phot1 Inhibition of ABCB19 Primes Lateral Auxin Fluxes in the Shoot Apex Required For Phototropism

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Abstract

It is well accepted that lateral redistribution of the phytohormone auxin underlies the bending of plant organs towards light. In monocots, photoreception occurs at the shoot tip above the region of differential growth. Despite more than a century of research, it is still unresolved how light regulates auxin distribution and where this occurs in dicots. Here, we establish a system in Arabidopsis thaliana to study hypocotyl phototropism in the absence of developmental events associated with seedling photomorphogenesis. We show that auxin redistribution to the epidermal sites of action occurs at and above the hypocotyl apex, not at the elongation zone. Within this region, we identify the auxin efflux transporter ATP-BINDING CASSETTE B19 (ABCB19) as a substrate target for the photoreceptor kinase PHOTOTROPIN 1 (phot1). Heterologous expression and physiological analyses indicate that phosphorylation of ABCB19 by phot1 inhibits its efflux activity, thereby increasing auxin levels in and above the hypocotyl apex to halt vertical growth and prime lateral fluxes that are subsequently channeled to the elongation zone by PIN-FORMED 3 (PIN3). Together, these results provide new insights into the roles of ABCB19 and PIN3 in establishing phototropic curvatures and demonstrate that the proximity of light perception and differential phototropic growth is conserved in angiosperms.

Introduction

Plants have evolved numerous ways to optimize photosynthetic light capture. Phototropism, the reorientation of growth towards light, is one of the most important of these adaptive processes [1]. Originally identified in monocot coleoptiles by Charles and Francis Darwin, phototropism is initiated by light perceived at the shoot tip generating a diffusible signal that influences differential elongation in the tissues below [2]. Subsequent studies have since shown that phototropism arises from increased growth on the shaded side of the stem [3], owing to an accumulation of the phytohormone auxin [4]. The role of PHOTOTROPIN (phot) blue-light receptors in establishing phototropic curvatures in the model dicot Arabidopsis thaliana is well established [5], as is evidence supporting auxin accumulation on the shaded side of photostimulated hypocotyls [6,7].

To date, little is known regarding the molecular events that transduce photoreceptor activation into auxin redistribution across the growing stem. Genetic analyses in Arabidopsis have identified several auxin transporter families [4]. Of these, members of the PIN-FORMED family, named after the influence phenotype of the pin1 mutant, are the primary mediators of directional auxin fluxes that regulate plant development [8]. PIN3 is proposed to mediate lateral auxin fluxes by differentially restricting auxin to the vascular cylinder [6]. Consistent with this mode of action, PIN3 exhibits a subcellular localization on the inner side of bundle sheath cells [6]. However, PIN3 also functions in apical hook opening [9], which may contribute to the delayed phototropism observed in etiolated pin3 seedlings [6,10].

Another transporter implicated in phototropism is ATP-BINDING CASSETTE B19 (ABCB19/MDR1/PGP19; herein abbreviated B19), which transports auxin out of the shoot apex and maintains long-distance auxin transport streams primarily by preventing cellular reuptake and diffusion into cells adjoining vascular tissues [11,12]. B19 is predominantly localized apolarily at the plasma membrane [12–14] and functions coordinately with PIN1 to mediate polar auxin flow from the shoot apex to the roots [15]. Additionally, b19 mutants exhibit enhanced phototropic curvature [10,15] that has been attributed to auxin accumulation in the hypocotyl elongation zone as a consequence of decreased...
Plants depend on sunlight for photosynthesis and adapt their growth to optimize light capture. Phototropism, the reorientation of growth towards light, is one important adaptive response. Modern studies of phototropism began with experiments in monocotyledonous grasses by Charles Darwin and led ultimately to the discovery of the plant growth hormone auxin, establishing the concept that light perception at the shoot apex triggers differential bending in the tissues below. In the past two decades, molecular-genetic analysis in the model flowering plant Arabidopsis thaliana has identified the principle photoreceptor for phototropism, phot1, as well as the major auxin transporters. Despite extensive efforts, how the photoreceptor regulates auxin transport so as to establish differential growth is still poorly understood, as is whether this process is conserved between monocots and dicots. Here, we introduce a new approach to the study of Arabidopsis phototropism in the absence of developmental events associated with seedling photomorphogenesis. In doing so, we show that the proximity of light perception and differential growth is conserved between monocots and dicots: in both plant types, differential growth is a consequence of lateral auxin movements across the shoot apex. Moreover, we identify two auxin transporters, PIN3 and ABCB19, that contribute to these movements, the latter serving to prime lateral auxin fluxes in the shoot apex. ABCB19 function is regulated by phot1, identifying it as a substrate for this class of photoreceptor kinase.

**Results**

**Hypocotyl Phototropism in Arabidopsis Following Dark Acclimation**

Phototropism to directional blue light is traditionally examined using etiolated seedlings. However, studies performed in this manner examine the combined processes of phototropism and seedling photomorphogenesis. For the experiments described here, it was desirable to discriminate auxin transport processes associated with phototropism from those involved in seedling photomorphogenesis. To isolate phototropic responses from developmental events, 3-d-old light-grown seedlings were subjected to a 24-h period of dark acclimation prior to phototropic stimulation. Light-dependent autophosphorylation of phot1 and phot2 was readily detectable following dark acclimation (Figure 1A), as was PIN1-B19-mediated polar transport [10,15]. A role for B19 in seedling photomorphogenesis has also been reported that is dependent on CRYPTOCHROME (cry) and PHYTOCHROME (phy) photoreceptors [16,17], both of which are known to impact phototropism in Arabidopsis [18,19].

We therefore sought to clarify the roles of B19 and PIN3 in establishing phototropic curvatures by discriminating auxin transport processes associated with phototropism from those involved in seedling photomorphogenesis. To address this, we employed an unconventional strategy to analyze phototropic responses in Arabidopsis post-photomorphogenesis by subjecting light-grown seedlings to dark acclimation. By adopting this revised experimental approach, we uncover new mechanistic information regarding the roles of B19 and PIN3 in effecting phototropism and demonstrate a direct link between photoreceptor activation and auxin transporter function. Furthermore, we provide clear evidence that lateral auxin fluxes required for phototropism in Arabidopsis seedlings are initiated in and above the hypocotyl apex and not in the region of differential growth, analogous to the auxin transport mechanisms proposed for monocot phototropism almost a century ago.

Figure 1. Phototropic response of dark-acclimated Arabidopsis. (A) Autophosphorylation activities for phot1 and phot2 in membranes isolated from wild-type (WT) and phot1-5 phot2-1 seedlings. Protein extracts were given a mock irradiation (darkness, D) or irradiated with 10,000 μmol m⁻² s⁻¹ white light (L) in the presence of radiolabeled ATP. (B) Phototropic response of seedlings (upper left) exposed to directional blue light (BL, 1 μmol m⁻² s⁻¹) for 12 h (upper right). Also shown is the phototropic bending of b19-1 (lower left) and pin3-4 (lower right) to blue light for 12 h and 24 h, respectively. (C) Phototropic time course for wild-type, phot1-5, b19-1, B19 overexpressing (b19oe), and pin3-4 seedlings. Hypocotyl deviation from the vertical was calculated (inset) by subtracting the measured angle (θ) from the vertical to determine the total angle of deviation (indicated by the dotted lines). Results represent the mean ± standard error, n = 10 seedlings. (D) Vertical hypocotyl growth within the first hour after treatment with directional blue light (1 μmol m⁻² s⁻¹) or continued darkness. Results represent the mean ± SD, n = 10 seedlings in three independent experiments. Asterisks represent significant differences from wild type among treatments (p < 0.05, t test).

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reproducible arching of the photostimulated hypocotyl along the vertical agar surface (Figure 1B). Hence, dark acclimation can be used to restore phototropism to its ground state, thereby offering a means to study hypocotyl phototropism after photomorphogenesis.

After dark acclimation, phototropism within the first 12 h of treatment with low fluence rate blue light (1 μmol m⁻² s⁻¹) was attributed solely to the action of phot1 (Figure 1C), as observed in etiolated seedlings under equivalent light conditions [20].

Similarly, hypocotyls of b19 mutants showed enhanced bending relative to wild type that was evident as early as 2–4 h after treatment (Figure 1C). Seedlings overexpressing B19 displayed reduced initial rates of curvature (Figure 1C) consistent with the expanded auxin efflux activity in these transgenic lines [21]. In contrast to etiolated seedlings (Figure S1A), bending of dark-acclimated pan3 hypocotyls was initially enhanced (Figure 1C), but was less than that of wild type when photostimulated beyond 12 h in both dark-grown and dark-acclimated seedlings (Figure S1B).

However, bending in pan3 mutants occurred at a higher position of the hypocotyl compared to wild type or b19 mutants (Figure 1B), suggesting that PIN3 is spatially important for establishing phototropic curvature in dark-acclimated seedlings.

Prior to bending, hypocotyls of dark-acclimated seedlings showed reduced vertical growth in response to directional blue light, as reported for etiolated Arabidopsis seedlings [22] and rice coleoptiles [23], which was dependent on phot1 and not cry or phy photoreceptors (Figure 1D). Hypocotyl growth inhibition was still apparent in pan3 mutants, whereas mutants lacking B19 showed comparable growth rates in dark and light but were reduced in comparison to phot1 mutants (Figure 1D). Pooling of auxin in the upper hypocotyl region, including the cotyledonary node, through an interruption of B19 function rather than PIN3 would concur with the observed hypocotyl growth arrest that precedes phototropism.

**phot1 Activation Reduces Polar Auxin Transport**

To determine whether the observed differences in vertical hypocotyl growth rates correlate with alterations in polar auxin flow, transport of the principle auxin indole-3-acetic acid (IAA) from the shoot apex was monitored in the presence or absence of directional blue light. Such an approach is not feasible with etiolated Arabidopsis seedlings since they lack a fully exposed shoot apex. ³H-IAA was applied to the shoot apex of dark-acclimated seedlings under safe light conditions. Seedlings were either retained in darkness or exposed to directional blue light (1 μmol m⁻² s⁻¹), and sections corresponding to the upper hypocotyl/cotyledonary node and the mid hypocotyl/elongation zone were harvested for radiotracer analysis. Measurements of ³H-IAA in these sections confirmed that directional blue light inhibits polar flow from the apical region (Figure 2A) to the mid hypocotyl (Figure 2B), resulting in IAA accumulation within the apical region. Moreover, blue-light-dependent inhibition of polar auxin transport was dependent on phot1, as phot1 mutants lacked this response (Figure 2A and 2B).

Apical auxin retention by phot1 showed little dependence on PIN3, PIN1, and ABCB1 (abbreviated to B1), a second member of the ABCB transporter family, while a loss of B19 function increased retention (Figure 2A and 2B). Conversely, overexpression of B19 reduced apical auxin (Figure S2). These findings suggest that B19 is a likely target of phot1 action to retain auxin within the upper hypocotyl within the first hour of blue-light exposure.

Quantification of endogenous IAA levels by gas chromatography–mass spectrometry showed a trend similar to that of the radiotracer studies (Figure 3A). phot1 mediated modest increases in

**3H-IAA accumulation in the apical tissues of dark-acclimated seedlings that included the upper hypocotyl, petioles, and cotyledons (Figure 3A). However, endogenous IAA levels were actually lower in the apical tissues of b19 seedlings after cotyledon removal (Figure 3B), whereas the levels of the IAA catabolism products oxindole-3-acetic acid (oxIAA) and oxIAA-glucose (oxIAA-Glc) were increased (Figure 3C). These findings suggest that sustained pooling of IAA in the shoot apex of b19 mutants is sufficient to induce its catabolic activity [24]. The enhanced rates of phototropism typically associated with b19 mutants most likely involve more dynamic or localized auxin accumulations than previously thought.**

**phot1 Interacts with B19**

To ascertain whether B19 represents a component of phot1 signaling, we performed co-immunoprecipitation using Arabidopsis expressing functional PHOT1/phot1-green fluorescent protein [GFP] [25]. B19 was found to co-immunoprecipitate specifically with phot1-GFP in darkness (Figure 4A). B19-phot1 interactions were also verified by mass spectrometry analysis of phot1-GFP and B19–hemagglutinin (HA) immunoprecipitates (Tables S1 and S2).

In vivo irradiation prior to immunoprecipitation of phot1-GFP
 attenuated its interaction with B19, implying that this interaction was transient upon phot1 kinase activation (Figure 4A), a property commonly associated with kinase-substrate interactions. A direct interaction between B19 and phot1 was further confirmed by bimolecular fluorescence complementation (BiFC). Reconstitution of a yellow fluorescent protein (YFP) signal at the plasma membrane was evident in tobacco epidermal cells co-expressing phot1 and B19 (Figure 4B), whereas no fluorescence was detected when either phot1 or B19 was expressed with empty vector controls (Figure S3).

The cytosolic C-terminus of B19 (B19C), encompassing its second nucleotide-binding domain (NBD), is important for its transporter activity and for mediating protein interactions [13,26]. We found that B19C interacts with the C-terminus of phot1, including its Ser/Thr kinase domain, both in yeast (Figure 4C) and in vitro co-immunoprecipitation assays (Figure 4D). Furthermore, domain mapping showed that the catalytic kinase subunit of phot1 was necessary for this interaction (Figure 4C). These data therefore suggest that B19 may function as a substrate for phot1 kinase activity. If this is the case, then substrate phosphorylation would be expected to be limited to B19 as no interaction was observed between phot1 kinase and the C-terminus of B1 in yeast (Figure 4C), despite its expression (Figure S4).

**phot1 Inhibits B19 Transporter Activity**

Our attempts to express functional phot1 and B19 simultaneously in either insect cells or *Schizosaccharomyces pombe* were unsuccessful, precluding the possibility of investigating substrate phosphorylation by co-expression analysis in these systems. As an alternative, functional B19 generated in membranes of *S. pombe* was mixed with active phot1 enriched from insect cells for in vitro phosphorylation analysis. Although the addition of *S. pombe* membranes reduced the level of phot1 autophosphorylation from insect cells, increased phosphorylation of B19 in the presence of phot1 was apparent using this approach (Figure 5A). Phosphorylation of B19 in the presence of phot1 was verified by immunoblot analysis (Figure 5B), suggesting that B19 is a substrate target for phot1 kinase activity.

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**Figure 3. IAA accumulation in dark-acclimated seedlings.** (A) Free IAA levels in wild-type (WT), phot1–5, and b19–1 seedlings. IAA measurements were determined as described [43] after 2.5 h directional blue light (BL, 1 μmol m$^{-2}$ s$^{-1}$) or continued darkness (D). Apical tissues including the upper hypocotyl, petioles, and cotyledons were excised for analysis, in addition to the mid hypocotyl (MH). (B) Free IAA determinations as in (A), but after cotyledon excision. Apical tissues including the upper hypocotyl and cotyledonary node with the lower half of the petioles were excised for analysis. (C) Accumulation of auxin catabolites oxIAA and oxIAA-Glc in dark-acclimated wild-type and b19–1 seedlings. Upper hypocotyls/cotyledonary nodes were excised and assayed. In each case, results represent the mean ± SD, n = 100 seedlings in three independent experiments.

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**Figure 4. phot1 interacts with B19.** (A) phot1-GFP and the plasma membrane marker fusion GST-Lti6b were immunoprecipitated from membranes from 3-d-old etiolated seedlings (D) or seedlings exposed to blue light (BL, 20 μmol m$^{-2}$ s$^{-1}$) for 5 min. Samples were subjected to immunoblot analysis with anti-GFP and anti-B19 antibodies. Input (left) represents solubilized microsomes used for immunoprecipitation (IP, right). (B) BiFC fluorescence images of phot1-YN and B19-YC co-expressed in tobacco leaf epidermis. Reconstitution of YFP fluorescence was visible at the plasma membrane. Scale bar = 20 μm. (C) The C-terminus of phot1 including its Ser/Thr kinase domain (amino acids 612–996) interacts with B19C (amino acids 1121–1252) in yeast. Growth on minimal selection medium (MS) selects for co-transformants, while growth on full selection medium (FS) selects for interacting proteins. Deletion of the catalytic subunit of phot1 kinase (shaded black, amino acids 663–961) results in a loss of interaction. phot1 kinase also interacts with the internal NBD of B19 (B19, amino acids 480–660). No interaction was detected between phot1 kinase and the C-terminus of B1 (B1C, amino acids 1049–1286). Vectors encoding murine p53 and the SV40 large T-antigen were included as a positive control. (D) In vitro binding of B19C to phot1 kinase (P1K). Equivalent amounts of GST-P1K and GST (indicated in the upper panel by Ponceau S staining) were incubated with c-Myc-tagged B19C synthesized by in vitro transcription/translation (Input). Binding of B19C was visualized by immunoblot analysis with anti-c-Myc antibodies. Sizes of molecular weight markers are indicated on the left (kilodaltons).

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B19 is a stable plasma membrane protein that does not exhibit the dynamic relocalization characteristics of PIN transporters [12]. We therefore rationalized that B19 phosphorylation by phot1 could impact its transporter activity as opposed to its subcellular trafficking. To address this, we examined Arabidopsis ABCB transporter function in HeLa cells, as reported previously [13]. Expression of B19 and B1 resulted in auxin efflux in this system, whereas expression of phot1 was without effect (Figure 5C). In each case, light treatment of HeLa cells resulted in a modest increase in auxin efflux activity. However, co-expression with phot1 reduced B19 transporter activity, particularly following irradiation. Introduction of a single point mutation (D806N) shown previously to inhibit phot1 kinase activity [27] abrogated this effect, indicating that phot1 activation leads to a loss of B19 transporter function. Consistent with our yeast two-hybrid analysis (Figure 4C), phot1 had no impact on B1-mediated auxin transport. While B19C interacts with phot1 kinase (Figure 4C and 4D), potential phosphorylation sites may include those residing within the first NBD of B19 [28], since this region was also found to interact with phot1 kinase (Figure 4C). One mechanism by which phot1 phosphorylation could impact B19 activity is by altering its interaction with key regulatory proteins. The immunophilin-type

Figure 5. phot1 phosphorylation and regulation of B19 auxin transporter activity. (A) In vitro phosphorylation of B19 in the presence of phot1 (upper panel). Functional B19-HA was expressed in S. pombe, whereas active phot1 was expressed in Spodoptera frugiperda (Sf9) insect cells. Membranes containing B19-HA were mixed with phot1 enriched by nickel affinity chromatography. Samples were given a mock irradiation (D) or irradiated with 10,000 μmol m⁻² s⁻¹ white light (L) after the addition of radiolabeled ATP. The asterisk indicates phot1 autophosphorylation. B19-HA is phosphorylated in the presence of phot1 (indicated by the black arrow). S. pombe membranes expressing no B19-HA (vector only) were included as a control. B19-HA and phot1 protein levels were monitored using anti-HA and anti-phot1 antibodies, respectively (lower panels). Sizes of molecular weight markers are indicated on the right (kilodaltons). (B) Phosphorylation of B19-HA by phot1 in (A) was confirmed by immunoprecipitation (IP) of B19-HA using anti-HA antibodies (right lane). All samples were treated with light. No B19 phosphosignal was detected in the absence of B19 (center lane) or when Sf9 extracts lacking phot1 were used as a control (left lane). (C) phot1 regulation of B19-mediated auxin efflux in HeLa cells. Cells were prepared under a green safe light then incubated in darkness (D) or exposed to white light (L, 4 μmol m⁻² s⁻¹) for the duration of the assay. Light-treated samples alone are shown for both B19 and B1 co-expression with kinase inactive phot1 (D806N). Data are expressed as net ³H-IAA efflux (calculated from net ³H-IAA retention in cells expressing indicated proteins versus that of empty vector controls). Results represent mean ± SD, n = 3. Three cell wells were used for each assay. The asterisk represents a significant difference between dark and light treatments (p < 0.05, t test). (D) To determine the effect of blue light on B19-TWD1 interactions in dark-acclimated seedlings, 3-d-old light-grown (L) Arabidopsis seedlings overexpressing TWD1 and functional B19-HA [12,13] were subjected to dark acclimation (DA) prior to irradiation with directional blue light (BL, 2 μmol m⁻² s⁻¹) for 2 h. B19-HA was immunoprecipitated with anti-HA antibody and subjected to immunoblot analysis with anti-B19 and anti-TWD1 antibodies. No attenuation of B19-TWD1 interactions was observed when B19 was immunoprecipitated from the phot1-5 mutant. The asterisk (*) indicates the position of B19. (E) Interaction analysis as in (D) but with 3-d-old dark-grown (D) seedlings. doi:10.1371/journal.pbio.1001076.g005
protein FKBP42/TWISTED DWARF 1 (TWD1) interacts directly with B19 [26] and is a positive regulator of B19-mediated auxin transport [21]. Co-immunoprecipitation of TWD1 with B19-HA was readily detected in dark-acclimated seedlings overexpressing TWD1 (Figure 5D). However, TWD1-B19 interactions were attenuated following 2 h of blue-light treatment. A similar effect of blue light was observed on TWD1-B19 interactions in etiolated seedlings (Figure 5E). In each case, a decrease in interaction between TWD1 and B19 following blue-light exposure was not observed in the phot1 mutant, suggesting that phot1 could act to repress B19 activity in Arabidopsis by disrupting its interaction with TWD1.

Lateral Auxin Fluxes Are Initiated above the Site of Phototropic Growth

Given the above results we conclude that localized pooling of auxin at and above the hypocotyl apex (Figure 2A), which coincides with reduced vertical growth at the onset of phototropism (Figure 1D), most likely results from a transient inactivation of B19 initiated by phot1 kinase activity. Consequently, auxin retention in the region immediately above the hypocotyl apex and not the elongation zone would serve to prime subsequent lateral fluxes required for phototropism [7,23]. In monocots, lateral auxin redistribution is proposed to occur at the coleoptile tip, the predominant site of asymmetric phot1 activation [29]. Our results therefore suggested that a similar mechanism of phototropic detection exists in Arabidopsis. To address this, we monitored cellular auxin accumulation using the synthetic transcriptional reporter DR5rev::GFP. Activity of the auxin-responsive promoter DR5 has been used to visualize the spatial pattern of auxin responses, and hence, indirectly, the distribution of auxin [30].

In wild type, DR5rev::GFP was readily detected in the epidermis and central vasculature of the cotyledonary node and upper hypocotyl (Figure 6A). This auxin accumulation pattern was attenuated in pin3–4 and b19–1 mutants (Figure 6B), where DR5rev::GFP signals were reduced or absent in the central vasculature and upper hypocotyl. Following blue-light exposure, DR5rev::GFP signals were asymmetrically localized in the epidermal cells on the shaded side of the hypocotyl (Figure 6C). This suggests that blue light is inhibiting the auxin response in the hypocotyl, reducing the accumulation of DR5rev::GFP in the epidermis and central vasculature. These results are consistent with the model that blue light inhibits auxin transport in the hypocotyl, preventing the lateral auxin fluxes required for phototropism.
hypo- 

cotyl (Figure 6A). Below in the elongation zone, DR5rev:GFP was associated predominantly with the central vasculature. Phototropic stimulation produced an asymmetric signal in the epidermis encompassing the cotyledonal node and upper hypocotyl after 3 h (Figure 6B) that was apparently phot- dependent (Figure S5A).

Heat map representation and quantification of the DR5rev:GFP fluorescence imaging confirmed that there was an increase in signal on the shaded versus the irradiated side of the hypocotyl apex and cotyledonal node (Figure 6C). Consistent with phot1 inhibition of B19-mediated auxin loading into, and retention within, provascular/vascular tissues, DR5rev:GFP signals decreased simultaneously in the vasculature of the elongation zone of wild-type 

cytocotyls (Figure 6B and 6C). However, no DR5rev:GFP asymmetry was detected in this epidermal region despite the onset of curvature. Thus, the primary site of asymmetric auxin accumulation following light perception by phot1 occurs at and above the hypocotyl apex, not in the region of differential growth.

Lateral Auxin Fluxes Occur in b19 and pin3 Mutants

Lateral DR5rev:GFP signals were still apparent in b19 and pin3 seedlings following phototropic stimulation (Figure 6B and 6C). Asymmetric auxin gradients were transient and no longer detectable following a 12-h period of irradiation, except in b19 mutants, where a weak asymmetric signal persisted in the epidermis of the shaded side of the hypocotyl (Figure S5B). More rapid diversion of auxin out of vascular tissues in and above the hypocotyl apex of b19 mutants (Figure 2A) may account for this sustained lateral signal.

In comparison to wild type, DR5rev:GFP signals were reduced in the central vascular tissue of b19 hypocotyls (Figure 6A), consistent with both IAA, oxIAA, and oxIAA-Glc determinations (Figure S) and the cellular localization of B19:PIN1-GFP (Figure S6). Dynamic pooling and catabolism of IAA within the central vasculature of b19 seedlings may contribute to a cellular auxin status that is too low for DR5rev:GFP to report. However, we cannot exclude the possibility that sustained IAA overaccumulation in b19 mutants could have an adverse effect on DR5rev:GFP reporter activity. In contrast to b19 mutants, a strong DR5rev:GFP signal was observed in the central vasculature of pin3 mutants (Figure 6A). Moreover, the DR5rev:GFP fluorescence observed to be more diffuse throughout the bundle sheath, consistent with the role of PIN3 in redirecting auxin back into the central cylinder [31]. Our radiotracer studies also suggested that PIN3, like PIN1, participates in auxin transport out of the hypocotyl apex and the region immediately above (Figure 2A). In addition, functional PIN3:PIN3-GFP fluorescence levels became reduced in and below the region of elongation in dark-acclimated seedlings following phototropic stimulation for 6 h (Figure 7A). By comparison, directional blue light had no effect on the level of PIN7:PIN7-GFP fluorescence even after 12 h of blue-light exposure (Figure 7B). The reduced abundance of PIN3 in the lower hypocotyl following phototropic stimulation, combined with the higher position of curvature associated with pin3 (Figure 1B) and pin3 b19 mutants (Table 1), led us to propose that PIN3 is involved in channeling auxin to the region of differential growth once lateral redistribution has been established in and immediately above the hypocotyl apex.

Discussion

Arabidopsis Sheds New Light on the Cholodny-Went Theory

In recent years, Arabidopsis has been used extensively as a genetic model to dissect the light-sensing and signaling mechanisms associated with phototropism [1]. However, research to date has focused on etiolated seedlings that simultaneously undergo photomorphogenesis in addition to phototropism when irradiated with directional blue light. In the present study, we show that it is possible to probe auxin fluxes associated with phototropism after such de- etiolation events are fully established. Dark-acclimated light-grown seedlings, like etiolated seedlings [20], are phototropically responsive (Figure 1A and 1B), but provide a more amenable system to study auxin transport processes in response to blue-light irradiation (Figure 2A and 2B). The open cotyledons of light-grown seedlings influence the curvature formation of photostimulated hypocotyls owing to their friction against the vertical agar surface, which must be uniformly maintained by experimental conditions. Nevertheless, the contribution of the cotyledons to the curvature response itself is likely to be minimal, as both phototropism and the establishment of lateral auxin fluxes can be detected in seedlings where the cotyledons have been removed prior to photostimulation (Figure 8A).

Figure 7. PIN3- and PIN7-GFP localization in dark-acclimated seedlings. (A) Functional PIN3:PIN3-GFP fluorescence in hypocotyls of seedlings exposed to blue light (BL, 1 μmol m⁻² s⁻¹) for 6 h or continued darkness. A blue-light-treated seedling is shown on the right and white arrows indicate a decrease in PIN3:PIN3-GFP signal. Scale bar = 200 μm. (B) Functional PIN7:PIN7-GFP fluorescence in hypocotyls of seedlings exposed to blue light (1 μmol m⁻² s⁻¹) for 12 h or continued darkness. Scale bar = 20 μm. Heat map representations of GFP signals are shown on the right. In each case data are representative of n>20 seedlings. doi:10.1371/journal.pbio.1001076.g007
With regard to phototropism, we draw two main conclusions from this work. First, the lateral auxin translocation system in dark-acclimated Arabidopsis seedlings complies with the Cholodny-Went theory of tropic curvature proposed for monocots in the late 1920s [32]. This hypothesis predicts that asymmetric light is perceived at the coleoptile tip and causes auxin to move from the irradiated to the shaded side. Auxin then moves down the coleoptile, where the higher concentration on the shaded side promotes differential growth towards light as a result of increased cell elongation. Although subject to many criticisms since it was first introduced, the Cholodny-Went theory has received considerable support from monocot studies [1,3]. However, the apex has generally not been regarded as the site of phototropic perception in dicots [33]. Rather, studies of the auxin transport processes associated with Arabidopsis phototropism have focused on possible redistribution of auxin out of the central vasculature at the hypocotyl elongation zone [6].

Here we demonstrate that the Cholodny-Went theory holds true for dicot phototropism, where the primary site of lateral auxin translocation resides at and above the hypocotyl apex (Figure 6). Quantification of DR5::GFP signals within the upper hypocotyl in wild type and pin2 mutants suggests that the increase in auxin in the epidermis of the shaded side of the hypocotyl coincides with a decrease in signal from the central vasculature (Figure 6C). This differs from the situation in monocots, where a net movement from the irradiated to the shaded side of the coleoptile is thought to establish the auxin asymmetry [23]. However, it is worth noting that this conclusion is drawn from radiotracer studies that monitor the effect of light on polar IAA transport within two separate halves of the coleoptile [1,3]. A spatial assessment of cellular auxin levels will be necessary to determine whether lateral auxin translocation in monocots is established differently from what is observed in dark-acclimated Arabidopsis.

### A Model for Dicot Phototropism

The results presented here also provide new mechanistic information about the development of hypocotyl phototropism in Arabidopsis (proposed schematically in Figure 8B). In particular, our studies uncover a direct connection between the primary photoreceptor for phototropism, phot1, and the auxin transporter B19. More specifically, B19 functions as a target for phot1 action in the shoot apical tissues to initially halt vertical growth (Figure 1D) and concentrate auxin within this region (Figure 2A and 2B). These findings are consistent with earlier reports showing that an inhibition of polar auxin transport [34], as well as light-mediated growth inhibition [22,23,35], precedes the onset of phototropic curvature.

B19 is directly regulated by phot1 kinase activity, identifying it as a likely substrate for this class of photoreceptor kinase (Figures 4 and 5A–5C). Regulation of B19 function in planta may require more complex phosphorylation events involving other kinases. Reconstitution of active B19 and phot1 in artificial membranes would aid analyses of their interactions, although such efforts have not been successful to date. Our data also suggest that in vivo interactions between phot1 and B19 are likely to be transient following irradiation (Figure 4A). This finding is not unexpected given that phot1 is rapidly internalized from the plasma membrane following blue-light excitation [25] and receptor autophosphorylation [36]. A transient interaction would also ensure that B19 transporter activity is not continually inhibited by phot1 action. One possible consequence of phot1 phosphorylation could be to disrupt an interaction between B19 and its positive regulator TWD1 (Figure 5D and 5E). If so, the FKBP38 mammalian ortholog of TWD1 [21] would also serve as a target of phot1 action on B19 transporter activity in HeLa cells (Figure 5C). Studies are now underway to test this potential mode of regulation (Figure S7).

### Table 1. Phototropic bending in auxin transport mutants of Arabidopsis.

| Genotype       | Mean Bending Angle (8 h) ± SD | Student’s t Test Compared to Col-0 | Notes                                         |
|----------------|-------------------------------|-----------------------------------|-----------------------------------------------|
| Col-0          | 26±2.4                        | ND                                |                                               |
| pin1–1         | 20±6.8                        | ND                                |                                               |
| pin1–5         | 23±3.6                        | ND                                |                                               |
| pin2 (agr1–1, ein1–1) | 27±2.3                     | ND                                |                                               |
| pin3–4         | 33±1.6                        | >p = 0.014                        | Bends at a higher position in the hypocotyl |
| pin4–1         | 25±2.8                        | ND                                |                                               |
| pin7 (NS48791) | 18±3.0                        | <p = 0.023                        |                                               |
| pin1–5 pin3–4  | 28±4.7                        | ND                                |                                               |
| pin3–4 b19–1   | 34±2.9                        | >p = 0.021                        | Bends at a higher position in the hypocotyl |
| b1–1           | 28±3.5                        | ND                                |                                               |
| b4–1           | 27±3.6                        | ND                                |                                               |
| b19–1          | 39±2.2                        | >p = 0.023                        |                                               |
| b1–1 b4–1 b19–1| 40±3.8                        | >p = 0.006                        |                                               |
| pin1–1 b1–1 b19–1 | 34±6.6                     | ND                                |                                               |
| aux1 lax2 lax3 | 21±3.3                        | p = 0.101                         |                                               |

Homologous dark-acclimated seedlings were analyzed for phototropic bending induced by 8 h directional blue light (1 μmol m⁻² s⁻¹). Results represent the mean ± SD, n = 10 seedlings in three independent experiments except for pin1–1 and pin1–1 b1–1 b19–1, where eight and five seedlings were used, respectively, in each experiment. For assays of pin1–1 and pin1–1 b1–1 b19–1, progeny of a single plant heterozygous for pin1–1 were used in the bending assays. Reduced-bending pin1–1 seedlings and increased-bending pin1–1 b1–1 b19–1 seedlings were selected and grown to maturity then examined for pin-formed inflorescences to determine if they were homoygous for pin1. Significant differences compared to wild type (Col-0) are indicated. ND, not determined.

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Following its transient pooling in apical perivascular tissues, auxin is subsequently translocated from the central vasculature to the epidermis at the shaded side, where PIN3 participates in its transport to the elongation zone below. Retention of auxin in the elongation zone is facilitated by a reduction in PIN3 abundance in and below this region of the hypocotyl after phototropic stimulation (Figure 7A). Without PIN3, auxin is unable to move downward to this region. Consequently, localized apical pooling results in faster initial rates of curvature (Figure 1C) at a higher region of the hypocotyl (Figure 1B). This is not apparent with etiolated pin3 seedlings, where curvature is reduced (Figure S1A), an outcome possibly arising from PIN3’s impact on apical hook opening [9] as reported for PIN1 [13].

Lateral Auxin Translocation Processes

The second conclusion to be drawn from this work is that the lateral translocation of auxin required for phototropism does not depend on B19 or PIN3 (Figures 1C and 6). Indeed, this process cannot be ascribed to any of the well-characterized auxin transporters, as mutants lacking these proteins are phototropic in our dark-acclimated system despite showing some altered rates in bending (Table 1). In light of these results, efforts to contrive a response mechanism involving these components are not convincing. Coordinated action between multiple transporters is therefore likely to drive the lateral redirection of auxin that is necessary to establish phototropic growth. PIN7 could be a factor in the lateral redistribution process, in conjunction with other transporters, as...
dark-acclimated pin7 seedlings are phototropic, but exhibit lower curvatures relative to wild type (Table 1).

The mechanism by which lateral auxin gradients are established is still unknown. As demonstrated here, phosphorylation of the transporters involved by phototropic receptors offers one possible mode of regulation. phot1 could control the subcellular trafficking of regulatory components of the lateral relocation process, as recent studies have identified ADP-ribosylation factors as direct interaction targets [37]. Lateral translocation may also be directed by differential proton extrusion, which represents a primary driving force for both auxin uptake and efflux [4]. Further studies of NON-PHOTOTROPIC HYPOCOTYL 3 (NPH3) will undoubtedly be important to improve our understanding of the processes involved, as mutants lacking this protein are phototropic [38] and fail to exhibit lateral auxin transport [23]. NPH3 is a plant-specific plasma-membrane-associated protein that directly interacts with phot1 [38]. Photocactivation of phot1 leads to dephosphorylation of NPH3, a signaling process that has been linked to the onset of phototropic curvature [39], but how NPH3 functions to orchestrate auxin transport regulation remains poorly understood. Members of the PHYTOCHROME KINASE SUBSTRATE (PKS) protein family are known to play a role in the establishment of phototropic curvatures [40], in addition to other phot-mediated responses [41]. We anticipate that adoption of this revised system to study phototropism will accelerate our understanding of the role of these, and as yet undiscovered, components of phototropic signaling.

**Materials and Methods**

**Plant Material and Growth**

The Arabidopsis thaliana ecotype Col-0 was used throughout this study. Details of each line can be found in Table S3. Unless otherwise stated, seedlings were grown on 0.8% agar plates containing ¼ Murashige Skoog basal salts (pH 5.5) supplemented with 0.5% sucrose for 3 d under white light (100 μmol m⁻² s⁻¹). During growth, plates were tilted 2° back from the vertical. Plates were transferred to darkness in light-tight gas exchange boxes equipped with directional 450-nm blue-light LED arrays (Roithner Lasertechnik) calibrated to 1 μmol m⁻² s⁻¹ and an infrared light source (Electro Optical Components) and tilted 1° forward for 24 h to dark acclimate. Subsequent growth in light or darkness was measured using a CCD video camera (Sony) equipped with a narrow band pass infrared filter (Electro Optical Components). Images were quantified using Image J software (US National Institutes of Health). Phototropism in etiolated seedlings was performed as described [42].

**Auxin Transport and Quantification**

Auxin transport and quantification was performed as described [13,26]. 3H-IAA at 10 μM (30 Gi mmol⁻¹), American Radiolabeled Chemicals) was dissolved in ethanol. 3H-IAA was adsorbed to the surface of 125-μm to 212-μm polystyrene beads (Sigma) and placed at the hypocotyl apex by touching the tips of the true leaves with a micromanipulator and liquid stream from a pipettor. All experiments were prepared under green or red safe lights. Seedlings were incubated in darkness or directional blue light (1 μmol m⁻² s⁻¹) for the times indicated. Sections were collected under white light, with cuts through the true leaves/cotyledonary petioles to remove sites of contact with polystyrene beads and hypocotyl immediately below the cotyledonary node. A second section, corresponding to the mid hypocotyl and elongation zone, was collected by making a second cut in the hypocotyl 2 mm below. Free IAA determination was performed as described [43] on sections made from the mid hypocotyl and apical tissues including the upper hypocotyl, petioles, and cotyledons. oxIAA and oxIAA-Glc levels were assayed by liquid chromatography–mass spectrometry using a Waters Micromass Q-TOF micro and Agilent HPLC-MSD/TOF system calibrated to a standard curve generated with a defined mixture of IAA, oxIAA, and oxIAA-Glc using ²⁶C-IAA (Cambridge Isotopes) as an internal standard.

**Auxin Efflux Assays in HeLa Cells**

Assays were performed as described [13,44] in darkness or under a green safe light unless otherwise stated. Cells were transfected with pTM1 vectors encoding B19 and PHOT1 in a ratio of 4:1, respectively. Co-expression of PHOT1 and B19 was verified by real time quantitative PCR and immunolocalization as described [13].

**Immunoprecipitation from Arabidopsis**

Membrane proteins were extracted and GFP immunoprecipitation performed as described [42]. Proteins were identified by liquid chromatography-tandem mass spectrometry using the Fingerprint Proteomics Facility (University of Dundee). B19 immunoprecipitation was performed as described [12,13], and proteins identified by using liquid chromatography-matrix-assisted laser desorption/ionization mass spectrometry analysis (Bindley Bioscience Center, Purdue University).

**Yeast Two-Hybrid Analysis**

Interaction assays were carried out as described [37]. Interactions between regions of B19 cDNA and PHOT1 cDNA were performed using pHAT17 prey and pGBK17 bait vectors, respectively (Clontech). The C-terminal region of B1 used for yeast two-hybrid analysis was as described [26].

**In Vitro Co-Immunoprecipitation**

Expression of GST-fusion proteins and GST was performed as described [37]. The TNT System (Promega) was used to generate c-Myc-tagged protein from 2-μg circular plasmid DNA (pGBK17). The MagneGST Protein Purification System (Promega) was used for in vitro GST pull-down assays in accordance to the manufacturer's instructions. Proteins were eluted in 1× SDS-PAGE loading buffer and subjected to SDS-PAGE and immunoblotting with anti-c-Myc antibodies (Roche).

**In Vitro Kinase Assays**

B19-HA was expressed in S. pombe [45]. Sf9 insect cells were used to express phot1 harboring a 6xHis epitope [27]. Sf9 extracts were passed through a HisSpinTrap column (GE Healthcare) in accordance with the vendor’s protocol. Eluted phot1 was desalted using a microcon 100 centrifugal filter (Millipore), and phosphorylation assays performed [27] in 20-μl reactions with 3 μg of phot1 extract and 10 μg of S. pombe membranes containing B19-HA. Immunoprecipitation of B19-HA following in vitro phosphorylation assays was performed as described [46] using anti-HA antibodies (Santa Cruz Biotechnology) and Immunobead Reagent (Bio-Rad). Kinase assays with plant membranes were performed as reported [42].

**Confocal Microscopy**

Images were collected using a Zeiss LSM 710 confocal laser-scanning microscope with a 40× water immersion objective (1.4 numerical aperture, C-Apochromatic) and argon laser. Excitation was fixed at 488 nm using the primary dichroic mirror, and the Meta detector was used to capture GFP (492–533 nm) emission.
Supporting Information

Figure S1 Phototropic responsiveness of pin3 mutants. (A) Phototropic response of 3-d-old etiolated wild-type (WT) and pin3-4 seedlings. Directional blue light (1 μmol m⁻² s⁻¹) was supplied for 12 h. Results represent the mean ± standard error, n = 10 seedlings. (B) Phototropism of dark-acclimated wild-type and pin3-4 seedlings. Seedlings were subjected to directional blue light (1 μmol m⁻² s⁻¹) for the times indicated. Results represent the mean ± standard error, n = 10 seedlings.

Figure S2 3H-IAA accumulation in dark-acclimated seedlings overexpressing B19 (b19oe). Seedlings were exposed to directional blue light (BL) (1 μmol m⁻² s⁻¹) or continued darkness (D). Upper hypocotyls including the cotyledonary node (UH) and mid hypocotyls including the elongation zone (MH) were excised after 2.5 h. Results represent the mean ± SD, n = 10 seedlings in three independent experiments.

Figure S3 Bright-field BiFC fluorescence images of phot1-YN and B19-YC co-expressed in tobacco epidermal cells. Reconstitution of YFP fluorescence between phot1-YN and B19-YC was visible at the plasma membrane (left). Only background YFP signals were detected for phot1-YN and YC (center) or YN and B19-YC (right). Scale bar = 20 μm and applies to all images.

Figure S4 Expression of phot1 kinase and the C-terminal NBD of B1 in yeast. Immunoblot analysis of yeast co-expressing phot1 kinase (P1K) and the C-terminal NBD of B1 (B1C). Protein extracts (10 μg) co-expressing phot1 kinase and either B1C (1) or empty vector controls (2) were probed with anti-GAL4 DNA-binding domain and activation domain antibodies to discriminate phot1 kinase and C-terminal B1 proteins, respectively.

Figure S5 Auxin accumulation in the upper hypocotyl/cotyledonary node of dark-acclimated hypocotyls. (A) DR5rev::GFP signals in phot1−5 phot2−1 seedlings exposed to directional blue light (BL, 1 μmol m⁻² s⁻¹) for 3 h. Signal intensity of DR5rev::GFP was noticeably lower in comparison to the other lines examined, giving rise to higher background plastid autofluorescence (represented by spots). (B) DR5rev::GFP signals in seedlings exposed to directional blue light for 12 h. Note the lack of signal in the vascular bundle in b19−1 seedlings and the reduced signal in the vascular bundle below the upper hypocotyl in pin3−4 seedlings. Data are representative of n > 5. In each case, scale bar = 200 μm.

Figure S6 B19-GFP localization in dark-acclimated seedlings. Functional B19:B19-GFP fluorescence is restricted to the central vasculature and epidermis (white arrows). Central spots represent plastid autofluorescence. Data are representative of n = 20. Scale bar = 100 μm.

Table S1 Liquid chromatography–tandem mass spectrometry analysis of phot1-GFP immunoprecipitates. Proteins identified with a Mascot score > 100 are shown. "PHOT1:phot1-GFP was immunoprecipitated from 3-d-old etiolated Arabidopsis seedlings kept in darkness (D) or exposed to a blue light (BL, 20 μmol m⁻² s⁻¹) for 5 min. "TAIR AGI numbers given where mass spectrometry data match a single accession; "various" denotes multiple isoforms were identified.

Table S2 Liquid chromatography–matrix-assisted laser desorption/ionization mass spectrometry analysis of B19 immunoprecipitates. Proteins identified with a Mascot score > 150 are shown. TAIR AGI numbers are given. B19:B19-HA was immunoprecipitated from 5-d-old Arabidopsis seedlings. Previously identified in B19 fractions [12].

Table S3 Lines used in this study.

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Author Contributions

The author(s) have made the following declarations about their contributions: Conceived and designed the experiments: JMC WAP ASM. Performed the experiments: JMC HY GR SS CET JL BT ME EK ORL JA ASM. Analyzed the data: JMC WAP ASM. Wrote the paper: ASM. Performed the experiments: JMC HY GR SS CET JL BT ME EK ORL JA ASM.

References

1. Holland JJ, Roberts D, Liscum E (2009) Understanding phototropism: from Darwin to today. J Exp Bot 60: 1969–1978.

2. Darwin C (1880) The power of movement in plants. London: John Murray Publishers.
3. Whippo CW, Haungarter RP (2006) Phototropism: bending towards enlightenment. Plant Cell 18: 1110–1119.

4. Vanneste S, Friml J (2009) Auxin: a trigger for change in plant development. Cell 136: 1003–1016.

5. Christie JM (2007) Phototropin blue-light receptors. Annu Rev Plant Biol 58: 21–45.

6. Friml J, Wimmierska J, Benková E, Mendgen K, Palme K (2002) Lateral relocation of auxin efflux regulator PIN3 mediates tropism in Arabidopsis. Nature 415: 896–899.

7. Esmoo CA, Timley AG, Ljung K, Sandberg G, Hearne LB, et al. (2006) A gradient of auxin and auxin-dependent transcription precedes tropic growth responses. Proc Natl Acad Sci U S A 103: 236–241.

8. Krček P, Skupa P, Libus J, Naramoto S, Tejos R, et al. (2009) The PIN-FORMED (PIN) protein family of auxin transporters. Genomics 10: 249–259.

9. Zádržková P, Petrášek J, Mravec J, Noh B, Murphy A, Spalding EP, et al. (2006) Role of PIN-mediated auxin efflux in apical hook development of Arabidopsis thaliana. Development 133: 607–617.

10. Nagashima A, Uchura Y, Sakai T (2008) The ABC subfamily B auxin transporter AAtABC19 is involved in the inhibitory effects of N-L-naphtylphthalamic acid on the phototropic and gravitropic responses of Arabidopsis hypocotyl. Plant Cell Physiol 49: 1250–1255.

11. Mravec J, Kubés M, Buhlach A, Gaykova V, Petrášek J, et al. (2008) Interaction of PIN and PGP transport mechanisms in auxin distribution-dependent development. Development 135: 3345–3354.

12. Ti tatupatana N, Bacleske JR, Bandyopadhyay A, Yang H, Mravec J, et al. (2009) ABCB19/PGP19 stabilizes PIN1 in membrane microdomains in Arabidopsis. Plant J 57: 27–44.

13. Blakeslee JJ, Bandyopadhyay A, Lee OR, Mravec J, Titapiwatanakun B, et al. (2007) Interactions among PIN-FORMED and P-glycoprotein auxin transporters in Arabidopsis. Plant Cell 19: 131–147.

14. Wu G, Cameron JN, Ljung K, Spalding EP (2010) A role for ABCB19-mediated polar auxin transport in seedling photomorphogenesis mediated by cryptochrome 1 and phytochrome B. Plant Cell 49: 1250–1255.

15. Noh B, Bandyopadhyay A, Peer WA, Spalding EP, Murphy AS (2003) Enhanced gravitropism and phototropism in plant nph1 mutants mislocalizing the auxin efflux protein PIN1. Nature 425: 999–1002.

16. Nagashima A, Suzuki G, Uchura Y, Saji K, Furukawa T, et al. (2008) Immunophilin-like TWISTED DWARF1 modulates auxin efflux activities of Immunophilin-like TWISTED DWARF1 and chloroplast relocation. Proc Natl Acad Sci U S A 105: 103–108.

17. Sakamoto K, Briggs WR (2002) Cellular and subcellular localization of phototropin 1. Plant Cell 14: 1723–1735.

18. Sealfin D, Komiya K, Bouchard R, Billson K, Berger J, et al. (2003) TWISTED DWARF1, a unique plasma membrane-anchored immunophilin-like protein, interacts with Phototropin 1 and functions as a molecular chaperone. J Biol Chem 278: 11645–11651.

19. Kowalyczk M, Bhalerao RP, Sandberg G (1998) Metabolism of indole-3-acetic acid in Arabidopsis. Plant Physiol 118: 205–206.

20. Sullivan S, Thomson CE, Lamont DJ, Jones MA, Christie JM (2008) Interaction specificity of PIN1 and PGP transport mechanisms in auxin distribution-dependent development. Development 135: 3345–3354.

21. Bouchard R, Bailly A, Blakeslee JJ, Oehring SC, Vincenzetti V, et al. (2006) Enhanced phototropic and gravitropic responses. Proc Natl Acad Sci U S A 103: 236–241.

22. Sakamoto K, Briggs WR (2002) Phototropin LOV domains exhibit distinct roles in regulating photoreceptor function. Plant J 32: 87–98.

23. New Light on the Mechanism of Dicot Phototropism

24. Sullivan S, Thomson CE, Lamont DJ, Jones MA, Christie JM (2008) Interaction specificity of PIN1 and PGP transport mechanisms in auxin distribution-dependent development. Development 135: 3345–3354.