediated organogenesis in Arabidopsis. Proc. Natl. Acad. Sci. USA 105, 21017-21022.
- Christie, J. M., Yang, H., Richter, G. L., Sullivan, S., Thomson, C. E., Lin, J., Titapiwatanakun, B., Ennis, M., Kaiserli, E., Lee, O. R. et al. (2011). phot1 inhibition of ABCB19 primes lateral auxin fluxes in the shoot apex required for phototropism. *PLoS Biol.* 9, e1001076.
- Clough, S. J. and Bent, A. F. (1998). Floral dip: a simplified method for Agrobacterium-mediated transformation of Arabidopsis thaliana. *Plant J.* 16, 735-743.
- de Lucas, M., Davière, J. M., Rodríguez-Falcón, M., Pontin, M., Iglesias-Pedraz, J. M., Lorrain, S., Fankhauser, C., Blázquez, M. A., Titarenko, E. and Prat, S. (2008). A molecular framework for light and gibberellin control of cell elongation. *Nature* **451**, 480-484.
- Dhonukshe, P., Huang, F., Galvan-Ampudia, C. S., Mähönen, A. P., Kleine-Vehn, J., Xu, J., Quint, A., Prasad, K., Friml, J., Scheres, B. et al. (2010). Plasma membrane-bound AGC3 kinases phosphorylate PIN auxin carriers at TPRXS(N/S) motifs to direct apical PIN recycling. *Development* **137**, 3245-3255.
- Ding, Z., Galván-Ampudia, C. S., Demarsy, E., Łangowski, L., Kleine-Vehn, J., Fan, Y., Morita, M. T., Tasaka, M., Fankhauser, C., Offringa, R. et al. (2011). Light-mediated polarization of the PIN3 auxin transporter for the phototropic response in Arabidopsis. *Nat. Cell Biol.* 13, 447-452.
- Feng, S., Martinez, C., Gusmaroli, G., Wang, Y., Zhou, J., Wang, F., Chen, L., Yu, L., Iglesias-Pedraz, J. M., Kircher, S. et al. (2008). Coordinated regulation of Arabidopsis thaliana development by light and gibberellins. *Nature* 451, 475-479.
- **Fode, B. and Gatz, C.** (2009). Chromatin immunoprecipitation experiments to investigate in vivo binding of Arabidopsis transcription factors to target sequences. *Methods Mol. Biol.* **479**, 261-272.
- Franklin, K. A., Lee, S. H., Patel, D., Kumar, S. V., Spartz, A. K., Gu, C., Ye, S., Yu, P., Breen, G., Cohen, J. D. et al. (2011). Phytochrome-interacting factor 4 (PIF4) regulates auxin biosynthesis at high temperature. *Proc. Natl. Acad. Sci. USA* 108, 20231-20235.
- Frigerio, M., Alabadí, D., Pérez-Gómez, J., García-Cárcel, L., Phillips, A. L., Hedden, P. and Blázquez, M. A. (2006). Transcriptional regulation of gibberellin metabolism genes by auxin signaling in Arabidopsis. *Plant Physiol.* 142, 553-563.
- Friml, J., Wiśniewska, J., Benková, E., Mendgen, K. and Palme, K. (2002). Lateral relocation of auxin efflux regulator PIN3 mediates tropism in Arabidopsis. *Nature* 415, 806-809.
- Friml, J., Vieten, A., Sauer, M., Weijers, D., Schwarz, H., Hamann, T., Offringa, R. and Jürgens, G. (2003). Efflux-dependent auxin gradients establish the apical-basal axis of Arabidopsis. *Nature* **426**, 147-153.
- Friml, J., Yang, X., Michniewicz, M., Weijers, D., Quint, A., Tietz, O., Benjamins, R., Ouwerkerk, P. B., Ljung, K., Sandberg, G. et al. (2004). A

Development 139 (21)

EVELOPMENT

- PINOID-dependent binary switch in apical-basal PIN polar targeting directs auxin efflux. *Science* **306**, 862-865.
- Fu, X. and Harberd, N. P. (2003). Auxin promotes Arabidopsis root growth by modulating gibberellin response. *Nature* 421, 740-743.
- Fujimori, T., Yamashino, T., Kato, T. and Mizuno, T. (2004). Circadian-controlled basic/helix-loop-helix factor, PIL6, implicated in light-signal transduction in *Arabidopsis thaliana*. Plant Cell Physiol. 45, 1078-1086.
- Gallego-Bartolomé, J., Minguet, E. G., Marín, J. A., Prat, S., Blázquez, M. A. and Alabadí, D. (2010). Transcriptional diversification and functional conservation between DELLA proteins in Arabidopsis. *Mol. Biol. Evol.* 27, 1247-1256.
- Gallego-Bartolomé, J., Arana, M. V., Vandenbussche, F., Zádníková, P., Minguet, E. G., Guardiola, V., Van Der Straeten, D., Benkova, E., Alabadí, D. and Blázquez, M. A. (2011). Hierarchy of hormone action controlling apical hook development in Arabidopsis. *Plant J.* 67, 622-634.
- **Galván-Ampudia, C. S. and Offringa, R.** (2007). Plant evolution: AGC kinases tell the auxin tale. *Trends Plant Sci.* **12**, 541-547.
- Grieneisen, V. A. and Scheres, B. (2009). Back to the future: evolution of computational models in plant morphogenesis. Curr. Opin. Plant Biol. 12, 606-614.
- Hornitschek, P., Kohnen, M. V., Lorrain, S., Rougemont, J., Ljung, K., López-Vidriero, I., Franco-Zorrilla, J. M., Solano, R., Trevisan, M., Pradervand, S. et al. (2012). Phytochrome interacting factors 4 and 5 control seedling growth in changing light conditions by directly controlling auxin signaling. *Plant J.* 71, 699-711.
- Huang, F., Zago, M. K., Abas, L., van Marion, A., Galván-Ampudia, C. S. and Offringa, R. (2010). Phosphorylation of conserved PIN motifs directs Arabidopsis PIN1 polarity and auxin transport. *Plant Cell* 22, 1129-1142.
- Huq, E., Al-Sady, B., Hudson, M., Kim, C., Apel, K. and Quail, P. H. (2004). Phytochrome-interacting factor 1 is a critical bHLH regulator of chlorophyll biosynthesis. *Science* 305, 1937-1941.
- Jacobs, W. P. and Case, D. B. (1965). Auxin transport, gibberellin, and apical dominance. Science 148, 1729-1731.
- Khanna, R., Shen, Y., Marion, C. M., Tsuchisaka, A., Theologis, A., Schäfer, E. and Quail, P. H. (2007). The basic helix-loop-helix transcription factor PIF5 acts on ethylene biosynthesis and phytochrome signaling by distinct mechanisms. *Plant Cell* 19, 3915-3929.
- King, K. E., Moritz, T. and Harberd, N. P. (2001). Gibberellins are not required for normal stem growth in Arabidopsis thaliana in the absence of GAI and RGA. *Genetics* 159, 767-776.
- Kleine-Vehn, J. and Friml, J. (2008). Polar targeting and endocytic recycling in auxin-dependent plant development. Annu. Rev. Cell Dev. Biol. 24, 447-473.
- Kuhn, H. and Galston, A. W. (1992). Physiological asymmetry in etiolated pea epicotyls: relation to patterns of auxin distribution and phototropic behavior. *Photochem. Photobiol.* 55, 313-318.
- Lehman, A., Black, R. and Ecker, J. R. (1996). HOOKLESS1, an ethylene response gene, is required for differential cell elongation in the Arabidopsis hypocotyl. *Cell* 85, 183-194.
- Leivar, P. and Quail, P. H. (2011). PIFs: pivotal components in a cellular signaling hub. *Trends Plant Sci.* 16, 19-28.
- Leivar, P., Monte, E., Al-Sady, B., Carle, C., Storer, A., Alonso, J. M., Ecker, J. R. and Quail, P. H. (2008). The Arabidopsis phytochrome-interacting factor PIF7, together with PIF3 and PIF4, regulates responses to prolonged red light by modulating phyB levels. Plant Cell 20, 337-352.
- Li, H., Johnson, P., Stepanova, A., Alonso, J. M. and Ecker, J. R. (2004). Convergence of signaling pathways in the control of differential cell growth in Arabidopsis. Dev. Cell 7, 193-204.
- Li, L., Ljung, K., Breton, G., Schmitz, R. J., Pruneda-Paz, J., Cowing-Zitron, C., Cole, B. J., Ivans, L. J., Pedmale, U. V., Jung, H. S. et al. (2012). Linking photoreceptor excitation to changes in plant architecture. *Genes Dev.* 26, 785-790.
- Lorrain, S., Allen, T., Duek, P. D., Whitelam, G. C. and Fankhauser, C. (2008). Phytochrome-mediated inhibition of shade avoidance involves degradation of growth-promoting bHLH transcription factors. *Plant J.* 53, 312-323.
- McGinnis, K. M., Thomas, S. G., Soule, J. D., Strader, L. C., Zale, J. M., Sun, T. P. and Steber, C. M. (2003). The Arabidopsis SLEEPY1 gene encodes a putative F-box subunit of an SCF E3 ubiquitin ligase. *Plant Cell* 15, 1120-1130.
- Monte, E., Tepperman, J. M., Al-Sady, B., Kaczorowski, K. A., Alonso, J. M., Ecker, J. R., Li, X., Zhang, Y. and Quail, P. H. (2004). The phytochrome-interacting transcription factor, PIF3, acts early, selectively, and positively in light-induced chloroplast development. *Proc. Natl. Acad. Sci. USA* 101, 16091-16098.
- Nozue, K., Covington, M. F., Duek, P. D., Lorrain, S., Fankhauser, C., Harmer, S. L. and Maloof, J. N. (2007). Rhythmic growth explained by coincidence between internal and external cues. *Nature* 448, 358-361.

- Oh, E., Yamaguchi, S., Hu, J., Yusuke, J., Jung, B., Paik, I., Lee, H. S., Sun, T. P., Kamiya, Y. and Choi, G. (2007). PIL5, a phytochrome-interacting bHLH protein, regulates gibberellin responsiveness by binding directly to the GAI and RGA promoters in Arabidopsis seeds. Plant Cell 19, 1192-1208.
- Oh, E., Kang, H., Yamaguchi, S., Park, J., Lee, D., Kamiya, Y. and Choi, G. (2009). Genome-wide analysis of genes targeted by PHYTOCHROME INTERACTING FACTOR 3-LIKE5 during seed germination in Arabidopsis. *Plant Cell* 21, 403-419.
- Park, E., Kim, J., Lee, Y., Shin, J., Oh, E., Chung, W. I., Liu, J. R. and Choi, G. (2004). Degradation of phytochrome interacting factor 3 in phytochrome-mediated light signaling. *Plant Cell Physiol.* 45, 968-975.
- Peng, J., Carol, P., Richards, D. E., King, K. E., Cowling, R. J., Murphy, G. P. and Harberd, N. P. (1997). The Arabidopsis GAI gene defines a signaling pathway that negatively regulates gibberellin responses. *Genes Dev.* 11, 3194-3205
- Raz, V. and Ecker, J. R. (1999). Regulation of differential growth in the apical hook of Arabidopsis. *Development* **126**, 3661-3668.
- Richter, R., Behringer, C., Müller, I. K. and Schwechheimer, C. (2010). The GATA-type transcription factors GNC and GNL/CGA1 repress gibberellin signaling downstream from DELLA proteins and PHYTOCHROME-INTERACTING FACTORS. Genes Dev. 24, 2093-2104.
- Santner, A. A. and Watson, J. C. (2006). The WAG1 and WAG2 protein kinases negatively regulate root waving in Arabidopsis. *Plant J.* 45, 752-764.
- Schwechheimer, C. (2011). Gibberellin signaling in plants—the extended version. Front. Plant Sci. 2, 107.
- Scott, T. K., Case, D. B. and Jacobs, W. P. (1967). Auxin-gibberellin interaction in apical dominance. *Plant Physiol.* **42**, 1329-1333.
- Shen, H., Moon, J. and Huq, E. (2005). PIF1 is regulated by light-mediated degradation through the ubiquitin-26S proteasome pathway to optimize photomorphogenesis of seedlings in Arabidopsis. *Plant J.* 44, 1023-1035.
- Sorefan, K., Girin, T., Liljegren, S. J., Ljung, K., Robles, P., Galván-Ampudia, C. S., Offringa, R., Friml, J., Yanofsky, M. F. and Østergaard, L. (2009). A regulated auxin minimum is required for seed dispersal in Arabidopsis. *Nature* 459, 583-586.
- Sun, J., Qi, L., Li, Y., Chu, J. and Li, C. (2012). PIF4-mediated activation of YUCCA8 expression integrates temperature into the auxin pathway in regulating Arabidopsis hypocotyl growth. *PLoS Genet.* **8**, e1002594.
- Teale, W. D., Paponov, I. A. and Palme, K. (2006). Auxin in action: signalling, transport and the control of plant growth and development. Nat. Rev. Mol. Cell Biol. 7, 847-859.
- **Ueguchi-Tanaka, M., Nakajima, M., Motoyuki, A. and Matsuoka, M.** (2007). Gibberellin receptor and its role in gibberellin signaling in plants. *Annu. Rev. Plant Biol.* **58**, 183-198.
- Ulmasov, T., Murfett, J., Hagen, G. and Guilfoyle, T. J. (1997). Aux/IAA proteins repress expression of reporter genes containing natural and highly active synthetic auxin response elements. Plant Cell 9, 1963-1971.
- Vandenbussche, F., Petrásek, J., Zádníková, P., Hoyerová, K., Pesek, B., Raz, V., Swarup, R., Bennett, M., Zazímalová, E., Benková, E. et al. (2010). The auxin influx carriers AUX1 and LAX3 are involved in auxin-ethylene interactions during apical hook development in Arabidopsis thaliana seedlings. *Development* 137, 597-606.
- Vriezen, W. H., Achard, P., Harberd, N. P. and Van Der Straeten, D. (2004). Ethylene-mediated enhancement of apical hook formation in etiolated Arabidopsis thaliana seedlings is gibberellin dependent. *Plant J.* **37**, 505-516.
- Willige, B. C., Ghosh, S., Nill, C., Zourelidou, M., Dohmann, E. M., Maier, A. and Schwechheimer, C. (2007). The DELLA domain of GA INSENSITIVE mediates the interaction with the GA INSENSITIVE DWARF1A gibberellin receptor of Arabidopsis. *Plant Cell* 19, 1209-1220.
- Willige, B. C., Isono, E., Richter, R., Zourelidou, M. and Schwechheimer, C. (2011). Gibberellin regulates PIN-FORMED abundance and is required for auxin transport-dependent growth and development in Arabidopsis thaliana. *Plant Cell* **23**, 2184-2195.
- Wisniewska, J., Xu, J., Seifertová, D., Brewer, P. B., Ruzicka, K., Blilou, I., Rouquié, D., Benková, E., Scheres, B. and Friml, J. (2006). Polar PIN localization directs auxin flow in plants. *Science* **312**, 883.
- Zádníková, P., Petrásek, J., Marhavy, P., Raz, V., Vandenbussche, F., Ding, Z., Schwarzerová, K., Morita, M. T., Tasaka, M., Hejátko, J. et al. (2010). Role of PIN-mediated auxin efflux in apical hook development of Arabidopsis thaliana. *Development* 137, 607-617.
- Zourelidou, M., Müller, I., Willige, B. C., Nill, C., Jikumaru, Y., Li, H. and Schwechheimer, C. (2009). The polarly localized D6 PROTEIN KINASE is required for efficient auxin transport in Arabidopsis thaliana. *Development* 136, 627-636.

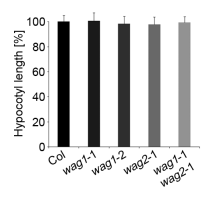
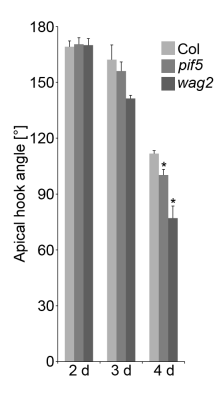


Fig. S1. WAG1 and WAG2 do not regulate hypocotyl elongation in the dark. Hypocotyl lengths of 4-day-old skotomorphogenic seedlings. Lengths are expressed relative to the wild-type control ( $n \ge 32$ ).



**Fig. S2 Apical hook opening of** wag2 and pif5 single mutants. Quantification of the apical hook angle of 2, 3 and 4 day (d)-old dark-grown seedlings of the wild-type and pif5 as well as wag2 mutants. Shown are the averages and the standard deviations of two biological replicates using minimum 20 seedlings per genotype per experiment. Asterisks indicate the significance (Student's t-test: \*0.01 < P < 0.05) in comparison with the wild type at the specific time point.

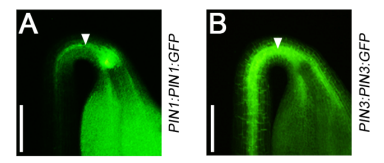


Fig. S3. Expression domains of PIN1:GFP and PIN3:GFP in the apical hook. Epifluorescence images of the apical hooks of PIN1:PIN1:GFP ( $\bf A$ ) and PIN3:PIN3:GFP ( $\bf B$ ) -expressing 3.5-day-old *Arabidopsis* seedlings. Arrowheads indicate the stele of the apical hook. Note that these strong fluorescence signals in the inner tissues were not detectable using confocal microscopy. Scale bar: 200  $\mu m$ .

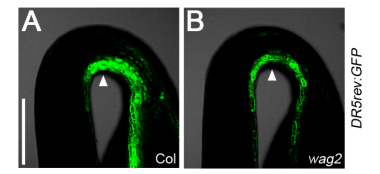


Fig. S4. Confocal images of the cotyledons of *DR5rev:GFP*-expressing 2-day-old skotomorphogenic wild-type and *wag2* seedlings. Arrowheads mark the expression maximum in the apical hook. Scale bar: 200 µm.

Table S1. List or primers used in this study

ChIP			
AT4G37770	ACS8	ACS8 FP	ATCGGAAATTCACATCGTGCCTA
		ACS8 RP	GATGTCAGAGAAGAATGAGCACGT
AT1G49240	ACT8	ACT8 LP	GCAGCATGAAGATTAAGGTCGTG
		ACT8 RP	TGTGGACAATGCCTGGACCTGCT
AT3G14370	WAG2	WAG2 Gboxl LP	GGCCCACAGATATGTGATTAG
		WAG2 Gboxl RP	TCAAAATATTATGAGGGATTCGC
AT3G14370	WAG2	WAG2 CDS 3a	GCGAATCTTGCGGAAGAGTCACG
		WAG2 CDS 5a	CTGACACCGATCTTGATCTCAGC
Genotyping			
LBb1.3		LBb1.3	ATTTTGCCGATTTCGGAAC
AT1G70940	PIN3	pin3-4 LP	TGCCACCTTCAATTCAAAAAC
		pin3-4 RP	TGATTTCTTGAGACCGATGC
AT1G53700	WAG1	WAG1-Asa	CTATCAAATCCTTGGCTTCAACC
		WAG1-Sa	CGATCTCAGCTTCACCTCCACAG
AT3G14370	WAG2	WAG2 5a	CTGACACCGATCTTGATCTCAGC
		WAG2-Asa	CCGTTACAGGCTCTGCCGCAAAC
GST constructs			
GST:WAG2(in)		attB1 WAG2	GGGGattB1TCATGGAACAAGAAGATTTCTATTTC
		attB2 WAG2	GGGGattB2TTAAACGCGTTTGCGACTCGC
GST:WAG2in		WAG2mut5a	GATTCGCTTTAGAGGTCATCGACCG
		WAG2mut3a	CGGTCGATGACCTCTAAAGCGAATC
GST:PIN1		PIN1LOOP FW	GGGG <i>attB1</i> GAGTACCGTGGAGCTAAGCTTTTG
		PIN1LOOP RV	GGGG <i>attB2</i> AGTTGGGATTACGAATAAGTTTC
GST:PIN2		PIN2LOOP FW	GGGG <i>attB1</i> GAGTTCCGTGGGGCTAAGCTTCTCATCTC
		PIN2LOOP RV	GGGG <i>attB2</i> AAGGGTTTCGAATGAGTTTTCTCAAACCCA
GST:PIN3		PIN3LOOP FW	GGGG <i>attB1</i> TTTCGTGGCGCCAAGATGCTC
		PIN3LOOP RV	GGGG <i>attB2</i> TAAGTGTTTGGGTTTCTGATGAGTTTC
GST:PIN4		PIN4LOOP FW	GGGG <i>attB1</i> TCGAGTACCGTGGCGCTAAGCTTCT
		PIN4LOOP RV	GGGG <i>attB2</i> CTGCAGTCAGTTTGGGTTTCTGATCAGCTTT
GST:PIN7		PIN7LOOP FW	GGGG <i>attB1</i> TCGAATACGAATACAGAGGAGCTAAGATCTTG
		PIN7LOOP RV	GGGG <i>attB2</i> TTAGTTTGGGTTTCTTATGAGTTTCCTC
Promoter GUS			
WAG1pro:GUS	1	WAG1+7	CTTCCATGGTTCCGGTGAA
		WAG1-7	TAAAATAATTATACTATGGAATTCCACATA
WAG2pro:GUS		WAG1-2303 WAG2+13	CTTCTTGTTCCATGGTTTTCTTCTTG
		WAG2-19	CGTAATATAAAAGAATTCTAACTAAG
Quantitative real-time PCR			
AT2G38120	AUX1	AUX1 3a	CATGCATAATCTCAACAGTAAC
		AUX1 5a	CCAGTAACATTTATTACATAAACG

AT5G55910	D6PK	D6PK 3a	CGAATTCTTCGACAAGCCTTCGG
		D6PK 5a	GTCCTGGTGGTGATTTGCATAC
AT4G26610	D6PKL1	D6PKL1 3a	CACATGTCCATCATCTCTAACAAG
		D6PKL1 5a	GCAATGAAGGTTATGGACAAAGG
AT5G47750	D6PKL2	D6PKL2 3a	CAGAGAGCATTATATGTCCGTC
		D6PKL2 5a	CTTTGGATCATCCATTTCTCCC
AT3G27580	D6PKL3	D6PKL3 3a	CCATAGCAAGAAGAACTTCAGC
		D6PKL3 5a	GCGAGGAAGAGCTTGTTAGAGC
AT3G50685		HKG 2step-LP	TTTAATCGGAGCGTTGGAAG
		HKG 2step-RP	TACAAAGACCAGCCCACGAT
AT5G01240	LAX1	LAX1 3b	CAGCCCATCAAGCACTTCAAACC
		LAX1 5b	GACGCCTGGTTTAGCTGTGCATC
AT2G21050	LAX2	LAX2 3a	GAACCTCAAACCACTGAATGAC
		LAX2 5a	CTAAGCTATCTGACATGTTTTGG
AT1G77690	LAX3	LAX3 3b	GGTGTAGACGCGAATCCGAACG
		LAX3 5b	GGTGCTTTACTTCACCGGAGCC
AT2G36910	PGP1	PGP1 3a	CGCCATGTAATGGATGAAATTAC
		PGP1 5a	GATGATGGAAGAAGTTCTCAAG
AT2G47000	PGP4	PGP4 3a	CCTCCTACAAATGTTGCTAGAAG
		PGP4 5a	CAAAGTCTCCAAAGTTGCTCTG
AT3G28860	PGP19	PGP19 3a	CAAGCTGCGAGAGGCCAAATAG
		PGP19 5a	GTCCTCGCTAACTTTGCTCAGC
AT3G45780	PHOT1	PHOT1 3a	GTTGGCATCAGGAAGTTCTCG
		PHOT1 5a	CTACAAGGTCCAGAGACTGATC
AT5G58140	PHOT2	PHOT2 3a	GATGCACGCTCGGTGAGCCTTG
		PHOT2 5a	GAGCTTCCAGATGCTAATACGC
AT2G34650	PID	PID 3a	GTCTAGCGAGACGAGTGAATCG
		PID 5a	CTCTCTCCGTCATAGACAACCTC
AT2G26700	PID2	PID2 3a	CCATCATGTGGAGATACTCTAAGG
		PID2 5a	CATCGGAAGTGTGTACCTCTGCC
AT1G73590	PIN1	PIN1 2stp 3	TCATCGTCTTTGTTACCGAAACT
		PIN1 2stp 5	CCTCCAGGGGAATAGTAACGACA
AT5G57090	PIN2	PIN2 2stp 3	GGTGGGTACGACGGAACA
		PIN2 2stp 5	GGCGAAGAAGCAGGAAGA
AT1G70940	PIN3	PIN3 2stp 3	CCGGCGAAACTAAATTGTTG
		PIN3 2stp 5	CCCAGATCAATCTCACAACG
AT2G01420	PIN4	PIN4 2stp 3	ATCAAGACCGCCGATATCAT
		PIN4 2stp 5	TTGTCTCTGATCAACCTCGAAA
AT1G23080	PIN7	PIN7 2stp 3	TCACCCAAACTGAACATTGC
		PIN7 2stp 5	TGGGCTCTTGTTGCTTTCA
AT5G25760	UBC21	UBC21 2step-LP	TCCTCTTAACTGCGACTCAGG
		UBC21 2step-RP	GCGAGGCGTGTATACATTTG
AT1G53700	WAG1	WAG1 2stp 3	AGATACTCCAAGGCGACGAG
		WAG1 2stp 5	GTTACCGATTTCCCCGGTTA
AT3G14370	WAG2	WAG2 2stp 3	CGAGGAGGCGAATGTACG
		WAG2 2stp 5	GACACCGATCTTGATCTCAGC