Control of vein-forming, striped gene expression by auxin signaling

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Abstract

Background: Activation of gene expression in striped domains is a key building block of biological patterning, from the recursive formation of veins in plant leaves to that of ribs and vertebrae in our bodies. In animals, gene expression is activated in striped domains by the differential affinity of broadly expressed transcription factors for their target genes and the combinatorial interaction between such target genes. In plants, how gene expression is activated in striped domains is instead unknown. We address this question for the broadly expressed MONOPTEROS (MP) transcription factor and its target gene ARABIDOPSIS THALIANA HOMEBOX8 (ATHB8).

Results: We find that ATHB8 promotes vein formation and that such vein-forming function depends on both levels of ATHB8 expression and width of ATHB8 expression domains. We further find that ATHB8 expression is activated in striped domains by a combination of (1) activation of ATHB8 expression through binding of peak levels of MP to a low-affinity MP-binding site in the ATHB8 promoter and (2) repression of ATHB8 expression by MP target genes of the AUXIN/INDOLE-3-ACETIC-ACID-INDUCIBLE family.

Conclusions: Our findings suggest that a common regulatory logic controls activation of gene expression in striped domains in both plants and animals despite the independent evolution of their multicellularity.

Keywords: Stripe formation, Gene regulatory network, Arabidopsis thaliana, Auxin, Leaf vascular patterning, MONOPTEROS, ARABIDOPSIS THALIANA HOMEBOX8, Incoherent feedforward loop, Vein network formation

Background

Narrow stripes of gene expression are fundamental units of biological patterning (e.g., [1–3]). Therefore, how multicellular organisms activate gene expression in narrow stripes is a central question in biology. In animals, where this question has been investigated extensively, broadly expressed transcription factors activate expression of their target genes in narrow stripes by (1) differential affinity of such transcription factors for their binding sites in target genes and (2) combinatorial interactions between transcription-factor-encoding target genes [4–7]. For example, the transcription factor Dorsal forms a ventral-to-dorsal gradient in Drosophila embryos (reviewed in [8]). Expression of Dorsal target genes with high-affinity Dorsal-binding sites is activated already at low levels of Dorsal, whereas expression of Dorsal target genes with low-affinity Dorsal-binding sites is activated only at high levels of Dorsal. However, this mechanism alone is insufficient to account for the expression of Dorsal target genes in stripes: interaction between Dorsal target genes themselves is also required: Dorsal activates expression of snail, which encodes a transcription factor that represses the expression of the Dorsal target gene ventral nervous system defective. Thus, expression of some Dorsal target genes such as ventral nervous system defective is repressed at high levels of Dorsal, at which snail is expressed, but activated at lower levels of Dorsal, at which snail is not expressed.

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In plants too, broadly expressed transcription factors activate expression of their target genes in narrow stripes (e.g., [9]); however, how these broadly expressed transcription factors do so is unclear. Here we addressed this question for the MONOPTEROS (MP) – ARABIDOPSIS THALIANA HOMEBOX8 (ATHB8) pair of Arabidopsis genes [10, 11]. ATHB8 expression is activated in single files of isodiametric ground cells of the leaf [12, 13]. ATHB8-expressing ground cells will elongate into procambial cells — the precursors to all vascular cells — and are therefore referred to as preprocambial cells [12–15]. Activation of ATHB8 expression in narrow preprocambial stripes depends on binding of the broadly expressed MP transcription factor to a low-affinity MP-binding site in the ATHB8 promoter [16]. However, the biological relevance of activation of ATHB8 expression by MP is unclear: whereas MP promotes vein formation [17], ATHB8 seems to have only transient and conditional functions in vein network formation [16, 18].

Here we show that ATHB8 promotes vein formation and that both levels of ATHB8 expression and width of ATHB8 expression domains are relevant to vein formation. Finally, we show that ATHB8 expression is restricted to narrow preprocambial stripes by a combination of (1) activation ofATHB8 expression through binding of peak levels of MP to a low-affinity MP-binding site in the ATHB8 promoter and (2) repression ofATHB8 expression by MP target genes of the Auxin/Indole-3-acetic-acid-inducible family.

**Results**

**Response of vein network formation to changes in ATHB8 expression and activity**

To understand how in plants broadly expressed transcription factors activate expression of their target genes in narrow stripes, we chose the MP – ATHB8 pair of Arabidopsis genes. During leaf development, the broadly expressed MP transcription factor directly activates ATHB8 expression in narrow preprocambial stripes that mark the position where veins will form [16], but the biological relevance of the interaction between the two genes is unclear.

That MP promotes vein formation is known [17], but the function of ATHB8 in this process is unresolved: athb8 mutants seem to have only transient and conditional defects in vein network formation, and the mutants have normal vein patterns [16, 18]. Therefore, we first asked whether ATHB8 had any permanent functions in vein network formation. To address this question, we characterized the vein networks in mature first leaves of the athb8-11 and athb8-27 loss-of-function mutants [19] (Table S1) — and of other genotypes in our study — by means of four descriptors: a cardinality index, a continuity index, and a connectivity index [20], and a cyllicity index.

The cardinality index is a proxy for the number of “veins” (i.e., stretches of vascular elements that contact other stretches of vascular elements at least at one of their two ends) in a network. The continuity index quantifies how close a vein network is to a network with the same pattern but in which at least one end of each “vein fragment” (i.e., a stretch of vascular elements that is free of contact with other stretches of vascular elements) contacts a vein. The cyllicity index is a proxy for the number of meshes in a vein network.

The cardinality index of both athb8-11 and athb8-27 was lower than that of wild type (WT) (Fig. 1A–C,K), suggesting that ATHB8 promotes vein formation. ATHB8 encodes a transcription factor member of the Homeodomain-Leucine Zipper III (HD-ZIP III) family [10]. To further test whether ATHB8 promoted vein formation and to test whether ATHB8 did so redundantly with other HD-ZIP III genes, we expressed microRNA165a (miR165a) — which targets all the HD-ZIP III genes [21] — by the Short-Root (SHR) promoter — which drives expression in the ATHB8 expression domain [22] (Additional File 1: Fig. S1A–D) — in both the WT and athb8-11 backgrounds.

The cardinality index of SHR:miR165a was lower than that of WT, and the cardinality index of SHR:miR165a; athb8-11 was lower than that of SHR:miR165a (Fig. 1D,E,K), supporting that ATHB8 promotes vein formation and suggesting that ATHB8 does so redundantly with other HD-ZIP III genes.

HD-ZIP III proteins bind DNA as homo- or heterodimers [23, 24]. Therefore, to further test whether ATHB8 promoted vein formation and whether ATHB8 did so redundantly with other HD-ZIP III genes, we generated a dominant-negative version of the ATHB8 transcriptional activator [25] by fusing the ATHB8 ORF to the sequence encoding the EAR (ethylene-responsive-element-binding-protein-associated amphiphilic repression) portable repressor domain [26]. In the resulting ATHB8:EAR, we introduced silent mutations that abolish miR165a-mediated downregulation [27]. We expressed the resulting mATHB8:EAR by the SHR promoter in both the WT and athb8-27 backgrounds.

The cardinality index of SHR::mATHB8:EAR was lower than that of WT, and the cardinality index of SHR::mATHB8:EAR;athb8-27 was lower than that of SHR::mATHB8:EAR (Fig. 1F,G,K), supporting that ATHB8 promotes vein formation and that ATHB8 does so redundantly with other HD-ZIP III genes.
Fig. 1 (See legend on next page.)
We next asked whether levels of *ATHB8* expression and width of *ATHB8* expression domains were relevant to vein formation. To address this question, we used *SHR::mATHB8*, which overexpresses *ATHB8* in its expression domain; *MP::ATHB8*, which expresses *ATHB8* in the broader *MP* expression domain (Additional File 1: Fig. S1E); and *MP::mATHB8*, which overexpresses *ATHB8* in the *MP* expression domain (Additional File 1: Fig. S1F).

The cardinality index of *SHR::mATHB8* was lower than that of WT; the cardinality index of *MP::ATHB8* was lower than that of *SHR::mATHB8*; and the cardinality index of *MP::mATHB8* was lower than that of *MP::ATHB8* (Fig. 1H–K). These results suggest that both levels of *ATHB8* expression and width of *ATHB8* expression domains are relevant to vein formation.

The continuity and connectivity indices of the genetic backgrounds with modified *ATHB8* expression or activity either were no different from those of their respective reference backgrounds or changed with no consistent relation to changes in *ATHB8* expression or activity (Fig. 1H–K). Therefore, the differences in cyclinity index of the genetic backgrounds with modified *ATHB8* expression or activity can be attributed to differences in their cardinality index (Fig. 1H–K), from which the cyclinity index is derived (see “Methods”).

In the root, *HD-ZIP III* genes promote differentiation of the xylem vascular tissue [28, 29]. We therefore asked whether changes in *ATHB8* expression or activity led to defects in leaf xylem differentiation.

Veins in *SHR::mir165a* had gaps in xylem differentiation, and those gaps were longer in *SHR::mir165a; athb8-11* (Fig. 1L,O,P). By contrast, the veins of the remaining genetic backgrounds with modified *ATHB8* expression or activity had no defects in xylem differentiation (Fig. 1L–N,Q–U). In conclusion, our results suggest that *ATHB8* promotes vein formation, both nonredundantly and redundantly with other *HD-ZIP III* genes; that levels of *ATHB8* expression and width of *ATHB8* expression domains are relevant to vein formation; and that *ATHB8* promotes xylem differentiation but only redundantly with other *HD-ZIP III* genes. By contrast, *ATHB8* is inconsequential to vein continuity and network connectedness.

### Relation between *ATHB8* expression domains and *MP* expression levels

Width of *ATHB8* expression domains is relevant to vein formation (Fig. 1). Therefore, we asked how *ATHB8* expression is activated in narrow preprocambial stripes by the broadly expressed MP. We hypothesized that *ATHB8* preprocambial expression is activated in narrow stripes by binding of peak levels of the broadly expressed MP to a low-affinity site in the *ATHB8* promoter. This hypothesis predicts that narrow stripes of *ATHB8* preprocambial expression correspond to peak levels of MP expression. To test this prediction, we simultaneously imaged expression of *ATHB8*:nCFP (nuclear CFP expressed by the *ATHB8* promoter) [14] and *MP::YFP* (MP:YFP fusion protein expressed by the *MP* promoter) in first leaves of the strong *mp-B4149* mutant [30], whose defects were rescued by *MP::YFP* expression (Additional File 1: Fig. S2A–C) (Additional File 2: Table S1) [14, 16, 19, 26, 27, 30–43].

*ATHB8* preprocambial expression can be reproducibly observed in midvein, first loops of veins (“first loops”), and second loops of first leaves, respectively 2, 3, and 4 days after germination (DAG) [16, 22, 44]. At these stages, *MP::MP::YFP* was expressed in *ATHB8*:nCFP-expressing cells at higher levels than in cells flanking...
ATHB8:nCFP-expressing cells (Fig. 2; Additional File 1: Fig. S3A,B).

To test whether the differential expression of MP::MP:YFP in ATHB8:nCFP-expressing cells and in cells flanking ATHB8:nCFP-expressing cells were an imaging artifact, we compared expression levels of nCFP driven by a ubiquitously active promoter (RIBO::nCFP) [31] in cells expressing ATHB8:nYFP [14] and in cells flanking ATHB8:nYFP-expressing cells. We focused our analysis on second loops of 4-DAG first leaves, in which ATHB8 preprocambial expression can be reproducibly observed [16, 22, 44].

Because levels of RIBO::nCFP expression in ATHB8::nYFP-expressing cells were no higher than those in cells flanking ATHB8::nYFP-expressing cells (Additional File 1: Fig. S3D,E; Additional File 1: Figure S4), we conclude that the differential expression of MP::MP:YFP in ATHB8::nCFP-expressing cells and in cells flanking ATHB8::nCFP-expressing cells is not an imaging artifact, and therefore that narrow stripes of ATHB8 preprocambial expression correspond to peak levels of MP expression.

Response of ATHB8 expression and vein network formation to changes in MP expression

The hypothesis — that ATHB8 preprocambial expression is restricted to narrow stripes by binding of peak levels of the broadly expressed MP transcription factor

![Fig. 2 ATHB8 and MP Expression Domains and Levels in Leaf Development. First leaves 2, 3, and 4 DAG. Column 1: schematics of leaves — imaged in columns 2–5 — illustrating onset of ATHB8 expression (red) — imaged in column 2 — associated with formation of midvein (2 DAG), first loop (3 DAG), or second loop (4 DAG) [16, 22, 44]. Magenta: epidermis; increasingly darker gray: progressively older ATHB8 expression domains. Columns 2–5: confocal laser scanning microscopy. Column 2: ATHB8:nCFP expression. Column 3: MP::MP:YFP expression; dashed magenta outline: MP::MP:YFP-expressing epidermal nuclei. Column 4: autofluorescence. Column 5: overlays of images in columns 2–4; red: ATHB8:nCFP expression; green: MP::MP:YFP expression; blue: autofluorescence. Column 6: ATHB8::nCFP and MP::MP:YFP expression levels (mean ± SE) in nuclei flanking ATHB8::nCFP-expressing nuclei (positions -2, -1, +1, and +2) relative to ATHB8::nCFP and MP::MP:YFP expression levels in nuclei co-expressing ATHB8::nCFP (position 0) during formation of midvein (top), first loop (middle), or second loop (bottom). Difference between ATHB8::nCFP expression levels in nuclei at position -2, -1, +1, or +2 and ATHB8::nCFP expression levels in nuclei at position 0, and between MP::MP:YFP expression levels in nuclei at position -2, -1, +1, or +2 and MP::MP:YFP expression levels in nuclei at position 0 was significant at P < 0.001 (*** by One-Way ANOVA and Tukey's Pairwise test. ATHB8::nCFP sample sizes: 35 (2 DAG), 33 (3 DAG), or 32 (4 DAG) leaves; position -2: 29 (2 DAG), 43 (3 DAG), or 49 (4 DAG) nuclei; position -1: 57 (2 DAG), 70 (3 DAG), or 66 (4 DAG) nuclei; position 0: 63 (2 DAG), 73 (3 DAG), or 69 (4 DAG) nuclei; position +1: 52 (2 DAG), 46 (3 DAG), or 58 (4 DAG) nuclei; position +2: 23 (2 DAG), 19 (3 DAG), or 37 (4 DAG) nuclei. MP::MP:YFP sample sizes: 35 (2 DAG), 33 (3 DAG), or 32 (4 DAG) leaves; position -2: 30 (2 DAG), 45 (3 DAG), or 50 (4 DAG) nuclei; position -1: 63 (2 DAG), 72 (3 DAG), or 67 (4 DAG) nuclei; position 0: 70 (2 DAG), 75 (3 DAG), or 70 (4 DAG) nuclei; position +1: 58 (2 DAG), 47 (3 DAG), or 59 (4 DAG) nuclei; position +2: 24 (2 DAG), 19 (3 DAG), or 38 (4 DAG) nuclei. Column 7: Local background levels (mean ± SE) in CFP and YFP confocal channels — measured in image regions containing no features of interest as in [14, 100] — relative to ATHB8::nCFP and MP::MP:YFP expression levels in nuclei co-expressing ATHB8::nCFP (position 0) during formation of midvein (top), first loop (middle), or second loop (bottom). Scale bars (shown, for simplicity, only in column 2): 5 μm.
to a low-affinity site in the \( \textit{ATHB8} \) promoter — predicts that loss of MP function will lead to extremely weak, or altogether absent, \( \textit{ATHB8} \) preprocambial expression, otherwise normally visible in second loops of 4-DAG first leaves [16, 22, 44]. To test this prediction, we quantified \( \textit{ATHB8}:\text{nYFP} \) expression levels in second loops of 4-DAG first leaves of the strong \( mp-\text{U55} \) mutant [16, 32].

Consistent with previous observations [16], \( \textit{ATHB8}:\text{nYFP} \) expression levels were greatly reduced in \( mp-\text{U55} \), leading to near-complete loss of \( \textit{ATHB8}:\text{nYFP} \) preprocambial expression (Fig. 3A,B,F). Moreover, consistent with previous observations [16, 17], near-complete loss of \( \textit{ATHB8} \) preprocambial expression in \( mp-\text{U55} \) developing leaves was associated with networks of fewer meshes and fewer, less frequently continuous, and less frequently connected veins in \( mp-\text{U55} \) mature leaves (Fig. 3G,H,K).

The hypothesis further predicts that lower levels of \( MP \) expression will lead to lower levels of \( \textit{ATHB8} \) preprocambial expression. To test this prediction, we quantified \( \textit{ATHB8}:\text{nYFP} \) expression levels in second loops of 4-DAG first leaves of the weak \( mp-11 \) mutant, in which an insertion in the MP promoter [33] leads to ~85% reduction in levels of WT \( MP \) transcript (Additional File 1: Figure S5).

In \( mp-11 \), \( \textit{ATHB8}:\text{nYFP} \) expression levels were lower and expression along the domain was more heterogeneous than in WT, leading to seemingly fragmented domains of weak \( \textit{ATHB8}:\text{nYFP} \) preprocambial expression (Fig. 3A,C,F). Moreover, as in \( mp-\text{U55} \), defects in \( \textit{ATHB8} \) expression in \( mp-11 \) developing leaves were associated with networks of fewer meshes and fewer, less frequently continuous, and less frequently connected veins in \( mp-11 \) mature leaves (Fig. 3G,I,K). However, the vein network and \( \textit{ATHB8} \) expression defects of \( mp-11 \) were weaker than those of \( mp-\text{U55} \) (Fig. 3A–C,G–I,K).

The hypothesis also predicts that higher levels of the broadly expressed MP will lead to higher levels of \( \textit{ATHB8} \) preprocambial expression in both vein and flanking cells, resulting in broader \( \textit{ATHB8} \) expression domains. To test this prediction, we overexpressed \( MP \) by its own promoter (MP::MP) — which led to ~10-fold increase in \( MP \) expression levels (Additional File 1: Figure S5) and which rescued defects of the strong \( mp-B4149 \) mutant (Additional File 1: Fig. S2A,B,D) (Additional File 2: Table S1) — and quantified \( \textit{ATHB8}:\text{nYFP} \) expression levels in second loops of 4-DAG MP::MP first leaves.

In MP::MP, \( \textit{ATHB8}:\text{nYFP} \) expression levels were higher in flanking cells, leading to broad bands of \( \textit{ATHB8}:\text{nYFP} \) expression; however, \( \textit{ATHB8}:\text{nYFP} \) expression levels were lower in vein cells (Fig. 3A,D,F). Nevertheless, broad bands of \( \textit{ATHB8} \) expression in MP::MP developing leaves were associated with abnormal vein networks in MP::MP mature leaves: veins ran close to one another for varying stretches of the narrow leaf laminae, then diverged, and either ran close to other veins or converged back to give rise to elongated meshes (Fig. 3G,J,K).

In conclusion, lower levels of \( MP \) expression lead to fragmented domains of \( \textit{ATHB8} \) preprocambial expression, and loss of \( MP \) function leads to near-complete loss of \( \textit{ATHB8} \) preprocambial expression. These observations are consistent with the hypothesis and suggest that \( MP \) expression levels below a minimum threshold are unable to activate \( \textit{ATHB8} \) preprocambial expression. However, that higher levels of \( MP \) expression fail to lead to higher levels of \( \textit{ATHB8} \) preprocambial expression in vein cells is inconsistent with the hypothesis and suggests that \( MP \) expression levels above a maximum threshold both activate and repress \( \textit{ATHB8} \) preprocambial expression. These observations are unaccounted for by the hypothesis; therefore, the hypothesis must be revised.

**Response of \( \textit{ATHB8} \) expression and vein network formation to changes in \( MP \) activity**

MP expression levels above a maximum threshold both activate and repress \( \textit{ATHB8} \) preprocambial expression (Fig. 3). Activation of \( \textit{ATHB8} \) preprocambial expression by MP is direct [16], but repression of \( \textit{ATHB8} \) preprocambial expression by MP need not be: \( MP \)-dependent repression of \( \textit{ATHB8} \) preprocambial expression could be mediated, for example, by an AUXIN/INDOLE-3-ACETIC-ACID-INDUCIBLE (AUX/IAA) protein such as BODENLOS (BDL)/IAA12 (BDL hereafter), whose expression is activated by MP and which binds to MP and inhibits its transcriptional activity [30, 45–47]. Were \( MP \)-dependent repression of \( \textit{ATHB8} \) preprocambial expression mediated by BDL, \( \textit{ATHB8} \) preprocambial expression would be reduced in the \( bdl \) mutant, in which the unstable BDL protein is stabilized [41]. To test this prediction, we quantified \( \textit{ATHB8}:\text{nYFP} \) expression levels in second loops of 4-DAG first leaves of the \( bdl \) mutant.

As in \( mp \), in \( bdl \) levels of \( \textit{ATHB8}:\text{nYFP} \) preprocambial expression levels were lower and expression along the domain was more heterogeneous than in WT, leading to seemingly fragmented domains of weak \( \textit{ATHB8}:\text{nYFP} \) preprocambial expression (Fig. 3A–C,F; Fig. 4A,B,I). Moreover, as in \( mp \), defects in \( \textit{ATHB8} \) expression in \( bdl \) developing leaves were associated with networks of fewer meshes and fewer, less frequently continuous, and less frequently connected veins in \( bdl \) mature leaves (Fig. 3G–I,K; Fig. 4J,K,O).

Were \( MP \)-dependent repression of \( \textit{ATHB8} \) preprocambial expression mediated by an AUX/IAA protein such as BDL, reducing or eliminating AUX/IAA-
Fig. 3 (See legend on next page.)
mediated inhibition of MP transcriptional activity would lead to higher levels of \textit{ATHB8} preprocambial expression in both vein and flanking cells, resulting in broader \textit{ATHB8} expression domains. To test this prediction, we turned the unstable BDL transcriptional repressor into a stabilized transcriptional activator as previously done for other AUX/IAA proteins [48–50]: we replaced the repressor domain of BDL [51] with the activator domain of the \textit{Herpes simplex} Virus Protein 16 (VP16) [35] and introduced a mutation that lengthens the half-life of BDL [45]. We expressed the resulting VP16:bdl\textDelta I by the MP promoter in the \textit{iiaa12-1} mutant, which lacks \textit{BDL} function [36], and the \textit{iiaa12-1;}\textit{tpl-1} double mutant, which in addition partially lacks the co-repressor function that mediates the AUX/IAA-protein-dependent repression of MP [52]. We quantified \textit{ATHB8};\textit{nYFP} expression levels in second loops of 4-DAG first leaves of the resulting MP::VP16:bdl\textDelta I;iiaa12-1;ltp1-1 background.

As in MP::MP, in both MP::VP16:bdl\textDelta I;iiaa12-1 and MP::VP16:bdl\textDelta I;iiaa12-1;ltp1-1 — but not in \textit{iiaa12-1} — \textit{ATHB8};\textit{nYFP} expression levels were higher in flanking cells (Fig. 3A,D,F; Fig. 4A,C,I; Additional File 1: Figure S6). Unlike in MP::MP, however, in both MP::VP16: bdl\textDelta I;iiaa12-1 and MP::VP16:bdl\textDelta I;iiaa12-1;ltp1-1, \textit{ATHB8};\textit{nYFP} expression levels were also higher in vein cells (Fig. 3A,D,F; Fig. 4A,C,I; Additional File 1: Figure S6). Accordingly, stronger \textit{ATHB8} expression domains in MP::VP16:bdl\textDelta I;iiaa12-1;ltp1-1 developing leaves were associated with stronger — though qualitatively similar — vein network defects in MP::VP16:bdl\textDelta I;iiaa12-1;ltp1-1 mature leaves: in the middle of these leaves, veins ran parallel to one another for the entire length of the narrow leaf laminae to give rise to wide midveins; toward the margin, veins ran close to one another for varying stretches of the laminae, then diverged, and either ran close to other veins or converged back to give rise to elongated meshes (Fig. 3G,J,K; Fig. 4L).

Next, we further tested the prediction that reducing or eliminating AUX/IAA-mediated inhibition of MP transcriptional activity would lead to higher levels of \textit{ATHB8} preprocambial expression in both vein and flanking cells, resulting in broader \textit{ATHB8} expression domains. As previously done [29, 53, 54], we created an irreplaceable version of MP by deleting its PHOX/BEM1 (PB1) domain, which is required for AUX/IAA-mediated inhibition [49, 53, 55, 56]. We fused the resulting MP\textDelta PB1 to a fragment of the rat glucocorticoid receptor (GR) [57] to confer dexamethasone (dex)-inducibility, expressed the resulting MP\textDelta PB1:GR by the \textit{MP} promoter, and quantified \textit{ATHB8};\textit{nYFP} expression levels in 4-DAG first leaves of the dex-grown MP::MP\textDelta PB1:GR background.

Consistent with previous observations [53, 58], in dex-grown MP::MP\textDelta PB1:GR, \textit{ATHB8};\textit{nYFP} expression was no longer restricted to narrow stripes; instead, \textit{ATHB8};\textit{nYFP} was expressed at higher levels in broad bands than spanned almost the entire width of the leaves (Fig. 4D,E,I). Accordingly, broader and stronger \textit{ATHB8} expression domains in dex-grown MP::MP\textDelta PB1:GR developing leaves were associated with veins running parallel to one another for the entire length of the narrow leaf laminae to give rise to midveins that spanned almost the entire width of dex-grown MP::MP\textDelta PB1:GR mature leaves (Fig. 4M–O).

Broader and stronger \textit{ATHB8} expression domains in dex-grown MP::MP\textDelta PB1:GR leaves may be the result of the leaves’ vein pattern defects, rather than of the reduction in AUX/IAA-mediated inhibition of MP-dependent activation of \textit{ATHB8} expression. To test this possibility, we leveraged two observations: (1) \textit{ATHB8} preprocambial expression is activated asynchronously in second loops during leaf development [13]; (2) by the time a vein has activated \textit{ATHB8} preprocambial expression, the vein’s position has been specified [59]. We therefore germinated and grew \textit{ATHB8};\textit{nYFP};MP::MP\textDelta PB1:GR in the absence of dex for 3.75 days, transferred the seedlings to dex-containing medium for 6 h, and quantified \textit{ATHB8};\textit{nYFP} expression levels in the newly formed second loops of 4-DAG first leaves. Because in 3.75-DAG first leaves, \textit{ATHB8} is expressed in midvein, first loops, and only one of the two second loops (Fig. 4F), the
Fig. 4 (See legend on next page.)
position of those veins can no longer be changed by dex-mediated activation of MP::MPAPB1:GR. As such, any activation of ATHB8 expression in the second loops formed after the dex-mediated activation of MP::MPAPB1:GR would only be the result of the reduction in AUX/IAA-mediated inhibition of MP-dependent activation of ATHB8 expression.

Consistent with what shown above (Fig. 4D,E,I), ATHB8::nYFP expression in the second loops formed after the dex-mediated activation of MP::MPAPB1:GR was no longer restricted to narrow stripes; instead, ATHB8::nYFP was expressed at higher levels in broad bands (Fig. 4F,G,I). These results are consistent with the interpretation that broader and stronger ATHB8 expression domains in dex-grown MP::MPAPB1:GR leaves (Fig. 4D,E,I) are the result of the reduction in AUX/IAA-mediated inhibition of MP-dependent activation of ATHB8 expression, rather than of the leaves’ vein pattern defects.

Our results suggest that MP-dependent repression of ATHB8 preprocambial expression is mediated by AUX/IAA proteins, including BDL (Fig. 4A–G,I). However, that BDL mediates MP-dependent repression of ATHB8 preprocambial expression is based upon the assumption that the lower levels of ATHB8 preprocambial expression in the dominant bdl mutant reflect hypermorphic, as opposed to neomorphic, effects of the bdl mutation. Were BDL indeed mediating MP-dependent repression of ATHB8 preprocambial expression, BDL expression domains would overlap with domains of ATHB8 preprocambial expression. To test whether that were so, we imaged expression of BDL::nGFP and BDL::BDL::GUS in 4-DAG first leaves.

Contrary to expectations, BDL::nGFP and BDL::BDL::GUS were only expressed in midvein and first loops and were expressed neither in second loops nor in their flanking cells (Fig. 4P,Q). We therefore asked whether the bdl mutation affected BDL expression. To address this question, we imaged GUS activity in 4-DAG first leaves of a BDL::BDL::GUS line that recapitulates the bdl phenotype [41, 60].

BDL::BDL::GUS was strongly expressed in midvein and first loops; in the top half of the leaf, BDL::BDL::GUS was also expressed in the inner nonvascular tissue, though expression was weaker than in midvein and first loops (Fig. 4R). In the bottom half of the leaf, BDL::BDL::GUS was strongly expressed in both epidermis and inner tissue, including the areas where second loops were forming.

Broad expression of BDL::BDL::GUS may be the result of the leaves’ vein network defects, rather than of an effect of the bdl mutation on BDL expression. To test this possibility, we generated a BDL::BDL::YFP line that expresses the transgene at low levels and that therefore leads to only very minor vein network defects (Fig. 4Q). We then imaged expression of BDL::BDL::YFP in 4-DAG first leaves.

The expression of BDL::BDL::YFP mirrored that of BDL::BDL::GUS, including expression in second loops and their flanking cells (Fig. 4S), suggesting that broad expression of BDL::BDL::GUS is the result of an effect of the bdl mutation on BDL expression, rather than of the leaves’ vein network defects. Moreover, these observations suggest neomorphic, as opposed to hypermorphic, effects of the bdl mutation on ATHB8 preprocambial expression.

In conclusion, our results are consistent with the hypothesis that MP expression levels above a maximum threshold both activate and repress ATHB8 preprocambial expression and that such MP-dependent repression of ATHB8 preprocambial expression is mediated by AUX/IAA proteins; such AUX/IAA proteins, however, are unlikely to include BDL.

Relation between ATHB8 expression domains and auxin levels
AUX/IAA proteins are degraded in response to the plant hormone auxin [41, 48, 61, 62]. Auxin-dependent degradation of AUX/IAA proteins releases MP from
inhibition, thus allowing MP to activate expression of its targets, including AUX/IAA genes and ATHB8 [16, 30, 46, 47, 53, 58, 63–67]. Therefore, narrow stripes of ATHB8 preprocambial expression should correspond to peak levels of sensed auxin. To test this prediction, we simultaneously imaged in midvein, first loops, and second loops of developing first leaves expression of ATHB8::nQFP (nuclear Turquoise Fluorescent Protein expressed by the ATHB8 promoter) and of the auxin ratiometric reporter R2D2 [42], which expresses an auxin-degradable nYFP and a non-auxin-degradable nRFp by the RIBOSOMAL PROTEIN S5A promoter, which is highly active in developing leaves [68]. In the R2D2 reporter, a high RFP/YFP ratio thus indicates high levels of auxin, whereas a low RFP/YFP ratio indicates low levels of auxin [42].

At all tested stages, the RFP/YFP ratio was higher in ATHB8::nQFP-expressing cells than in cells flanking ATHB8::nQFP-expressing cells (Fig. 5), suggesting that domains of ATHB8 preprocambial expression correspond to peak levels of sensed auxin.

Response of ATHB8 expression to manipulation of MP-binding site affinity

The hypothesis that MP expression levels below a minimum threshold are unable to activate ATHB8 preprocambial expression predicts that reducing the affinity of MP for its binding site in the ATHB8 promoter will lead to extremely weak, or altogether absent, ATHB8 preprocambial expression.

To test this prediction, we mutated the MP-binding site in the ATHB8 promoter (TGTCTG) to lower (TGTCAG) or negligible (TAGCTG) affinity for MP binding [16, 69–71], and imaged nYFP expressed by the native or mutant promoters in second loops of 4-DAG first leaves.

Mutation of the MP-binding site in the ATHB8 promoter to negligible affinity for MP binding led to greatly reduced levels of nYFP expression (Fig. 6A,B,F), resembling near-complete loss of ATHB8:nYFP preprocambial expression in mp-U55 [16] (Fig. 3A,B,F). Mutation of the MP-binding site in the ATHB8 promoter to lower affinity for MP binding led to lower levels of nYFP expression (Fig. 6A,C,F). Furthermore, expression along the domains was more heterogeneous than when nYFP was expressed by the native promoter (Fig. 6A,C,F), leading to seemingly fragmented domains of weak nYFP expression similar to those in mp-11 (Fig. 3A,C,F) and bdl (Fig. 4A,B,I).

The hypothesis that MP expression levels above a maximum threshold both activate and repress ATHB8 preprocambial expression predicts that increasing the affinity of MP for its binding site in the ATHB8 promoter will lead to higher levels of ATHB8 preprocambial expression in flanking cells, leading to broader ATHB8 expression domains, and to levels of ATHB8 preprocambial expression in vein cells that are no lower — though not necessarily any higher — than those in WT.

To test this prediction, we mutated the MP-binding site in the ATHB8 promoter (TGTCTG) to higher (TGTCCTC) affinity for MP binding [16, 69, 70], and imaged nYFP expressed by the native or mutant promoter in second loops of 4-DAG first leaves.

Mutation of the MP-binding site in the ATHB8 promoter to higher affinity for MP binding led to higher levels of nYFP expression in flanking cells (Fig. 6A,D,F), resulting in broad bands of nYFP expression similar to those in MP::MP (Fig. 3A,D,F) and, to a lesser extent, MP::VP16:bdld;iaa12-1 (Additional File 1: Fig. S6B), MP::VP16:bdld;iaa12-1:tpl-1 (Fig. 4A,C,I), and dex-grown MP::MPΔPB1:GR (Fig. 4D–G,I). However, unlike in MP::MP — in which ATHB8:nYFP expression levels in vein cells were lower than in WT (Fig. 3A,D,F) — and unlike in MP::VP16:bdld;iaa12-1, MP::VP16:bdld;iaa12-1:tpl-1, and dex-grown MP::MPΔPB1:GR — in which those levels were higher (Fig. 4A,C–G,I; Additional File 1: Fig. S6A,B) — nYFP expression levels in vein cells were unchanged by mutation of the MP-binding site in the ATHB8 promoter to higher affinity for MP binding (Fig. 6A,D,F), suggesting that MP levels are normally nonlimiting for ATHB8 preprocambial expression.

In conclusion, our results are consistent with the hypothesis that MP expression levels below a minimum threshold are unable to activate ATHB8 preprocambial expression and that MP expression levels above a maximum threshold both activate and repress ATHB8 preprocambial expression.

Discussion

A long-standing problem in biology is how gene expression is activated in narrow stripes by broadly expressed transcription factors (e.g., [72, 73]). Here we addressed this problem for plants by means of the MP – ATHB8 pair of Arabidopsis genes.

Consistent with interpretation of similar findings in animals (e.g., [74–76]), our results suggest that levels of expression of the MP transcription factor above a maximum threshold both activate and repress ATHB8 preprocambial expression. MP-dependent activation of ATHB8 expression is direct [16] and — we found — mediated by binding of MP to a low-affinity site in the ATHB8 promoter. By contrast, we found that MP-dependent repression of ATHB8 expression is indirect and mediated by members of the AUX/IAA family, which are themselves direct targets of MP [47, 64]. AUX/IAA proteins inhibit MP transcriptional activity and are degraded at peak levels of the plant hormone.
Fig. 5  ATHB8 Expression Domains and Auxin Levels. First leaves 2, 3, and 4 DAG. Columns 1–3: confocal laser scanning microscopy. Column 1: ATHB8::nQFP expression (red) associated with formation of midvein (2 DAG), first loop (3 DAG), or second loop (4 DAG) [16, 22, 44]. Column 2: Ratio of RPS5A::mDII:nRFP expression to RPS5A::DII:nYFP expression. Look-up table visualizes expression ratio levels: high RPS5A::mDII:nRFP/RPS5A::DII:nYFP ratio (green) indicates high auxin levels; low RPS5A::mDII:nRFP/RPS5A::DII:nYFP ratio (blue) indicates low auxin levels. Column 3: overlays of images in columns 1 and 2; blue: low RPS5A::mDII:nRFP/RPS5A::DII:nYFP ratio, i.e. low auxin levels; yellow: co-expression of ATHB8::nQFP (red) and high RPS5A::mDII:nRFP/RPS5A::DII:nYFP ratio (green), i.e. high auxin levels. Column 4: Ratio of RPS5A::mDII:nRFP expression levels to RPS5A::DII:nYFP expression levels (mean ± SE) in nuclei flanking ATHB8::nQFP-expressing nuclei (positions -2, -1, +1, and +2) relative to ratio of RPS5A::mDII:nRFP expression levels to RPS5A::DII:nYFP expression levels in nuclei co-expressing ATHB8::nQFP (position 0) during formation of midvein (top), first loop (middle), or second loop (bottom). Difference between ratio of RPS5A::mDII:nRFP expression levels to RPS5A::DII:nYFP expression levels in nuclei at position -2, -1, +1, or +2 and ratio of RPS5A::mDII:nRFP expression levels to RPS5A::DII:nYFP expression levels in position 0 was significant at P < 0.01 (**) or P < 0.001 (***) by One-Way ANOVA and Tukey’s Pairwise test. Sample sizes: 26 (2 DAG), 27 (3 DAG), or 29 (4 DAG) leaves; position -2: 56 (2 DAG), 42 (3 DAG), or 60 (4 DAG) nuclei; position -1: 52 (2 DAG), 37 (3 DAG), or 58 (4 DAG) nuclei; position 0: 74 (2 DAG), 85 (3 DAG), or 102 (4 DAG) nuclei; position +1: 44 (2 DAG), 44 (3 DAG), or 62 (4 DAG) nuclei; position +2: 42 (2 DAG), 25 (3 DAG), or 44 (4 DAG) nuclei. Scale bars (shown, for simplicity, only in column 2): 5 μm
auxin [30, 41, 48, 53, 58, 61–63, 66, 67] such as those we found corresponding to narrow stripes of ATHB8 preprocambial expression. As such, our results suggest that an incoherent type-I feedforward loop [77] restricts activation of ATHB8 preprocambial expression to narrow stripes: auxin activates MP, which in turn activates expression of intermediate-loop AUX/IAA genes; and MP and AUX/IAA genes jointly regulate expression of ATHB8, which converts the auxin signal input into vein formation output (Additional File 1: Figure S7).

Our finding that ATHB8 promotes vein formation both nonredundantly and redundantly with other HD-ZIP III genes is consistent with the observation that excess vein formation in the acaulis5 mutant depends on the function of ATHB8 and of the ATHB8-related REVOLUTA and ATHB15/CORONA genes [25].
Nevertheless, precisely how ATHB8 promotes vein formation remains unclear. Delayed vein formation in athb8 mutants [16] suggests that ATHB8 promotes timely vein formation, possibly preventing premature termination of vein formation by mesophyll differentiation [13]. Furthermore, because the athb8 mutation enhances the defects in coordination of cell polarity and vein patterning induced by the inhibition of the polar, cell-to-cell transport of auxin [16], it is possible that ATHB8 belongs to that auxin signaling pathway that controls coordination of cell polarity and vein patterning redundantly with polar auxin transport [78]. However, these possibilities remain to be tested.

Given the defects in ATHB8 preprocambial expression we observed in the bdl mutant, our finding that the AUX/IAA protein BDL is unlikely to be mediating MP-dependent repression of ATHB8 preprocambial expression is perhaps unexpected but certainly not unprecedented. Not only in veins — as we found — but in embryos too, the bdl mutation leads to expression of the bdl protein at stages earlier and in domains broader than those at and in which the BDL protein is expressed [63]. Furthermore, a mutation in the CRANE/IAA18 gene that, just like the bdl mutation, stabilizes the resulting mutant protein also leads to expression of the crane-2/iaa18-1 mutant protein at stages earlier and in domains broader than those at and in which the CRANE/IAA18 protein is expressed [66, 79]. These observations reinforce the need for caution when interpreting phenotypes of dominant mutants as hypermorphic — as opposed to neomorphic — as it has often been done for dominant aux/iaa mutations. In the future, it will be interesting to identify which AUX/IAA proteins mediate MP-dependent repression of ATHB8 preprocambial expression; as interesting as that identification will be, however, it will also be unlikely to change the logic of the regulatory network that we propose restricts ATHB8 preprocambial expression to narrow stripes.

In the future, it will also be interesting to understand what generates peak levels of sensed auxin and of MP expression in the leaf. One possibility is that those peaks are the result of the polar, cell-to-cell transport of auxin, which seems to converge on positions of peak MP expression [20, 53, 80–85]. Consistent with this possibility, abnormal positions of MP expression domains in developing auxin-transport-inhibited leaves foreshadow the abnormal positions of veins in mature auxin-transport-inhibited leaves [59, 78, 82, 86]. One other possibility is that peak levels of MP expression arise from MP’s self-activation — as proposed to happen during embryogenesis [47] and flower formation [87] — and the levels of MP expression we measured in MP::MP are consistent with this possibility. Yet another possibility is that — as proposed to happen during xylem differentiation in the leaf [25] or as the ATHB8-related PHABULOSA does in the root [88] — ATHB8 controls MP expression, such that interpretation of positional information feed back on generation of that information, as it often happens in animals (reviewed in [89]). Broader expression domains of an MP expression reporter in athb8 leaves [16, 90] are consistent with such a possibility. All these possibilities will have to be considered in future work to test whether the gene regulatory network our results suggest is required for restriction of ATHB8 preprocambial expression to narrow stripes is also sufficient for it.

Finally, it will be interesting to understand whether the incoherent feedforward loop we propose restricts activation of ATHB8 preprocambial expression to narrow stripes also controls the striped expression of ATHB8 in other organs and the striped expression of other genes in plants.

Conclusions

Our results suggest a mechanism by which in plants a broadly expressed transcription factor — MP — activates expression of a target gene — ATHB8 — in narrow stripes. The very same regulatory mechanism that controls activation of ATHB8 preprocambial expression in single files of cells is most frequently used in animals to generate stripes of gene expression [91], suggesting unexpected conservation of regulatory logic of striped gene expression in plants and animals despite the independent evolution of their multicellularity. Nevertheless, in animals, such regulatory logic typically leads to activation of target gene expression in a stripe that is outside the expression domain of the activating transcription factor (e.g., [74–76, 92]), whereas ATHB8 expression is activated in a stripe that is a subset of the MP expression domain. It will be interesting to understand whether these are plant- and animal-specific outputs of the same conserved regulatory logic.

Methods

Plants

Origin and nature of lines, genotyping strategies, and oligonucleotide sequences are in Additional File 2: Table S1, Additional File 2: Table S2, and Additional File 2: Table S3, respectively. Seeds were sterilized and sowed in [93]. Stratified seeds were germinated and seedlings were grown at 22 °C under continuous light (~90 μmol m⁻² s⁻¹). To induce MPΔPB1:GR translocation to the nucleus, seeds were sown on, or 3.75-DAG seedlings were transferred to, dex-supplemented medium (30 μM final concentration). Plants were grown at 25 °C under fluorescent light (~100 μmol m⁻² s⁻¹) in a 16-h-light/8-h-dark cycle and transformed as in [93]. For each construct generated in this study (see Additional File 2: Table S1), the progeny of at least 10 independent
transgenic lines were inspected to identify the most representative leaf expression pattern or vein network phenotype. Detailed analysis was performed on the progeny of two homozygous lines per construct. Such representative lines were selected because of strong expression or phenotype emblematic of the profile observed across the entire transgenic series and resulting from single transgene insertion. The same ATHB8::nYFP line (generated in WT background) [16] was introduced in all genetic backgrounds by crossing.

RT-qPCR
Total RNA was extracted with Qiagen’s RNeasy Plant Mini Kit from 4-day-old seedlings grown in half-strength Murashige and Skoog salts, 15 g l⁻¹ sucrose, 0.5 g l⁻¹ MES, pH 5.7, at 23 °C under continuous light (~80 μmol m⁻² s⁻¹) on a rotary shaker at 50 rpm. DNA was removed with Invitrogen’s TURBO DNA-free kit, and RNA was stabilized by the addition of 20 U of Thermo Fisher Scientific’s Superase-In RNase Inhibitor. First-strand cDNA was synthesized from ~100 ng of DNase-treated RNA with Thermo Fisher Scientific’s RevertAid Reverse Transcriptase according to the manufacturer’s instructions, except that 50 pmol of Thermo Fisher Scientific’s Oligo(dT)₁₈ Primer, 50 pmol of Thermo Fisher Scientific’s Random Hexamer Primer, and 20 U of Superase-In RNase Inhibitor were used. qPCR was performed with Applied Biosystems’ 7500 Fast Real-Time PCR System on 2 μl of 1:3-diluted cDNA with 5 pmol of each gene-specific primers (Additional File 2: Table S3), 2.5 pmol of gene-specific probe (Additional File 2: Table S3), and Applied Biosystems’ TaqMan 2X Universal PCR Master Mix in a 10-μl reaction volume. Probe and primers were designed with Applied Biosystems’ Primer Express. Relative MP transcript levels were calculated with the 2⁻ΔΔCt method [94] using ACTIN2 transcript levels for normalization.

Imaging
For confocal laser scanning microscopy, developing leaves were mounted and imaged as in [95], except that emission was collected from ~1.5–5.0-μm-thick optical slices. In single-fluorophore marker lines, YFP was excited with the 514-nm line of a HeNe laser, and emission was collected with a BP 520–555 filter. In multiple-fluorophore marker lines, CFP, QFP, and auto-fluorescent compounds were excited with the 458-nm line of a 30-mW Ar laser, YFP was excited with the 514-nm line of a 30-mW Ar laser, and RFP was excited with the 543-nm line of a HeNe laser. CFP and QFP emission were collected with a BP 475–525 filter, YFP emission was collected with a BP 520–555 filter; RFP emission was collected between 581 and 657 nm, and autofluorescence was collected between 604 and 700 nm. Signal intensity levels of 8-bit grayscale images acquired at identical settings were quantified in the Fiji distribution of ImageJ [96–99]. To visualize RFP/YFP ratios, the histogram of the YFP images was linearly stretched in the Fiji distribution of ImageJ such that the maximum gray value of the YFP images matched that of the corresponding RFP images, and the RFP images were divided by the corresponding YFP images. GUS activity in developing leaves was detected as in [13]. Stained leaves were fixed, cleared, and mounted as in [13], and mounted leaves were imaged with a Zeiss AxioImager.M1 microscope equipped with a QImaging MicroPublisher 5.0 RTV camera. Mature leaves were fixed, cleared, and mounted as in [54, 78], and mounted leaves were imaged as in [33]. Image brightness and contrast were adjusted by linear stretching of the histogram in in the Fiji distribution of ImageJ.

Vein network analysis
The cardinality, continuity, and connectivity indices were calculated as in [20]. Briefly, the number of “touch points” (TPs, where a TP is the point where a vein end contacts another vein or a vein fragment), “end points” (EPs, where an EP is the point where an “open” vein — a vein that contacts another vein only at one end — terminates free of contact with another vein or a vein fragment), “break points” (KPs, where a KP is each of the two points where a vein fragment terminates free of contact with veins or other vein fragments), and “exit points” (XPs, where an XP is the point where a vein exits leaf blade and enters leaf petiole) in dark-field images of cleared mature leaves was calculated with the Cell Counter plugin in the Fiji distribution of ImageJ. Because a vein network can be understood as an undirected graph in which TPs, EPs, KPs, and XPs are vertices, and veins and vein fragments are edges, and because each vein is incident to two TPs, a TP and an XP, a TP and an EP, or an XP and an EP, the cardinality index — a measure of the size (i.e., the number of edges) of a graph — is a proxy for the number of veins and is calculated as [(TPs + XPs — EPs)/2] + EPs, or (TPs + XPs + EPs)/2. The continuity index quantifies how close a vein network is to a network with the same number of veins, but in which at least one end of each vein fragment contacts a vein and is therefore calculated as the ratio of the cardinality index of the first network to the cardinality index of the second network: [(TP + XP + EP)/2]/[(TP + XP + EP + KP)/2], or (TP + XP + EP)/(TP + XP + EP + KP). The connectivity index quantifies how close a vein network is to a network with the same number of veins, but in which both ends of each vein or vein fragment contact other veins, and is therefore calculated as the ratio of the number of “closed” veins — those veins which contact vein fragments or other veins at both ends — in
the first network to the number of closed veins in the second network (i.e., the cardinality index of the second network): \[(TP + XP − EP)/(TP + XP + EP + KP)/2\], or \((TP + XP − EP)/(TP + XP + EP + KP)\). Finally, because the number of meshes in a vein network equals the number of closed veins, the cyclic index — a proxy for the number of meshes in a vein network — is calculated as: \((TP + XP − EP)/2\).

**Abbreviations**

ATHB: ARABIDOPSIS THALIANA HOMEOBOX; AUX/IAA: AUXIN/INDOLE-3-ACETIC-ACID-ININDUCIBLE; BDL: BODENLOS; CFP: Cyan fluorescent protein; DAG: Days after germination; dex: Dexamethasone; EAR: Ethylene-responsive-Acetate-Inducible; BDL: BODENLOS; CFP: Cyan fluorescent protein; GFP: Green fluorescent protein; GR: Glucocorticoid receptor; GUS: Green Fluorescent Protein; MP: MONOPTEROS; PB1: PHOX/BEM1; QFP: Turquoise fluorescent protein; REP: Red fluorescent protein; RPS5A: RBOSOMAL PROTEIN S5A; SHR: SHORT-ROOT; VP16: Virus protein 16; WT: Wild type; YFP: Yellow fluorescent protein

**Supplementary Information**

The online version contains supplementary material available at https://doi.org/10.1186/s12915-021-01143-9.

Acknowledgements

We thank the Arabidopsis Biological Resource Center for seeds of athb8-37, iaa12-1, and iaa12-1. Dolf Weijers for seeds of mp-B4149 and R202, Dolf Weijers and Gerd Jürgens for BDLC::GFP and BDLC::BDLGUS seeds, Dolf Weijers and Raju Data for BDLC::BDLC::GFP seeds, Hiroo Fukuda and Kyoko Ohashi-Ito for SHR::mATHB8 DNA, and Zachary Nimchuk for nQFP DNA. We thank Neil Harris and Przemek Prusinkiewicz for helpful comments on the manuscript.

Authors’ contributions

AK, JLD conceptualization, formal analysis, validation, investigation, visualization, methodology, writing — original draft, writing — review and editing; ES: conceptualization, resources, formal analysis, supervision, funding acquisition, validation, investigation, visualization, methodology, writing — original draft, project administration, writing — review and editing. All authors read and approved the final manuscript.

**Funding**

This work was supported by Discovery Grants of the Natural Sciences and Engineering Research Council of Canada (NSERC) (RGPIN-2016-04736 to ES). AK was supported, in part, by an NSERC CGS-M Scholarship. JLD was supported, in part, by an NSERC CGS-M scholarship and an NSERC PGS-D Scholarship. TJD was supported, in part, by an NSERC CGS-D Scholarship and an Alberta Ingenuity Student Scholarship. The funders had no role in study design, data collection and interpretation, or the decision to submit the work for publication.

**Availability of data and materials**

All data generated or analyzed during this study are included in this published article and its supplementary information files, or are available from the corresponding author on reasonable request.

**Declarations**

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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**Received**

23 October 2020 Accepted: 3 September 2021 Published online: 24 September 2021

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