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Identification of pathways directly regulated by SHORT VEGETATIVE PHASE during vegetative and reproductive development in *Arabidopsis*

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Abstract

Background: MADS-domain transcription factors play important roles during plant development. The *Arabidopsis* MADS-box gene *SHORT VEGETATIVE PHASE* (*SVP*) is a key regulator of two developmental phases. It functions as a repressor of the floral transition during the vegetative phase and later it contributes to the specification of floral meristems. How these distinct activities are conferred by a single transcription factor is unclear, but interactions with other MADS domain proteins which specify binding to different genomic regions is likely one mechanism.

Results: To compare the genome-wide DNA binding profile of *SVP* during vegetative and reproductive development we performed ChIP-seq analyses. These ChIP-seq data were combined with tiling array expression analysis, induction experiments and qRT-PCR to identify biologically relevant binding sites. In addition, we compared genome-wide target genes of *SVP* with those published for the MADS domain transcription factors *FLC* and *AP1*, which interact with *SVP* during the vegetative and reproductive phases, respectively.

Conclusions: Our analyses resulted in the identification of pathways that are regulated by *SVP* including those controlling meristem development during vegetative growth and flower development whereas floral transition pathways and hormonal signaling were regulated predominantly during the vegetative phase. Thus, *SVP* regulates many developmental pathways, some of which are common to both of its developmental roles whereas others are specific to only one of them.

Keywords: MADS-box, gene regulation, transcription factors, post transcriptional regulation, ChIP-seq, floral transition, floral development, *Arabidopsis thaliana*

Background

In plants organs are formed post-embryonically from populations of undifferentiated cells called meristems. In these meristems, stem cell activity is kept at the central zone whereas at the peripheral part of the meristem primordia arise in which cells differentiate into organs. In flowering plants like *Arabidopsis thaliana* during the vegetative phase the primordia that derive from the shoot apical meristem (SAM) develop into leaves [1,2].

The change to the subsequent generative phase is called floral transition, which is regulated by multiple flowering pathways that are controlled by environmental and endogenous cues. During the floral transition, the SAM undergoes a change in fate and becomes an inflorescence meristem (IM). The *Arabidopsis* IM is an indeterminate meristem and develops multiple determinate floral meristems (FMs) in a spiral manner, which in turn produce a precise number of floral organs arranged in a whorled pattern [1,3,4]. The reprogramming of meristems is regulated by a complex gene regulatory network in which transcription factors represent important key players.

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In *Arabidopsis* the photoperiod, thermosensory, and vernalization/autonomous pathways that respond to environmental signals, and the aging and gibberellic acid pathways that respond to the developmental and physiological state of the plant regulate the floral transition [5]. Many transcription factors encoding genes have been shown to be involved in the regulation of these pathways including those belonging to the MADS-box gene family [6,7]. One of these MADS-box genes controlling flowering time is *SHORT VEGETATIVE PHASE* (*SVP*) [8].

MADS-domain transcription factors have been identified in all eukaryotic kingdoms and in *Arabidopsis thaliana* they are involved in most important developmental processes [9-12]. MADS-domain factors activate or repress transcription by direct binding to short sequences called CArG-boxes that correspond to a 10 nucleotide sequence CC(A/T)₆GG present in the regulatory sequences of target genes. However, this motif can be quite variable allowing some mismatches [10,13]. Moreover MADS-domain proteins form homo and/or heterodimers and are also suggested to form tetrameric MADS-domain complexes [14]. The variety of interactions that many MADS-domain factors can make suggests that they may regulate different subsets of genes during different phases of development and might reflect an enormous regulatory potential [15]. Furthermore, their association with others co-factors probably also influences the affinity and specificity of the complex for specific target sequences [16,17].

During the vegetative phase *SVP* acts as a repressor of flowering since the *svp* mutant flowers very early [8]. *SVP* mediates flowering responses by perceiving signals from different endogenous and environmental flowering pathways such as the thermosensory, autonomous, and GA pathways [6,18]. *SVP* regulates the expression of three floral pathway integrator genes (FPI) that are *FLOWERING LOCUS T* (*FT*), *TWIN SISTER OF FT* (*TSF*), and *SUPPRESSOR OF OVEREXPRESSION OF CONSTANS 1* (*SOC1*) which all promote flowering [18,19]. To maintain plants in the vegetative phase, *SVP* represses the expression of *FT* and *TSF* in the phloem and *SOC1* in the SAM by directly binding to CArG boxes in *FT* and *SOC1* [6,18,19]. During the vegetative phase, *SVP* interacts with another central repressor of flowering time that is *FLOWERING LOCUS C* (*FLC*) and their function is mutually dependent. In fact it has recently been demonstrated that the *SVP-FLC* dimer acts to directly repress *FT* in the leaves and *SOC1* in the SAM [18]. During the floral transition, *SVP* expression gradually decreases until the *SVP* protein completely disappears from the IM [20]. In plants competent to flower, inputs deriving from the flowering pathways converge to repress *SVP* and *FLC* expression [18,19]. During the vegetative phase *SVP* plays an opposite role to its phylogenetically closest related MADS-box gene *AGAMOUS LIKE 24* (*AGL24*), which is a central promoter

of flowering [21,22]. Both *SVP* and *AGL24* directly regulate *SOC1* by binding its promoter on the same binding sites but they have an opposite effect on *SOC1* expression [23].

Interestingly, after the floral transition both *SVP* and *AGL24* are co-expressed in the floral meristem during stage 1 and 2 of flower development [24]. Analysis of the *svp agl24* double mutant, especially at higher temperatures, and the *svp ap1 agl24* triple mutants showed that *AGL24* and *SVP* play redundant roles during these early stages of flower development [20,24,25]. Combining the *svp agl24* double mutant with a weak *ap1* allele showed that *AGL24* and *SVP* together with *AP1* repress floral homeotic genes that control petal, stamen and carpel identity [25]. Protein interaction and genetic studies revealed that *SVP* and *AGL24* are able to form dimers with *AP1* and that this dimer is able to recruit the LEUNIG-SEUSS co-repressor complex [15,25]. Combining the *svp agl24* double mutant with a strong *ap1* allele showed that they are also controlling floral meristem identity since this triple mutant forms on the flanks of the IM new IMs instead of FMs resulting in a cauliflower like curd just as observed in the *ap1 cauliflower (cal)* double mutant [24,26]. Recently Simonini *et al.* [17] have shown that the co-repressor complex composed of LUG, SEU, and *SVP* is also able to repress the ovule identity gene *SEEDSTICK* (*STK*) in a complex together with BASIC PENTACYSTEINE transcription factors.

SVP is a key factor for *Arabidopsis* development and acts both during vegetative and reproductive phases where it plays different roles probably by interacting with different partners to regulate specific sets of target genes. Even though *SVP* is a gene of interest since its first characterization [8], still little is known about the mode of action and the network of genes controlled by this MADS-domain transcription factor. A powerful tool to study *in vivo* the genome-wide DNA-binding patterns of transcription factors is the ChIP-seq technology that consists in ultra-high throughput Solexa (Illumina) sequencing of DNA samples obtained by chromatin immunoprecipitation (ChIP). This technique has been used for a few years to identify direct target genes. At first for human transcription factors like NRSF, STAT1, PPAR γ , and FOXA2 [27-30] and recently this technology has been reported for the identification in *Arabidopsis* of genome wide targets of different MADS-domain proteins such as, SEPALLATA3 (*SEP3*), *AP1*, *FLC*, and *SOC1* [13,31-33] and another important transcriptional regulator such as *AP2* [34]. Moreover genome wide binding site analysis is also possible using the ChIP on chip method, as was done for *AGAMOUS LIKE 15* (*AGL15*), *LEAFY* (*LFY*), *SVP*, and *SOC1* [35-37].

Here we report the use of the ChIP-seq approach to identify genome wide binding sites for *SVP*, during two distinct developmental phases: the vegetative and reproductive

phase. This study allowed us to identify new pathways that are regulated by SVP in vegetative and reproductive tissues and to investigate genome-wide interaction dynamics of a transcription factor during different phases of development.

Results

Genome-wide mapping of SVP binding sites during vegetative and reproductive development

For genome-wide identification of the *in-vivo* binding sites of the SVP MADS-box transcription factor ChIP was performed followed by single end-read sequencing with the Solexa/Illumina GA platform. For the ChIP experiments Arabidopsis *svp* mutant plants expressing epitope tagged SVP were used [20]. The full genomic region of SVP including 3 kb upstream of the start codon was cloned as a C-terminal fusion with GREEN FLUORESCENT PROTEIN (GFP) [38]. Since SVP plays important roles during two distinct non-overlapping phases of development, namely the floral transition [8] and the early stages (stages 1 and 2) of flower development [20,24,25,39], studying the genome-wide binding sites of SVP provides an opportunity to compare the pathways directly regulated by SVP during these two developmental phases. Therefore vegetative phase material was harvested from 2-week-old seedlings grown under short-day conditions, whereas reproductive phase inflorescences with developing flowers of stage 1 to 11 [40] were harvested to analyze its targets during flower development.

Several independent ChIP experiments were performed. As control the same tissues were harvested from wild-type plants that did not express SVP-GFP. ChIP experiments that showed relatively high enrichment for known SVP binding regions (*FT* for the vegetative tissues and *AG* for reproductive tissues) were used to select samples for sequencing (see Additional data file 1, Figure S1) [6,20].

Distribution of SVP binding sites across the genome and within genes

For both vegetative and reproductive tissues as well as for the control, two independent ChIP reactions were sequenced. As in similar experiments [13,31], sequence reads obtained from duplicate experiments for each of the three samples were pooled. Only reads mapping to a unique position on the genome were considered for further analysis. This resulted in about 3 million uniquely mapped reads for the two experiments using inflorescence material, 5 million for experiments performed using vegetative material, and 6 million for control experiments (Additional data file 1, Table S1).

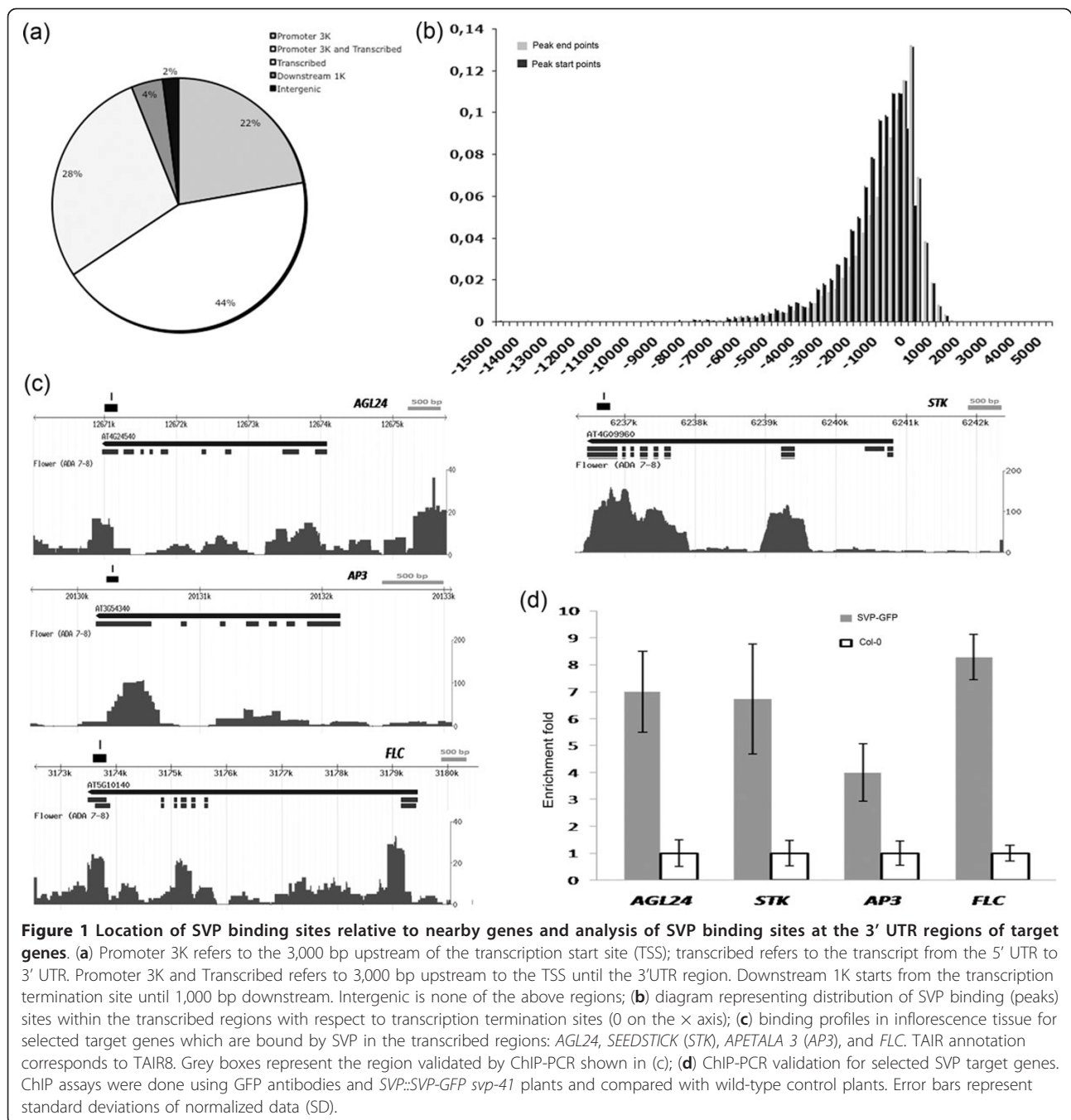
The regions enriched for binding sites were then identified with a strategy broadly similar to the one previously employed for SEP3 and AP1 [13,31], and implemented in the CSAR tool [41]. At a Bonferroni-corrected *P* value of 0.01 this resulted in about 13,000 regions in inflorescence

tissues and 25,000 in seedlings, reduced to about 8,000 and 15,000, respectively, at threshold 10^{-4} , and about 1,300 in both experiments at threshold 10^{-5} (see material and methods and Additional data file 2, Table S2). The overall distribution of SVP-binding sites across the genome in both tissues does not change significantly, and shows that 40% of the sites are located within the 3 Kb upstream of the gene, 27% in the transcribed region, whereas 4% are inside the 1 Kb downstream regions (Figure 1a). Regions falling within the transcribed regions tend to be located towards the 3' UTR/transcription termination (Figure 1b). A similar observation was made on the genome-wide distribution of SEP3 MADS-box protein binding sites [13]; moreover in Kaufmann *et al.* [31] they found that AP1 is able to bind the 3' region of *TERMINAL FLOWER 1 (TFL1)* which is an important shoot identity gene [42]. *TFL1* 3' region is indeed required for proper *TFL1* expression. To confirm binding sites of SVP a set of target genes containing predicted binding sites at the 3' end was selected and analyzed in detail. This set included *AGL24*, *SEEDSTICK (STK)*, *APETALA3 (AP3)*, and *FLOWERING LOCUS C (FLC)*. As shown in Figure 1c, these genes show peaks of enrichment in the inflorescence ChIP-seq data near their 3'UTR regions and, for *STK* and *FLC*, these regions correspond to predicted SVP binding sites (3'UTR is indicated by the striped rectangle). The enrichments on the 3' UTR were analyzed in independent ChIP-qPCR assays confirming that binding at the 3'UTR is significant (Figure 1d).

Candidate target genes were then identified by associating each gene with an overall *P* value calculated from the product of the *P* values of the single binding regions located across the whole gene, encompassing the 3 kb upstream of the transcription start site to 1 kb downstream of the transcribed region. Thus, genes could be ranked according to the overall *P* values obtained. Starting from the ranked gene lists, we selected as high-confidence targets 2,982 genes in seedlings (with a cumulative gene *P* value $< 1.26E-23$) and 2,993 genes in inflorescences (cumulative gene *P* value $< 3.16E-15$) (Additional data file 2, Table S2). The cut-offs on these lists were selected to maximize the number of known targets while excluding the maximum number of genes that were demonstrated to be false positives based on validations with ChIP-qPCR.

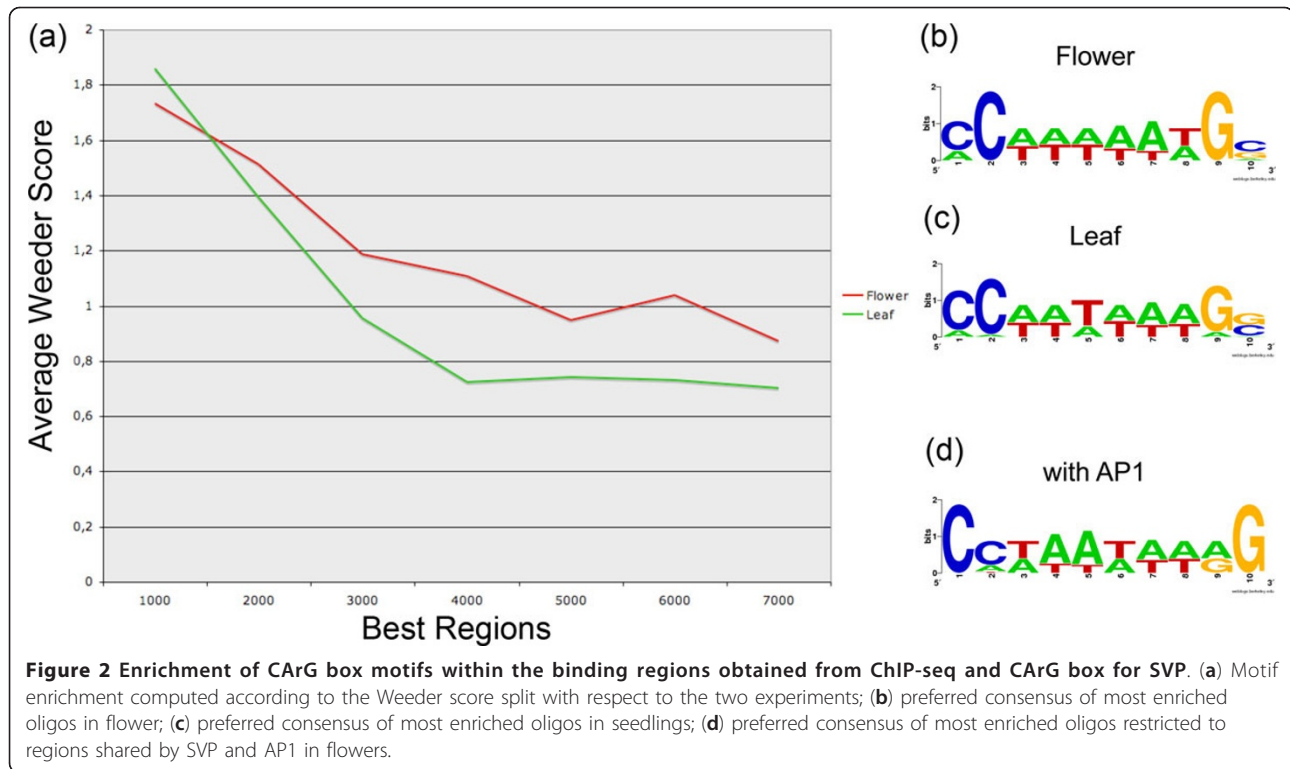
Binding motifs of the SVP protein

MADS-domain proteins are known to bind to different CArG box sequences, including the SRF-type (CC[A/T]6GG), the MEF2-type (C[A/T]8G), and other intermediate motifs (CC[A/T]7G/C[A/T]7GG) [10,43-46]. In order to assess the enrichment of CArG box motifs within the binding regions obtained from ChIP-seq, and to determine whether there is a preferred form of CArG box for



SVP, we ran a tailored version of the motif finder Weeder [47] in order to evaluate separately the enrichment within the regions of each oligonucleotide which could be considered a valid instance of a CArG box given the consensus described before and also including NC[A/T]6GN. Oligonucleotides found to be enriched in the regions were then clustered together to form the motif maximizing the enrichment score. Motif enrichment was computed according to the Weeder score, which compares the number of occurrences within the ChIP enriched

regions to an expected value derived from its number of occurrences genome-wide, computing a log ratio of the fold enrichment. The results are summarized in Figure 2a, split with respect to the two experiments performed and to the ranking of the ChIP regions according to their enrichment *P* value (best 1,000 regions, best 2,000, and so on). Enrichment clearly increases according to peak rank, with higher CArG box enrichment to be found within the peaks more enriched in the ChIP-seq experiments. Enrichment seems to be slightly higher in flower-enriched



regions with respect to leaf-enriched regions. Also, sequence alignment of most enriched oligos in flowers shows NC[A/T]6GN (shown in the sequence logo of Figure 2b and 2c) as a preferred consensus, which differs slightly from the already known forms briefly discussed above but closely resembles the one presented in Tao *et al.* [37]. Finally, oligo analysis restricted to regions shared by SVP and AP1 shows a more canonical CARG box, which is present in the regions with a much higher enrichment (about eight-fold enrichment with respect to the four-fold enrichment in the other regions; Figure 2d).

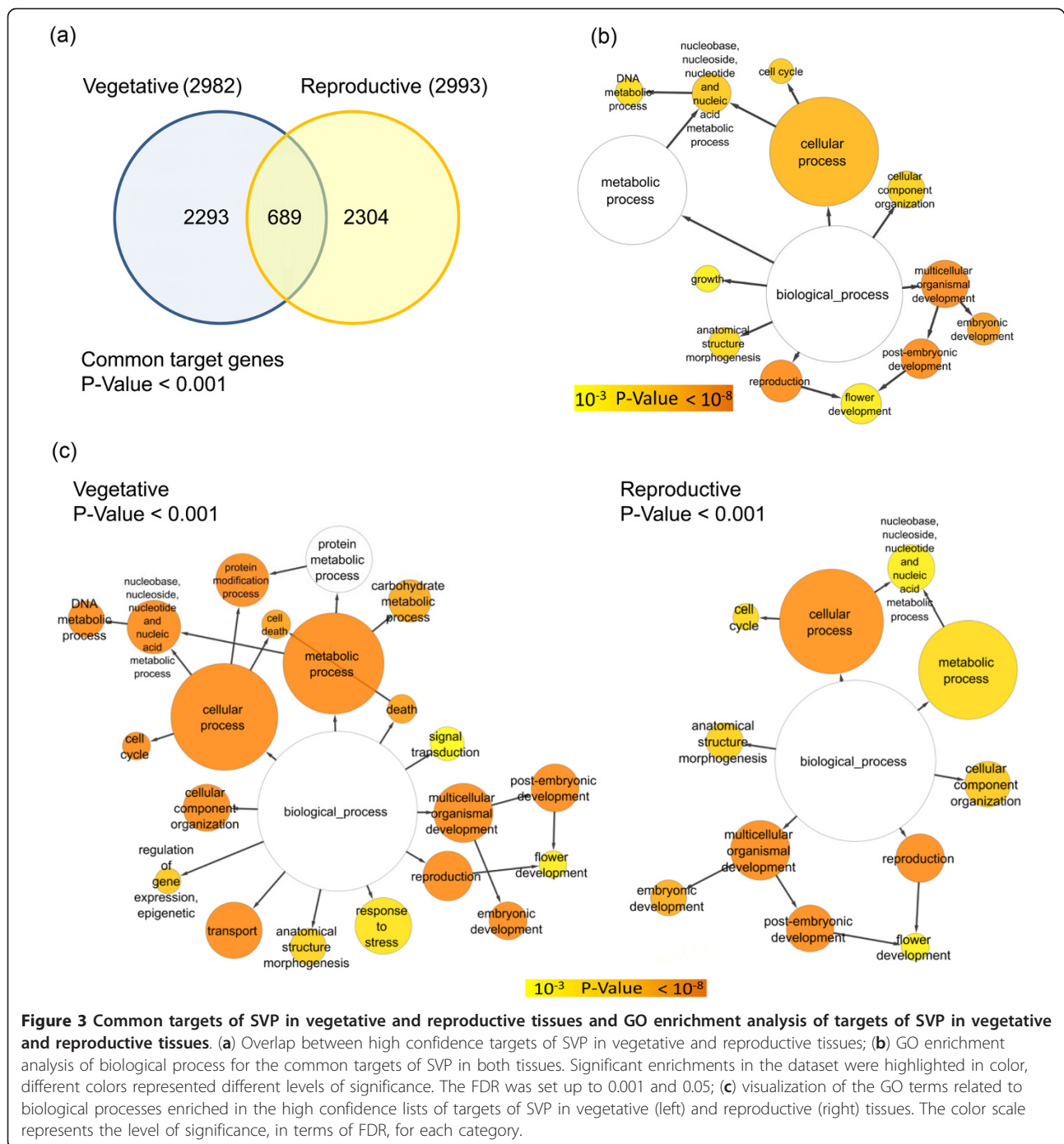
Comparison of SVP binding behaviour during vegetative and reproductive stages

During the vegetative stage SVP acts as a repressor of the floral transition [6,8,18,19], while later it plays an important role during floral meristem specification and organogenesis by regulating expression of organ identity genes [20,25,48]. Here SVP binding sites were identified in seedlings and inflorescences to compare its behaviour at these two stages. A small number of direct target genes of SVP were previously identified in both vegetative and reproductive tissues [6,18,24]. Binding of SVP to these known sites was confirmed in the ChIP-seq data in both conditions, although in some cases (for example *SOC1* in vegetative tissues, see below) the enrichment after the IP was not sufficient to exceed the *P* value threshold employed.

The high confidence lists of putative targets of SVP in vegetative and reproductive tissues show a significant overlap, even if this does not imply a perfect overlap of binding regions for common target genes, as shown in the next section. In total 689 genes appear in both lists, which represents a highly significant overlap (*P* value < 1E-200) (Figure 3a and Additional data file 2, Table S2). The GO analyses reveal that the biological processes enriched in both stages are related to development, cell cycle, and DNA metabolism. These may define a set of genes that reflect the core role of SVP during plant development (Figure 3b).

SVP directly binds to flowering-time genes of different regulatory pathways

Mutations in *SVP* cause early flowering, illustrating a role for SVP in repressing the floral transition, a process controlled by several regulatory pathways [6,8]. Consistent with this function, GO terms related to development, such as 'reproduction' and 'flower development', are significantly overrepresented in the list of putative SVP targets (Figure 3). Moreover, SVP represses flowering by reducing the mRNA levels of *FT* and *TSF* [6,19] key components of the photoperiodic pathway, and of the floral integrator *SOC1* [18]. In the ChIP-seq data, *FT* is indeed bound by SVP, but with a low *P* value (9.5×10^{-7}) (data not shown). Similarly, ChIP-chip experiments performed by Tao and collaborators were not sensitive enough to detect the binding of SVP to the *FT* locus [37]. Recent



work demonstrated that SVP also regulates flowering time independently of *FT* and *SOC1* [18,19]. Thus, we searched the list for known flowering-time regulators. Surprisingly, SVP bound genes involved in several different pathways (Additional data file 1, Table S3), including the circadian clock and photoperiodic pathway, represented by *GIGANTEA* (*GI*) and *PSEUDO-RESPONSE REGULATOR 7*

(*PRR7*), the autonomous pathway, represented by genes such as *FLOWERING LATE KH MOTIF* (*FLK*) and *FLOWERING LOCUS D* (*FLD*), genes encoding components of chromatin associated complexes, such as *CURLY LEAF* (*CLF*), *SWINGER* (*SWN*), and *VERNALIZATION2* (*VNR2*), and the light signaling pathway represented by *PHYTOCHROME A* (*PHYA*).

SVP and the regulation of growth regulator signaling during vegetative development

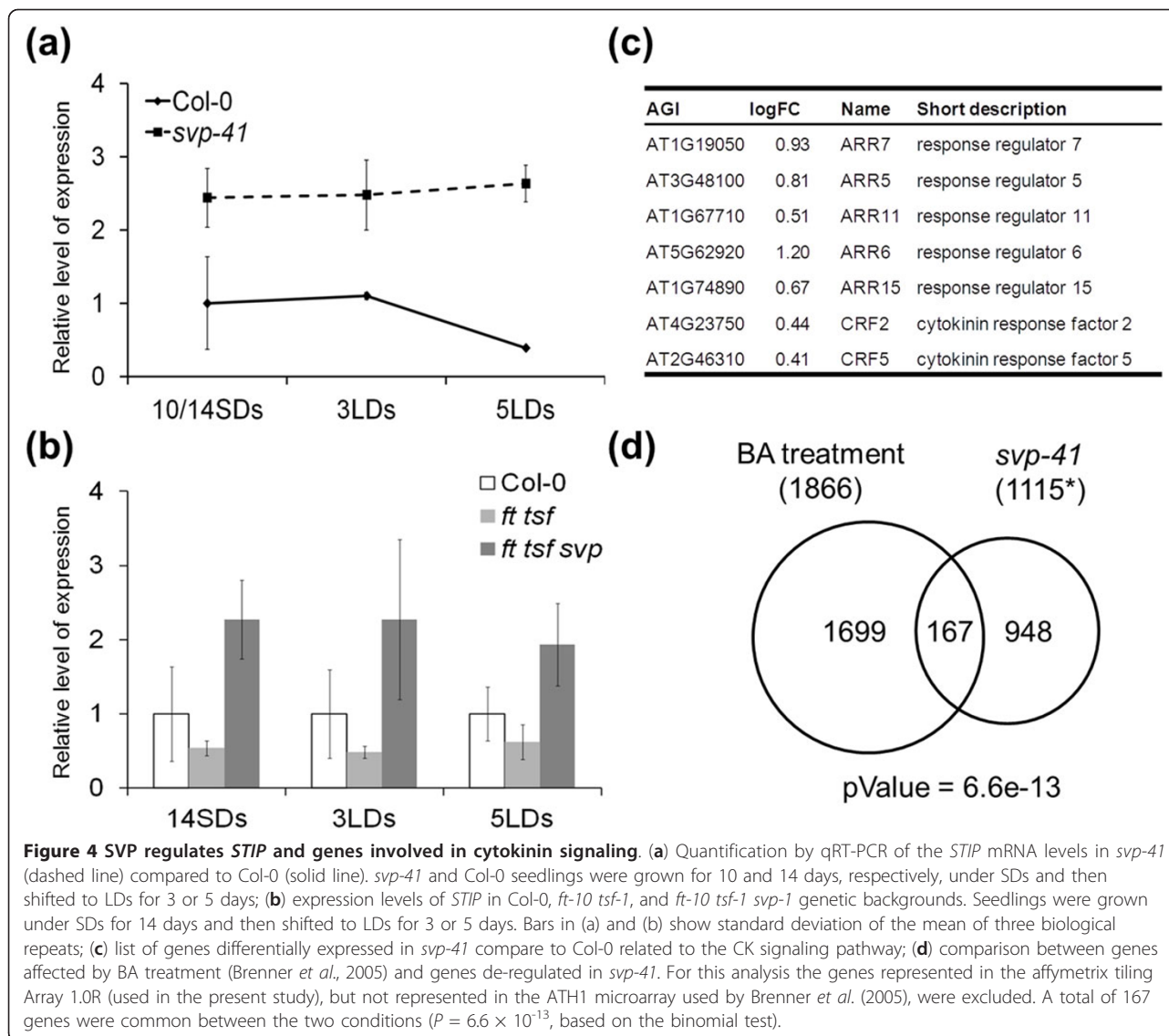
Growth regulators play different roles in flowering-time control and their molecular links to floral homeotic genes have been extensively reported [13,31,32]. SVP targets related to growth regulator signaling, response, transport and metabolism were identified in the ChIP-seq data (Additional data file 3, Table S4). For example, SVP binds directly to *STIP* (*STIMPY*), which was recently described as a component of the cytokinin (CK) signaling pathway [49], during the vegetative phase. The expression levels of this gene were tested in *svp-41* mutants and Col-0. The qRT-PCR experiments showed that *STIP* mRNA was present at significantly higher levels in *svp-41* mutants compared to Col-0 at all time points tested (Figure 4a). We also quantified the expression levels of *STIP* mRNA in *ft-10 tsf-1 svp-41*, which harbours null alleles of *FT* and *TSF* and *SVP* [19]. In *ft-10 tsf-1 svp-41* the expression levels of *STIP* were up-regulated compared to *ft-10 tsf-1* double mutants and Col-0 wild-type (Figure 4b), indicating that SVP controls this gene independently of the FT TSF photoperiodic signals. The effect of SVP on *STIP* expression might indirectly influence the expression of other genes involved in cytokinin signaling. To investigate this possibility a transcriptome analysis was performed by hybridizing RNA extracted from seedlings of wild-type Col-0 and *svp-41* to Affymetrix tiling arrays. The results of these experiments demonstrated that 1,381 genes were differentially expressed (FDR ≤ 0.05) in *svp-41* compared to Col-0 seedlings (Additional data file 4, Table S5). For some of these genes the change in expression in *svp-41* compared to Col-0 was also confirmed by qRT-PCR (Additional data file 1, Figure S2). A GO term test indicated that there is a significant enrichment of genes included in the category 'response to hormonal stimuli' (Additional data file 1, Figure S3 and Table S6). Interestingly seven genes upregulated in *svp-41* mutant were related to cytokinin signaling (Figure 4c). These genes belong to two different groups of cytokinin response genes: the type-A *ARABIDOPSIS RESPONSE REGULATORS* (*ARRs*) and the *CYTOKININ RESPONSE FACTORS* (*CRFs*). These two groups of genes are also transcriptionally activated by *STIP* [49], suggesting that the control of *STIP* by *SVP* has a broad effect on the cytokinin signaling pathway. Indeed, the effect of SVP on CK signaling was also reflected by the significant overlap (P value = 6.6×10^{-13}) between the lists of differentially expressed genes in *svp-41* mutant and the available expression-profiling data of seedlings treated with the CK benzyladenine (BA) [50] (Figure 4d and Additional data file 5, Table S7).

The ChIP-seq and tiling array data also suggested links between SVP and other growth regulators. For instance, SVP bound several genes involved in auxin signal

transduction, such as *BIG*, which encodes a putative auxin transporter required for normal auxin efflux and inflorescence development (Additional data file 3, Table S4) [51,52]. Another gene bound by SVP is *CORONATINE INSENSITIVE 1* (*COI1*), which encodes the jasmonate receptor (Additional data file 3, Table S4) [53,54]. Therefore SVP might affect auxin and jasmonate homeostasis by directly binding to genes encoding key components of their signaling cascade pathways. In agreement with this conclusion, our Tiling array data showed that members of the *SAUR-like auxin-responsive* family were up-regulated in *svp-41* mutant (Additional data file 3, Table S4 and Additional data file 1, Figure S2). In addition, six of the *JASMONATE ZIM-domain* (*JAZ*) genes (*JAZ1*, 5, 6, 7, 8, and *10*), which are part of the jasmonate signaling pathway and are transcriptionally activated by the hormone, were increased in expression in the mutant compared to Col-0 (Additional data file 3, Table S4 and Additional data file 1, Figure S2).

Common targets of SVP and FLC during vegetative development

MADS-domain proteins form multimeric complexes that are proposed to be important in determining their DNA binding specificity. Co-immunoprecipitation analysis and yeast two-hybrid assays demonstrated that SVP interacts with the related MADS-domain protein FLC and genetic data indicate that this interaction is likely functionally important in the control of flowering time [18,55]. Moreover, SVP associates with the promoter region of *SOC1* and the intron of *FT* where FLC also binds [18,39]. Recently the genome wide targets of FLC were identified using ChIP-seq technology [32]. Of these FLC putative targets, 112 were also detected in our experiment as being bound by SVP in vegetative tissue (P value = 1.9×10^{-6}) (Additional data file 1, Figure S4a). Nine of the FLC putative targets were previously validated by ChIP-qPCR and six of them shown to change in expression in *flc-3* mutants [32]. Of these confirmed FLC targets, four were selected to test by ChIP-qPCR if they were also bound by SVP (Figure 5b, c). Of these four FLC targets, three were bound by SVP in a similar location. One of these was *JAZ6*, which was bound by FLC in its promoter region and its expression is increased in *flc-3* [18]. *JAZ6* expression was also upregulated in *svp-41* (Figure 5a), however it was not enriched in our ChIP-seq experiment, and this was confirmed by independent ChIP-qPCR analysis, suggesting that the changes in *JAZ6* expression caused by SVP are not an effect of direct binding (Figure 5c). A second confirmed FLC target, *AGL16*, was not enriched in the SVP ChIP-seq data, however the region bound by FLC showed a low but consistent enrichment in ChIP-qPCR of SVP. This experiment suggests that SVP is weakly bound to the same region of *AGL16* as FLC, and



the low enrichment might explain why it was not detected in the ChIP-seq experiment. *AGL16* expression was not changed in *svp-41* compared to Col, similar to what was observed in *flc-3*. A third confirmed FLC target was SVP, and ChIP-qPCR confirmed that SVP binds to the same region in its own promoter as FLC. These ChIP-qPCR experiments demonstrate that there is a strong but not complete overlap in the targets of FLC and SVP.

SVP auto-regulates its gene expression in vegetative tissue and flowers

The ChIP-seq data indicated that SVP binds to its own genomic region in vegetative tissue and flowers. However, regions actually bound in both tissues may differ. This differential binding was confirmed by independent ChIP-qPCR experiments on two specific regions named I and II (Figure 6 a-c), located approximately 2,000 bp

upstream of the 5'UTR and in the terminal part of the SVP first intron, respectively. As shown in Figure 6b and 6c, SVP binds site I in floral tissue but not in vegetative tissue, whereas site II is bound in both tissues. Whether binding of SVP influenced its own expression was tested in different ways. In addition to the microarray experiment described above, another transcriptome analysis was performed by hybridizing RNA extracted from inflorescences of wild type Col-0 and *svp-41 agl24 ap1-12* to affymetrix tiling arrays. In this experiment 246 genes were differentially expressed (FDR ≤ 0.05) in *svp-41 agl24 ap1-12* compared to Col-0 inflorescences (Additional data file 4, Table S5). The tiling array expression data showed that SVP mRNA was downregulated in the *svp-41* single mutant in vegetative tissues (logFC -1.13; $P=0.001$) as well as in inflorescences of the *svp-41 agl24-2 ap1-12* triple mutant (logFC -0.86;

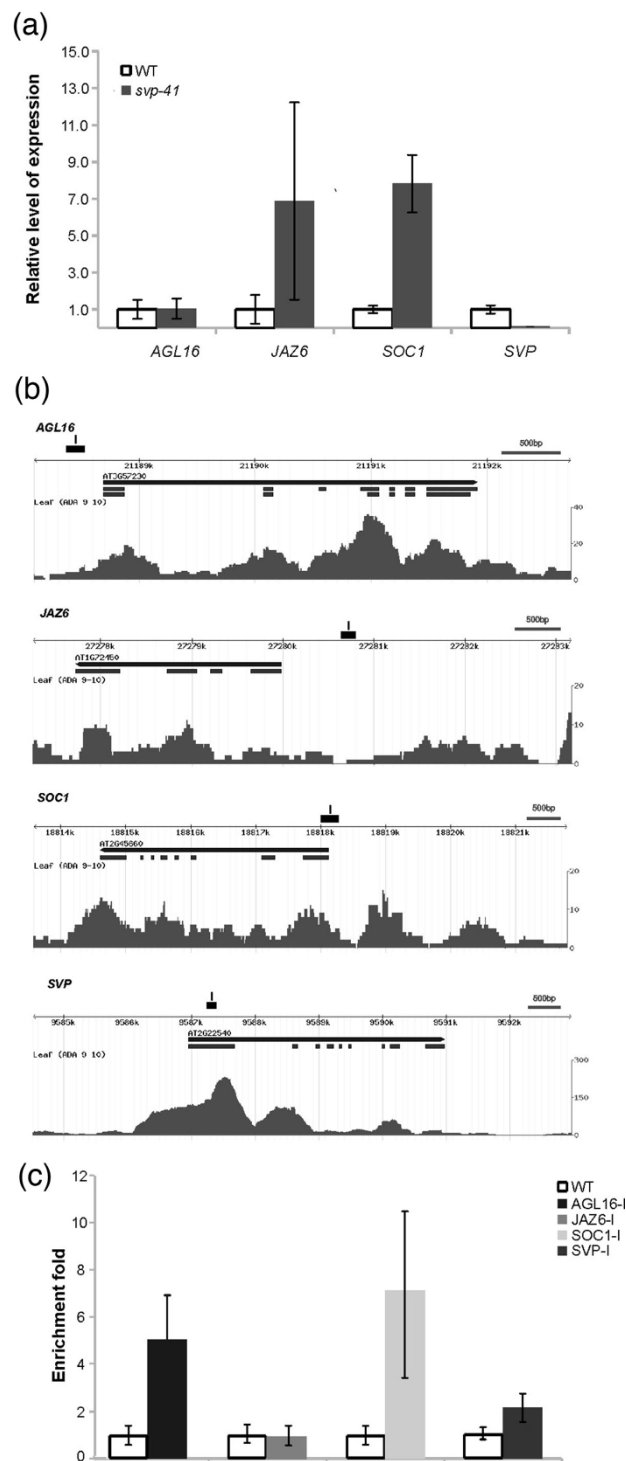


Figure 5 Common targets of SVP and FLC. (a) Expression of known direct targets of FLC in *svp-41*. Data represent expression of selected genes in microarray experiment with FDR <0.05. The expression level of each gene in *svp-41* was normalized to the level of wild type Col-0. Error bars represent SDs of normalized data; (b) binding profiles of ChIP-seq experiment for the selected genes. TAIR8 annotation corresponds to TAIR8. Grey boxes represent the region validated by ChIP-PCR which are shown in panel (c); (c) ChIP-PCR validation of selected genes using anti-GFP antibodies using seedlings of wild type Col-0 and SVP::SVP-GFP *svp-41* lines. Results are expressed relative to actin. Error bars represent SD.

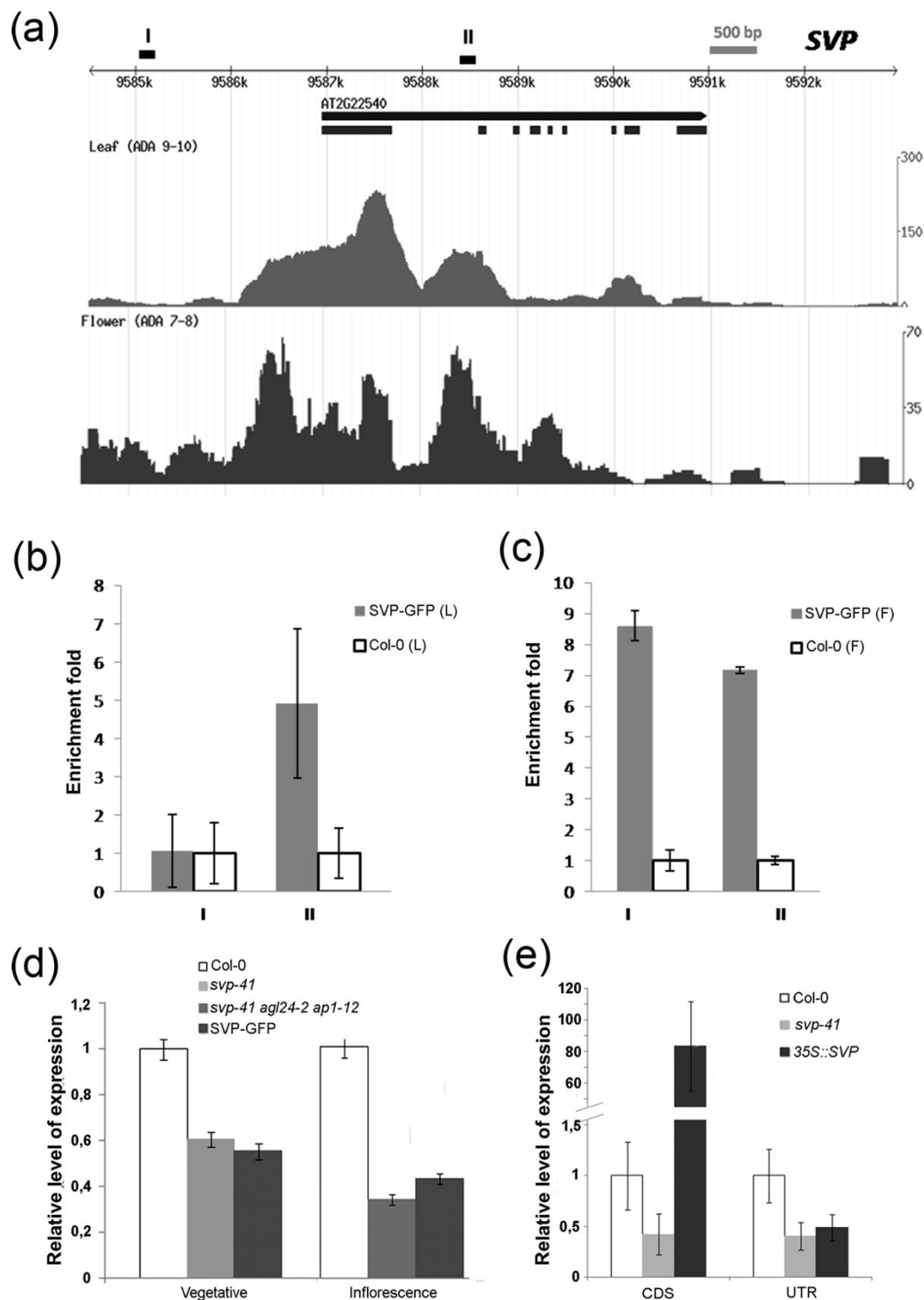


Figure 6 SVP binds and regulates itself. (a) Binding profiles for SVP on *SVP* genomic locus in seedlings (upper panel) and inflorescence (lower panel) tissues. TAIR annotation corresponds to TAIR8. Grey boxes represent the region validated by ChIP-PCR in panels (b) and (c); (b) and (c) ChIP-PCR validations for two specific regions named I and II. ChIP assays were done using GFP antibodies and *SVP::SVP-GFP* *svp-41* plants and compared to wild-type control plants. ChIP-PCR validation in vegetative (b) and reproductive tissue (c); (d) qRT-PCR expression analysis using primers for the *SVP* 3'UTR region. RNA was extracted from wild-type Col-0, *svp-41*, and *SVP::SVP-GFP* *svp-41* seedlings and from wild-type Col-0, *svp-41* *agl24-2* *ap1-12* triple mutant, and *SVP::SVP-GFP* *svp-41* inflorescences; (e) qRT-PCR expression analyses using primers for the *SVP* 3'UTR region and coding region. RNA was extracted from wild-type Col-0, *svp-41*, and 35S::*SVP* seedlings. In all graphs error bars represent the standard deviation of normalized data (SD).

$P=0.02$). This downregulation was validated by qRT-PCR using independent *svp-41* single mutant, *svp-41 agl24-2 ap1-12* triple mutant and wild-type cDNA samples obtained from RNA extracted from seedlings and inflorescences (Figure 6d). Since *svp-41* is a deletion mutant in which two base pairs are deleted in the second exon resulting in a frame-shift of the open reading frame [8], this reduction in mRNA level might be due to non-sense-mediated decay [56]. To investigate this possibility, we performed qRT-PCR assays using primers designed on the 3'UTR region of the endogenous *SVP* gene, which is not present in the *SVP::SVP-GFP* fusion construct. RNA was extracted from wild-type, *svp-41* and *SVP::SVP-GFP svp-41* seedlings and from wild-type, *svp-41 agl24-2 ap1-12* and *SVP::SVP-GFP svp-41* inflorescences (Figure 6d). The results confirmed a reduction in mRNA level also in *SVP::SVP-GFP svp-41* tissues suggesting that indeed this reduction in *SVP* mRNA level seems to depend on the mRNA instability in the mutant background. As an alternative approach the abundance of *SVP* mRNA expressed from the endogenous gene was tested in plants in which *SVP* was overexpressed from a *35S::SVP* transgene. A qRT-PCR strategy was used in which the cDNA expressed from the transgene and endogenous gene can be distinguished (Figure 6e). This experiment demonstrated that *SVP* mRNA expressed from the endogenous locus is reduced in *35S::SVP* plants. Taken together our data suggest that *SVP* directly regulates its own expression, and that it probably acts to repress its own transcription.

Genes involved in meristem development are targets of *SVP* at two developmental stages

Genes involved in meristem development were enriched as *SVP* targets in both vegetative material and flowers. *SVP* is expressed in the SAM during the vegetative stage [6,8,18,19]. In addition it plays an important role during floral meristem specification and organogenesis [25,48]. Consistent with this idea a significant enrichment of *SVP* target genes related to post-embryonic developmental processes was detected in the ChIP-seq results of both vegetative and reproductive samples (Figure 3c). Due to the expression pattern of *SVP*, putative targets with annotated functions in meristem development were screened for directly (Additional data file 3, Table S4). The *CLV-WUS* feedback loop plays a central role in maintaining meristematic activities [57]. In the ChIP-seq data *CLV1* and *CLV2*, two important players in *WUS* regulation, are targets of *SVP* in vegetative tissues and *CLV1* is also bound during reproductive development. Additionally, according to the ChIP-seq data, the HD-ZIP III encoding genes *PHABULOSA (PHB)*, *PHAVOLUTA (PHV)*, *REVOLUTA (REV)*, and *HOMEODOMAIN GENE 8 (ATHB8)*, which regulate post-embryonic

meristem initiation [58], are also bound by *SVP* in vegetative tissue. Furthermore, *PHB* which is a regulator of the size of the *WUS*-expression domain [59], is also bound by *SVP* in the floral meristem. In order to test whether the binding of *SVP* to some of these genes affects their spatial pattern of expression we performed RNA *in-situ* hybridization experiments. A broader expression pattern of *PHB* and *CLV1* was observed in shoot apical meristems of *svp-41* mutants than Col-0 wild-type plants grown for 2 weeks under SDs (vegetative phase) (Figure 7a, b, d, e). However, these differences might be due to the larger size of the *svp-41* meristem compared to Col-0 at this stage. Thus, the patterns of expression of *PHB* and *CLV1* were also compared in 10-day-old *svp-41* mutants and 2-week-old Col-0 plants, which have SAMs of similar size. Confirming our previous result *PHB* and *CLV1* mRNA were detected in a broader region of the *svp-41* (10 SDs) SAM compared to Col-0 (Figure 7c and 7f). These results together with the ChIP-seq data suggest that *SVP* directly regulates the expression pattern of these genes. Furthermore, *KANADII (KAN1)* and *KAN2*, involved in the establishment of abaxial-adaxial polarity in lateral organs produced from the apical meristem, resulted also to be direct targets of *SVP* in inflorescences. It has been hypothesized that complementary regions of action of the class III HD-ZIP genes and *KANADI* genes leads to the establishment of adaxial and abaxial domains in developing lateral organs. The possible role of *SVP* and other MADS-domain proteins in the regulation of part of these genes in reproductive tissues is presented below.

Genome wide targets of *SVP* during flower development and comparison with the targets of *AP1* and *SEP3*

During the early stages of flower development (stage 1 and 2) *AP1* interacts with *SVP* and the dimer recruits the SEU-LUG repressor complex to control the expression of homeotic genes to maintain the floral meristem in an undifferentiated state [25]. At late stage 2, when *SVP* expression is switched off, *AP1* interacts with *SEP3* to control sepal and petal identity. Recently, genome-wide binding studies for *SEP3* and *AP1* during inflorescence development were published [13,31] providing the opportunity to compare these datasets with the one obtained here for *SVP*.

A total of 265 common putative targets for both *SVP* and *AP1* were identified (P value $<7.2E-06$) (Additional data file 6, Table S8 and Additional data file 1, Figure S4). This overlap is expected because *SVP* and *AP1* act redundantly during floral meristem specification where their expression domains overlap [24]. Interestingly transcription factors are enriched among common targets. In addition *SVP* binds to *AP1*, suggesting that it regulates a

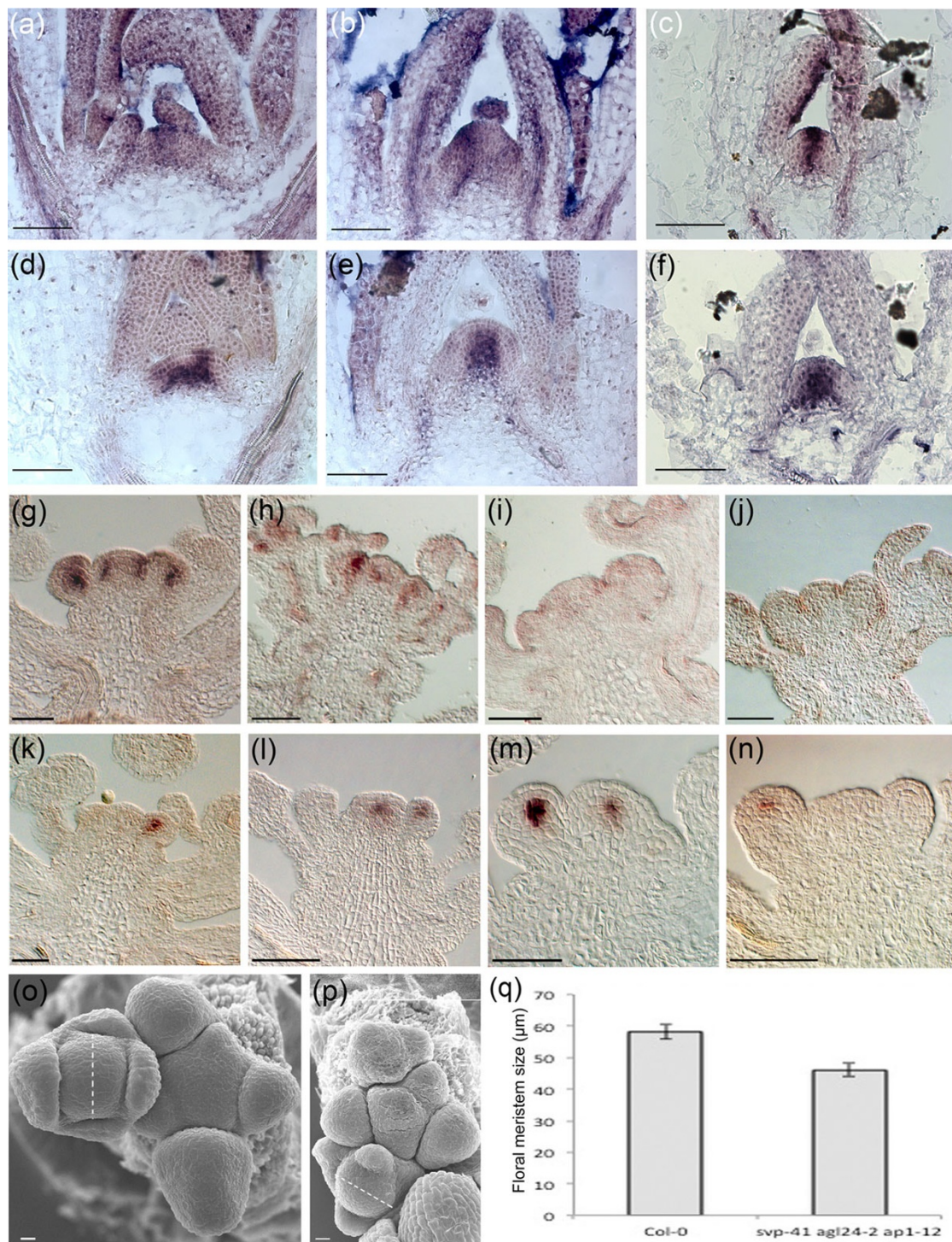


Figure 7 Expression analysis of meristem developmental genes by *in-situ* hybridization analysis in vegetative and reproductive tissues and floral meristem size analysis. (a-c) Patterns of expression of *PHB*: (a) 14-day-old wild-type, (b) 14-day-old *svp-41*, and (c) 10-day-old *svp-41* mutant; (d-f) patterns of expression of *CLV1*: (d) 14-day-old wild-type, (e) 14-day-old *svp-41* mutant, and (f) 10-day-old *svp-41* mutant; in both *svp-41* 10 and 14-day-old seedlings the *PHB* and *CLV1* mRNA were detected in a broader region of the SAM compared to Col-0; (g, h) expression of *ARF3* in wild type and *svp-41 agl24-2 ap1-12* inflorescence respectively; (i, j) *KAN1* expression pattern in wild-type and *svp-41 agl24-2 ap1-12* inflorescences; (k, l) *CLV1* expression in wild-type and *svp-41 agl24-2 ap1-12* inflorescence; (m, n) expression profile of *WUS* in wild-type and *svp-41 agl24-2 ap1-12* inflorescences, its expression seems to be higher in the wild-type FM than in the triple mutant FMs at the same developmental stage. The scale bar represents 50 μm. (o) View of wild-type inflorescence; (p) view of *svp-41 agl24-2 ap1-12* inflorescences; central zone of triple mutant FMs at stage 3 were compared to those of wild-type plants. The scale bar represents 10 μm. (q) Diagram showing the difference in FMs size between the wild-type and *svp-41 agl24-2 ap1-12* triple mutant central dome, error bars represent standard error (SE).

functionally redundant gene as well as itself. The overlap between the targets of SVP with those published for SEP3 [13] revealed 413 (P value $<5.91E-10$) genes that are bound by both of these MADS domain transcription factors (Additional data file 6, Table S8 and Additional data file 1, Figure S4). *KAN1*, *CLV1*, *PHB*, and *ARF3* also named *ETTIN*, that are present in the subset of genes bound by SVP and AP1, are also present in the list of genes regulated by both SVP and SEP3.

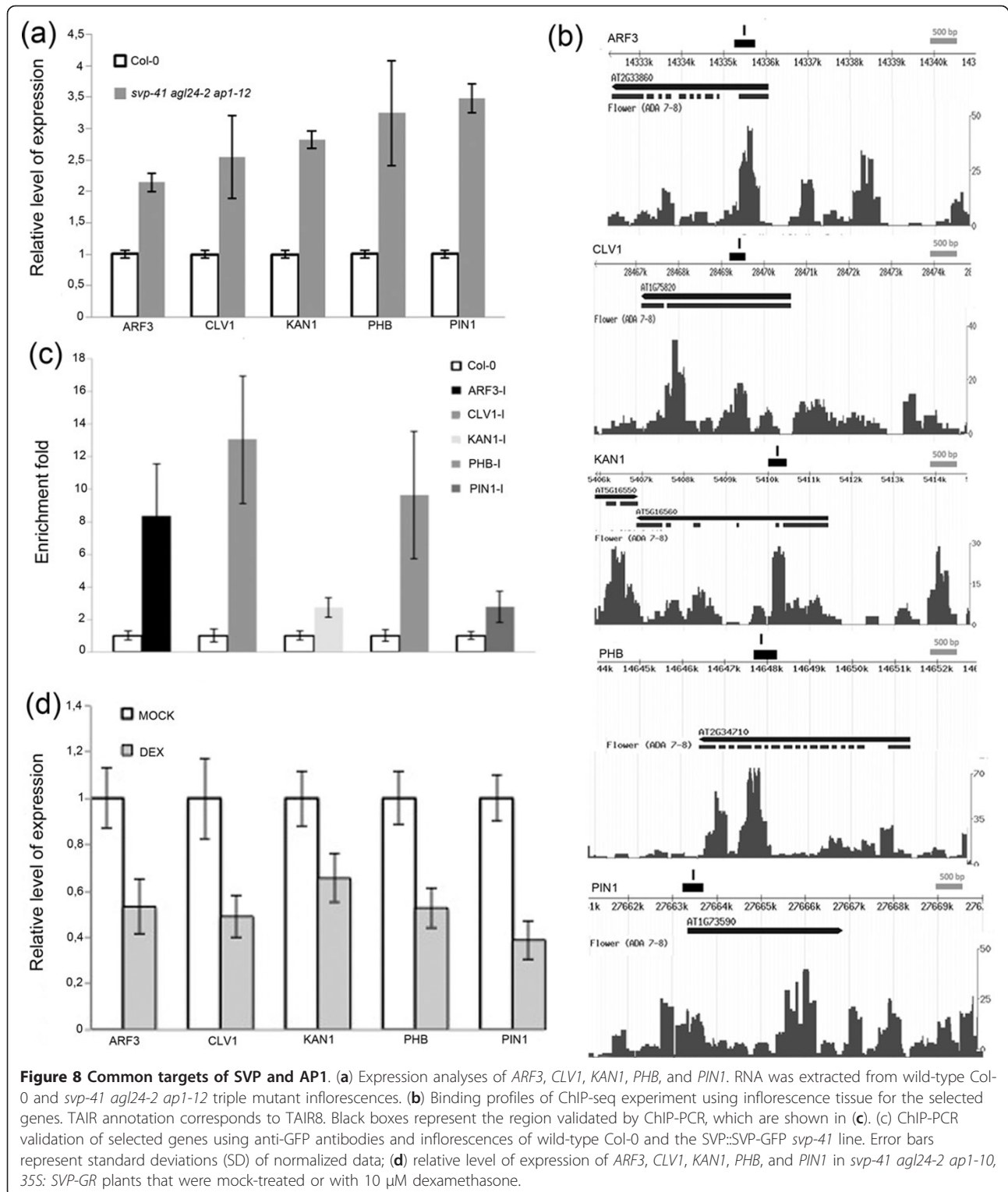
Transcriptome data obtained from the tiling array hybridization experiments using RNA extracted from inflorescences of Col-0 and the *svp-41 agl24 ap1-12* mutant showed that the number of differentially expressed genes were considerably fewer than those found by comparing the vegetative tissue of *svp-41* and Col-0 wild-type plants (Additional data file 4, Table S5). However, the number of deregulated genes might be underestimated in this analysis because the whole inflorescence of *svp-41 agl24 ap1-12* mutant plants were used, whereas *SVP* expression is restricted to stage 1-2 FMs only. Therefore, altered expression of several targets might not be detected in this material. To overcome this we also checked the expression of putative SVP target genes by a qRT-PCR approach, collecting the most inner parts of Col-0 and *svp-41 agl24 ap1-12* inflorescences, avoiding the already opened flowers. Both *KAN1* and *PHB* mRNAs were increased in abundance in the *svp-41 agl24-2 ap1-12* mutant background compared to wild-type (Figure 8a) and the enrichment of these genes observed in the CHIP-seq experiment was confirmed by means of independent CHIP-qPCR analysis (Figure 8b and 8c), suggesting a direct regulation of these genes by both SVP and AP1 during flower development. Interestingly also *CLV1*, which plays an important role in establishing and maintaining floral meristem identity [60], is a direct target of both SVP and AP1 in reproductive tissue and its expression was increased in the *svp-41 agl24-2 ap1-12* triple mutant compared to wild-type (Figure 8a-c). Another transcription factor encoding gene that is bound by SVP and AP1 and upregulated in *svp-41 agl24-2 ap1-12* inflorescences is *ARF3* (Figure 8a-c). ARFs are proteins that are activated by convergent auxin flow. Dynamic changes in auxin fluxes are mediated by PIN proteins and interestingly SVP and AP1 can interact with the genomic region of *PIN1*. Analysis by qRT-PCR showed increased levels of *PIN1* mRNA in *svp-41 agl24-2 ap1-12* inflorescences in comparison to the wild-type control, suggesting a direct role of SVP and AP1 in its regulation which was confirmed by independent CHIP-qPCR experiments (Figure 8a-c). We further examined the expression of *ARF3*, *CLV1*, *KAN1*, *PHB*, and *PIN1* in response to SVP activation using the functional steroid-inducible system. The *svp-41 agl24-2 ap1-10* triple mutant was transformed with a

construct in which the 35S promoter directs a fusion between SVP and a part of the rat glucocorticoid receptor (GR), as reported previously [61]. The *svp-41 agl24-2 ap1-10* mutant forms cauliflower like curds since its unable to establish FM identity and therefore it proliferates IMs instead. The obtained transgenic plants showed upon induction with the steroid dexamethasone (DEX) rescue of the development of FMs and flowers that resembled those of the *agl24-2 ap1-10* double mutant (Additional data file 1, Figure S5). We treated the inflorescences twice, at time 0 and again after 8 h with DEX and collected the material after 24 h from the first treatment. This time point was selected according to Smyth et al. [40], since they showed that the duration of stage 1 of flower development is 24 h. *ARF3*, *CLV1*, *KAN1*, *PHB*, and *PIN1* expression levels were all decreased after DEX treatment of *svp-41 agl24-2 ap1-10* 35S::SVP-GR inflorescences, confirming that SVP acts as a repressor of those genes (Figure 8d).

To investigate the changes in expression profiles of some of these target genes, we performed *in-situ* hybridization experiments using wild type and *svp-41 agl24-2 ap1-12* inflorescences (Figure 7g-n). For *ARF3*, *KAN1*, and *CLV1* the expression pattern was not changed suggesting that the upregulation of these genes is not due to ectopic expression. Interestingly *in situ* using a specific probe for *WUS* clearly showed that in comparison to wild-type, in stage 2 FMs this gene was lower expressed in the *svp-41 agl24-2 ap1-12* triple mutant. Since *svp-41 agl24-2 ap1-12* flowers show reduced numbers of floral organs compared to wild-type or any of the single mutants [25], we wondered if these defects were caused by changes in meristem size. Therefore the central zone of FMs at stage 3 of flower development of the *svp-41 agl24-2 ap1-12* triple mutant and wild-type were compared. The size of the central zone is defined by the distance between the opposite lateral sepals (Figure 7o-q). The *svp-41 agl24-2 ap1-12* FMs were significantly smaller, as compared to those of wild-type plants (Table 1 and Figure 7q). Taken together all these data suggest a role of SVP in the control of FM size, probably by modulating the expression of genes involved in the CLV-WUS pathway.

SVP binds in reproductive tissues to genes encoding post-translational regulators

Interestingly, the high confidence list of SVP target genes in inflorescence tissue exhibits a significant enrichment of genes related to Cullin-RING ubiquitin ligase complexes, mainly involved in post-translational regulation of substrate proteins by attaching poly-ubiquitin chains that target the substrate for 26S proteasome degradation [62,63]. The substrate specificity of CUL4-RING-LIGASES (CRL4s) is exerted by proteins that contain a DWD box (DDB1-binding WD-40 box)



or a WDxR sub-motif [64-67]. Proteins with these motifs are referred to as potential DCAF (DDB1-CUL4 ASSOCIATED FACTOR) proteins [67], which may target proteins for ubiquitination [64,68]. However, they

have also been implicated in chromatin mediated transcriptional control [69]. In Arabidopsis, 119 different putative DCAF proteins have been identified [67] and our ChIP-seq experiments suggest that nearly half of

Table 1 Floral meristem size

	Floral meristem mean \pm SE (μ m)
Col-0	$n=8$ 58.1 \pm 2.2
<i>svp-41 agl24-2 ap1-12</i>	$n=8$ 46.7 \pm 2

Col-0 vs. *svp-41 agl24-2 ap1-12*: Two sample T-test, $t = 3.9200$, $DF = 14$, $P=0.0015$.

them (47 of 119) are targets of SVP in both tissues tested and more than half of these (26 of 47) are putative SVP targets in reproductive tissues (Additional data file 1, Table S9).

Among the putative DCAF floral SVP targets to which a function in floral development had not previously been ascribed (Additional data file 1, Table S9), we selected *WDR55* as a case study for detailed analyses of its function as a SVP target in flower development.

The regulation of *WDR55* by SVP forms as an alternative pathway for the regulation of *AG*

WDR55 was shown to interact with DDB1A, suggesting a regulative role through a putative CUL4-DDB1^{*WDR55*} E3 complex, and plays a major role in *Arabidopsis* reproductive development. *WDR55* is required for gametogenesis and embryogenesis and is suggested to be involved in auxin-dependent regulation of embryo development [70].

In order to verify that *WDR55* expression requires *SVP*, we performed qRT-PCR analyses on *svp* double and triple mutant combinations. Compared to wild-type, *WDR55* transcripts were reduced in abundance in the double mutant *svp-41 agl24-2* (30°C) and in the *svp-41 agl24-2 ap1-12* mutant background (Figure 9a). The binding of SVP to *WDR55*, as observed in the ChIP-seq experiment, was confirmed by means of independent ChIP-qPCR analysis (Figure 9b), suggesting that changes of *WDR55* expression in *svp-41 agl24-2* and *svp-41 agl24-2 ap1-12* are due to the direct action of SVP during flower development.

A recent report describes two mutant alleles of *WDR55* that demonstrate a requirement of *WDR55* in gametophyte development and function, as well as for setting up the embryo body plan. The weaker of these alleles, *wdr55-2*, displayed close to mendelian ratios of mutant seeds (22.7%) and no homozygous plants could be identified, although a small fraction (2%) could be expected from the genetic data [70]. In order to screen for the theoretical presence of homozygous plants in the progeny, we allowed a large number of seeds from heterozygous *wdr55-2* plants to germinate for a prolonged period on MS-2 agar plates containing glufosinate (BASTA) selection. Indeed, we identified a class of late germinating, small seedlings that initially were smaller than the glufosinate sensitive seedlings (3.6%, $n = 1,035$). However, this

class was BASTA resistant and thus carried the *wdr55-2* mutation.

Generally, *wdr55-2* seedlings supported growth, but were severely delayed compared to wild-type. In particular, *wdr55-2* inflorescences were smaller than wild-type and had fewer flowers. Upon inspection we found that the mutant floral organs were generally smaller and often morphologically distinct from wild-type (Figure 9 and Additional data file 1, Figure S6). The sepals were thinner and often fused at early stages and did not separate completely at maturation (Figure 9d and 9e, Additional data file 1, Figure S6b, c and Table S10). The petals were smaller and thinner, as well as being non-uniform in size (Additional data file 1, Figure S6e and Table S10). The stamens were smaller and never occurred in sixes as in wild-type Col (Additional data file 1, Figure S6 and Table S10). The *wdr55-2* flowers also displayed homeotic transformations (Figure 9g, i). We observed unfused carpels (Figure 9f), carpeloid sepals (Figure 9g), petals that resemble stamens filaments and carpeloid filaments with ectopic papillar cells (Figure 9i) at a moderate frequency. New flowers appeared to grow out from whorl 1 or 2 at a low frequency (Figure 9j) and most of the flowers appeared to be asymmetric in flower organ organization (Figure 9h).

Due to the homeotic transformations observed in *wdr55-2* flowers, we checked the expression of the organ identity genes *APETALA3* (*AP3*), *PISTILLATA* (*PI*), and *AGAMOUS* (*AG*) by *in-situ* hybridization (Figure 9k-n and Additional data file 1, Figure S7). The *in-situ* analysis shows that in the *wdr55-2* mutant, the expression pattern of both *AP3* and *PI* is maintained as wild-type plants (Additional data file 1, Figure S7).

AG is expressed in the inner part of the floral meristem where stamen and carpel primordia develop. During flower development *AG* expression is restricted to whorls 3 and 4 (Figure 9k). The *in-situ* analysis shows that in the *wdr55-2* mutant, *AG* is expressed in chimeric organs that develop in the second whorls (Figure 9l) as well as in carpeloid-sepals developing in first whorls (Figure 9m) where stigmatic tissues and carpeloid structures are detectable. *AG* is expressed already in early stages of flower development, in particular stage 1 (Figure 9n), but the architecture of inflorescences in *wdr55-2* makes precise staging difficult.

SEU, *LUG*, *AP1*, and *SVP* are involved in *AG* regulation, and by mutation ectopic *AG* expression is found [25,71-73]. *SEU* and *LUG* are thought to be cadastral genes, and are involved in the control of expression boundaries of floral homeotic genes [71,73] and they interact to repress *AGAMOUS* (*AG*) in the outer two whorls of the flower [72,73]. The SVP-AP1 dimer binds the LUG-SEU repressor and directly regulates *AG* expression during early stages of flower development

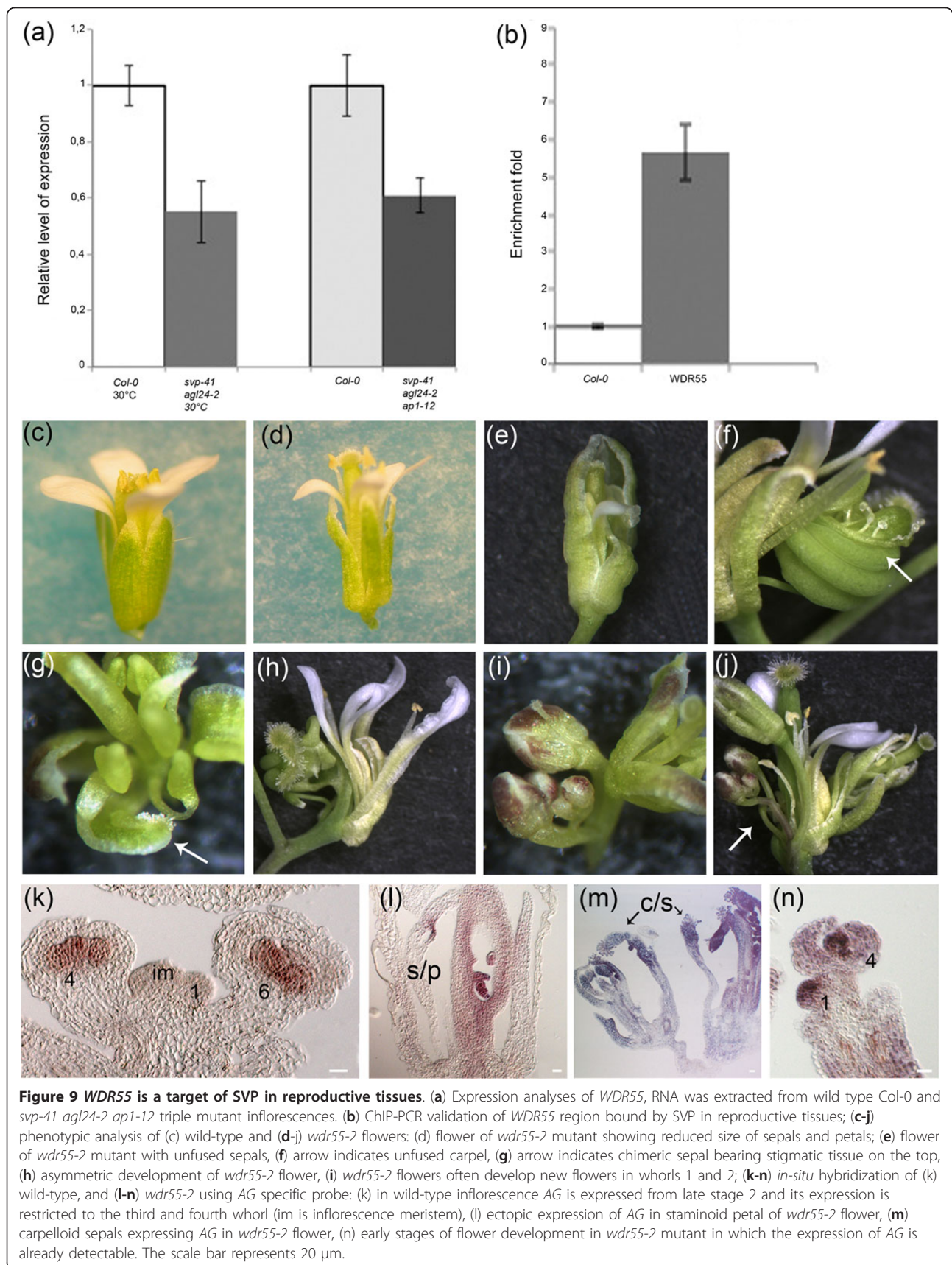


Figure 9 *WDR55* is a target of SVP in reproductive tissues. **(a)** Expression analyses of *WDR55*, RNA was extracted from wild type Col-0 and *svp-41 agl24-2 ap1-12* triple mutant inflorescences. **(b)** ChIP-PCR validation of *WDR55* region bound by SVP in reproductive tissues; **(c-j)** phenotypic analysis of (c) wild-type and (d-j) *wdr55-2* flowers: (d) flower of *wdr55-2* mutant showing reduced size of sepals and petals; (e) flower of *wdr55-2* mutant with unfused sepals, (f) arrow indicates unfused carpel, (g) arrow indicates chimeric sepal bearing stigmatic tissue on the top, (h) asymmetric development of *wdr55-2* flower, (i) *wdr55-2* flowers often develop new flowers in whorls 1 and 2; **(k-n)** *in-situ* hybridization of (k) wild-type, and (l-n) *wdr55-2* using *AG* specific probe: (k) in wild-type inflorescence *AG* is expressed from late stage 2 and its expression is restricted to the third and fourth whorl (im is inflorescence meristem), (l) ectopic expression of *AG* in staminoid petal of *wdr55-2* flower, (m) carpeloid sepals expressing *AG* in *wdr55-2* flower, (n) early stages of flower development in *wdr55-2* mutant in which the expression of *AG* is already detectable. The scale bar represents 20 μ m.

[20,25]. To investigate the regulation of *AG* through *WDR55* further, a Yeast-2-Hybrid (Y2H) was performed with *SEU*, *LUG*, *AP1*, and *SVP*. Upon repeated testing, however, *WDR55* did not interact with any of these proteins (data not shown). This could be due to weak interactions, and thus not detectable in our Y2H system, or *WDR55* does not directly interact on a protein level with these *AG* regulators.

Taken together, our data suggest a role of *WDR55* in floral development. In particular it seems to control the pattern of *AG* expression independently from *LUG-SEU* repressor complex, indicating an additional pathway by which *SVP* repress *AG* expression. However, the function of *WDR55* in flowers does not seem to be restricted to the regulation of the boundaries of *AG* expression as exemplified by the *ag-1 wdr55-2* double mutant (Additional data file 1, Figure S8).

Discussion

The MADS-domain factor *SVP* has different functions during development. An 'early' function as a repressor of the floral transition and a 'later' function in floral meristem identity specification [6,8,18,20,24,25,48]. These two functions are also reflected by *SVP* expression, which is present in the leaves and SAM during the vegetative phase, is repressed in the meristem when plants switch to reproductive development and then reappears in the floral meristem during the early stages of flower development [8,24]. Whether *SVP* regulates different or similar sets of genes during these two phases of development is unknown. We employed ChIP-seq analysis to study the genome-wide binding behavior of *SVP* during these phases. *SVP* was found to bind to approximately 3,000 genes at both stages of development. Some genes were regulated by *SVP* at both stages of development, such as those in pathways regulating meristem development, whereas others were specific to one of the stages. One mechanism by which these differences in target gene specificity are likely to occur is through interactions between *SVP* and other MADS domain protein partners generating complexes with different specificities. Consistent with this idea, comparison of the targets of *SVP* and two of its partners, *AP1* and *FLC*, showed similarities and differences.

Genome-wide ChIP-Seq experiments reveal several roles for *SVP* in modulating vegetative development

SVP bound to approximately 3,000 genes during vegetative development. GO terms analysis of these genes identified functional categories such as 'reproduction' and 'flower development' as being significantly over-represented in the list of putative *SVP* targets (Figure 3c). Similar results were previously found by Tao *et al.* [37]. These authors performed ChIP-chip experiments and identified a total of

328 genes bound by *SVP* during floral transition [37]. Comparison of the *SVP* target list of Tao *et al.* [37] and the list of targets of *SVP* at the vegetative stage presented here showed that only 15 genes are in common between the two datasets (Additional data file 7, Table S11). This discrepancy might occur for several reasons. First, Tao *et al.* made use of hybridization to Tiling arrays (ChIP-chip) to identify the genomic regions bound by *SVP* whereas in the present study these regions were identified by direct sequencing. As described previously, the set of peaks identified by the two technologies can be significantly different [74]. Second, in the ChIP-chip experiments of Tao and collaborators [37] *SVP* was expressed from the constitutive *CaMV35S* promoter whereas for the experiments shown here *SVP-GFP* was expressed from the native *SVP* promoter. MADS-domain transcription factors (including *SVP*) are expressed in specific tissues and interact with different partners to bind DNA in a tissue-specific manner [18], so the ectopic expression of *SVP* in all plant tissues and cell-types, as in *35S::SVP* plants, may affect the detection of the binding of this protein to genomic regions in a cell-specific context. Third, Tao *et al.* [37] identified *SVP* targets in 9-day-old seedlings grown under LDs. In the current study the vegetative tissue was harvested from *SVP::SVP-GFP svp-41* plants grown for 2 weeks under SDs (see Material and Methods). *SVP* interacting proteins might be expressed differently under these two conditions and therefore affect the capacity and/or selectivity of *SVP* to bind certain genomic regions.

Previously *SVP* was shown to delay flowering by directly repressing transcription of *FT* and *SOC1*, and reducing the mRNA level of the *FT* paralogue *TSF* [6,18,19]. Here, direct binding to *TSF* was not detected suggesting *SVP* might repress its transcription indirectly. *FT* and *TSF* are components of the photoperiodic flowering pathway, while *SOC1* is activated by *FT* in the SAM and acts as a point of convergence of other pathways [75-77]. Analysis of the flowering-time genes present in the high confidence list of *SVP* targets in vegetative tissue detected other genes acting in the photoperiodic flowering pathway or in the circadian clock that acts upstream of it. Notably, *GI* and *PRR7* are targets of *SVP* and both are involved in the photoperiodic induction of flowering and circadian clock regulation [78-80]. Both genes are positive regulators of *CO*, which in turn activates *FT* transcription under long photoperiods. Also the increase in *SVP* protein accumulation in the *lhy cca1* double mutant in continuous light, points to a link between *SVP* regulation and light/clock signaling [55].

The ChIP-seq data suggest that *SVP* likely also affects flowering by other mechanisms. The *FT* gene is a target for *PRC2* and carries the chromatin mark H3K27me3 [81,82]. Therefore the regulation of *PRC2* components by *SVP* may have an indirect effect on *FT* expression.

Mutations in components of PRC2, such as *CLF* that was also identified as a SVP target, cause ectopic expression of MADS-domain proteins that can then promote earlier flowering by mechanisms that remain unclear [83]. Furthermore, PRC2 and other chromatin-related targets of SVP reduce the expression of *FLC* [84], which encodes another MADS-domain protein that is a strong repressor of flowering and physically interacts with SVP [18,55,85]. This complex of FLC and SVP also binds directly to SVP, as discussed later, likely leading to repression of *SVP* transcription. Thus SVP appears to influence flowering time through several pathways that include chromatin regulation and feedback regulation on its own expression, as well as direct binding to genes encoding components of the circadian clock, photoperiodic flowering pathway and floral integrators.

SVP binds to genes involved in hormonal pathways

Our ChIP-seq data revealed numerous putative direct targets of SVP involved in hormonal pathways. SVP binds to genes involved in auxin, GA, cytokinin, and jasmonate homeostasis (Additional data file 3, Table S4). One of these direct targets is *STIP*, a gene involved in the maintenance of the pluripotency and proliferation of meristematic tissue in *Arabidopsis* [86]. Overexpression of *STIP* was shown to partially restore the SAM of the cytokinin insensitive *ahk2-2 ahk3-3 cre1-12* triple mutants, indicating that *STIP* acts downstream of CKs in the establishment of the SAM during early seedling development [49]. Several studies detected a role for cytokinins in the promotion of the floral transition [87]. For instance, the mutant *altered meristem program 1 (amp1)* contains elevated levels of cytokinins and flowers earlier than wild-type plants [88]. Interestingly, the *amp1* mutant rescues the late-flowering phenotype of the *gi* mutant, demonstrating that CK is implicated in the LD pathway downstream of *GI* [50]. Our qRT-PCR experiments showed that *STIP* mRNA is induced in *svp-41* and in *ft-10 tsf-1 svp-41* (Figure 6a, b). This result indicates that SVP represses *STIP* independently or downstream of the two major photoperiod outputs *FT* and *TSF*. In addition, the induction of *STIP* in *svp-41* correlates with increased mRNA expression of several cytokinin response genes, belonging to the *type-A ARR*s and *CRF*s transcription factor families (Figure 6c), in agreement with the proposed role of *STIP* in the CK signaling pathway [49]. Moreover, a significant number of genes deregulated in *svp-41* were also found to be differentially expressed in response to BA (Figure 6d). These results suggest that in the *svp-41* mutant the up-regulation of *STIP* leads to the activation of the CK signaling pathway.

Additional targets of SVP encode hormonal receptors such as *COI1* that may also explain changes in gene expression of signaling components of jasmonate (*JAZ*s

genes). Furthermore the auxin responsive genes SAURs increase in expression in *svp-41* mutants, and these changes may be caused by altered auxin signaling, as SVP binds directly to genes related to auxin transport, such as *BIG* [51]. These effects suggest that the developmental role of SVP is likely to involve complex regulation of hormonal signaling pathways.

Common targets of the dimerizing MADS-box factors

FLC and SVP

MADS-box factors form multimeric complexes that are proposed to be important in determining their DNA binding specificity and thereby their function [15,89]. SVP interacts with FLC and they are proposed to repress flowering as part of a complex that binds to the *SOC1* and *FT* genes [6,18,55,77]. To determine how extensive the overlap in target genes between FLC and SVP is, we compared the vegetative SVP ChIP-seq dataset with the one recently published for FLC [32]. The 112 genes in common between FLC and SVP high confidence targets included *CYTOKININ RESPONSE 1 (CRE1/CHASE)*, supporting a role for both proteins in regulating cytokinin signaling, as discussed above for SVP. However, the ChIP-seq and ChIP-qPCR experiments suggest that SVP and FLC bind to different regions of the gene, with SVP binding in an exon and FLC in the promoter. By contrast, SVP and FLC bound to the same region on the *SVP* promoter suggesting that the heterodimer composed of SVP and FLC could control SVP expression by means of a feedback loop. Taken together this comparison suggests that FLC and SVP do bind to many genes in similar positions, supporting the idea that they often bind to targets as a heterodimer, however some targets appear to be bound by only one of the proteins, indicating that they also have unique targets. Such a conclusion is consistent with the genetic data, which demonstrated that *svp flc* double mutants flower earlier than either single mutant [18,55].

SVP is linked to meristem function during two phases of development

Analysis of the subset of SVP targets that is common to vegetative and reproductive development showed an enrichment of genes involved in meristem function. During vegetative development the SAM continuously produces new cells that sustain plant growth by producing leaves and lateral branches, whereas after its formation the FM enlarges in an undifferentiated state until late stage 2, after which floral organ formation is initiated. *WUS* has a central role in development of both of these stages, participating in the maintenance of the vegetative, inflorescence, and floral meristems [59]. The ChIP-seq analysis showed that SVP binds to regulators of different stages of meristem development and some of these

converge on the regulation of *WUS*. The *WUS* expression domain is restricted to a small group of L3 cells in the center of the meristem by the action of the *CLAVATA* (*CLV*) genes [57]. Our data show that SVP binds *CLV1* in both vegetative and reproductive tissues and *CLV2* in vegetative tissue. Besides the *CLAVATA* pathway, other genes that restrict *WUS* expression, for instance HD-ZIPIII and *SPLAYED* (*SYD*) [58,90] are also targets of SVP. In vegetative tissues SVP binds four of the five HD-ZIPIII genes described in Arabidopsis, *PHB*, *PHV*, *REV*, and *ATHB8*, and during flower development SVP binds *PHB*. Interestingly, we observed that the patterns of expression of *CLV1* and *PHB* become broader in the SAM of *svp-41* mutants compared to Col-0 (Figure 7). These data suggest that SVP influences meristem development by directly binding to genes that act at different levels in the regulatory hierarchy. *SVP* mRNA abundance in the SAM falls as it undergoes conversion from a vegetative to an inflorescence meristem and this correlates with the meristem becoming more domed and increasing in size [18,19]. Reduced activity of SVP in the inflorescence meristem might therefore alter the activity of meristem maintenance pathways to compensate for size differences between the vegetative and inflorescence meristem.

Similarly, floral meristem activity is under control of the MADS-box gene *AG*, which represses *WUS* expression after stage 6 of flower development [91]. SVP and AP1 both repress *AG* expression in the floral meristem, which in turn prevents the repressive activity of *AG* on *WUS*. Interestingly, our data show that SVP control *CLV1* activity since it binds directly to its locus, in the *svp-41 agl24 ap1-12* triple mutant *CLV1* is upregulated (Figure 8a) and the induction of SVP-GR result in the downregulation of *CLV1*; however the pattern of *CLV1* expression is retained (Figure 7 k and l) suggesting a direct role of SVP in the regulation of *CLV1* mRNA quantity, but not in the spatial boundary. Since *CLV1* is also involved in repressing *WUS* activity, the deregulation of *CLV1* could be the cause of the downregulation of *WUS* expression that we detected by *in situ* (Figure 7m, n). Together these data show that SVP and AP1 secure *WUS* expression in the floral meristem via two pathways: the direct repression of *AG* and through direct repression of *CLV1*. This hypothesis is further strengthened by the observation that in the *svp-41 agl24 ap1-12* triple mutant a reduction in floral organ number was observed [25], which is probably due to a decrease in meristem size resulting from increased *CLV1* activity. Indeed the analysis of floral meristem size that we performed in this study revealed that in the triple mutant the FMs are smaller compared to the wild-type (Figure 7q and Table 1) indicating a direct correlation between SVP action and different *WUS* regulatory pathways.

Common targets of AP1 and SVP

SVP together with *AGL24* and AP1 controls floral meristem identity and these proteins are important to prevent early expression of floral homeotic genes, such as *AP3*, *PI*, *SEP3*, and *AG* in the floral meristem [20]. This repression of floral organ identity genes involves recruitment of the LUG-SEU repressor complex by the AP1-SVP heterodimer [25]. As soon as the sepal primordia start to differentiate from the FM *SVP* expression disappears, probably due to interaction between AP1 and *SEP3*, as the latter starts to be expressed during late stage 2 of flower development [92]. Comparison of the gene lists obtained by ChIP-seq experiments for SVP and AP1 [31] identified a significant number of common target genes. Since SVP is strictly expressed in the floral meristem (stages 1 and 2 of flower development), many of these common targets are likely regulated during FM formation rather than specification of floral organ identity. Notably among these common targets transcription factors are enriched. These transcription factors include those involved in meristem maintenance and development. *PHB*, *KANI*, and *ARF3* are all bound by both SVP and AP1 and are upregulated in *svp-41 agl24 ap1-12* inflorescences and the induction of SVP-GR result in the downregulation of *PHB*, *KANI*, and *ARF3* suggesting that SVP modulate their activity. *PHB*, *KANI*, and *ARF3* are involved in the regulation of meristem development and floral organ formation [58,93-95]. Interestingly the activity of ARFs proteins is controlled by convergent auxin flow that is controlled by PIN proteins and SVP and AP1 bound the genomic region of *PIN1*, which is expressed in the IM as well as in the FM. Indeed the expression level of *PIN1* is repressed by SVP. Taken together, these data suggest that there are interactions between the different regulatory networks that control FM formation and differentiation.

Analysis of the *SEP3* ChIP-seq dataset revealed that *CLV1*, *PHB*, *KANI*, and *ARF3* are also bound by *SEP3*, which also interacts with AP1 [15]. The expression profiles of SVP and *SEP3* are mutually exclusive, suggesting a different modulation of the expression of the same target genes by SVP and *SEP3* during floral meristem specification and floral meristem differentiation.

SVP targets are enriched in post-transcriptional and post-translational regulators

Multiple layers of regulation of gene expression play important roles in plant development. Post-transcriptional regulation can enhance and extend the effects of transcriptional regulation. The observation that SVP targets are enriched in genes encoding post-transcriptional and post-translational regulators indicates that SVP may affect gene expression not only by directly binding to target genes and modulating their transcription, but also by indirectly influencing post-transcriptional regulation.

Protein ubiquitination influences the stability and localization of proteins, resulting in the modulation of their biological functions. Defects in ubiquitination pathways can result in abnormal floral organ identity as suggested by the functional analyses of the *DCAF1* and *CYP71* genes, which are part of Cullin-RING ubiquitin ligase complexes [67,96].

SVP binds to a large number of DCAF encoding genes in FMs suggesting that SVP could be involved in the control of both proteasome and epigenetically mediated regulation of floral processes (Additional data file 1, Table S9). Several SVP targets are linked to chromatin-mediated regulation, such as two uncharacterized WD40 proteins containing Bromodomains, known to bind acetylated lysine residues in histones [97]. Thus SVP likely controls developmental processes by regulating gene expression directly through transcriptional regulation and indirectly by modulating transcription of genes encoding post-transcriptional and post-translational regulators.

It was recently reported that the WDR protein WDR55 is a putative DCAF and may function in a CUL4 - DDB1^{WDR55} E3 ligase complex [70]. Interestingly we discovered that *WDR55* is a target of SVP, which bound its genomic locus in inflorescence tissues. Moreover *WDR55* results downregulated in *svp-41 agl24* and *svp-41 agl24 ap1-12* compared to the wild-type inflorescences indicating that SVP acts as a direct activator of *WDR55* expression in the floral meristem.

The role of WDR55 in floral organ ontogenesis

The analyses of the mutant *wdr-55-2* showed variable phenotype in flower development such as reduced number of organs, asymmetric and reduced sepal and petal size, and occasionally chimeric organs such as petaloid stamens and carpelloid stamen or sepals. *In-situ* hybridization analysis revealed that *AG* was misexpressed in the *wdr55-2* flower. In wild-type, *AG* expression is always restricted to the two inner whorls (whorls 3 and 4). In homozygous *wdr55-2* mutant flowers *AG* expression is detectable earlier than in wild-type and in all floral whorls. This strongly suggests that WDR55 is involved in both spatial and temporal regulation of *AG*. The SVP-AP1 heterodimer is thought to recruit LUG-SEU and regulate *AG* expression in early stages of flower development [25]. We tested if WDR55 could bind any of these proteins but were not able to show any interaction.

Taken together the overall data indicate that SVP repress *AG* expression through two different pathways, the first is via the interaction with the co-repressor complex containing LUG-SEU and the dimer SVP-AP1 [25] and the second by SVP controlling the expression level of *WDR55*. The floral phenotype of the *wdr55-2*

mutant is variable and did not result in the deregulation of *AG* in all the flowers, this suggests that SVP in the *wdr55-2* background is, although less efficient, still able to repress *AG* directly probably via the LUG-SEU pathway.

Conclusions

In summary, our data indicate that the SVP genome-wide binding profiles during two distinct developmental stages show a significant overlap and that this subset of genes includes a wider set of important regulators of plant development than was previously realized. However, there is also a large group of SVP target genes that are not bound at both stages, clearly reflecting distinct functions during vegetative and reproductive phases. The specificity of SVP binding to DNA is probably influenced by interaction with different MADS-domain partners, such as FLC and AP1. A related observation was made for the *Drosophila* MADS domain protein MEF2 that is expressed widely during development, but has specific targets at different stages dependent on the presence of interacting transcription factors [98]. The presented data provide new insights into the enormous diversity of pathways that are regulated by SVP and forms a basis for detailed analysis of the roles of SVP in regulating specific genes and pathways in combination with different interacting proteins.

Materials and methods

Plant material and growth conditions

For ChIP and microarray analysis of vegetative phase, *SVP::SVP-GFP*, *svp-41* single mutant (for plasmid construction see [20]) and wild-type seedlings were grown 14 days under short-day (SD) conditions (8 h light/16 h dark) at 22°C. For ChIP and microarray analysis of the reproductive phase, *SVP::SVP-GFP svp-41*, triple mutant *svp-41 agl24-2 ap1-12* and wild-type plants were grown under long-day (LD) conditions (LD; 16 h light/8 h dark) at 22°C. For the GR induction study the triple mutant *svp-41 agl24-2 ap1-10* was used [24]. All the plants were from the same Columbia ecotype. The *SVP::SVP-GFP svp-41* transgenic line and triple mutant *svp-41 agl24-2 ap1-12* have been previously described [20,25]. *ft-10 tsf-1 svp-41* and *ft-10 tsf-1* were described previously in Jang *et al.* [19]. The *wdr55-2* (WiscD-Lox430F06) line is in the Col-0 ecotype and is a T-DNA insertion mutant obtained from the Nottingham Arabidopsis Stock Centre [99]. Seeds were surface sterilized using EtOH, bleach and Tween20 before germinated on MS media [100] supplemented with 2% sucrose (MS-2) and glufosinate-ammonium for BASTA selection of *wdr55-2* plants. All seeds were stratified on MS-2 plates at 4°C O.N. before being transferred to 18°C for

about 12 days until germination. The seedlings were eventually transferred to soil and grown at 18°C under LD conditions (16 h).

ChIP assays

For ChIP experiments, the commercial antibody GFP: Living Colors_ full-length A.v. polyclonal antibody was used (Clontech [101]). Chromatin was prepared from inflorescences (2 weeks after bolting) and from 14-day-old seedlings of *svp*, grown under SD conditions. Wild-type plants (inflorescences and seedlings) were used as negative controls. ChIP assays were performed as previously described by [20] and in Additional data file 1, Methods S1 with a minor modification in the sonication step. DNA samples were sonicated six times 30 s each with amplitude 30 to 40, with intervals of 1 min (100-500 bp range fragments obtained).

We used as a positive control for the ChIP in the reproductive phase a region of the *AG* second intron (*AG.V*) that previously has been demonstrated to bind SVP-GFP [20]. For the vegetative phase we used regions in *FT* bound by SVP [18] (Additional data file 1, Figure S1). Enrichment fold to evaluate the quality of each ChIP sample was tested by qRT-PCR as described in Additional data file 1, Methods S2, all the primers used for ChIP-qPCR are in Additional data file 1, Table S12).

Sample preparation for ChIP-seq Illumina/Solexa sequencing

Two independent ChIP experiments (enrichment fold controlled by real-time PCR) were used for vegetative and reproductive ChIP-seq assays, respectively. We used one ChIP DNA sample for each library preparation and these were run on the Genome Analyzer. The DNA quantification of immunoprecipitated DNA was performed with the Quant-iT dsDNA HS Assay Kit (Invitrogen). Libraries for Solexa sequencing were prepared following the Illumina kit protocol, with some modifications. The first step 'Perform End Repair' was repeated twice, adding fresh enzymes and incubating 1 h longer than indicated by the protocol. Two units of undiluted Klenow enzyme was used. The incubation time of the step 'Ligate adapters to DNA fragments' was prolonged to 1 h instead of 15 min. Each library was validated quantifying the DNA with Quant-iT dsDNA HS Assay Kit (Invitrogen).

Read mapping and identification of enriched regions

Sequence reads were mapped to the unmasked Arabidopsis genome (TAIR8 build) using the Seqmap tool [102], allowing at most two mismatches at any position. Trimming unmapped reads at the 5' or 3' end led to marginal improvements in the number of reads mapped, and this step was therefore skipped. Reads belonging to

duplicate experiments in each of the three conditions were pooled together. Only reads mapping to a unique position on the genome were considered for further analysis. This resulted in about 3 million uniquely mapped reads for the two inflorescences experiments, 5 million for seedlings experiments, and 6 million for control experiments. In each experiment, uniquely mapped reads were extended by 300 bps along the 5'→3' direction. This resulted in a base pair by base pair coverage map of the genome, that is, giving for each base pair the number of extended sequence reads that contained it. Only base pairs covered by reads mapping on both strands were considered valid for further analysis. Enrichment was then calculated in each valid base pair by comparing, for each IP experiment, the coverage in the experiment to the coverage in the control used as expected value, and computing an enrichment *P* value with a negative binomial distribution. In each comparison, the coverage of the two samples was normalized according to the number of reads obtained in each. Enriched regions were then defined as regions consisting of consecutive base pairs characterized by calculated *P* values <0.01 and not interrupted by a gap of 100 or more base pairs that were either non-valid or with a *P* value >0.01. The *P* value associated with each of these regions was defined as the minimum *P* value among the base pairs belonging to the region. Regions <150 bps were then discarded regardless of the *P* value. The number of remaining candidate-enriched regions was finally used to compute a Bonferroni corrected *P* value to be associated to the regions themselves. The overall strategy we followed in our analysis for the identification of enriched regions is highly similar to the one adopted in the SEP3 and AP1 ChIP-Seq experiments [13,31] and in the CSAR peak-finding tool [41], which has been shown to be better suited for ChIP-Seq experiments in Arabidopsis. *P* values for enrichment were computed by using a negative binomial distribution instead of the Poisson, as the former provides a better fit to count data from ChIP-Seq experiments [103]. Also, we employed a more conservative Bonferroni correction for multiple testing aimed at minimizing the number of false positive predictions.

Starting from regions with corrected *P* values <0.01, potential target genes were then identified by associating with each gene an overall *P* value given by the product of the *P* values associated with the single binding regions located in its gene locus, from 3 kbps upstream of the transcription start site to 1 kbp downstream of the transcribed region. Protocols of ChIP, DNA extraction, sequencing preparation, data processing, and all the associated files to this study can be found in the GEO (Gene Expression Omnibus) database (ID: GSE33120).

Tiling array experiments

The vegetative tissue samples were obtained from aerial parts of the *svp-41* single mutant and wild-type seedlings grown for 2 weeks under SD conditions (8 h light/16 h dark) and harvested at zeitgeber 8 (ZT8). For the reproductive tissue sampling we used wild-type and *svp-41 agl24-2 ap1-12* triple mutant inflorescences grown for 2 weeks under SD conditions and then moved to LD conditions (16 h light/8 h dark). The inflorescences were collected at 2 weeks after bolting at ZT8. RNA from three independent biological replicates was extracted using the RNA Plant Mini kit, QIAGEN (www1.qiagen.com/) and quantified by NanoDrop; 1 µg of total RNA was reverse transcribed into cDNA using an oligo(dT)-T7 primer, and was then converted into cRNA and linearly amplified by T7 *in-vitro* transcription reaction using the standard Ambion protocol (MessageAmp aRNA Kit, Ambion). cRNA was then reverse transcribed with random primers to dUTP-containing ds cDNA (WT ds cDNA Synthesis Kit, catalog no. 900813; Affymetrix). Fragmentation and labeling was performed with the GeneChip WT double-stranded DNA Terminal Labeling Kit (catalog no. 900812, Affymetrix). After fragmentation, 7.5 µg of ds-cDNA was hybridized for 16 h at 45°C on GeneChip Arabidopsis Tiling 1.0R Array. GeneChips were washed and stained with Fluidics Script FS450_0001 in the Affymetrix Fluidics Station 450. Then, the GeneChips were scanned using the GeneChip Scanner 3000 7G. Data were processed in R as described in [104]. Probe-level data were pre-processed using the RMA algorithm implemented in the Bioconductor package Affy. Linear models and empirical Bayes methods from the Limma package of Bioconductor were applied to derive a *P* value, false discovery rate (FDR; *P* adjusted), and mean of log₂-based ratio across replicates. The data were deposited in the GEO (Gene Expression Omnibus) database (ID: GSE32397).

Gene Ontology analysis

The Bingo 2.44 plug-in [105] implemented in Cytoscape v2.81 [106] was used to determine and visualize the GO enrichment according to the GOSlim categorization. A hypergeometric distribution statistical testing method was applied to determinate the enriched genes and the Benjamini and Hochberg FDR correction was performed in order to limit the number of false positives. The FDR was set up to 0.001 and 0.05 for the ChIP-seq and expression data, respectively. In addition to Bingo 2.44, further GO annotation analysis of the targets of SVP was performed by using TAIR bioinformatics resources [107].

cDNA preparation and qRT-PCR analysis

Expression analyses in the vegetative phase was performed using the *svp-41* single mutant, *35S::SVP* and

wild-type seedlings grown for 2 weeks under SD conditions; for the reproductive phase we used wild-type and *svp-41 agl24-2 ap1-12* triple mutant inflorescences grown for 2 weeks under SD conditions and then moved to LD conditions. The inflorescences were collected at 2 weeks after bolting.

Total RNA from three biological replicates was extracted with the LiCl method, and its integrity was checked on agarose gels. The samples were treated with DNase (TURBO DNA-free; Ambion [108]) and reverse transcribed according to the ImProm-II_ Reverse Transcription System (Promega [109]) instructions. Sequence primers for RT-PCR amplification are listed in Additional data file 1, Table S13. Ten-fold dilutions of cDNA were tested in RT-PCR and qRT-PCR experiments using reference genes.

Enrichment folds were detected using a SYBR Green assay (Bio-Rad [110]). The real-time PCR assay was performed in triplicate using a Bio-Rad C1000 Thermal Cycler optical system or LightCycler480 (ROCHE) thermal cycler. For expression analyses normalized expression was calculated using the delta-delta Ct method (DDC(t)). For ChIP experiments, relative enrichment was calculated as described in Additional data file 1, Methods S2. For the expression analysis ubiquitin, PEX4, and PP2a-F were used as reference genes.

In-situ hybridization

In-situ hybridization has been performed as described in Additional data file 1, Method S3. The *WUS* antisense probe has been cloned according to Brambilla *et al.* [111]. The *ARF3* antisense probe has been cloned in the pGEM-T easy using the primers FW-CCCATCTGTATCATCATCACC and REV- CTCTCATTGCATAGATGTCC. The *KANI* antisense probe has been cloned in the pGEM-T easy using the primers FW- AAGACCACTAA-CAAGCCTGC and REV- CATTCTCGTGCCAATC TGGTC. The *CLV1* antisense probe has been cloned according to Clark *et al.* [60]. The *PHB* antisense probe has been cloned in the pGEM-T easy using the primers FW-GGTAGCGATGGTGCAGAGG and REV- CGAAC-GACCAATTCACGAAC. Sections were observed using a Zeiss Axiophot D1 microscope (Zeiss [112]) equipped with differential interface contrast (DIC) optics. Images were captured on an Axiocam MRc5 camera (Zeiss) using the AXIOVISION program (version 4.4).

Scanning electron microscopy

SEM has been performed as described in Additional data file 1, Method S4.

Inducible expression experiments

The *p35S::SVP-GR* construct was produced as follows: the coding region of *SVP* was amplified from inflorescence

cDNA using primers Fw-CGTTGCCATGGCGAGAGAA AAGAT and Rev- ATTGTTCGGATCCCCACCACCA-TACGG containing NcoI and BamHI sites, respectively, cloned into pGEM-T easy (Promega), digested with NcoI and BamHI and ligated into pBluescript SK (Stratagene) containing a portion of the rat glucocorticoid hormone binding domain (a.a 508-795 [61]) to produce *pSK-SVP-GR*. The *AG-GR* fragment was amplified from the *pSK-SVP-GR* using the primers For and Rev and subcloned into the pTOPO vector (Life Technology). Finally *SVP-GR* was subcloned into the Gateway destination vector pB2GW7.0 [113] containing the 35S promoter. *p35S::SVP-GR* was transformed in *svp-41 agl24-2 ap1-10* background (*ap1-10* heterozygous) and the T1 generation was selected for BASTA resistance.

After bolting, inflorescences of *35S::SVP-GR svp-41 agl24-1 ap1-10* plants were treated with a solution containing 10 μ M dexamethasone (Sigma-Aldrich), 0.01% (v/v) ethanol, and 0.015% (v/v) Silwet L-77. Mock treatment consist of 0.01% (v/v) ethanol, and 0.015% (v/v) Silwet L-77.

For each time point, tissue from eight plants was collected. Tissue was removed as close to the surface of the inflorescence as possible to ensure an enrichment of FM cells.

Appendix

Accession numbers

Arabidopsis Genome Initiative locus identifiers for the genes mentioned in this article are as follows: *AGL24* [TAIR:AT4G24540], *STK* [TAIR:AT4G09960], *AP3* [TAIR:AT3G54340], *FLC* [TAIR:AT5G10140], *SVP* [TAIR:AT2G22540], *JAZ6* [TAIR:AT1G72450], *AGL16* [TAIR:AT3G57230], *SOCI* [TAIR:AT2G45660], *CLV1* [TAIR:AT1G75820], *PINI* [TAIR:AT1G73590], *ARF3/ETT* [TAIR:AT2G33860], *KAN1* [TAIR:AT5G16560], *PHB* [TAIR:AT2G34710], *JAZ7* [TAIR:AT2G34600], *SADHU* [TAIR:AT3G42658], *JAZ8* [TAIR:AT1G30135], *GA2ox6* [TAIR:AT1G02400], *ARR6* [TAIR:AT5G62920], *ARR7* [TAIR:AT1G19050], *DDF1* [TAIR:AT1G12610], *GA2ox2* [TAIR:AT1G30040], *miR167* [TAIR:AT1G31173], *ACD6* [TAIR:AT4G14400], *API1* [TAIR:AT1G69120], *WDR55* [TAIR:AT2G34260], *VRN2* [TAIR: AT4G16845], *CLF* [TAIR: AT2G23380], *SWN* [TAIR: AT4G02020], *GI* [TAIR: AT1G22770], *FLK* [TAIR: AT3G04610], *FLD* [TAIR: AT3G10390], *PRR7* [TAIR: AT5G02810], *PHYA* [TAIR: AT1G09570], *STIP* [TAIR: AT2G33880], *ARR11* [TAIR: AT1G67710], *ARR5* [TAIR: AT3G48100], *ARR15* [TAIR: AT1G74890], *CRF2* [TAIR: AT4G23750], *CRF5* [TAIR: AT2G46310], *PHV* [TAIR: AT1G30490], *REV* [TAIR: AT5G60690], *ATHB8* [TAIR: AT4G32880], *ATBARD1* [TAIR: AT1G04020], *KAN2* [TAIR: AT1G32240], *LMI1* [TAIR: AT5G03790], *DCAF1* [TAIR:

AT4G31160], *JAZ5* [TAIR: AT1G17380], *JAZ10* [TAIR: AT5G13220], *JAZI* [TAIR: AT1G19180]

Additional material

Additional data file 1: contains: **Figure S1:** Analysis of chromatin sample used for ChIP-seq experiments. **Figure S2:** qRT-PCR validation of differentially expressed genes between Col-0 and *svp-41* plants at the vegetative phase. **Figure S3:** GO enrichment analysis of differentially expressed genes between Col-0 and *svp-41* plants at the vegetative stage. **Figure S4:** Venn diagram containing the overlapping set of putative targets between SVP and FLC and SVP, AP1, and SEP3. **Figure S5:** Biologically active SVP-GR fusion. **Figure S6:** Flower organs of in *wdr55-2 -/-* mutants show reduced size and asymmetric positioning. **Figure S7:** *In-situ* hybridization of wild-type and *wdr55-2* inflorescence using *AP3* and *Pl* probes. **Figure S8:** Flower morphology of *wdr55-2 ag-1* mutant. **Table S1:** Summary of sequencing and mapping. **Table S3:** List of putative targets of SVP related to flowering time. **Table S6:** List of genes differentially expressed in *svp-41* compare to *Col-0* and related to auxin, cytokinin, or jasmonate homeostasis. **Table S9:** List of WDxR motif containing proteins found in SVP DNA binding screen. **Table S10:** Flower organ count from *wdr55-2 -/-* mutants. **Table S12:** Primer pairs used for ChIP-qPCR assays. **Table S13:** Primer pairs used for the qRT-PCR expression analysis. **Methods S1:** ChIP protocol. **Methods S2:** qRT-PCR. **Methods S3:** *In-situ* hybridization. **Methods S4:** Scanning electron microscopy.

Additional data file 2: contains Table S2: High confidence targets of SVP in vegetative and reproductive tissues; list of the targets of SVP bound in both vegetative and reproductive tissues; lists of binding regions of SVP in vegetative and reproductive tissues.

Additional data file 3: contains Table S4: Lists of putative SVP targets with annotated functions in: meristem development in vegetative and reproductive tissues; response to hormonal stimuli such as auxin, cytokinin, ethylene, abscisic acid, jasmonate, and gibberellins in vegetative tissue.

Additional data file 4: contains Table S5: Tiling array expression data obtained using RNA extracted from: wild-type Col-0 and *svp-41* plants at the vegetative stage, inflorescences of wild-type Col-0 and *svp-41 agl24 ap1-12* and overlap between tiling array and ChIP-seq data.

Additional data file 5: contains Table S7: Lists of differentially expressed genes in *svp-41* mutant and the available expression-profiling data of seedlings treated with the CK benzyladenine (BA).

Additional data file 6: contains Table S8: putative targets for both SVP and AP1 and putative targets for both SVP and SEP3.

Additional data file 7: contains Table S8: Comparison of the SVP target list of Tao et al. [37] and the list of high confidence targets of SVP in vegetative tissue presented in this study.

Abbreviations

CARG-boxes: MAD5-domain factors binding consensus; ChIP-qPCR: chromatin immunoprecipitation followed by quantitative real-time polymerase chain reaction; ChIP-seq: chromatin immunoprecipitation combined with high throughput DNA sequencing; ChIP: chromatin immunoprecipitation; CK: cytokinin; CSAR: ChIP-seq analysis in R; FM: floral meristem; GA: gibberellin; GEO: Gene Expression Omnibus; GO: Gene Ontology; GR: glucocorticoid receptor; IM: inflorescence meristem; qRT-PCR: quantitative real-time polymerase chain reaction; SAM: shoot apical meristem; Y2H: Yeast-2-Hybrid

Authors' contributions

VG did the ChIP experiments, biological analysis, and writing; FA did the transcriptome analysis, biological analysis, and writing; AS did the ChIP-seq and writing; RFG, SS, JLM, ST, and KNB performed the biological analysis; FZ and GMP were responsible for bioinformatics; GP did the bioinformatics and writing; LC conducted the research design; PEG, GC, and MMK did the

research design and writing. All authors read and approved the final manuscript.

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