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Antagonistic Interaction of BLADE-ON-PETIOLE1 and 2 with BREVIPEDICELLUS and PENNYWISE Regulates Arabidopsis Inflorescence Architecture^{1[C][W][OA]}

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The transition to flowering in many plant species, including *Arabidopsis thaliana*, is marked by the elongation of internodes to make an inflorescence upon which lateral branches and flowers are arranged in a characteristic pattern. Inflorescence patterning relies in part on the activities of two three-amino-acid loop-extension homeodomain transcription factors: BREVIPEDICELLUS (BP) and PENNYWISE (PNY) whose interacting products also promote meristem function. We examine here the genetic interactions between BP-PNY whose expression is up-regulated in stems at the floral transition, and the lateral organ boundary genes BLADE-ON-PETIOLE1 (BOP1) and BOP2, whose expression is restricted to pedicel axils. Our data show that *bp* and *pnw* inflorescence defects are caused by BOP1/2 gain of function in stems and pedicels. Compatible with this, inactivation of BOP1/2 rescues these defects. BOP expression domains are differentially enlarged in *bp* and *pnw* mutants, corresponding to the distinctive patterns of growth restriction in these mutants leading to compacted internodes and clustered or downward-oriented fruits. Our data indicate that BOP1/2 are positive regulators of KNOTTED1-LIKE FROM ARABIDOPSIS THALIANA6 expression and that growth restriction in BOP1/2 gain-of-function plants requires KNOTTED1-LIKE FROM ARABIDOPSIS THALIANA6. Antagonism between BOP1/2 and BP is explained in part by their reciprocal regulation of gene expression, as evidenced by the identification of lignin biosynthetic genes that are repressed by BP and activated by BOP1/2 in stems. These data reveal BOP1/2 gain of function as the basis of *bp* and *pnw* inflorescence defects and reveal how antagonism between BOP1/2 and BP-PNY contributes to inflorescence patterning in a model plant species.

Flowering plants display a remarkable variety of inflorescence architectures selected to optimally display flowers for pollination and seed dispersal. Formation of the aerial parts of a plant is controlled by the shoot apical meristem (SAM), a cluster of pluripotent stem cells located at the apex of the primary shoot. The SAM produces a series of reiterative modules known as phytomers to generate the aerial parts of the plant.

Each phytomer comprises an internode (stem) subtending a node, which is a leaf associated with a potential axillary meristem (Steeves and Sussex, 1989). Elaboration of the different parts of a module (leaves, internodes, and axillary meristems) varies according to the phase of development and between species to generate architectural diversity (Sussex and Kerk, 2001).

Arabidopsis thaliana has distinct vegetative and reproductive phases. During vegetative development, the SAM generates leaf primordia on its flanks; both internode and axillary meristem formation are inhibited, resulting in a compact rosette of leaves. At the end of the vegetative phase, endogenous and environmental cues promote the transition to flowering. The SAM responds to floral inductive signals by acquiring inflorescence meristem (IM) fate. During reproductive development, internodes elongate and axillary meristems proliferate at the expense of leaves to generate lateral branches and flowers in a regular spiral pattern on the inflorescence (Bowman and Eshed, 2000; Fletcher, 2002; Barton, 2010). While the pathways that promote floral fate of axillary meristems and repress leaf development are well studied, less is known about the formation and patterning of internodes.

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Internode patterning is a key determinant of inflorescence architecture, with variations in the length and pattern of internode elongation contributing to diversity in inflorescence height and organization of secondary branches and flowers on the primary stem. Formation of internodes is associated with the proliferation and elongation of cells in the region underlying the central zone of the meristem, termed the rib zone (Steeves and Sussex, 1989; Fletcher, 2002). Following their elongation, internodes are gradually fortified through the differentiation of interfascicular fibers with secondary thickened cell walls, which provides mechanical support (Nieminen et al., 2004; Ehling et al., 2005).

Internode patterning is dependent in part on the overlapping activities of two three-amino-acid loop-extension homeodomain transcription factors: the class I KNOTTED1-like homeobox (KNOX) protein BREVIPEDICELLUS (BP; formerly KNOTTED1-LIKE FROM ARABIDOPSIS THALIANA1 [KNAT1]) and the BEL1-like (BELL) protein PENNYWISE (PNY; also called BELLRINGER, REPLUMLESS, and VAMAANA) whose interacting products also promote meristem maintenance (Douglas et al., 2002; Venglat et al., 2002; Byrne et al., 2003; Smith and Hake, 2003; Bhatt et al., 2004; Rutjens et al., 2009; for review, see Hamant and Pautot, 2010). Mutations in BP cause short internodes and downward-pointing pedicels (Douglas et al., 2002; Venglat et al., 2002) whereas mutations in PNY cause irregular internode elongation, resulting in clusters of lateral organs (branches and flowers) spaced along the inflorescence (Byrne et al., 2003; Smith and Hake, 2003). In *bp pny* double mutants internodes are shorter than in either single mutant, signifying that BP and PNY have only partly overlapping roles in internode elongation and patterning (Smith and Hake, 2003). In both mutants, defects in vascular differentiation also occur, resulting in changes in how lignin is deposited in stems (Douglas et al., 2002; Mele et al., 2003; Smith and Hake, 2003). Previous genetic studies have shown that two class I KNOX genes, *KNAT2* and *KNAT6*, are misexpressed in *bp* and *pny* mutant stems and pedicels. Inactivation of these genes, primarily *KNAT6*, rescues *bp* and *pny* defects in inflorescence architecture (Ragni et al., 2008) however this is the extent of our current knowledge.

Here, we examine genetic interactions between BP-PNY and BLADE-ON-PETIOLE1 (BOP1) and BOP2, two BTB-ankryin transcriptional coregulators that are expressed in lateral organ boundaries (Ha et al., 2004; Hepworth et al., 2005). *BOP1/2* expression is limited to the pedicel axil in inflorescence stems where their function is to promote the formation of a vestigial abscission zone (McKim et al., 2008). *BOP1/2* are indirect transcriptional repressors of BP in leaves (Ha et al., 2007; Jun et al., 2010) but their genetic interactions with BP, and its partner PNY, during reproductive development have yet to be examined. We show here that BP and PNY are transcriptional repressors of *BOP1/2*, preventing expression in stems and pedicels. Consistent with this, inactivation of the *BOP* genes rescues *bp*

and *pny* inflorescence defects. We further show that *BOP1/2* exert their activity in part through the boundary gene *KNAT6*, which functions in the same genetic pathway. Finally, we show that *bp* and *pny* inflorescence defects are mimicked by *BOP1/2* gain of function. To explain this, we provide evidence that the reciprocal functions of BP and *BOP1/2* in the inflorescence are likely a consequence of their antagonistic regulation of downstream target genes, such as those involved in lignin biosynthesis that are repressed by BP and activated by *BOP1/2* in stems. These data redefine *bp* and *pny* phenotypes as the consequence of *BOP1/2* gain of function, shedding light on how interactions between BP-PNY and *BOP1/2* influence inflorescence architecture in a model plant species.

RESULTS

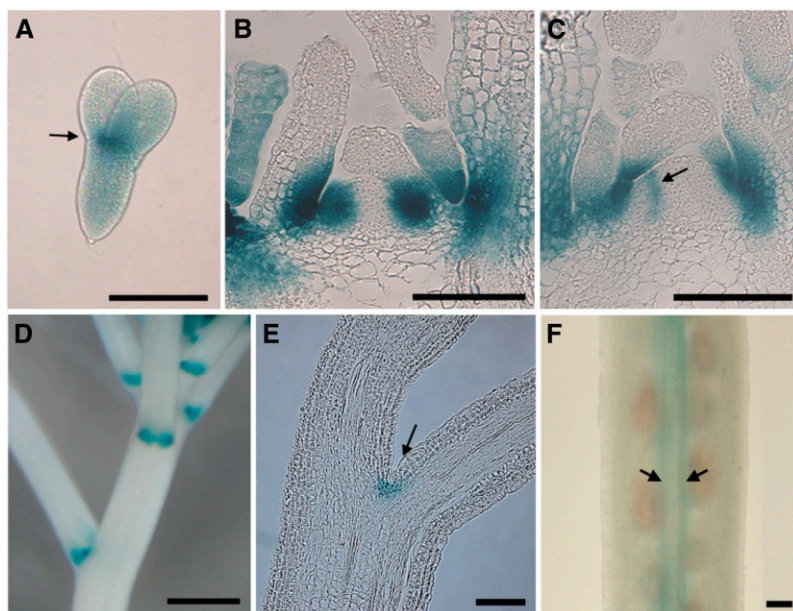
Expression of *BOP1* and *BOP2* in Lateral-Organ Boundaries

Previous analysis of *BOP* expression by in situ hybridization or through use of *BOP1:GUS* or *BOP2:GUS* reporter genes is consistent in showing that the *BOP* genes are expressed in lateral-organ boundaries formed during embryonic, vegetative, and reproductive development (Ha et al., 2004; Hepworth et al., 2005; Norberg et al., 2005; McKim et al., 2008; Xu et al., 2010). We have consolidated these data (Fig. 1). Using a *BOP2:GUS* reporter gene, expression was verified at the base of cotyledons in mature embryos (Fig. 1A; Ha et al., 2004). During postembryonic vegetative development, *BOP2* expression was first associated with the boundary at stage 2 of leaf development, when primordia first appear as morphologically distinct from the SAM (Fig. 1, B and C, arrow indicates stage 2 leaf). As leaves expand, expression associates with the adaxial base of leaves, which gives rise to the petiole (Fig. 1, B and C; Norberg et al., 2005). Expression is also observed in the axil of pedicels (Fig. 1, D and E; McKim et al., 2008) and in the valve margins of fruit (Fig. 1F). Importantly, *BOP1/2* expression is excluded from the IM and the replum of fruits, representing structures with meristematic function. While analysis of loss-of-function *bop1 bop2* mutants has revealed that *BOP1/2* transcriptionally repress meristematic genes in leaves (Ha et al., 2007) and floral primordia (Xu et al., 2010) relatively little is known about how *BOP1/2* gain of function perturbs plant architecture.

A Spectrum of Inflorescence Architecture Defects Caused by *BOP1/2* Gain of Function

Previous phenotypic analysis of *BOP1* or *BOP2* overexpression in plants has drawn attention to *bp*- and *pny*-like defects in inflorescence architecture, either short plants with floral pedicels pointing downward (Ha et al., 2007) or short bushy plants with irregular internodes (Norberg et al., 2005). Comparison of the strong activation-tagged *bop1-6D* line to *bp*

Figure 1. *BOP2:GUS* expression pattern in boundaries. A, Mature embryo; expression at base of cotyledons (arrow). B to C, Shoot apex of a short-day-grown seedling, longitudinal section; expression begins in stage 1 leaf primordia and localizes to the boundary of stage 2 leaves (arrow). As primordia expand, *BOP2* expression associates with the adaxial base of leaves, which elongate to form the petiole. D, Inflorescence; horseshoe pattern of expression in the axils of floral pedicels. E, Pedicel, longitudinal section; expression in the axil (arrow). F, Silique; expression in the valve margins (arrows). Scale bars, 0.1 mm except D, 0.5 mm. [See online article for color version of this figure.]



pn1 double mutants revealed remarkably similar inflorescence architectures (Fig. 2, A–C), suggesting that *BOP1/2* might antagonize both activities. This also suggested that *BOP1/2* gain of function might elicit a spectrum of inflorescence defects. To examine this further, we generated transgenic plants overexpressing *BOP1* or *BOP2* in Columbia-0 (*Col-0*) and Landsberg *erecta* (*Ler*) backgrounds and scored for defects in inflorescence architecture (Fig. 2, D–I; Table I). This analysis showed that in *Ler* plants, downward-pointing siliques was the prevalent phenotype (up to 45% of transformants) whereas in *Col-0* plants, clustered siliques was the prevalent phenotype (up to 20% of transformants). Compatible with this, the *erecta* mutation enhances the phenotypic severity of *bp* mutants (Douglas et al., 2002). Taken together, these findings suggest that *BOP* gain of function has variable effects on inflorescence architecture conditioned in part by ecotype. These defects may result from the antagonism of *BP* and/or *PNY* expression or activity. To examine this further, we tested the effect of *bop1 bop2* loss of function on expression of *bp* and *pn1* mutant phenotypes in a *Col* background.

Inactivation of *BOP1/2* Partially Rescues the *bp* Phenotype

To first examine *BOP1/2* interactions with *BP*, we generated *bop1 bop2 bp-1* and *bop1 bop2 bp-2* triple mutants and analyzed their phenotypes relative to wild-type and parental controls. *bp* mutants are characterized by short internodes, reduced apical dominance, and downward-pointing siliques (Douglas et al., 2002; Venglat et al., 2002). This analysis showed that inactivation of the *BOP* genes largely rescues *bp* inflorescence defects (Fig. 3, A–D; Supplemental Fig.

S1) similar to inactivation of *KNAT2* and *KNAT6* (Ragni et al., 2008). Quantitative phenotypic analyses were performed on 24 plants per genotype, by measuring the average height, internode length, and number of rosette paraclades for wild type and mutants (Fig. 4, A–D). These analyses confirmed that *bop1 bop2* loss of function counteracted the short stature of *bp-1* and *bp-2* plants (Fig. 4A) and partially restored apical dominance in *bp-1* mutants (Fig. 4B). Whereas *bp-1* mutants have a significant number of short internodes in the 1- to 5-mm range, the distribution in *bop1 bop2 bp-1* triple mutants was similar to wild type (Fig. 4C). Whereas *bp-1* pedicels point downward at an average angle of 84.9° relative to the primary stem, the average angle in *bop1 bop2 bp-1* triple mutants was 47.7°, similar to wild type (Fig. 4D). Also, the average pedicel angle in *bop1 bop2* double mutants was steeper than wild type (34.7° versus 50.3°), showing that *BOP1/2* regulate pedicel orientation as well as abscission zone formation at the stem-pedicel junction (Fig. 4D; Supplemental Fig. S2; McKim et al., 2008). No rescue occurred in *bp-2 bop1* or *bp-2 bop2* double mutants (data not shown), indicating that *BOP1* and *BOP2* have redundant functions.

Inactivation of *BOP1/2* Completely Rescues the *pn1* Phenotype

Given that *BP* and *PNY* coregulate internode patterning, we next examined the interaction of *BOP1/2* with *PNY* by generating *bop1 bop2 pn1* triple mutants. *pn1* mutants are characterized by clusters of siliques due to irregular internode elongation, defects in phyllotaxy, reduced apical dominance, and replumless fruits (Byrne et al., 2003; Roeder et al., 2003; Smith and Hake, 2003; Bhatt et al., 2004). Inactivation of the

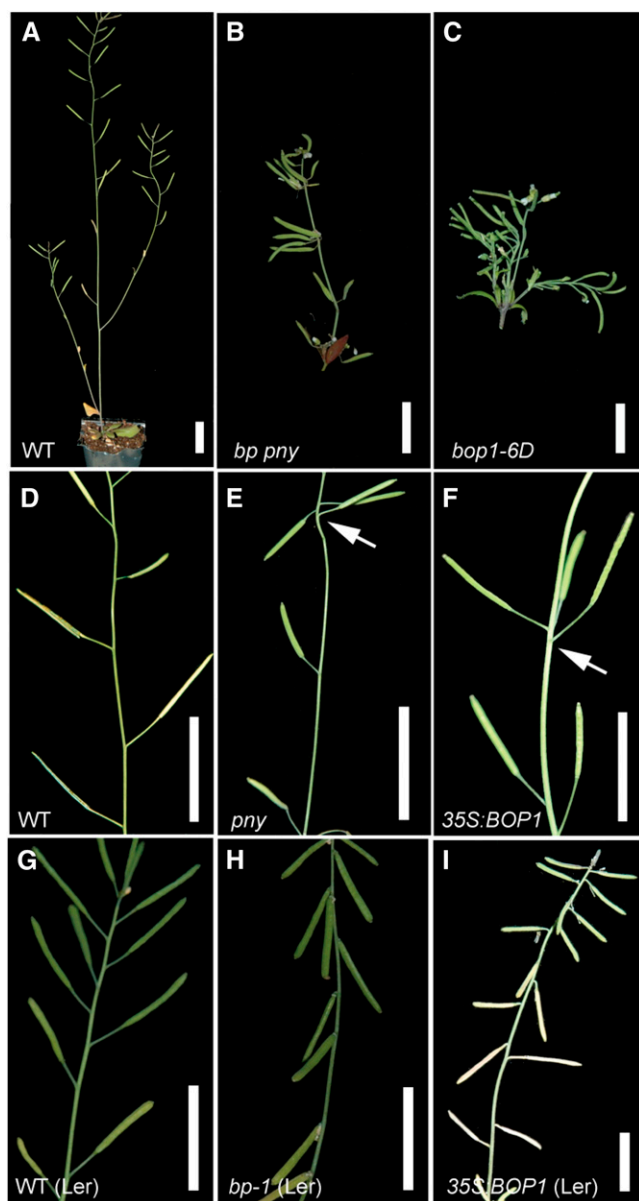


Figure 2. *BOP1* gain of function causes *bp*- and *pny*-like defects in inflorescence architecture. Representative inflorescences are shown for: A, Col wild type; B, *bp-2 pny* double mutant. C, *bop1-6D*; an activation-tagged *BOP1* overexpression line (with four 35S CaMV enhancers). Compact internodes similar to *bp-2 pny*. D, Col. E, *pny* mutant. F, 35S:*BOP1* transformant in Col (one 35S CaMV enhancer) with clustered siliques as in *pny* (arrows in E and F). G, *Ler* wild type. H, *bp-1* in *Ler*. I, 35S:*BOP1* transformant in *Ler*; downward-pointing siliques as in *bp-1*. Scale bars, 1 cm. [See online article for color version of this figure.]

BOP genes also rescued *pny* inflorescence defects (Fig. 3, A, E, and F). Quantitative phenotypic analyses were performed on 24 plants per genotype to further monitor this rescue, by measuring the average height, internode length, and number of rosette paraclades for wild type and mutants. These analyses confirmed that loss-of-function *bop1 bop2* restored the stature of *pny* plants and the number of rosette paraclades to wild

type (Fig. 5, A and B). Whereas *pny* mutants have a significant number of internodes in the 1- to 5-mm range, the distribution in *bop1 bop2 pny* triple mutants was similar to wild type (Fig. 5C). To quantify rescue of phyllotactic patterning in *bop1 bop2 pny* triple mutants, we measured divergence angles between successive floral pedicels on the primary stem (Fig. 5D; see Peaucelle et al., 2007). Whereas the distribution of divergence angles in *pny* was largely random (mean of 175°), the distribution in *bop1 bop2 pny* triple mutants was similar to wild type (mean of 142° versus 141°). Surprisingly, partial loss of *BOP* function was sufficient to rescue the *pny* phenotype since *pny bop1* and *pny bop2* mutant inflorescences also resembled wild type (data not shown).

A final defining characteristic in *pny* mutants is a replumless fruit (Roeder et al., 2003). Scanning electron microscopy (SEM) showed that inactivation of *BOP1/2* also rescues replum formation in *pny* fruits (Supplemental Fig. S3, A–D), similar to inactivation of *KNAT6* and consistent with coexpression of *BOP1/2* and *KNAT6* in valve margins (Fig. 1F; Supplemental Fig. S4; Ragni et al., 2008). We further examined the pattern of lignin deposition in fruit cross sections (Supplemental Fig. S3, E–H). In *pny* mutants, lignin (pink color) was detected throughout the junction between the valves, reflecting lack of the replum. In *bop1 bop2* and *bop1 bop2 pny* triple mutants, lignin formed only at the valve margins as in wild type. Collectively, these data demonstrate complete rescue of *pny* defects, supporting the model that *BOP1/2* antagonize *BP* and *PNY* activities in the inflorescence. These data further suggest that *BOP1/2* and *KNAT6* control similar developmental processes, based on their similar interactions with *BP* and *PNY* and their overlapping expression patterns in lateral organ boundaries (Ragni et al., 2008; see also Fig. 1; Supplemental Fig. S4).

BOP1/2 Expression Domains Are Expanded in *bp* and *pny* Mutants

Ragni et al. (2008) showed that *BP* and *PNY* prevent *KNAT2* and *KNAT6* expression in stems and pedicels and that loss-of-function *knat6* (and *knat2 knat6*) rescues *bp* and *pny* defects. This prompted us to examine if *BOP1/2* expression domains are likewise expanded in *bp* and *pny* mutants, using the *BOP2:GUS* reporter gene (Fig. 6, A–O). In *bp* mutants, *BOP2* expression was expanded in stems and pedicels, particularly below nodes. Expression on the abaxial side of nodes is consistent with localized growth restriction, causing pedicels to point downward. Staining was also seen in stripes of abnormal epidermal tissue that extend below the node and become ectopically lignified in mature *bp* stems (Fig. 6, F–I; Venglat et al., 2002; Mele et al., 2003). Stem cross sections from just below the node confirmed *BOP2* misexpression in the stem cortex beneath the epidermis and in phloem regions associated with the primary vascular bundle (Fig. 6J). In *pny* mutants, *BOP2* expression was also expanded

Table 1. Summary of inflorescence defects in plants overexpressing *BOP1* or *BOP2*

Transgene	Ecotype	Plants with	Plants with	Total No. of Transformants
		Downward-Oriented Siliques	Clustered Siliques	
		%		
<i>35S:BOP1</i>	Col	0.0	20.6	175
<i>35S:BOP2</i>	Col	0.0	10.0	80
<i>tCUP4:BOP1</i>	Col	0.0	61.1	18
<i>35S:BOP1</i>	Ler	44.5	22.0	164
<i>35S:BOP2</i>	Ler	27.6	20.4	196

in stems and pedicels above and below nodes, compatible with growth impairment, causing irregular internodes and silique clustering (Fig. 6, K–N). Stem cross sections near *pny* nodes confirmed *BOP2* misexpression throughout the stem cortex (Fig. 6O). *BOP1:GUS* expression in *bp* and *pny* mutants showed a similar pattern (Supplemental Fig. S5). In summary, the misexpression patterns of *BOP1/2* differ in *bp* and *pny* mutants, bearing resemblance to the distinct inflorescence defects that characterize these mutants.

BOP1/2 Promote *KNAT6* Expression

Given that *BOP1/2* and *KNAT6* are both required for *bp* and *pny* phenotypes and *BOP1/2* gain of function produces *bp*- and *pny*-like phenotypes, we compared *KNAT6:GUS* expression in various *BOP* gain-of-function lines: *bp*, *pny*, and *35S:BOP2* or *bop1-6D*. Misexpression of *KNAT6:GUS* in stems was confirmed for all genotypes (Fig. 7, A–D). However, the reporter gene was not expressed in boundaries of the IM, indicating that some of its control sequences were missing (data not shown). We therefore used *in situ* hybridization to further examine *KNAT6* expression in the inflorescence apex and stem (Fig. 7, E–T). In the *bp* mutant, *KNAT6* transcript was misexpressed in the stem cortex and vascular tissue (Fig. 7, J and N) and beneath the node in a stripe pattern (Fig. 7R) similar to misexpression of *BOP2* (Fig. 6, I and J). In the *pny* mutant, *KNAT6* was misexpressed in the vascular tissue of elongated stems similar to *bop1-6D* mutants (Fig. 7, K, L, O, P, S, and T). Both mutants formed extra vascular bundles, resulting in a dense vascular ring with little interfascicular space (Fig. 7, O and P; Smith and Hake, 2003). *KNAT6* transcript levels were also monitored in internodes and pedicels by quantitative reverse transcription (qRT)-PCR. These data confirmed 2- to 3-fold higher levels of *KNAT6* transcript in *bp-2*, *pny*, and *bop1-6D* plants relative to wild type and *bop1 bop2* controls (Fig. 7U). Higher levels of *KNAT6* transcript are consistent with an expanded domain of *KNAT6* expression in *bop1-6D/35S:BOP2* stems. We therefore concluded that *BOP1/2* promote *KNAT6* expression. Consistent with this, *KNAT6* transcript levels were slightly lower in *bop1 bop2 bp* and *bop1 bop2 pny* internodes and pedicels relative to *bp-2* and *pny* single mutants (Fig. 7U). No similar up-regulation was observed for *KNAT2* in *bop1-6D* plants (data not shown).

BOP1/2 Exert Their Function through *KNAT6*

Given that *BOP1/2* promote *KNAT6* expression, we reasoned that *BOP1/2* may exert all or part of their function through *KNAT6*. To examine this, we tested the effect of *kнат6* loss of function on the phenotype of a strong *35S:BOP2* gain-of-function line with short compact inflorescences (Norberg et al., 2005). In this experiment, plants that were homozygous for the *35S:BOP2* transgene were crossed to wild type or to lines homozygous for *kнат2*, *kнат6*, or *kнат2 kнат6* mutations. The phenotypes of progeny were scored in the F1 generation. To rule out transgene silencing, we took the additional step of confirming *BOP2* overexpression in F1 populations (Supplemental Fig. S6). These ex-

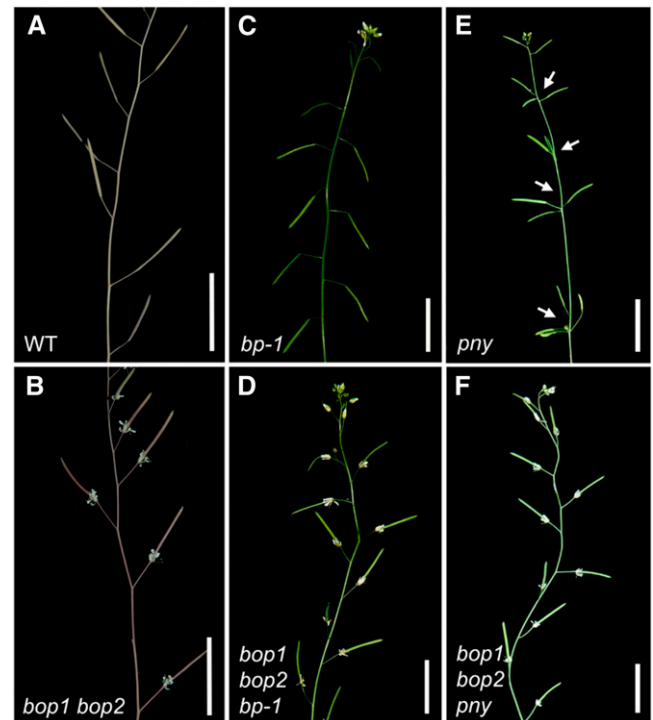


Figure 3. Phenotypic suppression of *bp* and *pny* inflorescence defects by *bop1 bop2*. A, Wild-type control. B, *bop1 bop2* mutant. C, *bp-1* mutant; downward-pointing siliques. D, *bop1 bop2 bp-1* mutant; partial rescue of *bp-1* phenotype. E, *pny* mutant; clustered siliques (arrows). F, *bop1 bop2 pny* mutant; similar to wild type. Scale bars, 2 cm. [See online article for color version of this figure.]

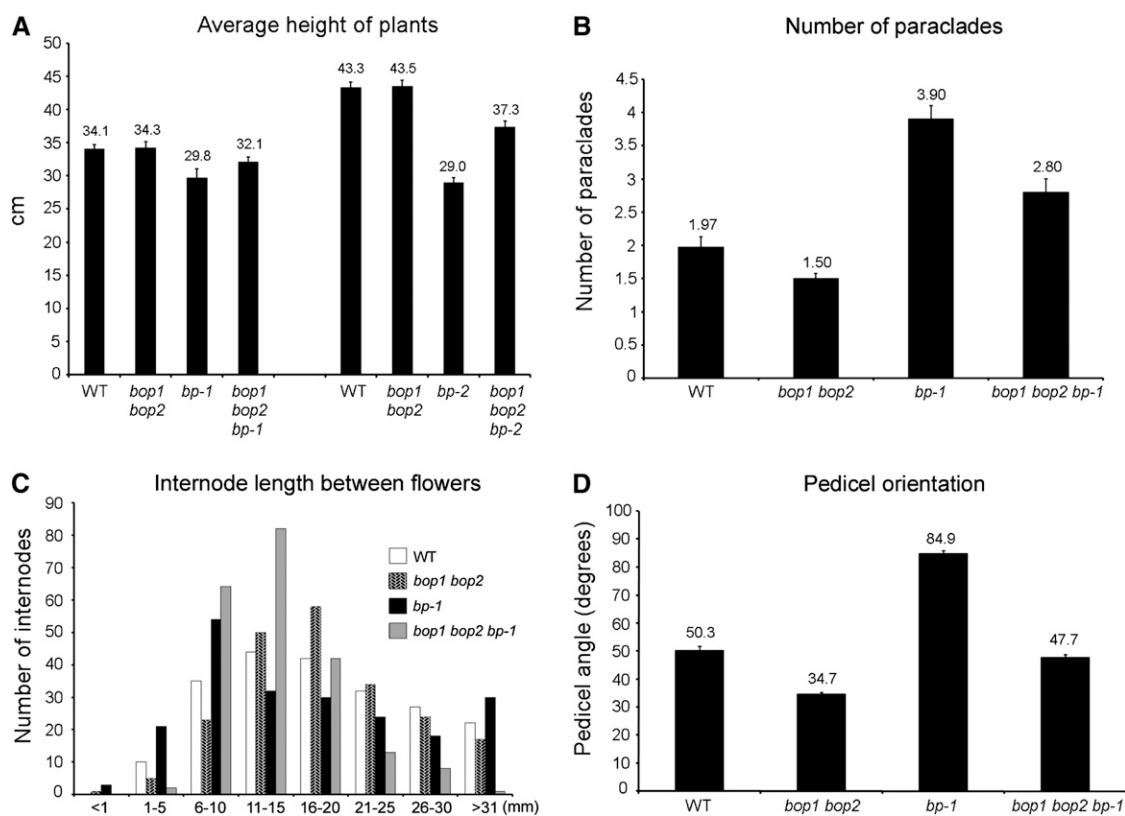


Figure 4. Quantitative analysis of *bp* phenotypic rescue by *bop1 bop2*. At least 24 plants for the indicated genotypes were analyzed. A, Average inflorescence height; inactivation of *BOP1/2* partially rescues the short stature of *bp-1* and *bp-2* mutants. B, Average number of paraclades; inactivation of *BOP1/2* partially restores apical dominance in *bp-1* mutants. C, Distribution of internode lengths between successive siliques on the primary inflorescence. Internodes between the first and 11th siliques (counting acropetally) were measured. Distribution of internode lengths in *bop1 bop2 bp-1* triple mutants is similar to wild type. D, Average orientation of pedicels; inactivation of *BOP1/2* corrects pedicel orientation in *bp-1* mutants. Error bars, se.

periments revealed that partial *knat6* loss of function (i.e. *knat6/+* or *knat2/+ knat6/+*) was sufficient to restore internode elongation in *35S:BOP2* plants (Fig. 8, A and C–E). In contrast, no rescue occurred in control crosses to wild type or *knat2* alone (Fig. 8, A, B, and E). Compatible with this, mutations in *knat2* alone do not rescue *bp* or *pnny* inflorescence defects (Ragni et al., 2008). These data indicate that BOP1/2 exert much of their function through KNAT6. Interestingly however, *35S:KNAT6* plants are not short and mimic *35S:BP* plants with lobed leaves (Supplemental Fig. S7A; see also Lincoln et al., 1994; Dean et al., 2004), indicating that the functions of BP and KNAT6 are redundant when BOP1/2 is not comexpressed. Thus, both BOP1/2 and KNAT6 are required to exert changes in inflorescence architecture.

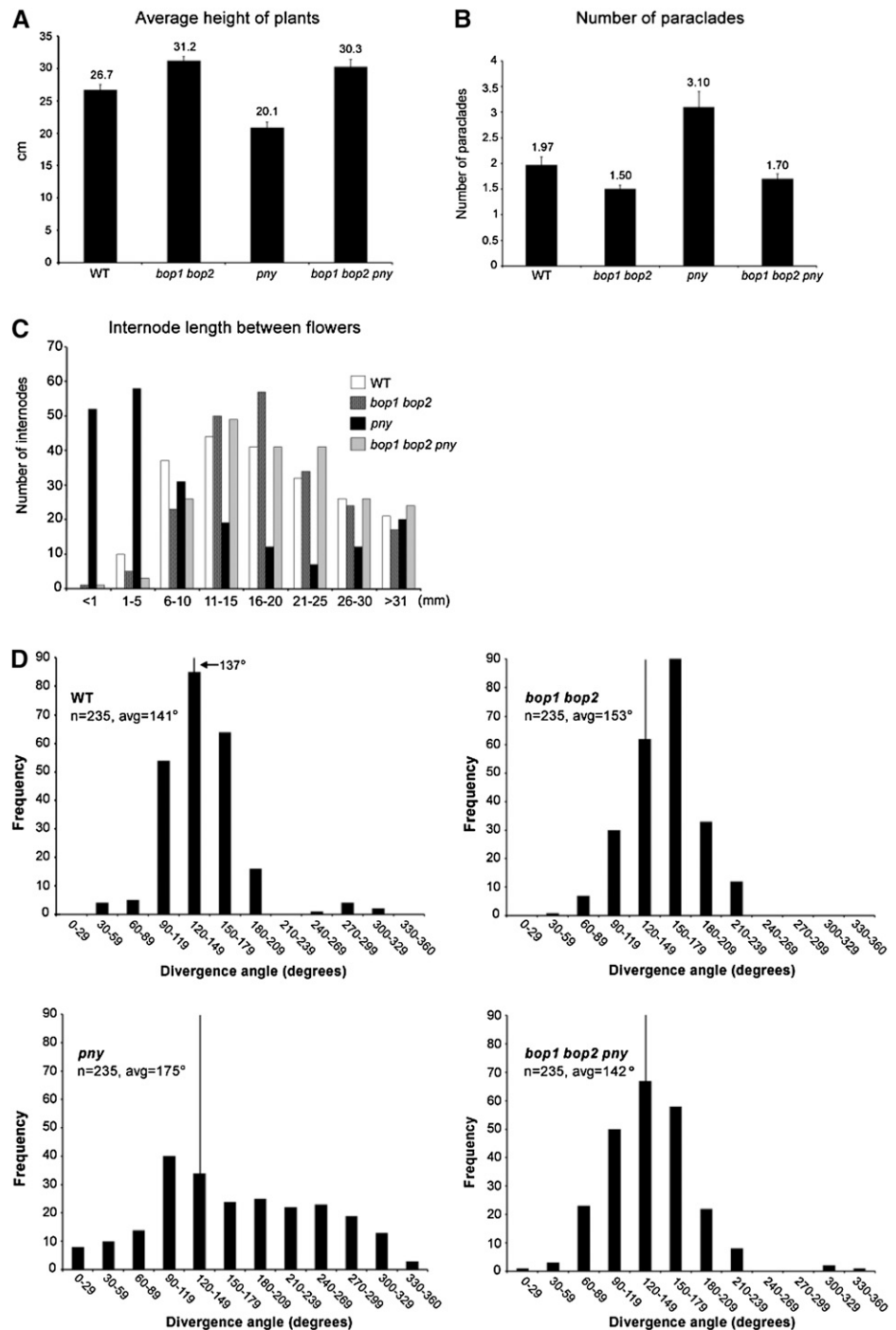
BOP1/2 and BP/PNY Are Antagonistic Regulators of Stem Lignification

We next sought to determine how BOP1/2 gain of function antagonizes BP and PNY activities in the stem. We initially considered that BOP1/2 might function through ASYMMETRIC LEAVES2 (AS2) to inhibit BP

and/or PNY expression in stems. BOP1/2 indirectly repress BP in leaves by promoting AS2 expression, whose product is a direct repressor of BP transcription (Guo et al., 2008; Jun et al., 2010). However, inactivation of AS2 failed to rescue the short stature of *35S:BOP2* plants (Ha et al., 2007) or *bp* and *pnny* inflorescence defects (Supplemental Fig. S7, B and C). Moreover, no decrease in BP or PNY expression was detected in the stem of BOP1/2 loss- or gain-of-function mutants (Supplemental Fig. S8). These data indicate that BOP1/2 control of stem architecture is largely independent of AS2 and transcriptional repression of BP. We therefore examined the model that BOP1/2 function downstream of BP-PNY and have reciprocal functions in the stem based on their compartmentalized expression domains in the inflorescence.

To examine this, we turned to work showing that BP is a negative regulator of lignin deposition in stems (Mele et al., 2003). In the primary inflorescence stem, formation of secondary cell walls is tightly regulated over developmental time (Ehltling et al., 2005). Cross sections were cut from the base of wild-type and mutant primary stems at the same developmental age and stained with phloroglucinol, which detects lignin

Figure 5. Quantitative analysis of *pnv* phenotypic rescue by *bop1 bop2*. At least 24 plants per genotype were analyzed. **A**, Average height of primary inflorescence; inactivation of *BOP1/2* rescues short stature of *pnv* mutants. **B**, Average number of rosette paraclades; inactivation of *BOP1/2* restores apical dominance in *pnv* mutants. **C**, Distribution of internode lengths between successive siliques on the primary inflorescence. Internodes between the first and 11th siliques (counting acropetally) were measured. The distribution of siliques in *bop1 bop2 pnv* mutants was similar to wild type. **D**, Distribution of divergence angles between siliques on the primary inflorescence. At least 10 successive angles between the first and 24th siliques (counting acropetally) were measured for $n \geq 14$ plants per genotype. The class containing the theoretical angle of 137° is indicated by a vertical line. Average angle, Avg. In *pnv* plants, distribution is uniform across all classes but in *bop1 bop2 pnv* plants, the distribution is similar to wild type.



deposition, a hallmark of secondary walls in vessel and fiber cells in the inflorescence stem. As expected, complex patterning changes were seen in *bp* mutants. Phloem fibers overlying primary vascular bundles were prematurely lignified. In addition, gaps were observed in the ring of interfascicular fiber cells with lignin abnormally deposited in the epidermis and cortex of these gaps. This pattern correlates with the position of abnormally differentiated stripes of tissue in *bp* stems that originate below nodes and extend

basipetally (Fig. 9, A–C; Douglas et al., 2002; Venglat et al., 2002; Mele et al., 2003). Loss-of-function *bop1 bop2* partially rescued *bp* defects, resulting in a pattern similar to wild type (Fig. 9, A, C, and D). Ectopic stem lignification also occurs in *pnv* stems, albeit in a different pattern than for *bp*, which may reflect differences in where or when *BOP1/2* are misexpressed. In *pnv* stems, vascular bundles were more crowded than in wild type, resulting in a dense vascular ring (Fig. 7O; Supplemental Fig. S9; Smith and Hake, 2003).

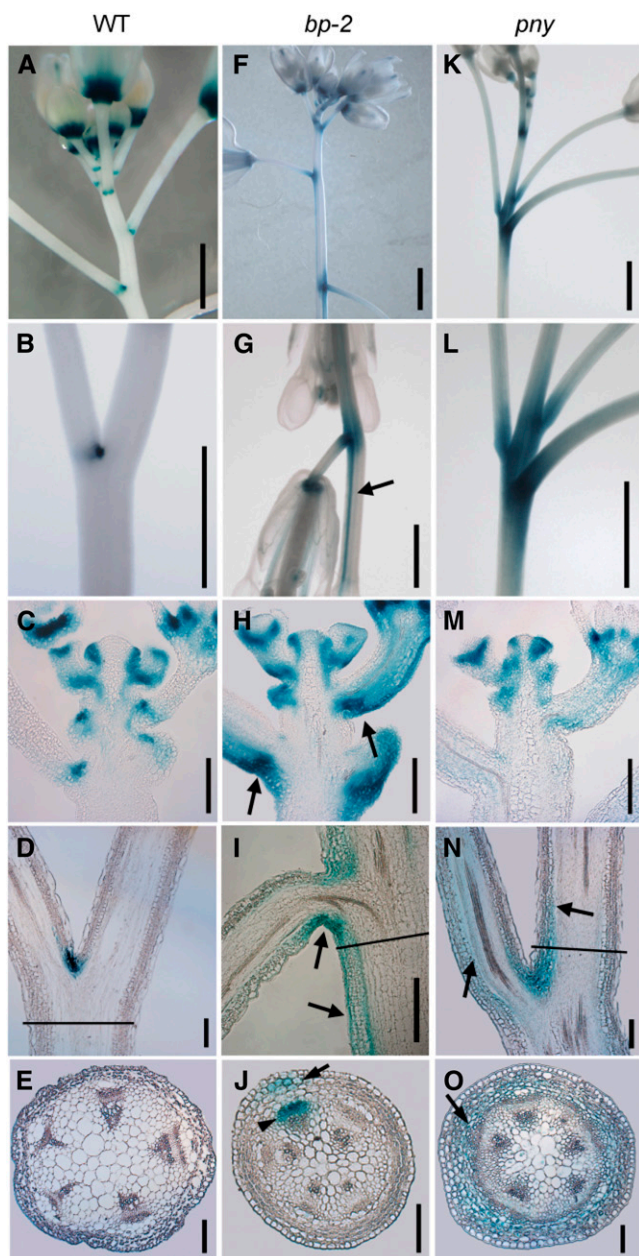


Figure 6. *BOP2:GUS* expression in wild-type, *bp*, and *pny* inflorescences. A to E, Wild type. A and B, Expression restricted to stem-pedical axil. C, Apex; no expression in the IM, internodes, or pedicels. D, Node. E, Stem; line in D shows plane of cross section. F to J, *bp-2* mutant. F to G, Expression expands beyond nodes, thin stripes of tissue extending basipetally below nodes stain strongly (arrow). H, Apex; misexpression on the abaxial side of nodes (arrows) and in pedicels. I, Node; misexpression on the abaxial side of the node (arrows). J, Stem; stripe of expression below node. Line in I shows plane of cross section. Arrow, cortical cells; arrowhead, phloem. K to O, *pny* mutant. K and L, Expression expands above and below nodes. M, Apex; staining strongest near the apex and in pedicels. N, Node; diffuse expression above and below the node (arrows). O, Stem; misexpression in stem cortex (arrow). Line in N shows plane of cross section. Scale bars, 1 mm except 100 μ m for C to E, H to J, and M to O. [See online article for color version of this figure.]

Loss-of-function *bop1 bop2* also rescued *pny* defects, resulting in a pattern similar to wild type (Supplemental Fig. S9, A, E, and G). Importantly, stem cross sections from *35S:BOP2* and *bop1-6D* plants showed expanded patterns of lignification, similar to *bp pny* double mutants (Fig. 9E; Supplemental Fig. S9; Smith and Hake, 2003). In *BOP1/2* overexpressing lines, the vascular ring was dense (similar to *pny* mutants) and phloem fiber cells overlying primary vascular bundles were prematurely lignified (similar to *bp* mutants). However, there were no gaps in the vascular ring, presumably due to uniformity in *BOP1/2* misexpression. In *bop1-6D* mutants, parts of the pith were lignified, never observed in wild-type stem development. Thus, *BOP1/2* gain of function induces lignified phloem and interfascicular fibers in a pattern reminiscent of the secondary growth that occurs in trees (Fig. 9E; Supplemental Fig. S9; Nieminen et al., 2004; Baucher et al., 2007). These data support the model *BOP1/2* function downstream of BP-PNY in the stem and have a reciprocal function associated with lignin biosynthesis.

BOP1/2 Activate Genes Repressed by BP

Microarray and electrophoretic mobility shift assay experiments have previously identified lignin biosynthetic genes that are directly repressed by *BP* in stems (Mele et al., 2003). Direct targets of *PNY* have not been identified to our knowledge. Therefore, qRT-PCR was used to examine whether lignin biosynthetic gene transcripts are reciprocally regulated by *BP* and *BOP1/2* in inflorescence stems (Fig. 9G). This approach confirmed up-regulation of all four genes previously identified by Mele et al. (2003) as up-regulated in mature *bp-2* stems (*Phe ammonia lyase1* [*PAL1*]; *cinnamate 4-hydroxylase1* [*C4H1*]; *4-coumarate CoA ligase1* [*4CL1*]; and *PRXR9GE*, a class III peroxidase) as well as several additional genes (*C3H1*; *caffeoyl CoA 3-O-methyltransferase1* [*CCoMT1*]; *cinnamyl alcohol dehydrogenase5* [*CAD5*]) in the lignin biosynthetic pathway (for review, see Boerjan et al., 2003). Mutation of *bop1 bop2* in *bp-2* restored all but one of these gene transcripts to near wild-type levels, supporting the observed promotive effect of *BOP1/2* on lignin deposition in stems. Four of the above genes were also up-regulated in *bop1-6D* stems (*C4H1*, *C3H1*, *CAD5*, and *PRXR9GE*), suggesting that *BOP1/2* directly or indirectly promotes the expression of genes in the lignin biosynthetic pathway. Similar results were obtained using internode tissue (data not shown). As reported by Mele et al. (2003), the class III peroxidase gene *PRXR9GE* showed the greatest fold-change (15- to 20-fold) over wild type in both *bp-2* and *bop1-6D* stems, suggesting that polymerization of monolignol subunits may be a key regulatory point in the developmental control of secondary wall formation. Collectively, these data support the model that *BOP1/2* and *BP* have reciprocal functions in the stem and show how antagonistic interactions between *BOP1/2* and

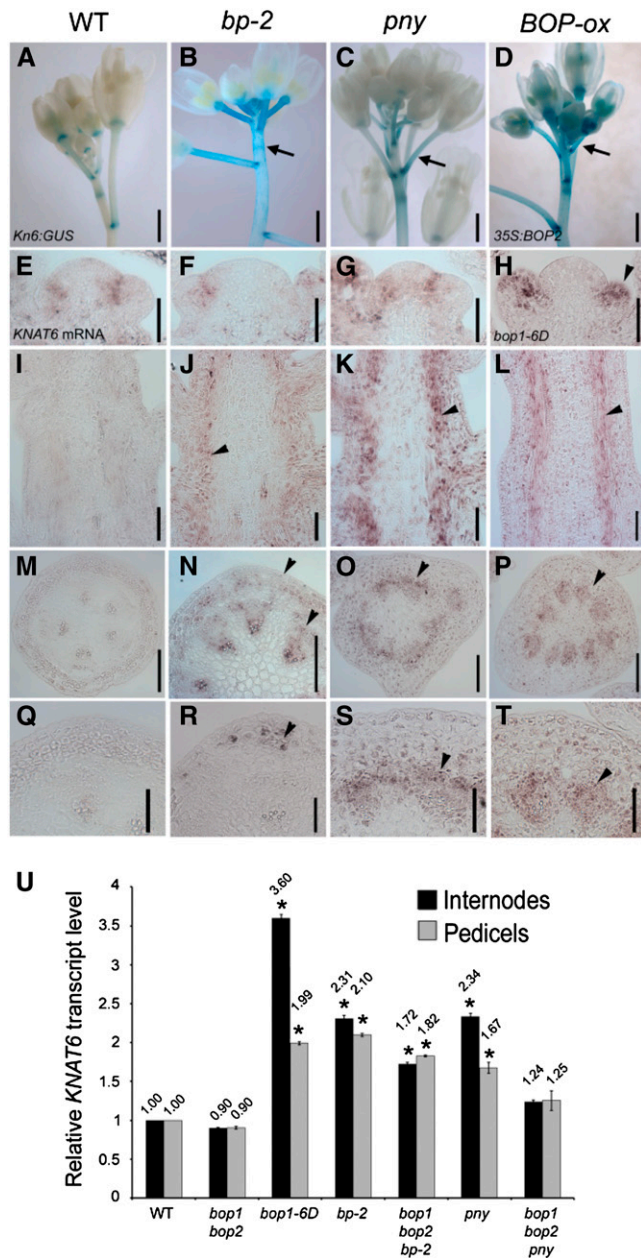


Figure 7. *KNAT6* expression in wild type, *bp-2*, *pny*, and *BOP* gain-of-function mutants. A to D, *KNAT6:GUS* expression. Inflorescences shown for: A, wild type; B, *bp-2*; C, *pny*; D, *35S:BOP2*. Expression localized to the pedicel axil in wild type (A) but misexpressed in stems and pedicels of mutants (B–D). E to T, *KNAT6* mRNA detected using in situ hybridization. Inflorescence apices shown for: E, Wild type; F, *bp-2*; G, *pny*; H, *bop1-6D*. Transcript is correctly localized to the IM-floral meristem boundary except in *bop1-6D* (H) where expression is throughout the adaxial area of floral meristems. Stem longitudinal sections shown for: I, wild type; J, *bp-2*; K, *pny*; L, *bop1-6D*. Transcript up-regulated in the cortex of mutant stems (J–L). In K and L, the vascular cambium area shows strong expression. Stem cross sections shown for: M, wild type; N, *bp-2*. Expression strongest in the cortex and vascular bundles (arrowheads). O, *pny*. Irregular vascular bundles; vascular cambium area shows the strongest expression (arrowhead). P, *bop1-6D*; strong expression in vascular bundles (arrowhead). Magnified stem cross sections shown for: Q, wild type. R, *bp-2*; stripe of expression in

BP-PNY are important for patterning of cell-type differentiation in stems as well as inflorescence architecture.

DISCUSSION

Internodes are elongated at the transition to flowering as a result of expanded rib meristem activity in the IM (Steeves and Sussex, 1989; Fletcher, 2002). The meristem expression of *BP* diminishes with the floral transition and becomes restricted to the cortex of the inflorescence stem and pedicel, where its activity together with *PNY* promotes internode elongation and vascular patterning (Lincoln et al., 1994; Douglas et al., 2002; Venglat et al., 2002; Smith and Hake, 2003). In this article, we used a genetics approach to examine how interactions between BP-PNY and the lateral-organ boundary regulators *BOP1/2* govern Arabidopsis inflorescence architecture. Our data show that a spectrum of *bp-* and *pny*-like defects in inflorescence architecture are caused by *BOP1/2* gain of function. Our key findings are that BP and PNY restrict *BOP1/2* expression to the pedicel axil together with *KNAT6* to prevent their misexpression in stems and pedicels, which causes altered growth patterns in *bp* and *pny* inflorescences. Our data also indicate that *BOP1/2* promote *KNAT6* expression and that both activities are required to inhibit internode elongation in stems (Fig. 10). Our analysis of gain-of-function mutants demonstrates that *BOP1/2* function downstream of BP-PNY in a reciprocal manner, accelerating the final steps of stem differentiation in opposition to BP.

BOP1/2 Differentially Regulate *KNOX* Activity in Leaves and the Inflorescences

Previous work has established that *BOP1/2* in leaves function together with *AS1/2* to maintain repression of the class I *KNOX* genes *BP*, *KNAT2*, and *KNAT6* during leaf development (Ori et al., 2000; Semiarti et al., 2001; Ha et al., 2003, 2007; Jun et al., 2010). In this context, *BOP1/2* indirectly represses *BP* transcription by promoting *AS2* expression in leaf petioles (Jun et al., 2010). Genetic assays show that *BOP1/2* also repress *BP* through an *AS2*-independent pathway that is as-yet undefined (Ha et al., 2007; data not shown). Our data reveals an opposite regulatory pattern in inflorescences with BP and PNY functioning as transcriptional repressors of *BOP1/2* and *KNAT6*. Comisexpression of *BOP1/2* and *KNAT6* permits their opposing function downstream of BP-PNY to restrict

cortex below node (arrowhead). S, *pny*; expression strongest in stripe of cells near vascular cambium (arrowhead). T, *bop1-6D*; expression in vascular bundles. U, qRT-PCR analysis of relative *KNAT6* transcript levels in wild-type and mutant internodes and pedicels. Asterisks, Significantly different from wild type (Student's *t* tests, $P < 0.0001$; except *pny*, $P < 0.001$). Scale bars, 50 μm except 0.5 mm for A to D and 100 μm for M to P. [See online article for color version of this figure.]

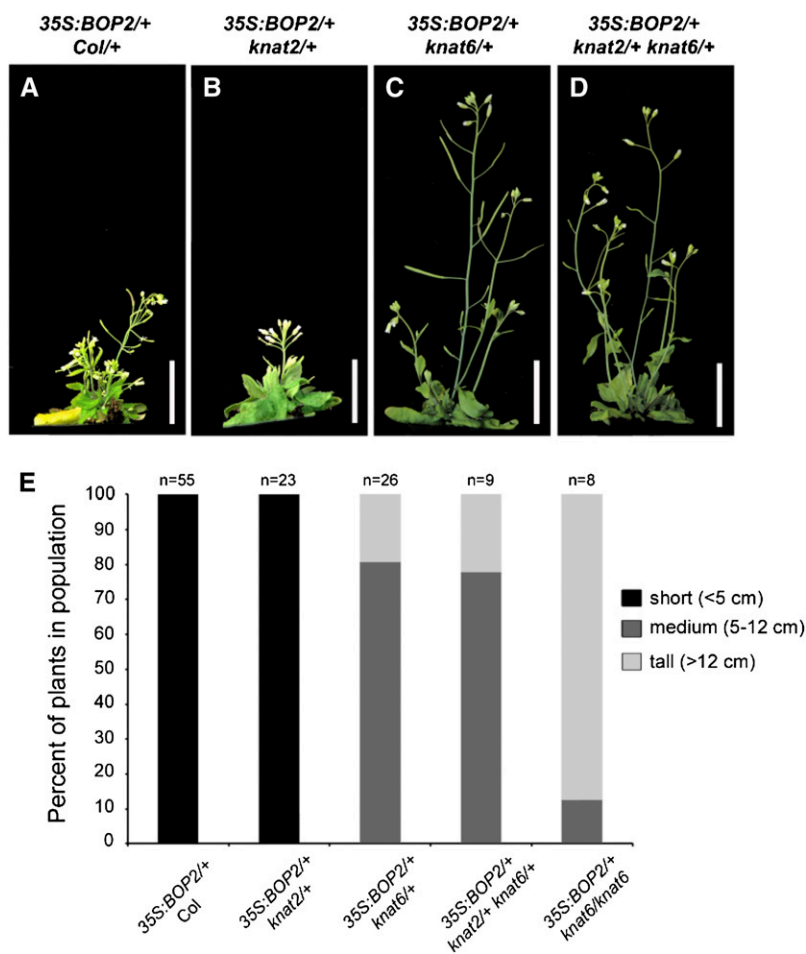


Figure 8. Inactivation of *KNAT6* rescues compact internodes caused by *BOP2* gain of function. Plants homozygous for a *35S:BOP2* transgene were crossed to wild-type control plants or to plants homozygous for mutations in *knat2*, *knat6*, or *knat2 knat6*. The inflorescences of representative F1 plants are shown. A, *35S:BOP2/+ Col*. B, *35S:BOP2/+ knat2/+*. C, *35S:BOP2/+ knat6/+*. D, *35S:BOP2/+ knat2/+ knat6/+*. E, Quantitative analysis of inflorescence height in populations of F1 plants for the genotypes indicated. Scale bars, 2 cm. [See online article for color version of this figure.]

growth and promote premature differentiation of the stem. These data are compatible with *BOP1/2* gain-of-function studies in moss. In this species, stabilization of *BOP1/2* transcripts causes premature gametophore differentiation (Saleh et al., 2011).

Misexpression of *BOP1/2* Restricts Growth to Create Variations in Inflorescence Architecture

Variations in inflorescence architecture are extensive among flowering plants, with the length and pattern of internode elongation and pedicel angle acting as key variables in the display of flowers (Steeves and Sussex, 1989; Sussex and Kerk, 2001). Short internodes and pedicels like those in *bp* mutants are reminiscent of species with spike-type inflorescences where internodes between successive flowers are short (Bell and Bryan, 2008). Conversely, long internodes separating whorls of flowers on the stem, like those in *pny* mutants, are reminiscent of species with verticillate-type inflorescences (Bell and Bryan, 2008). Our data show that a spectrum of inflorescence architectures ranging from short internodes, to downward-pointing pedicels, to clusters of flowers may be produced by differentially regulating the pattern and degree of *BOP*

gain of function in stems. In *bp* mutants, ectopic *BOP1/2* expression on the abaxial side of nodes leads to growth restriction and downward-pointing pedicels. *BOP1/2* are also misexpressed in the stem cortical tissue where BP-PNY normally function, thereby inhibiting internode elongation and causing cells to differentiate prematurely. In *pny* mutants, *BOP1/2* are strongly misexpressed in the pedicels and stem cortex of young internodes, leading to irregular elongation of internodes and the clustering of flowers in whorls. These defects are phenocopied to various degrees by ectopically expressing *BOP1/2* under the control of single or multiple 35S Cauliflower mosaic virus (CaMV) enhancers, indicating that *BOP1/2* function downstream of BP-PNY in an antagonistic manner. However, *BOP1/2* gain of function does not reduce *BP* or *PNY* transcript levels in the stem (Supplemental Fig. S8), indicating that *BOP1/2* likely oppose BP-PNY function posttranscriptionally.

BOP1/2 and *KNAT6* Function in the Same Genetic Pathway

Our genetic assays and expression data show that misexpression of *BOP1/2* is the cause of inflorescence

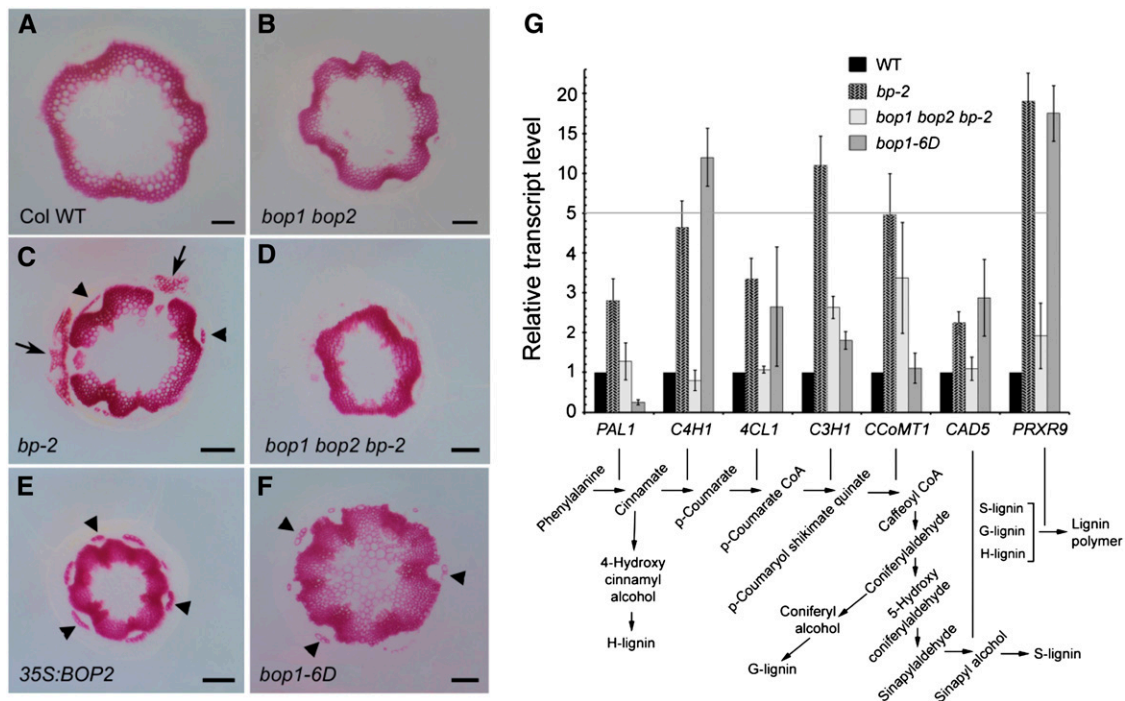


Figure 9. Lignification pattern and lignin biosynthetic gene expression in wild-type and mutant stems. A to F, Cross sections from the base of fully elongated stems were stained with phloroglucinol-HCl to reveal lignin. Representative sections are shown for: A, wild type; B, *bop1 bop2*. C, *bp-2*; gaps in the vascular ring (arrows) are associated with stripes of ectopically lignified epidermal/cortical tissue. Arrowheads, Premature lignification of phloem fiber cells in primary vascular bundles. D, *bop1 bop2 bp-2*; similar to wild type. E, *35S:BOP2*; dense vascular ring compared to wild type. Arrowheads, Premature lignification of phloem fiber cells, similar to *bp-2* mutants. F, *bop1-6D*; similar to *35S:BOP2* but pith is also lignified. Scale bars, 100 μ m. G, qRT-PCR analysis of lignin biosynthesis genes in stem tissue (same stage as above). Error bars, SE of three biological replicates. Position of genes in the lignin biosynthetic pathway is depicted below (adapted from Mele et al., 2003; Zhou et al., 2009). [See online article for color version of this figure.]

patterning defects in *bp* and *pn1* mutants. For reasons that are unclear, inactivation of *BOP1/2* partially suppresses *bp* defects but completely suppresses *pn1* defects. This difference may be related to the slightly different roles that *bp* and *pn1* play in internode development (Hake and Smith, 2003; Peaucelle et al., 2011). These data extend the work of Ragni et al. (2008) who showed an identical pattern of rescue for *bp* and *pn1* defects by inactivation of *KNAT6* (and to a lesser extent *KNAT2*), genes that are misexpressed in an overlapping domain with *BOP1/2* in *bp* and *pn1* stems (Figs. 6 and 7). These studies place *BOP1/2* and *KNAT6* in the same genetic pathway controlling inflorescence architecture. Compatible with this, *BOP1/2* gain of function promotes *KNAT6* expression. However, both activities are required to restrict internode elongation since inactivation of *KNAT6* restores internode elongation in *35S:BOP2* plants and *35S:KNAT6* internodes are not short (Fig. 8; Supplemental Fig. S7; Dean et al., 2004). Despite several attempts with *35S:BOP1-GR* transgenic plants and chromatin immunoprecipitation assays, we have yet to determine if *BOP1/2* directly regulate *KNAT6*. Given that BP and *KNAT6* are highly related proteins with the same gain-of-function phenotype (Lincoln et al., 1994; Chuck et al., 1996; Dean et al., 2004) they

may regulate some of the same genes. However, *KNAT6* with *BOP1/2* function in opposition to BP. A physical complex between *BOP1/2* and *KNAT6* was not detected in yeast (*Saccharomyces cerevisiae*; data not shown). We therefore favor a model in which *BOP1/2* bind independently to the same promoters as *KNAT6* or induce the expression of a *KNAT6* cofactor to exert their effect.

In fruits, *BOP1/2* and *KNAT6* likewise function in the same genetic pathway as evidenced by rescue of replum formation in *pn1* mutants by either *bop1 bop2* or *knat6* loss of function (Ragni et al., 2008; this study). *BOP1/2* and *KNAT6* may also share a role in floral-organ abscission based on recent evidence that IDA-dependent signaling inhibits BP activity, allowing *KNAT2* and *KNAT6* to promote abscission (McKim et al., 2008; Shi et al., 2011). Thus, antagonism between BP-PNY and a genetic pathway involving *BOP1/2* and *KNAT6* is likely to be a conserved module in plant development.

ARABIDOPSIS THALIANA HOMEBOX GENE1 Is a Potential *KNAT6* Cofactor

The BELL homeodomain protein encoded by *ARABIDOPSIS THALIANA HOMEBOX GENE1* (*ATH1*) is another potential member of the *BOP1/2* and *KNAT6*

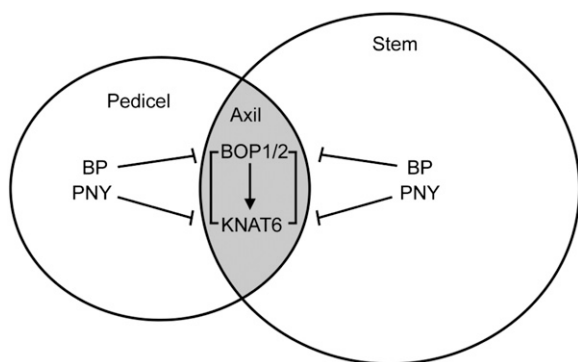


Figure 10. Summary of genetic interactions between BP-PNY, BOP1/2, and KNAT6 in the inflorescence. BP and PNY in the stem and pedicels are transcriptional repressors of *BOP1/2* and *KNAT6*, limiting their expression to the pedicel axil. BOP1/2 gain-of-function mutants phenocopy *bp* and *pny* mutants because BOP1/2 function downstream of BP-PNY in an antagonistic manner. BOP1/2 are positive regulators of *KNAT6* expression that depend in part on KNAT6 activity to exert changes in inflorescence architecture.

genetic pathway that will be important to investigate. KNOX homeodomain proteins like *KNAT6* perform many of their functions as heterodimers with BELL proteins (e.g. Byrne et al., 2003; Bhatt et al., 2004; Kanrar et al., 2006; Rutjens et al., 2009). These partnerships can influence protein-protein interactions, nuclear localization of the KNOX partner, and/or binding-site selection (Smith et al., 2002; Hackbusch et al., 2005; Cole et al., 2006; Rutjens et al., 2009). Interestingly, loss-of-function *ath1-1* rescues *pny* inflorescence defects (like *bop1 bop2* and *knat6*) whereas *ATH1* gain of function causes short internodes (Gómez-Mena and Sablowski, 2008; Rutjens et al., 2009). Given that *ATH1* transcripts are highly up-regulated in *bop1-6D* internodes (data not shown), an *ATH1*-*KNAT6* complex may restrict stem growth. Short internodes are typical of defects in gibberellic acid (GA) biosynthesis (Achard and Genschik, 2009; Schwechheimer and Willige, 2009) of which *BP* is a repressor (Hay et al., 2002). However, GA 20-oxidase transcript levels in *35S:BOP2* and *bop1-6D* stems are not significantly different from wild type (data not shown) and spray treatment of *35S:BOP2* plants with GA did not restore internode elongation (data not shown), making it uncertain if defects in GA biosynthesis or catabolism are at play.

BP and BOP1/2 Antagonistically Regulate Secondary Cell Wall Biosynthesis

Lignin biosynthesis is one of the major components of secondary cell wall formation, essential for water transport and the structural support of plants. In *Arabidopsis*, abundant interfascicular fibers with secondarily thickened cell walls develop in the primary stem as the inflorescence matures (Nieminen et al., 2004; Ehlting et al., 2005). In *bp* mutants, lignin deposition in interfascicular fibers and phloem occurs prematurely, showing that part of the function of *BP* is to

delay terminal cell differentiation, potentially until internode elongation is complete (Mele et al., 2003). However, these defects are alleviated by *bop1 bop2* mutation, showing that BOP1/2 promotes terminal cell fate and is a developmental regulator of lignin formation. Although *BOP1/2* are not normally expressed in *Arabidopsis* stems, boundaries such as the valve margin of fruits and the base of floral organs or leaves following abscission are lignified in aid of cell separation and scar fortification, respectively (Sexton, 1976; Lewis et al., 2006; Lee et al., 2008). Interestingly, publically available poplar (*Populus* spp.) microarray data indicates that two potential *BOP* orthologs are highly expressed in xylem (<http://www.bar.utoronto.ca>), which suggests a conserved role for BOP1/2 in promoting secondary growth in trees.

Mele et al. (2003) identified four lignin biosynthetic genes whose expression was up-regulated in *bp-9* stems. Our study confirmed up-regulation of these genes (*PAL1*, *C4H1*, *4CL1*, *PRXR9GE*) as well as several others (*C3H1*, *CAD5*, *HCT*) in *bp-2* and *bop1-6D* stems and internodes. Given that *BP* binds directly to the promoters of lignin genes (Mele et al., 2003) it will be interesting to confirm biochemically whether BOP1/2 and *BP* directly regulate a common set of genes to exert their antagonistic functions. Of the lignin genes surveyed, the class III cell wall peroxidase transcript *PRXR9GE* shows the most dramatic up-regulation in *bp-2* and *bop1-6D* lines (15-fold or more) relative to wild-type control plants. Class III peroxidases are one of several classes of cell wall enzymes that use hydrogen peroxide as an oxidant to generate monolignol phenoxy radicals, thus allowing the spontaneous coupling of monolignols into their polymer form (Boerjan et al., 2003; Passardi et al., 2004). Peroxidase activity is low in seedlings and increases with age in the aerial parts of the plant (Mele et al., 2003; Cosio and Dunand, 2010). Thus, the final step of lignin formation may be a key point of developmental control, making the transcriptional regulation of *PRXR9GE* an interesting case study.

In conclusion, our data establish BOP1/2 gain of function as the basis of *bp* and *pny* inflorescence defects. Our study shows that ectopic BOP1/2 activity in stems both restricts growth and promotes terminal cell differentiation, dramatically altering inflorescence architecture. Future studies will establish the molecular basis of antagonism between BP-PNY and BOP1/2. Ultimately, this work will provide important insight into how changes in the interplay between KNOX-BELL factors in the meristem and BOP1/2 in lateral organ boundaries drives species variation in inflorescence architecture.

MATERIALS AND METHODS

Plant Material and Growth Conditions

Plants were grown in growth chambers on agar plates and/or in soil at 21°C in 24-h light (100 $\mu\text{mol m}^{-2}\text{s}^{-1}$). Wild type was the Col-0 ecotype of *Arabidopsis* (*Arabidopsis thaliana*) unless stated otherwise. The double mutant

bop1-3 bop2-1 was described previously (Hepworth et al., 2005). Mutant alleles of *as2-1* (CS3117), *pnny-40126* (SALK_40126), *knat2-5* (SALK_099837), and *knat6-2* (SALK_054482) were obtained from the Arabidopsis Biological Resource Center and described previously (Byrne et al., 2000; Iwakawa et al., 2002; Smith and Hake, 2003; Belles-Boix et al., 2006). Mutant alleles of *bp-1* and *bp-2* (introgressed from RLD into Col-0) were provided by Raju Datla (Venglat et al., 2002). The strong *35S:BOP2* line and activation-tagged overexpression line *bop1-6D* were kindly provided by O. Nilsson (Norberg et al., 2005). The reporter lines *KNAT2:GUS* (C24 ecotype) and *KNAT6:GUS* (Wassilewskija ecotype) were gifts from Veronique Pautot (Dockx et al., 1995; Belles-Boix et al., 2006). The reporter line *BLR:GUS* (*Ler* ecotype, here called *PNY:GUS*) was provided by Mary Byrne (Byrne et al., 2003). Control crosses to Col determined that ecotype does not affect the expression pattern of GUS reporter genes. The reporter line *BOP2:GUS* is described elsewhere (Xu et al., 2010). All mutant combinations were constructed by crossing and confirmed by PCR genotyping where possible.

Primers and Genotyping

Primers used for genotyping, plasmid construction, and transcript analysis are listed in Supplemental Table S1. The strategy for genotyping *bop1-3*, *bop2-1*, *pnny-40126*, *knat2-5*, and *knat6-2* Salk T-DNA insertion mutants was as described (www.signal.salk.edu). For genotyping *bp-2*, primers bp-2dCAPS-F1 and bp-2dCAPS-R1 were used to amplify products from wild-type and *bp-2* genomic DNA. The product from Col wild type is slightly larger than the corresponding product from *bp-2*, allowing their resolution on a 3.5% agarose gel.

Construction of *35S:BOP1*, *35S:BOP2*, and *tCUP4:BOP1* Transgenic Lines

To create pBAR/35S:BOP1/2 constructs, a fragment containing one copy of the viral 35S promoter was excised from p35S:BOP2 (Norberg et al., 2005) by digestion with *EcoRI* and *BamHI* and cloned into the corresponding site of pBAR1 (a gift from the Dangl Lab, University of North Carolina) to create the intermediate plasmid pBAR/35S. Primer pairs B1-1/B1-2 and B2-1/B2-2 incorporating *BamHI* restriction sites were used to amplify *BOP1* and *BOP2* coding sequences, respectively, from cloned cDNA templates. The resulting PCR products were digested with *BamHI* and ligated into the corresponding site in pBAR to generate pBAR/35S-BOP1 and pBAR/35S-BOP2. The *EntCUP4* promoter is an alternative constitutive promoter (Malik et al., 2002). To create pCUP4:BOP1, a DNA fragment containing the *BOP1* coding sequence was amplified by PCR from cloned cDNA template using *EcoRI*-BOP1-F1 and *BOP1*-RR as the primers. The resulting fragment was digested with *EcoRI* and *BamHI* and ligated into the corresponding sites of pBAR1 to generate the intermediate plasmid pBAR1/BOP1. A 0.5-kb DNA fragment containing the *EntCUP4* promoter was then amplified by PCR using pEntCUP4-nos-GUS as a plasmid template and *EcoRI*-tCUP-F1 and *EcoRI*-tCUP-R1 as the primers. The resulting fragment was digested with *EcoRI* and ligated into the corresponding sites of pBAR/BOP1 to create pCUP4:BOP1. Wild-type plants were transformed by floral dipping (Clough and Bent, 1998) using the *Agrobacterium* strain C58C1 pGV101 pMP90 (Koncz and Schell, 1986). Basta-resistant transformants were selected on soil using the herbicide Finale (AgrEvo). Phenotypes were scored in the T1 generation.

Phenotypic Analysis of Inflorescence Structure

Quantitative phenotypic analyses of 6-week-old plants were performed as described (Ragni et al., 2008). Phyllotaxy measurements were obtained as previously described (Peaucelle et al., 2007). The divergence angle between the insertion points of two successive floral pedicels along the main inflorescence was measured. Divergence angles were measured for the first 15 siliques of each inflorescence (counting acropetally) according to the orientation that resulted in the smallest average divergence angle. Angle of pedicel orientation was determined using a protractor to measure the angle of pedicel attachment relative to the stem. Orientation was measured for the first 11 siliques of each inflorescence (counting acropetally).

In Situ Hybridization and Localization of GUS Activity

Tissues were fixed and analyzed for GUS activity essentially as described by Sieburth and Meyerowitz (1997). Tissues were stained for 2 to 18 h at 37°C

and cleared overnight with 70% ethanol prior to imaging. Alternatively, stained tissues were embedded in Paraplast Plus (Sigma). Sections (10 μm) cut with a microtome were affixed to glass slides and dewaxed with tert-butanol and xylene prior to imaging. In situ hybridizations were performed as described (Xu et al., 2010). Primers used to make *BP* and *KNAT6* antisense probes are listed in Supplemental Table S1.

SEM

Samples were prepared for SEM as described in Hepworth et al. (2005). Images were collected using a Vega-II XMU variable pressure SEM (Tescan).

Lignin Staining

Tissue sections (25 μm) were cut from paraffin-embedded mature green siliques to analyze replum patterning or from elongated internodes between the third and fourth siliques on the primary stem to analyze stem patterning. Tissue sections affixed to glass slides were dewaxed and dehydrated prior to addition of 2% phloroglucinol (in 95% ethanol) followed by 6N HCl for color development. For the analysis of lignin at stem bases, cross sections were cut from the base of 32-d-old flowering plants with a razor blade and placed in 3 mL of 2% phloroglucinol solution. After 5 min, five drops of concentrated HCl were added. Two minutes were allowed for color development and images were immediately collected.

qRT-PCR

Total RNA was isolated from leaves, pedicels, internodes, or the base of bolting stems (bottom 2.5 cm of 32-d-old flowering plants) using Trizol reagent (Invitrogen). cDNA was generated using 1 μg of total RNA as the template and Superscript III RT (Invitrogen) as the polymerase. qPCR was performed in triplicate using 2 μL of 10-fold diluted cDNA as the template in reactions containing SYBR Green and IQ Supermix (BioRad) using a Rotor-Gene 6000 thermocycler (Qiagen). Annealing conditions were optimized for each primer pair and data quality was verified by melting curve analysis. Relative transcript levels were calculated as described (Murmu et al., 2010). Values were normalized to *GAPC* and then to the wild-type control. For Figure 9G only, cDNA was generated using 2 μg of total RNA as the template and diluted 20-fold. *ACTIN2* was used as a normalization control. Reactions were performed in triplicate using an annealing temperature of 55°C. All experiments were repeated at least twice with independently isolated RNA with similar results obtained. Primers for the analysis of lignin genes are given in Supplemental Table S2.

Sequence data for genes described in this article can be found in the GenBank/EMBL data libraries under the accession numbers: At2g41370 (*BOP1*), At3g57130 (*BOP2*), At1g70510 (*KNAT2*), At1g23380 (*KNAT6*), At4g08150 (*BP*), At5g02030 (*PNY*), At3g04120 (*GAPC*), At2g37040 (*PAL1*), At2g30490 (*C4H1*), At1g51680 (*4CL1*), At2g40890 (*C3H1*), At4g34050 (*CCoMT1*), At4g34230 (*CAD5*), At3g21770 (*AtPRXR9GE*), and At3g18780 (*ACT2*).

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Figure S1. Phenotypic suppression of *bp-2* inflorescence defects by *bop1 bop2*.

Supplemental Figure S2. Comparison of stem-pedicel junctions in wild type and mutants.

Supplemental Figure S3. Loss-of-function *bop1 bop2* restores replum formation in *pnny* mutants.

Supplemental Figure S4. *KNAT6:GUS* and *KNAT2:GUS* expression patterns in wild-type plants.

Supplemental Figure S5. *BOP1:GUS* expression pattern in wild type and mutants.

Supplemental Figure S6. Quantitative analysis of *BOP2* transcript in *35S:BOP2* lines crossed to wild type and mutants.

Supplemental Figure S7. Inflorescence phenotype of *35S:KNAT6* transgenic plants and the double mutants *bp-2 as2-1* and *pnv as2-1*.

Supplemental Figure S8. Comparison of *BP* and *PNY* expression levels in wild type, *bop1 bop2*, and *bop1-6D* inflorescence stems.

Supplemental Figure S9. Analysis of stem lignification pattern in wild type and mutants with *pnv*.

Supplemental Table S1. List of general primers.

Supplemental Table S2. List of primers for qPCR analysis of lignin genes.

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