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Regulation of *FT* splicing by an endogenous cue in temperate grasses

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Appropriate flowering timing is crucial for plant reproductive success. The florigen, FLOWERING LOCUS T (FT), interacts with 14-3-3 proteins and the bZIP transcription factor FD, functioning at core nodes in multiple flowering pathways. There are two FT homologues, FT1 and FT2, in *Brachypodium distachyon*. Here we show that *FT2* undergoes age-dependent alternative splicing (AS), resulting in two splice variants (*FT2 α* and *FT2 β*). The *FT2 β* -encoded protein cannot interact with FD or 14-3-3s but is able to form heterodimers with *FT2 α* and FT1, thereby interfering with the florigen-mediated assembly of the flowering initiation complex. Notably, transgenic plants overproducing *FT2 β* exhibit delayed flowering, while transgenic plants in which *FT2 β* is silenced by an artificial microRNA display accelerated flowering, demonstrating a dominant-negative role of *FT2 β* in flowering induction. Furthermore, we show that the AS splicing of *FT2* is conserved in important cereal crops, such as barley and wheat. Collectively, these findings reveal a novel posttranscriptional mode of *FT* regulation in temperate grasses.

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The correct timing of the transition from the vegetative to the reproductive stage is a critical point during plant life cycles. Plants require appropriate environmental conditions, including optimal photoperiod and temperature, to stimulate flowering at a specific time of the year. Additionally, plants also must develop to a certain age with fixed amounts of hormones to achieve flowering¹. As the encoder of the mobile flower-promoting signal florigen, the *FLOWERING LOCUS T (FT)* gene has been shown to have key roles in integrating external and endogenous cues to control the onset of flowering.

FT proteins are a clade of the phosphatidylethanolamine-binding protein (PEBP) family, which act as highly conserved regulators in plants that relay signals from upstream to downstream in flowering pathways. In *Arabidopsis thaliana*, FT messenger RNA (mRNA) gradually increases with developmental time under a long-day (LD) photoperiod². After synthesis in leaves, FT protein travels through the vasculature to the shoot apical meristem (SAM) and interacts with a bZIP transcription factor, named FD, which is expressed specifically in the SAM^{3,4}. Subsequently, the FT/FD complex activates a number of MADS box genes, including *APETALA1*, *FRUITFULL* and *SUPPRESSOR OF OVEREXPRESSION OF CONSTANS 1* to stimulate floral organ initiation⁵. The 14-3-3 proteins, a family of regulatory molecules conserved in animal and plants, can directly interact with the FT homologue Heading date 3a in rice apical shoots, forming a functional complex that translocates to the nucleus⁶. Thus, 14-3-3 proteins may function as scaffold proteins to associate FT with FD to trigger flowering.

Numerous FT transcriptional regulators have been identified to date. For example, in *A. thaliana*, CONSTANS and GIGANTEA activate FT expression in the photoperiod pathway⁷⁻⁹, while FLOWERING LOCUS C and SHORT VEGETATIVE PHASE form a MADS-box complex that negatively mediates FT expression in the flowering vernalization-related pathway¹⁰. Two well-known plant-conserved microRNAs (miRNAs), miR156 and miR172, directly or indirectly act at the FT region through their targets, mediating age-dependent flowering processes¹¹⁻¹³. Compared with the extensive knowledge of FT control at the transcriptional level, our understanding of FT regulation at the posttranscriptional level is limited.

Temperate grasses, including wheat and barley, are major sources of food and biofuel worldwide. Some specialized components, lacking orthologues in *A. thaliana* and rice, have been discovered in temperate grasses and can affect FT gene expression in day-length and vernalization flowering pathways¹⁴⁻¹⁶. The typical *Pooideae* grass *Brachypodium distachyon* has been considered as a suitable model system for investigating gene function in temperate cereals because of its small genome, simple growth requirements and short life cycle¹⁷. Phytochrome C, rather than phytochrome A or B, is an essential light receptor for photoperiodic flowering in wheat and *B. distachyon*, suggesting that there are different evolutionary features of LD flowering responses in temperate grasses¹⁸⁻²⁰. Moreover, a *Pooideae*-specific miRNA, miR5200, has recently been identified that directly silences two FT orthologues and has a crucial role in the photoperiod-mediated flowering pathway in *B. distachyon*, further implying that a distinct flowering control mechanism exists in temperate cereals¹⁴.

Alternative splicing (AS) is a universal mechanism that produces multiple mRNAs from one gene through the variable selection of splice sites during mRNA generation^{21,22}. AS can modulate gene expression levels by introducing a premature termination codon, resulting in the degradation of a special RNA-splicing isoform through a nonsense-mediated decay pathway²³. Additionally, different transcript isoforms may produce truncated

proteins that may have different subcellular localization, stability level or function²³. Several studies have implicated the involvement of AS in plant development, circadian rhythm, starch metabolism, hormone signaling and abiotic stress response, as well as pathogen defense²⁴⁻³⁶, suggesting that this mechanism of gene regulation has a significant role in plant adaptation and evolution. Despite the revelation by next-generation sequencing that AS occurs in >60% of *A. thaliana* and 42% of *B. distachyon* genes, the biological significance of most AS events in plants is still largely unknown^{21,37-39}.

Previously, we demonstrated that two orthologous FT genes, *FTL1 (Bradi2g07070)* and *FTL2 (Bradi1g48830)*, have miR5200-targeting sites and are regulated by miR5200 in *B. distachyon*¹⁴. Because in *B. distachyon*, more amino acids of FTL2, compared with FTL1, are identical with those of florigen VRN3 in wheat and barley, we re-designated FTL1 as FT2 and FTL2 as FT1 to be clearer when analyzing the phylogenetic relationships among the FTs of temperate grasses⁴⁰.

In this study, we show that AS of FT2 has an important role in controlling florigen activity by producing a competitive repressor in *B. distachyon*. FT2 β , which only lacks a short section of the N-terminal PEBP domain, can attenuate the activities of the flowering initiation complex by interacting with full-size FT2 α and FT1. The FT2 β /FT2 α ratio progressively decreased with plant growth, resulting in a slow gradual increase in the florigen activity level. The overexpression of FT2 β in transgenic plants resulted in delayed flowering, whereas plants with downregulated FT2 β activity through artificial miRNA-induced repression, exhibited accelerated flowering, further demonstrating that FT2 β acts as a dominant-negative repressor in flowering time control in *B. distachyon*. Together, these results reveal a novel molecular mode of FT posttranscriptional regulation in temperate grasses.

Results

FT2 is subject to AS generating two isoforms. In *B. distachyon*, FT1 and FT2 share high amino-acid sequence identity and their overproduction gives rise to extremely early bolting phenotypes, indicating that they perform redundant roles in flowering initiation^{14,15,40}. During the study of the regulatory mechanism of FT1 and FT2 in flowering, we interestingly found that two PCR products were generated by a pair of FT2 primers, implying the existence of FT2 AS in *B. distachyon* (Fig. 1a). By comparing the two FT2 cDNA sequences, we discovered that one splicing product was missing 84 nucleotides at the 5'-end of the first exon compared with a typical FT gene. Thus we designated this form as FT2 β and the full-size transcript as FT2 α (Fig. 1b). A protein sequence alignment showed that the missing 28 amino acids came from the 5' terminal region of the PEBP domain of FT2 β , which is more variable compared with the middle and 3' regions (Fig. 1c; Supplementary Fig. 1a). However, no FT1 splicing variant was detected in *B. distachyon*, possibly because the gene structure of FT1 is different from that of FT2 (Supplementary Fig. 1b,c).

Using specific primers to amplify each FT2 RNA splicing product, we detected the presence of both variants in *B. distachyon* in most cases (Fig. 1a). Most AS events in plants are shown to be affected by ambient temperature fluctuations^{24,25,28,41}. Similarly, the ratios of FT2 β /FT2 α were altered when plants were grown at high temperatures (Supplementary Figs 2 and 3).

To determine the role, if any, of different FT2 variants in the control of flowering, we attempted to independently overproduce FT2 α and FT2 β in *B. distachyon*. As previously described by us and others, the ectopic expression of intact FT2 (FT2 α) in *B. distachyon* leads to very early flowering and arrested vegetative

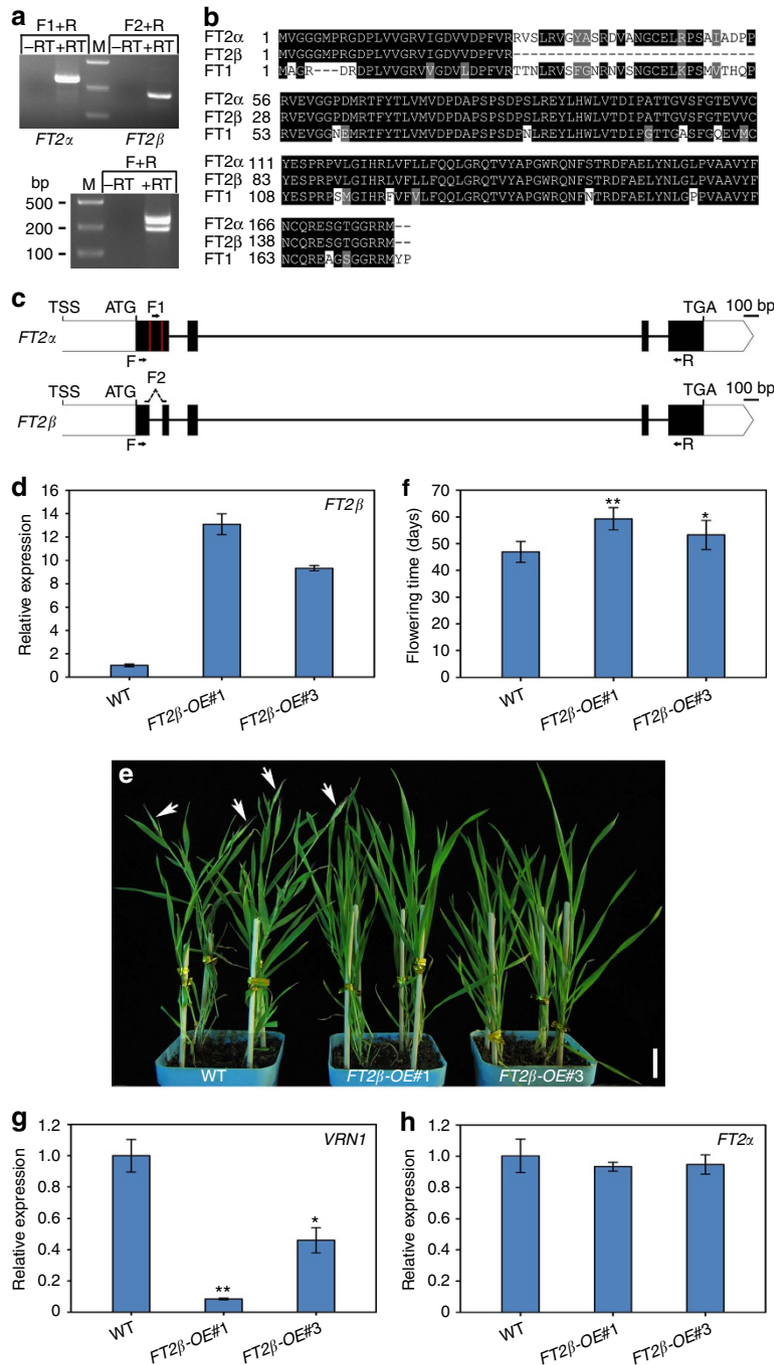


Figure 1 | Identification of FT2AS in *B. distachyon*. (a) Detection of alternatively spliced FT2 transcripts by RT-PCR. F1 and F2 are specific forward primers for FT2α and FT2β amplification, respectively, while primer F is used for FT2α and FT2β PCR simultaneously. R is the reverse primer. RT, reverse transcription. M, marker. bp, base pairs. (b) Protein sequence alignment of different FT2 alternative isoforms and FT1. (c) Schematic genomic structure of FT2 splicing variants. White boxes indicate untranslated regions and black boxes indicate exons. TSS indicates the transcriptional start site. (d) qRT-PCR analysis of FT2β expression in wild-type Bd21 and the indicated FT2β-OE transgenic plants. Three four-week-old *B. distachyon* plants were pooled and collected for RNA isolation. *UBC18* was used as an internal control for normalization of qRT-PCR results. qRT-PCR analyses were performed in three biological replicates with similar results. The point represents the mean value of three technical replicates in a representative biological experiment. Error bars indicate s.d. (e) Representative phenotype of FT2β overexpressing transgenic plants. White arrows point to spikes. Scale bar, 2 cm. (f) Flowering time of wild-type and the indicated FT2β-OE transgenic plants. Error bars indicate s.d. (n = 13). Student's *t*-test, ***P* < 0.01, **P* < 0.05. (g) qRT-PCR analysis of downstream flowering gene *VRN1* expression in wild-type and the indicated FT2β-OE transgenic plants. The four-week-old whole plant tissues including leaves and shoot apex were collected for *VRN1* examination. qRT-PCR analyses were performed in three biological replicates with similar results. The point represents the mean value of three technical replicates in a representative biological experiment. Error bars indicate s.d. Student's *t*-test, ***P* < 0.01, **P* < 0.05. (h) qRT-PCR analysis of FT2α expression levels in wild-type and the indicated FT2β-OE transgenic plants.

growth^{14,40}, demonstrating the positive florigen activity of FT2 α . Next, we generated FT2 β overexpression (FT2 β -OE) transgenic plants and determined the flowering time in their T₁ generations under LD conditions. Wild-type and transgenic plants displayed similar rates of vegetative development. However, eight independent positive FT2 β -OE plants exhibited significantly late heading dates compared with wild-type and the null segregants of the transgenic plants (Supplementary Fig. 4).

Because FT2 β -OE#1 and FT2 β -OE#3 displayed a severe and mild delay of flowering time, respectively, we chose their T₃ homozygous plants for further molecular and phenotypic analyses. Quantitative real-time PCR (qRT-PCR) showed that FT2 β was approximately 12- and 9-fold higher in FT2 β -OE#1 and FT2 β -OE#3 plants, respectively, than in the parental wild-type (Fig. 1d; Supplementary Fig. 5a,b). FT2 β -OE#1 bolted 12 days (59 versus 47 days) and FT2 β -OE#3 bolted more than 6 days (53 versus 47 days) later than wild-type plants under LD conditions (Fig. 1e,f). Furthermore, the expression of VRN1, an APETALA1 orthologue acting downstream of FT in temperate grasses, was dramatically reduced in FT2 β -OE#1 and FT2 β -OE#3 plants (Fig. 1g; Supplementary Fig. 5c,d). The FT2 β -OE plants exhibited wild-type levels of FT2 α mRNA, excluding the possibility that the late flowering of FT2 β -OE was caused by the suppression of FT2 α expression (Fig. 1h; Supplementary Fig. 5e,f). Thus FT2 α and FT2 β may have different effects on floral transition in *B. distachyon*.

FT2 β does not interact with FD or 14-3-3 proteins. In *A. thaliana* and rice, during transport from leaves to SAM, florigen interacts with several 14-3-3 proteins in the cytoplasm and subsequently translocates to the nucleus to bind FD and form a flowering activation complex.

The interaction of TaFDL2, the FD-like protein in wheat, with FT1 and FT2 has been previously demonstrated^{14,42}. Thus we cloned its orthologue from *B. distachyon* and assigned the same name as in wheat (Supplementary Fig. 6). To address whether FT2 α and FT2 β in *B. distachyon* can form a flowering initiation complex with FD, we first used yeast two-hybrid assays to determine their interactions with FDL2. Strong interactions were detected between FT2 α and FDL2 in yeast cells. However, there was no detectable binding of FT2 β to FDL2 in yeast (Fig. 2a). To verify these results, we examined the associations between the two FT2 isoforms and FDL2 using bimolecular fluorescence complementation (BiFC) assays with a split yellow fluorescent protein (YFP) system in *Nicotiana benthamiana* leaf mesophyll cells. As shown in Fig. 2b, there was a strong fluorescent signal in the nucleus when FT2 α was transiently co-expressed with FDL2. By contrast, no YFP fluorescence could be detected in leaves co-expressing FT2 β and FDL2, suggesting that FT2 β cannot bind FDL2 *in vivo*. Co-immunoprecipitation (Co-IP) assays further confirmed a strong binding between FT2 α and FDL2 but no interaction between FT2 β and FDL2 (Fig. 2c).

Similar to Heading date 3a in rice, wheat FT1 can also interact with two 14-3-3 proteins, GF14b and GF14c^{6,42}. Thus we selected several *B. distachyon* 14-3-3 proteins including GF14b and GF14c to determine their abilities to interact with FT2 α and FT2 β (Fig. 2d; Supplementary Fig. 7a). Both GF14b and GF14c interacted with FT2 α in the yeast two-hybrid and BiFC assays (Fig. 2d,e). In contrast, FT2 β did not bind to any 14-3-3 proteins (Fig. 2d,e; Supplementary Fig. 7b). Thus 14-3-3 proteins may not form bridges with FT2 β to trigger flowering in *B. distachyon*. Together, these observations suggest that FT2 splicing variants may have different actions, with one forming a flowering activation complex as a result of direct physical interactions with 14-3-3s and FD, while the other has lost this ability.

FT2 β attenuates the bindings of florigen to FD and 14-3-3s. Genetic engineering experiments have shown that unnatural splicing variants can act as dominant-negative mutants to interrupt the activities of wild-type isoforms^{24,25,28}. FT2 β 's inability to bind to FD and 14-3-3s implied that it may interfere with flowering activation complex formation. To test this possibility, we expressed FT2 β in yeast cells, which contained BD-FT2 α and AD-FDL2 fusions (Fig. 3a). β -Galactosidase (β -Gal) activity assays revealed that the expression of FT2 β substantially repressed FT2 α -FD binding (Fig. 3b). Likewise, the interaction between FT2 α and GF14b in yeast three-hybrid systems was significantly reduced when FT2 β was expressed (Fig. 3b).

To further investigate the negative effects of FT2 β on flowering activation complex formation, we transiently co-expressed FT2 β with split YFP-tagged FT2 α and FDL2 proteins in *N. benthamiana* leaf mesophyll cells. Compared with the GUS protein, which was expressed instead of FT2 β as a control, the numbers of fluorescent cell nuclei, as well as the intensity of fluorescence, generated by the interaction of FT2 α and FDL2 was reduced when FT2 β was co-expressed (Fig. 3c,d). Meanwhile, in line with the yeast three-hybrid assay results, competition BiFC experiments also showed a repressive effect of FT2 β on the association of FT2 α with GF14b in the cytoplasm of *N. benthamiana* leaf cells (Fig. 3e,f).

Because FT1 and FT2 may have redundant roles in flowering initiation^{4,14,40}, we investigated whether FT2 β could affect the efficiency of flowering induction complex formed by FT1, in addition to that formed by FT2 α . Indeed, FT2 β had an inhibitory effect in the competition BiFC assay (Supplementary Fig. 8). Collectively, these results indicate that FT2 β functions as a repressor of the formation of the flowering initiation complex containing FT1, FT2 α , 14-3-3s and FD.

FT2 β forms heterodimers with FT2 α and FT1. FT2 β suppresses the interactions of florigen proteins with FD and 14-3-3s; however, we did not detect any physical interactions of FT2 β with FDL2 or 14-3-3s (Fig. 2). Thus we next investigated another possible mechanism.

It has been suggested that heterodimers may be generated by different splicing isoforms^{24,25,28}, thus we investigated whether FT2 β was able to form a heterodimer with FT2 α to produce a nonfunctional protein complex. To test this possibility, we first performed a yeast two-hybrid analysis and observed a strong interaction between FT2 α and FT2 β (Fig. 4a). The formation of a heterodimer between FT2 α and FT2 β was also detected in the nuclei as well as in the cytoplasm of *N. benthamiana* leaf cells using BiFC assays (Fig. 4b). Furthermore, a yeast two-hybrid analysis showed an interaction of FT2 β with FT1, suggesting that FT2 β is able to form heterodimers with both florigen proteins (Supplementary Fig. 9).

Interestingly, yeast two-hybrid and BiFC assays revealed that both FT2 α and FT1, but not FT2 β , could form homodimers (Fig. 4a–c; Supplementary Fig. 9). Thus we propose that the negative effect of FT2 β on flowering time results from the binding of FT2 β to FT2 α and FT1, which prevents them from interacting with FD and 14-3-3s, thereby reducing the number of functional flowering initiation complexes.

FT2 AS is controlled by an endogenous cue. Having discovered a molecular mode by which FT2 β represses flowering, we wondered when FT2 β contributes to the regulation of floral initiation in *B. distachyon*. Unlike in *A. thaliana*, temperature has a very limited impact on the timing of phase transition from the vegetative to reproductive stage in *B. distachyon*⁴³; therefore, FT2 splicing may have little, if any,

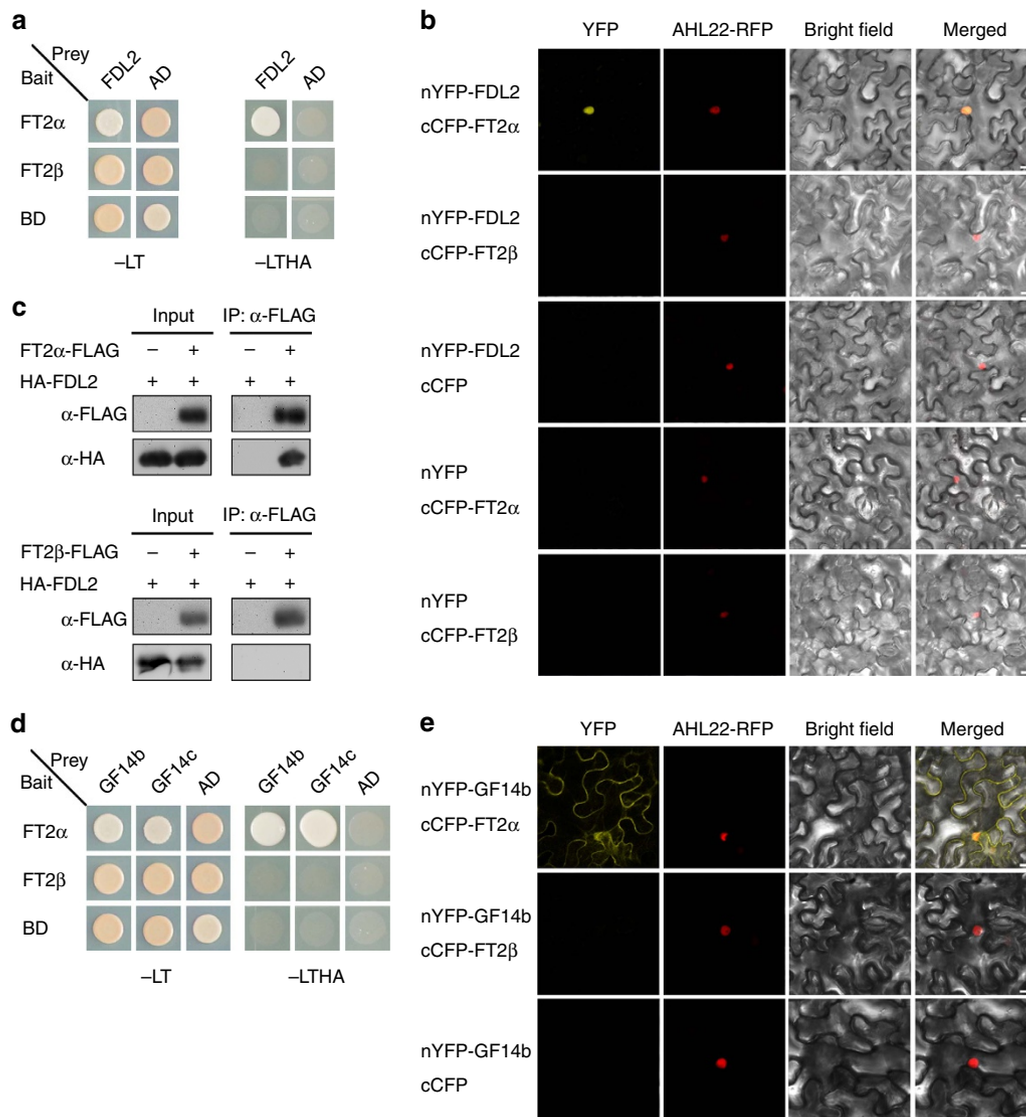


Figure 2 | Interaction analysis of FT2 α and FT2 β with FD-like and 14-3-3 proteins. (a) Interaction analysis of FT2 α and FT2 β with FD-like protein (FDL2) in yeast cells. Yeast cell growth on selective media without Leu, Trp, His and Ade (–LTHA) indicates positive interactions. **(b)** BiFC interaction assays of FT2 α and FT2 β with FDL2. Partial YFP fusion constructs were expressed transiently in *N. benthamiana* leaves. AHL22-RFP was used as a nuclear marker. Scale bar, 20 μ m. **(c)** *In vivo* Co-IP assays of two FT2 splicing isoforms with FDL2. FLAG-tag FT2 α or FLAG-tag FT2 β were co-expressed with HA-tag FDL2 in *N. benthamiana* leaves. **(d)** Yeast two-hybrid analyses of interactions between FT2 isoforms and two 14-3-3 proteins (GF14b and GF14c). **(e)** BiFC interaction assays of FT2 α and FT2 β with 14-3-3 protein GF14b. Scale bar, 20 μ m.

relevance to temperature-mediated flowering, even though the FT2 β /FT2 α ratio can change in response to the environmental temperature.

Previously, we reported that FT1 and FT2 transcription level gradually increase with development under LD conditions in *B. distachyon*¹⁴. Thus we asked whether FT2 AS changes with plant age. Using a qRT-PCR analysis with splicing-variant-specific amplification primers, the ratio of FT2 β /FT2 α abundance was found to decline during plant growth (Fig. 5a,b; Supplementary Figs 10 and 11). Although the transcription levels of FT2 α and FT2 β gradually increased throughout plant development as a result of photoperiodic induction, the proportions of different FT2-splicing isoforms changed with increasing plant age. FT2 β accumulated to a much higher level than FT2 α during the first 4 weeks, but FT2 α rose much faster than FT2 β afterwards (Fig. 5a; Supplementary Figs 10 and 11), thereby resulting in a gradual decrease in the FT2 β /FT2 α ratio as development progressed (Fig. 5b).

To further determine the effects of increasing age on FT2 AS, we examined the diurnal expression rhythms of FT2 α and FT2 β in 4- and 7-week-old *B. distachyon* leaves. As shown in Fig. 5c and Supplementary Fig. 12, the FT2 β expression level was higher than that of FT2 α in 4-week-old plants during the day-night cycle. By contrast, more FT2 α accumulated than FT2 β in 7-week-old plants (Fig. 5d; Supplementary Fig. 12), further confirming the development-dependent change in FT2 AS.

Therefore, in addition to the transcriptional control of FT, the posttranscriptional regulation of FT by AS may also contribute to flowering at the appropriate time in *B. distachyon*. We propose that a high level of FT2 β in young plants may prevent precocious flowering, which allows plants to accumulate sufficient biomass to produce enough seeds to propagate the next generation.

Down regulation of FT2 β induces early flowering. We previously found that the overexpression of miR5200, which

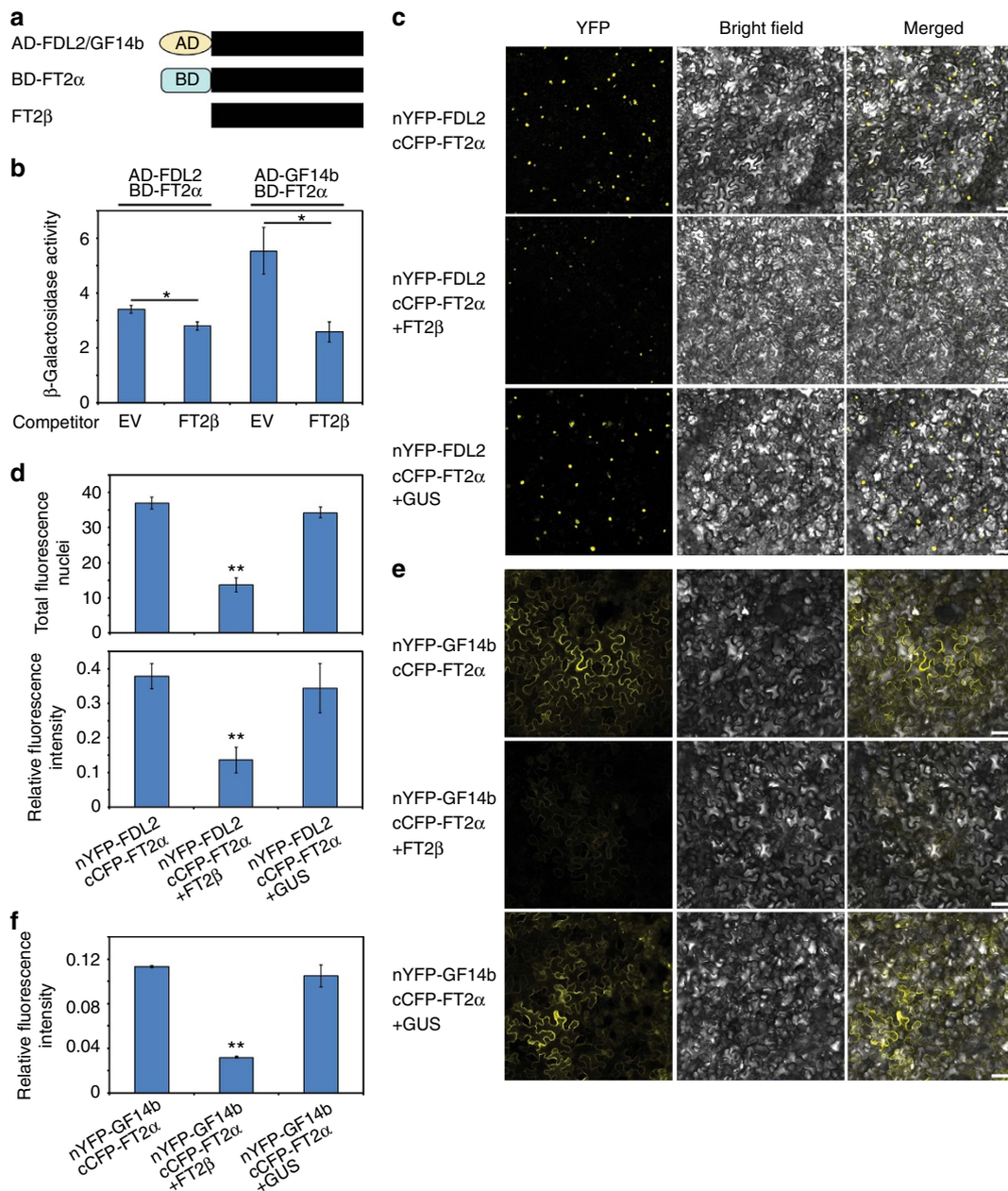


Figure 3 | Attenuation of FT2 α -binding capacity with FD-like and 14-3-3 proteins by FT2 β . (a) Constructs used for yeast three-hybrid assays. FT2 β was used as the competitor to the interacting proteins with activation domain (AD) and DNA-binding domain (BD). (b) Relative quantification of inhibition activities of FT2 β in yeast three-hybrid assays. β -Gal activities were measured in the presence or absence of FT2 β (Student's *t*-test, **P* < 0.05). Bars indicate s.e.m. of three replicates. EV, empty vector. (c) Representative photographs of fluorescence signals in BiFC assays for determination of FT2 β effects on FT2 α and FDL2 interactions in *N. benthamiana* leaf cells. GUS was used as a control in the absence of FT2 β . Scale bar, 50 μ m. (d) Total fluorescence nuclei and relative fluorescence intensity in BiFC assays used for analyses of FT2 β inhibition activities to FT2 α and FDL2 interactions. Bars indicate s.e.m. of three replicates (Student's *t*-test, ***P* < 0.01). (e) Representative photographs showing the fluorescence signals in BiFC assays for determination of FT2 β effects on FT2 α and 14-3-3 interactions in *N. benthamiana* leaf cells. GUS was used as a control in the absence of FT2 β . Scale bar, 50 μ m. (f) Relative fluorescence intensity in BiFC assays used for determination of FT2 β -inhibitory activities to FT2 α and 14-3-3 interactions. Bars indicate s.e.m. of three replicates (Student's *t*-test, ***P* < 0.01).

simultaneously targets both *FT1* and *FT2*, led to a severe delay of flowering onset in *B. distachyon*¹⁴, suggesting that the suppression of *FT2 α* expression may result in late flowering. To further characterize the role of FT2 β in flowering control, we attempted to use artificial miRNAs to knock down *FT2 β* expression, because artificial miRNA can specifically silence one splicing variant but not the other.

In plants, most of miRNAs start with a 5'uridine (U) for their correct sorting into ARGONAUTE 1 (AGO1) silencing

complex⁴⁴. However, there is no adenine (A) nucleotide available for the design of artificial miRNA with a 5'U across the *FT2 β* transcripts that skips the exon nucleotides of *FT2 α* (Supplementary Fig. 13a). Even beginning with a 5'A, miR172 can be properly loaded into AGO1-silencing effectors as a result of its specific secondary structure⁴⁴. Thus we designed an artificial miRNA using *MIR172a* as a backbone to downregulate *FT2 β* (Fig. 6a; Supplementary Fig. 13a,b). We successfully obtained eight independent transgenic Bd21-3 plants and termed them

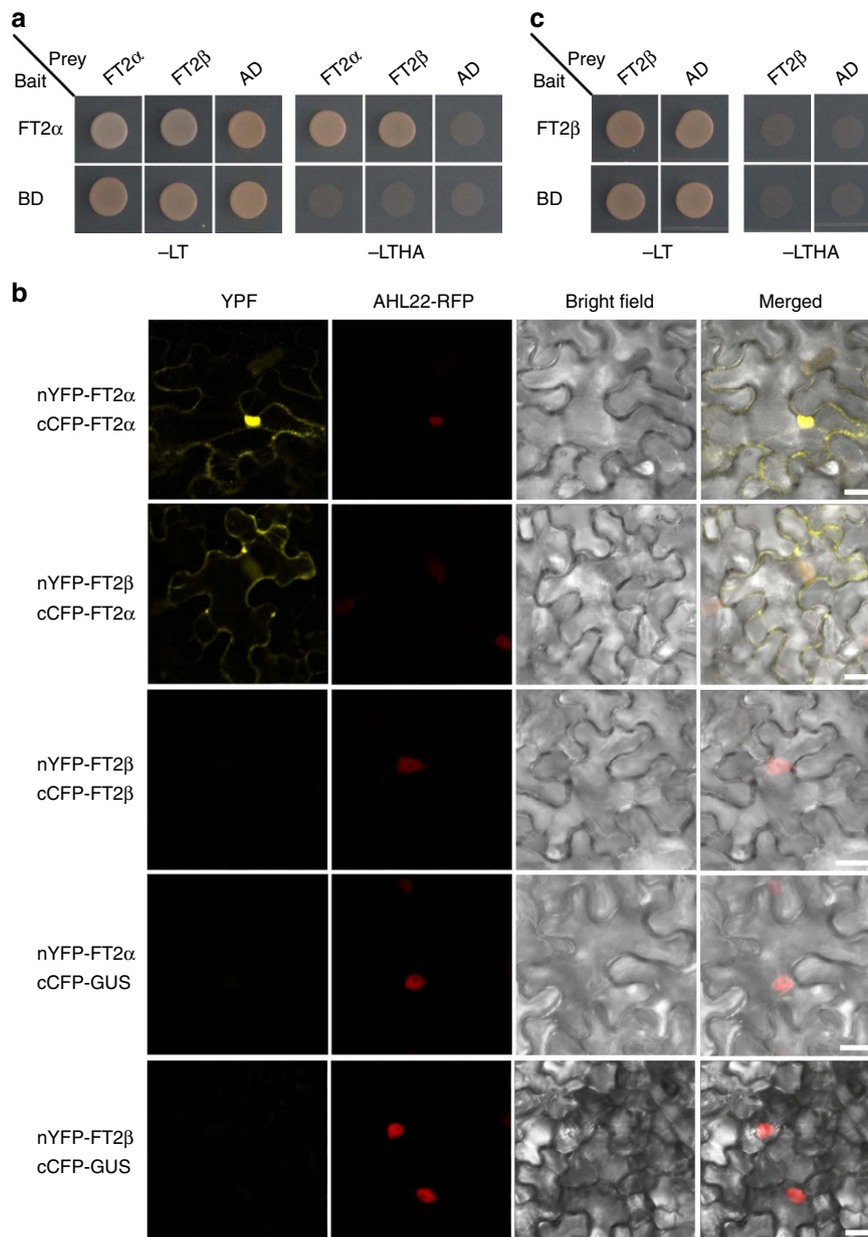


Figure 4 | Homodimer and heterodimer formation analyses between FT2 α and FT2 β . (a) FT2 α can form homodimer and heterodimer with FT2 β in yeast cells. (b) BiFC assays for FT2 α and FT2 β homodimer and heterodimer formation determination. GUS was used as a control in the absence of FT2 α or FT2 β . Scale bar, 20 μ m. (c) FT2 β cannot form homodimer in yeast cells. Yeast cell growth on selective media without Leu, Trp, His and Ade (–LTHA) indicates positive interactions.

amiRFT2 β . RNA analysis revealed high accumulation of artificial miRNA and are resultant decrease of FT2 β expression in typical transgenic plants (Fig. 6b,c; Supplementary Fig. 14a,b). In addition, we did not find reductions in the gene expression levels of FT2 α or other potential *amiRFT2 β* targets in *amiRFT2 β* plants (Fig. 6d; Supplementary Figs 14c,d and 15), suggesting that our artificial miRNA, which was introduced into plants, can specifically act on the desired transcripts.

An analysis of the flowering time of these transgenic plants, in the T₃ generation, under LD conditions revealed early heading (Fig. 6e). The heading date was earlier by an average of 8 days in *amiRFT2 β* compared with those of control plants (54 versus 62 days, on average) (Fig. 6f). Consistent with these phenotypic observations, *VRN1* expression was two fold to four fold higher in *amiRFT2 β* than in wild-type plants (Fig. 6g; Supplementary

Fig. 14e,f). The extent of early flowering phenotype in different *amiRFT2 β* lines correlated with the mature artificial miRNA accumulation as well as the reduced FT2 β expression, suggesting that the acceleration of flowering by reduced FT2 β activity is likely due to the formation of more flowering activation complex and the resulting enhanced *VRN1* expression.

To help exclude the possibility that the early flowering in *amiRFT2 β* is due to the unspecific effects of the artificial miRNA, we generated *amiRFT2 α* transgenic lines, which specifically silenced FT2 α expression by introducing a different artificial miRNA that targeted the intron region of FT2 α where it is different from that of FT2 β (Supplementary Fig. 13). We observed a severe delay of flowering (94 versus 68 days) and a reduction in *VRN1* expression in strong *amiRFT2 α* lines (Supplementary Figs 16 and 17). Weak *amiRFT2 α* lines exhibited

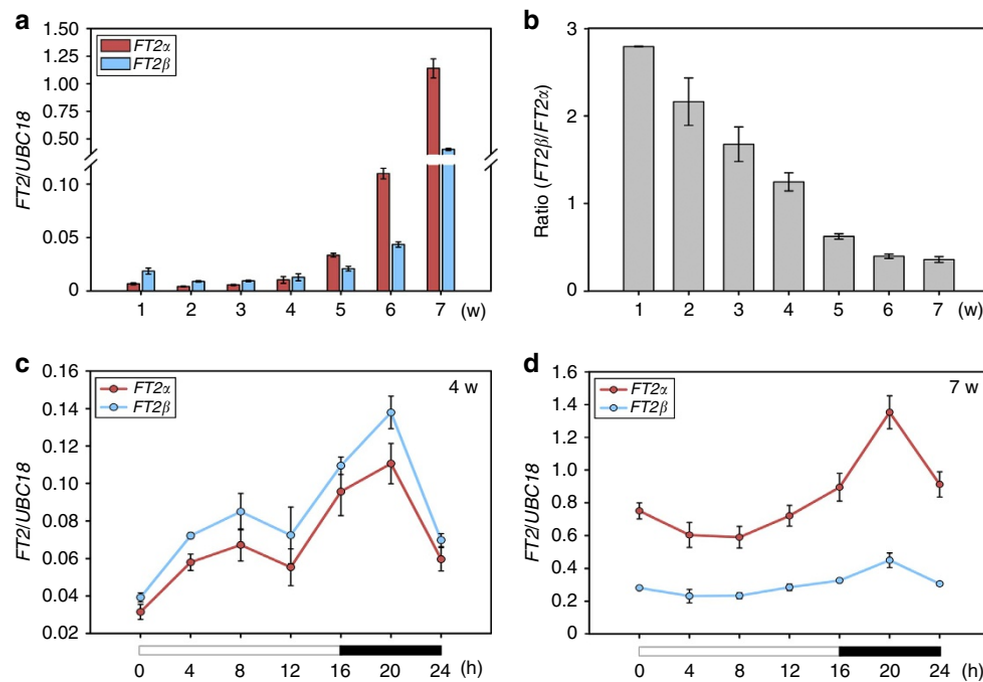


Figure 5 | Age-dependent FT2 AS. (a) Expression analysis of FT2 α and FT2 β along with different age. Upper flat leaves from five Bd21 plants with the indicated age were pooled and collected for RNA isolation. Each qRT-PCR analysis was performed in three biological replicates with similar results. The point represents the mean value of three technical replicates in a representative biological experiment. The transition from vegetative to reproductive stage of Bd21 plants occurs at 4–5 weeks. Standard curve for FT2 α , FT2 β and UBC18 quantification is shown in Supplementary Fig. 2. The absolute value of FT2 α , FT2 β and UBC18 is shown in Supplementary Fig. 10a. Error bars indicate s.d. w, weeks. (b) FT2 β /FT2 α ratio during plant development. (c) and (d) Diurnal expression analysis of FT2 α and FT2 β in 24-h period when plants were 4 and 7 weeks old. The absolute value of FT2 α , FT2 β and UBC18 at each time point is shown in Supplementary Fig. 12. Upper flat leaves from five Bd21 plants at the indicated time were pooled and collected for RNA isolation. The white and black bars along the horizontal axes represent light and dark periods, respectively. The numbers below the horizontal axes indicate the time in hours. Error bars indicate s.d. h, hours, w, weeks.

comparable heading dates with those of wild-type plants (Supplementary Figs 16 and 17). This may be because of the limited decrease in FT2 α expression or the redundant activities of FT1 and FT2 in *B. distachyon*. The lack of early flowering phenotypes in all four *amiFT2 α* lines suggested that the precocious heading of *amiFT2 β* was due to the compromised FT2 β activity rather than the unspecific effects of the artificial miRNA.

Taken together, the acceleration and delay in heading dates in *amiFT2 β* and *amiFT2 α* plants, respectively, further indicated that FT2 β and FT2 α have antagonistic roles in the flowering processes of *B. distachyon*.

Dynamic change in FT2 AS is prevalent in temperate grasses.

In light of the opposite roles of FT2 splicing isoforms in *B. distachyon* flowering control demonstrated above, we asked whether the age-dependent modulation of FT2 AS was conserved among other plants. Using RT-PCR, similar splicing variants of FT2 orthologues in *A. thaliana*, rice and maize were not detected, even though the gene structures of the FT2 homologous genes in rice and maize are the same as that in *B. distachyon* (Supplementary Fig. 18).

As *B. distachyon* belongs to *Pooideae*, we investigated whether AS of FT2 is present in other temperate grasses. Indeed, we found two splicing variants of FT2 in *Aegilops tauschii*, wheat and barley, which differed by the loss of nucleotides that encode the same amino acids as in *B. distachyon* (Supplementary Fig. 19). Additionally, we observed that wheat and barley FT2 β were expressed at much lower levels than FT2 α in 8-week-old plants,

whereas FT2 α transcripts were less abundant in 2-week-old plants (Fig. 7a,b). These results suggest that the change in the AS of FT2 regulated by development is conserved in temperate grasses.

Discussion

AS is a common gene regulatory mechanism that greatly increases the complexity of transcripts and proteins encoded by an organism's genome. From high-throughput transcriptome studies in plants, it is now clear that AS is much more prevalent than previously estimated³⁷. However, when, how and why these splicing variants are generated remains, to a large extent, unknown^{22,23}. In this study, we reported that the highly conserved flowering integrator gene, FT, is also subjected to AS in temperate grasses. FT2 β can heterodimerize with FT2 α and FT1 in plant leaves and is unable to bind FD and 14-3-3s at the shoot apex, resulting in its interference with the formation of flowering inductive complexes. Thus it has a dominant-negative role in controlling flowering onset. The importance of change in FT2 AS during development was demonstrated by the alteration in heading date timing, which results from artificial-miRNA induced reduction in FT2 β activity in *B. distachyon* (Fig. 6).

AS is thought to regulate functional gene actions in two ways: by directly altering mRNA stability or by encoding distinct protein isoforms, which may influence physical interactions between the original protein and its associates²³. The latter is frequently associated with AS that is regulated by environmental stress or temperature response^{24,25,28,45,46}. Although we detected that FT2 β /FT2 α ratio was influenced by ambient temperature

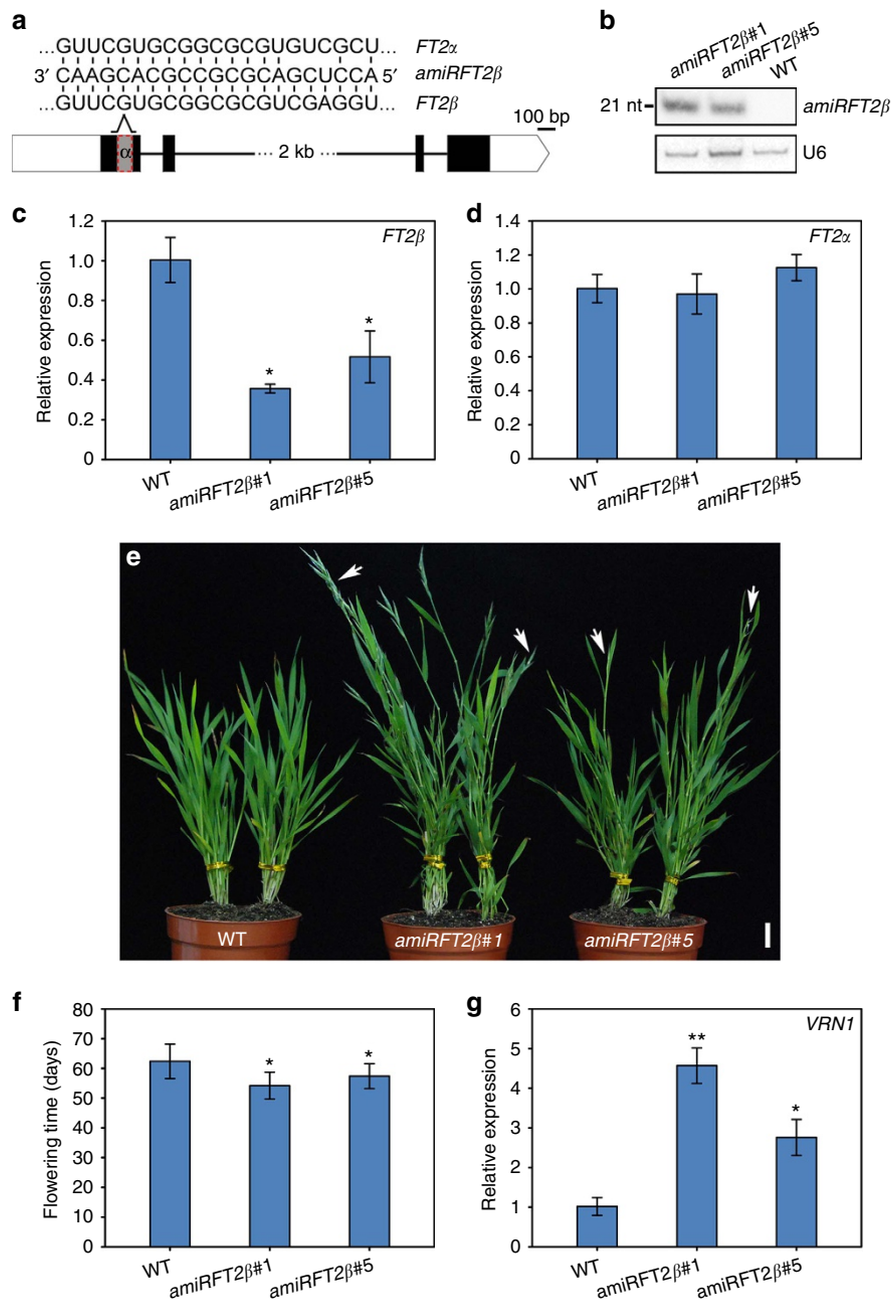


Figure 6 | Interruption of *FT2β* activity by artificial miRNA results in early flowering. (a) Schematic diagram of *amiRFT2β*. (b) Northern blotting analysis of artificial miRNA accumulation in *amiRFT2β* transgenic plants. U6 was used as a loading control for RNA gel blot. (c) qRT-PCR analysis of *FT2β* expression in wild-type Bd21-3 and the indicated *amiRFT2β* transgenic plants. *UBC18* was used as an internal control for normalization of qRT-PCR results. Three six-week-old *B. distachyon* plants were pooled and collected for RNA isolation. Each qRT-PCR analysis were performed in three biological replicates with similar results. The point represents the mean value of three technical replicates in a representative biological experiment. Error bars indicate s.d. Student's *t*-test, **P* < 0.05 (d) qRT-PCR analysis of *FT2α* expression in wild-type and the indicated *amiRFT2β* transgenic plants. (e) Representative photograph of flowering phenotypes in *amiRFT2β* transgenic and wild-type Bd21-3 plants. White arrows point to spikes. Scale bar, 2 cm. (f) Flowering time of wild-type and the indicated two lines of *amiRFT2β* transgenic plants. Error bars indicate s.d. (*n* = 10). Student's *t*-test, **P* < 0.05 (g) qRT-PCR analysis of flowering downstream gene *VRN1* expression in wild-type and the indicated *amiRFT2β* transgenic plants. Total RNA from 5-week-old whole-plant tissues, including leaves and shoot apex, were used for *VRN1* examination. Student's *t*-test, ***P* < 0.01, **P* < 0.05.

changes, the AS of *FT2* may not be involved in temperature-mediated flowering control in *B. distachyon*, as increased ambient temperatures have limited influences on flowering signal induction in temperate grasses⁴³.

Interestingly, the *FT2β*/*FT2α* ratio was progressively reduced during development, indicating that *FT2* AS is regulated by an endogenous cue rather than an external cue for fine-tuning plant

flowering. *FT2* AS may be modulated by some special splicing factors that abundance-, localization- or posttranslational modification-related changes as plant grow. Because miR156 and miR172 participate in the age-dependent regulation of flowering in diverse plants^{12,13}, it will be interesting to explore whether alterations in *FT2* growth-related AS in *B. distachyon* is controlled by these two miRNAs during flowering processes.

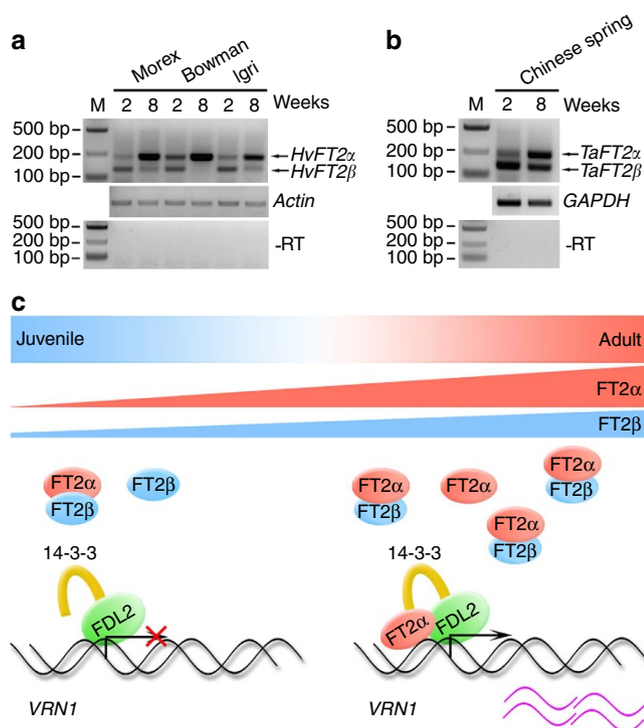


Figure 7 | Dynamic changes in FT2 AS with plant development is prevalent in barley and wheat. (a) RT-PCR analysis of *FT2α* and *FT2β* expression in barley with the indicated age. Primers designed for simultaneous amplification of *FT2α* and *FT2β* in a single PCR reaction were used. Total RNA from three typical varieties of barley was isolated for reverse transcription. M, marker. (b) RT-PCR analysis of *FT2α* and *FT2β* expression in wheat with different ages. Total RNA from Chinese Spring was isolated for reverse transcription. (c) A working model for two *FT2* splicing isoforms during plant flowering-induction processes in temperate grasses. Although both two *FT2* splicing transcripts increase along with plant development, their increasing rates are not consistent, resulting in distinct *FT2β/FT2α* ratio with different age. *FT2β* and *FT2α* can form heterodimers. Thus when plants are vegetative, the formation of flowering-induction complex is blocked, because *FT2β* expression level is higher than that of *FT2α*. When plants grow to reproductive stage, more *FT2α* accumulates than *FT2β*, and the flowering initiation complex can be formed in the SAM to induce *VRN1* expression, ensuring eventual plant flowering at right growth phase.

FT2 and *FT1* in *B. distachyon* are thought to have redundant roles on florigen activity because of their high amino-acid identities, similar temporal and spatial expression patterns, and same early flowering phenotype if overproduced. Thus it seems likely that *FT2* and *FT1* may have an analogous regulatory mechanism^{14,15}. However, *FT2* and *FT1* are not regulated by identical schemes at the posttranscriptional level. First, their regulation by miR5200 is different. The base-pairing of miR5200 with the *FT2* target site is not as perfect as that of *FT1*, resulting in the cleavage of *FT2* by miR5200 at a non-canonical miRNA cleavage site¹⁴. Second, *FT2*'s function is modulated by AS, which is altered during development to repress flowering in the early stage, while no *FT1* AS events occur during the life cycle. Because both AS- and miRNA-mediated gene silencing are posttranscriptional gene regulatory mechanisms in eukaryotes, we propose that posttranscriptional modulation may be the major force for differentiating and fine-tuning different FT-like protein activities in *Pooideae* plants during evolution.

In *A. thaliana*, BROTHER OF FT AND TFL1 (BFT), a member of the FT/TFL1 protein family, mediates the link between

flowering and stress signaling⁴⁷. BFT is expressed at low levels, irrespective of day length, but can be dramatically induced under highly saline conditions. BFT and FT have similar abilities to bind FD. Thus, when plants encounter salt stress, BFT causes delayed flowering by competing with FT for FD binding⁴⁷. Similarly, TFL1 represses FT activity through its interactions with FD, which is involved in the prevention of flowering in immature meristems of *A. thaliana*, *A. alpina* and even apple trees^{48,49–52}. In *B. distachyon*, although developmentally regulated *FT2β* also interferes with flowering initiation complex formation, its molecular mechanism is different from that of BFT in *A. thaliana* because *FT2β* does not bind FD. In a sense, *FT2β* functions as a competitor of FD, because both *FT2β* and FD can dimerize with *FT2α*. *FT2β/FT2α* ratio is high in young plants and this prevents the rapid progression towards flowering. If there was no *FT2β* generated at the vegetative stage, then the florigen activity conferred by either *FT2α* or *FT1* would immediately exceed the threshold for the reproductive phase transition, which may lead to precocious flowering and detrimental effects on seed production.

We found that full-length *FT2* in *B. distachyon* can form homodimers and *FT2-FT1* heterodimers. Similar findings that FT interacts with itself in *A. thaliana* suggest a possible biological relevance for the multimeric forms of FT proteins in plants⁴⁸. Hence, further studies on when and where FT proteins form polymers *in vivo* will be important for determining their exact roles in accelerating flowering.

Plants contain complex regulatory machineries to optimize the seasonal timing of flowering. Our present discovery of AS-mediated *FT2* regulation in temperate grasses not only extends the manner of flowering control by FT but also reveals florigen control at both the transcriptional and posttranscriptional levels during the process of floral transition (Fig. 7c). In respect that multiple alternatively spliced forms of FT orthologous genes have been detected in *Platanus acerifolia*⁵³, it will be interesting to determine whether the blocking flowering complex formation by splice variants is the same FT regulatory mechanism in trees.

Methods

Plant materials and growth conditions. *B. distachyon* used in this study were grown under LD conditions (16 h light/8 h dark) in growth chambers with temperatures of 22 °C during the day and 16 °C at night. For most experiments, the *B. distachyon* accession Bd21 was used. To minimize the crosstalk effects of photoperiod and vernalization on Bd21-3, seeds of transgenic and control plants were placed on moistened filter paper in Petri dishes and subjected to 4 °C cold treatment for 1 week before planting under the indicated conditions. Unless stated otherwise, all plant seedlings or leaves were harvested at Zeitgeber time 16 for use in the experiments.

Constructs and plant transformation. The *FT2* (*FT2α* and *FT2β*) coding sequences and the miR172a backbone of the artificial miRNA were amplified from *B. distachyon* cDNA. *FT2α*, *FT2β* and artificial miRNAs constructs were all under the control of the maize (*Zea mays*) *UBIQUITIN (UBI)* promoter for overexpression in Bd21 or Bd21-3 plants. The sequences of the primers for *B. distachyon FT2α*, *FT2β* and artificial miRNA amplification, as well as for binary vector construction, are listed in Supplementary Table 1. Transformation of *B. distachyon* was mediated by *Agrobacterium tumefaciens* strain AGL1 using compact embryogenic calli derived from immature embryos⁵⁴. Independent transgenic lines were genotyped for T-DNA using *HPT II*-specific forward and reverse primers. The confirmed positive T₃ transgenic plants were used in flowering time determination and gene expression assays.

Plant flowering time measurements. The flowering times of the indicated plants were measured as the number of days from the date of planting in soil to the day when the first spike emerged. At least 10 plants of each transgenic line were recorded to calculate the average flowering time. Statistical significance was determined using Student's *t*-test.

RNA expression analysis. Total RNA was isolated from at least three pooled *B. distachyon* plants using TRIzol reagent (Invitrogen). The total RNA sample was pretreated with RNase-free DNase I (Promega) prior to cDNA synthesis to remove DNA and then reverse transcribed by EasyScript reverse transcriptase (TransGen Biotech) using oligo (dT) primers. RT-PCR analyses of gene transcripts were essentially carried out as previously described²⁵. qRT-PCR reactions were performed with SYBR Premix Ex Taq (Takara) and run on an ABI 7,500 Fast system with the following procedure: 95 °C for 1 min, then 40 cycles of 95 °C for 10 s, 62 °C for 10 s, and 72 °C for 30 s, followed by a melting dissociation program.

For the quantification of *FT2 α* and *FT2 β* transcripts, *FT2 α* and *FT2 β* cDNAs were cloned into pGADT7 vectors, and an absolute standard curve of each transcript was made using 10-fold serial dilutions from 10⁻¹⁸ to 10⁻²³ mol μ l⁻¹. The gene expression levels (Fig. 5 and Supplementary Fig. 3) were calculated based on the ratio of absolute quantifications of *FT2 α* and *FT2 β* to the corresponding absolute value of *UBC18*. For gene relative expression analysis, *UBC18* mRNA was examined in parallel and used for data normalization, and the relative expression levels were calculated using the $\Delta\Delta$ Ct method.

All of the qRT-PCR analyses were performed at least in three independent biological replicates with three technical repetitions. For artificial miRNA detection, 2 μ g total RNA was used for RNA gel blots. ³²P-end-labelled oligonucleotides complementary to the artificial miRNA sequences were used as probes. The sequences of the primers and small RNA blot probes are listed in Supplementary Table 1.

Yeast assays. Yeast two-hybrid assays were conducted using the Matchmaker GAL4 Two-Hybrid System (Clontech). The PCR products of *FT2 α* , *FT2 β* , *FT1*, *FDL2* and diverse *14-3-3s* from *B. distachyon* cDNA were independently cloned into the pGBKT7 and pGADT7 vectors. Transformation of AH109 cells was performed according to the general protocol. The transformed yeast cells were spread on the selective medium SD/-Leu/-Trp/-His/-Ade plus 2.5 mM 3-amino-triazole to determine the interactions.

For the yeast three-hybrid assay, bait and prey were co-transformed with or without the competitor *FT2 β* into yeast AH109 cells. The interaction strengths were quantified by a liquid β -Gal assay using chlorophenol red- β -D-galactopyranoside (CPRG) as substrates according to the Yeast Protocol Handbook (Clontech). Briefly, the yeast cells were broken open using several freeze/thaw cycles and mixed with CPRG buffer. When the colour of the sample turned to red, 3.0 mM ZnCl₂ was added to stop the reaction. The β -Gal units were calculated by the supernatant absorbance at 578 and 600 nm.

BiFC assays. To generate the constructs for the BiFC assays, full-length cDNA fragments of *FT2 α* , *FT2 β* , *FT1*, *FDL2* and *GF14b* were amplified and cloned into the pDONRZeo (Invitrogen) vector for fusion with the N-terminus of YFP or the C-terminus of CFP by LR reaction. The resulting plasmids were introduced into *A. tumefaciens* EHA105, which were then infiltrated into 4-week-old *N. benthamiana* leaves for transient expression. In brief, the *A. tumefaciens* strains were resuspended to OD₆₀₀ = 0.8 in infiltration buffer (150 μ M acetosyringone, 10 mM MgCl₂ and 10 mM MES). The resuspensions containing each BiFC construct as well as the p19 plasmid were mixed at 1:1:1 ratio and then infiltrated into leaves with a syringe.

For the protein-binding competition assay, the *A. tumefaciens* harbouring the *FT2 β* plasmid was co-infiltrated with an equal volume of the indicated construct. *A. tumefaciens* harbouring *GUS* in place of *FT2 β* was infiltrated in parallel as a control. The tobacco leaves were imaged under Zeiss LSM 700 confocal microscope 3 days after infiltration. Excitation of YFP was performed using an argon laser line at 514 nm and red fluorescent protein (RFP) with 563 nm, while capture of YFP fluorescence was at 518–555 nm and RFP at 568–636 nm. Primers used to construct BiFC plasmids are shown in Supplementary Table 1.

Co-IP assays. Full-length coding sequences of *FT2 α* , *FT2 β* and *FDL2* were cloned into tagging plasmids under the control of Cauliflower Mosaic Virus 35S promoter. The 4-week-old *N. benthamiana* leaves were infiltrated with an *A. tumefaciens* strains harbouring *FT2 α* -FLAG plus HA-FDL2 or *FT2 β* -FLAG plus HA-FDL2. The infected leaves were collected 3 days after infiltration. Total protein was then extracted with IP buffer (50 mM Tris-HCl pH 7.5, 150 mM NaCl, 1 mM EDTA, 1% Triton X-100, 10% glycerol, 1 mM phenylmethyl sulfonyl fluoride, 1 mM NaVO₄, 50 μ M MG132 and 1 \times protease inhibitor cocktail) and centrifuged three times at 16,000g at 4 °C. The supernatant was incubated with Anti-Flag M2 Affinity Gel (Sigma, A2220) at 4 °C for 3 h with gentle shaking. After centrifugation at 600g at 4 °C for 1 min, the agarose beads were washed five times with washing buffer (50 mM Tris-HCl pH 7.5, 150 mM NaCl, 1 mM EDTA, 0.1% Triton X-100). The precipitate was suspended, boiled in SDS loading buffer, separated by an SDS-PAGE and detected by anti-HA (Sigma, H6908, 1:5,000 dilution) or anti-FLAG antibody (Sigma, H7425, 1:3,000 dilution). The full scan picture of Co-IP assay results is shown in Supplementary Fig. 20.

Data availability. The sequence of *FT2 β* has been submitted to Genbank under accession code (KP637176). The authors declare that all other data that support the

findings of this study are available in the manuscript and its Supplementary files or are available from the corresponding author upon request.

References

- Yu, S. *et al.* Gibberellin regulates the *Arabidopsis* floral transition through miR156-targeted SQUAMOSA promoter binding-like transcription factors. *Plant Cell* **24**, 3320–3332 (2012).
- Kobayashi, Y., Kaya, H., Goto, K., Iwabuchi, M. & Araki, T. A pair of related genes with antagonistic roles in mediating flowering signals. *Science* **286**, 1960–1962 (1999).
- Abe, M. *et al.* FD, a bZIP protein mediating signals from the floral pathway integrator FT at the shoot apex. *Science* **309**, 1052–1056 (2005).
- Li, C. & Dubcovsky, J. Wheat FT protein regulates VRN1 transcription through interactions with FDL2. *Plant J.* **55**, 543–554 (2008).
- Wigge, P. A. *et al.* Integration of spatial and temporal information during floral induction in *Arabidopsis*. *Science* **309**, 1056–1059 (2005).
- Taoka, K. *et al.* 14-3-3 proteins act as intracellular receptors for rice Hd3a florigen. *Nature* **476**, 332–335 (2011).
- Sawa, M. & Kay, S. A. GIGANTEA directly activates *Flowering Locus T* in *Arabidopsis thaliana*. *Proc. Natl Acad. Sci. USA* **108**, 11698–11703 (2011).
- Wenkel, S. *et al.* CONSTANS and the CCAAT box binding complex share a functionally important domain and interact to regulate flowering of *Arabidopsis*. *Plant Cell* **18**, 2971–2984 (2006).
- Tiwari, S. B. *et al.* The flowering time regulator CONSTANS is recruited to the FLOWERING LOCUS T promoter via a unique cis-element. *New Phytol.* **187**, 57–66 (2010).
- Li, D. *et al.* A repressor complex governs the integration of flowering signals in *Arabidopsis*. *Dev. Cell* **15**, 110–120 (2008).
- Mathieu, J., Yant, L. J., Murdter, F., Kuttner, F. & Schmid, M. Repression of flowering by the miR172 target SMZ. *PLoS Biol.* **7**, e1000148 (2009).
- Wu, G. *et al.* The sequential action of miR156 and miR172 regulates developmental timing in *Arabidopsis*. *Cell* **138**, 750–759 (2009).
- Wang, J. W., Czech, B. & Weigel, D. miR156-regulated SPL transcription factors define an endogenous flowering pathway in *Arabidopsis thaliana*. *Cell* **138**, 738–749 (2009).
- Wu, L. *et al.* Regulation of FLOWERING LOCUS T by a microRNA in *Brachypodium distachyon*. *Plant Cell* **25**, 4363–4377 (2013).
- Ream, T. S. *et al.* Interaction of photoperiod and vernalization determines flowering time of *Brachypodium distachyon*. *Plant Physiol.* **164**, 694–709 (2014).
- Turner, A., Beales, J., Faure, S., Dunford, R. P. & Laurie, D. A. The pseudo-response regulator Ppd-H1 provides adaptation to photoperiod in barley. *Science* **310**, 1031–1034 (2005).
- Brkljacic, J. *et al.* *Brachypodium* as a model for the grasses: today and the future. *Plant Physiol.* **157**, 3–13 (2011).
- Woods, D. P., Ream, T. S., Minevich, G., Hobert, O. & Amasino, R. M. PHYTOCHROME C is an essential light receptor for photoperiodic flowering in the temperate grass, *Brachypodium distachyon*. *Genetics* **198**, 397–408 (2014).
- Chen, A. *et al.* Phytochrome C plays a major role in the acceleration of wheat flowering under long-day photoperiod. *Proc. Natl Acad. Sci. USA* **111**, 10037–10044 (2014).
- Pearce, S., Kippes, N., Chen, A., Debernardi, J. M. & Dubcovsky, J. 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**, 141 (2016).
- Syed, N. H., Kalyana, M., Marquez, Y., Barta, A. & Brown, J. W. Alternative splicing in plants—coming of age. *Trends Plant Sci.* **17**, 616–623 (2012).
- Reddy, A. S., Marquez, Y., Kalyana, M. & Barta, A. Complexity of the alternative splicing landscape in plants. *Plant Cell* **25**, 3657–3683 (2013).
- Staiger, D. & Brown, J. W. Alternative splicing at the intersection of biological timing, development, and stress responses. *Plant Cell* **25**, 3640–3656 (2013).
- Seo, P. J., Kim, M. J., Ryu, J. Y., Jeong, E. Y. & Park, C. M. Two splice variants of the IDD14 transcription factor competitively form nonfunctional heterodimers which may regulate starch metabolism. *Nat. Commun.* **2**, 303 (2011).
- Seo, P. J. *et al.* A self-regulatory circuit of CIRCADIEN CLOCK-ASSOCIATED1 underlies the circadian clock regulation of temperature responses in *Arabidopsis*. *Plant Cell* **24**, 2427–2442 (2012).
- James, A. B. *et al.* Alternative splicing mediates responses of the *Arabidopsis* circadian clock to temperature changes. *Plant Cell* **24**, 961–981 (2012).
- Shikata, H. *et al.* Phytochrome controls alternative splicing to mediate light responses in *Arabidopsis*. *Proc. Natl Acad. Sci. USA* **111**, 18781–18786 (2014).
- Pose, D. *et al.* Temperature-dependent regulation of flowering by antagonistic FLM variants. *Nature* **503**, 414–417 (2013).
- Chung, H. S. & Howe, G. A. A critical role for the TIFY motif in repression of jasmonate signaling by a stabilized splice variant of the JASMONATE ZIM-domain protein JAZ10 in *Arabidopsis*. *Plant Cell* **21**, 131–145 (2009).

30. Yan, K. *et al.* Stress-induced alternative splicing provides a mechanism for the regulation of microRNA processing in *Arabidopsis thaliana*. *Mol. Cell* **48**, 521–531 (2012).
31. Jiang, J., Zhang, C. & Wang, X. A recently evolved isoform of the transcription factor BES1 promotes brassinosteroid signaling and development in *Arabidopsis thaliana*. *Plant Cell* **27**, 361–374 (2015).
32. Nicolas, M., Rodriguez-Buey, M. L., Franco-Zorrilla, J. M. & Cubas, P. A recently evolved alternative splice site in the BRANCHED1a gene controls potato plant architecture. *Curr. Biol.* **25**, 1799–1809 (2015).
33. Li, P., Tao, Z. & Dean, C. Phenotypic evolution through variation in splicing of the noncoding RNA COOLAIR. *Genes Dev.* **29**, 696–701 (2015).
34. Kesari, R. *et al.* Intron-mediated alternative splicing of *Arabidopsis* P5CS1 and its association with natural variation in proline and climate adaptation. *Proc. Natl Acad. Sci. USA* **109**, 9197–9202 (2012).
35. Wang, Z. *et al.* ABA signalling is fine-tuned by antagonistic HAB1 variants. *Nat. Commun.* **6**, 8138 (2015).
36. Gou, J. Y. *et al.* Wheat stripe rust resistance protein WKS1 reduces the ability of the thylakoid-associated ascorbate peroxidase to detoxify reactive oxygen species. *Plant Cell* **27**, 1755–1770 (2015).
37. Marquez, Y., Brown, J. W., Simpson, C., Barta, A. & Kalyna, M. Transcriptome survey reveals increased complexity of the alternative splicing landscape in *Arabidopsis*. *Genome Res.* **22**, 1184–1195 (2012).
38. Filichkin, S. A. *et al.* Genome-wide mapping of alternative splicing in *Arabidopsis thaliana*. *Genome Res.* **20**, 45–58 (2010).
39. Mandadi, K. K. & Scholthof, K. B. Genome-wide analysis of alternative splicing landscapes modulated during plant-virus interactions in *Brachypodium distachyon*. *Plant Cell* **27**, 71–85 (2015).
40. Lv, B. *et al.* Characterization of FLOWERING LOCUS T1 (FT1) gene in *Brachypodium* and wheat. *PLoS ONE* **9**, e94171 (2014).
41. Kim, J. Y., Ryu, J. Y., Baek, K. & Park, C. M. High temperature attenuates the gravitropism of inflorescence stems by inducing SHOOT GRAVITROPISM 5 alternative splicing in *Arabidopsis*. *New Phytol.* **209**, 265–279 (2016).
42. Li, C., Lin, H. & Dubcovsky, J. Factorial combinations of protein interactions generate a multiplicity of florigen activation complexes in wheat and barley. *Plant J.* **84**, 70–82 (2015).
43. Boden, S. A., Kavanova, M., Finnegan, E. J. & Wigge, P. A. Thermal stress effects on grain yield in *Brachypodium distachyon* occur via H2A.Z-nucleosomes. *Genome Biol.* **14**, R65 (2013).
44. Mi, S. *et al.* Sorting of small RNAs into *Arabidopsis* argonaute complexes is directed by the 5' terminal nucleotide. *Cell* **133**, 116–127 (2008).
45. Filichkin, S. A. *et al.* Environmental stresses modulate abundance and timing of alternatively spliced circadian transcripts in *Arabidopsis*. *Mol. Plant* **8**, 207–227 (2014).
46. Sureshkumar, S., Dent, C., Seleznev, A., Tasset, C. & Balasubramanian, S. Nonsense-mediated mRNA decay modulates FLM-dependent thermosensory flowering response in *Arabidopsis*. *Nat. Plants* **2**, 16055 (2016).
47. Ryu, J. Y. *et al.* The *Arabidopsis* floral repressor BFT delays flowering by competing with FT for FD binding under high salinity. *Mol. Plant* **7**, 377–387 (2014).
48. Matsoukas, I. G., Massiah, A. J. & Thomas, B. Starch metabolism and antiflorigenic signals modulate the juvenile-to-adult phase transition in *Arabidopsis*. *Plant Cell Environ.* **36**, 1802–1811 (2013).
49. Hanano, S. & Goto, K. *Arabidopsis* TERMINAL FLOWER1 is involved in the regulation of flowering time and inflorescence development through transcriptional repression. *Plant Cell* **23**, 3172–3184 (2011).
50. Wang, R. *et al.* Aa TFL1 confers an age-dependent response to vernalization in perennial *Arabis alpina*. *Plant Cell* **23**, 1307–1321 (2011).
51. Kotoda, N. *et al.* Molecular characterization of FLOWERING LOCUS T-like genes of apple (*Malus x domestica* Borkh.). *Plant Cell Physiol.* **51**, 561–575 (2010).
52. Jaeger, K. E., Pullen, N., Lamzin, S., Morris, R. J. & Wigge, P. A. Interlocking feedback loops govern the dynamic behavior of the floral transition in *Arabidopsis*. *Plant Cell* **25**, 820–833 (2013).
53. Zhang, J. *et al.* The FLOWERING LOCUS T orthologous gene of *Platanus acerifolia* is expressed as alternatively spliced forms with distinct spatial and temporal patterns. *Plant Biol. (Stuttg.)* **13**, 809–820 (2011).
54. Alves, S. C. *et al.* A protocol for Agrobacterium-mediated transformation of *Brachypodium distachyon* community standard line Bd21. *Nat. Protoc.* **4**, 638–649 (2009).
55. Wu, L. *et al.* DNA methylation mediated by a microRNA pathway. *Mol. Cell* **38**, 465–475 (2010).

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Author contributions

L.W. and L.M. conceived and supervised the research. L.W., L.M., Z.Q. and J.W. designed the experiments. Z.Q., J.W., G.S., F.C., N.F., X.K., S.G., K.C. and L.W. performed the experiments. Z.Q., J.W., A.L., L.M. and L.W. analyzed the data. L.W., L.M. and Z.Q. wrote the article.

Additional information

Supplementary Information accompanies this paper at <http://www.nature.com/naturecommunications>

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