RESEARCH ARTICLE

Mutagenesis of a Quintuple Mutant Impaired in Environmental Responses Reveals Roles for CHROMATIN REMODELING4 in the Arabidopsis Floral Transition

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Short Title: Role of CHR4 in the floral transition

One Sentence Summary: A genetic screen employed to identify genes that regulate flowering independently of environmental cues revealed a role for the chromatin remodeler CHR4 in promoting floral identity.

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ABSTRACT

Several pathways conferring environmental flowering responses in Arabidopsis thaliana converge on developmental processes that mediate the floral transition in the shoot apical meristem. Many characterized mutations disrupt these environmental responses, but downstream developmental processes have been more refractory to mutagenesis. Here, we constructed a quintuple mutant impaired in several environmental pathways and showed that it possesses severely reduced flowering responses to changes in photoperiod and ambient temperature. RNA-seq analysis of the quintuple mutant showed that the expression of genes encoding gibberellin biosynthesis enzymes and transcription factors involved in the age pathway correlates with flowering. Mutagenesis of the quintuple mutant generated two late-flowering mutants, quintuple ems 1 (qem1) and qem2. The mutated genes were identified by isogenic mapping and transgenic complementation. The qem1 mutant is an allele of the gibberellin 20-oxidase gene ga20ox2, confirming the importance of gibberellin for flowering in the absence of environmental responses. By contrast, qem2 is impaired in CHROMATIN REMODELING4 (CHR4), which has not been genetically implicated in floral induction. Using co-immunoprecipitation, RNA-seq and ChIP-seq, we show that CHR4 interacts with transcription factors involved in floral meristem identity and affects the expression of key floral regulators. Therefore,
CHR4 mediates the response to endogenous flowering pathways in the inflorescence meristem to promote floral identity.

INTRODUCTION

Lateral shoot organs initiate from cells on the flanks of the shoot apical meristem (SAM), and the identity of the formed organs changes during development (Bowman and Eshed, 2000). In Arabidopsis thaliana, the transition from vegetative leaf initiation to flower production occurs in response to several environmental and endogenous cues. Important environmental signals that control flowering include seasonal fluctuations in temperature and day length, which are mediated by the photoperiodic and vernalization pathways, whereas ambient changes in temperature also influence flowering time (Srikanth and Schmid, 2011; Andres and Coupland, 2012). In addition, endogenous signals such as gibberellins (GAs) and the age of the plant contribute to the floral transition in the absence of inductive environmental cues (Wilson et al., 1992; Wang et al., 2009).

Three intersecting environmental pathways that promote flowering have been well characterized. The photoperiodic pathway promotes flowering under long days (LDs) but not under short days (SDs), in which plants flower much later. Exposure to LDs stabilizes the CONSTANS transcription factor (Valverde et al., 2004), which in turn activates transcription of FLOWERING LOCUS T (FT) and TWIN SISTER OF FT (TSF) in the leaf vascular tissue (Kardailsky et al., 1999; Kobayashi et al., 1999; Suarez-Lopez et al., 2001; An et al., 2004; Yamaguchi et al., 2005). The FT and TSF proteins, which are related to phophatidyl-ethanolamine binding proteins, move to the SAM (Corbesier et al., 2007; Jaeger and Wigge, 2007; Mathieu et al., 2007), where they physically interact with the bZIP transcription factor FD (Abe et al., 2005; Wigge et al., 2005; Abe et al., 2019). In the SAM, the FT–FD protein complex promotes the transcription of genes encoding floral activators, such as SUPPRESSOR OF OVEREXPRESSION OF CO1 (SOC1) and FRUITFULL (FUL), which induce the floral transition, as well as APETALA1 (AP1) and LEAFY (LFY), which promote floral meristem identity (Schmid et al., 2003; Wigge et al., 2005; Torti et al., 2012; Collani et al., 2019). Because they represent the mobile signal linking leaves and the shoot apical meristem, FT and TSF are essential for the photoperiodic flowering response, and ft tsf double mutants are daylength-insensitive (Yamaguchi et al., 2005; Jang et al., 2009).
The seasonal cue of exposure to winter cold mediates flowering via the vernalization pathway, which represses transcription of the floral repressor FLOWERING LOCUS C (FLC) (Michaels and Amasino, 1999; Sheldon et al., 1999). FLC is a MADS-box transcription factor that forms regulatory complexes with other MADS-box floral repressors, such as SHORT VEGETATIVE PHASE (SVP) (Li et al., 2008). Thus, vernalization reduces FLC transcription and promotes flowering via the endogenous and photoperiodic pathways, whereas mutants for FLC are essentially insensitive to vernalization. The genome-wide binding sites of FLC and SVP include those in several genes that promote flowering within the photoperiodic pathway, such as FT and SOC1 (Searle et al., 2006; Lee et al., 2007; Li et al., 2008; Deng et al., 2011; Mateos et al., 2015; Richter et al., 2019). Because FLC is stably repressed by exposure to cold, plants can flower through the photoperiodic pathway when they are exposed to LDs after cold exposure. Also, genes within the endogenous pathway that are repressed by FLC, such as SQUAMOSA PROMOTER BINDING PROTEIN-LIKE15 (SPL15), can promote flowering during vernalization (Deng et al., 2011; Hyun et al., 2019).

Arabidopsis also flowers rapidly when exposed to high temperatures, and this response can overcome the delay in flowering observed under SDs at lower growth temperatures (Balasubramanian et al., 2006). FT and TSF are transcribed at high temperature under SDs and promote early flowering; thus their transcriptional repression under SDs at lower temperatures is overcome at high temperatures (Kumar et al., 2012; Galvao et al., 2015; Fernandez et al., 2016). Accordingly, MADS-box repressors of FT and TSF, particularly FLOWERING LOCUS M and SVP, do not accumulate under SDs at high temperature, and mutations in these genes reduce the flowering response to high temperature (Lee et al., 2007; Lee et al., 2013; Pose et al., 2013; Airoldi et al., 2015). The reduced activity of these repressors also enhances the response of the meristem to low levels of FT and TSF transcription in the leaves (Fernandez et al., 2016). Triple mutants for FT, TSF and SVP are insensitive to higher temperatures under SDs (Fernandez et al., 2016).

In addition to these environmental pathways, there are several endogenous flowering pathways. A set of genes was ascribed to the autonomous flowering pathway, because they caused late-flowering under LDs and SDs and were therefore considered to promote flowering independently of photoperiodic cues (Koornneef et al., 1991). Mutations in all these genes caused elevated levels of FLC mRNA, and
the encoded proteins contribute to \textit{FLC} expression at the transcriptional and post-transcriptional levels (Whittaker and Dean, 2017). The late-flowering phenotype of autonomous pathway mutants can therefore be suppressed by mutations in \textit{FLC} (Michaels and Amasino, 2001). In addition, gibberellin (GA) is an important contributor to endogenous flowering regulation, because mutations or transgenes that strongly reduce GA levels almost abolished flowering under non-inductive SDs (Wilson et al., 1992; Galvao et al., 2012; Porri et al., 2012). Finally, microRNA156 (miR156) negatively regulates the floral transition and is developmentally regulated such that its abundance decreases progressively with increasing plant age (Wu and Poethig, 2006; Wang et al., 2009). This miRNA negatively regulates the accumulation of several SPL transcription factors, including SPL3, SPL9 and SPL15, which promote the floral transition, particularly under non-inductive SDs (Gandikota et al., 2007; Wang et al., 2009; Yamaguchi et al., 2009; Hyun et al., 2016; Xu et al., 2016). Thus, miR156/SPL modules have been associated with an endogenous flowering pathway related to plant age.

Here, we extend our understanding of the genetic basis of the floral transition by screening specifically for genes that regulate flowering independently of the environmental pathways. To this end, we constructed a high-order quintuple mutant, \textit{svp-41 ftc-3 ft-10 tsf-1 soc1-2}, which shows reduced sensitivity to environmental flowering signals because it is impaired in responses to photoperiod and high temperature. Using RNA-seq, we characterized the expression of flowering-related genes in this mutant, and we employed a forward genetics approach to identify genes controlling flowering time in this background. This allowed us to define a role for CHROMATIN REMODELING4 (CHR4) in promoting the floral transition.

\textbf{RESULTS}

\textbf{Phenotypic and molecular characterization of a quintuple mutant strongly impaired in responses to environmental cues}

To assess the flowering time of Arabidopsis plants in which the major environmental pathways were inactivated, we constructed the quintuple mutant \textit{svp-41 ftc-3 ft-10 tsf-1 soc1-2} (hereafter referred to as the quintuple mutant). The quintuple mutant showed a dramatically reduced flowering response to day length compared to Col-0. Under long days (LDs), the quintuple mutant bolted later and after forming more vegetative rosette leaves than the wild type (Col-0) (\textbf{Figure 1A and B}). However,
under short days (SDs) at 21°C, the mutant bolted much earlier than Col-0 in terms of days to flowering and rosette leaf number (Figure 1A and B). Bolting of the quintuple mutant was delayed by fewer than 10 days in SDs compared to LDs, whereas bolting in Col-0 was delayed by approximately 50 days. Similarly, the quintuple mutant formed ~5 more rosette leaves under SDs than LDs, whereas Col-0 formed over 40 more rosette leaves. The flowering time of the quintuple mutant was also insensitive to higher ambient temperatures under SDs when considering bolting time, but it displayed partial insensitivity in terms of rosette leaf number (Figure 1A and B). Finally, GA₄ treatment accelerated flowering of Col-0 under SDs (Wilson et al., 1992) but had a smaller effect on the flowering time of the quintuple mutant (Supplemental Figure 1A and B). These results are consistent with the idea that the GA response and signaling are activated in the quintuple mutant, as previously shown for svp-41 mutants (Andres et al., 2014). Overall, the quintuple mutant showed strongly impaired responses to environmental signals such as day-length and ambient temperatures, in terms of time to bolting and the number of rosette leaves formed. These data suggest that in the quintuple mutant, the floral transition occurs via endogenous mechanisms such as the GA or age pathway.

In addition to effects on bolting time and vegetative rosette leaf number, the quintuple mutant produced more cauline leaves than Col-0 in all conditions tested (Supplemental Figure 1C). The quintuple mutant formed on average 4.5-fold more cauline leaves than Col-0 under LDs and 2.3-fold more under SDs. The increased cauline leaf number in the mutant compared with Col-0 suggests that the mutant is also impaired in the determination of floral meristem identity after floral induction and bolting, such that more phytomers contain cauline leaves and axillary shoots than in Col-0.

We then compared the developmental stage of the shoot apex of the quintuple mutant to that of Col-0 by performing in situ hybridizations for FUL transcript on apical cross-sections of SD-grown plants of different ages (Figure 1C). FUL encodes a MADS-box floral activator that is partially genetically redundant with SOC1. FUL mRNA accumulates in the SAM during the early stages of the floral transition (Ferrandiz et al., 2000; Melzer et al., 2008; Torti et al., 2012). In the apices of SD-grown plants, FUL mRNA accumulated approximately one-week earlier in the quintuple mutant than in Col-0 (Figure 1C), which is consistent with the earlier flowering phenotype of the mutant.
Because the quintuple mutant flowers earlier under SDs and major regulators of flowering are inactivated, the transcriptional network associated with the floral transition is probably differentially expressed in the mutant compared to Col-0. To define these differences, we performed RNA-seq on apices of the quintuple mutant and Col-0 through a developmental time course under SDs. Apical samples were harvested from both genotypes 3, 4, 5 and 6 weeks after sowing. In vegetative apices of both genotypes at 3 weeks after sowing, only 46 genes were differentially expressed (adjp-value < 0.05) (DEGs) between the quintuple mutant and Col-0. At 4, 5 and 6 weeks, when the mutant flowered more rapidly than Col-0 (Figure 1C), 486, 736 and 568 genes, respectively, were differentially expressed in the mutant compared with Col-0 (Supplemental Data Set 1). At these time points, approximately 45%, 14% and 33% of the DEGs, respectively, were more highly expressed in the quintuple mutant vs. Col-0 (Supplemental Data Set 1). The mRNAs of SPL3, SPL4, SPL5, SPL9, SPL12 and SPL15, which are regulated by miR156 and contribute to the endogenous age-related flowering pathway, were more abundant in the quintuple mutant, which is consistent with promotion of flowering by the age pathway (Figure 1D and E). Moreover, the floral activators FD, FDP and AGAMOUS-LIKE6 (AGL6) were more highly expressed in the mutant vs. Col-0 (Figure 1D and E, Supplemental Figure 1D and Supplemental Data Set 1), and the expression of the floral repressors MADS-AFFECTING FLOWERING 4 (MAF4) and MAF5 was attenuated in the quintuple mutant (Supplemental Data Set 1). Moreover, genes encoding enzymes involved in GA biosynthesis and catabolism were differentially expressed in the quintuple mutant (Figure 1D).

A sensitized mutant screen in the quintuple mutant background identifies two loci that promote flowering
We then employed the quintuple mutant as a sensitized background for mutagenesis screening to identify genes that regulate flowering independently of environmental pathways. This approach was expected to identify mutations in endogenous components, because the major environmental floral response pathways are already impaired in the mutant, and mutations in the autonomous pathway should not be recovered, because FLC is inactive in the quintuple mutant. We screened the M$_2$ generation for mutants with altered flowering behaviour (Methods). Two mutants showing delayed floral transition in the quintuple mutant background, quintuple ems 1
(qem1) and quintuple ems 2 (qem2), were selected for detailed studies because they exhibited strong and reproducible phenotypes in the M₃ generation. Both lines segregated the mutant phenotype in a 3:1 ratio in the BC1F₂ generation (Methods), suggesting that a single recessive mutation was responsible for the phenotypes of both mutants. Plants segregating the qem1 or qem2 phenotype in the respective BC1F₂ populations were then bulk-harvested. Fast-isogenic mapping (Methods) (Hartwig et al., 2012) localized qem1 and qem2 with high confidence to different regions on chromosome 5 (Supplemental Figure 2 and Figure 2).

The qem1 mutation localized to the same region of chromosome 5 as the gibberellin 20-oxidase gene GA20ox2 (Supplemental Figure 2C and Supplemental Table 1). Mutation of GA20ox2 delays flowering and has a stronger effect in the svp-41 background (Rieu et al., 2008; Plackett et al., 2012; Andres et al., 2014). In qem1, a single nucleotide polymorphism was identified in the first exon of GA20ox2 that was predicted to cause an amino-acid substitution in the protein (ser137asn). To confirm that this mutation causes the late-flowering phenotype of qem1, we performed molecular complementation. Introducing the Col-0 genomic GA20ox2 locus into qem1 strongly reduced leaf number and flowering time, so that the transgenic lines flowered at a similar time or earlier than the quintuple mutant (Supplemental Figure 2D and E), confirming that the mutation in GA20ox2 was responsible for the delayed flowering of qem1. This result is consistent with the RNA-seq data showing that GA20ox2 mRNA is more highly expressed in the quintuple mutant background than in Col-0 (Supplemental Figure 1D and with the previous observation that svp-41 mutants contain higher levels of bioactive GAs than the wild type (Andres et al., 2014). Therefore, the GA pathway likely plays a decisive role in promoting the floral transition in the quintuple mutant.

The qem2 mutant was later flowering and initiated more rosette and cauline leaves than the quintuple mutant (Figure 2A and B), indicating a delay in the floral transition and impaired floral meristem identity. The region of chromosome 5 to which qem2 mapped contained no previously described flowering-time genes (Figure 2C and Table 1). Three high-confidence polymorphisms predicted to cause non-synonymous mutations in the coding sequences At5g43450, At5g44690 and At5g44800 were identified (Table 1). At5g43450 encodes a protein with similarity to ACC oxidase, At5g44690 encodes a protein of unknown function, and At5g44800 encodes the CHD3-like ATP-dependent chromatin-remodelling factor CHR4. In
Arabidopsis, CHR4 is most closely related to PICKLE (PKL), which represses flowering via the GA pathway (Fu et al., 2016; Park et al., 2017) and promotes flowering via the photoperiodic pathway though FT activation (Jing et al., 2019a; Jing et al., 2019b). Both PKL and CHR4 are homologous to SWI/SWF nuclear-localized chromatin remodelling factors of the CHD3 family (Ogas et al., 1999), and CHR4 is also named PICKLE RELATED1 (PKR1) (Aichinger et al., 2009). The chr4 mutant shows no obvious mutant phenotype under standard growth conditions (Aichinger et al., 2009). However, CHR4 function has been implicated in floral organ development because it interacts with the MADS-domain transcription factors AGAMOUS (AG), APETAL3 (AP3), PISTILLATA (PI), SEPALLATA3 (SEP3), and AP1, as revealed by immunoprecipitation of these factors (Smaczniak et al., 2012). Therefore, we hypothesized that the mutation in CHR4 caused the qem2 mutant phenotype. We tested this by introducing pCHR4:CHR4 and pCHR4:CHR4-VENUS constructs into qem2. The increased leaf number phenotype of qem2 was reduced to a similar number as in the progenitor quintuple mutant in all transformed lines (Figure 2D). Thus, we conclude that the later-flowering qem2 phenotype was caused by the mutation in CHR4.

Phenotypic characterization of chr4 and its effects on gene expression during floral induction

The qem2 mutant contains a mutation in the SNF2-related helicase/ATPase domain of CHR4, resulting in the substitution of a conserved alanine (ala) residue by valine (val) (ala713val) (Figure 3A). To analyze the chr4 mutant phenotype in the Col-0 background, we characterized the T-DNA insertion allele chr4-2 (SAIL_783_C05), containing a T-DNA insertion within the coding sequence between the helicase/ATPase domain and the DNA-binding domain (Figure 3A). The T-DNA insertion also causes a reduction in CHR4 mRNA levels (Supplemental Figure 3).

We compared the leaf number, bolting time, and flowering time of qem2 and chr4-2 with those of their respective progenitors under LDs (Supplemental Figure 4A, B) and SDs (Figures 3B–E). The qem2 mutant formed approximately 20 more rosette leaves and 30 more cauline leaves than the quintuple mutant under both LDs and SDs (Figure 3B, C and Supplemental Figure 4A, B). Despite having more rosette leaves, the bolting time of qem2 was similar to that of its progenitor (Figure 3D), whereas time to first open flower was markedly delayed in qem2 (Figure 3D, F).
which is consistent with the increased number of cauline leaves. The phenotypic difference between Col-0 and chr4-2 was less severe than that between qem2 and the quintuple mutant. Under LDs, chr4-2 and Col-0 initiated a similar number of leaves (Supplemental Figure 4B). Under SDs, chr4-2 and Col-0 had a similar rosette leaf number, but chr4-2 bolted earlier and produced more cauline leaves (Figure 3B–E). CHR4 function appeared to be more important for flowering control in the quintuple mutant background, suggesting it might preferentially regulate flowering via the GA and aging pathways.

The chr4-2 and qem2 mutants bolted slightly earlier than their progenitors but initiated a similar number or more rosette leaves (Figure 3B and D), suggesting that they might have a shorter plastochron and initiate rosette leaves more rapidly. To determine the plastochron, we counted rosette leaves weekly until the plants bolted under SDs. Early in rosette development, chr4-2 and qem2 produced leaves at a similar rate as their progenitors, but later in rosette development, the mutants produced leaves more rapidly than the progenitors, leading to a steep increase in leaf number (Figure 3G and H). More rapid leaf initiation can be related to an enlarged SAM (Barton, 2010); therefore, we compared the SAMs of chr4-2 and qem2 to those of Col-0 and the quintuple mutant, respectively, after 4 and 5 weeks of growth under SDs (Supplemental Figure 5). The SAMs of plants carrying either chr4 mutant allele were larger than those of their progenitors, but this was most pronounced for qem2 compared with the quintuple mutant (Supplemental Figure 5).

The transition to flowering in Arabidopsis can be conceptualised as two sequential steps in which the inflorescence meristem acquires different identities. After the transition from a vegetative meristem, the inflorescence meristem (I₁) initially forms cauline leaves and axillary branches, and after transition from I₁ to I₂, it initiates floral primordia (Ratcliffe et al., 1999). Rosette leaf number and days to bolting can be used as a proxy for the I₁ transition, whereas the number of cauline leaves produced on the flowering stem and days to the first open flower indicate when the I₁ to I₂ transition occurs. Cauline leaves can be distinguished from rosette leaves due to their smaller size and more pointed shape, so that the increased number of leaves on the inflorescence stem can be explained by a delayed I₂ transition rather than by enhanced internode elongation between rosette leaves. Compared to Col-0, chr4-2 is not delayed in the transition from vegetative meristem to I₁ but is delayed in the transition from I₁ to I₂. By contrast, compared to the
quintuple mutant, *gem2* mutants were strongly delayed in both the transition to I₁ and to I₂ (Figure 3B–E).

In Arabidopsis, AP1 confers floral meristem identity and is a marker for the I₁ to I₂ transition; therefore, we performed *in situ* hybridizations to monitor the appearance of AP1 mRNA through a developmental time course (Figure 4). At 5 weeks after germination, no AP1 expression was detected in any of the genotypes, indicating that the plant meristems were still vegetative. AP1 mRNA was detected at 6 weeks in Col-0 and chr4-2. In *gem2* mutants, AP1 mRNA appeared more than 1 week later than in the quintuple mutant, which is consistent with observation that more cauline leaves formed in *gem2* (Figure 4).

We then performed RNA-seq along a developmental time course to identify the genome-wide effects of CHR4 on gene expression during the floral transition. We examined the transcriptomes of shoot apices of Col-0, the quintuple mutant, chr4-2, and *gem2* plants grown for 3, 4, 5, or 6 weeks under SDs and compared the chr4-2 and *gem2* transcriptomes to those of Col-0 and the quintuple mutant, respectively (Supplemental Data Set 2). The analysis focused on 237 genes previously reported to regulate the floral transition in Arabidopsis (Bouche et al., 2016). In total, 26 of these genes were significantly differentially (adjp-value < 0.05 and log2FC |1|) expressed genes (DEGs) between chr4-2 and Col-0 (Figure 5A), and 18 were DEGs between *gem2* and the quintuple mutant (Figure 5C). Nine genes were common to the two lists (AGL79, BRANCHED1 (BRC1), FUL, SEP3, AGL17, SPL4, BROTHER OF FT AND TFL1 (BFT), EARLY FLOWERING 4 (ELF4) and MAF4). The expression of SPL4, which encodes a component of the age-dependent flowering pathway, increased at several time points in the chr4 and *gem2* mutants compared to their respective progenitors (Figure 5A–D). In particular, SPL4 was most highly expressed in 4-week-old *gem2* and in 5-week-old chr4-2 plants (Figure 5B–D). FUL was also more highly expressed in both mutants at later time points (Figure 5B and D) and is a direct target of SPL9, SPL15, and SPL3 during the floral transition (Wang et al., 2009; Yamaguchi et al., 2009; Hyun et al., 2016). Indeed, a corresponding small increase in mRNA levels of SPL9 and SPL15 was also observed in the CHR4 mutants (Supplemental Data Set 2). The earlier increase in expression of SPL4, SPL9, and SPL15 is consistent with the earlier bolting observed in the mutants, as *gem2* bolted around two days and chr4-2 around 10 days earlier than their respective progenitors (Figure 3D).
We detected elevated expression of *TERMINAL FLOWER1* (*TFL1*) in chr4-2 (Figure 5A and B) and *BFT* in *qem2* (Figure 5C). Overexpression of *TFL1* and *BFT*, which both encode proteins related to phosphatidylethanolamine-binding proteins, reduces *AP1* and *LFY* expression and delays floral organ initiation (Ratcliffe et al., 1998; Yoo et al., 2010). Consistent with this finding, *LFY* mRNA was also less abundant in *qem2* (Figure 5D). During the inflorescence meristem transition from I$_1$ to I$_2$, increased LFY activity induces floral meristem identity by directly activating *AP1* transcription and reducing GA levels, such that SPL9 recruits DELLA proteins to the regulatory region of *AP1* (Weigel et al., 1992; Wagner et al., 1999; Yamaguchi et al., 2014). Therefore, in the absence of CHR4 function, attenuated *LFY* transcription likely contributes to a delay in the transition to the I$_2$ phase, as reflected by the increased number of cauline leaves in *qem2*.

**CHR4 protein localisation in planta and identification of in vivo protein interactors of CHR4**

Chromatin remodelers are often recruited to target genes by specific transcription factors. Therefore, to further understand its mode of action during the floral transition, we identified proteins that interact with CHR4. We used the transgenic plants described above that express a fusion of VENUS fluorescent protein and CHR4 expressed from its native promoter (*pCHR4:CHR4-VENUS*). We analyzed the expression pattern of this CHR4-VENUS protein by confocal microscopy and compared it to the results of *in situ* hybridization analysis of *CHR4* mRNA. CHR4-VENUS was localized to the nucleus and its spatial pattern was similar to the mRNA pattern detected by *in situ* hybridization in the SAM, floral organs, and young leaves (Supplemental Figure 6).

To identify protein interactors, we immunoprecipitated CHR4-VENUS protein from inflorescence tissue and 5-week-old SD apical-enriched tissue using anti-GFP antibodies and used p35S-YFP transgenic plants as a negative control. Proteins that specifically co-immunoprecipitated with CHR4-VENUS were identified by protein mass spectrometry (Methods). In total, 136 and 342 proteins were significantly (FDR = 0.01) enriched in inflorescences and 5-week-old SD apex enriched tissue, respectively. The CHR4-interacting proteins in inflorescences included the floral homeotic MADS-domain transcription factors AP1, SEP3, PI and AP3 (Table 2 and Supplemental Data Set 3). The reciprocal experiment of immunoprecipitating AP1
was performed with gAP1:GFP plants and CHR4 was detected among the coimmunoprecipitated proteins (Supplemental Data Set 3 and Supplemental Figure 7). Taken together, these results confirm the previous finding that CHR4 could be co-immunoprecipitated with AG, AP3, PI, SEP3 and AP1 (Smaczniak et al., 2012). Moreover, SEP1 and SEP2 were also found here to be interaction partners of CHR4 in inflorescence tissues (Table 2, Supplemental Data Set 3, Supplemental Figure 7). In addition to floral homeotic proteins, other MADS-domain proteins were found to interact with CHR4 in inflorescences, including AGL6 and the fruit- and ovule-specific protein SHATTERPROOF2 (SHP2) (Favaro et al., 2003) (Table 2, Supplemental Data Set 3, Supplemental Figure 7).

Other classes of transcription factors involved in the floral transition were identified in CHR4 complexes. Notably, SPL2, SPL8, and SPL11 were found to be interaction partners in inflorescences, whereas SPL13 was identified as a partner in inflorescences and enriched apices (Table 2, Supplemental Data Set 3, Supplemental Figure 7). Furthermore, TARGET OF EARLY ACTIVATION TAGGED1 (TOE1), an AP2-domain transcription factor that represses the floral transition (Aukerman and Sakai, 2003), also interacted with CHR4 in enriched apices. A further list of transcription factors and chromatin remodellers identified as CHR4 interactors is provided in Table 2 and Supplemental Data Set 3.

These experiments demonstrated that CHR4 associates in vivo with several transcription factors of the MADS, SPL, and AP2 classes that contribute to the floral transition and floral meristem identity.

Genome-wide effects of CHR4 on histone modifications and gene expression
Proteins from the CHD3 group that includes CHR4 can participate in different chromatin remodelling pathways and either repress or activate gene expression, depending on the factors with which they associate. For example, PKL associates with genes enriched in trimethylation of histone H3 lysine 27 (H3K27me3), which is related to gene repression (Zhang et al., 2008; Zhang et al., 2012), and maintains this epigenetic state (Carter et al., 2018). In addition, PKL reduces H3K27me3 at specific target genes in particular tissues and environments (Jing et al., 2013). Changes in H3K27me3 and H3K4me3 were also reported in the rice (Oryza sativa) mutant of a CHR4 homologue (Hu et al., 2012). To test whether CHR4 regulates gene expression by influencing histone modifications, we compared global
H3K27me3 and H3K4me3 levels in Col-0 and chr4-2 plants (Supplemental Figure 8). No clear difference in the global frequency of these histone marks was observed between the two genotypes, suggesting that CHR4 does not affect the total accumulation of these histone modifications.

To test whether CHR4 affects the deposition of these histone marks at specific loci, we performed chromatin immunoprecipitation sequencing (ChIP-seq) to compare genome-wide H3K27me3 and H3K4me3 levels in Col-0 and chr4-2. H3K27me3 and H3K4me3 ChIP-seq experiments were performed on three biological replicates for each genotype (see Methods). In total, 10,194 H3K27me3-marked regions and 15,992 H3K4me3-marked regions were identified in the two genotypes (Supplemental Data Set 4). Quantitative comparison with DANPOS2 (Chen et al., 2013) revealed a subset of regions with significant differences (FDR < 0.05) in H3K27me3 or H3K4me3 levels between Col-0 and chr4-2. In total, 857 regions were differentially marked with H3K27me3 and 1,032 regions were differentially marked with H3K4me3 (Supplemental Data Set 4). Notably, hypermethylated as well as hypomethylated regions were identified in chr4-2 (Figure 6A). The genes differentially marked with H3K27me3 included regulators of key hormonal pathways involved in the floral transition, such as GIBBERELLIN 3-OXIDASE1 (GA3ox1) and GA3ox4, which encode GA biosynthesis enzymes. Genes encoding components of auxin signalling (ETTIN (ETT) and AUXIN RESISTANT 1 (AUX1)) and an enzyme that catabolises cytokinin (CYTOKININ OXIDASE 5 (CKX5)) were also differentially marked with H3K27me3 in chr4-2 (Supplemental Data Set 4). Genes differentially marked with H3K4me3 included the regulators of the floral transition SPL15, FLORAL TRANSITION AT THE MERISTEM1 (FTM1) (Torti et al., 2012), and JUMONJI DOMAIN-CONTAINING PROTEIN30 (JMJ30) (Jones et al., 2010; Yan et al., 2014) (Supplemental Data Set 4). In addition, 39 genes differentially marked by both H3K27me3 and H3K4me3 were detected, including the flowering-time regulators miR156D and AGL19 (Supplemental Data Set 4, Figure 6C).

We also examined the extent to which the differentially marked genes were also differentially expressed. H3K27me3 is associated with gene repression, and therefore, genes with higher H3K27me3 levels in chr4-2 compared to Col-0 were expected to be expressed at lower levels in chr4-2 than in Col-0. Indeed, a significant overrepresentation (Representation factor: 6.2, p-value < 1.317e-11) of downregulated genes was observed among those marked with increased levels of
H3K27me3 in chr4-2 (Figure 6B). Among the downregulated and hypermethylated genes in chr4-2 was AHL3, encoding an AT-hook protein that regulates vascular tissue boundaries in roots (Zhou et al., 2013) (Figure 6C). By contrast, H3K4me3 is associated with gene activation and therefore, genes marked with higher H3K4me3 levels in chr4-2 compared to Col-0 were expected to be expressed at higher levels. Indeed, a significant overrepresentation (Representation factor: 10.8, p-value < 2.176e-48) of upregulated genes between those marked with higher levels of H3K4me3 was observed (Figure 6B). Among the upregulated and hypermethylated genes in chr4-2 are CHR23, which is involved in stem-cell maintenance at the SAM (Sang et al., 2012) and SPL15, a promoter of the floral transition at the shoot meristem (Hyun et al., 2016) (Figure 6C). Moreover, spl15 produced fewer cauline leaves than the wild type (Schwarz et al., 2008), indicating a premature transition to the I2 phase of flower initiation. On the other hand, plants expressing a miR156-resistant transcript of SPL15 (rSPL15), which leads to an increase in SPL15 protein accumulation, produced more cauline leaves than the wild type (Hyun et al., 2016), indicating a delay in the transition to the I2 phase of flower initiation, as observed in qem2 mutants.

In conclusion, CHR4 affects H3K27me3 and H3K4me3 levels at a subset of loci in the genome, and changes in both histone modifications in chr4-2 are significantly correlated with changes in gene expression. Notably, a significant increase in H3K4me3 was detected at the SPL15 locus, and a higher level of SPL15 mRNA was found in chr4-2; these findings are consistent with the premature bolting and delay in the transition to the I2 phase of flower initiation observed in chr4-2.

DISCUSSION

We performed an enhanced genetic screen to identify regulators of the floral transition, and in particular, to focus on endogenous flowering pathways at the shoot meristem. To this end, we generated a quintuple mutant background strongly impaired in floral responses to environmental stimuli. Mutagenesis of these plants identified a chromatin remodeler, CHR4, which plays important roles in the floral transition, especially in response to endogenous flowering pathways and during the transition from forming cauline leaves with axillary branches (I1) to forming floral primordia (I2).
The quintuple mutant is strongly impaired in environmental flowering responses and flowers via endogenous pathways

The quintuple mutant showed strongly reduced flowering responses to long photoperiods and high ambient temperature. This insensitivity is consistent with the loss of function of *FT* and *TSF*, which confer photoperiodic responses, and the loss of function of *FT*, *TSF* and *SVP*, which are involved in responses to high ambient temperature (Yamaguchi et al., 2005; Kumar et al., 2012; Fernandez et al., 2016). Therefore, the floral transition in the quintuple mutant is likely promoted by endogenous flowering pathways. In support of this conclusion, RNA-seq analysis detected higher mRNA levels of several *SPL* genes in the mutant vs. Col-0. Some of these genes, such as *SPL15* and *SPL4*, are negatively regulated by miR156, which decreases in abundance as plants proceed from the juvenile to the adult phase (Wu and Poethig, 2006; Gandikota et al., 2007; Hyun et al., 2016). Therefore, these SPLs were previously considered to be components of an age-related flowering pathway (Wang et al., 2009; Hyun et al., 2017). However, the mRNA of *SPL8*, which is not regulated by miR156 but has overlapping functions with the miR156-targeted SPL genes (Xing et al., 2010), also increased in abundance in the quintuple mutant, suggesting a broader deregulation of this class of transcription factors in this genetic background.

Transcriptome profiling of the quintuple mutant also detected differential expression of genes encoding enzymes involved in GA biosynthesis, such as *GA20ox2*. Higher *GA20ox2* mRNA expression was detected in the quintuple mutant compared to Col-0 under SDs. The accumulation of GA₄ under SDs in Col-0 plants coincides with the floral transition and increased abundance of the mRNAs of floral meristem identity genes such as *LFY* (Eriksson et al., 2006). Although the GA biosynthesis pathway is complex and includes many enzymatic steps (Yamaguchi, 2008), *GA20ox2* appears to be important for controlling the floral transition, especially under SDs (Rieu et al., 2008; Plackett et al., 2012; Andres et al., 2014). SVP reduces *GA20ox2* transcript levels and GA levels at the shoot apex as part of the mechanism by which it represses flowering (Andres et al., 2014). We therefore propose that increased *GA20ox2* transcription in the quintuple mutant contributes to its higher GA levels and earlier floral transition under SDs. In support of this notion, the *qem1* mutation was found to be an allele of *GA20ox2* and to delay flowering of the quintuple mutant.
The proposed role for SPLs and GA in causing early flowering of the quintuple mutant is consistent with the previous finding that SPL proteins mediate some of the effects of GA during reproductive development (Porri et al., 2012; Yu et al., 2012; Yamaguchi et al., 2014; Hyun et al., 2016) and that SPL8 regulates several GA-mediated developmental processes (Zhang et al., 2007). Furthermore, SPL9 and SPL15 interact with DELLA proteins, which are negative regulators of GA responses that are degraded in the presence of GA (Daviere and Achard, 2013). SPL15 promotes the transcription of target genes that induce flowering, such as FUL and miR172b, and activation of these genes by SPL15 is repressed by interaction with DELLAs (Hyun et al., 2016). In Col-0, the role of SPL15 in flowering is particularly important under SDs, when floral induction occurs independently of environmental cues and is dependent on endogenous processes such as the GA pathway (Hyun et al., 2019). By contrast, the DELLA-SPL9 interaction can negatively or positively affect transcription, depending on the target genes and the developmental context (Yamaguchi et al., 2009; Yu et al., 2012). Taken together, these results demonstrate that the floral transition in the sensitized quintuple mutant background involves the interdependent functions of GA and SPL proteins.

A chromatin remodeller was identified as a regulator of the floral transition in the sensitized screen

The genetic framework for flowering-time control in Arabidopsis is based on analysis of late-flowering mutants identified after mutagenesis of early-flowering accessions (Koornneef et al., 1998). However, important regulators were not identified in these screens, but were readily found as early-flowering mutants from mutagenising late-flowering lines (Michaels and Amasino, 1999) or as late-flowering suppressor mutants after mutagenesis of transgenic plants or mutants requiring vernalization (Chandler et al., 1996; Onouchi et al., 2000). Here, we extended this approach by mutagenising a quintuple mutant background that flowered almost independently of environmental cues. Until recently, the molecular characterization of mutations isolated in such complex backgrounds using classical genetic approaches would have been extremely time-consuming and laborious, but this process has been simplified by the implementation of bulk-segregant analysis after backcrossing the mutant to the progenitor followed by whole-genome resequencing (Abe et al., 2012; Hartwig et al., 2012; Schneeberger, 2014).
The second characterized mutation identified in the quintuple mutant background, *qem2*, is an allele of *CHR4*. This gene encodes a chromatin remodeller that was previously identified as a member of protein complexes that include AP1 and other MADS-box transcription factors (Smaczniak et al., 2012), but its role in flowering had not been demonstrated genetically. Nevertheless, several chromatin modifiers and remodellers contribute to the regulation of the floral transition (Farrona et al., 2008), such as BRAHMA (BRM), a member of the SWI/SNF complex involved in nucleosome sliding and/or eviction, and the H3K27me3-specific histone demethylase RELATIVE OF EARLY FLOWERING6 (REF6), which acts cooperatively with BRM to regulate gene expression during floral development (Farrona et al., 2004; Lu et al., 2011; Wu et al., 2012; Li et al., 2016; Richter et al., 2019). Also, the SWI2/SNF2-RELATED1 (SWR1) complex protein PHOTOPERIOD-INSENSITIVE EARLY FLOWERING1 (PIE1) is involved in H2A.Z deposition and delays the floral transition (Noh and Amasino, 2003; March-Diaz et al., 2008; Coleman-Derr and Zilberman, 2012). Interestingly, PKL and PIE1 were previously proposed to act in the same pathway to define and maintain genomic domains with elevated H3K27me3 levels, suggesting that CHR4 may contribute at different levels within this process (Carter et al., 2018). Taken together, mass spectrometry identified several proteins in association with CHR4 that are involved in regulating histone modifications as well as multiple transcription factors with specific roles in floral meristem identity or the floral transition, suggesting that CHR4 functions in different multimeric complexes that regulate flowering.

**CHR4 affects the expression of flowering genes by modulating H3K4me3 and H3K27me3 levels and affects different stages of the floral transition**

The most closely related protein to CHR4 is another CHD3-like family member, PKL, which orchestrates deposition of H3K27me3 and facilitates nucleosome retention (Zhang et al., 2008; Zhang et al., 2012; Jing et al., 2013; Carter et al., 2018). In rice, loss of function of the CHR4 homologue CHR729 results in changes in the abundance of H3K27me3 and H3K4me3 at approximately 56% and 23%, respectively, of loci marked by these modifications (Hu et al., 2012). Similarly, we observed variation in H3K27me3 or H3K4me3 levels at a subset of loci marked by these modifications in *chr4-2*, indicating a conserved function between rice and
Arabidopsis. Notably, we observed higher levels of H3K4me3 at the SPL15 locus in chr4-2 vs. the wild type.

The floral transition is considered to be a dual-step process: in the first step, the inflorescence meristem produces cauline leaves and axillary branches (I1), and in the second phase, it forms floral primordia (I2) (Ratcliffe et al., 1999). Detailed phenotypic analysis of chr4 mutants showed that CHR4 affects both these phases but with opposite effects. The chr4 mutation accelerates the transition from the vegetative meristem to I1 but delays the I1 to I2 transition. The premature transition to I1 was reflected by earlier bolting, and this correlated with increased abundance of SPL15, SPL4 and FUL mRNA expression. These genes are associated with early bolting and flowering, and SPL15 in particular caused premature bolting when its expression was increased by mutations that rendered its mRNA insensitive to miR156 (Hyun et al., 2016). SPL15 also promotes the meristematic transition from vegetative to inflorescence meristem (Hyun et al., 2016). Moreover, spl15 mutants produced fewer cauline leaves than the wild type (Schwarz et al., 2008), whereas rSPL15 transgenmic plants produced more cauline leaves (Hyun et al., 2016), indicating that SPL15 extends the I1 phase. We propose that the higher expression of SPL15 in chr4 promotes earlier bolting and extends the I1 phase. This increased activity of SPL15 could also be enhanced in chr4 by increased activity of the GA biosynthetic pathway, as the resulting reduction in DELLA activity would be predicted to allow SPL15 to more effectively activate transcription of its target genes, leading to premature bolting and more cauline leaves.

Mutant chr4 plants also produced more cauline leaves and required more time to open the first flowers than their progenitors, indicating a delay in the I2 transition. These mutants also exhibited higher levels of TFL1 and BFT mRNAs; the overexpression of these genes delays the I2 transition by repressing AP1 and LFY expression (Ratcliffe et al., 1998; Yoo et al., 2010). Consistent with this conclusion, the onset of AP1 transcription occurred later in qem2 than in the quintuple mutant progenitor, and LFY mRNA was less abundant in qem2 than in the quintuple mutant in the RNA-seq time-course at week 6 in SDs. The chr4 mutant phenotype is strongly enhanced in the quintuple mutant background, probably explaining why chr4 was recovered in the sensitized mutant screen but was not previously identified by mutagenesis of Col-0 plants, where it exhibited a strong effect only under SDs. We
propose that CHR4 contributes to the floral transition in response to GA signalling and that the increased dependency of the quintuple mutant on the GA pathway to promote flowering increases the impact of CHR4 loss of function on the floral transition. Similarly, the stronger phenotype of chr4-2 in Col-0 under SDs than LDs is consistent with a specific role in the floral transition mediated by GA.

In conclusion, the combination of forward genetics and functional gene characterization identified CHR4 as a regulator of different stages of the floral transition. Immunoprecipitation of CHR4 suggested that it acts in distinct protein complexes that contain different transcription factors as well as other chromatin remodelling proteins. The contribution of CHR4 within distinct complexes presumably explains its pleiotropic effects, even during flowering, where it affects both bolting and floral identity during the transition from I₁ to I₂. Our genome-wide analyses represent the first step in understanding the mechanism by which CHR4 affects these phenotypes by identifying genes whose expression is altered by H3K27me3 or H3K4me3 in chr4 mutants. Further studies are now required to link the specific protein complexes in which CHR4 contributes to histone changes on defined targets. Attempts to perform ChIP-seq on pCHR4:CHR4-VENUS lines did not succeed, but pursuing this approach in the future would define the genome-wide sites with which CHR4 associates and help define its effects on the histone marks at direct target genes. Such approaches would help determine the mechanisms by which CHR4 regulates gene expression and allow this mechanism to be compared with that of PKL, which cooperates with PIE1 and CLF at target genes to maintain elevated H3K27me3 levels (Carter et al., 2018).

METHODS

Plant materials, growth conditions, and phenotypic analysis

For all studies, Arabidopsis thaliana Columbia (Col-0) ecotype was used as the wild type (WT). To construct the svp-41 flc-3 ft-10 tsf-1 soc1-2 quintuple mutant, svp-41 flc-3 FRI plants (Mateos et al., 2015) were first crossed to svp-41 ft-10 tsf-1 soc1-2 ful-2 plants (Andres et al., 2014). The F₁ plants were self-fertilized and the F₂ progeny were genotyped for each mutation except ful-2, which was scored phenotypically. Approximately 1,000 F₂ plants were grown in soil under LD conditions and DNA was extracted from those that flowered later than Col-0. Genotyping was
performed to identify plants that carried all mutations, lacked the FRI introgression, and were homozygous for FUL in the F₃ generation. chr4-2 corresponds to SAIL_783_C05. Homozygous mutant plants were selected by PCR using specific primers (Supplemental Data Set 5).

Seeds were immersed in 0.1% melt universal agarose (Bio-Budget Technologies GmbH) for three days at 4°C in darkness for stratification. Plants were grown in soil under controlled conditions of LDs (16 h light/8 h dark) and SDs (8 h light/16 h dark) at 21°C or 27°C. The light intensity was 150 µmol-m⁻²-s⁻¹ under all conditions. The growth-chamber is equipped with fluorescent tube bulbs from Philips (F17T8/TL841 ALTO-T8) to supply wavelengths from 430 to 650 nm, and supplemented with LEDs to provide light in the far-red spectrum. As a proxy for flowering time, the number of rosette and cauline leaves on the main shoot was counted as well as the number of days to bolting and first flower opening.

Ethylmethanesulfonate (EMS) treatment of seeds

For EMS treatment, 200 mg (~10,000) seeds of the quintuple mutant were wrapped in Miracloth and immersed in 0.1% KCl solution on a shaker at 4°C for 14 h. The seeds were washed with ddH₂O and treated with 100 mL 30 mM EMS diluted in ddH₂O on a magnetic stirrer in a fume hood overnight (8–9 h). The seeds were washed twice with 100 mL 100 mM sodium thiosulfate for 15 min and three times with 500 mL ddH₂O for 30 min. After washing, the seeds were immersed in 2 L 0.1% agarose. Approximately 50 seeds in 10 mL agarose were sown as the M₁ generation in 9 × 9 cm pots using plastic pipettes. The M₁ plants were grown and self-fertilized, and seeds were harvested in bulks of 50 M₁ plants. One hundred and forty-six M₂ bulked families were screened for plants showing altered flowering time.

GA treatment

The GA₄ stock (Sigma, Cat. G7276-5MG) was prepared in 100% ethanol with a final concentration of 1 mM. GA treatments were performed by spraying 2-week-old plants under SDs with either a GA solution (10 µM GA₄, 0.02% Silwet 77) or a mock solution (1% ethanol, 0.02% Silwet 77). Spraying was performed twice weekly until the plants bolted.

Selection of mutants and sequencing
Approximately 10 $M_2$ generation seeds from each $M_1$ plant were sown. Screening for potential mutants was initially performed under LD greenhouse conditions, and all plants were grown together with the quintuple mutant and Col-0 plants as a reference. Individuals that flowered later or earlier than the quintuple mutant in the $M_2$ population were selected. These $M_2$ putative mutants were self-fertilized and rescreened in the $M_3$ generation. Approximately 24 $M_3$ progeny of each potential mutant were grown under the same conditions to test the heritability of the phenotype. $M_3$ plants were backcrossed to the quintuple mutants to generate $BC_{1F_1}$ seeds. The $BC_{1F_2}$ offspring of such a cross formed the isogenic mapping population.

Approximately 70 plants showing the mutant phenotype were selected from a population of ~300 $BC_{1F_2}$ plants. One leaf sample of each selected plant was harvested and pooled. Leaf material from the quintuple plants was also harvested as a control. Genomic DNA was extracted from both pools and sent for Illumina sequencing with a depth of approximately 80-fold coverage. Reads were aligned to the TAIR10 reference genome using SHORE (Schneeberger et al., 2009). SHOREmap (Schneeberger et al., 2009; Sun and Schneeberger, 2015) was used to identify polymorphisms, and those present in approximately 100% of reads in the identified mutant but absent from the progenitor were identified as candidates for the causal mutation.

**In situ hybridization**

*In situ* hybridization was performed as previously described (Bradley et al., 1993), with minor modifications. Instead of Pronase, proteinase K (1 mg/mL in 100 mM Tris, pH 8, and 50 mM EDTA) was used for protease treatment by incubating at 37°C for 30 min. Post-hybridization washes were performed in 0.1× SSC instead of the original 2× SSC with 50% formamide. The sequences of primers used to generate the probes are listed in Supplemental Data Set 5. For each genotype and time point, 3 independent apices were analyzed.

**RNA extraction and RNA-seq analysis**

Total RNA was extracted from 15 shoot apices after removing all visible leaves under a binocular for each of the three independent biological replicates using an RNeasy Plant Mini Kit (Qiagen) and treated with DNase (Ambion) to remove residual genomic DNA. Library for sequencing was prepared using an Illumina TruSeq library preparation kit according to the manufacturer’s protocol. Sequencing was performed
using Illumina the HiSeq3000 platform in 150-bp single reads. For each sample, approximately 15,000,000 reads were generated. FastQC was used to assess quality control parameters (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/). To estimate expression levels, the RNA-seq reads were mapped to the A. thaliana TAIR10 (Lamesch et al., 2012) reference genome (ftp://ftp.arabidopsis.org/home/tair) using TopHat2 under default settings (Kim et al., 2013), except that only a single alignment was permitted per read and the coverage-based junction search was disabled (settings: -g 1 –no-coverage-search). Samtools was used to sort and index BAM alignment files and to calculate BAM file statistics (Li et al., 2009). HTSeq was used to tabulate the number of reads mapping to each genomic feature, with counts tabulated only for genes that completely overlapped a given feature (Anders et al., 2015). We used the Wald test implemented in DESeq2 to detect differentially expressed genes for pair-wised comparison. To visualise the expression levels of candidate genes, the expression level for each gene was calculated as transcripts per million (TPM).

**ChIP-seq experiment and data analysis**

Three independent biological replicates for each genotype were generated. For each sample, 1 g plant material was used per biological replicate. Material was collected from plants grown in SD at 21°C for 5 weeks (5–6 h after lights on). Using jeweler’s forceps, leaves with elongated petioles were removed to obtain SAM-enriched tissues. ChIP experiments were performed following a previously published protocol (Kaufmann et al., 2010) with minor modifications. Samples were sonicated in a water bath Bioruptor (Diagenode) four times for 5 min each of 15 sec on and 15 sec off, with a 1-min incubation between each sonication treatment. After the preclearing step, the sample was split into three aliquots: the first aliquot was incubated with anti-H3K27me3 antibody (Active Motif, Cat. 39155, Lot. 25812014), the second one was incubated with anti-H3K4me3 antibody (Millipore, Cat.17–614, Lot.1973237) and the third one with anti-H3 antibody (ab1791, Abcam). Samples were prepared for Illumina sequencing using the Ovation Ultralow V2 DNA-Seq Library Preparation Kit from Tecan Genomics according to the manufacturer’s protocol. H3K27me3 and H3K4me3 enrichment was tested by ChIP-qPCR before and after library preparation. Libraries were analyzed on the Bioanalyzer and quantified with the qBit before sequencing on the HiSeq3000. Samples were sequenced in a 150-bp single reads run.
FASTQ files were mapped to the *A. thaliana* genome TAIR10 using Bowtie (Langmead et al., 2009) with default parameters. Clonal reads were removed using a customised python script. Reproducibility between biological replicates was assessed using the Spearman correlation for the genome-wide read distribution at each pair of replicates using DeepTool (Ramirez et al., 2014). The “multiBamSummary” function was used with default parameters except for “bin size”, which was set to 1 kb and the “plotCorrelation” function of deepTools2 in Galaxy (http://deeptools.ie-freiburg.mpg.de/) (Supplemental Figure 9). H3K27me3 and H3K4me3 modified regions were identified with DANPOS2 (Chen et al., 2013). The “Dpeak” function in DANPOS2 was used with default parameters, except for the parameter – I (read extension length), which was set to 300 bp, the mean size of the DNA in the samples after sonification. Genomic regions were associated with genes if located within the start and the end of the gene using a customised python script.

**Plasmid construction**

Cloning of the *CHR4* locus was performed based on polymerase incomplete primer extension (Klock and Lesley, 2009) with modifications for large fragments and multiple inserts. All PCR amplifications were performed with Phusion Enzyme (New England BioLabs) following the manufacturer’s recommendations. The constructs pCHR4:CHR4-pDONR207 (18.4 kb) and pCHR4:CHR4:9AV-pDONR207 (19 kb) were generated as follows. Primers Q810 and Q811 were used to amplify the *CHR4* promoter (3.6 kb) and the PCR products were cloned into pDONR207 by BP reaction to generate the pCHR4-pDONR207 construct. The primer pairs Q058 and Q814, and Q815 and Q816 were used to amplify a fragment containing 9xala-VENUS (9AV) (0.7 kb) and the 3’UTR of CHR4 (3.8 kb), respectively. Overlap PCR with primers Q058 and Q816 was performed to fuse the amplicons. The primers Q817 and Q818 were used to linearize the construct pCHR4-pDONR207. The amplicons were mixed with linearized pCHR4-pDONR207 to construct the plasmid pCHR4:9AV:3’URTCHR4-pDONR207. The obtained plasmid was linearized with primers Q835 and Q836 and mixed with the coding sequence of *CHR4* (8.5 kb) amplified with primers Q819 and Q820 to construct the plasmid pCHR4:CHR4:9AV-pDONR207 (called *pCHR4:CHR4-VENUS* in the text). All primers used for molecular cloning are listed in Supplemental Data Set 5. Subsequently, the plasmids were cloned into the binary vector pEarleyGate301 (Earley et al., 2006) by LR reaction and transformed into *E.*
coli DH5-α-cells before being transformed into Agrobacterium tumefaciens GV3101 cells (Van Larebeke et al., 1974).

**Plant transformation and selection**

Plants (Col-0 and svp flc ft tsf soc1) were transformed by the floral-dip method (Clough and Bent, 1998). Transformants were selected by spraying twice with BASTA. The progenies were grown on plates with 1× Murashige and Skoog (MS) medium (Murashige and Skoog, 1962) containing sucrose and 10 µg mL⁻¹ phosphinotricin (PPT) to test for segregation and to select for single locus insertion lines and homozygosity in the following generations. Alternatively, the nondestructive ppt leaf assay was used to assess resistance to PPT. One young leaf per plant was harvested and placed on a plate with 1 MS without sucrose with 10 µg mL⁻¹ PPT. The plates were incubated for four days.

**Confocal microscopic analyses**

To visualise VENUS expression in shoot meristems, the method of (Kurihara et al., 2015) was used with minor modifications. Shoot apices were collected and placed in ice-cold 4% paraformaldehyde (PFA; Sigma-Aldrich) prepared in phosphate-buffered saline (PBS) at pH 7.0. The samples were vacuum infiltrated twice for 10 min each time, transferred to fresh 4% PFA, and stored at 4°C overnight. The next day, the samples were washed in PBS twice for 10 min each and cleared with ClearSee (10% xylitol, 15% sodium deoxycholate and 25% urea) at room temperature for ~1 week. The samples were then transferred to fresh ClearSee solution with 0.1% Renaissance 2200 and incubated in the dark overnight. The shoot meristems were imaged by confocal laser scanning microscopy (Zeiss LSM780) using settings optimised to visualise VENUS fluorescent proteins (laser wavelength, 514 nm; detection wavelength, 517–569 nm) and Renaissance 2200 (laser wavelength, 405 nm; detection wavelength, 410–510 nm).

**Sample preparation and LC-MS/MS data acquisition**

Three independent biological replicates for each genotype (gCHR4-VENUS and p35S-YFP), each consisting of 1 g plant material were generated. For inflorescence tissues, plants were grown in LD at 21°C, whereas SAM-enriched tissue samples were collected from plants growing in SD at 21°C for 5 weeks (5–6 h after lights on). Using jeweler’s forceps, leaves with elongated petioles were removed to obtain SAM-
enriched tissues. Nuclei were isolated according to a published protocol (Kaufmann et al., 2010). Samples were sonicated in a Bioruptor (Diagenode) water bath four times, 5 min each of 15 sec on and 15 sec off, with a 1-min incubation between each sonication treatment. Sonicated samples were centrifuged twice at 4°C for 10 min. The supernatants were transferred to a clean tube. After adding 40 µL GFP-trap Agarose beads from Chromotek (gta-20) and 10 µL Benzonase, the samples were incubated at 4°C for 2 hr. After incubation, the GFP-trap beads were washed four times with 1 mL wash buffer (750 µL 5M NaCl, 1.25 mL Tris-HCl pH 7.4 in 25 mL H₂O). Immunoprecipitated samples enriched with GFP-trap beads were submitted to on-bead digestion. In brief, dry beads were re-dissolved in 25 µL digestion buffer 1 (50 mM Tris, pH 7.5, 2M urea, 1 mM DTT, 5 µg µL⁻¹ trypsin) and incubated for 30 min at 30°C in a Thermomixer with 400 rpm. Next, the beads were pelleted and the supernatant was transferred to a fresh tube. Digestion buffer 2 (50 mM Tris, pH 7.5, 2M urea, 5 mM CAA) was added to the beads. After mixing and centrifugation, the supernatant was collected and combined with the previous one. The combined supernatants were incubated overnight in the dark at 32°C in a Thermomixer at 400 rpm. The digestion was stopped by adding 1 µL trifluoroacetic acid (TFA) and the samples were desalted with C18 Empore disk membranes according to the StageTip protocol (Rappsilber et al., 2003).

Dried peptides were re-dissolved in 2% acetonitrile (ACN), 0.1% TFA (10 µL) for analysis and measured without dilution. The samples were analyzed using an EASY-nLC 1200 (Thermo Fisher) coupled to a Q Exactive Plus mass spectrometer (Thermo Fisher). Peptides were separated on 16-cm frit-less silica emitters (New Objective, 0.75 µm inner diameter), packed in-house with reversed-phase ReproSil-Pur C18 AQ 1.9 µm resin (Dr. Maisch). Peptides (0.5 µg) were loaded onto the column and eluted for 115 min using a segmented linear gradient of 5% to 95% solvent B (0 min: 5%B; 0–5 min -> 5%B; 5–65 min -> 20%B; 65–90 min ->35%B; 90–100 min -> 55%; 100–105 min ->95%, 105–115 min ->95%) (solvent A 0% ACN, 0.1% FA; solvent B 80% ACN, 0.1%FA) at a flow rate of 300 nL min⁻¹. Mass spectra were acquired in data-dependent acquisition mode using the TOP15 method. MS spectra were acquired in the Orbitrap analyzer with a mass range of 300–1750 m/z at a resolution of 70,000 FWHM and a target value of 3 × 10⁶ ions. Precursors were selected with an isolation window of 1.3 m/z. HCD fragmentation was performed at a normalized collision energy of 25. MS/MS spectra were acquired with a target value
of $10^5$ ions at a resolution of 17,500 FWHM, a maximum injection time of 55 ms, and a fixed first mass of m/z 100. Peptides with a charge of +1, greater than 6, or with an unassigned charge state were excluded from fragmentation for MS2. Dynamic exclusion for 30s prevented repeated selection of precursors.

**Data analysis**

Raw data were processed using MaxQuant software (version 1.5.7.4, http://www.maxquant.org/) (Cox and Mann, 2008) with label-free quantification (LFQ) and iBAQ enabled (Tyanova et al., 2016). MS/MS spectra were searched by the Andromeda search engine against a combined database containing A. thaliana sequences (TAIR10_pep_20101214; ftp://ftp.arabidopsis.org/home/tair/Proteins/TAIR10_protein_lists/) and sequences of 248 common contaminant proteins and decoy sequences. Trypsin specificity was required and a maximum of two missed cleavages allowed. Minimal peptide length was set to seven amino acids. Carbamidomethylation of cysteine residues was set as fixed and oxidation of methionine and protein N-terminal acetylation as variable modifications. Peptide-spectrum-matches and proteins were retained if they were below a false discovery rate of 1%. Statistical analysis of the MaxLFQ values was carried out using Perseus (version 1.5.8.5, http://www.maxquant.org/). Quantified proteins were filtered for reverse hits and hits “identified by site”, and MaxLFQ values were log$_2$-transformed. After grouping the samples by condition, only proteins that had two valid values in one of the conditions were retained for subsequent analysis. Two-sample t-tests were performed with a permutation-based FDR of 5%. Alternatively, quantified proteins were grouped by condition and only hits that had three valid values in one of the conditions were retained. Missing values were imputed from a normal distribution (0.3 width, 2.0 downshift, separately for each column). Volcano plots were generated in Perseus using an FDR of 1% and an S0 = 1. The Perseus output was exported and further processed using Excel. ANOVA tables are shown in Supplemental Data Set 6.

**Accession Numbers**

The sequence of the genes and loci described here can be obtained from TAIR using the following gene identifiers: CHR4 (AT5G44800), SVP (AT2G22540), FLC (AT5G10140), SOC1 (AT2G45660), FT (AT1G65480), TSF (AT4G20370), GA20ox2 (AT5G51810) and SPL15 (AT3G57920).
The Illumina sequencing data have been deposited to the GEO with the dataset identifier GSE140728. The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE (Vizcaíno et al., 2016) partner repository with the dataset identifier PXD016457.

Supplemental Data

Supplemental Figure 1. svp flc ft tsf soc1 probably flowers as a result of endogenous pathways.

Supplemental Figure 2. Molecular genetic analysis of qem1.

Supplemental Figure 3. CHR4 expression in Col-0 and chr4-2.

Supplemental Figure 4. CHR4 loss-of-function phenotype in LDs.

Supplemental Figure 5. Shoot apical meristem size.

Supplemental Figure 6. CHR4 expression profile and protein localization.

Supplemental Figure 7. Volcano plot of protein–protein interactions.

Supplemental Figure 8. Global accumulation H3K27me3 and H3K4me3 marks in Col-0 and chr4-2.

Supplemental Figure 9. Spearman correlation for ChIP-seq samples.

Supplemental Table 1. Candidate SNPs annotated in genes by SHOREmap for qem1.

Supplemental Data Set 1. Whole-genome expression profiling experiments comparing the profiles of the genotypes Col-0 and svp flc ft tsf soc1 grown for 3, 4, 5 or 6 weeks under SD conditions.

Supplemental Data Set 2. Whole-genome expression profiling experiments comparing the profiles of the genotypes Col-0 vs chr4-2 and svp flc ft tsf soc1 vs. qem2 grown for 3, 4, 5 or 6 weeks under SD conditions.

Supplemental Data Set 3. IP-MS results for CHR4-VENUS and AP1-GFP pull-down: list of CHR4-interacting proteins.

Supplemental Data Set 4. Comparative analysis of H3K27me3 and H3K4me3 ChIP-seq results in Col-0 and chr4-2 obtained with DANPOS2.

Supplemental Data Set 5. List of primers used in the study.

Supplemental Data Set 6. ANOVA tables.
**Table 1.** Candidate SNPs in *qem2* annotated in genes.

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1 Chr: chromosome. 2 Position: position of the mutated nucleotide. 3 R: nucleotide in the reference genome (*svp flc ft tsf soc1*). 4 M: nucleotide in *qem2*. 5 N: number of reads supporting the mutation. 6 AF: allele frequency. 7 Sh: SHORE Score (max. 40). 8 Region: region of the locus where the mutation was identified. 9 Gen ID: gene identifier. 10 Type: type of mutation (nonsynonymous or synonymous). 11 AR: amino acid in the reference genome (*svp flc ft tsf soc1*). 12 AM: amino acid in *qem2*.

**Table 2.** List of CHR4 interacting proteins.

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<th>Gene ID</th>
<th>Name</th>
<th>No. Of Unique Peptides (IP1-IP2-IP3)</th>
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<th>log2 ratio</th>
<th>p-value</th>
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**TRANSCRIPTION FACTORS**

- AT1G69120 | AP1 |
- AT5G20240 | PI |
- AT3G54340 | AP3 |
- AT5G15800 | SEP1 |
- AT3G02310 | SEP2 |
- AT2G45650 | AGL6 |
- AT2G42830 | SHP2 |
- AT3G13960 | GRF5 |
- AT4G37740 | GRF2 |
- AT5G43270 | SPL2 |
- AT1G02065 | SPL8 |
- AT1G27360 | SPL11 |
- AT5G50670 | SPL13 |
- AT2G28550 | TOE1 |
- AT3G02150 | TCP13 |

**CHROMATIN REMODELLER**

- AT2G46020 | BRM |
- AT1G08600 | ATRX |
- AT5G04240 | ELF6 |
- AT2G28290 | SYD |
- AT2G25170 | PKL |
- AT3G12810 | PIE1 |
- AT5G18620 | CHR17 |
- AT3G06400 | CHR11 |
- AT3G48430 | REF6 |

* SAMs with younger leaves at 5w-SD-stage
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<th>Gene ID</th>
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<th>p-value</th>
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**TRANSCRIPTION FACTORS**

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**CHROMATIN REMODELLER**

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**General transcriptional coregulators**

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ACKNOWLEDGEMENTS

We thank Anne Harzen for support in the mass-spectrometry experiments. We thank René Richter, Franziska Turck and John Chandler for critical comments to the manuscript. A.P. was supported by an EMBO long-term fellowship (AFL-2017-74). X. Y. was supported by the China Scholarship Council (201804910196). The laboratory of G.C. is funded by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) under Germany’s Excellence Strategy (EXC 2048/1 Project ID: 390686111) and is supported by a core grant from the Max Planck Society.

AUTHOR CONTRIBUTIONS

S.Q., A.P. and G.C. conceived and designed the experiments. S.Q., A.P., X.Y. and F.A. performed the experiments. S.Q, A.P., K.S., H.S., B.S., S.S. and H.N. analysed the data. A.P, Q.S., and G.C wrote the manuscript.

REFERENCES


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chromatin remodeling ATPases cause embryogenesis and stem cell maintenance defects.

Plant J 72, 1000-1014.


Figure 1. Phenotypic and molecular characterization of the quintuple mutant svp flc ft tsf soc1.

(A) Days to bolting and (B) leaf number of plants grown under LD-21°C, SD-21°C and SD-27°C compared with Col-0. At least 17 plants were analyzed for each genotype. The data were analyzed with one-way ANOVA using Tukey’s HSD as a post-hoc test. Different letters indicate significant differences ($p \leq 0.05$). Whiskers represent a distance of 1.5 times the interquartile range.

(C) In situ hybridization analysis of FUL mRNA accumulation in shoot apical meristems of different genotypes grown in short days (SDs). Plants were harvested each week between 2 and 6 weeks after germination. Scale bar = 50 μm.

(D) Transcriptional profile comparisons in apices of svp flc ft tsf soc1. The analysis focuses on genes implicated in flowering time control. The data are represented as a heatmap to highlight upregulated (red) and downregulated genes (blue). Gene expression changes are represented as log2-fold changes.

(E) Box plots from RNA-seq data showing differential expression of SPL9, SPL15, FD, FUL and AGL6 in the apices of svp flc ft tsf soc1 and Col-0 under SDs. The Y axis shows transcripts per kilobase million (TPM). The X axis shows time of sampling as weeks after sowing. Whiskers represent distance from the lowest to the largest data point.
Figure 2. Molecular genetic analysis of qem2. (A) Leaf number at flowering of plants grown under LDs. Twelve plants were analyzed per genotype. The data were compared with one-way ANOVA using Tukey’s HSD as a post-hoc test. Different letters indicate significant differences ($p \leq 0.05$). Whiskers represent the distance of 1.5 times the interquartile range. (B) Images of qem2 and svp flc ft tsf soc1 plants approximately 50 days after germination, showing that qem2 produces more leaves than svp flc ft tsf soc1 under LDs. (C) Allele frequency (AF) estimates for EMS-induced mutations. Local AFs indicate that the qem2 mutation localized to chromosome (chr) 5. (D) Leaf number for svp flc ft tsf soc1, qem2, gCHR4 qem2 and gCHR4-VENUS qem2 plants under LDs. At least 11 plants per genotype were analyzed. The data were compared with one-way ANOVA using Tukey’s HSD as a post-hoc test. Different letters indicate significant differences ($p \leq 0.05$). Whiskers represent a distance of 1.5 times the interquartile range.
Figure 3. Characterization of CHR4. (A) Schematic representation of the CHR4 locus showing the position of the mutation in qem2 and the T-DNA insertion site (chr4-2). The CHR4 protein domains are illustrated: a plant homeodomain (PHD) zinc finger (blue), a chromo domain (red), a SNF2-related helicase/ATPase domain (green) and a DNA-binding domain (yellow). The EMS-induced protein sequence change is located within the SNF2-related helicase/ATPase domain. (B) Leaf number, (C) cauline leaf number, (D) days to bolting and flowering and (E) number of days from bolting to flowering of Col-0, chr4-2, svp flc ft tsf soc1 and qem2 plants grown under short days (SDs). At least 17 plants were analyzed for each genotype. The data were compared with one-way ANOVA using Tukey HSD as a post-hoc test. Different letters indicate significant differences ($p \leq 0.05$). Whiskers represent a distance of 1.5 times the interquartile range. (F) 12-week-old plants growing in SDs. Red arrows indicate first open flower. Scale bar = 10 cm (G) Rosettes of Col-0, chr4-2, svp flc ft tsf soc1 and qem2 plants after 38 days and 43 days of growth in SDs. Scale bar = 1 cm (H) Rosette leaf number of Col-0, chr4-2, svp flc ft tsf soc1 and qem2 plants grown under SDs from 3 weeks to 7 weeks. 18 plants were analyzed for each genotype. Error bars represent standard deviation of the mean. * indicates significant differences (p-value < 0.05) between Col-0 and chr4-2 (blue) or svp flc ft tsf soc1 and qem2 (red).
Figure 4. Temporal and spatial patterns of expression of the floral meristem identity gene AP1 in Col-0, chr4-2, svp flc ft tsf soc1 and qem2. In situ hybridization analysis of AP1 mRNA accumulation in the shoot apical meristems of plants under SDs. The genotypes analyzed are shown together with the number of weeks (w) after germination when material was harvested. For each time point and genotype, three independent apices were examined with similar results. Scale bar = 50 μm.
Figure 5. Transcriptional changes in chr4 mutants. (A) Transcriptional profile comparisons represented as a heatmap to highlight genes implicated in flowering time control that are significantly upregulated (red) or downregulated (blue) in chr4-2 compared to WT. Gene expression changes are represented as log2-fold change. (B) Box plots from RNA-seq data showing FD, TFL1, FUL, and SPL4 transcript levels in apices of chr4-2 and Col-0 under SDs. The Y axis shows transcripts per kilobase million (TPM). The X axis shows time of sampling as weeks after sowing. Whiskers represent distance from the lowest to the largest data point. (C) Transcriptional profile comparisons represented as a heatmap to highlight genes implicated in flowering time control that are significantly upregulated (red) or downregulated (blue) in qem2 compared to svp flc ft tsf soc1. (D) Box plots from RNA-seq data showing FUL, SPL4, LFY and BRC1 transcript levels shown as transcripts per kilobase million (TPM) in apices of qem2 and svp flc ft tsf soc1 under SDs. The Y axis shows transcripts per kilobase million (TPM). The X axis shows time of sampling as weeks after sowing. Whiskers represent distance from the lowest to the largest data point.
Figure 6. Histone modification variation in chr4-2. (A) Scatterplots showing H3K27me3 and H3K4me3 enrichment between Col-0 and chr4-2 in apices of five-week-old plants grown under SDs. Blue and orange dots represent significantly more highly methylated regions at FDR = 0.05 in Col-0 and chr4-2, respectively. (B) Venn diagram showing the overlap between differentially expressed genes (DEGs) and genes differentially marked by H3K27me3 and H3K4me3. (C) H3K27me3 and H3K4me3 profiles and expression of AHL3, AGL19, CHR23 and SPL15.
Mutagenesis of a Quintuple Mutant Impaired in Environmental Responses Reveals Roles for CHROMATIN REMODELING4 in the Arabidopsis Floral Transition
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