

# Floral Induction in Arabidopsis by FLOWERING LOCUS T Requires Direct Repression of *BLADE-ON-PETIOLE* Genes by the Homeodomain Protein PENNYWISE<sup>1</sup>[OPEN]

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Flowers form on the flanks of the shoot apical meristem (SAM) in response to environmental and endogenous cues. In Arabidopsis (*Arabidopsis thaliana*), the photoperiodic pathway acts through *FLOWERING LOCUS T* (*FT*) to promote floral induction in response to day length. A complex between *FT* and the basic leucine-zipper transcription factor *FD* is proposed to form in the SAM, leading to activation of *APETALA1* and *LEAFY* and thereby promoting floral meristem identity. We identified mutations that suppress *FT* function and recovered a new allele of the homeodomain transcription factor *PENNYWISE* (*PNY*). Genetic and molecular analyses showed that ectopic expression of *BLADE-ON-PETIOLE1* (*BOP1*) and *BOP2*, which encode transcriptional coactivators, in the SAM during vegetative development, confers the late flowering of *pny* mutants. In wild-type plants, *BOP1* and *BOP2* are expressed in lateral organs close to boundaries of the SAM, whereas in *pny* mutants, their expression occurs in the SAM. This ectopic expression lowers *FD* mRNA levels, reducing responsiveness to *FT* and impairing activation of *APETALA1* and *LEAFY*. We show that *PNY* binds to the promoters of *BOP1* and *BOP2*, repressing their transcription. These results demonstrate a direct role for *PNY* in defining the spatial expression patterns of boundary genes and the significance of this process for floral induction by *FT*.

Plants produce new organs from a population of pluripotent cells in meristems whose function is related to stem cells in animals. Meristems are located at different positions of the plant body and give rise to different organs. The shoot apical meristem (SAM) produces leaves and flowers at the tips of stems, whereas the axillary

meristems give rise to lateral structures (Bowman and Eshed, 2000). By reprogramming these pluripotent cells at the meristems, plants can readily modify their development in response to changes in environmental conditions.

Flowers develop from floral meristems (FMs) that are formed on the flanks (floral primordium) of the SAM in response to environmental and endogenous cues (Pidkowich et al., 1999). Major environmental signals are the seasonal fluctuations in temperature and day length that are used by plants to anticipate optimal conditions for reproduction. Changes in temperature and day length are integrated into flowering-signaling networks by the thermosensory and vernalization/autonomous pathways and the photoperiodic pathway, respectively (Martinez-Zapater and Somerville, 1990; Lee and Amasino, 1995; Valverde et al., 2004; Andrés and Coupland, 2012). On the other hand, the plant hormone GA and the age of the plant constitute the internal signals affecting flowering in many plant species (Wilson et al., 1992; Fowler et al., 1999; Yu et al., 2012). *FLOWERING LOCUS T* (*FT*) is a key component of the photoperiodic pathway. *FT* encodes a small globular protein that shares high homology with mammalian phosphatidylethanolamine-binding proteins/Raf-1 kinase inhibitory protein (Kardailsky et al., 1999; Kobayashi et al., 1999; Nakamura

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F.A. and M.R.-B. performed most of the experiments; R.M.-G. generated the pPNY::Venus:PNY construct; S.J. contributed to the ethyl methanesulfonate mutagenesis; J.A. and P.N. provided the genome resequencing data; V.P. and K.S. analyzed the Next Generation Sequencing data; F.A. and G.C. conceived the project, designed the experiments, interpreted the data, and wrote the article.

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et al., 2014; Romera-Branchat et al., 2014). In *Arabidopsis thaliana*, *FT* is induced by long days (LDs) and has been placed at the core of the photoperiodic pathway, downstream of the *GIGANTEA* and *CONSTANS* genes (Suárez-López et al., 2001; Valverde et al., 2004; Yoo et al., 2005). *FT* mRNA is expressed specifically in the companion cells of the phloem, and its protein moves systemically to the shoot apex through the phloem sieve elements (Corbesier et al., 2007; Jaeger and Wigge, 2007; Mathieu et al., 2007; Tamaki et al., 2007). According to recent studies in *Cucurbita moschata* and *Arabidopsis*, FT protein is unloaded into the surrounding shoot meristem tissue from the terminal phloem (Yoo et al., 2013). Once FT is unloaded into the shoot meristem, it is thought to physically interact with two basic leucine zipper (bZIP) transcription factors called FD and FD PARALOG (FDP), which are expressed in this tissue (Abe et al., 2005; Wigge et al., 2005). However, recent work in rice (*Oryza sativa*) suggested that this interaction is not direct and is mediated by 14-3-3 proteins (Taoka et al., 2011). Consistent with this model, the loss of function of *FD* and *FDP* strongly suppresses the early flowering of transgenic plants overexpressing *FT* (Abe et al., 2005; Wigge et al., 2005; Jaeger et al., 2013). In *Arabidopsis*, the FT-FD complex is believed to induce the transcription of genes encoding several floral-promoting proteins, such as the MADS-box transcription factors SUPPRESSOR OF OVEREXPRESSION OF CONSTANS1 (SOC1) and FRUITFULL (FUL), which accelerate flowering, as well as APETALA1 (AP1), also a MADS-box transcription factor, and LEAFY (LFY), which promote FM identity (Schmid et al., 2003; Teper-Bamnolker and Samach, 2005; Wigge et al., 2005; Corbesier et al., 2007). Indeed, the FT-FD complex directly binds to the promoter of *AP1*, whose expression at the floral primordia is associated with FM formation (Wigge et al., 2005). Therefore the FT-FD complex is predicted to be active in the incipient floral primordia to induce the expression of *AP1* and promote flowering. In addition to these genes, the FT-FD complex also promotes the transcription of the family genes encoding the SQUAMOSA BINDING PROTEIN LIKE (SPL) transcription factors. Recent studies using chromatin immunoprecipitation (ChIP) assays showed that *SPL3*, *SPL4*, and *SPL5* loci are bound by FD, which transcriptionally regulates these genes (Jung et al., 2012). In turn, SPL proteins control the expression of *FUL*, *LFY*, and *AP1* genes by directly binding to their promoters (Wang et al., 2009; Yamaguchi et al., 2009). These data reflect the high degree of complexity implicit to the genetic networks controlling floral induction in the SAM.

The three-amino acid-loop-extension (TALE) homeodomain superclass comprises transcription factors involved in the SAM function. The BEL1-like homeodomain (BELL) and the KNOTTED-like homeodomain (KNOX) are TALE proteins that share similar structure and function (Hamant and Pautot, 2010; Hay and Tsiantis, 2010; Arnaud and Pautot, 2014). Members of the two families can form heterodimers to regulate

various developmental processes. The *BELL* family comprises 13 members (Smith et al., 2004). *PENNYWISE* (*PNY*), also known as *BELLRINGER*, *REPLUMLESS* (*RPL*), *VAAMANA*, or *LARSON*, encodes a BELL protein that plays roles in organ patterning by affecting internode length, phyllotaxis, and fruit replum development (Byrne et al., 2003; Roeder et al., 2003; Smith and Hake, 2003). In fruits, *PNY* is required for the replum formation, where it acts as a transcriptional repressor of *SHATTERPROOF* (*SHP*) MADS-box genes. The repressive activity of *PNY* restricts *SHP* gene expression to the valve margin domain. In the absence of a functional *PNY* (in *rpl* mutants), *SHP* genes are ectopically expressed, and the replum cells take on valve margin fates (Roeder et al., 2003). *PNY* mutants also display dramatic defects in inflorescence development (Smith and Hake, 2003; Bao et al., 2004). Interestingly, these defects are corrected by the lack of *KNOTTED*-like from *Arabidopsis thaliana*6 (*KNAT6*) and *KNAT2*, two related *KNOX* genes. In the *pnny* mutant, *KNAT2* and *KNAT6* expression domains are enlarged, indicating that *PNY* regulates inflorescence development at least partially by limiting their spatial pattern of expression (Ragni et al., 2008). These data suggest that a major molecular function of *PNY* is to maintain the spatial expression of organ patterning genes restricted to specific domains.

Recent studies elucidated that *PNY* is also involved in the acquisition of competence to respond to floral inductive signals. In these studies, it was shown that simultaneous loss of function of *PNY* and its paralog *POUND-FOOLISH* (*PNF*) completely blocks the floral transition. Indeed, the double mutant *pnny pnf* is not able to undergo the floral transition, even under LD conditions (Smith et al., 2004). Moreover, overexpression of *FT* from the constitutive *Cauliflower mosaic virus 35S* promoter barely activates flowering of the *pnny pnf* double mutant (Kanrar et al., 2008). At the molecular level, the concurrent loss of function of *PNY* and *PNF* affects the ability of FT to activate the transcription of *AP1*, *LFY*, and probably *SPLs* (Lal et al., 2011). Therefore, *PNY* has somehow been integrated into the *FT*-signaling pathway. However, the genetic and molecular mechanism underlying the effect on *PNY* on the *FT* flowering pathway remains unknown.

Here, we show that *PNY* operates in the *FT*-signaling pathway by restricting the spatial pattern of expression of *BLADE-ON-PETIOLE1* (*BOP1*) and *BOP2* genes in the SAM. *PNY* directly binds to *BOP1* and *BOP2*, which encode two BTB (for Bro ad-complex, Tramtrack, Bric-à-brac)-ankyrin transcriptional coactivators, which function at lateral organ boundaries in the determination of leaf, flower, inflorescence, and root nodule architecture (Ha et al., 2004, 2007; Hepworth et al., 2005; Norberg et al., 2005; Karim et al., 2009; Xu et al., 2010; Couzigou et al., 2012; Khan et al., 2012b, 2014). We found that *BOP1* and *BOP2* are also involved in flowering-time regulation by repressing the expression of *FD* in the shoot meristem. These data indicate that *PNY* has an unexpected function during plant development in regulating the pattern of expression of

flowering-time genes in the shoot meristem by repressing *BOP1/2* gene transcription.

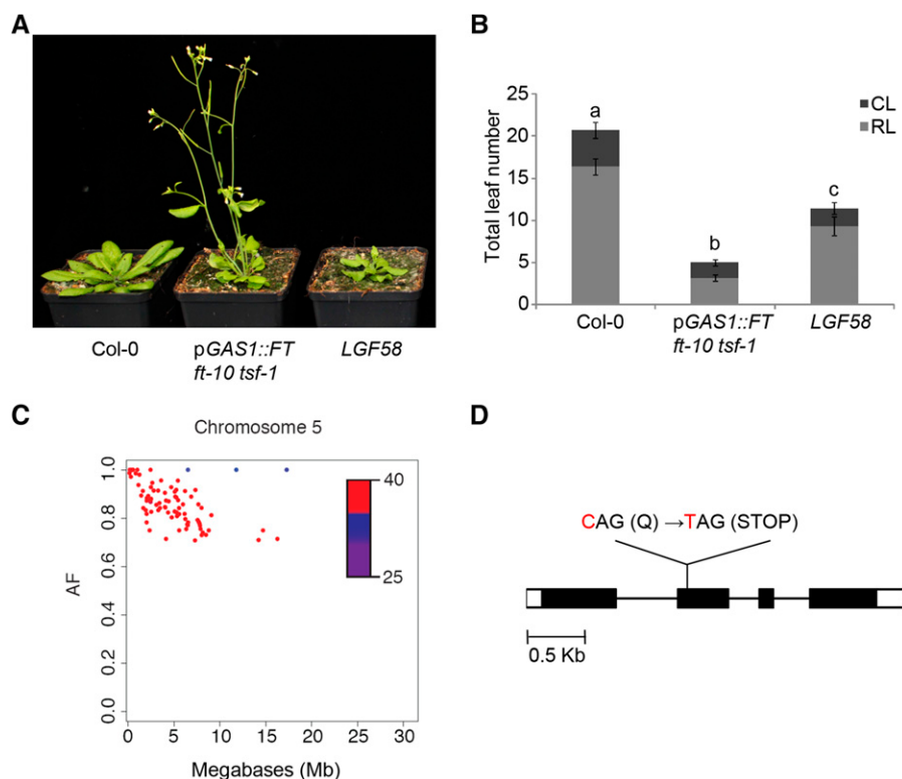
## RESULTS

### A Sensitized Forward Genetic Screen followed by Fast Isogenic Mapping Identifies *PNY* as a Regulator of the FT-Signaling Pathway

A sensitized genetic screen was designed to identify genes affecting the ability of FT to activate flowering. In the double mutant *ft-10 tsf-1*, which carries null mutations in *FT* and its closest homolog *TWIN SISTER OF FLOWERING LOCUS T (TSF)*, the floral promotion activity of FT is abolished, causing late flowering and insensitivity to photoperiod (Yamaguchi et al., 2005; Jang et al., 2009). FT function in *ft-10 tsf-1* can be restored using the transgene *pGAS1::FT* (GALACTOL SYNTHASE1 [*GAS1*] promoter fused to FT), which is active only in phloem companion cells of the minor veins (Jang et al., 2009). Thus, *pGAS1::FT ft-10 tsf-1* plants show early flowering compared with *ft-10 tsf-1* double mutants under LDs and short days (SDs). These plants were used to screen for mutations that suppress promotion of flowering by FT. Seeds of *pGAS1::FT ft-10 tsf-1* were mutagenized with ethyl methanesulfonate (EMS), and late-flowering plants were screened in the M2 generation under SD. Recovered mutants are hereafter called *late flowering in pGAS1::FT ft-10 tsf-1 (lgf)*. Early flowering of *pGAS1::FT ft-10 tsf-1* plants grown under SD is entirely dependent on movement of FT from the leaves to the SAM, so *lgf* mutations were

expected to define genes required for FT function or transport. Around 35,000 M2 plants were screened, and several *lgf* mutants were selected (Supplemental Fig. S1). The *lgf58* mutation most strongly suppressed the early flowering conferred by misexpression of FT (Fig. 1; Supplemental Fig. S1). This mutant also showed other phenotypic abnormalities, such as short stature and lanceolate leaves (Supplemental Fig. S1).

The fast isogenic mapping approach was used to identify the *lgf58* mutation (Hartwig et al., 2012; Schneeberger, 2014). A mapping population was created by backcrossing *lgf58* to *pGAS1::FT ft-10 tsf-1* and self fertilizing the resulting F1 plants. A total of 566 BC1F2 plants were grown, and 174 of them exhibited the late-flowering phenotype of *lgf58* (an approximate ratio of 3:1), suggesting that a single recessive mutation was responsible for the effect. To construct the pool, an individual leaf from each of the 174 plants showing the mutant phenotype was collected. Genomic DNA extracted from the pooled material and the progenitor *pGAS1::FT ft-10 tsf-1* were sequenced using Illumina technology. By applying SHOREmap (Schneeberger et al., 2009b; Sun and Schneeberger, 2015), candidate loci, at which mutant alleles were strongly overrepresented in the pool, were identified on the top arm of chromosome 5 (Fig. 1; Supplemental Fig. S2; for more details, see also "Materials and Methods"). Three loci were selected as high-confidence candidates because they carried a nonsynonymous mutation and showed an allele frequency equal to 1.0 and a quality score equal to 40 (Supplemental Table S1; "Materials and Methods"). Among the highest probability candidates,

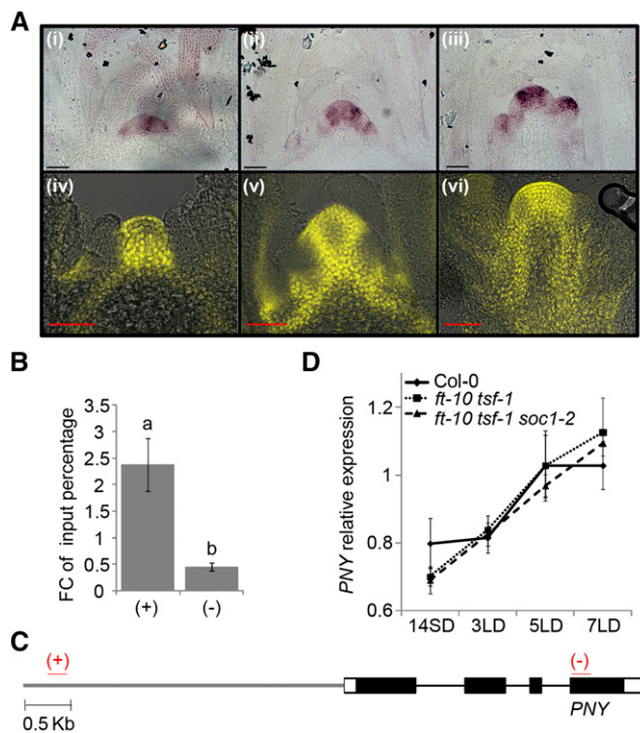


**Figure 1.** Identification and cloning of a functional suppressor of FT. Phenotypic comparison between *lgf58*, *pGAS1::FT ft-10 tsf-1* and Columbia-0 (Col-0) wild-type plants (A) and their flowering time under LD ( $n = 10$ ; B). CL, Cauline leaves; RL, rosette leaves. Letters shared in common between the genotypes indicate no significant difference (Student's  $t$  test,  $P < 0.05$ ). C, Graphic showing the allelic frequency estimations at EMS-induced mutations (AF; y axis) across chromosome 5 (Mb; x axis) of *lgf58*. AFs were calculated by dividing the number of reads supporting the mutant allele by the number of all reads aligning to a given marker. The color code indicates the resequencing consensus (SHORE) score. EMS mutations showing a SHORE score higher than 25 were selected. AFs in chromosome 5 were higher compared with other regions in the genome (see also Supplemental Fig. S2). D, Scheme of the *PNY* locus showing the position of the mutation and the sequence change found in *lgf58*.

the gene *AT5G02030* contained a mutation in the second exon predicted to produce a premature stop codon (Fig. 1; Supplemental Table S1). *AT5G02030* encodes the BELL protein PNY. Mutations in *PNY* cause defects in plant architecture, abnormalities in the fruit replum, and late flowering (Byrne et al., 2003; Roeder et al., 2003; Smith and Hake, 2003; Bao et al., 2004; Smith et al., 2004). Double mutants containing mutations in *PNY* and its paralog *PNF* fail to undergo the transition from vegetative to reproductive phase, even in the presence of high levels of *FT* mRNA expressed from the 35S promoter (Kanrar et al., 2008). Similar phenotypes were also observed in *lgf58* (Fig. 1; Supplemental Fig. S1) and in Col-0 plants carrying the newly isolated mutant allele of *PNY* (hereafter called *pnny-58*) segregated away from pGAS::*FT ft-10 tsf-1* (Supplemental Fig. S3). Previously, *pnny pnf* double mutants were proposed to be blocked in the floral transition due to impairment of the ability of FT to activate the transcription of downstream target genes, such as *AP1*. In agreement with this idea, the mRNA levels of *AP1*, *LFY*, and *SPL4*, which are transcriptionally activated downstream of FT, were dramatically reduced in the *lgf58* mutant compared with pGAS::*FT ft-10 tsf-1*. However, as also described earlier for *pnny pnf* double mutants, *SOC1* mRNA expression was unaltered in *lgf58* (Supplemental Fig. S1; Kanrar et al., 2008). Col-0 plants carrying the *pnny-58* mutant allele grown under LD also showed late flowering compared with wild-type plants and were more extreme than the previously reported mutant allele *pnny-40126* (Smith and Hake, 2003; Ragni et al., 2008; Supplemental Fig. S3). This delay in flowering was corrected by introducing a transgenic copy of the wild-type genomic *PNY* locus into the mutant plants (pPNY::*Venus:PNY pnny-58*), confirming that the mutation *pnny-58* was responsible for the late-flowering phenotype (Supplemental Fig. S4). Moreover, the early flowering of pGAS::*FT ft-10 tsf-1* was also reduced by combining it with the mutant allele *pnny-40126*, but the effect on flowering was less severe than for *lgf58* (Supplemental Fig. S4). These results demonstrate that the sensitized suppressor screen identified a novel allele of *PNY* that causes a stronger delay in flowering than those previously described, and which strongly reduces the ability of *FT* to activate flowering.

### Regulation of *PNY* during Photoperiodic Induction of Flowering

Loss of *PNY* function reduced the capacity of FT to activate flowering in response to inductive LD photoperiods or in transgenic pGAS::*FT* plants expressing higher levels of *FT* mRNA. *PNY* mRNA is expressed in the SAM (Smith and Hake, 2003), but whether it is regulated in response to changes in day length is unknown. *PNY* mRNA distribution was analyzed by in situ hybridization in different environments (Fig. 2). Under SD, *PNY* mRNA was expressed in the central zone of the SAM and excluded from the leaf boundaries



**Figure 2.** Pattern of expression of *PNY* during photoperiod flowering. A, *PNY* mRNA (i–iii) and protein accumulation (iv–vi). Plants were grown for 14 d under SD (i and iv; vegetative stage) and then shifted to LD for 3 (ii and v; floral transition) and 5 additional days (iii and vi; flower development). Scale bars = 50  $\mu$ m. B, ChIP-quantitative PCR (qPCR) showing binding of *SOC1*:GFP on *PNY* promoter. The y axis represents the fold change (FC) of enrichment (percentage of input) between the qPCR results using positive primers (flanking a *CArG*-box motif) and the negative ones (flanking the coding sequence of *PNY*). Letters shared in common between the genotypes indicate no significant difference (Student's *t* test,  $P < 0.05$ ). C, Localization of the primers used for ChIP-qPCR of *SOC1*:GFP on the *PNY* promoter. Region (+) contains a *CArG*-box cis-motif, which was enriched after immunoprecipitation of *SOC1*:GFP. Region (–) was used as a negative control. D, Study of the *PNY* expression in plants shifted from SDs to LDs. Plants were grown for 2 weeks under SDs and then shifted to LDs for 3, 5, and 7 additional days. RNA was extracted from dissected shoot apices. Error bars indicate SD.

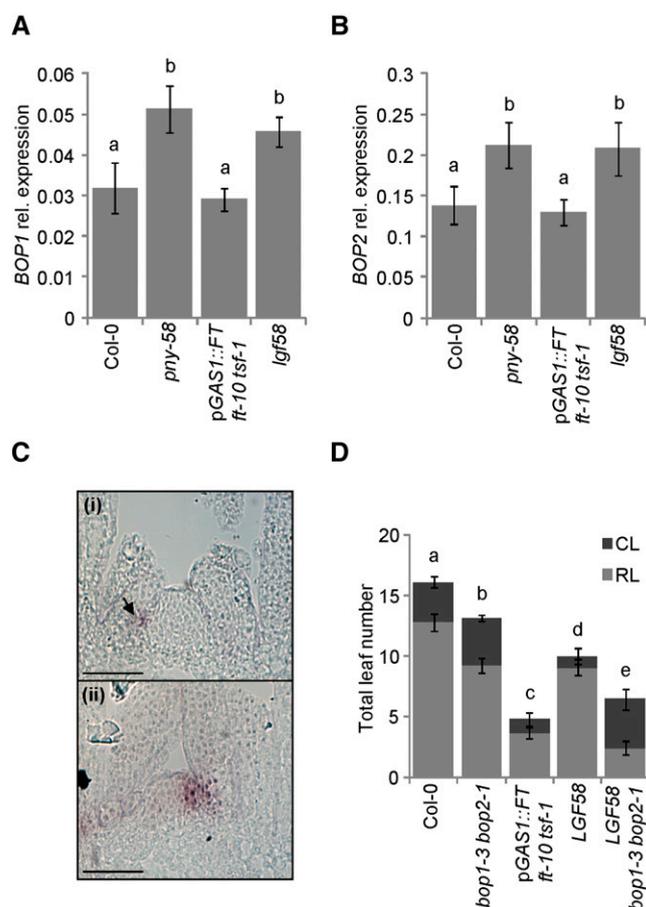
(Fig. 2). After transferring plants to LD, *PNY* mRNA was detected more broadly, but was still not detected in leaf boundaries, nor in floral primordia. The pattern of *PNY* protein expression was also tested by constructing pPNY::*Venus:PNY pnny-40126* plants (Fig. 2; Supplemental Fig. S5). The transgene complemented the defects of the *pnny-40126* mutant. By confocal microscopy, *Venus:PNY* was detected in the same domains shown by in situ hybridization to express *PNY* mRNA in the shoot meristem (Fig. 2). Similarly, its pattern of expression broadened during photoperiodic induction of flowering, and was excluded from leaf boundaries and floral primordia (Fig. 2). Therefore, expression of *PNY* mRNA and its translated product is increased by photoperiod in specific domains of the apex.

*SOC1* is an important mediator of the FT-signaling pathway, and its mRNA expression pattern in the SAM

overlaps with that of *PNY* (Jang et al., 2009; Torti et al., 2012). A recent genome-wide study identified the *PNY* promoter as a direct target of *SOC1* (Immink et al., 2012). Therefore, binding of *SOC1* to the *PNY* locus was tested directly by ChIP-qPCR. In this way, binding of *SOC1* to the *PNY* promoter was confirmed. *SOC1:GFP* bound to the *PNY* promoter at approximately 3,400 bp from its start codon ATG, in a region containing a putative CARG box. This position was very close to the observed binding peak of *SOC1* by ChIP sequencing (Immink et al., 2012). No binding of *SOC1:GFP* was found in a region comprising the fourth exon of *PNY*, which was used as a negative control (Fig. 2). These data suggest that *SOC1* might contribute to photoperiodic induction of *PNY* during flowering. Whether *PNY* expression was reduced by mutations in *FT*, *TSF*, or *SOC1* was also tested. However, no differences in *PNY* mRNA levels were detected in the *ft-10 tsf-1*, *ft-10 tsf-1 soc1-2*, and *soc1-2* mutants compared with Col-0 during the floral transition (Fig. 2; Supplemental Fig. S6). These results suggested that *PNY* expression is increased by exposure to LDs, and that *SOC1* might contribute to this, although at the times tested, no detectable difference in *PNY* expression could be associated with the loss of function of *SOC1*, *FT*, or *TSF* by reverse transcription (RT)-qPCR.

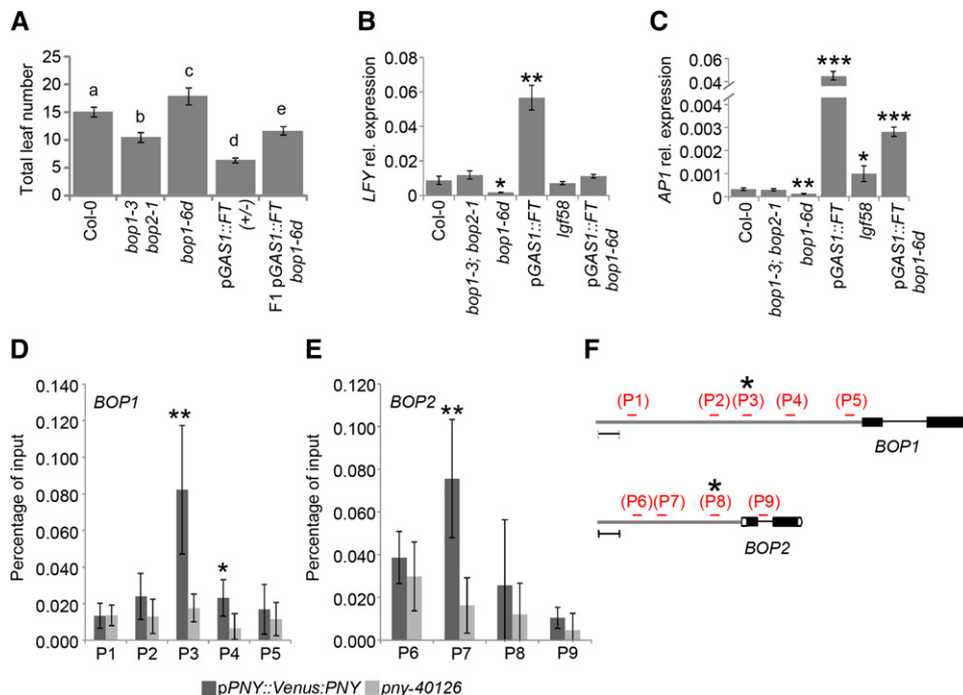
### Suppression of FT Function by *pny* Is Caused by Ectopic Expression of *BOP* Genes

Mutations in *PNY* and other TALE transcription factors, such as *BREVIPEDECELLUS* (*BP*), impair Arabidopsis architecture, particularly shortening internodes and altering silique position and orientation (Byrne et al., 2003; Roeder et al., 2003; Smith and Hake, 2003). These defects in plant architecture in *pny* mutants are associated with broader expression of *BOP* genes in the inflorescence stem and pedicels and are suppressed in *pny bop1 bop2* triple mutants (Khan et al., 2012b). The role of *BOP* genes in the late flowering of *pny* mutants has not been examined, so we tested whether this aspect of the *pny-58* phenotype was due to an increase of *BOP* gene expression in the SAM prior to floral induction. In 7-d-old plants grown under LDs, *BOP1* and *BOP2* mRNA levels were higher in *pny-58* mutants compared with the wild type, as tested by RT-qPCR (Fig. 3). Similarly, these genes were more highly expressed in *lgf58* plants than in *pGAS1::FT ft-10 tsf-1* (Fig. 3). In situ hybridization experiments detected broader expression of *BOP2* in the SAM of vegetative 7-d-old *pny-58* plants than in Col-0 (Fig. 3; Supplemental Fig. S7) and in 10-d-old plants that were undergoing the floral transition (Supplemental Fig. S7). To test whether this increase in *BOP1/2* expression during vegetative development contributed to the late flowering of *pny-58* mutants, the *pny-58 bop1-3 bop2-1* triple mutant was constructed. These plants flowered much earlier than *pny-58* at a similar time to *bop1-3 bop2-1*, which were slightly earlier flowering than Col-0 (Supplemental Fig.



**Figure 3.** *BOP1/2* genes are important for flowering regulation mediated by *PNY*. Expression of *BOP1* (A) and *BOP2* (B) of shoot apices dissected from plants grown under LDs for 7 d. Letters shared in common between the genotypes indicate no significant difference (Student's *t* test,  $P < 0.05$ ). C, In situ hybridizations with *BOP2* probe in shoot meristems of Col-0 (i) and *pny-58* (ii). Plants were grown for 7 LDs and stayed at the vegetative stage. Expression of *BOP2* in Col-0 plants was observed at the boundaries between leaves and the SAM (black arrow). Scale bars = 50  $\mu$ m. D, Flowering time of the *BOP1/2* and *PNY* mutant combinations grown under LDs. CL, Cauline leaves; RL, rosette leaves. Error bars in A, B, and D indicate SD.

S7). Therefore, *BOP1* and *BOP2* are required for the late flowering of *pny-58* mutants. Similarly, introduction of *bop1-3 bop2-1* into the *pGAS1::FT pny-58 (lgf58)* line restored flowering to a similar time to *pGAS1::FT ft-10 tsf-1* (Fig. 3; Supplemental Fig. S7), but the *pGAS1::FT bop1-3 bop2-1 pny-58* line (*lgf58 bop1 bop2*) produced a higher number of cauline leaves than *pGAS1::FT ft-10 tsf-1*, causing a slightly increased number of total leaves. This increased number of cauline leaves could be caused by the down-regulation of *LFY*, because *lfy* mutants produce more cauline leaves (Weigel et al., 1992), and *BOP1/2* promote *LFY* expression in the meristem (Karim et al., 2009). Thus, the levels of *LFY* mRNA in *bop1 bop2* at different developmental stages was quantified by RT-qPCR. Compared with the wild type, the expression of *LFY* in the *bop1 bop2* double



**Figure 4.** *BOP1/2* genes interfere with FT function and are directly bound by PNY. A, Flowering time of plants misexpressing *BOP* genes in the presence of high levels of *FT*. Letters shared in common between the genotypes indicate no significant difference (Student's *t* test,  $P < 0.05$ ). B, Expression of *LFY* of dissected shoot apices of plants grown for 7 LDs. Asterisks indicate statistical differences between Col-0 and other genotypes (Student's *t* test; \*\*,  $P = 0.0004$ ; \*,  $P = 0.006$ ). C, Expression of *AP1* of dissected shoot apices of plants grown for 7 LDs. Asterisks indicate statistical differences between Col-0 and other genotypes (Student's *t* test; \*\*\*,  $P = 0.001$ ; \*\*,  $P = 0.006$ ; \*,  $P = 0.03$ ). ChIP-qPCR of PNY on the promoters of *BOP1* (D) and *BOP2* (E). x axis indicates the primers used for its qPCR. Asterisks indicate statistical differences between pPNY::Venus:PNY and *pnv-40126* (Student's *t* test; \*\*,  $P = 0.0001$ ; \*,  $P = 0.02$ ). F, Localization of the primers used for the ChIP-qPCR experiment on the *BOP1* (top scheme) and *BOP2* (bottom scheme) loci. \*, AP1 binding genomic regions according to Kaufmann et al. (2010). Scale bars = 0.5 kb. Error bars in A to E indicate SD.

mutant was slightly increased during vegetative development (7–10 LDs) but reduced at the reproductive stage (17 LDs; Fig. 6).

Taken together these data support the hypothesis that late flowering of *pnv-58* mutants is due to higher and ectopic expression of *BOP1/2* in the vegetative apex.

Since mutations in *PNY* suppress the capacity of FT to induce flowering, *BOP1/2* up-regulation should also delay the floral transition promoted by this protein. To explore this further, a dominant activation tagging allele of *BOP1* (*bop1-6D*) was used (Norberg et al., 2005). The *bop1-6D* mutants flowered later than Col-0 (Fig. 4). To test whether this occurs through the FT floral-promoting pathway, the double transgenic pGAS1::FT *bop1-6D* was constructed. Flowering time experiments demonstrated that *bop1-6D* delayed flowering of pGAS1::FT (Fig. 4). This result supports the idea that ectopic *BOP* expression impairs activity of the FT-signaling pathway. Indeed, the expression of *LFY* and *AP1*, which is activated in the SAM downstream of FT (Schmid et al., 2003; Moon et al., 2005), was dramatically reduced by overexpression of *BOP1* (Fig. 4).

These data together indicate that the repression of *BOP* genes in the shoot meristem by *PNY* is required for FT to efficiently promote the floral transition.

#### PNY Represses *BOP1* and *BOP2* Transcription by Directly Binding to Their Promoters

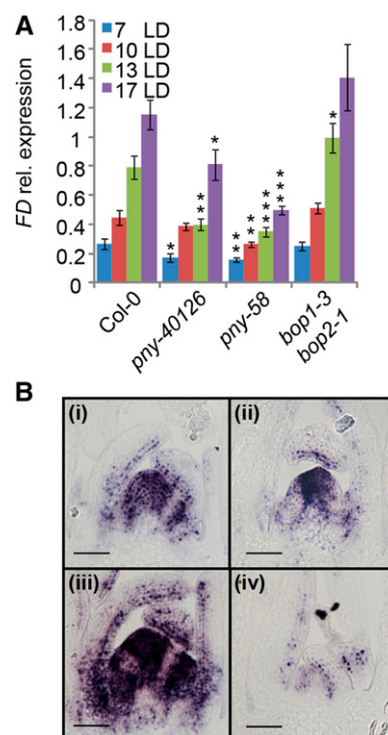
The increase of *BOP1* and *BOP2* expression in *pnv-58* mutants suggested that PNY might directly bind to the promoters of these genes. This possibility is further supported by proteomics analysis, which detected PNY and AP1 in the same transcriptional complex (Smaczniak et al., 2012), and by ChIP sequencing, which identified *BOP1* and *BOP2* as putative direct targets of AP1 (Kaufmann et al., 2010). Whether PNY binds to the same *BOP1/2* promoter regions as AP1 was therefore tested. ChIP-qPCR was performed on chromatin extracted from inflorescences of pPNY::Venus:PNY transgenic plants. The chromatin was immunoprecipitated using a GFP antibody, which detects Venus:PNY on western blots (Supplemental Fig. S7), followed by qPCR with combinations of primers spanning regions of the *BOP1* and *BOP2* promoters. Regions of the *TARGET OF EAT1*

(*TOE1*) and *LFY* promoters, which were shown to be directly bound by the AP1/PNY complex (Smaczniak et al., 2012), were used as positive controls. As expected, Venus:PNY bound to the *TOE1* and *LFY* loci within the same region that was reported for AP1 (Supplemental Fig. S7; Kaufmann et al., 2010). The binding of Venus:PNY to *BOP1* and *BOP2* was then tested. Venus:PNY bound to the *BOP1* promoter at two different positions (P3 and P4). One of these positions (P3) was the same as that reported for AP1 (Fig. 4; Kaufmann et al., 2010). An enrichment of chromatin immunoprecipitated by Venus:PNY was also detected within the *BOP2* promoter. In this case, the enrichment was found in a region located around 1 kb upstream of the one predicted for AP1 (Fig. 4). Notably, some potential PNY binding sites within these genomic regions were identified. For example, the P3 genomic region contains a core motif (ATGGAT) reported as a binding site for BEL1-LIKE HOMEODOMAIN (BLH1; Staneloni et al., 2009). Within the region P7, the two motifs AAATTACCA and AATTATCCT, which are similar to those previously identified as binding sites of BELLRINGER (BLR) in the *AGAMOUS* intronic region (AAATTAAAT, AAAT-TAGTC, and ACTAATTT; Bao et al., 2004; Smaczniak et al., 2012), were also found. However, only shorter versions of these motifs (AATTAT, AATTT, and AAATT) were identified within the P4 genomic region.

Collectively, these data indicate that PNY directly binds and represses the expression of *BOP1/2*.

#### Loss of PNY Function and BOP Overexpression Reduce the mRNA of FD, a Component of the FT-Signaling Pathway, in the Shoot Meristem

PNY loss of function caused a strong reduction in the expression of several genes, such as *SPLs*, *LFY*, and *AP1*, that are expressed at the apex during flowering downstream of FT (Supplemental Fig. S1). These results were in agreement with previous reports on *pnv* *pnf* double mutants (Smith et al., 2004; Kanrar et al., 2008). However, the molecular mechanisms that cause the reduction of expression of these genes and a delay in flowering in *pnv* mutants are not clear. FD directly interacts with FT (Abe et al., 2005; Moon et al., 2005), and mutations in *FD* and *PNY* were found to delay flowering of *pGAS::FT* transgenic plants to a similar extent (Supplemental Fig. S8), suggesting that they might influence the FT-signaling pathway at common positions. Furthermore, analysis of the mRNAs of *SPLs*, *LFY*, and *AP1* by RT-qPCR showed that the *fd-3* mutation suppressed the expression of these genes in a *pGAS1::FT* background to a similar extent as *pnv-58* (Supplemental Fig. S8). These observations suggested that mutations in *PNY* might affect the FT-signaling pathway by reducing *FD* expression to impair photoperiodic floral induction by FT. To test this possibility, the expression of *FD* mRNA in *pnv* mutants was analyzed. *FD* mRNA level was tested by RT-qPCR and was reduced in shoot apices of *pnv-58* and *pnv-40126* mutants compared with Col-0 wild-type plants (Fig. 5). Because *PNY* regulates



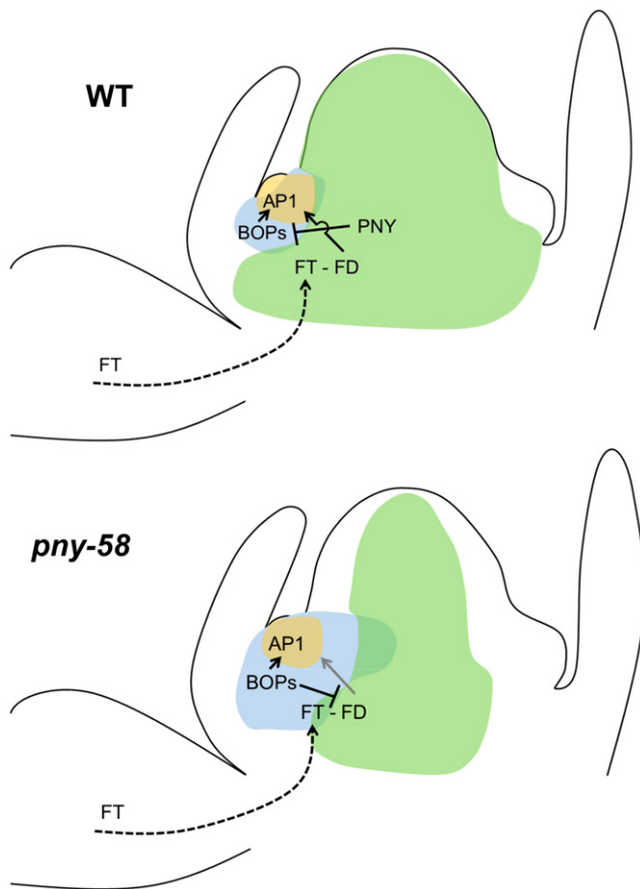
**Figure 5.** *BOP1/2* genes regulate the pattern of expression of *FD*. A, *FD* expression levels in different plants misexpressing *BOP1/2* and *PNY*. RNA was extracted from shoot apices of plants grown during 7 LDs (vegetative stage), 10 to 13 LDs (floral transition), and 17 LDs (reproductive stage). Asterisks indicate statistical differences between Col-0 and other genotypes (Student's *t* test; \*\*\*,  $P = 0.05$ ; \*\*,  $P = 0.01$ ; \*,  $P = 0.001$ ). B, In situ hybridization of plants grown for 10 LDs showing the expression pattern of *FD* in Col-0 (i), *pnv-58* (ii), *bop1-3 bop2-1* (iii), and *bop1-6D* (iv). Bar = 50  $\mu\text{m}$ . Error bars in A to C indicate SD.

flowering through the transcriptional repression of *BOP* genes, the effect of *bop* mutations on *FD* mRNA level was tested. *FD* mRNA was slightly higher in *bop1-3 bop2-1* double mutants compared with Col-0 (Fig. 5). Moreover, in situ hybridization experiments showed that the *FD* mRNA was reduced in the meristem-leaf boundaries of the *pnv-58* (Fig. 5). That this was due to ectopic expression of *BOP* genes was supported by the dramatic reduction in *FD* expression observed in meristems of *bop1-6D* (Fig. 5). By contrast, *FD* mRNA appeared slightly increased in the SAM of *bop1-3 bop2-1* double mutants (Fig. 5). These results suggest that *PNY* controls flowering at least partially through repression of *BOP* gene expression to allow *FD* mRNA to increase in the meristem.

Taken together, these data indicate that the activity of the FT pathway during photoperiodic induction of flowering requires repression of *BOP* genes by *PNY*.

#### DISCUSSION

We performed a sensitized mutant screen to identify genes required for FT signaling during photoperiodic



**Figure 6.** Model explaining the spatial regulation of flowering-related genes by PNY and *BOP1/2*. It has been suggested that FT is delivered from the phloem to the proximity of the SAM (dashed line; Yoo et al., 2013). Once FT is in the shoot meristem, it is supposed to interact with FD to activate the transcription of *AP1* (yellow shade). After the floral induction, *BOP1/2* activate the transcription of *AP1* mRNA in the FM. We showed that PNY directly represses *BOP1/2* expression (blue shade) in the shoot meristem, so that, in the absence of PNY (*pny-58*, bottom), *BOP1/2* pattern of expression becomes broader. We also found that *BOP1/2* repress *FD* expression before floral transition. Thus, the ectopic expression of *BOP1/2* in *pny* mutants leads to the reduction of the expression domain of *FD* mRNA (green shade) in the shoot meristem. Consequently, FT-FD complex formation might be impaired (gray arrow) and the transcriptional activation of *AP1* mRNA reduced.

flowering. Among all of the mutations recovered in this screen, a novel allele of the homeobox gene *PNY* caused the strongest delay in flowering. This *pny-58* allele delayed flowering both in the *pGAS1::FT ft-10 tsf-1* background used for the screen and in Col-0. Expression and genetic analyses indicated that the late flowering of *pny* mutants was caused by ectopic expression of *BOP1* and *BOP2* in the shoot meristem during vegetative development. Thus, the repression of expression of *BOP* genes is a major aspect of the contribution of *PNY* to flowering time control. This conclusion is in agreement with previous reports that repression of *BOP* genes by *PNY* is necessary for wild-type inflorescence development because ectopic *BOP* gene expression

causes abnormalities such as short internodes and reduced apical dominance (Norberg et al., 2005; Ha et al., 2007; Khan et al., 2012a, 2012b). Consistent with *BOP* repression being a fundamental function of *PNY*, we found that *PNY* binds directly to the promoters of both *BOP* genes. In the meristems of *pny* mutants or *bop1-6d* plants carrying a gain-of-function allele of *BOP1*, *FD* mRNA was strongly reduced, and this likely contributed to the reduced responsiveness to FT. We propose that restriction of *BOP* expression to the proximal regions of lateral organs, particularly its exclusion from the shoot meristem by *PNY*, is required for wild-type levels of *FD* expression and thus efficient floral induction in response to FT (Fig. 6).

### Significance of Defining Organ Boundaries for FT Signaling and *FD* Expression

In plant meristems, lateral organ boundaries separate the meristematic zone containing undifferentiated cells from the lateral organs containing differentiated cells (Rast and Simon, 2008; Khan et al., 2014). The *BOP1* and *BOP2* genes are expressed at the base of lateral organs, adjacent to the boundary (Ha et al., 2004; Hepworth et al., 2005; Norberg et al., 2005). They help define the boundary region by repressing homeobox genes, which maintain the meristematic region, by directly activating *ASYMMETRIC LEAVES2* and other genes that specify the boundary (Ha et al., 2007; Jun et al., 2010). We observed that, in *pny* mutants, the *BOP* genes are ectopically expressed in the vegetative meristem, as was previously described for older inflorescence meristems (Norberg et al., 2005; Ha et al., 2007; Khan et al., 2012a, 2012b). Thus, in wild-type plants during vegetative development, *PNY* contributes to positioning of the boundary between the vegetative meristem and leaves by repressing *BOP* gene expression in the meristem. Indeed, we observed that, in the absence of functional *PNY* (*pny-58*), *BOP2* expression becomes broader in the meristem. Similarly, Khan et al. (2015) found that the *BOP1* spatial pattern of expression was enlarged in the inflorescence meristem of *pny pnf*. This restriction of *BOP* expression and proper localization of the boundary is required for correct timing of the floral transition, because *pny* mutants are late flowering, and this is suppressed in the *pny bop1 bop2* triple mutant (Fig. 3; Supplemental Fig. S7). The sensitized screen used to identify the *pny-58* mutation illustrated the importance of *PNY* and the *BOP* genes downstream of FT in the photoperiodic flowering pathway, as previously shown by the capacity of *pny pnf* double mutants to suppress the early-flowering phenotype caused by *35S::FT* (Kanrar et al., 2008).

At the meristem, FT is proposed to activate downstream genes by directly interacting with the bZIP *FD* transcription factor (Abe et al., 2005; Wigge et al., 2005). This relationship between FT-related proteins and *FD* is highly conserved in higher plants, having also been observed in rice and tomato (*Solanum lycopersicum*; Pnueli et al., 2001; Taoka et al., 2011). We found that the

spatial pattern of expression of *FD* in the SAM is regulated by the *PNY*, *BOP1*, and *BOP2* genes. Ectopic *BOP* gene expression in the *pnny* mutant or the gain-of-function *bop1-6d* mutation strongly reduced *FD* transcription (Fig. 5). Whether this repression is due to direct recruitment of BOP proteins to the *FD* gene or an indirect effect of BOP proteins, for example, by activating transcription of boundary genes, remains unclear. Nevertheless, the reduction in *FD* mRNA likely explains the impaired sensitivity of the meristem to FT. Consistent with this idea, *LFY* and *AP1* mRNA levels were reduced in *GAS1::FT ft-10 tsf-1 pnny-58 (lgf58)* plants as observed for *GAS1::FT fd-3* and in *bop1-6d* mutants (Fig. 4; Supplemental Figs. S1 and S8). These results demonstrate that ectopic expression of *BOP* genes in the vegetative meristem reduces *AP1* and *LFY* expression during the early stages of floral transition, as expected for plants with reduced *FD* activity, although later in the floral primordial, BOP proteins have a direct role in the activation of *AP1* (Karim et al., 2009; Xu et al., 2010). Surprisingly, the expression levels of *LFY* were reduced during flower development in the *bop1 bop2* double mutant compared with wild-type plants (Supplemental Fig. S8; Karim et al., 2009). We interpret these data as indicating a dual role for BOP1/2 in floral development. They might act as transcriptional repressors of *LFY* and *AP1* during the early stages of floral transition (probably mediated through *FD*) and promote the expression of these two genes during floral development.

The finding that ectopic expression of BOP function represses *FD* in the meristem suggests that, in wild-type plants, *BOP* gene expression in the boundary region of lateral organs might also repress *FD*, thus reducing its expression in lateral organs and restricting it to the meristem. Interestingly, *FD* expression is excluded from a strip of cells adjacent to lateral organs that might represent the boundary domain (Fig. 5B; Wigge et al., 2005), although higher resolution analysis allowing direct comparison of boundary gene expression with *FD* will be required to test this suggestion. In response to FT signaling, downstream genes are expressed in specific spatial domains of the meristem. Spatial patterning of *FD* expression in the apex may impose spatial constraints on FT signaling by ensuring, for example, that activation of the FT pathway does not occur in boundary regions.

#### Genetic and Molecular Interactions between PNY and Other Homeodomain Transcription Factors in the Regulation of Flowering

*PNY* is a member of the TALE homeodomain transcription factor family. These proteins are divided into two classes, referred to as KNOX and BELL. *PNY* is a member of the BELL class and interacts in the meristem with KNOX proteins, particularly BP and SHOOT MERISTEMLESS, to form heterodimers that regulate transcription. PNF and ARABIDOPSIS THALIANA

HOMEBOX1 (*ATH1*) are other BELL class proteins expressed in the meristem that have been implicated in flowering-time control (Smith et al., 2004; Proveniers et al., 2007). TALE transcription factors were shown to directly regulate genes encoding components of hormonal pathways or other transcription factors (Bolduc et al., 2012; Arnaud and Pautot, 2014). We demonstrate that *PNY*, presumably acting as a heterodimer with KNOX proteins expressed in the meristem, acts directly to repress genes encoding the transcriptional coactivators *BOP1* and *BOP2*.

*ATH1* and *KNAT6*, a KNOX class protein, are expressed at lateral organ boundaries during inflorescence development, where their activation requires *BOP1/2*. During vegetative development, *ATH1* is expressed in the SAM and acts as a floral repressor. A recent study reported that *BOP1* regulates the expression of *ATH1* by directly binding to its promoter (Khan et al., 2015). Therefore, the ectopic expression of *BOP1/2* in the meristem of *pnny* mutants might increase *ATH1* expression, contributing to the late-flowering phenotype. *ATH1* delays flowering at least partially by activating expression of the floral repressor *FLOWERING LOCUS C (FLC)* in the SAM. *FLC* represses *FD* by directly binding to its promoter (Searle et al., 2006). Thus, the repression of *FD* by *BOP1/2* could at least partially be due to increased *ATH1* activity, leading to misexpression of *FLC* mRNA in the SAM (Proveniers et al., 2007). Alternatively, *BOP1/2* might interact directly with the promoter of *FD*. Further studies must be done to discriminate between these possible scenarios.

*PNY* and the related BELL protein PNF are genetically redundant in the promotion of flowering. Nevertheless, *pnny-58* was clearly late flowering in the single mutant, as described for other *pnny* alleles named *blr* (Byrne et al., 2003). The *pnny pnf* double mutant did not flower in any environmental condition tested and was assumed to be impaired in the competence to flower (Smith et al., 2004). Genetic and molecular analyses of the double mutant indicated that expression of *LFY* and *AP1* was strongly reduced in the inflorescence apices of these plants, but *FT* expression in leaves was unaffected (Kanrar et al., 2008). The conclusion that *PNY* PNF acts between FT and *LFY* was supported by the observation that *pnny pnf 35S:LFY* plants produced flowers, but *pnny pnf 35S:FT* plants did not (Kanrar et al., 2008). These results are in agreement with our observation that a primary effect of *pnny* on flowering is reduction of *FD* mRNA, which is required for FT to promote flowering. Similarly, Lal et al. (2011) described a reduction in *SPL4* and *SPL5* expression in *pnny pnf* apices, and activation of both of these genes at the shoot meristem is dependent on FT and *FD* (Torti et al., 2012). However, in contrast to our data, Kanrar et al. (2008) found that *FD* mRNA was present in the meristem of *pnny pnf* plants at levels similar to those found in the wild type; therefore, the mechanism by which FT activity was impaired by *pnny pnf* was unclear. This discrepancy with our data might be due to the age of the plants examined, as we studied *pnny* mutants during vegetative development just prior

to floral induction and found a clear decrease in *FD* mRNA likely due to ectopic *BOP1/2* expression, whereas Kanrar et al. (2008) examined the inflorescence meristem of plants 20 d after floral induction had occurred in the wild-type controls. In support of our data, Jaeger et al. (2013) also reported a reduction of *FD* mRNA levels in *pnv* *pnf* plants grown under inductive LDs, and we found that *FD* was strongly repressed in *bop1-6d* plants that express *BOP1* in the meristem. Therefore, taken together, the data suggest that flowering is delayed in *pnv* and *pnv* *pnf* mutants, at least partially, by reducing FT signaling, and our data indicate that this occurs due to reduced *FD* expression in the vegetative meristem caused by ectopic expression of *BOP1/2*, are directly repressed in wild-type plants by PNY (Fig. 6).

## MATERIALS AND METHODS

### Plant Materials and Growth Conditions

The wild type was the Col-0 ecotype of *Arabidopsis* (*Arabidopsis thaliana*). The transgenic plants *pGAS1::FT*, *pGAS1::FT ft-10 tsf-1*, and *pSOC1::SOC1:GFP soc1-2* were previously described in Jang et al. (2009) and Immink et al. (2012). The mutant alleles used were *pnv-40126* (Smith and Hake, 2003), *bop1-3 bop2-1* (Hepworth et al., 2005), *fd-3* (Abe et al., 2005), *soc1-2* (Lee et al., 2000), and *ft-10 tsf-1 soc1-2* (Torti et al., 2012). The activation-tagged overexpressing line *bop1-6D* was described in Norberg et al. (2005). Plants were grown in climatic chambers under LD (16-h light/8-h dark) or SD (8-h light/16-h dark) conditions with a light intensity of 150 mmol m<sup>-2</sup> s<sup>-1</sup>, 21°C, and 70% relative humidity.

### Molecular Cloning of pPNY::Venus:PNY

Cloning of locus *PNY* was based on polymerase incomplete primer extension (PIPE; Klock and Lesley, 2009) with modifications for large fragments and multiple inserts. All PCR amplifications were done with Phusion Enzyme (New England BioLabs) following the manufacturer's recommendations. Amplification of the *PNY* coding sequence was done from genomic DNA covering from the 5'-untranslated region (5'UTR) until the 3'UTR (primers A1-F/A2-R), obtaining a PCR product of 3.5 kb. The promoter was amplified from -5,553 region until 78 nucleotides of exon 1 (primers A3-F/A4-F) obtaining a PCR product of 5.7 kb. PCR products were independently cloned into pDONR201 (Invitrogen) by BP reaction generating constructs *csPNY-pENTR201* and *pPNY-pENTR201*, respectively. Insertion of the sequence of fluorescent protein Venus (Nagai et al., 2002; Heisler et al., 2005) at the amino terminus was done by amplifying three different PCR elements adding overlap sequences among them: *I-PIPE-1* (Venus), *I-PIPE-2* (coding sequence PNY), and *vector-PIPE* (promoter PNY-pENTR201). Elaboration of *I-PIPE-1* was done by amplifying Venus and adding a linker of nine Ala to exon1 (primers A5-F/A6-R) obtaining a product of 772 nucleotides. Generation of *I-PIPE-2* was done using *csPNY-pENTR201* as template, producing a PCR fragment of 3.3 kb containing the region from exon1 until 3'UTR (primers A7-F/A8-R). Finally, *vector-PIPE* was generated using as template *pPNY-pENTR201*, obtaining a PCR fragment of 7.8 kb (primers A9-F/A10-R) comprising promoter-5'UTR plus *pENTR201* backbone. For the assembly of the different fragments, equimolar amounts of each *I-PIPE* element were mixed while keeping a ratio of 1:10 to *vector-PIPE*. The mixture was cloned into chemical competent DH5- $\alpha$  cells. The final construct (11.8 kb) was verified by digestion analysis and sequencing. Subsequently, the construct was cloned into the binary vector *pEarleyGate301* (Earley et al., 2006) by LR reaction and transformed into *Agrobacterium tumefaciens* GV3101 cells. *pnv-40126* plants were transformed by the floral-dip method (Clough and Bent, 1998). The list of primers used for the molecular cloning can be found in the Supplemental Table S2.

### Mutagenesis and Genetic Screen

Approximately 10,000 seeds (200 mg) of *pGAS1::FT ft-10 tsf-1* were wrapped in miracloth and imbibed for 14 h in 50 mL of 0.1% (w/v) KCl (50 mg of KCl in 50 mL of dH<sub>2</sub>O) at 4°C on a shaker. Then, the seeds were treated with 30 mM

EMS for 12 h. After the treatment, the seeds were washed twice with 100 mL of 100 mM sodium thiosulfate followed by two additional washes with 500 mL of water. The seeds were transferred to a flask containing 2 L of water and distributed in 200 pots by pipetting (50 seeds/pot). M1 generation was grown in a greenhouse under LD conditions. M2 seeds from each pot were harvested together and treated as a pool. Approximately 500 M2 seeds from each pool were used for the genetic screen. The screening of the M2 seeds was performed in climatic chambers under SDs. Mutants showing late flowering compared with *pGAS1::FT ft-10 tsf-1* (*lgf* mutants) were selected, and the phenotypes were confirmed in the M3 generation.

### Flowering Time Measurements

Flowering time was scored as the number of leaves at bolting. The number of rosette leaves was determined when the shoot reached approximately 0.5-cm length. The cauline leaf number was defined when the shoot was totally elongated. At least 10 individual plants were scored by genotype. All experiments were independently repeated at least twice.

### Resequencing and Mapping Strategy

*lgf58* homozygous mutant was crossed with *pGAS1::FT ft-10 tsf-1* to generate the BC1F2 mapping population. A total of 174 late-flowering mutants out of 566 F2 plants were selected. One leaf sample of each was harvested and pooled. Leaf material from the original *pGAS1::FT ft-10 tsf-1* parental was also harvested. Genomic DNA (gDNA) from 1 g of the pooled and the *pGAS1::FT ft-10 tsf-1* leaf material was extracted using a DNeasy Plant Maxi Kit (Qiagen). Four micrograms of gDNA was sent to the Cologne Center of Genomics (Cologne, Germany) for sequencing. Sequencing was performed on an Illumina HiSeq2000 instrument. Up to four independent gDNA samples (i.e. *pGAS1::FT ft-10 tsf-1* and various *lgf* mutants, including *lgf58*) were resequenced in a single HiSeq2000 flow cell lane with a read length of 100 bp (paired end) by using barcoding (multiplexing). After sequencing and applying the quality controls, we obtained a total of 109,948,920 reads from *pGAS1::FT ft-10 tsf-1* and 87,972,438 reads from *lgf58* plants. A total of 101,934,960 (92%) reads from *pGAS1::FT ft-10 tsf-1* and 85,609,714 (97%) reads from *lgf58* were aligned to the reference sequence TAIR10 (*Arabidopsis* Genome, 2000) by applying SHORE (Schneeberger et al., 2009b) and GenomeMapper (Schneeberger et al., 2009a), representing an average coverage of 73 $\times$  and 57 $\times$  the respective resequenced genome (*pGAS1::FT ft-10 tsf-1* and *lgf58* population). Before identifying single nucleotide polymorphisms (SNPs) from the alignment of the pooled *lgf58* plants short read data, an SNP analysis of *pGAS1::FT ft-10 tsf-1* was applied to identify all fixed SNPs of this pool. After removing these SNPs from the SNP analysis of *LGf58*, we obtained a total of 20,137 putative differences from the reference sequence. Of those, 1,174 were further removed as they were located in the mitochondria and chloroplast genome. From the remaining 18,963 putative differences, 2,212 revealed the canonical EMS mutation (G/C:A/T). From those, we obtained a final set of 137 mutations by relaxed filtering for reliable mutations (at least a SHORE quality score of 25 and a minimum allele frequency of 0.7 for the mutated allele). The list of top candidates can be found in Supplemental Table S1. Short reads are available through the European Nucleotide Archive under accession number PRJEB10593.

### Protein Extraction and Immunoblotting Assays

Total protein was extracted from inflorescences, ground in liquid nitrogen and homogenized in denaturing buffer (100 mM Tris-HCl [pH 7.5], SDS 3%, 10 mM dithiothreitol, and 1% protein inhibitor), mixed by vortexing, and rotated for 10 min at 4°C. Cell debris was removed by centrifugation at 14,500 rpm for 10 min at 4°C. Proteins were quantified by the bicinchoninic acid method, and 30  $\mu$ g of proteins was loaded in a gel preceded by boiling for 5 min. Anti-rabbit GFP antibody (Abcam ab290) was used for the western blot. The blot was incubated with SuperSignal Femto West Substrate (Thermo Fisher Scientific) following the manufacturer's protocol and detected with a LAS-4000 Mini-image analyzer (Fujifilm). Coomassie Brilliant Blue was used as the loading control.

### ChIP and qPCR

ChIP was performed as previously described (Andrés et al., 2014; Mateos et al., 2015). GFP antibody from Abcam (ab290) was used to immunoprecipitate the chromatin. Inflorescences (containing flowers until stage 13) of

approximately 4-week-old *pPNY::Venus:PNY* and *pnv-40126* plants grown under LD were collected at zeitgeber time 3 for the ChIP assays. The percentage of input method was used for data normalization (Haring et al., 2007). qPCR values obtained from the immunoprecipitated samples were divided by the qPCR values of a 1:10<sup>2</sup> dilution of the input sample. For validation of the PNY binding to *BOP1* and *BOP2*, several pair of primers spanning the *BOP1* and *BOP2* promoters were tested (Supplemental Table S2). Two biological replicates were performed for each ChIP assay. Only one of the replicates is shown.

### qPCR Methods for RNA Expression Analysis and Genotyping of *pnv-58*

RNA expression analyses were performed as described in Andrés et al. (2014). Total RNA was extracted from plant tissue by using the RNeasy Plant Mini Kit (Qiagen) and treated with DNA-free DNase (Ambion) to remove residual genomic DNA. The RNA was then quantified by using the Nanodrop ND-1000. One microgram of total RNA was used for RT (Superscript II; Invitrogen). Levels of mRNA were quantified by qPCR in a LightCycler 480 instrument (Roche) using the *PEROXIN4* gene (*AT5G25760*) as a standard. Three biological replicates were performed for each RT-qPCR assay. The average of the three replicates is shown. The list of primers used for expression analyses can be found in Supplemental Table S2. Graphs were obtained from three independent technical replicates, although all RT-qPCRs were repeated at least twice and showed identical results.

Genotyping of the mutant allele *pnv-58* was performed by high-resolution melting (Wittwer et al., 2003). *pnv-58* allele carried a single nucleotide change (C > T). Primers flanking this mutation (K617 and K618; Supplemental Table S2) were used to amplify (by PCR) a 79-bp amplicon from gDNA extracted from leaf material (Plant DNeasy Kit; Qiagen). The PCR products were diluted five times in water. Three microliters of the PCR product dilutions was used as a template for the qPCR in a LightCycler 480 instrument (Roche). Primers K617 and K618 were used for the amplification. The PCR conditions were as follows: 95°C for 3 min (pre-incubation) and 22 cycles of 95°C for 20 s, 60°C for 20 s, and 72°C for 20 s. For the melting curve generation, the temperature was increased from 65°C to 97°C (ramp rate 0.11°C/s and 5 acquisitions/°C). The mutant and wild-type alleles could be differentiated by analyzing the melting peaks. The *pnv-58* and wild-type alleles produced a melting peak at 77.66°C ± 0.03°C and 78.33°C ± 0.03°C, respectively.

### In Situ Hybridization and Microscopy Techniques

In situ hybridizations were performed as described in Torti et al. (2012). The *FD* probe was synthesized as described in Searle et al. (2006). The list of primers used to generate the other probes can be found in Supplemental Table S2. For *Venus:PNY* visualization in shoot meristems, a method described previously (Wang et al., 2014) was used, with small modifications. Shoot apices were collected and placed on ice-cold 2.5% paraformaldehyde (PFA; Sigma-Aldrich) prepared in phosphate-buffered saline at pH 7.0. Samples were vacuum infiltrated for 30 min, transferred to fresh 2.5% PFA, and stored at 4°C overnight. The second day, the samples were incubated for 30 min in 1% PFA supplemented with 10%, 20%, and 30% Suc. Then, samples were embedded in OCT (Sakura Finetek) at -20°C. Sections of 35 μm were made using a cryotome (Frigocut 2800; Reichert Jung). Sections containing a visible meristem were selected and mounted with ProLong Diamond Antifade (Invitrogen). The shoot meristems were imaged by confocal laser scanning microscopy (Zeiss LSM780). The confocal laser scanning microscopy settings were optimized for the visualization of *Venus* fluorescent proteins (laser wavelength, 514 nm; detection wavelength, 517–569 nm).

### Statistical Analysis

All of the statistical analyses were performed by using SigmaStat 3.5 software.

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession numbers AT5G02030 (*PNY*), AT3G57130 (*BOP1*), AT2G41370 (*BOP2*), AT1G65480 (*FT*), AT4G35900 (*FD*), AT2G45660 (*SOC1*), AT1G69120 (*AP1*), AT5G61850 (*LFY*), and AT1G53160 (*SPL4*).

### Supplemental Data

The following supplemental materials are available.

**Supplemental Figure S1.** Phenotype of the mutants recovered in the genetic screen and characterization of *lgf58*.

**Supplemental Figure S2.** Scheme of the cloning-by-sequencing of the *lgf58* mutant.

**Supplemental Figure S3.** Phenotypical characterization of mutants carrying the *pnv-58* allele.

**Supplemental Figure S4.** Flowering time of plants carrying mutations that suppress FT function.

**Supplemental Figure S5.** Functional characterization of *pPNY::Venus:PNY* transgenic plants.

**Supplemental Figure S6.** Effect of photoperiod on *PNY* pattern of expression.

**Supplemental Figure S7.** Interaction between *PNY* and *BOP1/2* genes in flowering time control.

**Supplemental Figure S8.** Effect of *BOP1/2* genes on the FT-signalling pathway.

**Supplemental Table S1.** Candidate loci identified by SHOREmap.

**Supplemental Table S2.** List of primers used in this work.

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