The APETALA2 homolog CaFFN regulates flowering time in pepper

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Abstract
Flowering time is an important agronomic trait that contributes to fitness in plants. However, the genetic basis of flowering time has not been extensively studied in pepper. To understand the genetics underlying flowering time, we constructed an F2 population by crossing a spontaneous early flowering mutant and a late-flowering pepper line. Using bulked segregant RNA-seq, a major locus controlling flowering time in this population was mapped to the end of chromosome 2. An APETALA2 (AP2) homolog (CaFFN) cosegregated with flowering time in 297 individuals of the F2 population. A comparison between the parents revealed a naturally occurring rare SNP (SNP2T > C) that resulted in the loss of a start codon in CaFFN in the early flowering mutant. Transgenic Nicotiana benthamiana plants with high CaFFN expression exhibited a delay in flowering time and floral patterning defects. On the other hand, pepper plants with CaFFN silencing flowered early. Therefore, the CaFFN gene acts as a flowering repressor in pepper. CaFFN may function as a transcriptional activator to activate the expression of CaAGL15 and miR156e and as a transcriptional repressor to repress the expression of CaAG, CaAP1, CaSEP3, CaSOC1, and miR172b based on a qRT-PCR assay. Direct activation of CaAGL15 by CaFFN was detected using yeast one-hybrid and dual-luciferase reporter assays, consistent with the hypothesis that CaFFN regulates flowering time. Moreover, the CaFFN gene association analysis revealed a significant association with flowering time in a natural pepper population, indicating that the CaFFN gene has a broad effect on flowering time in pepper. Finally, the phylogeny, evolutionary expansion and expression patterns of CaFFN/AP2 homologs were analyzed to provide valuable insight into CaFFN. This study increases our understanding of the involvement of CaFFN in controlling flowering time in pepper, thus making CaFFN a target gene for breeding early maturing pepper.

Introduction
The timing of the transition from vegetative growth to flowering is a major factor that ensures that plants produce enough progeny for their sustainable development1–3. Thus, flowering time is an important agronomic trait that contributes to fitness in annual plants. Genetic analysis of the model plant Arabidopsis thaliana has identified numerous genes involved in flowering-time control that may participate in the photoperiod, vernalization, ambient temperature, aging, autonomous or gibberellin pathways4, each of which plays a pivotal role in flowering time and promotes function in response to exogenous or endogenous signals. Downstream of these pathways are a small number of floral integrator genes, including FLOWERING LOCUS T (FT) and SUPPRESSOR OF OVEREXPRESSION OF CONSTANS 1 (SOC1)5–9. ‘Integrators’ activate floral meristem identity (FMI) genes, such as LEAFY (LFY), APETALA1 (API), APETALA2 (AP2), FRUITFULL (FUL), and CAULIFLOWER (CAL), which encode proteins that promote floral development1. Among these FMI genes, the transcription factor AP2 directly promotes the expression of the floral repressor AGAMOUS-LIKE15 (AGL15) and directly represses the expression of the flowering promoting transcripts SOCI, API, SEPALLATA3 (SEP3),...
AGAMOUS (AG), and FUL. In addition, AP2 positively regulates miR156 and negatively regulates miR172 to form a complex direct feedback loop that influences AP2 expression.

Solanaceae includes species such as potato (Solanum tuberosum), tomato (S. lycopersicum), pepper (Capsicum annum), eggplant (S. melongena), and tobacco (Nicotiana tabacum) that are of agricultural interest or research interest. Among these species, tomato has been more extensively studied for the control of flowering time11–13. As a member of the Solanaceae family, pepper (C. annum L.) has a sympodial shoot structure. The pepper shoot apical meristem (SAM) first produces stems and leaves alternately in the vegetative phase and then, upon the transition to the reproductive phase, terminates in an inflorescence meristem (IM) that subsequently develops into the first flower; plant growth continues from the upper most axillary meristem14,15. Flowering time in pepper can be best measured by counting the number of leaves (or nodes) on the primary stem (Nle) from the cotyledon until the first flower, which has also been described as the number of first flower nodes (FFNs)16–18. In pepper, several genes controlling flowering time have been identified by screening an EMS-mutagenized population. Of these genes, mutants of CaJOINTLESS (CaJ), CaBLIND (CaBL) and CaS were late flowering14,15,19, and mutants of fasciculate (fa) and E-62 were early flowering20,21. These genes have a pleiotropic effect on plant growth in addition to affecting flowering time in pepper; for example, CaS is required for flower formation, and CaBL regulates axillary meristem initiation15,19. However, most of these mutant genes are unlikely to be utilized in pepper breeding; previous studies have reported that mutants of CaS were characterized by complete inhibition of flower formation15,19, and mutants of CaBL had no axillary shoot development in the vegetative phase and only two sympodial units in the reproductive phase19.

Pepper germplasm resources exhibit extensive natural variation in flowering time22. Genetic analysis showed that the flowering time trait in pepper was a quantitative trait commonly controlled by a few major genes and some minor genes, as well as environmental factors such as temperature and photoperiod16,17,23,24. Quantitative trait loci (QTLs) affecting flowering time have been identified on different chromosomes using different populations17,18,23–26. For example, Tan et al.17 mapped one major and five minor QTLs affecting FFNs on chromosomes P2, P7, P10 and P11 based on an interspecific genetic map, and 12 genes were recommended as important candidate genes for the major QTL Nle2.2. Zhang et al.18 identified 23 candidate genes on chromosome 12 within an interval of 3.98 Mb using combined specific-locus amplified fragment sequencing (SLAF-seq) and bulked segregant analysis (BSA). Zhu et al.26 detected three flowering time QTLs based on an interspecific F2 population, in which an AP2 protein encoding gene Capana02g000700 was predicted to be the candidate gene for the major QTL. However, further studies to identify and validate the function of candidate genes controlling flowering time in pepper have been limited to date.

The early flowering pepper B9431, which can blossom and bear fruit at the first node on the main stem, is an ideal natural material for determining the genetic control of flowering time in pepper. In this study, an F2 population with 297 plants derived from an intraspecific cross between inbred lines B9431 (early flowering) and A145 (late flowering) was developed. The loci controlling flowering time in this population was mapped by the bulked segregant RNA-Seq (BSR-Seq) method. The heterologous overexpression assay in N. benthamiana and the virus-induced gene silencing (VIGS) assay in pepper A145 were performed to verify the function of the candidate gene CaFFN. Similarly, a real-time fluorescent quantitative PCR assay, a yeast one-hybrid assay and a dual-luciferase reporter assay were conducted to determine how the CaFFN gene influences the flowering time of pepper. Furthermore, CaFFN gene association analysis was performed on 167 pepper inbred lines to test the effect of this gene on the control of natural variations in flowering time. The phylogeny, evolutionary expansion and expression patterns of CaFFN/AP2 homologs were analyzed to provide useful insight into CaFFN.

Results

Genetic analysis of the flowering time trait in pepper

A natural precocious variant was discovered in our breeding work, from which a stable early flowering inbred line named B9431 was created after years of selfing and selective breeding (Fig. 1a). B9431 can flower after the development of one leaf on the primary stem. To investigate the inheritance of flowering time in pepper, an F2 population including 297 plants developed from a cross between B9431 and A145 was studied. Flowering time, in this study, was measured by counting the number of leaves on the primary stem between the cotyledon and the first flower (referred to as the number of FFNs) according to previous studies16–18. The average number of FFNs on parent B9431 was 2.3 (ranging from 1 to 4), parent A145 was 14 (ranging from 13 to 15), and the F1 progeny was 8.5 (ranging from 8 to 10); that on the F2 population ranged from 2 to 14. The F2 population showed a bimodal distribution (Fig. 1b). Based on a distribution of the F2 population, in which an AP2 protein encoding gene Capana02g000700 was predicted to be the candidate gene for the major QTL. However, further studies to identify and validate the function of candidate genes controlling flowering time in pepper have been limited to date.

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BSR-Seq identified the major locus underlying flowering time in pepper

Thirty plants with the minimum number of FFNs from the F₂ population were chosen to construct an ‘early flowering pool’, and 30 plants with the maximum number of FFNs were chosen to construct a ‘late-flowering pool’. The total RNA of the two pools was extracted for Illumina high-throughput sequencing, and the clean data were aligned to the C. annuum cv. CM334 genome chromosomes (release 1.55) (http://peppergenome.snu.ac.kr/). The frequencies for the single-nucleotide polymorphisms (SNPs) thus detected were calculated for each pool. The average difference in the SNP frequency (index) between the two pools was calculated across a 1.0 Mb genomic interval using a 3.0 Mb sliding window, and the results were plotted against all twelve chromosomes of the reference genome (Fig. 1c). The figure shows an obvious peak in the SNP delta values at the distal end of chromosome 2, which indicates one strong major locus governing flowering time in this region.

Identification of a candidate gene with a point mutation at the start codon in the early flowering parent B₉₄₃₁

At the end of chromosome 2, the AP2 transcription factor-encoding gene CA02g14540 caught our attention, since its likely ortholog AP2 in Arabidopsis has numerous functions, including participation in floral transition and development. The sequences of this gene in the two parents of the F₂ population were determined by sequencing the PCR products amplified with the primers listed in Supplementary Table S1. A point mutation at the start codon ATG (referred to as SNP2T > C) was discovered in the early flowering parent B₉₄₃₁ (Fig. 2a). This SNP led to the loss of the original start codon of CaFFN and resulted in a putative frameshift mutation of CaFFN in B₉₄₃₁. To investigate the potential of this gene to be a candidate gene controlling flowering time in this study, a cleaved amplified polymorphic sequence (CAPS) marker CSF2 (Fig. 2b, Supplementary Table S1) was developed based on SNP2T > C. The 297 individuals in the F₂ population of B₉₄₃₁ × A₁₄₅ were genotyped using marker CSF2. The results showed that all early flowering (number of FFNs ranging from 2 to 4) plants had the same genotype as B₉₄₃₁, and all late-flowering (number of FFNs ranging from 6 to 14) plants had the same genotype as A₁₄₅ or the F₁ (Fig. 2b), indicating cosegregation of CSF2 with flowering time. These results demonstrated that CA02g14540 was a candidate gene (referred to as the CaFFN gene) underlying flowering time in the F₂ population of B₉₄₃₁ × A₁₄₅.
Two *N. benthamiana* transgenic lines expressing *CaFFN* exhibited a notable delay in flowering time

To investigate whether *CaFFN* might function in controlling flowering time, transgenic *N. benthamiana* lines expressing pepper *CaFFN* under the control of the CaMV 35S promoter were generated. However, no obvious changes in flowering time or flower patterning were observed in the 32 transgenic lines, even though the expression of the *CaFFN* gene was detected by qRT-PCR. *CaFFN* is an AP2-like gene with a putative conserved miR172 target site. It is thus speculated that a modification of the miR172 target site would increase the expression of the *CaFFN* gene was detected by qRT-PCR. For this reason, 22 transgenic lines transformed with modified *CaFFN* cDNA containing six mismatches ( synonymous mutations) to miR172 (miR172-resistant *CaFFN* mutant, 35S::CaFFNm3) were generated (Fig. 2a). Among them, two transgenic lines (lines 10 and 16) exhibited a notable delay in flowering time (Fig. 3a).

Levels of *CaFFN* mRNA were high in the late-flowering 35S::CaFFNm3 lines

To further explore why these two transgenic lines behaved differently, 14 transgenic lines transformed with 35S::CaFFNm3 were used for qRT-PCR analysis. Primers FFN-RT-jF4 and CaFFN-sNR2 (Supplementary Table S1) specific to *CaFFN* were designed to examine the expression level of *CaFFN* in *N. benthamiana*. The results revealed that the expression level of the *CaFFN* gene was higher in these two phenotypic lines than in other plants (Fig. 3c). These two transgenic lines produced numerous leaves in an extended period of vegetative growth and flowered two and a half months later than control plants. In addition to the delayed-flowering phenotype, sterility and floral patterning defects, which were characterized by shorter filaments and poorly developed petals with a reduced size, were observed in line 10 (L10) and line 16 (L16) (Fig. 3b). L16, with the highest level of *CaFFN* mRNA, exhibited the most dramatic change; it produced the smallest petals and an even shorter style (Fig. 3b). These results provide evidence that the *CaFFN* gene has a function in regulating flowering time and floral development.

Silencing of the *CaFFN* gene promotes early flowering in pepper

To study the functional roles of *CaFFN* in flowering time in pepper, loss-of-function experiments were
performed on parental line A145, in which CaFFN was silenced by VIGS. Agrobacterium carrying the tobacco rattle virus (TRV)-based vector TRV2::CaFFN mixed with Agrobacterium carrying the TRV1 vector were infiltrated into pepper seedlings at the four-leaf stage. Seedlings injected with TRV2::PDS- and TRV1-containing Agrobacterium were used as controls to measure the effect of gene silencing, which induced a photobleaching phenotype. Seedlings injected with TRV2- and TRV1-containing Agrobacterium were used as negative controls. The infiltrated seedlings were grown at 20 ± 1 °C and 90 ± 5% relative humidity to enhance silencing efficiency. Plants injected with TRV2::PDS showed photobleaching (Fig. 4a), suggesting successful gene silencing in pepper A145. Plants injected with TRV2::CaFFN flowered earlier (the average number of FFNs was eight, with a standard error of 0.30) than control plants (the average number of FFNs was 14, with a standard error of 0.26) (Fig. 4a). Consistently, qRT-PCR results showed an obvious decline in the expression level of CaFFN in TRV2::CaFFN plants compared to the control plants (Fig. 4b). The result that silencing of the CaFFN gene leads to early flowering in TRV2::CaFFN plants further shows the function of CaFFN as a flowering repressor in pepper.

Gene expression changes in response to CaFFN silencing

The above results in this study indicated that CaFFN is a repressor of flowering. However, the mechanism underlying flowering time alteration caused by CaFFN remains to be determined. As the closest homolog to AP2, determining whether the network surrounding AP2 is conserved and applicable to CaFFN is worthy of our attention. For this purpose, the likely orthologs of AG, AGL15, AP1, SEP3, and SOC1 (CaAG [CA01g12980], CaAGL15 [CA02g27000], CaAP1 [CA02g27000], CaSEP3 [CA11g09730], and CaSOC1 [CA01g19500]) were searched by reciprocal best-hit analysis in the C. annuum cv. CM334 genome (release 1.55) (http://peppergenome.snu.ac.kr/). The expression levels of these flowering time and floral organ development genes were analyzed by qRT-PCR relative to the housekeeping gene CaGAPDH [CA03g24310]. The results showed that the expression levels of CaAG, CaAP1, CaSEP3, and CaSOC1 were significantly higher in CaFFN-silenced plants than in the control (P < 0.01). In contrast, the expression level of CaAGL15 in CaFFN-silenced plants was significantly lower than that in the control (P < 0.01) (Fig. 4b). The expression levels of microRNAs were determined relative to Cal6 [CA07g02150]. MiR172b exhibited an increased level and miR156e represented a reduced level in CaFFN-silenced plants (Fig. 4b). The results suggested the role of CaFFN as a transcriptional activator to promote the expression of CaAGL15 and miR156e and as a
Fig. 4 (See legend on next page.)
transcriptional repressor to repress the expression of CaAG, CaAP1, CaSEP3, CaSOC1, and miR172b.

The CaFFN transcription factor directly activates the floral repressor gene CaAGL15

The finding that the expression level of the floral repressor gene CaAGL15 was reduced in CaFFN gene-silenced pepper plants prompted us to examine whether CaAGL15 is directly regulated by CaFFN. We checked the ability of CaFFN to bind to the cis-elements of CaAGL15 using a yeast one-hybrid assay. The full-length cDNA of CaFFN was cloned in frame with the GAL4 activation domain (AD) in the pGADT7 vector. The cis-elements of CaAGL15 were inserted into the pAbAi vector. The minimal inhibitory concentration of aureobasidin A (AbA) was 400 ng/ml for bait strains of CaFFN and the bait vector grew in SD/-Leu selection medium supplemented with the minimal inhibitory concentration of AbA, while yeast transformed with pGADT7-lam and pGADT7-pAbAi-P53 were used as positive and negative controls, respectively. Superscripts represent the concentration of aureobasidin A (AbA). The ability of CaFFN to bind to the promoter of CaAGL15 was tested by dual luciferase assay. The LUC/REN fold value of the empty vector P62-SK plus promoter was set to 1.0. The values are the means ± SD (Student’s t test, **p < 0.01, ***p < 0.001). Hypothetical models for the transcriptional regulatory network involving CaFFN according to the transcriptional relationships in this study and reports in Arabidopsis.

Arrows represent activation, and bars represent inhibition. CaFFN bound to the cis-elements of CaAGL15 in the yeast one-hybrid (Y1H) assay. pGADT7-p53/pAbAi-p53 and pGADT7-lam/pAbAi-P53 were used as positive and negative controls, respectively. The results showed that CaFFN directly interacted with CaAGL15, since yeast transformed with pGADT7-CaFFN and the bait vector grew in SD/-Leu selection medium supplemented with the minimal inhibitory concentration of AbA, while yeast transformed with pGADT7-lam and pAbAi-P53, as a negative control, did not (Fig. 4c).

In addition, a dual-luciferase assay was performed in N. benthamiana leaves. Recombinant pGREEN0800-CaAGL15 containing the promoter region of CaAGL15 (reporter) and the fused protein with CaFFN (effector) were coinfiltrated into leaves. Cotransformation of CaAGL15 and the CaFFN protein resulted in a twofold LUC/REN ratio compared with the control (Fig. 4d). The results indicated that CaFFN can bind to the CaAGL15 promoter in vivo and transactivate the expression of the reporter gene.

CaFFN was significantly associated with flowering time in the pepper population

In this study, the CaFFN gene was shown to be a flowering repressor in pepper. To determine the distribution of the SNP2T > C mutation of the CaFFN gene in the pepper population, the CaFFN sequences of 383 pepper accessions were surveyed in PepperPan, a pepper pangenome browser. The results showed that SNP2T > C was a rare SNP that occurred in only three additional accessions, CXJ21 (B33), CXJ22 (B35), and CXJ70 (A100) (Fig. 5a), and these three accessions were early flowering materials, with FFN numbers of 3, 3, and 5, respectively, suggesting that SNP2T > C could affect flowering time in multiple genetic backgrounds. The rare frequency of SNP2T > C also implies that the mutation occurred very recently and has not been widely used in breeding programs.

Due to the low number of accessions with nucleotide C at position 2, SNP2T > C could not be validated by association analysis. To further investigate whether CaFFN is an important regulator of flowering time in the pepper
population, the numbers of FFNs in 164 pepper inbred lines were investigated. The numbers of FFNs exhibited wide phenotypic variations, ranging from 2 to 23, with an average of 10.5. Combined with the genotypic SNP/Indel data from the CaFFN sequenced region in 164 pepper accessions, an association analysis was performed to quantify significant associations. A SNP 50 bp upstream of the start codon (SNP-50G > A) was significantly associated with FFNs ($P = 3.25E−07 < 0.01$), explaining 14.5% of the phenotypic variation. The results demonstrated that CaFFN was significantly associated with flowering time in the pepper population.

A sequence comparison between the CaFFN alleles of 53 pepper accessions did not reveal clear causative polymorphisms in the coding region. It is speculated that the 5′-UTR may influence the number of FFNs by affecting the expression of the CaFFN gene. To test this hypothesis, 28 randomly selected accessions were divided into two haplotypes according to SNP-50G > A, and the expression level of the CaFFN gene was determined by qRT-PCR (Fig. 5b). Statistical analysis showed that SNP-50G > A significantly correlated with the expression level of the CaFFN gene ($P < 0.05$), suggesting that the expression level of CaFFN might be influenced by its 5′-UTR.

**Characterization of the CaFFN/AP2 homologs in pepper**

CaFFN has two AP2 domains and is a member of the AP2 family gene. AP2 family genes with two or more AP2 domains play essential roles in plant growth and development, as shown in this study. To systematically analyze the AP2 family genes in pepper, the HMM profile with the AP2 domain (PF00847) was used as a query, and a total of 24 AP2 family genes were obtained in the genome database of the cultivated pepper Zunla-1 (Fig. 6). Duplication events are known to be the major driving force in the expansion of gene families. To explore the major driving force for the expansion of the AP2 family in pepper, the duplication types of each gene were detected by the MCScanX package, including singleton, dispersed, proximal, tandem and whole-genome duplication/triplication (WGD/T) or segmental duplication. Combined with the data of the retained triplicate genes from WGT in pepper, the duplication types of each gene were detected by the MCScanX package, including singleton, dispersed, proximal, tandem and whole-genome duplication/triplication (WGD/T) or segmental duplication. Combined with the data of the retained triplicate genes from WGT in pepper, the duplication types of each gene were detected by the MCScanX package, including singleton, dispersed, proximal, tandem and whole-genome duplication/triplication (WGD/T) or segmental duplication. Combined with the data of the retained triplicate genes from WGT in pepper, the duplication types of each gene were detected by the MCScanX package, including singleton, dispersed, proximal, tandem and whole-genome duplication/triplication (WGD/T) or segmental duplication. Combined with the data of the retained triplicate genes from WGT in pepper, the duplication types of each gene were detected by the MCScanX package, including singleton, dispersed, proximal, tandem and whole-genome duplication/triplication (WGD/T) or segmental duplication.

**Discussion**

In this study, the locus controlling flowering time in the F2 population derived from an intraspecific cross between the early flowering pepper B9431 and the late-flowering pepper A145 was mapped to the end of chromosome 2. This locus was reported to be a flowering time locus in several previous reports. Borovsky et al. provided supporting genetic evidence for the putative function of CaAP2 in this chromosome region as a flowering repressor, but direct proof of the function of CaAP2 is still lacking. The CaAP2 gene (referred to in this study as the CaFFN gene) with a spontaneous natural point mutation (SNP/T > C) at the start codon in B9431 cosegregated with the FFN trait in the F2 population of 297 individuals; this is additional genetic evidence for the function of CaFFN/AAP2 as a flowering repressor. To provide more direct proof of the function of the CaFFN gene, transgenic *N. benthamiana* lines expressing the CaFFN gene were generated, and the flowering repressor function of this gene was verified by two 35S::CaFFNm3 transgenic lines with a high level of CaFFN mRNA that flowered two and a half months later than control plants. The function of the CaFFN gene was also verified by VIGS of the CaFFN gene in the pepper A145, which flowered earlier with a six-node reduction in the number of FFNs.

Putative miR172 target sites in AP2-like genes are conserved in eudicots, monocots, and early land plants, indicating that the expression of AP2-like genes could be regulated by miR172. Moltschka et al. generated transgenic *N. benthamiana* lines expressing *Arabidopsis* wild-type AP2 (35S::AP2) and the miR172-resistant AP2 mutant (35S::AP2m3). Nearly all 35S::AP2 plants were nonphenotypic with barely detectable levels of AP2 mRNA; however, 35S::AP2m3 plants accumulated high levels of AP2 mRNA and exhibited floral patterning defects such as the proliferation of numerous petals, stamens, and carpels. In our study, all 35S::CaFFN plants were nonphenotypic with a low level of CaFFN mRNA. However, only two 35S::CaFFNm3 plants that accumulated high levels of CaFFN mRNA exhibited an obvious late-flowering phenotype and floral patterning defects. It
is inferred that a visible flowering-time alteration in *N. benthamiana* requires CaFFN mRNA to reach a certain threshold. Further analyses are needed to clarify the reason for the different expression levels of 35S::CaFFNm3 plants. In contrast to 35S::AP2m3 plants with numerous petals, stamens, and carpels, these two 35S::CaFFNm3 plants exhibited short filaments, poorly developed petals, and a sterile phenotype. These results demonstrated that
the CaFFN gene has a flowering repressor function as an AP2 gene in Arabidopsis and likely results in differentiation in the floral patterning of pepper.

In Arabidopsis, AP2 is involved in various developmental processes at the shoot apex. The expression patterns of AP2 and CaFFN in SAM were consistent with their function as flowering repressors, which showed a high expression level at the vegetative meristem stage and a low expression level at the transition meristem stage. Similar expression patterns were found for nine additional homologs in Arabidopsis and tomato, indicating that these genes may have similar/overlapping functions. Previous research has demonstrated that six AP2 domain-containing genes, including AP2, showed functional redundancy of their role in flowering time in Arabidopsis; the early flowering phenotype of the ap2 mutant was largely obscured by the effects of the other five members of the euAP2 lineage. In contrast, all of the CaFFN mutants in C. annuum, including E-62, B9431, B33, B35, and A100, exhibited an obvious early flowering phenotype, which indicated that the flowering time control function of this gene had little redundancy with other genes in C. annuum. This result was further confirmed by the expression patterns of CaFFN/AP2 homologs in the SAM of C. annuum, which differed from that of most of the detected CaFFN/AP2 homologs (Fig. 6), representing a functional differentiation of these homologs. On the other hand, no floral patterning defects were discovered in E-62, B9431, B33, B35, and A100, indicating that the function of floral patterning regulation of the CaFFN gene was redundant and could be compensated by other gene(s) in pepper.

Capsicum annuum, as a paralog of CaFFN that has arisen from an ancient WGT (~47.31 Mya), might have some functional redundancy with CaFFN since it presented a similar expression pattern as CaFFN in the SAM of pepper (Fig. 6). The possible function of Capsicum annuum in flower development and transcriptional repression of flowering time was also reported in a previous study. However, Capsicum annuum 02g000700 only may have a weak effect on flowering repressor in C. annuum, since its expression level in the early flowering inbred line CA1 (C. annuum) was much lower than that in the late flowering inbred line 740 (C. chinense). This is a reasonable explanation for why the early flowering phenotype of the CaFFN mutants in C. annuum was not obscured by Capsicum annuum 02g000700.

To investigate how the CaFFN gene participates in the complicated network to affect flowering time and floral organ morphogenesis, the transcription profiles of several key flowering time and organ development genes were assayed by qRT-PCR. The results suggested that CaFFN controls flowering time and floral development by regulating genes that are involved in the aging pathway (Fig. 4b): CaFFN functions as a transcriptional repressor to suppress the expression of the floral activator genes CaAP1, CaSEP3, and CaSOC1 as well as the key floral identity gene CaAG; it also functions as a transcriptional activator to activate the expression of the floral repressor gene CaAGL15. In addition, CaFFN negatively regulated miR172 and positively regulated miR156, as indicated by their expression relationship. Moreover, yeast one-hybrid assays and dual-luciferase reporter assays revealed that CaFFN directly activates the floral repressor gene CaAGL15. These results provide the first insight into the mechanisms employed by CaFFN to regulate flowering time and floral patterning in pepper.

Genome-wide association analysis (GWAS) of 36 agronomic traits, including the FNF trait, in 287 pepper accessions was carried out based on SLAF-seq data, but no peak regions associated with the FNF trait were detected. It is unclear whether the CaFFN gene has a broad effect on flowering time in the pepper population. In our study, CaFFN gene association analysis was performed in 167 pepper inbred lines. A SNP 50 bp upstream of the start codon was significantly associated with the number of FNNs, explaining 14.5% of the phenotypic variation. This result indicated that the CaFFN gene has a broad effect on flowering time in the pepper population. This broad effect is not related to the SNP2T > C mutation at the start codon of CaFFN, which is a rare mutation in the pepper population, but is most likely due to the variation(s) in the 5′-UTR of the CaFFN gene. In addition to the result that the SNP in the 5′-UTR (SNP-50G > A) of CaFFN was significantly associated with the FNF trait, the results that SNP-50G > A significantly correlated with the expression level of the CaFFN gene indicated by qRT-PCR also support the speculation that the 5′-UTR may influence the number of FNNs by affecting the expression of the CaFFN gene in the pepper population. Whether SNP-50G > A is the causative mutation that influences the expression of CaFFN or a SNP tightly linked with the causative mutation needs further research.

The control of flowering time in pepper is an attractive objective for breeding aimed at creating locally adapted cultivars. In pepper, several genes, including CaJ, CaBL, CaS, FA, and CaAP2, were reported to be flowering regulators. Only FA and CaAP2/CaFFN were described as flowering repressor genes, and the effect of the FA gene on flowering time was minor. Despite a certain degree of reduction in shoot and fruit size and the extreme flowering time phenotype in the homozygous mutant of the CaFFN gene, it is still an ideal breeding material for genetic improvement. Mutated CaFFN could significantly reduce the number of FNNs with minimal negative impact on yield in multiple genetic backgrounds when heterozygous, making it very useful for hybrid pepper breeding. This gene was used to select early
maturing pepper varieties in our pepper breeding experiments.

**Materials and methods**

**Plant materials**

The precocious pepper B₉₄₃₁ was crossed with the late-flowering inbred line A₁₄₅ (from the Asian Vegetable Research and Development Center). The F₁ hybrid was selfed to generate a segregating F₂ population. A total of 297 individuals of the F₂ population were used for mapping. The flowering time of pepper was measured by counting the number of leaves on the primary stem between the cotyledon and first flower. All plants were grown in a greenhouse in Nanchang (28°33′N, 115°56′E, China).

**BSR-seq**

BSR-Seq combined bulked-segregant analysis and RNA-Seq and was performed to map loci for target traits. For the genetic analysis, two pools were constructed; each pool contained 30 individuals from the F₂ segregating population with extreme phenotypes. For the phenotyping of the FFN trait, total RNA of leaves at the blooming period was extracted using RNAiso Plus reagent (Takara, Dalian, China). RNA was quantified and assessed using a Qubit Fluorometer (Invitrogen, Carlsbad, USA), NanoDrop spectrophotometer (Thermo Fisher Scientific, Wilmington, USA) and Agilent 2100 bioanalyzer (Agilent Technologies, Palo Alto, USA). Pair-end sequencing was carried out on an Illumina HiSeq X-ten instrument. Clean RNA-seq data were mapped to the C. annuum cv. CM334 genome chromosomes (release 1.55, [http://peppergenome.snu.ac.kr/](http://peppergenome.snu.ac.kr/)) using BWA software. SNPs were screened using SAM tools. The differences in allele frequencies of the 12 pepper chromosomes between the two pools were calculated and plotted.

**SNP marker development and genotyping**

To develop a SNP marker for the CaFFN gene, which was speculated to be the candidate gene controlling flowering time in pepper, the genomic sequences of the CaFFN gene in B₉₄₃₁ and A₁₄₅ were amplified (with primers listed in Supplementary Table S1) and sequenced. The polymorphic SNP (SNP2T > C) at position 2 of the CaFFN gene between the B₉₄₃₁ and A₁₄₅ genotypes was converted into the CAPS marker CSF2 for genotyping. PCR was carried out in B₉₄₃₁, A₁₄₅, the F₁ population, and the F₂ population (297 individuals), and each reaction had a total volume of 10 μl containing 50 ng genomic DNA, 0.2 μM primer FFN1-5UF4, 0.2 μM primer FFN1-eIR2 (Supplementary Table S1) and 5 μl 2×Tag Master Mix (Novoprotein Scientific Inc., Shanghai, China). PCR was performed with the following conditions: denaturing for 1.5 min at 94 °C, followed by 38 cycles of at 94 °C for 20 s, 55 °C for 20 s, and 72 °C for 1 min, and a final elongation at 72 °C for 5 min. The amplified PCR products were digested with the restriction enzyme Tail (Thermo Fisher Scientific, Waltham, USA) directly in a final volume of 10 μl consisting of 0.65 μl 10 × R Buffer, 3.5 μl PCR product, 0.2 μl Tail and 5.65 μl ddH₂O. The mixture was incubated at 65 °C for 15 h and 80 °C for 20 min and subjected to gel electrophoresis on a 2% agarose gel.

**Plant transformation of N. benthamiana**

CaFFN gene overexpression vectors were generated by cloning into the Xhol + Xbal digestion sites of the vector pHellsgate 8. In this study, two different overexpression vectors were constructed: one included the wild-type CaFFN cDNA of A₁₄₅, (35S::CaFFN), and the other included mutant CaFFN cDNA with six mismatches to mir172 that did not change the amino acid sequence (synonymous mutations) (35S::CaFFNm3). The recombinant plasmid was introduced into N. benthamiana by Agrobacterium-mediated transformation as described previously. The plants were screened in medium supplemented with 75 mg/L kanamycin (the negative control could not regenerate) in a plant growth chamber at 24 °C under 16 h light/8 h dark cycles. The transgenic seedlings were identified by the primers FFN-RT-F and HR-gate8-RV (Supplementary Table S1).

**VIGS in pepper A₁₄₅**

For CaFFN silencing analysis, the specific coding regions of the CaFFN gene were used for VIGS vector construction. The sequence specificity was confirmed by genome-wide homologous sequence searching by BLAST against sequences in the Zunla databases ([http://peppersequence.genomics.cn/page/species/blast.jsp](http://peppersequence.genomics.cn/page/species/blast.jsp)). Two fragments of ~200 bp in different regions of the CaFFN gene were inserted into the pTRV2 vector at the EcoRI site to construct two vectors, TRV2::CaFFN-1 and TRV2::CaFFN-2. In addition, a fragment of ~100 bp of the CaPDS gene was inserted into pTRV2 to construct a positive control vector, TRV2::CaPDS. The resulting vectors TRV2::rec and TRV1 were transformed into A. tumefaciens (strain GV3101). A. tumefaciens cells harboring TRV1 and TRV2::CaFFN-1/TRV2::CaFFN-2, TRV2 (as a negative control) or TRV2::CaPDS, as a positive control (resuspended in induction medium at a 1:1 ratio, OD₆₀₀ = 0.6), were coinfiltrated into seedlings of pepper A₁₄₅ at the four-leaf stage. The infiltrated seedlings were grown in a phytotron in the dark for 24 h and then at 20 ± 1 °C under 16 h light/8 h dark cycles with a relative humidity of ~90 ± 5%.

**Quantitative reverse transcription PCR**

To determine the relative transcription levels of selected genes, total RNA was extracted from plant tissue using
RNAiso Plus reagent (Takara, Dalian, China) according to the manufacturers’ instructions. Single-stranded cDNA was synthesized using TransScript® II One-Step gDNA Removal and cDNA Synthesis SuperMix (TransGen Biotech., Beijing, China) for quantitative reverse transcription PCR (qRT-PCR) analysis of target genes or using TransScript® miRNA First-Strand cDNA Synthesis SuperMix (TransGen Biotech., Beijing, China) for qRT-PCR analysis of target microRNAs. Quantitative real-time PCR was performed on a Bio-Rad CFX Connect Real-Time System (Bio-Rad Laboratories, Hercules, USA) using NovoStart® SYBR qPCR SuperMix Plus (Novoprotein Scientific Inc., Shanghai, China). Gene expression was calculated relative to CaGAPDH or CaU6 in pepper or relative to NbbTUB in tobacco by the 2^ΔΔCT method. The amplification program was performed at 95 °C for 2 min, followed by 95 °C for 10 s and 60 °C for 30 s (45 cycles). The primers used for qRT-PCR are listed in Supplementary Table S1. In each case, three technical replications were performed for each of at least three independent biological replicates.

**Yeast one-hybrid (Y1H)**

To investigate the interaction between the CaFFN transcription factor and CaAGL15, the full-length CaFFN open reading frame (ORF) sequence was transformed into the pGADT7 vector to provide the prey protein, and the cis-elements (~2000 bp upstream of the start codon isolated from pepper A145) of the CaAGL15 gene were inserted into the pAbAi vector as bait sequences. The linearized bait plasmid pAbAi-CaAGL15 was transformed into the Y1H Gold yeast strain, and the minimal inhibitory concentration of AbA was determined for the bait strain with the empty pGADT7 vector. Then, the prey plasmid pGADT7-CaFFN was transformed into the bait strain to detect the DNA-protein interaction by incubating them at 30 °C for 3 days on SD/-Leu medium supplemented with the minimal inhibitory concentration of AbA.

**Dual-luciferase reporter assay**

The cis-element of CaAGL15 (~2000 bp upstream of the start codon isolated from pepper A145) was inserted into pGREEN0800-LUC to generate the reporter plasmid pGGreen0800-CaAGL15. The full-length ORF of CaFFN was inserted into the pGreenII 62-SK vector to generate the effector vector. The effector and reporter Agrobacterium cultures were mixed together and infiltrated into N. benthamiana leaves. Empty pGreenII 62-SK (EV) was cotransformed with the CaAGL15 reporter as a no-interaction control. Firefly luciferase and Renilla luciferase were quantified at 3 days after infiltration using Dual Luciferase Reporter Assay System reagents (Promega, Fitchburg, United States). Three biological replicates were conducted for each combination.

**Association analysis of CaFFN genes**

A forward primer FFN-solcs-e4R1 and a reverse primer FFN-solcs-5UF1 and a reverse primer were used to amplify a fragment of the CaFFN gene in 164 inbred lines from core pepper collections. Each amplified fragment was sequenced using the FFN-solcs-5UF1 primer, and the sequences obtained were aligned using MUSCLE. SNPs and InDels were identified, and variants were filtered according to the MAF (MAF ≥ 5%). The variants from 164 pepper inbred lines and FFN phenotypes were used in the association analysis, which was conducted using cMLM in TASSEL v3.0. The population structure (Q matrix) and familial kinship (K matrix) derived from our previous studies were taken into account to avoid spurious associations. The associated variant(s) with the FFN trait were identified at the probability level of 0.01.

**Analysis of AP2 family genes**

The genome database of the cultivated pepper Zunla-1 was used to identify the identification of AP2 family genes. The hidden Markov model (HMM) profile of the AP2 domain (Pfam accession No. PF00847) was used as a query to BLAST against the Zunla-1 protein database. AP2 family members were identified by checking gene structures in Pfam to confirm the presence of two or more AP2 domains. BLASTP alignment (E < 1e−5, top five matches) was carried out across the whole-genome protein sequences, and the output results were loaded into MCScanX (MATCH_SIZE: 5) to identify collinear blocks and to distinguish the singleton, dispersed, proximal, tandem, and WGD/segmental duplication types of these genes. The coding sequences of the AP2 family genes and four additional genes (Capana01g000159, Capana10g001789, Capana03g000560, and Capana10g001776) were used to create a codon alignment by DAMBE. A phylogenetic tree of CaFFN/AP2 homologs was constructed using the maximum likelihood (ML) method in MEGA, and branch reliability was assessed with 1000 bootstrap replications. The Ks (synonymous substitutions per site) values were calculated in PAML. The formula T = Ks/2R was used to estimate the divergence time, where R is 1.5 × 10−8 synonymous substitutions per site per year. The expression data of 19 CaFFN/AP2 homologs were obtained from this study (primers are listed in Supplementary Table S1) and previous studies. The expression data were scaled to construct the heatmap by the Pheatmap package (https://CRAN.R-project.org/package=pheatmap) in R.

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