#### 1 **Short Title:** 2 Targets of the flowering time regulator FD 3 4 **Author for Contact:** 5 Markus Schmid 6 7 **Article title:** 8 FT modulates genome-wide DNA-binding of the bZIP transcription factor FD in 9 Arabidopsis thaliana 10 11 **Author names:** Silvio Collani<sup>1,2</sup>, Manuela Neumann<sup>2</sup>, Levi Yant<sup>2,#</sup>, Markus Schmid<sup>1,2,3,\*</sup> 12 13 14 **Author affiliations:** 15 <sup>1</sup>Umeå Plant Science Centre, Department of Plant Physiology, Umeå University, SE-16 901 87 Umeå, Sweden. 17 <sup>2</sup>Max Planck Institute for Developmental Biology, Department of Molecular Biology, 18 72076 Tübingen, Germany. 19 <sup>3</sup>Beijing Advanced Innovation Centre for Tree Breeding by Molecular Design, Beijing 20 Forestry University, Beijing 100083, People's Republic of China. 21 22 One sentence summary: 23 Genomic and biochemical analyses identify targets of the flowering time regulator FD 24 at the genome-wide scale and shed light on the requirement for interaction with the 25 florigen FLOWERING LOCUS T. 26 27 **Author contributions:** 28 S.C., L.Y., and M.S. designed the experiments. L.Y. established some of the FD:GFP 29 reporter lines and performed initial ChIP (-seq) and flowering time analyses. M.N. 30 cloned phosphomic and non-phosphorable versions of FD and analyzed their effect on 31 flowering time. S.C. performed the EMSA studies, flowering times analysis and carried 32 out and analyzed the ChIP-seq and RNA-seq experiments. S.C. and M.S. wrote the 33 manuscript with input from all authors. M.S. agrees to serve as the corresponding

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#### **ABSTRACT**

The transition to flowering is a crucial step in the plant life cycle that is controlled by multiple endogenous and environmental cues, including hormones, sugars, temperature, and photoperiod. Permissive photoperiod induces the expression of FLOWERING LOCUS T (FT) in the phloem companion cells of leaves. The FT protein then acts as a florigen that is transported to the shoot apical meristem (SAM), where it physically interacts with the bZIP transcription factor FD and 14-3-3 proteins. However, despite the importance of FD for promoting flowering, its direct transcriptional targets are largely unknown. Here, we combined ChIP-seq and RNAseq to identify targets of FD at the genome scale and assess the contribution of FT to DNA binding. We further investigated the ability of FD to form protein complexes with FT and TFL1 through the interaction with 14-3-3 proteins. Importantly, we observe direct binding of FD to targets involved in several aspects of plant development not understood to be directly related to the regulation of flowering time. Our results confirm FD as central regulator of the floral transition at the shoot meristem and provides evidence for crosstalk between the regulation of flowering and other signaling pathways.

#### INTRODUCTION

The floral transition represents a crucial checkpoint in the plant life cycle at which the shoot apical meristem (SAM) ceases to produce only leaves and begins producing reproductive organs. As the commitment to this developmental phase transition is usually irreversible for a given meristem, plants have evolved several pathways to integrate environmental and endogenous stimuli to ensure flowering is induced at the correct time. A rich literature has identified hormones, sugars, temperature, and day length (photoperiod) as main factors in flowering time regulation (reviewed in Srikanth and Schmid, 2011; Romera-Branchat et al., 2014; Song et al., 2015). Photoperiod in particular has been shown to regulate flowering time in many plant species and, depending on the light requirements, short day (SD), long day (LD) and day-neutral plants have been distinguished. In *Arabidopsis thaliana*, LD promotes flowering but plants will eventually flower even under non-inductive SD.

It has long been known that in day-length responsive species, inductive photoperiod is

mainly perceived in leaves where it results in the formation of a long-distance signal,

83 or florigen, that moves to the SAM to induce the transition to flowering (An et al., 2004; 84 Corbesier et al., 2007; Mathieu et al., 2007). The molecular nature of florigen has eluded identification for the better part of a century. However, recently FLOWERING 85 86 LOCUS T (FT) and related genes, which encode phosphatidylethanolamine-binding 87 proteins (PEBP), have been identified as evolutionarily conserved candidates 88 (Corbesier et al., 2007; Mathieu et al., 2007). Under inductive photoperiod, FT is 89 expressed in leaf phloem companion cells (PCC) and there is good evidence that the FT 90 protein is loaded into the phloem sieve elements and transported to the SAM (reviewed 91 in Srikanth and Schmid, 2011; Song et al., 2015). At the SAM, FT interacts with FD 92 and 14-3-3 proteins and the resulting florigen-activation complex (FAC) is thought to 93 control the correct expression of flowering time and floral homeotic genes to promote 94 the transition of the vegetative meristem into a reproductive inflorescence meristem 95 (Abe et al., 2005; Wigge et al., 2005; Taoka et al., 2011). 96 FD belongs to the bZIP transcription factor (TF) family (Jakoby et al., 2002) and is 97 mainly expressed at the SAM (Abe et al., 2005; Schmid et al., 2005; Wigge et al., 2005). 98 It has been proposed that, in order to interact with FT and 14-3-3 proteins, FD must be 99 phosphorylated at threonine 282 (T282) (Abe et al., 2005; Wigge et al., 2005; Taoka et 100 al., 2011). Recently, two calcium-dependent kinases expressed at the SAM, CALCIUM 101 DEPENDENT PROTEIN KINASE 6 (CPK6) and CPK33, have been shown to 102 phosphorylate FD (Kawamoto et al., 2015). FD interacts not only with FT but also with 103 other members of the PEBP protein family. Interestingly, some of the six PEBP proteins 104 encoded in the A. thaliana genome regulate flowering in opposition (Kim et al., 2013). 105 FT and its paralog TWIN SISTER OF FT (TSF) promote flowering. Mutations in tsf 106 enhance the late flowering phenotype of ft in LD, but in addition TSF also has distinct 107 roles in SD (Yamaguchi et al., 2005). Other members of the PEBP protein family, most 108 prominently TERMINAL FLOWER 1 (TFL1), oppose the flower-promoting function 109 and TSF, and repress flowering. The Arabidopsis ortholog of 110 CENTRORADIALIS (ATC) has been shown to act as a SD-induced floral inhibitor 111 that is expressed mostly in the vasculature, but was undetectable at the SAM (Huang et 112 al., 2012). Furthermore, ATC has been suggested to move long distances and can 113 interact with FD to inhibit APETALA1 (API) expression. ATC has thus been proposed 114 to antagonize the flower-promoting effect of FT (Huang et al., 2012). Similarly, 115 orthologs of ATC in rice (RCNs) have been recently showed to antagonize the function 116 of FT-like protein (Kaneko-Suzuki et al., 2018). Finally, BROTHER OF FT (BFT), 117 which like ATC is strongly expressed in the leaf vasculature, can interact with FD in

119 expression, thereby delaying flowering (Yoo et al., 2010; Ryu et al., 2014). 120 TFL1 differs from FT only in 39 non-conserved amino acids but as mentioned above 121 has an opposite biological function: TFL1 represses flowering while FT is a floral 122 promoter (Ahn et al., 2006). It has been demonstrated that substitutions of a single 123 amino acid (TFL1-H88; FT-Y85) or exchange of the segment B encoded by the fourth 124 exon are sufficient to impose TFL1-like activity onto FT, and vice versa (Hanzawa et 125 al., 2005; Ahn et al., 2006; Ho and Weigel, 2014). Similar to FT, TFL1 also interacts 126 with FD, both in yeast-2-hybrid assays as well as in plant nuclei (Wigge et al., 2005; 127 Hanano and Goto, 2011). Together, these findings suggest that activating FD-FT and 128 repressive FD-TFL1 complexes compete for binding to the same target genes (Ahn et 129 al., 2006). This hypothesis is further supported by the observation that TFL1 apparently 130 acts to repress transcription (Hanano and Goto, 2011) whereas FT seems to function as 131 a transcriptional (co-)activator (Wigge et al., 2005). However, evidence that these 132 protein complexes in fact share interactors such as 14-3-3 proteins or control the same 133 targets remains sparse. 134 FD has been reported as direct and indirect regulator of important flowering time and 135 floral homeotic genes such as SUPPRESSOR OF OVEREXPRESSION OF CONSTANS 136 1 (SOC1), SQUAMOSA PROMOTER BINDING PROTEIN-LIKE 3 (SPL3), SPL4, 137 SPL5, LEAFY (LFY), AP1, and FRUITFULL (FUL). Several flowering time pathways contribute to SOC1 regulation. Indeed, it has been proposed that expression of SOC1 138 139 can be directly promoted by the FD-FT complex (Lee and Lee, 2010). However, SOC1 expression can also be activated independently from FD-FT probably through the 140 141 SPL3, SPL4, and SPL5 proteins (Moon et al., 2003; Wang et al., 2009; Lee and Lee, 142 2010), which have been shown to be directly or indirectly activated by the FD-FT 143 complex (Jung et al., 2012). The activation of floral homeotic genes such as API and 144 FUL in response to FD-FT activity at the SAM can at least in part be explained by the 145 direct activation of the floral meristem identify gene LFY through SOC1 (Moon et al., 146 2005; Yoo et al., 2005; Jung et al., 2012). In addition, it has also been proposed that the 147 FD-FT complex can promote the expression of AP1 and FUL by directly binding to 148 their promoters (Abe et al., 2005; Teper-Bamnolker and Samach, 2005; Wigge et al., 2005). Taken together, these results support a central role for FD in integrating different 149 150 pathways to ensure the correct timing of flowering. However, FD targets have not yet 151 been identified at the genome scale, nor has the requirement for protein complex 152 formation for FD function in A. thaliana been addressed systematically.

the nucleus, interfering with FT function under high salinity by inhibiting AP1

Here we identify direct and indirect targets of FD at the genome scale using ChIP-seq and RNA-seq in wildtype as well as in ft-10 tsf-1 double mutants. Our results demonstrate that FD can bind to DNA in vivo even in the absence of FT/TSF. However, FD binding to a subset of targets, which includes many important flowering time and floral homeotic genes, was reduced in the ft-10 tsf-1 double mutant, strongly supporting a role for FT/TSF in modulating the binding of FD to DNA and the expression of functionally important target genes. In addition, we report the effects of FD phosphorylation on protein complex formation with FT and TFL1 via 14-3-3 proteins in vitro and show how phosphorylation of FD affects flowering time in planta. Finally, our ChIP-seq experiments identified hundreds of previously unknown FD target genes, both in the PCCs as well as at the SAM. For example, we observed that FD directly binds to and regulates the expression of genes in hormone signaling pathways. These newly identified FD target genes represent a precious resource not only to enhance our knowledge of the photoperiod pathway but also to better understand the integration of different signaling pathways at the transcriptional level. Taken together, our findings support a role for FD as a central integrator of flowering time and provide important novel data to guide future research on the integration of diverse signaling pathways at the SAM.

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## RESULTS

# FD binds G-box motifs when expressed in PCCs

FD is normally expressed at the shoot apical meristem (SAM) whereas its interaction partner FT is expressed in leaf phloem companion cells (PCC). As most 14-3-3 proteins are ubiquitously expressed at moderate to high levels and have also been detected in PCCs (Schmid et al., 2005; Deeken R. et al., 2008), we reasoned that expression of FD from the PCC-specific *SUC2* promoter would maximize FAC complex formation and enable us to investigate the role of FT in modulating FD transcriptional activity.

We performed ChIP-seq on independent biological duplicates in a stable *pSUC2::GFP:FD* reporter line, which shows no discernible phenotype when comparted to Col-0 background. A *pSUC2::GFP:NLS* line, in which the GFP protein is fused to the nuclear localization signal (NLS) was used as a control. A total of 2068 and 3236 genomic regions showing significant enrichment (peaks) were identified in the first and second replicate, respectively (Fig. S1A).

188 In the individual replicates, the majority of the peaks mapped to promoter regions 189 (65,1% and 63.8%, respectively), followed by intergenic regions (16% and 16.8%), 190 transcriptional terminator sites (9.2% and 10.7%), exons (6.4% and 5.6%) introns 191 (2.4% and 2.3%), 5'-UTRs (0.5% and 0.3%), and 3'-UTRs (0.4% and 0.5%) (S1D). 192 The relative enrichment of peaks mapping to promoter regions is in agreement with 193 what is expected from a transcriptional regulator. In both replicates, the majority of the 194 peaks are located between 600 bp and 300 bp upstream the nearest transcription start 195 site (TSS) (Fig. S1G, J). Overlapping results from the two biological replicates 196 identified 1754 high-confidence peaks shared in both experiments (Fig. S1A, 197 Supplemental Data Set 1). Similar to the individual experiments, these high-confidence 198 peaks mostly mapped to the promoter regions (66.8%) (Fig. 1A). Only this subset of 199 peaks, which includes important flowering time and flower development genes such as 200 AP1, FUL, LFY, SOC1, SEP1, SEP2, SEP3, was used for further analysis. 201 De novo motif analysis of the 1754 high-confidence peaks using MEME-ChIP (Machanick and Bailey, 2011) revealed that peak regions showed a strong enrichment 202 203 of G-boxes (CACGTG), which is a canonical bZIP binding site (Fig. S1M). The subset 204 of 1754 peak regions was associated with 1676 unique genes, with 68 genes containing 205 more than one FD binding site. Taken together, these results demonstrate that, when 206 misexpressed in the PCCs, FD is capable of binding to G-box elements in a large 207 number of genes that are involved in diverse aspects of the plant life cycle.

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## FT and TSF enhance binding of FD to DNA

209 210 To test whether FT and its paralog TSF are required for FD to bind to DNA, the 211 pSUC2::GFP:FD reporter and pSUC2::GFP:NLS control constructs were transformed 212 into the ft-10 tsf-1 mutant background. Results from two independent biological 213 replicates show that FD is capable of binding to DNA even in the absence of FT and 214 TSF. Most peaks (63% and 62.1% in the first and second biological replicate, 215 respectively) mapped to promoter regions within 600 bp and 300 bp nucleotides upstream the nearest TSS (Fig. S1E, H, K). Overall, these results are very similar to 216 217 those observed for pSUC2::GFP:FD in Col-0 (Fig. 1A, Fig. S1E, H, K, N). Comparison 218 between the two biological replicates identified 2696 common peaks in ft-10 tsf-1 219 mutant that mapped to 2504 unique genes (Fig. S1B, Supplemental Data Set 2). 220 Surprisingly, overlapping the sets of genomic regions bound by FD with high-221 confidence in WT (1754) and ft-10 tsf-1 (2696) backgrounds identified 1530 shared 222 peaks (Fig. 1B, Supplemental Data Set 3), suggesting that FD is capable of binding to

223 the majority of its targets in the absence of FT and TSF. Motif enrichment analysis of 224 sequence comprising the 1530 shared peaks revealed that FD maintained a strong 225 preference for binding to G-box motifs (Fig. 1C). 226 Analysis of differentially bound (DB) regions revealed that, although FT and TSF were 227 not required for FD to bind DNA, their presence increased the enrichment of FD on a subset of target loci and this difference in binding was sufficient to discriminate Col-0 228 229 and ft-10 tsf-1 (Fig. 1D). A total of 885 DB regions with a FDR < 0.05 were observed 230 between WT and ft-10 tsf-1 and almost all of these loci showed higher enrichment in 231 WT (Fig. 1E, Supplemental Data Set 4). Interestingly, this subset includes important 232 floral homeotic genes such as API, SEPI, SEP2, and FUL, as well as two members of 233 the SPL gene family, SPL7 and SPL8. We also found FD bound to the second exon of 234 LFY, a master regulator of flower development (Fig. 1F). In addition, we detected 235 binding to loci encoding genes involved in the regulation of gibberellic acid 236 biosynthesis and degradation such as GA2OX4, GA2OX6, and GA3OX1 as well as to 237 three key components of the circadian clock, CCA1, LHY, and TIC (Supplemental Data 238 Set 4). 239 To test the robustness of our results and for any possible bias due to the use of the 240 different genetic backgrounds Col-0 and ft-10 tsf-1 as controls, peaks were called again 241 using pSUC2::GFP:NLS in Col-0 as single negative control. Analysis identified 917 242 DB loci (Fig. S2), which is comparable to the 885 DB loci from the previous analysis 243 (Fig. 1E). In addition, affinity test analysis clustered by genotype rather than the control 244 used (Fig. S2), ruling out strong bias due to the usage of different genetic backgrounds 245 for peak calling. 246 Importantly, FD is capable of inducing the known FAC target gene AP1 in leaves when 247 expressed under the pSUC2 promoter, suggesting that a functional FAC can be formed 248 in the phloem companion cells when FD is present (Fig. S3A). The finding that API 249 expression could only be observed in the Col-0 background but not in pSUC2::GFP:FD 250 ft-10 tsf-1 further supports this idea. However, in contrast to AP1, we failed to detect 251 induction of SOC1 in the PCCs of pSUC2::GFP:FD (Fig.S3A), suggesting that other 252 co-factor(s) are specifically expressed at the SAM might be required to fully activate 253 FD target gene expression.

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## FD phosphorylation is required for complex formation and to promote flowering

To confirm the binding of FD to G-boxes we performed electrophoretic mobility shift

assays (EMSAs) using the bZIP domain of the A. thaliana FD protein (FD-C) and a

258 probe consisting of a 30bp fragment from the SEP3 promoter containing a G-box that 259 we had identified as FD target region in our ChIP-seq (Fig. 1F). We observed weak 260 binding of FD-C, but failed to detect higher order complexes when 14-3-3, FT, or both 261 were added (Fig. 2A). In contrast, a clear supershift with 14-3-3 and FT was observed 262 when a phosphomimic variant of FD-C, FD-C T282E, was used (Fig. 2B). 263 Interestingly, TFL1, which is similar to FT in structure (Ahn et al., 2006) but delays 264 flowering, was capable of forming a complex with 14-3-3 and wildtype FD-C (Fig. 2A). 265 Similar results were obtained with the full-length version of FD (Fig. S4A). Taken 266 together, these results demonstrate that A. thaliana FD is capable of binding to DNA without FT, confirming results from our ChIP-seq experiments. Furthermore, this 267 268 indicates that the unphosphorylated form of FD, in complex with 14-3-3 proteins, can 269 interact with TFL1. 270 To investigate the importance of FD phosphorylation in vivo we complemented the fd-271 2 mutant with pFD::FD, pFD::FD-T282E, and pFD::FD-T282A (which cannot be 272 phosphorylated) and determined flowering time of homozygous transgenic plants. 273 Plants transformed with the WT version of FD rescued the late flowering phenotype of 274 fd-2, indicating that the rescue construct was fully functional (Fig. 3). In contrast, plants 275 transformed with the T282A version flowered with the same number of leaves as fd-2, 276 demonstrating that FD needs to be phosphorylated to induce flowering. Interestingly, 277 plants transformed with the T282E phosphomimic version of FD flowered even earlier 278 than WT (Fig. 3), indicating that control of FD phosphorylation is important for its 279 function in vivo. To test whether serine 281 (S281), which is located next to T282, 280 constitutes a potential FD phosphorylation site, we complemented fd-2 with pFD::FD-281 S281E and pFD::FD-S281E/T282E constructs. Interestingly, these lines flowered as 282 early as plants transformed with the phosphomimic version T282E (Fig. 3), indicating 283 that S281 may be a FD phosphorylation site but that mimicking double-phosphorylation 284 of S281/T282 does not accelerate flowering any further. These in vivo results are in 285 agreement with our in vitro EMSA results and confirm that phosphorylation of FD is 286 required for its function and must be finely regulated in order to avoid either premature 287 or delayed flowering. It should be noted, however, that the phosphomimic version of 288 the C-terminal fragment of FD (as used in the EMSA analyses) is insufficient to fully 289 rescue the late flowering of fd-2 (Fig. S3B), suggesting that the N-terminal region of 290 FD, even though it does not contain any known functional domains, nevertheless 291 contributes to FD function.

## Targets of FD at the SAM

293 294 The rationale for carrying out the initial ChIP-seq experiments in PCCs was to 295 maximize the likelihood of FAC formation and to study the contribution of FT/TSF to 296 FD DNA binding. However, since our ChIP-seq and EMSA results indicated that FD-297 FT interaction is not required for FD to bind to DNA, we decided to determine direct 298 targets of FD in its natural context at the SAM. 299 To this end we performed ChIP-seq using a fd-2 mutant complemented with a 300 pFD::GFP:FD construct (Fig. S3C). We performed ChIP-seq using two independent 301 biological replicates from apices of 16-day-old plants grown in LD conditions. In the 302 two replicates, we could identify 703 and 1222 FD-bound regions, respectively, of 303 which 595 were shared between the replicates (Fig. S1C, Supplemental Data Set 5). Of 304 these, 69.7% mapped to core promoter regions within 300 to 600 bp upstream of the 305 nearest TSS, 15.8% to intergenic regions, followed by lesser percentages to TTS 306 (6.2%), exons (5.9%), introns (1.8%) and 5'-UTRs (0.5%) (Fig. 1G, Fig. S1F, I, L). 307 Similar to the situation in our PCC-specific ChIP-seq analyses we found G-box 308 sequences as the most overrepresented transcription factor binding sites under the peak 309 regions (Fig. 1H, Fig. S1O). The 595 peak regions shared between the replicates 310 mapped to 572 individual genes, which we consider high-confidence in vivo targets of FD at the SAM and include important flowering-related genes such as API, FUL, 311 312 SOC1, and SEP3. 313 The precise location of the FD binding site in the API promoter has been discussed 314 controversially (Wigge et al., 2005; Benlloch et al., 2011). Taking into account all six 315 ChIP-seq datasets, we were able to extract a 64 bp sequence covering the peak summits 316 on the AP1 promoter (Fig. 4A, B). Interestingly, this sequence lies about 100 bp 317 downstream of the C-box that had previously been implicated in FD binding to API 318 (Wigge et al., 2005), but contains several other palindromic sequences. However, none 319 of these sequences is a bona fide G-box. We selected three potential binding sites within 320 the 64 bp sequence and tested them, along with the upstream C-box, by EMSA for FD 321 binding (Fig. 4C, S4B). Results show that only the phosphomimic version of FD-C 322 (FD-C T282E) in combination with 14-3-3 can bind to all four DNA sequences tested. 323 Furthermore, we detected a supershift for all palindromic sites tested, included the C-324 box, when we added TFL1. In contrast, for FT an additional shift resembling the pattern 325 obtained with the G-box in SEP3 promoter was only observed for "site 2" (Fig. 4C, 326 2B). Closer inspection of the nucleotide sequences of the probes used for the G-box in the SEP3 promoter and the "site 2" in the AP1 promoter revealed that the possible FD 327

- binding site in the AP1 promoter (GTCGAC) is also present in the SEP3 promoter,
- where it overlaps with the G-box (Fig. 4D). Interestingly, in the context of the SEP3
- probe, full-length FD and FD-C tolerated mutating the core of the G-box from CG to
- 331 GC, whereas CG to TA mutations as well as converting the G-box to a C-box
- 332 (GACGTC) abolished binding (Fig. S4D). To further test if "site 2" on the API
- promoter constitutes a real FD binding site, we mutated its core from CG to TA and
- 334 checked whether this was sufficient to abolish binding. Results show that indeed the
- binding of FD to this mutated version of "site 2" was strongly reduced, except in the
- presence of TFL1 (Fig. S4E).
- Taken together, our findings exclude the C-box as the FD binding site in the AP1
- promoter. Furthermore, our results suggest that in vitro FD can bind to other motifs
- besides the G-box, possibly through interaction with partners other than 14-3-3 and
- 340 FT/TSF, and we characterized possible a new binding site (GTCGAC) that could
- constitute the true FD binding site in the *AP1* promoter.

## Differentially expressed genes at the SAM and direct targets of FD

- To test which of the 595 high confidence binding sites we had identified by ChIP-seq
- at the SAM were in fact transcriptionally regulated by FD we performed RNA-seq on
- apices from fd-2 mutants and the pFD::GFP:FD fd-2 rescue line. 21 day-old SD-grown
- 347 seedlings were shifted to LD to induce synchronous flowering and apices were
- harvested on the day of the transfer to LD (T0), as well as 1, 2, 3, and 5 days after the
- shift (T1, T2, T3, T5) from three independent biological replicates.
- Differentially expressed (DE) genes were determined for each time point and genes
- with an adjusted p-value (padj) lower than 0.1 were selected as significantly DE. We
- 352 identified in total 1759, 583, 2421, 924, and 153 DE genes in T0, T1, T2, T3, and T5,
- respectively, corresponding to 4189 unique genes (Fig. 5A, Supplemental Data Set 6).
- 354 PCA analysis showed that the first and second principal components, which explain
- 355 37% and 21% of the total variance, corresponded to the different time points and
- genotypes, respectively (Fig. S5A). The best separation between the genotypes in the
- 357 PCA was observed at T3 and T5, indicating that FD contributes to the transcriptional
- 358 changes at the SAM mainly after exposure to two long days. This observation is in
- agreement with the expression profile of FD, which in the *pFD::GFP:FD* rescue line
- 3(0 · 1 0 T3 (F; 05P) I · 1 FP · 1 · 11 · 1 · 1 · 0/3
- increased after T2 (Fig. S5B). In contrast, FD expression remained low in the fd-2
- mutant, indicating the validity of our experimental approach (Fig. S5B).

362 Next, we intersected the list of genes that were bound by FD at the SAM (572) with the 363 list of DE genes (4189). In total, 135 (23.6%) of the 572 FD-bound genes were significantly DE at the SAM during the transition to flowering at least at one timepoint, 364 365 indicating that these genes are transcriptionally regulated by FD, which is more than 366 expected by chance (Fig. 5B, C, Supplemental Data Set 7). Among these 135 directly 367 bound and differentially expressed genes we observed several previously known FD-368 regulated flowering time and floral homeotic genes including API, FUL, and SOCI 369 (Fig. 1F, S6A). In addition, this set of 135 high-confidence FD targets contained also 370 the MADS box gene SEP3, the promoter of which is bound by FD and which is downregulated in fd-2 mutant (Fig. 1F, S6A). Interestingly, we did not observe binding of 371 372 FD to any of the other members of SEPALLATA gene family in ChIP-seq samples from 373 the SAM, although we did detect FD binding in promoter regions of SEP1 and SEP2, 374 but not SEP4,in ChIP-seq from seedlings in which FD had been misexpressed from the 375 SUC2 promoter. One possible explanation for this is that the ChIP-seq at the SAM 376 apparently worked less efficiently and identified fewer FD targets than in leaves (595 377 vs. 1754), which might result in a larger number of false negatives. In agreement with 378 this interpretation, SEP1 is down-regulated in the fd-2 mutant background (Fig. S6), 379 indicating that FD directly or indirectly regulates the expression of SEP1 at the SAM. 380 Interestingly, we also found FD bound to TPR2, a member of the TOPLESS (TPL)-381 related gene family. TPL and its family members (TPR1, TPR2, TPR3 and TPR4) are 382 strong transcriptional co-repressors that interact with other proteins throughout the 383 plant to modulate gene expression (Causier et al., 2012). TPR2 is down-regulated in the fd-2 mutant throughout the floral transition from T0 to T5 (Fig. S6), indicating FD 384 385 might regulate development at the SAM through TPR2 in a photoperiod-independent 386 manner. 387 Gene Ontology (GO) analysis of these 135 genes that were bound and differentially 388 expressed by FD revealed significant enrichment in several biological process 389 categories (Fig. S7), including "flower development" and "maintenance of 390 inflorescence meristem identity", as expected for a flowering time regulator such as 391 FD. More surprisingly, however, genes related to the "response to hormone" category 392 were also significantly overrepresented (Supplemental Data Set 8). Among the 27 genes 393 in this category are four genes that are best known for their role in jasmonate signaling 394 (MYC2, JAZ3, JAZ6 and JAZ9), three genes directly connected to auxin signaling 395 (ARF18, WES1, and DFL1), four genes involved in abscisic acid signaling (ALDH3311, 396 ATGRDP1, HAII and PP2CA), and the flowering-related gene SOCI, which is wellknown to be regulated by gibberellins (Supplemental Data Set 8). Closer inspection of the expression profiles of these 27 candidate genes revealed that *ARF18* showed a trend similar to *SOC1*, being strongly induced after T2 in Col-0 but not in *fd-2*. The four jasmonate-related genes showed a peculiar expression profile in *fd-2*: an increase from T0 to T1, a decrease in T2, another increase in T3, and decreasing again in T5. Since this peculiar expression profile was observed in three *JAZ* genes, we checked the remaining genes in this family and found that 11 out of 13 displayed a similar pattern (Fig. S6). Furthermore, this profile was also observed in three other genes, *DMR6*, *ESP* and *TOE2*, that have previously been implicated in pathogen resistance and the jasmonate pathway (Fig. S7B). Taken together, these results suggest that FD plays an active role not only in the regulation of flowering time but also functions to connect different hormone signaling pathways.

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## Validation of FD targets

We selected a subset of putative FD direct target genes and determined their expression in early flowering FD overexpression lines (p35S::FD) and Col-0. To minimize any bias due to the early flowering of p35S::FD, experiments were carried out in vegetative 7-day-old LD-grown seedlings. For validation, we selected genes known to play a major role in floral transition, genes that according to Gene Ontology are involved in flowering time and floral development, and other genes that showed a marked differential expression in fd-2 but for which a role in flowering time regulation had not previously been studied in detail. qRT-PCR assays confirmed that both SOC1 and AP1 were strongly up-regulated in p35S::FD (Fig. 6). Although we had only found SEP3 to be bound by FD in the SAM ChIP-seq analysis, we tested expression of all four SEPALLATA genes (SEP1 – SEP4) in the p35S::FD line. SEP3 was the only SEP gene that was strongly induced in seedlings in response to FD overexpression, while SEP1 showed only moderate induction. In contrast, expression of SEP2 and SEP4 did not show differences between p35S::FD and Col-0 (Fig. 6). Interestingly, SEP1, SEP2, and SEP3 were also bound by FD in PCC-specific ChIP-seq in seedlings and SEP1 and SEP3 exhibited strong differential expression in RNA-seq (Fig. S6). AS1, which has been demonstrated to be involved in flowering time by regulation of FT expression in leaves (Song et al., 2012), did not show significant difference in expression between Col-0 and p35S::FD. We also tested two FRIGIDA-like genes, FRI-like 4a and FRIlike 4b, of which FRI-like 4b showed a decreased expression in p35S::FD. In addition, we also tested two genes, MYC2 and AFR1, which were bound by FD in both the pSUC2

and *pFD* ChIP-seq experiments, differentially expressed at the SAM, but not differentially bound in *ft-10 tsf-1* mutant, *i.e.* not directly influenced by the presence of FT and TSF, for their contribution to flowering time regulation. *MYC2* showed no differences in expression in *p35S::FD* compared to Col-0, whereas *AFR1* was upregulated in *p35S::FD* (Fig. 6). To genetically test the role of these two genes in the regulation of flowering we isolated T-DNA insertion lines and determined their flowering time under LD at 23°C. Both *myc2* and *afr1* were significantly early flowering, both as days to flowering and total leaf number, compared to WT (Fig. 7), confirming their role in regulating the floral transition.

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#### **DISCUSSION**

FD was originally identified as a component of the photoperiod-dependent flowering pathway in A. thaliana based on the late flowering phenotype of the loss-off-function mutant (Koornneef et al., 1991). FD, which encodes a bZIP transcription factor, is expressed at the SAM prior to the floral transition but does not seem to induce flowering on its own. Later, it was demonstrated that FD physically interacts with FT, the florigen, and that this interaction is important for its function as a promoter of flowering (Abe et al., 2005; Wigge et al., 2005). In addition, FD was found to also interact with TFL1, which is normally expressed in the SAM and antagonizes the function of FT as floral activator. This and other findings led to the hypothesis that FD is held in an inactive state in the vegetative SAM through interaction with TFL1. When FT is induced in the PCCs and transported to the SAM in response to inductive photoperiod, FT competes with TFL1 for interaction with FD, eventually resulting in the formation of transcriptionally active FD-FT complexes (Ahn et al., 2006). However, the exact molecular mechanisms of FD action and its genome-wide targets remained largely unknown. Here we employed biochemical, genomic, and transcriptomic approaches to clarify the role of FD in the regulation of flowering transition in A. thaliana. We found that neither FT nor TSF are required for FD to bind to DNA but that their presence increases the enrichment of FD on a subset of target loci, which encode for known flowering time and floral homeotic genes such as API, SEP1, SEP2, and FUL. Our data are compatible with the model described by (Ahn et al., 2006), according to which FT acts as a transcriptional coactivator. Without FT, FD would still be capable of binding to DNA but would not activate transcription. This hypothesis is supported

467 by our finding that FD is capable of binding to DNA by itself in vitro. However, we 468 can not exclude the possibility that the binding of FD to DNA we observed in the 469 pSUC2::GFP:FD ft-10 tsf-1 reporter line could be mediated by the floral repressors 470 BFT and ATC, both of which are expressed in the leaf vasculature and are known to 471 interact with FD (Ryu et al., 2014; Huang et al., 2012; Yoo et al., 2010). 472 In this context, our EMSA results are of particular interest as they demonstrate that, at 473 least in vitro, TFL1 is capable of interacting with unphosphorylated FD via 14-3-3 474 proteins, suggesting that the transcriptionally inactive ternary FD/14-3-3/TFL1 475 complex could be the ground state at the SAM. Only after FD has been phosphorylated 476 can FT, together with 14-3-3 proteins, form an active FAC to induce flowering. This 477 requirement for phosphorylation of T282 of FD adds another safeguard to the system 478 that might help to prevent disastrous premature induction of flowering. Our results 479 clearly suggest that phosphorylation is important for FD function and add to our 480 understanding concerning the role of FD phosphorylation, which had mostly been based on the analyses of a FD/14-3-3/Hd3a complex in rice using a short FD peptide (Taoka 481 482 et al., 2011; Kaneko-Suzuki et al., 2018). 483 Which kinases regulate phosphorylation of FD in vivo has been a matter of debate, but 484 recently two calcium-dependent kinases, CPK6 and CPK33, have been shown to 485 phosphorylate FD (Kawamoto et al., 2015). Building on this, we show that expression 486 of a non-phosphorable version of the FD protein (T282A) under the control of the pFD 487 promoter failed to rescue the late flowering of fd-2. In contrast, expression of a 488 phosphomimic version of FD (T282E) resulted in early flowering when expressed in 489 fd-2. Similar results were obtained using a S281E phosphomimic FD. These results 490 indicate that the phosphorylation of FD must be tightly controlled to prevent premature 491 flowering. Interestingly, both CPK6 and CPK33 are more strongly expressed in 492 transition apices than they are in vegetative apices (Schmid et al., 2005), which would 493 be in agreement with an activation of FD by these two kinases during floral induction. 494 Somewhat surprisingly, we observed that the C-terminal part of the FD protein, which 495 includes the bZIP domain and the phosphorylation site, was sufficient to trigger 496 complex formation with FT, TFL1 and 14-3-3 proteins. This suggests that the N-497 terminal region of FD, which is predicted to be highly unstructured and contains a 498 stretch of 25 amino acids containing 19 serine residues, might be dispensable for 499 FD/14-3-3/FT complex formation. However, the N-terminal region of FD is 500 evolutionarily conserved, indicating that it may contribute to FD function. This notion 501 is supported by our observation that expression of the C-terminal part of FD in plants 502 only partially restored the late flowering of fd-2 mutants. 503 Part of the flowering promoting activity of FD can probably be expressed through its 504 effect on members of the SEP gene family of MADS-domain transcription factors, 505 which are required for the activity of the A-, B-, C-, and D-class floral homeotic genes 506 (reviewed in Theissen et al., 2016). In addition to its function as a floral homeotic gene, 507 SEP3 has also been reported to promote flowering by accumulation in leaves under FT 508 regulation (Teper-Bamnolker and Samach, 2005) and as downstream target of the 509 miR156-SPL3-FT module in response to ambient temperature (Hwan Lee et al., 2012). 510 However, how SEP3 is regulated at the SAM has remained unclear. Interestingly, we 511 found that FD bound strongly to the SEP3 promoter and SEP3 is downregulated in the 512 fd-2 mutant. As FD also binds to the promoter and activated expression of the A-class 513 gene API, FD activity might be sufficient to induce formation of sepals, which form 514 the outmost floral whorl, and which according to the quartet model require the 515 formation of a SEP/AP1 complex (Theissen et al., 2016). However, it should be noted 516 that fd mutants do not display notable homeotic defects, indicating that FD is clearly 517 not the only factor regulating SEP3 and AP1 expression. Furthermore, binding of FD 518 to AP1 is unlikely to be mediated by a C-box as previously suggested (Wigge et al., 519 2005; Taoka et al., 2011) as the summits of the ChIP-seq peaks do not cover this region 520 of the API promoter. Interestingly, this region contains several palindromic sequences, 521 one or more of which most likely mediate FD binding to the API promoter. 522 Another interesting outcome of our analyses is the indication that FD might contribute 523 to the regulation of other processes in the plant besides flowering. In particular, we 524 found that FD directly regulated the expression of genes involved in several hormone 525 signaling pathways. For example, we observed FD binding to the promoter of MYC2, a 526 bHLH transcription factor that plays a key role in jasmonate response. It has been 527 shown that MYC2 forms a complex with JAZ proteins and the TPL co-repressor, and 528 that this interaction is dependent on NINJA proteins (Pauwels et al., 2010). In this 529 context it is noteworthy that FD also bound directly to the promoter of TPR2 promoter 530 and that TPR2 was strongly downregulated in fd-2. This finding indicates that FD not 531 only regulates MYC2 but also at least some of the interacting TPL-like transcriptional 532 co-repressors. Finally, we also observed strong binding of FD to (and misexpression 533 of) a number of JAZ genes in either PCCs and/or the SAM in our ChIP-seq and RNA-534 seq data. Taken together, this indicates that FD may control the expression of three core components of jasmonate signaling: MYC2, TPR2, and several JAZ genes. These results 535

536 support earlier findings that had reported a link between jasmonate signaling 537 components and flowering time regulation. JAZ proteins have been shown to regulate 538 flowering in leaves through the direct interaction with the floral repressors TOE1 and 539 TOE2, which is also bound by FD and differentially expressed in fd-2, and the 540 regulation of FLC that negatively regulate FT expression (Zhai et al., 2015). Moreover, 541 MYC2 has also been reported to affect flowering time by regulating FT expression in 542 leaves (Zhai et al., 2015; Wang et al., 2017). However, previous publications had 543 reported contradictory results concerning the flowering phenotype of the myc2 mutant, 544 ranging from late flowering (Gangappa and Chattopadhyay, 2010) to early flowering 545 (Wang et al., 2009) or no obvious effect (Major et al., 2017). In our conditions the myc2 546 mutant exhibited early flowering compared to Col-0, in agreement with the report from 547 Wang and colleagues (Wang et al., 2009) (Fig. 7). We also identified ARF18, a member 548 of the auxin response factor family, as a direct target of FD. Notably, the expression of 549 ARF18 is strongly induced after T2 in Col-0 but not in fd-2 and this pattern is the same 550 of known direct FD targets, e.g.: AP1 and SOC1. Moreover, ARF18 is also induced at 551 the SAM during the floral transition (Schmid et al., 2005) providing further evidence 552 for a possible link between FD and ARF18. In summary, our findings suggest a link 553 between the photoperiodic pathway gene FD and hormone signaling pathways. 554 Although further experiments will be necessary to better understand this connection, 555 we hypothesize that linking hormone signaling to flowering time through FD regulation 556 might allow plants to fine tune their flowering time response to abiotic and biotic 557 stresses. 558 Apart from connecting FD with hormone signaling we characterized another target gene 559 in more detail, AFR1. This gene encodes a putative histone deacetylase subunit and had 560 previously been shown to negatively affect the expression of FT in the leaves. Further, 561 afr1 mutations cause early flowering (Fig. 7)(Gu et al., 2013). Our results suggest that 562 FD might modulate flowering through ARF1-mediated regulation of chromatin. 563 However, such regulation would most likely not be mediated by FT, as FT is normally 564 not expressed at the SAM. 565 Taken together, our results support the role of FD as a key regulator of photoperiod-566 induced flowering and the expression of A- and E-class floral homeotic genes in A. 567 thaliana. Furthermore, FD might play an important role in coordinating the crosstalk 568 between the photoperiod pathway and hormone signaling pathways and provide a 569 convergence point for diverse environmental and endogenous signaling pathways.

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## MATERIAL AND METHODS

# 574 Plant materials and growth conditions

- 575 Arabidopsis thaliana accession Col-0 was used as wild-type. Mutants investigated in
- 576 this study are: fd-2 (SALK 013288), ft-10 (GABI 290E08), tsf-1 (SALK 087522),
- 577 myc2 (SALK 017005), arf1 (SALK 026979) (Tab. S1). Seeds were stratified for 3
- days in 0.1% agar in the dark at 4°C and directly planted on soil. Plants were grown on
- soil under long days (16 hours of light and 8 hours of night) or under short days (8 hours
- of light and 16 hours of night) at 23°C and 65% relative humidity. Plants used for
- flowering time measurements were grown in a randomized design to reduce location
- 582 effects in the growth chambers.

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## DNA vectors and plant transformation

- 585 DNA vectors used in this study are listed in table S2. Coding sequences were amplified
- by PCR from cDNA and cloned into either pGREEN-IIS vectors for flowering time
- 587 studies or pET-M11 vectors for protein expression. Final constructs were transformed
- by electroporation in Agrobacterium tumefaciens and Arabidopsis plants of accession
- Col-0 and fd-2 were transformed by the floral dip method. Basta treatment (0.1% v/v)
- was used for screening for transgenic lines.

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## ChIP and ChIP-seq

- 593 Approximately 1.5 grams of seedlings (pSUC2::GFP:FD and pSUC2::GFP:NLS in
- both Col-0 and ft-10 tsf-1) or 300 mg of manually dissected apices (pFD::GFP:FD in
- 595 fd-2; Col-0) from 16 day old plants grown on soil under long day 23°C were harvested
- and fixed in 1% formaldehyde under vacuum for 1 hour. ChIP was performed as
- 597 previously described (Kaufmann et al., 2010) with the following minor changes:
- sonication was performed using a Covaris E220 system (conditions: intensity 200 W,
- 599 duty 20, cycles 200, time 120 seconds), incubation time with antibody was increased to
- over-night, incubation time with protein-A agarose beads was increased to 4 hours,
- purification of DNA after de-cross linking was performed with MinElute Reaction
- 602 Cleanup Kit (Qiagen).
- Anti-GFP from AbCam (ab290) was used for immuno-precipitation. ChIP-seq libraries
- were prepared using TruSeq ChIP Library Preparation Kit (Illumina) and BluePippin
- was used for gel size selection of fragments between 200 bp and 500 bp. Final

606 concentration and size distribution of the libraries were tested with Qubit and

607 BioAnalyzer (Agilent High Sensitivity DNA Kit). Libraries were sequenced on an

608 Illumina HiSeq3000 system using the 50bp single end kit.

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## RNA extraction, RNA-seq and expression analysis

- For RNA-seq, Col-0 and fd-2 plants were grown for 21 days under short days at 23°C
- and then shifted to long days 23°C. RNA was extracted from manually dissected apices
- collected the day of the shift (T0) and 1, 2, 3, 5 days after shifting (T1, T2, T3 and T5
- respectively) using the RNeasy Plant Kit (Qiagen) according to the manufacturer's
- 615 instructions. RNA integrity and quantification were determined on a BioAnalyzer
- system. 1 µg of of RNA was used to prepare libraries using the TruSeq RNA Library
- Prep Kit (Illumina). All libraries were quality controlled and quantified by Qubit and
- Bioanalyzer and run on a Illumina HiSeq3000 with 50bp single end kit.
- Validation of the selected FD targets was performed in 7 day old seedlings grown on
- 620 soil under long days at 23°C.
- 621 RNA was extracted using the RNeasy Plant Kit (Qiagen) according to the
- manufacturer's instructions. cDNA was synthetized using the RevertAid RT Reverse
- Transcription Kit (ThermoScientific) according to the manufacturer's instructions.
- 624 qRT-PCRs were performed on a CFX96 Touch Real-time PCR Detection System
- 625 (BioRad) using LightCycler 480 SYBR Green I Master (Roche). Oligonucleotides used
- as primers for qRT-PCR are listed in table S3.

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## ChIP-seq and RNA-seq analysis

- Raw data from ChIP-seq were trimmed of adapters and aligned to the A. thaliana
- genome (TAIR10 release) using bwa (Li and Durbin, 2010). MACS2 was used to call
- peaks using default parameters (Zhang et al., 2008). Mapped reads from samples
- expressing GFP:NLS under the same promoter of the GFP:FD (e.g.: pSUC2) in
- 633 seedlings experiments or Col-0 without any vector in apices experiments were used for
- out using the R package normalization. Differentially bound analyses were carried out using the R package
- "DiffBind" using default parameters (Stark, 2011; Ross-Innes et al., 2012).
- 636 For the analysis of RNA-seq data, sequencing reads mapping to rRNAs were filtered
- out using Sortmerna (Kopylova et al., 2012) and the remaining reads were trimmed of
- adapters using Trimmomatic (Bolger et al., 2014). Alignment to the A. thaliana genome
- was performed with STAR (Dobin et al., 2013) and read counted with HTSeqCount

640	(Anders et al., 2015). Differential expression analysis was performed using DESeq2
641	with default parameters (Love et al., 2014).
642	
643	Electrophoretic Mobility Shift Assays (EMSA)
644	Coding sequences of both the wild-type version as well as the phosphomimic variant
645	(T282) of FD and its C-terminal domain (FD-C, amino acids: 203-285), 14-3-3 v
646	(At3g02520; GRF7), FT, and TFL1 were amplified by PCR to generate N-terminal 6X-
647	His-tag CDS which were cloned into pETM-11 expression vector by restriction enzyme
648	cloning. All plasmids were transformed into Escherichia coli strain Rosetta plysS and
649	proteins were induced with 1mM IPTG at 37°C over-night. Cell lysis was performed
650	by sonication and proteins were purified using His60 columns (Clontech) and eluted in
651	50 mM of sodium phosphate buffer pH 8.0, 300 mM NaCl, 300 mM Imidazole. EMSA
652	was performed using 5'-Cy3-labeled, double-stranded oligos of 30 bp covering the G-
653	box contained in the SEP3 promoter as a probe (Eurofins). For probe synthesis, single
654	strand oligos were annealed in annealing buffer (10 mM Tris pH 8.0, 50 mM NaCl, 1
655	mM EDTA pH 8.0). Binding reactions were carried out in buffer containing 10 mM
656	Tris pH 8.0, 50 mM NaCl, 10 $\mu M$ ZnSO <sub>4</sub> , 50 mM KCl, 2.5% glycerol, 0.05% NP-40 in
657	a total volume of 20 $\mu$ l. The binding reaction was kept in dark at room temperature for
658	20 minutes and then loaded in native 8% polyacrylamide gel and run in 0.5X TBE at
659	4°C in dark. Results were visualized using a Typhoon imaging system.
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663	ACCESSION NUMBERS
664	RNA-seq and ChIP-seq data have been deposited at European Nucleotide Archive
665	(ENA) under accession number PRJEB24873 and PRJEB24874, respectively.
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669	SUPPLEMENTARY MATERIAL
670	Supplemental Figures
671	Figure S1. ChIP-seq summary statistics for the different biological replicates:
672	pSUC2::GFP:FD in Col-0 (A, D, G, J) and ft-10 tsf-1 mutant background (B,
673	E, H, K), pFD::GFP:FD in fd-2 mutant background (C, F, I, L).

674 Figure S2. Verification of comparability of controls used for normalization of FD 675 (pSUC2::GFP:FD) ChIP-seq in WT and ft-10 tsf-1 seedlings. 676 Figure S3. Effect of misexpression of FD on gene expression and flowering time. 677 Figure S4. Electrophoretic mobility shift assays (EMSAs) to test FD binding to the 678 SEP3 and AP1 promoters. 679 Figure S5. Summary of RNA-seq results. 680 Figure S6. Expression profile of selected FD target genes. 681 Figure S7. Gene Ontology (GO) analysis on the subset of 135 direct genes of FD. 682 **Supplemental Tables** 683 684 Table S1. List of mutants and oligos for genotyping used in the study. 685 List of vectors used in the study. Table S2. List of oligos used for qRT-PCR in the study. 686 Table S3. 687 688 **Supplemental Data Sets** 689 List of 1754 FD-bound peaks identified in seedlings Supplemental Data Set 1. 690 expressing *pSUC2::GFP:FD* in Col-0. 691 Supplemental Data Set 2. List of 2427 FD-bound peaks identified in seedlings 692 expressing *pSUC2::GFP:FD* in *ft-10 tsf-1*. 693 Supplemental Data Set 3. List of 1514 FD-bound peaks detected in seedlings 694 expressing pSUC2::GFP:FD in either Col-0 or ft-10 tsf-1. 695 **Supplemental Data Set 4.** List of 917 peaks that were differential bound in 696 seedlings expressing *pSUC2::GFP:FD* in either Col-0 or *ft-10 tsf-1*. 697 List of 595 shared FD-bound peaks in apices **Supplemental Data Set 5.** 698 pFD::GFP:FD fd-2 rescue line. 699 **Supplemental Data Set 6.** List of differentially expressed genes. 700 List of 135 potential direct targets of FD. Supplemental Data Set 7. 701 List of 27 genes related to "response to hormone" Supplemental Data Set 8. 702 category within the subset of the 135 direct target of FD. 703 704 705 706 **ACKNOWLEDGEMENTS** 

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# 723 FIGURE LEGENDS

- 724 Figure 1. Identification of FD targets by pSUC2::GFP:FD ChIP-seq in Col-0 and ft-
- 725 10 tsf-1 and pFD::GFP:FD ChIPseq in fd-2.
- 726 **(A)** Annotation of high-confidence peaks found in two biological replicates in Col-727 0 and *ft-10 tsf-1*.
- 728 **(B)** 4-set venn diagram representing the overlapping peaks among all the biological
- replicates from Col-0 and ft-10 tsf-1. The majority of the peaks (1514) are
- shared between the two genetic backgrounds.
- 731 **(C)** Nucleotide logo of the predicted FD binding site.
- (D) Correlation heatmap calculated on a binding matrix based on ChIP-seq reads
- counts for Col-0 and *ft-10 tsf-1* samples (affinity scores). The presence/absence
- of FT and TSF is sufficient to discriminate the two genetic backgrounds.
- 735 (E) Differentially bound (DB) peaks between Col-0 and ft-10 tsf-1. Red dots
- indicate differentially bound peaks with a FDR < 0.05.
- 737 **(F)** Reads from Col-0, *ft-10 tsf-1* and control sample mapped against selected
- flowering related genes.
- 739 (G) Annotation of high-confidence peaks identified by ChIPseq in two biological
- replicates in *pFD::GFP:FD fd-2*.
- 741 **(H)** Nucleotide logo of the predicted FD binding site at the SAM.

- Figure 2. The C-terminal part of FD (FD-C) binds to a G-box in the *SEP3* promoter in vitro.
- (A) Electrophoretic mobility shift assay (EMSA) of the wild-type form of FD-C in combination with 14-3-3v, FT, and TFL1. FD-C weakly binds the probe on its own but it is not able to form a complex with 14-3-3v and FT. However, FD-C forms a complex with 14-3-3v and TFL1 capable of binding the G-box.
  - **(B)** Phosphomimic version of FD-C (FD-C\_T282E) in combinations with 14-3-3v, FT and TFL1. The phosphomimic version of FD-C binds the G-box alone and interacts with 14-3-3v, which facilitates interaction with FT and TFL1. Both wild-type and phosphomimic version of FD-C require 14-3-3v for interaction with FT or TFL1.
    - Asterisk (\*) indicate shifted probe.

- **Figure 3.** Phosphorylation of FD at threonine 282 (T282) modulates flowering time in *A. thaliana*.
  - Expression of wildtype (WT) pFD::FD rescues the late flowering phenotype of fd-2. Mutation of T282 to alanine (T>A) in  $pFD::FD\_T282A$ , which prevents phosphorylation, abolishes rescue of fd-2. Mutations mimicking constitutive phosphorylation of T282 (T>E), S281 (S>E), or both (ST>EE) induce early flowering. Results are shown for two independent homozygous lines per construct. Statistical significance was calculated using an unpaired t-test compared to Col-0. \*\*\* indicates a significance level p < 0.01.

- **Figure 4.** Mapping of the FD binding site in the AP1 promoter.
  - (A) Normalized reads from six ChIP-seq experiments mapped on the AP1 locus. The result shows that the C-box is upstream of all peak summits.
  - (B) Nucleotide sequence encompassing the six peak summits shows several palindromic regions representing putative binding sites of *FD* on *AP1* promoter. The distance between the closest potential FD binding site under the ChIP-seq peaks and the C-box is 92 bp. Black triangles indicate the summits of the six separate ChIP-seq experiments. Putative FD binding sites are underlined and numbered from 1 to 4.
- **(C)** Electrophoretic mobility shift essay (EMSA) of the phosphomimic version of 776 FD-C (FD-C\_T282E) in combinations with 14-3-3v, FT and TFL1 using the 777 four putative binding sites reported in panel B. Free probes are not visible

778	because gels were running longer to maximize the distance between shifted
779	probes. Coloured squares indicate shifted probes.
780	<b>(D)</b> Comparison of the probes used for EMSA: the G-box in SEP3 promoter (Fig. 2)
781	and the binding site 2 in AP1 promoter. The putative FD binding site in AP1
782	promoter is also conserved in SEP3 promoter and overlaps with the G-box.
783	
784	Figure 5. RNA-seq results at the shoot apical meristem.
785	(A) Scatter blot of differentially expressed (DE) genes between the fd-2 mutant and
786	pFD::GFP-FD fd-2 (control) at 5 time points before and during the transition
787	to photoperiod-induced flowering. T0 - T5 indicate day of sample collection
788	before (T0) and 1, 2, 3, 5 days after shifting plants to long day. Red dots
789	indicate DE genes with a padj < 0.1.
790	(B) Venn diagrams showing the overlap between FD target genes identified by
791	ChIP-seq and DE genes found by RNA-seq at the SAM at each time point.
792	(C) Venn diagram showing the overlap between FD target unique genes identified
793	by ChIP-seq and DE unique genes in at least one time point found by RNA-
794	seq at the SAM. A total of 135 genes were classified as putative direct targets
795	of FD. Statistical significance was calculated using the Fisher's exact test.
796	Asterisk (*) indicates a significance level $p = 1.03E-07$ .
797	
798	<b>Figure 6.</b> Validation of FD targets in Col-0 and <i>p35S::FD</i> .
799	qRT-PCR analysis of 12 putative direct targets of FD. RNA was isolated from
800	7 day old seedlings to minimize any bias due to the early flowering of the
801	p35S::FD line. Error bars represent ±SD from three biological replicates.
802	
803	Figure 7. Flowering time of myc2 and afr1.
804	Flowering time of homozygous of myc2 and afr1 T-DNA insertion lines was
805	scored as days to flowering (A) and total leaves (B). Statistical significance
806	was calculated using unpaired t-test compared to Col-0. *** and ** indicate a
807	significance level $p < 0.01$ and $p < 0.05$ , respectively.
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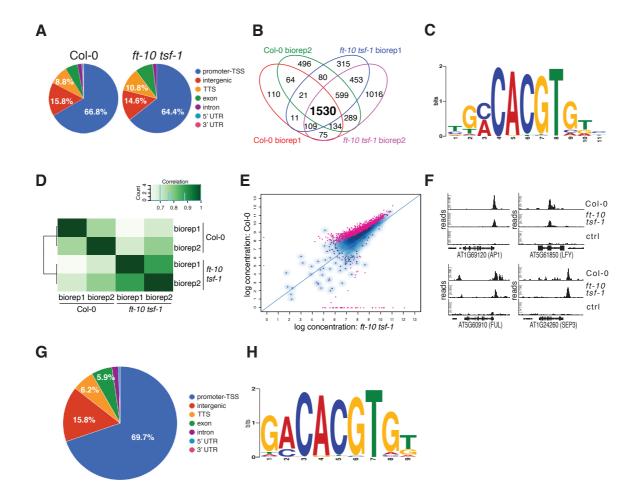
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**Figure 1.** Identification of FD targets by *pSUC2::GFP:FD* ChIP-seq in Col-0 and *ft-10 tsf-1* and *pFD::GFP:FD* ChIPseq in *fd-2*.

- (A) Annotation of high-confidence peaks found in two biological replicates in Col-0 and *ft-10 tsf-1*. TSS: transcription start site; TTS: transcription terminator site.
- **(B)** 4-set venn diagram representing the overlapping peaks among all the biological replicates from Col-0 and *ft-10 tsf-1*. The majority of the peaks (1530) are shared between the two genetic backgrounds.
- (C) Nucleotide logo of the predicted FD binding site.
- **(D)** Correlation heatmap calculated on a binding matrix based on ChIP-seq reads counts for Col-0 and *ft-10 tsf-1* samples (affinity scores). The presence/absence of FT and TSF is sufficient to discriminate the two genetic backgrounds.
- (E) Differentially bound (DB) peaks between Col-0 and ft-10 tsf-1. Red dots indicate differentially bound peaks with a FDR < 0.05. Blue dots represent peaks that were not significantly differentially bound.
- **(F)** Reads from Col-0, *ft-10 tsf-1* and control (ctrl) sample mapped against selected flowering-related genes.
- **(G)** Annotation of high-confidence peaks identified by ChIPseq in two biological replicates in *pFD*::*GFP*:*FD fd*-2.
- **(H)** Nucleotide logo of the predicted FD binding site at the SAM.

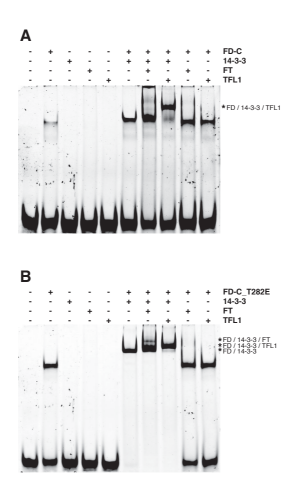
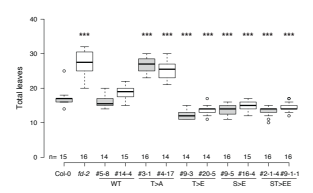


Figure 2. The C-terminal part of FD (FD-C) binds to a G-box in the SEP3 promoter in vitro.

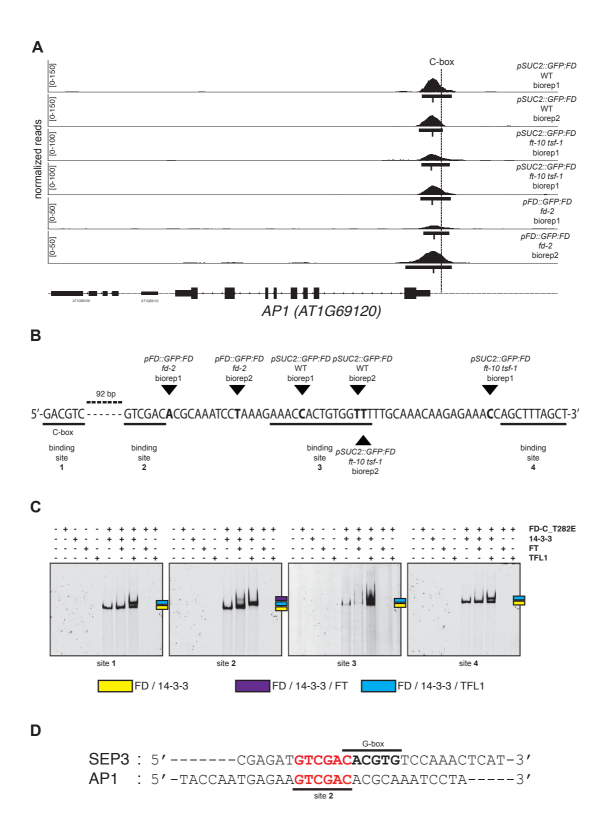
- (A) Electrophoretic mobility shift assay (EMSA) of the wild-type form of FD-C in combination with 14-3-3v, FT, and TFL1.
- **(B)** EMSA using the phosphomimic version of FD-C (FD-C\_T282E) in combination with 14-3-3v, FT and TFL1.

First lanes to the left in (A) and (B) show the free probe without any added proteins. Plus signs (+) above the lanes indicate which proteins were used in a specific EMSA reaction. Asterisk (\*) indicates higher order complexes.



**Figure 3.** Phosphorylation of FD at threonine 282 (T282) modulates flowering time in A. thaliana.

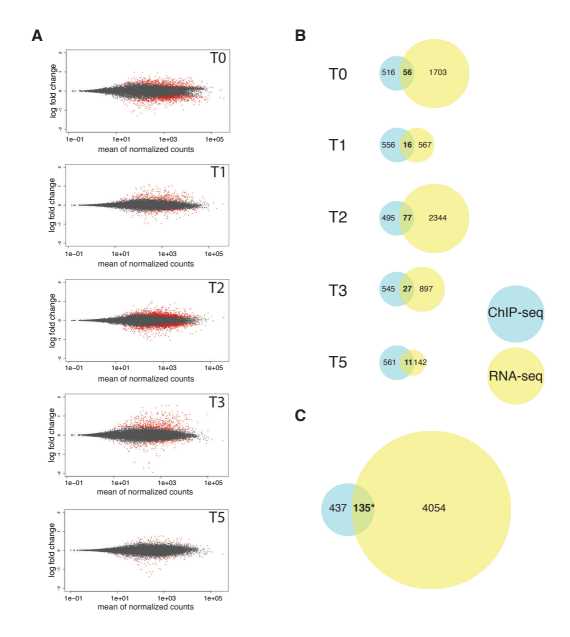
Box plot reporting the flowering time of control plants (Col-0), the fd-2 muntant, and the fd-2 mutant transformed with either wildtype (WT) FD cDNA under the control of the FD promoter (pFD::FD), non-phosphorable FD (T>A, pFD::FD\_T282A), or phosphomimic versions of FD (T>E, pFD::FD\_T282E; S>E, pFD::FD-S281E; ST>EE; pFD::FD\_S281E/T282E) as number of leaves formed by the main meristem (total leaves). Results are shown for two independent homozygous lines per construct. Number of plants (n) analyzed per genotype is indicated. Statistical significance was calculated using an unpaired t-test compared to Col-0. \*\*\* indicates a significance level p < 0.01.



**Figure 4.** Mapping of the FD binding site in the AP1 promoter.

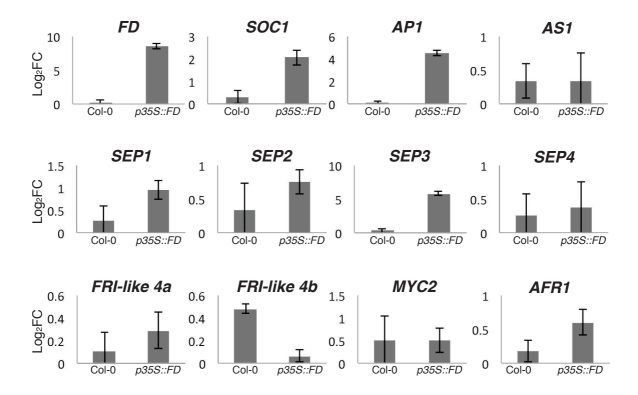
(A) Normalized reads from six ChIP-seq experiments mapped on the *AP1* locus. Horizontal bars below the peaks indicate regions of significant enrichment. The position of the summit is indicated for each peak. The C-box previously implicated in FD binding is located upstream (to the right) of peak summits in all six experiments.

- **(B)** Nucleotide sequence encompassing the six peak summits shows several palindromic regions representing putative binding sites of FD on the *AP1* promoter. The distance between the closest potential FD binding site under the ChIP-seq peaks and the C-box is 92 bp. Black triangles indicate the summits of the six separate ChIP-seq experiments. Putative FD binding sites are underlined and numbered from 1 to 4.
- (C) Electrophoretic mobility shift essay (EMSA) of the phosphomimic version of FD-C (FD-C\_T282E) in combination with 14-3-3v, FT and TFL1 using the four putative binding sites reported in panel B. Free probes are not visible because gels were running longer to maximize the distance between shifted probes. Coloured squares indicate shifted probes.
- **(D)** Comparison of the probes used for EMSA: the G-box in *SEP3* promoter (Fig. 2) and the binding site 2 in *AP1* promoter. The putative FD binding site (site 2) in the *AP1* promoter, which is also conserved in *SEP3* promoter where it overlaps with the G-box, is marked in red



**Figure 5.** RNA-seq results at the shoot apical meristem.

- (A) Scatter plots of differentially expressed (DE) genes between the *fd-2* mutant and *pFD::GFP-FD fd-2* (control) at 5 time points before and during the transition to photoperiod-induced flowering. T0 T5 indicate day of sample collection before (T0) and 1, 2, 3, 5 days after shifting plants to long day. Red dots indicate DE genes with a padj < 0.1.
- **(B)** Venn diagrams showing the overlap between 572 unique FD target genes identified by ChIP-seq and DE genes found by RNA-seq at the SAM at five time points before (T0) and during (T1 T5) the transition to flowering.
- (C) Venn diagram showing the overlap between 572 unique FD target genes identified by ChIP-seq at the SAM (two biological replicates) and genes found to be differentially expressed at the SAM by RNA-seq at least at one time point (T0, T1, T2, T3, T5; three biological replicates per time point). A total of 135 genes were classified as putative direct targets of FD. Statistical significance was calculated using the Fisher's exact test. Asterisk (\*) indicates a significance level p = 1.03E-07.



**Figure 6.** Validation of FD targets in Col-0 and *p35S::FD*.

RT-qPCR analysis of 12 putative direct targets of FD. RNA was isolated from 15-20 7-day-old Col-0 and p35S::FD seedlings. Error bars represent  $\pm$ SD from three biological replicates.

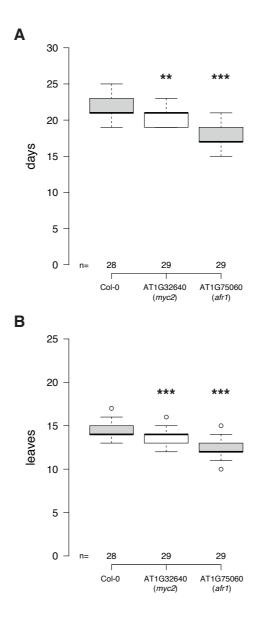


Figure 7. Flowering time of myc2 and afr1.

Flowering time of homozygous myc2 and afr1 T-DNA insertion lines was scored as days to flowering (A) and total leaves (B). Number (n) of plants per genotype is indicated. Statistical significance was calculated using an unpaired t-test compared to Col-0. \*\*\* and \*\* indicate a significance level of p < 0.01 and p < 0.05, respectively. Unfilled circles represent outliers.