Mapping and identification of genetic loci affecting earliness of bolting and flowering in lettuce

Leah Rosental1,4 · David W. Still2,3 · Youngsook You3 · Ryan J. Hayes1,5 · Ivan Simko1

Received: 16 April 2021 / Accepted: 25 June 2021 / Published online: 1 July 2021
© This is a U.S. government work and not under copyright protection in the U.S.; foreign copyright protection may apply 2021

Abstract

Key message Photoperiod and temperature conditions elicit different genetic regulation over lettuce bolting and flowering. This study identifies environment-specific QTLs and putative genes and provides information for genetic marker assay.

Abstract Bolting, defined as stem elongation, marks the plant life cycle transition from vegetative to reproductive stage. Lettuce is grown for its leaf rosettes, and premature bolting may reduce crop quality resulting in economic losses. The transition to reproductive stage is a complex process that involves many genetic and environmental factors. In this study, the effects of photoperiod and ambient temperature on bolting and flowering regulation were studied by utilizing a lettuce mapping population to identify quantitative trait loci (QTL) and by gene expression analyses of genotypes with contrasting phenotypes. A recombinant inbred line (RIL) population, derived from a cross between PI 251246 (early bolting) and cv. Salinas (late bolting), was grown in four combinations of short (8 h) and long (16 h) days and low (20 °C) and high (35 °C) temperature. QTL models revealed both genetic (G) and environmental (E) effects, and GxE interactions. A major QTL for bolting and flowering time was found on chromosome 7 (qFLT7.2), and two candidate genes were identified by fine mapping, homology, and gene expression studies. In short days and high temperature conditions, qFLT7.2 had no effect on plant development, while several small-effect loci on chromosomes 2, 3, 6, 8, and 9 were associated with bolting and flowering. Of these, the QTL on chromosome 2, qBFr2.1, co-located with the Flowering Locus T (LsFT) gene. Polymorphisms between parent genotypes in the promotor region may explain identified gene expression differences and were used to design a genetic marker which may be used to identify the late bolting trait.

Introduction

The phase transition from vegetative to the reproductive stage is a critical step in the life cycle of flowering plants that affects the successes of flowering, fruit set, and seed production. In lettuce (Lactuca sativa L.), the transition from the vegetative to reproductive stage occurs in the meristem concealed within the rosette. During the transition, the vegetative shoot apical meristem cap is elongated, and then, the microscopic floral primordia are formed (Chen et al. 2018a, b). Subsequently, rapid elongation of the stem internodes occurs, a process called bolting, and the inflorescence expands followed by flowers opening over the course of several days. Since lettuce is a crop grown for its leaf rosettes, early bolting is an undesirable trait. Determining the genetic and environmental factors affecting bolting would increase the ability to develop cultivars with greater resistance to premature bolting.
The vegetative–flowering transition is regulated by a complex network of genetic and environmental factors. In the well-studied Arabidopsis (*Arabidopsis thaliana* (L.) Heynh) model, more than 100 genes have been implicated in the control of flowering time (Komeda 2004; Putterill et al. 2004; Amasino 2010; Bouché et al. 2016). Internal regulatory pathways converge with pathways integrating environmental signals to control the timing of meristem differentiation. Endogenous factors influence floral induction and include the gibberellin (GA) pathway, circadian clock, and plant age (Fornara et al. 2010; Song et al. 2015; Bao et al. 2020). Environmental cues influencing flowering time include day length, light quality, ambient temperature, and vernalization (Michaels and Amasino 2000; Samach and Coupland 2000; Blázquez et al. 2003; Song et al. 2013; Romera-Branchat et al. 2014; Capovilla et al. 2015; Cho et al. 2017).

Lettuce is a facultative long-day plant, and cultivars and accessions exhibit wide variability in the rate of transition to flowering (Ryder 1983; Lafta et al. 2017). Very early cultivars (Ryder 1996) start to bolt and flower seven weeks after planting in long days in a greenhouse while very late cultivars may flower as late as four months after planting. Besides genetic factors, environmental factors substantially affect the rate of transition to the reproductive phase (Ito et al. 1963; Jenni et al. 2013). Earlier bolting (less than 90 days) can be induced in certain genotypes by increased day length (Waycott 1995) or by high ambient temperatures (Rappaport et al. 1956), thus leading to a strong genotype x environment (GxE) interaction (Lafta et al. 2017; Wien 2020).

For lettuce production, a prolonged vegetative phase is beneficial for accumulation of leaf biomass and to maintain quality. Premature bolting results in elongated cores and/or cracked heads that are not suitable for the market. In addition, the initiation of flower signaling causes biochemical changes to form accumulation of latex in the leaves which makes them undesirably bitter (Simonne et al. 2002). Premature bolting can lead to partial or complete yield loss, particularly in early fall planting periods in hot production regions. Therefore, the genetic basis for regulation of bolting and flowering has been the focus of numerous studies over the years (Thompson 1943; Rappaport et al. 1956; Ryder 1983; Ryder and Milligan 2005; Han et al. 2021).

Ryder (1983, 1988) identified two genes controlling early flowering (*Ef*) *Ef-I* and *Ef-2*, with earliness being incompletely dominant. The author later described four more *Ef* genes in additional crosses (Ryder and Milligan 2005). More recently, multiple quantitative trait loci (QTLs) for bolting and flowering time, as well as for stem length, which correlates with earliness of bolting, have been identified on lettuce chromosomes 2 and 7 (Jeukens and Lindhout 2004; Hartman et al. 2012, 2013; Han et al. 2021) in crosses between cultivated lettuce and its wild relatives. In a recombinant inbred line (RIL) population developed from crossing late and early bolting lettuce cultivars and evaluated in four summer experiments, three QTLs were identified on chromosomes 2, two QTLs on chromosome 5, and one each on chromosomes 7 and 8 (Jenni et al. 2013). Linkage mapping performed on another RIL population developed from crossing two cultivars with similar time to bolting revealed QTLs on chromosomes 1, 4, and 5 (Mamo et al. 2019). Perhaps due to the polygenic nature of the trait, transgressive segregation was observed in several bi-parental populations (Silva et al. 1999; Jenni et al. 2013; Mamo et al. 2019). A genome-wide association study (GWAS) identified several single nucleotide polymorphism (SNP) markers linked to bolting (on chromosomes 1, two on 7, and one that was not mapped) and flowering (on chromosomes 5, 7, 8 and one that was not mapped; Kwon et al. 2013). Although bolting and flowering are closely related processes, mapping of each phenotype separately resulted in only partial overlap of the associated SNPs. Another GWAS study performed on ~500 accessions identified a major QTL for developmental rate on chromosome 7, while minor QTLs with substantial QTL x environment interaction were detected on most of the other chromosomes (Søhatip Kandel et al. 2020).

In lettuce, the homolog of the Arabidopsis *FLOWERING LOCUS T* (*FT*) gene had only one copy and *LsFT* expression levels increased with higher ambient temperature and induced earlier differentiation to flowering (Fukuda et al. 2011). The response of *LsFT* to heat treatment was also characterized in *FT* RNAi plants (Chen et al. 2018a). In these lines, there was also a reduction in expression of putative downstream genes (based on homology to Arabidopsis *AP2*, *AP3*, and *LFY*). Another study showed association between the bolting time and the expression levels of additional Arabidopsis homologs in lettuce: *LsFVEL*, *LsFLDL*, *LsLDL*, *LsLFY*, *LsAP1L* (Fukuda et al. 2017). The lettuce homolog of *AtSOC1* (*LsSOC1*), an important floral integrating gene, was found to be expressed in lettuce shoot apical meristems during the reproductive phase. Silencing of this gene greatly delayed flowering (Chen et al. 2018a, b). *LsSOC1* was also shown to greatly increase upon heat treatment which promoted flowering. Han (2016) found many genes that were differentially expressed between two lettuce cultivars with different bolting time. The authors implicated a group of MADS-box and GA genes as flowering regulators (Ning et al. 2019). GA and related genes were shown to be related to bolting (Fukuda et al. 2009; Umetsu et al. 2011) specifically in response to high temperatures (Liu et al. 2018). The similarities in expression and function of these central genes suggest the main structure of the signaling pathway to be conserved between lettuce and Arabidopsis. There are many examples where homologs of Arabidopsis genes function differently in crop species (Blümel et al. 2015) and
paralogs that broadened natural variation in flowering time (Blackman et al. 2010). In other cases, several genes have been found in crops that have no Arabidopsis homologs. Therefore, focusing solely on Arabidopsis homologues could lead to identifying new players or overlook the central genes that are involved in regulation of bolting and/or flowering in lettuce.

While studying functions of gene homologs and molecular genetic manipulation sheds light on the molecular mechanism of flowering in lettuce, their use for traditional breeding is limited. For conventional breeding that uses crossing and selection, information about beneficial alleles that exist in plant accessions, including marker-assisted selection (MAS) (GuoYou et al. 2010; Simko 2013), is most needed.

To identify novel genes for lettuce flowering time, we used a recombinant inbred lines (RIL) population segregating for flowering time. The RIL population was created from a cross between PI 251246 and cv. Salinas (Yoong et al. 2016). Parental line PI 251246 is a primitive type lettuce accession cultivated for seed oil that skips the rosette stage and flowers within six weeks after germination (Grube and Ryder 2004). Parental line cv. Salinas, an iceberg type, is relatively late to bolting, 95 days under long-day conditions and ~132 days under short-day conditions (Waycott 1995).

The wide range of flowering phenotypes and environmental sensitivities observed in the RIL population, and careful evaluation in controlled environments, revealed multiple environment-specific QTLs and interactions among them. Fine mapping in conjunction with gene sequencing and expression studies allowed identification of candidate genes and the development of a MAS assay that can be used by breeders to confer bolting resistance.

Material and methods

RIL population, plant growing conditions, and phenotyping of phenological stages

A F7 RIL population, consisting of 161 families, was evaluated under controlled environmental conditions. The RIL was derived from single seed descent following a cross between PI 251246 and cv. Salinas. For the first 14 days, plants were grown in controlled environment chambers under a photoperiod regime consisting of 8 h light and 16 h dark at 20 °C for 14 days to allow seedling establishment. Light was provided by a combination of fluorescent and incandescent electric bulbs. After 14 days, plants were transferred to growth chambers under four light and temperature conditions: long day and low temperature (LD-LT; 16 h light/8 h dark at 20 °C), long days and high temperature (LD-HT; 16 h light/8 h dark at 35 °C), short days and low temperature (SD-LT; 8 h light/16 h dark at 20 °C), and short days and high temperature (SD-HT; 8 h light/16 h dark at 35 °C). In each of these four environmental conditions, the phenological stage of three plants per RIL and both parents was evaluated weekly using a rating scale of 1 = rosette; 2 = bolting-visible internode elongation; 3 = visual buds; 4 = expanded inflorescence; 5 = flowering–opening of first flower; 6 = more than half of buds flowered; 7 = open involucres. Also evaluated was the number of days to bolting ($D_{BLT}$; days after germination to reaching stage 2) and days to flowering ($D_{FLT}$; days after germination to reaching stage 5). The population-wide response to environmental conditions was statistically evaluated with two-tailed Student’s t-test using the population mean and the variance that were calculated from the values of all RILs grown at each environment. The experiment-wise p-value lower than 0.05 was considered statistically significant; thus, the test-wise error rate was adjusted for multiple comparisons using Bonferroni correction. It has been shown that $D_{BLT}$ and $D_{FLT}$ share both common and unique pathways (Kwon et al. 2013). To potentially identify regulation of flowering distinct from bolting, the residuals from the linear regression of $D_{FLT}$ to $D_{BLT}$ ($D_{BF}$) were calculated for each RIL and used alongside other traits as an independent phenotype for QTL analysis.

Genotyping and linkage map construction

DNA was extracted from each RIL family (DNeasy Plant Mini Kit, Cat No. 69104, Qiagen Inc., Valencia, CA), digested with the EcoT22I enzyme, and subjected to genotyping by sequencing (GBS; Elshire et al. 2011) by the Cornell Institute of Biotechnology (http://www.biotech.cornell.edu/brc/genomics-facility). SNP calling and filtering were performed using the TASSEL SNP discovery pipeline (Glaubitz et al. 2014). A linkage map based on informative markers was generated using an online version of MSTmap (http://mstmap.org/). Fully linked markers, with no recombination between them, were collapsed to a single combined marker. The linkage map for QTL analysis was re-generated to verify the correct placing and centimorgan (cM) distance determination of the combined markers and the original ones.

QTL mapping and model selection

QTL mapping and model selection were performed in R using the R/qtl package (Broman et al. 2003). Simple interval QTL mapping (SIM) was performed using the Haley-Knott regression method (Feenstra et al. 2006). The significance LOD threshold for each environmental condition was determined by 1,000 permutations. QTLs were deemed significant when exceeding $\alpha=0.05$ LOD threshold and suggestive when exceeding $\alpha=0.1$ LOD threshold. QTL naming followed the nomenclature convention using $q+trait$
abbreviation + chromosome number + QTL number on the particular chromosome. Trait abbreviations followed those suggested by Han et al. (2021).

All significant and suggestive QTLs were considered in the model fitting for $D_{BLT}$ and $D_{FLT}$ in each environmental condition separately. Initially, additive and interactive effects of all QTLs were considered in the model. Subsequently, a backward selection was applied to eliminate QTLs out of the model when they had no significant additive effect or interaction. The Bayesian information criterion (BIC; Broman and Speed 2002) with a $\delta$ value of 2 to minimize false positives was calculated for each evaluated model. Finally, the best model was selected based on the lowest BIC, i.e., the model explaining the highest variance with the least number of parameters.

**Fine mapping**

To enable fine mapping of the $qBLT7.2$—$qFLT7.2$ QTL region, an $F_2$ population was developed from a cross between two RILs (RIL 68 and RIL 69) with high similarity of marker alleles across the whole genome, but harboring contrasting marker alleles in the region of interest.

Genomic polymorphisms between RIL 68 and RIL 69 in the region of interest were identified by SNP mining of RNA-seq data generously provided by Fei-Yian Yoong (Yoong et al. 2016). To identify additional polymorphisms, Sanger sequencing was performed (Eton Bioscience Inc., San Diego, California) on PCR amplification products that targeted genes and inter-gene segments within the region of interest. Polymorphisms identified along the $qBLT7.2$—$qFLT7.2$ region were used to design high-resolution DNA melting (HRM) markers (Simko 2016; Supplementary Table 1). Testing for recombination and genotyping of all recombinants with the HRM markers was performed with the LightScanner instrument (BioFire Diagnostics LLC, Salt Lake City, Utah). PCR conditions and florescent detection were according to manufacturer’s instructions. Specifically, PCR annealing temperature was 66 °C, and elongation temperature was held at 72 °C for 30 s and LightScanner temperature scan ranged between 72 and 96 °C.

To identify recombinants, 2,094 $F_2$ plants, all derived from a single $F_1$ plant from the RIL 68 × RIL 69 cross, were evaluated. Of these, 773 plants were found to have recombination between two co-dominant markers flanking the QTL region. Recombinant plants were genotyped with 18 additional markers, designed throughout the $qBLT7.2$ / $qFLT7.2$ region. The phenological development of recombinants was evaluated weekly on plants grown under long days and mild temperature (18–25 °C) conditions in which the parental RILs (RIL 68 and RIL 69) significantly differed in their flowering time. Differences between parent genotypes and $F_2$ plants used in the fine mapping study were statistically evaluated with two-tailed Student’s $t$-test using the mean and the variance calculated from the values of at least five plants. The experiment-wise $p$-value lower than 0.05 was considered statistically significant; thus, the test-wise error rate was adjusted for multiple comparisons using Bonferroni correction.

$qBFr2.1$ allele effect validation

In order to validate the effect of the $P$ allele (originating from PI 251246 parent) on flowering time, $F_3$ plants from the RIL 68 × RIL 69 cross were evaluated. Plants selected by HRM markers were homozygous for $P$ alleles in $qBFr2.1$ and for $S$ alleles (originating from cv. Salinas parent) in other detected $D_{FLT}$ QTLs. For further validation, a series of back crosses of the $F_3$ plants to cv. Salinas was conducted, selecting for the $P$ allele with the HRM marker LZ001. $F_3$ and backcross plants were evaluated for $D_{FLT}$ in the greenhouse during spring under ~LD-LT conditions.

**Gene expression assays**

To examine the range of gene expression responses of flowering related genes, the RIL parental lines (PI 251246 and cv. Salinas, along with the RILs used for fine mapping, RIL 68 and RIL 69, were used. The plants were grown under the four photoperiod and temperature conditions (LD-LT, LD-HT, SD-LT, SD-HT) and evaluated weekly using the phenological rating scale. In previous experiments a ~14-day difference between appearance of a first bud and a first flower was consistently observed. Furthermore, the appearance of the bud demonstrated that the transition from a vegetative to a reproductive phase occurred. Therefore, in this experiment $D_{BUD}$ (days after germination to reaching stage 3) was used to quantify developmental progress. The response to environmental conditions of the genotypes used in the gene expression study was statistically evaluated with two-tailed Student’s $t$-test using the mean and the variance calculated from the values of at least nine plants. The experiment-wise $p$-value lower than 0.05 was considered statistically significant; thus, the test-wise error rate was adjusted for multiple comparisons using Bonferroni correction.

At selected time points, leaf and stem tip samples were harvested for RNA extraction. Leaf samples were taken from the distal part of the second-to-newest fully expanded leaf in the rosette. Stem tip samples included the apical meristem with minimal wrapping of young leaves. To improve uniformity of results, all samples were collected between hour two and three of the growth chamber light period. For each sample, three independent pools of tissue were collected from three individual plants and flash-frozen in liquid nitrogen. Total RNA was extracted from each sample using

© Springer
the RNeasy Plant Mini Kit (Cat No. 74904, Qiagen Inc., Valencia, CA)) according to the manufacturer's instructions.

A total of 31 candidate genes within the qBLT7.2 / qFLT7.2 QTL region were identified using the lettuce reference genome (ver. 4; Reyes-Chin-Wo et al. 2017). Genes whose predicted function was deemed unlikely to be involved in reproductive transition regulation were excluded from further consideration, resulting in 12 candidate genes retained. To better understand the effects of the QTL alleles on the reproductive transition regulatory network, 15 genes known to be involved in flowering time regulation in Arabidopsis were tested for expression (Supplementary Table 2). Of these, eleven have been identified in lettuce (Han et al. 2016; Huo et al. 2016; Ning et al. 2019) including MADS-box genes. The MADS-box genes assessed in this work are arbitrarily named MAD1-7 because of their high sequence homology. Additionally, three normalization genes (PP23, PP21, and TIP4) were chosen from the literature based on their low constitutive expression in various tissues and in different experimental conditions. PP23, PP21, and TIP4 encode for Protein Phosphatase 2a subunit 3, Protein phosphatase 2A-1, and TIP41-like protein, respectively (Czechowski et al. 2005; Borowski et al. 2014; Sgamma et al. 2016).

The GenomeLab GeXP Analysis System (SCIEX, USA) was used to assay mRNA levels in tissues with multiplex primer design (Supplementary Table 2). Gene expression primers for multiplexed reactions were designed using GenomeLab eXpress Profiler software. The multiplexed expression panel consisted of 27 candidate genes plus the three normalization genes. Complementary primer binding sites for both genotypes were verified by sequencing. The cDNA for GeXP was synthesized from 50 ng of total RNA using the GenomeLab GeXP Start Kit. PCR and multiplex detection were performed according to manufacturer’s instructions (Hayashi et al. 2007). Gene expression was quantified as area under the peak. All peaks under a background noise threshold of 500 counts were removed, and remaining peaks were normalized to the intensity of the internal control Kan’ gene. Since TIP4 was not stably expressed in the experiment, in order to select optimal genes for normalization, the full panel of 30 genes was subject to an expression stability test following the GeNorm algorithm (Vandesompele et al. 2002). The geometric mean of expression levels of the selected three most stable genes, PP21, PP23, and 7b_8, was used for normalization of expression levels of all target genes.

Lactuca sativa FLOWERING LOCUS T gene expression was independently evaluated by RT- real-time PCR, using the same RNA samples as were used for the GeXP analysis. Reverse transcription was performed using 170 ng of total RNA with the High-Capacity RNA-to-cDNA™ Kit (Cat no: 4387406, Thermo Fisher Scientific Inc., Foster City, California). The normalization gene PP23 was selected for real-time PCR based on having lowest variation in the GeXP panel. Target and normalization genes were detected in the same well by a duplex TaqMan reaction using reporter dyes FAM and VIC, respectively, and custom-designed primers and probes (Supplementary Table 2), with an annealing temperature of 60 °C (Custom TaqMan® Gene Expression Assays, Thermo Fisher Scientific Inc., Foster City, California). Color compensation to account for differences in using different dyes was applied to the raw data. Dose curve experiments revealed amplification efficiency for both genes and genotypes ranging from 1.98 to 2.02; thus, an efficiency of 2 was used for calculation of target to reference concentration ratio (conc. ratio = 2^((Cp target−Cp ref)). Each sample was tested in triplicate, and their average was used for further statistical analysis.

Gene expression differences between the genotypes and controlled environment condition were compared using Tukey’s Honest Significant Difference (HSD) test. For comparison of environmental condition effect on gene expression, values of the four environmental conditions at a specific time point were compared by Tukey’s HSD test for each genotype. For comparisons of genotype effect on gene expression, values of the four genotypes at a specific time point were compared by Tukey’s HSD for each condition. An indication of a significant increase or decrease was made by Student’s t-test, comparing gene expression values of the specific genotype and environmental condition between the time points described in the text. Significance was determined at α = 0.05 corrected for the number of comparisons in each time point.

**Candidate gene homology, sequencing, and polymorphism annotation**

Candidate genes were identified by positioning GBS markers to the lettuce reference genome (Reyes-Chin-Wo et al. 2017) and using blastp to search the Arabidopsis Information Resource (TAIR) protein database with the predicted lettuce protein sequence. Reciprocally, the protein of the best Arabidopsis match was used to search the lettuce TSA database in NCBI using tblastn (McGinnis and Madden 2004).

Candidate genes and their promotor regions were sequenced using PCR amplification of overlapping gene segments. The PCR products were purified and subjected to Sanger sequencing (Eton Bioscience Inc., San Diego, California). To better understand the biological significance of differences discovered in promotor regions, an in silico annotation based on known Arabidopsis promotor motifs was performed using AgRIS (Davuluri et al. 2003) and PLACE (Higo et al. 1999).
Results

QTLs associated with day length and temperature

Under all environmental conditions, $D_{BLT}$ were lowest for PI 251246 and highest for cv. Salinas with no transgressive segregation observed among RILs. In contrast, transgressive segregation for $D_{FLT}$ was observed with several RIL families displaying both earlier and later $D_{FLT}$ compared to PI 251246 and cv. Salinas, respectively (Fig. 1).

Both $D_{BLT}$ and $D_{FLT}$ of RILs grown under HT were significantly shorter ($p < 0.0001$, $t$-test) than under LT (Table 1). Day length did not significantly affect $D_{BLT}$, but $D_{FLT}$ was significantly shorter ($p < 0.0001$, $t$-test) in LD compared to SD. The range of days between bolting and flowering also differed among the four environmental conditions, and both HT and LD significantly ($p < 0.0001$, $t$-test) reduced the observed range. These results support the idea that $D_{BLT}$ and $D_{FLT}$ share both common and unique pathways (Kwon et al. 2013; Chen et al. 2018b) and hence support the use of residuals of the linear correlation between them, $D_{BFr}$, as an additional phenotypic trait.

QTL analysis using the RIL population revealed a total of two bolting QTLs across the four growing conditions, accounting for 13.6–42.9% of the total phenotypic variance explained (PVE) (Fig. 2, Table 2). One significant flowering QTL was identified, which accounted for 24.0–51.9% of the PVE in various conditions. In addition, three QTLs were identified for $D_{BFr}$, which accounted for 13.0–37.4% of the PVE.

A major QTL for $D_{BLT}$, $qBLT7.2$, was found under LD-LT conditions while the same locus showed only a minor effect at LD-HT and SD-LT and no observable effect at SD-HT conditions. This locus also had a large effect on $D_{FLT}$, $qFLT7.2$, in LD conditions under both high and low temperature (PVE = 52.0% and 47.3% under LD-LT and LD-HT, respectively), but had a lower effect under SD-LT conditions (PVE = 24.1%). The QTL effect on $D_{BFr}$ was relatively small under LD-LT and SD-LT conditions (13% and 9% of PVE, respectively), but high under LD-HT (PVE = 37.5%). These results show that under LD conditions alleles at this locus affect bolting and flowering differently depending on the ambient temperature.

Three other QTLs associated with $D_{BLT}$, $D_{FLT}$ and $D_{BFr}$ were identified, one on chromosome 8 ($qBLT8.1$), one each on chromosome 2 ($qBFr2.1$), and chromosome 6 ($qBFr6.1$). The allele contributing to the delayed flowering phenotype at $qBFr2.1$ originated from the early flowering parent PI 251246.

Three additional locations contain ‘suggestive’ QTLs (0.1 ≥ $\alpha$ ≥ 0.05, Table 1). Under LD-LT, a suggestive QTL ($sqBLT7.1$) was detected on chromosome 7. Under

Fig. 1 Distribution of $D_{BLT}$ in the RIL population grown in four controlled conditions: LD-HT, LD-LT, SD-HT, and SD-LT. $D_{BLT}$ of population parents, cv. Salinas (Sal), and PI 251246 (PI) in each condition are marked by vertical line if evaluated.
The early flowering parent, PI 251246, contributed alleles for delayed flowering for the latter two QTLs.

QTL models

To identify QTLs that explained the highest amount of phenotypic variation for $D_{BLT}$ and $D_{FLT}$, Bayesian information criterion were used (Broman and Speed 2002). The PVE explained by the models ranged from 13.5% to 83.8%, of a single QTL model for bolting under SD-HT and a five QTL model for flowering under LD-LT, respectively (Table 3). An identical five QTL model was the best fit for both flowering under LD-HT and LD-LT; all other conditions tested had unique models (Table 3, Supplementary Table 3).

The models show that QTLs which were detected only at a suggestive (0.1 ≥ $\alpha$ ≥ 0.05) level using SIM can have a significant effect on flowering through interaction with other QTLs (i.e., epistasis). Furthermore, the extent of their effect and interaction is affected by the light and temperature conditions. The most obvious condition-dependent interaction was seen between $qFLT2.1$ and $qFLT7.2$ (Fig. 3), where at LD-HT and SD-LT alleles in these positions have constant effects, while at SD-HT only $qFLT2.1$ alleles affect $D_{FLT}$.

At LD-LT, the $qFLT2.1$ alleles have an effect on flowering only when the homozygous combination of alleles originating from cv. Salinas is present at the $qFLT7.2$ locus, thus demonstrating epistatic effect. In this population, under all environmental conditions, the alleles maintained directionality of their effect on bolting and flowering.

Fine mapping of $qBLT7.2$ / $qFLT7.2$ region

Under field and controlled environment studies, $qBLT7.2$ / $qFLT7.2$ was consistently observed to have the largest PVE of all QTLs detected. Therefore, this locus was selected for

---

Table 1: Descriptive statistics of the RIL population for $D_{BLT}$ and $D_{FLT}$ grown under four different controlled environment conditions

| Day length | Temperature | Condition abbreviation | $D_{BLT}$ Range | Mean | SD | $D_{FLT}$ Range | Mean | SD | $D_{BFr}$ Range | Mean | SD |
|------------|-------------|------------------------|-----------------|------|----|-----------------|------|----|-----------------|------|----|
| Long       | High        | LD-HT                  | 13–27           | 20.6 | 3.8| 34–111          | 58.8 | 19.9| 14–84           | 38.3 | 17.6|
| Long       | Low         | LD-LT                  | 20–40           | 25.3 | 6.8| 41–153          | 73.2 | 26.7| 21–119          | 43.8 | 20.7|
| Short      | High        | SD-HT                  | 13–48           | 23.3 | 7.9| 41–111          | 69.6 | 16.1| 7–91            | 43.9 | 14.0|
| Short      | Low         | SD-LT                  | 20–41           | 24.6 | 6.3| 69–160          | 100.3| 26.2| 35–133          | 76.2 | 24.1|

SD—standard deviation

$^a$ $D_{BLT}$—days after germination to bolting

$^b$ $D_{FLT}$—days after germination to first flower

$^c$ $D_{BFr}$—the residuals from the correlation between bolting and flowering, representing the degree by which flowering is regulated independently of bolting
further study. $F_2$ plants with recombination in the $qBLT7.2$ / $qFLT7.2$ region were grown in the greenhouse and evaluated for bolting and flowering. Under greenhouse conditions, the parents of the $F_2$ cross, RIL 68 and RIL 69, significantly differed ($p < 0.001$, $t$-test) in flowering time by an average of 40 days. Of 21 markers tested (Supplementary Table 1), five closely linked markers, LZ431, LZ481, LZ261, LZ299, and LZ425 displayed complete association with the flowering phenotype (Fig. 4). The phenological stage of $F_2$ plants heterozygous in these markers was significantly higher ($p < 1e^{-7}$, $t$-test) than that of plants with homozygous S allele in all closely linked markers. The QTL region of interest was re-defined based on these markers and included a 1.15-Mbp-long region (chromosome 7: 207,702,411—209,071,520 bp) which contained 31 predicted genes. Genes whose function, predicted by Arabidopsis homology annotations (Reyes-Chin-Wo et al. 2017), was unlikely to be involved in development regulation (such as glycolysis or chloroplast targeted genes) were not pursued for further analysis.

### Table 2 QTLs for $D_{BLT}$, $D_{FLT}$ and $D_{BFr}$ detected in PI 251246 × cv. Salinas RIL population

| Day length | Temperature | QTL     | Chr | Peak LOD location (cM) | 1-LOD support interval (cM) | Marker closest to peak LOD | Peak LOD | PVE$^b$ | Allel additive effect$^c$ (days) |
|------------|-------------|---------|-----|------------------------|-----------------------------|----------------------------|----------|--------|---------------------------------|
| Long       | High        | $qBLT7.2$ | 7   | 106.5                  | 86.7–106.5                  | cS7_242106278               | 3.3       | 10.7   | 1.2                             |
|            |             | $qFLT7.2$ | 7   | 94.1                   | 93.2–98.3                   | cS7_209035609               | 15.5      | 47.2   | 13.0                            |
|            |             | $qBFr7.2$ | 7   | 94.1                   | 93.2–98.3                   | cS7_209035609               | 11.1      | 37.4   | 10.2                            |
| Low        |             | $sqBLT7.1$| 7   | 0                      | 0–5                         | cS7_1980750                 | 2.8       | 10     | 2.2                             |
|            |             | $qBLT7.2$ | 7   | 94.1                   | 93.2–98.3                   | cS7_209035609               | 16        | 42.9   | 4.4                             |
|            |             | $qFLT7.2$ | 7   | 94.1                   | 93.2–98.3                   | cS7_209035609               | 18.8      | 51.9   | 18.7                            |
|            |             | $qBFr7.2$ | 7   | 94.1                   | 79.3–98.3                   | cS7_209035609               | 3.4       | 13     | 6.2                             |
| Short      | High        | $qBFr2.1$ | 2   | 18.1                   | 15.6–26                     | cS2_40045553                | 3.4       | 14.4   | -5.3                            |
|            |             | $sqBFr3.1$| 3   | 57.7                   | 53–63                       | cS3_113143153               | 2.7       | 9.6    | -4.4                            |
|            |             | $qBFr6.1$ | 6   | 100.3                  | 97.5–104.4                  | cS6_220422004               | 3.7       | 13.8   | 5.4                             |
|            |             | $qBLT8.1$ | 8   | 34.6                   | 28.9–38.5                   | S8_71245709                 | 4.7       | 13.6   | 2.9                             |
|            |             | $sqFLT9.1$| 9   | 17.9                   | 17–24                       | cS9_31330927                | 3         | 11.7   | -5.3                            |
| Low        |             | $qBLT7.2$ | 7   | 94.1                   | 91.9–98.3                   | cS7_209035609               | 5.9       | 20.2   | 2.7                             |
|            |             | $qFLT7.2$ | 7   | 91.9                   | 87.5–93.2                   | cS7_200076206               | 4.9       | 24     | 13.6                            |

$^a$Suggestive QTLs (sq) are significant at $\alpha = 0.1$ (determined by 1000 permutations)

$^b$PVE—percent of variance in mapped phenotype explained by the QTL

$^c$directionality of effect: positive allele contributed by cv. Salinas; negative allele contributed by PI 251246

### Table 3 Genetic models to explain the $D_{BLT}$ and $D_{FLT}$ phenotype under each environmental condition

| Trait       | Day length | Temperature | Model LOD | PVE$^a$ | $p$-value model | Positions included in model$^b$ |
|-------------|------------|-------------|-----------|---------|-----------------|--------------------------------|
| Bolting     | Long       | High        | 16.2      | 46.7    | 8.88E-04        | p2.1×p3.1×p6.1×p7.2×p8.1+p9.1$^c$ |
|             | Low        | High        | 17.3      | 47.5    | 1.11E-16        | p7.1×p7.2                     |
|             | Low        | Low         | 3.7       | 13.6    | 3.53E-05        | p8.1                           |
|             | Low        | Low         | 16.5      | 46      | 4.03E-04        | p2.1×p6.1×p7.2×p8.1×p9.1      |
| Flowering   | Long       | High        | 39.4      | 83.1    | <2.00E-16       | p2.1×p6.1×p7.1×p7.2×p9.1     |
|             | Low        | High        | 45.1      | 83.8    | <2.00E-16       | p2.1×p6.1×p7.1×p7.2×p9.1     |
|             | Low        | Low         | 23.0      | 61.1    | 2.45E-07        | p2.1×p3.1×p6.1×p7.1×p9.1     |
|             | Low        | Low         | 13.4      | 77      | 4.93E-05        | p2.1×p6.1×p7.2×p9.1           |

$^a$PVE—percent of variance in mapped phenotype explained by the complete model

$^b$Positions of markers included in the models correspond to QTL names found in Table 1. All QTLs were initially included in the full model followed by removal of the QTL with the smallest contribution to the model (backward elimination)

$^c$Positions where the interaction among QTLs was significant are marked with ‘x’, while positions preceded by + have an additive effect
Locus qBLT7.2 / qFLT7.2 candidate genes, homology, and sequencing

The predicted annotation of twelve genes further studied in the fine mapped region was verified by Arabidopsis protein homology (Supplementary Table 2), and three genes were found homologous to genes known to be involved in flowering time regulation.

CONSTANS LIKE-9 (COL9) was found to be the best matching homolog for the Lsat_1_v5_gn_7_97240.1 lettuce gene. In Arabidopsis, COL9 delays flowering time and is a FT, SOC1, and CO repressor (Cheng and Wang 2005). The full coding region and part of the UTRs of COL9 in cv. Salinas and PI 251246 were sequenced using both genomic and cDNA. A single polymorphism between sequences of the two genotypes was found in the fourth intron, but none in the coding region. The COL9 gene-specific marker used for fine mapping, LZ431, mapped to the center of the QTL region and matched exactly with the flowering time phenotype (Fig. 4d).

PHYTOCHROME C (PhyC) was found to be the best match for Lsat_1_v5_gn_7_96941.1. PhyC encodes a phytochrome protein that is known to be involved in flowering time regulation in Arabidopsis (Pearce et al. 2016). Sequencing of PhyC revealed six SNP polymorphisms in the coding sequence between cv. Salinas and PI 251246, all in the first exon. A gene-specific marker, LZ425, based on a SNP in the first exon of this gene, matched exactly with the flowering time phenotype (Fig. 4d). Of the six described SNPs, five were silent mutations, but one in position 118 (cv. Salinas, reference genome) causes a change of isoleucine to phenylalanine in PI 251246. This mutation is located within the histidine kinase A domain (SM00388), as annotated by the SMART database (Letunic and Bork 2018), and serves as a dimerization and phosphoacceptor domain of histidine kinases (Vierstra and Davis 2000). The isoleucine residue, corresponding to isoleucine in position 901 in Arabidopsis, is conserved in PhyC genes in angiosperms (supplementary information). It is also conserved in histidine kinase A domains of more diverse organisms, including prokaryotes (it is either isoleucine, leucine, or alanine in over 60% of matched sequences in the SMART database). While the functional effect of this amino acid change is not known, it could alter the gene function in PI 251246.

AT4G14540.1, which is a nuclear factor Y, subunit B3, transcription regulator involved in photoperiod regulation (Kumimoto et al. 2008), was the best matching homolog for Lsat_1_v5_gn_7_96781.1. However, a gene-specific marker
was not fully linked to the flowering phenotype so this gene was not considered to be a likely candidate.

Expression of candidate genes from qBLT7.2/qFLT7.2 region

To identify genes whose gene expression pattern corresponds to QTL effects in various conditions, PI 251246, cv. Salinas, RIL 68, and RIL 69 were sampled at multiple time points during development under the same environmental conditions used for QTL mapping. Of the four genotypes, PI 251246 was the earliest to bolt in all environmental conditions, bolting 21 days after germination. Soon after bolting, buds were visible on these plants (35 \( D_{BUD} \) in SD-LT and 28 \( D_{BUD} \) in the other three conditions). The three other genotypes bolted and flowered later than PI 251246 (Fig. 5). In these genotypes, \( D_{BLT} \) was significantly lower in HT conditions compared to LT \((p < 0.0001, t\)-test), while day length did not have a consistent effect on \( D_{BLT} \). What seems to be an exception in cv. Salinas is probably caused by low light levels due to aged bulbs at the beginning of the growth period in the SD-LT growth chamber. Only the cv. Salinas plants in the chamber were etiolated, and therefore, accurate
evaluation of bolting was not possible. Instead, the presented data are corresponding to the first observation of stem internode elongation that is closely associated with bolting.

While PI 251246 $D_{\text{BUD}}$ was early and relatively unchanged in the four conditions, RIL 69 was also early, but mean $D_{\text{BUD}}$ ranged 28–44 across the four environmental conditions. RIL 68 and cv. Salinas had significantly ($p < 0.001$, $t$-test) higher $D_{\text{BUD}}$ than PI 25146 and RIL 69, and both displayed a wide range of mean $D_{\text{BUD}}$ in the four environmental conditions (71–109 and 67–124 for RIL 68 and cv. Salinas, respectively). For all genotypes, both HT and LD led to lower $D_{\text{BUD}}$ compared to LT ($p < 0.0001$, $t$-test) and SD ($p < 0.05$, $t$-test), respectively. These data indicate that tested temperature conditions had a larger effect than day length on development rate.

Gene expression studies focused on identifying genes which displayed expression patterns consistent with $D_{\text{BLT}}$ and $D_{\text{BUD}}$ phenotypes in response to specific environmental conditions, or genes whose expression pattern differed between PI 251246 and RIL 69, with $P$ allele (originating from PI 251246), and cv. Salinas and RIL 68, with $S$ allele (originating from cv. Salinas). PI 251246 and RIL 69 reached budding stage in only a few weeks and therefore had only the earlier sampling points available for gene expression assays.

Of the twelve targeted genes located in the $q\text{FLT7.2}$ locus (Supplementary Table 2), most did not display any consistent significant patterns between environmental conditions or genotypes (Supplementary Fig. 1). Interestingly, many genes in the study displayed an increase in gene expression in tips of RIL 68 between 42 and 49 days after germination under HT conditions region ($7b_1$, $7b_3$, $7b_4$, $7b_5$, $7b_7$, and $7b_{11}$, putatively annotated as U-box protein, HNRNPA1_3, COL9, unknown protein1, unknown protein2, and FAR1, respectively) also displayed a decrease in expression in RIL 69 tips under SD-LT during the 42 days of sampling. $7b_1$ (U-box protein) had significantly lower expression under SD-LT later in the experiment.

Notably, expression of $\text{COL9}$ ($\text{Lsat}_1\_\text{v5}_5\_\text{gn}_7\_97240.1$) in cv. Salinas and RIL 68 leaves was significantly higher in HT conditions compared to LT conditions, in which $D_{\text{BUD}}$ was higher. On the other hand, in PI 251246 and RIL 69 expression of this gene was more effected by day length, as plants grown under LD displayed significantly higher expression than SD grown plants. In addition, at 14 days after germination leaf expression was higher in PI 251246 and RIL 69 compared to cv. Salinas and RIL 68 (Fig. 6). Additionally, $\text{PhyC}$ ($\text{Lsat}_1\_\text{v5}_5\_\text{gn}_7\_96941.1$) gene expression was significantly higher in cv. Salinas and RIL 68 compared to PI 251246 and RIL 69 in tips under all conditions (Fig. 7). In leaves, expression of the $\text{PhyC}$ gene significantly increased in PI 251246 and RIL 69 at 21 days after germination in SD-LT, in which $D_{\text{BUD}}$ is higher, compared to the other environmental conditions.

Based on gene expression, homology, and sequencing, the strongest candidate genes within the fine mapped $q\text{BLT7.2} / q\text{FLT7.2}$ region include $\text{PhyC}$ and $\text{COL9}$. The lettuce $\text{PhyC}$ homolog found in the QTL region has a mutation causing a change of a conserved isoleucine to phenylalanine in a domain that may influence the protein function. Moreover, the correspondence between

**Fig. 6** Relative expression levels of $\text{COL9}$ ($\text{Lsat}_1\_\text{v5}_5\_\text{gn}_7\_97240.1$) in leaves over the time course of plant development. Shapes indicate genotype: PI 251246 (full circle), RIL 69 (empty circle), RIL 68 (empty square), and cv. Salinas (full square). Colors indicate growth conditions: LD-HT (dark red), LD-LT (dark blue), SD-HT (red), and SD-LT (blue). Error bars indicate standard error of three replicates.
expression pattern and the QTL marker alleles further suggests potential involvement of COL9 and PhyC in regulation of the flowering phenotype.

**QTL qDBFr2.1**

Under SD-HT conditions, the QTL associated with the residuals from the linear regression of $D_{FLT}$ to $D_{BLT}$ designated as qDBFr2.1, mapped to chromosome 2 and accounted for 14.4% of the phenotypic variation observed. In this QTL, the late flowering allele was contributed by the PI 251246 parental line (Table 2). When considering multi-QTL models, qDBtFr2.1 was included in two of the four environmental conditions for $D_{BLT}$ and all four conditions for $D_{FLT}$ (Table 3). Within this genomic region, only a single gene, Lsat_1_v5_gn_2_17881, was found to have an Arabidopsis homolog known to be related to flowering, the lettuce Flowering Locus T (LsFT) gene. FT is a widely conserved gene in dicots and monocots and is a key integrator of floral induction signaling pathways (reviewed in: Ballerini and Kramer 2011). To validate the location of this gene in the QTL, a gene-specific HRM marker (LZ001) mapped to the center of the qDBFr2.1 QTL.

**LsFT gene expression**

At 14 days after germination, when all plants were grown under SD-LT, gene expression levels of LsFT were assessed on leaves and tips of cv. Salinas, PI 251246, RIL 68, and RIL 69. PI 251246 had the highest LsFT expression in leaves, with lower expression in RIL 69, and very little to no expression was observed in RIL 68 and cv. Salinas (Fig. 8). In tips, LsFT expression was highest in cv. Salinas, but only lowly expressed in PI 251246, RIL 69 and RIL 68 (Fig. 8).

In leaves of PI 251246 sampled 21 days after germination, expression of LsFT increased about fourfold higher than observed at 14 days after germination under LD-HT (Fig. 9). LsFT expression of leaves in RIL 69 increased by four to five fold at 21 and 28 days after germination under LD-HT, LD-LT, respectively (Fig. 9). In cv. Salinas, LsFT expression increased about tenfold above that observed at 14- days under LD-HT (Fig. 9). LsFT expression remained essentially unchanged in RIL 68 up through 80 days after germination (Fig. 9).

For tips, LsFT increased sharply in PI 251246 by 21 days after germination under all four environmental conditions (Fig. 9). In RIL 69, LsFT increased under LD-HT and SD-LT (but no data for LD-HT, SD-HT) at 28 and 42 days after germination, respectively (Fig. 9). In RIL 68 and cv. Salinas, LsFT expression fluctuated under all environmental conditions, but remained essentially unchanged across all sampling dates (Fig. 9).
Sequencing of genomic and cDNA revealed no polymorphisms between cv. Salinas and PI 251246 in the coding sequence of the gene. There were, however, numerous SNPs and indels in introns and the promoter region. The region up to 600 bp upstream of the transcription start site had relatively few polymorphisms. However, the region between 660 and 1,186 bp upstream of the start codon had many SNPs and indels in PI 251246 compared to cv. Salinas. Based on in silico motif annotation (Davuluri et al. 2003) and (Higo et al. 1999), two GATA motifs (agatag or agataa) and a SORLIP1 motif (agccac) were found in cv. Salinas but were absent in PI 251246. In Arabidopsis, these motifs are related to light response (Terzaghi and Cashmore 1995; Adrian et al. 2010). None of the predicted motifs were found only in PI 251246 while absent in cv. Salinas (Fig. 10).

**Discussion**

This work identified multiple QTLs affecting bolting and flowering time, which is not surprising considering the complex regulatory network controlling flowering time in the plant kingdom (reviewed in: Komeda 2004; Putterill et al. 2004; Andrés and Coupland 2012). The phenotypic divergence and genetic distance between the RIL population parents allowed detection of multiple QTLs. The current finding that most QTLs were involved in epistatic interactions strengthens the previous results documenting epistatic effect of QTLs on flowering time (Ryder and Milligan 2005; Schwartz et al. 2009).

The critical effect of day length and temperature on plant development (Koornneef et al. 1998; Song et al. 2013) and lettuce flowering (Ito et al. 1963; Jenni et al. 2013; Lafta et al. 2017) is well established. This study aimed to dissect the effect of these environmental factors and their combinations over genetic regulation of development. The QTL detected under the SD-HT condition were not detected under the other three environmental conditions tested, suggesting a different bolting and regulatory network (Table 2). While under both LD conditions and also in SD-LT one major
Fig. 9 Relative expression levels of *LsFT* in plant leaves (top row) and tips (bottom row) over the time course of plant development. Shapes indicate genotype: PI 251246 (full circle), RIL 69 (empty circle), RIL 68 (empty square), and cv. Salinas (full square). Colors indicate growth conditions: LD-HT (dark red), LD-LT (dark blue), SD-HT (red), and SD-LT (blue). Error bars indicate standard error of three replicates. Each panel uses a y-axis scale to optimize visualization of changes during development of each genotype.

Fig. 10 Genomic region of *LsFT*. Light green bars annotate exons and light gray bars annotate UTRs. Red lines annotate sequence polymorphisms between cv. Salinas and PI 251246. Blue blocks annotate predicted transcription factor binding sites in cv. Salinas that are affected by polymorphisms in PI 251246. Nucleotides are numbered relative to the translation start site.
QTL, $q_{FLT7.2}$, dominated and explained much of the phenotypic variance (47.2%), the QTL was not detected under SD-HT. Instead, other QTLs, with lower PVE under other conditions, had significant additive effects. Alternative regulation of flowering under SD-HT was also found in Arabidopsis (Kumar et al. 2012; Fernández et al. 2016). Similarly, in fava bean ($Vicia faba$) under SD-HT many small QTLs were revealed compared to LD conditions (Catt et al. 2017).

Despite the extremely divergent bolting and flowering times of cv. Salinas and PI 251246, transgressive segregation was observed for $D_{FLT}$. In two detected QTLs and one suggestive QTL, $q_{BFr2.1}$, $sq_{BFr3.1}$, and $sq_{FLT9.1}$, late flowering alleles were contributed by the very early flowering parent. Presence of these alleles, and their interaction with other loci, could explain the transgressive segregation. If these alleles are not yet present in domesticated lettuce cultivars, they can contribute to development of lettuce breeding lines and cultivars with late flowering. For bolting, on the other hand, no transgressive segregation was observed, and either the $q_{BLT7.2}$ allele from PI 251246 determined early bolting or, under SD-HT, the PI 251246 allele in $q_{BLT8.1}$. QTLs for flowering, stem length, and plant development were previously found on this chromosome (Hartman et al. 2012; Jenni et al. 2013). Furthermore, an $L. serriola$ accession used in previous bolting studies (Hartman et al. 2012) had the same $P$ allele for marker LZ001, linked to the $q_{BFr2.1}$, as PI 251246.

The two QTLs focused on in this study, $q_{BFr2.1}$ and $q_{FLT7.2}$, had epistatic relations, and the effect of the $q_{FLT7.2}$ QTL was specific to the environmental condition. In SD-HT, when $q_{FLT7.2}$ had no additive effect, no epistatic effect was observed in the models and interaction plot (Fig. 3). However, in LD condition, the QTL $P$ allele determined early flowering while $S$ allowed $q_{BFr2.1}$ QTL alleles to affect $D_{FLT}$. Since $D_{FLT}$ in the $F_{2}$ plants heterozygous in $q_{FLT7.2}$ was early, similar to the $P$ genotype, $S$ can be defined as the recessive allele and could perhaps indicate loss of function. This suggests that in environmental conditions and genotypes where $q_{FLT7.2}$ is functional and has an effect, it overrides any effect $q_{BFr2.1}$ may have. Consequently, only in conditions or alleles where $q_{FLT7.2}$ is not functional, $q_{BFr2.1}$ (and perhaps other loci, as seen in SD-HT) conveys their effect. In other words, $q_{BFr2.1}$ is not ‘induced’ only under SD-HT, but rather is manifested in these conditions due to lack of $q_{FLT7.2}$ effect.

This observation is supported by the gene expression pattern of $LsFT$, the putative gene underlying $q_{BFr2.1}$ QTL. During most of development $LsFT$ expression seems to be governed more by $q_{FLT7.2}$ allele than by the $q_{BFr2.1}$ allele. Specifically, RIL 68 expression in tips and leaves is similar to cv. Salinas and not like PI 251246 and RIL 69, with whom it shares $P$ allele in $q_{BFr2.1}$. The latter genotypes have higher $LsFT$ expression which increases in the first weeks of development and have an early flowering phenotype. This suggests that $q_{FLT7.2}$ is an $LsFT$ expression inducer and supports the detailed work describing the effect of $LsFT$ expression to induce flowering in lettuce (Fukuda et al. 2011, 2017).

In accordance with $FT$’s known role as floral integrator (Kim et al. 2013), other genetic and environmental factors influence $FT$ expression levels. In this study, under HT conditions, $FT$ had higher expression, as found previously in lettuce (Fukuda et al. 2011; Han et al. 2016) and Arabidopsis (Schwartz et al. 2009; Kim et al. 2012). In Chrysanthemum ($Chrysanthemum morifolium$) leaves, on the other hand, heat caused reduction in FT-like gene expression and delayed flowering (Nakano et al. 2013). However, the increase under HT conditions in leaves is not translated to an increase in tips in genotypes with $q_{FLT7.2} S$ allele. The lack of epistasis in the SD-HT QTL model stands in contrast with the apparent influence of $q_{FLT7.2}$ on $LsFT$ gene expression in this condition, as well.

| Range  | Mean | SD  |
|--------|------|-----|
| Cv. Salinas | 100  | 100 | 0 |
| PI 251246 | 30   | 30  | 0 |
| RIL 68 | 88–92 | 90  | 2.82 |
| RIL 69 | 65   | 65  | 0  |
| BC$_{1}$S$_{1}$-1 | 88–150 | 111.2 | 18.66 |
| BC$_{1}$S$_{1}$-2 | 86–140 | 102.5 | 17.21 |
| BC$_{1}$S$_{1}$-3 | 85–153 | 104  | 21.43 |
| BC$_{2}$ | 92–128 | 111.4 | 13.56 |

Table 4. $D_{FLT}$ of backcross families for $q_{BFr2.1}$ effect validation under a LD-LT greenhouse environment.
The only time point when cv. Salinas LsFT gene expression matched its late flowering allele in qBFr2.1 locus was in the tips of 14-day old plants. At this point, all plants were grown in SD-LT conditions, considered to be least inductive of flowering. The exclusive high expression in cv. Salinas’s tips, not seen in leaves, may initiate an induction of flowering in the absence of qFLT7.2. The inconsistency in LsFT expression between leaves and tips, also observed in later stages in cv. Salinas and RIL 68, is especially interesting given previous reports that FT moves from leaves to the meristem within the tip where its protein is activated (Corbesier et al. 2007; Putterill and Varkonyi-Gasic 2016).

FT has a long promoter with many cis regulatory elements which affect its expression (Schwartz et al. 2009; Adrian et al. 2010; Liu et al. 2014), such as those identified to be different between cv. Salinas and PI 251246. The lettuce FT gene is located in a relatively gene-poor region, with only one gene detected within a 3 Kbp upstream transcription start site with no others for 116 Kbp, as seen in other plant species (Adrian et al. 2010). The modified elements in PI 251246, GATA motif, and SORLIP1 motif, may influence the expression pattern differences seen between tips of cv. Salinas and the P allele genotypes at 14 days.

As opposed to the most obvious candidate gene underlying the qBFr2.1 QTL, fine mapping was required to shorten the candidate list for qFLT7.2. After narrowing the interval to a 1.15 Mbp region, two genes remained plausible candidates. Still, the possibility that both genes have a role in the control of flowering in lettuce, either together or each at different conditions, cannot be excluded. Lsat_1_v5_gn_7_96941.1 was identified as an Arabidopsis PhyC homolog. PhyC belongs to a family of phytochrome factors (Sharrock and Quail 1989), which have important roles in multiple aspects of plant growth and development, including germination, chloroplast development, photomorphogenesis, shade avoidance, and photoperiod-dependent flowering (Kendrick and Kronenberg 1994; Franklin et al. 2003; Nishida et al. 2013). In Arabidopsis, PhyC is not required for long-day induction of flowering time (Franklin et al. 2003). It is, however, involved in day length perception and delay of flowering in SD, together with PhyA and PhyB (Monte et al. 2003). In wheat and barley, on the other hand, PhyC was found to have a key role in LD-induced flowering (Nishida et al. 2013; Pearce et al. 2016).

Phytochrome family proteins bind a bilin chromophore and share a basic structure (Pham et al. 2018), with distinct and partially overlapping roles in development control and signal transduction. The N terminus of phytochromes has a kinase domain, and kinase activity was shown to be important for plant photo- response (Shin et al. 2016). Gene sequencing of both parents revealed a mutation in PI 251246 located in a conserved amino acid within the histidine kinase A domain. While the effect of the change on protein function is not known, it could possibly effect photoperiod-dependent flowering time regulation.

The closest Arabidopsis homolog of Lsat_1_v5_gn_7_97240.1 is COL9. The CO-like gene family three-group structure is conserved in various plant groups (Wang et al. 2019). Although most genes in the family promote flowering in Arabidopsis, COL9 was found to suppress FT, SOC1, and CO and delay flowering in Arabidopsis (Cheng and Wang 2005). Functional homologs of COL9 affect flowering time in Arabidopsis (Cheng and Wang 2005), rice (Wu et al. 2018), maize (Maldonado et al. 2019) and were also identified in Ambrosia artemisiifolia (Mátyás et al. 2019). In this study, COL9 expression was higher in early flowering genotypes with the P allele, which could suggest a flowering promoting role. On the other hand, expression in S allele genotypes was higher in LT conditions, in which flowering is later. This contrasts with a previous study in which LsCOL9 expression was found to significantly increase under flowering inducing heat conditions (Liu et al. 2018). These discrepancies may be due to unidentified polymorphisms in the genotype’s promoters, leading to differing sensitivity to the environmental conditions. So the effect of COL9 gene expression on the flowering phenotype is not yet well defined. Taken together, either LsCOL9 or LsPhyC or both could be qBLT/qFLT7.2 causal gene involved in lettuce flowering.

In summary, this work contributes to efforts of breeding cultivars less prone to premature bolting through improved resolution and understanding of the major qBLT/qFLT7.2 QTL. This QTL was detected in multiple experimental populations and has a determining effect on bolting and flowering rates in most lettuce production environments.

Another main contribution is the identification of the qBFr2.1 allele from PI 251246 not currently present in the rosette-forming lettuce genepool, as demonstrated with marker LZ001 on a diverse group of cultivars. Since the primitive oil seed cultivar (PI 251246) allele has a significant effect in delaying flowering in SD-HT conditions and in delaying bolting and flowering in certain field conditions (You, Still, Rosental, Hayes and Simko—unpublished data), introgression of the allele into commercial germplasm may lead to development of breeding lines with delayed bolting. The allele may be particularly useful when breeding cultivars specifically adapted for lettuce production in full plantings in the southwestern USA, an environment especially susceptible to premature bolting and flowering. The LZ001 marker can be used to facilitate breeding by MAS.

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s00122-021-03898-9.
Data availability

DWS. California State University Agricultural Research Institute to RJH and approved the final manuscript.

data analysis, and drafting of the manuscript. All authors edited and approved the final manuscript.

Author contribution statement

LR planned and performed experiments, collected and analyzed data, and drafted the manuscript. YY guided and assisted in planning and analyzing GeXP experiment. RJH and DS conceived the project, obtained funding, supervised the experiments, and drafted the manuscript. IS supervised the experiments, data analysis, and drafting of the manuscript. All authors edited and

Funding

This research was supported by grant # 14–01-008 from the California State University Agricultural Research Institute to RJH and DWS.

LR is a fellow of the Arian de Rothschild Women Doctoral Program.

Data availability

data transparency

Declarations

Conflicts of interest

The authors state that there is no conflict of interest.

References

Adrian J, Farrona S, Reimer JJ et al (2010) cis-Regulatory elements and chromatin state coordinately control temporal and spatial expression of flowering locus tin arabidopsis. Plant Cell Online 22:1425–1440. https://doi.org/10.1105/tpc.110.074682

Amasino R (2010) Seasonal and developmental timing of flowering. Plant J 61:1001–1013. https://doi.org/10.1111/j.1365-313X.2010.04148.x

Andrés F, Coupland G (2012) The genetic basis of flowering responses to seasonal cues. Nat Rev Genet 13:627–639. https://doi.org/10.1038/nrg3291

Ballerini ES, Kramer EM (2011) In the light of evolution: a reevaluation of conservation in the CO-FT regulon and its role in photoperiodic regulation of flowering time. Front Plant Sci 2:1–13. https://doi.org/10.3389/fpls.2011.00081

Bao S, Hua C, Shen L, Yu H (2020) New insights into gibberellin signaling in regulating flowering in Arabidopsis. J Integr Plant Biol 62:118–131. https://doi.org/10.1111/jipb.12892

Blackman BK, Strasburg JL, Raduski AR et al (2010) The role of recently derived ft paralogs in sunflower domestication. Curr Biol 20:629–635. https://doi.org/10.1016/j.cub.2010.01.059

Blázquez MA, Aha JH, Weigel D (2003) A thermosensory pathway controlling flowering time in Arabidopsis thaliana. Nat Genet 33:168–171. https://doi.org/10.1038/ng1085

Blümel M, Dally N, Jung C (2015) Flowering time regulation in crops—what did we learn from Arabidopsis? Curr Opin Biotechnol 32:121–129. https://doi.org/10.1016/j.copbio.2014.11.023

Borowski JM, Galli V, da Silva MR et al (2014) Selection of candidate reference genes for real-time PCR studies in lettuce under abiotic stress. Planta 239:1187–1200. https://doi.org/10.1007/s00425-014-2041-2

Bouché F, Lobet G, Tocquin P, Périlleux C (2016) FLOR-ID: an interactive database of flowering-time gene networks in Arabidopsis thaliana. Nucleic Acids Res 44:D1167–D1171. https://doi.org/10.1093/nar/gkv1054

Broman KW, Speed TP (2002) A model selection approach for the identification of quantitative trait loci in experimental crosses. J R Stat Soc B 64:641–656. https://doi.org/10.1111/j.1467-9868.2011.00815.x

Broman KW, Wu H, Sen Š, Churchill GA (2003) R/qtl: QTL mapping in experimental crosses. Bioinformatics 19:889–890. https://doi.org/10.1093/bioinformatics/btg112

Capovilla G, Schmid M, Posé D (2015) Control of flowering by ambient temperature. J Exp Bot 66:59–69. https://doi.org/10.1093/jxb/eru416

Catt SC, Braich S, Kaur S, Paul JG (2017) QTL detection for flowering time in fava bean and the responses to ambient temperature and photoperiod. Euphytica 213:1–13. https://doi.org/10.1007/s10681-017-1910-8

Chen Z, Han Y, Ning K et al (2018a) Inflorescence development and the role of lfsf in regulating bolting in lettuce (Lactuca sativa L.). Front Plant Sci 8:1–10. https://doi.org/10.3389/fpls.2017.02248

Chen Z, Zhao W, Ge D et al (2018b) LCM-seq reveals the crucial role of LsSOC1 in heat-promoted bolting of lettuce (Lactuca sativa L.). Plant J 95:516–528. https://doi.org/10.1111/tpj.13968

Cheng XF, Wang ZY (2005) Overexpression of Cole9, a CONSTANS-LIKE gene, delays flowering by reducing expression of CO and FT in Arabidopsis thaliana. Plant J 43:758–768. https://doi.org/10.1111/j.1365-313X.2005.02491.x

Cho LH, Yoon J, An G (2017) The control of flowering time by environmental factors. Plant J 90:708–719. https://doi.org/10.1111/tpj.13461

Corbesier L, Vincent C, Jang S et al (2007) FT protein movement contributes to long-distance signaling in floral induction of Arabidopsis. Science. https://doi.org/10.1126/science.1141752

Czechowski T, Stitt M, Altmann T et al (2005) Genome-wide identification and testing of superior reference genes for transcript normalization. Plant Physiol 139:5–17. https://doi.org/10.1104/pp.105.067343.1

Davuluri RV, Sun H, Palaniswamy SK et al (2003) AGRIS: arabidopsis Gene Regulatory Information Server, an information resource of Arabidopsis cis-regulatory elements and transcription factors. BMC Bioinformatics 4:1–11. https://doi.org/10.1186/1471-2105-4-25

Elshire RJ, Glaubitz JC, Sun Q et al (2011) A robust, simple genotyping-by-sequencing (GBS) approach for high diversity species. PLoS ONE 6:1–10. https://doi.org/10.1371/journal.pone.0019379

Feenstra B, Skovgaard IM, Broman KW (2006) Mapping quantitative trait loci by an extension of the Haley-Knott regression method. Theor Appl Genet 110:2077–2084. https://doi.org/10.1007/s00122-006-0697-6

Fernández V, Takahashi Y, Le Gourrierec J, Coupland G (2016) Photoperiodic and thermosensory pathways interact through CONSTANS-LIKE gene, in lettuce. J Plant Physiol 168:1602–1607. https://doi.org/10.1016/j.jplph.2016.04.024

Franklin KA, Davis SJ, Stoddart WM et al (2003) Mutant analyses define multiple roles for phytochrome c in arabidopsis photomorphogenesis. Plant Cell Online 15:1981–1989. https://doi.org/10.1105/tpc.017193

Fornara F, de Montaigu A, Coupland G (2010) SnapShot: control of flowering by ambient temperature. Plant J 61:1001–1013. https://doi.org/10.1111/j.1365-313X.2010.04148.x

Fukuda M, Matsuo S, Kikuchi K et al (2009) The endogenous level of GA1is upregulated by high temperature during stem elongation in Arabidopsis thaliana. Nucleic Acids Res 37:9206–9219. https://doi.org/10.1093/nar/gkp759

Fukuda M, Matsuo S, Kikuchi K et al (2011) Isolation and functional analysis of LsFT gene, in lettuce. J Plant Physiol 168:1602–1607. https://doi.org/10.1016/j.jplph.2011.02.004

Fukuda M, Matsu o S, Kikuchi K et al (2009) The endogenous level of GA1is upregulated by high temperature during stem elongation in lettuce through LsGA3ox1 expression. J Plant Physiol 166:2077–2084. https://doi.org/10.1016/j.jplph.2009.06.003

Fukuda M, Matsu o S, Kikuchi K et al (2011) Isolation and functional characterization of the FLOWERING LOCUS T homolog, the LsFT gene, in lettuce. J Plant Physiol 168:1602–1607. https://doi.org/10.1016/j.jplph.2011.02.004
Fukuda M, Yanai Y, Nakano Y et al (2017) Isolation and gene expression analysis of flowering-related genes in lettuce (Lactuca sativa L.). Hortic J 86:340–348. https://doi.org/10.2503/hortj.0KD-036

Glaziev IC, Caststevens TM, Lu F et al (2014) TASSEL-GBS: a high capacity genotyping by sequencing analysis pipeline. PLoS ONE. https://doi.org/10.1371/journal.pone.0090346

Grube R, Ryder E (2004) Identification of Lettuce (Lactuca sativa L.) Germplasm with Genetic Resistance to Drop Caused by Sclerotinia minor. J Am Soc Hortic Sci 129:70–76

GuoYou Y, Ogbonnaya F, van Ginkel M (2010) Marker-assisted recurrent backcrossing in cultivar development. Molecular plant breeding: principle, method and application. Studium Press LLC, Houston, pp 295–319

Han Y, Chen Z, Lv S et al (2016) MADS-box genes and gibberellins regulate bolting in lettuce (Lactuca sativa L.). Front Plant Sci 7:1–14. https://doi.org/10.3389/fpls.2016.01889

Han R, Truco MJ, Lavelle DO, Michelmore RW (2021) A comparative analysis of flowering time regulation in lettuce. Front Plant Sci 12:1–14. https://doi.org/10.3389/fpls.2021.632708

Hartman Y, Hooftman DAP, Uwimana B et al (2012) Genomic regions in crop-wild hybrids of lettuce are affected differently in different environments: implications for crop breeding. Evol Appl 5:629–640. https://doi.org/10.1111/j.1752-4571.2012.00240.x

Hayashi E, Aoyama N, Wu Y et al (2007) Multiplexed, quantitative gene expression analysis for lettuce seed germination on GenomoLab™ GeXP Genetic Analysis System. Beckman-Coulter Application Information-10295A. https://sciex.jp/content/dam/SCIEX/pdf-technology/All-A-10295A-c.pdf

Higo K, Ugawa Y, Iwamoto M, Korenaga T (1999) Plant cis-acting regulatory DNA elements (PLACE) database: 1999. Nucleic Acids Res 27:297–300. https://doi.org/10.1093/nar/27.1.297

Huo H, Wei S, Bradford KJ (2016) DELAY OF GERMINATION1 (DOG1) regulates both seed dormancy and flowering time through microRNA pathways. Proc Natl Acad Sci 113:E2199–E2206. https://doi.org/10.1073/pnas.160055811

Ito H, Kato T, Konno Y (1963) Factors associated with the Flower Induction in Lettuce. Tohoku J Agric Res 14:51–65

Jenni S, Truco MJ, Michelmore RW (2013) Quantitative trait loci associated with tipburn, heat stress-induced physiological disorders, and maturity traits in crisphead lettuce. Theor Appl Genet 126:3065–3079. https://doi.org/10.1007/s00122-013-2193-7

Jeuken MJW, Lindhout P (2004) The development of lettuce backcross inbred lines (BILs) for exploitation of the Lactuca saligna (wild lettuce) germplasm. Theor Appl Genet 109:394–401. https://doi.org/10.1007/s00122-004-1643-7

Kendrick RE, Kronenberg GMH (1994) Photomorphogenesis in plants. Springer, Netherlands. https://doi.org/10.1007/978-94-011-1884-2

Kim JJ, Lee JH, Kim W et al (2012) The microRNA156–SQUAMOSA promoter binding protein-like3 module regulates ambient temperature-responsive flowering via flowering locus in Arabidopsis. Plant Physiol 159:461–478. https://doi.org/10.1104/pp.111.192369

Kim W, Park TI, Yoo SJ et al (2013) Generation and analysis of a complete mutant set for the Arabidopsis FT/FUL family shows specific effects on thermo-sensitive flowering regulation. J Exp Bot 64:1715–1729. https://doi.org/10.1093/jxb/ert036

Komeda Y (2004) Genetic regulation of time to flower in arabisponitiana. Annu Rev Plant Biol 55:521–535. https://doi.org/10.1146/annurev.arplant.55.031903.141644

Koomneef M, Alonso-Blanco C, Peeters AJ, Soppe W (1998) Genetic control of flowering time in Arabidopsis. Annu Rev Plant Biol 49:345–370. https://doi.org/10.1002/9781119409144.ch22

Kumar SV, Lucyshyn D, Jaeger KE et al (2012) Transcription factor PIF4 controls the thermosensory activation of flowering. Nature 484:242–245. https://doi.org/10.1038/nature10928

Kumamoto RW, Adam L, Hymus GJ et al (2008) The nuclear factor Y subunits NF-YB2 and NF-YB3 play additive roles in the promotion of flowering by inductive long-day photoperiods in Arabidopsis. Planta 228:709–723. https://doi.org/10.1007/s00045-007-773-6

Kwon S, Simko I, Hellier B et al (2013) Genome-wide association of 10 horticultural traits with expressed sequence tag-derived SNP markers in a collection of lettuce lines. Crop J 1:25–33. https://doi.org/10.1016/j.cj.2013.07.014

Lafta A, Turini T, Sandoyva G, Mou B (2017) Field Evaluation of green and red leaf lettuce genotypes in the imperial, san joaquin, and salinas valleys of California for heat tolerance and extension of the growing seasons. HortScience 52:40–48

Lee ON, Fukushima K, Park HY, Kawabata S (2021) QTL analysis of stem elongation and flowering time in lettuce using genotyping-by-sequencing. Genes 12(6):947. https://doi.org/10.3390/genes12060947

Letunic I, Bork P (2018) 20 years of the SMART protein domain annotation resource. Nucleic Acids Res 46:D493–D496. https://doi.org/10.1093/nar/gkx922

Liu L, Adrian J, Pankin A et al (2014) Induced and natural variation of promoter length modulates the photoperiodic response of FLOWERING LOCUS T. Nat Commun 5:1–9. https://doi.org/10.1038/ncomms5558

Liu X, Lv S, Liu R et al (2018) Transcriptomic analysis reveals the roles of gibberellin-regulated genes and transcription factors in regulating bolting in lettuce (Lactuca sativa L.). PLoS ONE 13:1–17. https://doi.org/10.1371/journal.pone.0191518

Maldonado C, Mora F, Bengosi Bertagna FA et al (2019) SNP- and haplotype-based GWAS of flowering-related traits in maize with network-assisted gene prioritization. Agronomy. https://doi.org/10.3390/agronomy9110725

Mamo BE, Hayes RJ, Truco MJ et al (2019) The genetics of resistance to lettuce drop (Sclerotinia spp.) in lettuce in a recombinant inbred line population from Reine des Glaces × Eruption. Theor Appl Genet 132:2439–2460. https://doi.org/10.1007/s00122-019-0298-7

Mátyás KK, Hegedűs G, Taller J et al (2019) Different expression of PHYTOCHROME B subunits NF-YB2 and NF-YB3 play additive roles in the control of flowering time in Arabidopsis. Annu Rev Plant Biol 70:345–370. https://doi.org/10.1146/annurev-arplant.55.031903.141644

McGinnis S, Madden TL (2004) BLAST: at the core of a powerful and diverse set of sequence analysis tools. Nucleic Acids Res 32:20–25. https://doi.org/10.1093/nar/gkh435

Michaels SD, Amasino RM (2000) Memories of winter: Vernalization and the competence to flower. Plant Cell Environ 23:1145–1153. https://doi.org/10.1046/j.1365-3040.2000.00643.x

Monte E, Alonso JM, Ecker JR et al (2003) Isolation and characterization of phyB mutants in arabidopsis reveals complex crosstalk between phytochrome signaling pathways. Plant Cell 15:1962–1980. https://doi.org/10.1105/tpc.012971

Nakano Y, Higuchi Y, Sumitomo K, Hisamatsu T (2013) Flowering retardation by high temperature in chrysanthemums: Involvement of FLOWERING LOCUS T-like 3 gene repression. J Exp Bot 64:909–920. https://doi.org/10.1093/jxb/ers370

Ning K, Han Y, Chen Z et al (2019) Genome - wide analysis of MADS - box family genes during flower development in lettuce. Plant Cell Environ 42:1868–1881. https://doi.org/10.1111/pec.13523
Nishida H, Ishihara D, Ishii M et al (2013) Phytochrome C is a key factor controlling long-day flowering in barley. Plant Physiol 163:804–814. https://doi.org/10.1104/pp.113.222570

Pearce S, Kippes N, Chen A et al (2016) RNA-seq studies using wheat PHYTOCHROME B and PHYTOCHROME C mutants reveal shared and specific functions in the regulation of flowering and shade-avoidance pathways. BMC Plant Biol 16:1–19. https://doi.org/10.1186/s12870-016-0831-3

Pham VN, Kathare PK, Huq E (2018) Phytochromes and phytochrome interacting factors. Plant Physiol 176:1025–1038. https://doi.org/10.1104/pp.17.01384

Putterill J, Varkonyi-Gasic E (2016) FT and florigen long-distance flowering control in plants. Curr Opin Plant Biol 33:77–82. https://doi.org/10.1016/j.pbi.2016.06.008

Putterill J, Laurie R, Macknight R (2004) It’s time to flower: the genetic control of flowering time. BioEssays 26:363–373. https://doi.org/10.1002/bies.20021

Rappaport L, Wittwer S, Tukey H (1956) Seed vernalization and flowering responses of Arabidopsis thaliana. Genetics 183:723–732. https://doi.org/10.1534/genetics.109.104984

Romera-Branchat M, Andrés F, Coupland G (2014) Flowering enhancer gene family in lettuce. Theor Appl Genet 133:1947–1966. https://doi.org/10.1007/s00344-013-1855-z

Ryder EJ (1983) Inheritance, linkage, and gene interaction studies in lettuce. J Am Soc Hortic Sci 108:985–991

Ryder EJ (1988) Early flowering in lettuce as influenced by a second flowering time gene and seasonal variation. J Am Soc Hortic Sci 113:456–460

Ryder EJ (1996) Ten lettuce genetic stocks with early flowering genes Ef-lef-1 and Ef-2ef-2. HortScience 31:473–475

Ryder EJ, Milligan DC (2005) Additional genes controlling flowering time gene and seasonal variation. J Am Soc Hortic Sci 120:460–467

Samach A, Coupland G (2000) Time measurement and the control of flowering in plants. BioEssays 22:38–47

Schwartz C, Balasubramanian S, Warthmann N et al (2009) Cis-regulatory changes at Flowering Locus T mediate natural variation in flowering responses of Arabidopsis thaliana. Genetics 183:723–732. https://doi.org/10.1034/genetics.109.104984

Sgambarra A, Mage J, Massiah A, Jackson S (2016) Selection of reference genes for diurnal and developmental time-course real-time PCR expression analyses in lettuce. Plant Methods 12:1-9. https://doi.org/10.1186/s13007-016-0121-y

Sharrock RA, Quail PH (1989) Novel phytochrome sequences in Arabidopsis thaliana: structure, evolution, and differential expression of a plant regulatory photoreceptor family. Genes Dev 3:1745–1757. https://doi.org/10.1101/gad.3.11.1745

Shin AY, Han YJ, Baek A et al (2016) Evidence that phytochrome functions as a protein kinase in plant light signalling. Nat Commun. https://doi.org/10.1038/ncomms11545

Silva EC, Maluf WR, Leal NR, Gomes LAA (1999) Inheritance of bolting tendency in lettuce Lactuca sativa L. Euphytica 109:1–7. https://doi.org/10.1023/A:1003698117689

Simko I (2013) Marker-Assisted Selection for Disease Resistance in Lettuce. In: Rajeev KV, Tuberosa R (eds) Translational Genomics for Crop Breeding, Volume I: Biotic Stress. John Wiley & Sons Inc, First Edit, pp 267–289

Simko I (2016) High-resolution DNA melting analysis in plant research. Trends Plant Sci 21:528–537. https://doi.org/10.1016/j.tplants.2016.01.004

Simonne A, Simonne E, Eitenmiller R, Coker CH (2002) Bitterness and composition of lettuce varieties grown in the southeastern United States. HortTechnology 12:721–726

Song YH, Ito S, Imaizumi T (2013) Flowering time regulation: photoperiod- and temperature-sensing in leaves. Trends Plant Sci 18:575–583. https://doi.org/10.1016/j.tplants.2013.05.003

Song YH, Shim JS, Kinmonth-Schultz HA, Imaizumi T (2015) Photoperiodic flowering: time measurement mechanisms in leaves. Annu Rev Plant Biol 66:441–464. https://doi.org/10.1146/annurev-arplant-043014-115555

Shiapiti Kandel J, Peng H, Hayes RJ et al (2020) Genome-wide association mapping reveals loci for shelf life and developmental rate of lettuce. Theor Appl Genet 133:1947–1966. https://doi.org/10.1007/s00122-020-03568-2

Terzaghi WB, Cashmore AR (1995) Light-regulated transcription. Annu Rev Plant Physiol Plant Mol Biol 46:445–474

Thompson RC (1943) Further studies on interspecific genetic relationships in Lactuca. J Agric Res 66:41–48

Umetu A, Sawada Y, Mitsuhashi W et al (2011) Characterization of a loss-of-function mutant of gibberelin biosynthetic gene lsga3ox1 in lettuce. Biosci Biotechnol Biochem 75:2398–2400. https://doi.org/10.1271/bbb.110475

Vandesompele J, De Preter K, Pattyn F et al (2002) Accurate normalization of real-time quantitative RT-PCR data by geometric averaging of multiple internal control genes. Genome Biol 3:34-41. https://doi.org/10.1186/gb-2002-3-7-research0034

Vierstra RD, Davis SJ (2000) Bacteriophytochromes: new tools for understanding phytochrome signal transduction. Semin Cell Dev Biol 11:511–521. https://doi.org/10.1006/scdb.2000.0206

Wang L, Xue J, Dai W et al (2019) Genome-wide identification, phylogenetic analysis, and expression profiling of constans-like (col) genes in vitis vinifera. J Plant Growth Regul 38:631–643. https://doi.org/10.1007/s00122-018-9878-8

Waycott M (1995) Photoperiodic response of genetically diverse lettuce accessions. J Am Soc Hortic Sci 120:460–467

Wien HC (2020) Lettuce. In: Wien HC, Stutzel H (eds) The physiology of vegetable crops, 2nd edn. CAB International, Wallingford, UK, pp 333–356

Wu W, Zhang Y, Zhang M et al (2018) The rice CONSTANS-like protein OsCOL15 suppresses flowering by promoting Ghd7 and repressing RID1. Biochem Biophys Res Commun 495:1349–1355. https://doi.org/10.1016/j.bbrc.2017.11.095

Yoong F-Y, O'Brien LK, Truco MJ et al (2016) Genetic variation for temperature-sensitive regulation of ETHYLENE RESPONSE FACTOR1 (ERF1). Plant Physiol 170:472–488. https://doi.org/10.1104/pp.15.01251

Publisher’s Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.