TRANSPARENT TESTA GLABRA 1 participates in flowering time regulation in Arabidopsis thaliana

Barbara A.M. Paffendorf¹, Rawan Qassrawi¹, Andrea M. Meys¹, Laura Trimborn¹ and Andrea Schrader¹,²

¹Botanical Institute, Department of Biology, University of Cologne, Cologne, Germany
²RWTH Aachen University, Institute for Biology I, Aachen, Germany

ABSTRACT

Pleiotropic regulatory factors mediate concerted responses of the plant’s trait network to endogenous and exogenous cues. TRANSPARENT TESTA GLABRA 1 (TTG1) is such a factor that has been predominantly described as a regulator of early developmental traits. Although its closest homologs LIGHT-REGULATED WD1 (LWD1) and LWD2 affect photoperiodic flowering, a role of TTG1 in flowering time regulation has not been reported. Here we reveal that TTG1 is a regulator of flowering time in Arabidopsis thaliana and changes transcript levels of different targets within the flowering time regulatory pathway. TTG1 mutants flower early and TTG1 overexpression lines flower late at long-day conditions. Consistently, TTG1 can suppress the transcript levels of the floral integrators FLOWERING LOCUS T and SUPPRESSOR OF OVEREXPRESSION OF CO1 and can act as an activator of circadian clock components. Moreover, TTG1 might form feedback loops at the protein level. The TTG1 protein interacts with PSEUDO RESPONSE REGULATOR (PRR)s and basic HELIX-LOOP-HELIX 92 (bHLH92) in yeast. In planta, the respective pairs exhibit interesting patterns of localization including a recruitment of TTG1 by PRR5 to subnuclear foci. This mechanism proposes additional layers of regulation by TTG1 and might aid to specify the function of bHLH92. Within another branch of the pathway, TTG1 can elevate FLOWERING LOCUS C (FLC) transcript levels. FLC mediates signals from the vernalization, ambient temperature and autonomous pathway and the circadian clock is pivotal for the plant to synchronize with diurnal cycles of environmental stimuli like light and temperature. Our results suggest an unexpected positioning of TTG1 upstream of FLC and upstream of the circadian clock. In this light, this points to an adaptive value of the role of TTG1 in respect to flowering time regulation.

INTRODUCTION

While a species adapts to ranges of abiotic and biotic conditions, the individual plant must cope with its daily local conditions. It achieves this by integrating various signaling pathways and the current status of the plant itself—for example, its developmental stage or the combination and availability of metabolites. Pleiotropic regulators aid in concerted responses and, thereby, regulate a subset of the plant’s trait network. Due to the depth
of insights achieved in the past decades of plant molecular biology, its model species *Arabidopsis thaliana* (*A. thaliana*) *(Koornneef & Meinke, 2010)* is well suited to analyze such pleiotropic regulators.

TRANSPARENT TESTA GLABRA 1 (TTG1) is one such pleiotropic regulator. The gene is expressed in all major organs of *A. thaliana* including the shoot apical meristem *(Walker et al., 1999)*. TTG1 encodes a WD40 repeat protein *(Walker et al., 1999)*. The integrity of the protein’s WD40 repeats is crucial to its function and its C-terminus is expected to be of high relevance for the protein’s proper folding and domain structure *(Zhang & Schrader, 2017)*. TTG1 is known as the head of an evolutionarily conserved gene regulatory network that controls five major traits of adaptive value: seed pigmentation (production of proanthocyanidin), accumulation of anthocyanidins (in seedlings), seed coat mucilage production, trichome and root hair patterning *(Zhang et al., 2003)*. Molecular mechanisms underlying the early developmental traits are under investigation since decades in *A. thaliana* and beyond. Already in 1981, the *ttg1* syndrome was described for induced *A. thaliana* mutants characterized by yellow seeds with transparent testa and by the absence of trichomes (glabrous leaves), anthocyanin accumulation as well as seed mucilage *(Koornneef, 1981)*. Few additional traits like the carbon partitioning between the seed oil, seed pigment and seed mucilage biosynthetic pathways were analyzed in dependence of TTG1 *(Chen et al., 2015; Li et al., 2018)*. However, surprisingly little is known about the role of TTG1 towards late developmental traits.

One of the most important developmental switches in the plant’s life cycle is the transition from the vegetative to the reproductive phase. The appropriate regulation of flowering time is essential for the reproductive success of plants and, therefore, a key determinant of plant fitness. Several genetically identified pathways that are involved in the regulation of flowering time are influenced by environmental (e.g., vernalization, ambient temperature and photoperiod) and endogenous (e.g., autonomous, gibberellin, circadian clock, age, sugar budget) signals *(Blumel, Dally & Jung, 2015)*. These interwoven regulatory mechanisms converge to the floral integrators *FLOWERING LOCUS T* (FT), *SUPPRESSOR OF OVEREXPRESSION OF CO1* (SOC1) and also *LEAFY* (LFY) *(Simpson & Dean, 2002)*.

CONSTANS (CO) and FT form an important module of the photoperiodic pathway. CO expression rises about 8 h after dawn with a peak at night *(Suarez-Lopez et al., 2001)*. The accumulating CO protein activates the florigen gene *FT* in leaves *(An et al., 2004; Song et al., 2015)*. In the night, CO is degraded through the COP1/SPA complex (CONSTITUTIVE PHOTOMORPHO-GENESIS 1/SUPPRESSOR OF PHYA-105) *(Jang et al., 2008; Laubinger et al., 2006; Liu et al., 2008)*. Hence, at long days, sufficient FT protein is formed in the leaves and moves to the shoot apical meristem where it induces flowering *(Andres & Coupland, 2012)*. Although CO is expressed under short-day (SD) conditions, it cannot sufficiently induce *FT* expression due to the extended night *(Valverde et al., 2004)*. Consequently, mutants of the photoperiod pathway flower late under long-day (LD) conditions and do not deviate in flowering time from the wild type at SD conditions. One such mutant is the *gigantea* (gi) mutant. GI is an activator of CO *(Sawa et al., 2007)*. Its protein levels are also regulated by COP1 in presence of EARLY FLOWERING 3 (ELF3) *(Yu et al., 2008)*.
A. thaliana is a facultative LD plant. Winter annual accessions flower late and are responsive to vernalization which reduces the FRIGIDA (FRI)-activated transcript levels of the floral repressor FLOWERING LOCUS C (FLC) through epigenetic modifications at the FLC locus (Deng et al., 2018; Hepworth & Dean, 2015). In the rapid-cycling summer annual accessions, either FRI is defective, which reduces FLC transcript levels, or the FLC allele is weak (Michaels et al., 2003). Low levels of FLC induce flowering as FLC is a suppressor of FT (Searle et al., 2006). FT activates the downstream transcription factors LEAFY (LFY) and APETALA1 (API) at the shoot apical meristem and thereby causes flowering when an FT threshold is passed (Turck, Fornara & Coupland, 2008).

SOC1 acts downstream of FT and upstream of LFY. Similarly to FT, it is directly targeted by the floral repressor FLC (Lee & Lee, 2010) which mediates signals from the autonomous and vernalization response pathways. Both pathways act through suppression of FLC expression (Simpson & Dean, 2002). SHORT VEGETATIVE PHASE (SVP) is an interaction partner of FLC (Li et al., 2008) and both are mediators of the ambient temperature pathway (Lee et al., 2007; Simpson & Dean, 2002). Ambient temperature adjusts flowering time in a way that cool temperature delays flowering, whereas warm temperature accelerates flowering (Balasubramanian et al., 2006; Blazquez, Ahn & Weigel, 2003). SVP itself also acts as a direct suppressor of FT and SOC1 (Li et al., 2008). Moreover, SVP can activate members of a group of additional FT suppressors, the APETALA2 (AP2) domain containing transcription factors TEMPRANILLO (TEM) 1 and TEM2 (RAV transcription factors with AP2/ERF and B3 DNA-binding domain), AP2 and the AP2-like transcription factors SCHLAFMÜTZE (SMZ), SCHNARCHZAPFEN (SNZ), TARGETs OF EARLY ACTIVATION TAGGED (EAT) (TOE) 1, TOE2 and TOE3 (Tao et al., 2012; Yant, Mathieu & Schmid, 2009). These AP2 domain containing factors act directly at the FT gene. TOE1 is able to bind to the FT promoter close to the CO-binding site (Zhang et al., 2015). SMZ also seems to affect FT expression directly, since FT was found as a target of SMZ in a ChIP-chip assay (Mathieu et al., 2009). TEM1 and TEM2 act as FT repressors by binding to its 5’ UTR. Furthermore, it is suggested that the balance between TEM and CO controls FT transcription and thereby is involved in determination of flowering (Castillejo & Pelaz, 2008).

The AP2 domain containing factors are not only connected with the ambient temperature pathway but also with the gibberellin signaling pathway by TEM1 and TEM2 (Osnato et al., 2012). Moreover, TOE1 interacts with the activating region of CO and the LOV domain of FLAVIN-BINDING KELCH REPEAT F-BOX1 (FKF1). This prevents CO from activating FT transcription and FKF1 from stabilizing CO (Zhang et al., 2015).

Upstream of CO, the circadian clock influences the flowering time regulatory pathway. Circadian oscillators are the key for a plant to synchronize with the external environmental cues providing an adaptive advantage (Dodd et al., 2005; Michael et al., 2003). The circadian clock and its feedback loops cause in general rhythmic gene expression within and downstream of the clock. A screen analyzing the MYELOBLASTOSIS (MYB), basic HELIX-LOOP-HELIX (bHLH) and basic region/leucine zipper factors in A. thaliana found that 20% of these are under the control of the clock (Hanano et al., 2008). In turn,
MYB3R2, bHLH69 and bHLH92 were found to alter clock parameters when overexpressed and therefore might position upstream of the clock (Hanano et al., 2008).

The core negative feedback loop of the clock is formed by the MYB-like proteins CIRCADIAN CLOCK ASSOCIATED 1 (CCA1) and LATE ELONGATED HYPOCOTYL (LHY) which are expressed in the morning and the evening-expressed PSEUDO RESPONSE REGULATOR (PRR) PRR1/TIMING OF CAB EXPRESSION 1 (TOC1) (Oakenfull & Davis, 2017). Several additional loops are formed within the central oscillator. PRR9, PRR7, PRR5 and PRR3 peak successively during the day (Matsushika et al., 2000) filling the gap between CCA1/LHY and TOC1. The PRRs act as suppressors of CCA1 and LHY (Nakamichi et al., 2010). Moreover, GI forms a predicted feedback loop with TOC1 (Locke et al., 2006) and the evening complex consisting of ELF4, ELF3 and LUX ARRHYTHMO (LUX) is required to maintaining circadian rhythms through regulating different key clock genes (Huang & Nusinow, 2016).

The PRR proteins have an N-terminal pseudo-receiver domain which is similar to the phospho-accepting receiver of the two-component response regulators but lacks the presumed phosphor-accepting aspartate. At their C-terminus, a CO, CO-like and TOC1 (CCT) motif is shared by the name-giving proteins (Makino et al., 2000; Matsushika et al., 2000; Strayer et al., 2000). PRRs act antagonistically with LHY/CCA1 on the downstream CO-FT module (Nakamichi et al., 2007). At the protein level, PRRs interact with and stabilize the CO protein enhancing CO-mediated FT transcription (Hayama et al., 2017).

Only few transcriptional activators of the circadian clock are known (Shim, Kubota & Imaizumi, 2017). One of these is LIGHT-REGULATED WD1 (LWD1). LWD1 and LWD2 are the closest homologs of TTG1 that regulate photoperiodic flowering (Wu, Wang & Wu, 2008). Double mutants flower early at LD conditions and exhibit increased FT transcript levels (Wu, Wang & Wu, 2008). The LWD genes are rhythmically expressed in dependence of PRR9 which forms a feed-back loop with LWD1 (Wang et al., 2011). LWD1 can bind to the promoter of PRR5, PRR9 and PRR1/TOC1. With TEOSINTE BRANCHED1-CYCLOIDEA-PCF20 (TCP20) and TCP22 it binds to the CCA1 promoter activating its expression (Wang et al., 2011; Wu et al., 2016). To date, there was no evidence suggesting an involvement of TTG1 in the regulation of the circadian clock and flowering time. A potential involvement of TTG1 in its transcriptional control has not been reported.

Here we reveal that TTG1 can modulate flowering time along with an initial embedding of TTG1 in the flowering time regulatory pathway. Most strikingly, TTG1 can suppress FT and SOC1 transcript levels and increase those of clock components while reducing their amplitude as observed within one day. PRR proteins can interact with TTG1 in yeast and, when co-expressed in planta, exhibit interesting subcellular localization patterns of and with TTG1. In the same systems, TTG1 also interacts with and modulates the localization of bHLH92. Flowering time results at LD conditions as well as the integrators’ transcript levels are in line with an increase of the FLC transcript level upon TTG1 overexpression. Together, we suggest that TTG1 acts in multilayered processes in flowering time regulation and it might act upstream of FLC and the clock.
MATERIALS & METHODS

Plant material and growth conditions

The used A. thaliana mutants ttg1-9, ttg1-11, ttg1-21, ttg1-22, glabra3 (gl3)-3, enhancer of gl3 (egl3)-19114, transparent testa 8 (tt8)-SALK, myc1-1, cop1-4 (all Col-0), ttg1-1 (Ler), ttg1-10 (Ws) (Table S7) have been described before (Alonso et al., 2003; Appelhagen et al., 2014; Jakoby et al., 2008; Koornneef, 1981; Larkin et al., 1994; Larkin et al., 1999; McNellis et al., 1994; Pesch et al., 2013; Rosso et al., 2003; Walker et al., 1999; Wester et al., 2009). Primers used for genotyping (including dCAPS primers) are listed in Table S8 (Appelhagen et al., 2011; Jaegle et al., 2016; Neff, Turk & Kalishman, 2002; Schrader et al., 2013). The floral dip method (Clough & Bent, 1998) was used to generate overexpression lines in Col-0 and cop1-4 background. T1 plants were BASTA selected and resistant plants were screened for YFP fluorescence using a Leica stereomicroscope (MZ FLIII) (Leica Microsystems, http://www.leica-microsystems.com). This analysis was repeated for plants homozygous for the insert being at least in T3 generation and overexpressing YFP-TTG1 (Table S4). Two walk-in plant chambers were used. Detailed conditions at the respective used areas are listed in Table S1. For flowering time experiments, single seeds were placed in parallelly prepared pots with soil and stratified for 7 d at 5°C before being transferred to the respective growth condition.

Seedlings for qRT-PCR experiments were sterilized with 70% (v/v) ethanol, 2% NaOCl, stratified at 5°C and grown on Murashige and Skoog medium containing 1% sucrose at “cold” LD conditions (see Table S1). Seedlings for circadian transcript profiles were snap frozen in liquid nitrogen (~100 mg) on day eight starting at ZT0 in 4 h intervals until ZT20. Samples used comparing transcript levels in the overexpression lines OE01-03 and OE19-21 were similarly harvested at ZT11 and ZT13 as well as parallelly grown seedlings for comparing protein levels in the same lines at ZT11 and ZT10, respectively. Three biological replicates were analyzed for circadian profiles and comparisons of transcript and protein levels among the TTG1 overexpression lines.

Phenotyping

Flowering time was recorded as the number of post-stratification days until bolting and the total number of leaves at the time point of bolting. Bolting was defined as the time at which the first bud was visible. Two (bHLH overexpression lines’ experiment) or three experiments (TTG1 overexpression lines, all ttg1 mutant sets) were conducted for each set of analyzed genotypes and for each condition with at least six individual plants per genotype and experiment. See Table S2 for details.

Constructs

Gateway™ (Invitrogen™, http://www.invitrogen.com) entry clones containing the coding DNA sequence (CDS) for the respective protein were generated using BP reaction with the previously published vectors TTG1 pAS2.1 and GL3 pcACT2 (Pesch et al., 2015), EGL3 pcACT2, TT8 pAS2.1, MYC1 pAS2.1 (vectors provided by M Pesch) or PCR products using primers for TOC1, PRR5, PRR7, PRR9, bHLH92, LWD1 and LWD2 CDS are listed in Table S8 and cDNA from Col-0 seedlings as
a template or using the vectors 35S:PRR7:CFP and 35S:PRR9:CFP (Hayama et al., 2017) as a template, respectively. The used entry vector in all cases was pDONR207 (Invitrogen). All entry vectors were sequenced. LWDs were amplified from the cDNA using GGGGACAGTTTGTACAAAAAAGCAGGCTTAATGGGAACGAGCAGCGATCC and GGGGACCACTTTGTACAAGAAAGCTGGGTTTCAAACCCTGAGAATT for LWD1 and GGGGACAGTTTGTACAAAAAAGCAGGCTTAATGGTTACGAGCAGCGATCA and GGGGACCACTTTGTACAAGAAAGCTGGGTTTCAGACCCGGAGAATC for LWD2. See Table S9 for the TTG1, LWD1 and LWD2 CDS which were also used in the alignment in Fig. S2. To generate the construct used for the overexpression lines OE01-03 in Col-0 and OE19-OE21 in cop1-4 background, TTG1 pDONR207 was recombined using Gateway™ LR Clonase™ (Invitrogen™, a version for C-terminal fusions was published before Feys et al., 2005). In analogy, the CDS in pDONR207 for GL3 (OE04-OE06), EGL3 (OE07-OE09), TT8 (OE10-OE12) and MYC1 (OE13-OE15) were recombined into pENSG-YFP and used to generate the overexpression lines numbered as given in brackets and expressing YFP-bHLH fusion proteins driven by the Pro35s.

For tobacco co-localization experiments, TTG1 pDONR207 and PRR5 pDONR207 were recombined into pNmR (Schrader et al., 2013) and PRR5, PRR7, PRR9, TOC1, bHLH92 (all in pDONR207) were recombined into pENSG-CFP (N. Medina-Escobar, a version for C-terminal fusions was published before Feys et al., 2005) and for the Y2H experiments, also in pAS2.1-attR and pACT-attR (Clontech, http://www.clontech.com, modified J.F. Uhrig). LWD1 pDONR207 and LWD2 pDONR207 were similarly used in combination with pAS2.1-attR and pACT-attR.

In analogy to YFPattB1-pBat-TL-B-p35s and RFP-HAattB1-pBat-TL-B-p35s previously created as negative controls (Schrader et al., 2013) CFPattB1 was amplified from pENSG-CFP with primers ANS393 and ANS235 and recombined in pBat-TL-B-p35s (Schrader et al., 2013) to obtain CFPattB1-pBat-TL-B-p35s.

qRT-PCR experiments
About 100 mg of seedlings was harvested for each RNA extraction. RNA extractions were done according to the manufacturer’s instructions (RNasey Plant Mini Kit, Qiagen, http://www.qiagen.com) using a Tissue Lyser (Qiagen) and followed by DNase I (ThermoFisher, https://www.thermofisher.com) treatment. RNA integrity was tested on a gel prior to cDNA synthesis (SuperScript™ III First-Strand Synthesis System, Invitrogen, or the RevertAid First Strand cDNA Synthesis Kit, ThermoFisher) and RNaseH treatment as suggested before (Martel, Grundemann & Schomig, 2002). A PCR using Elongation factor 1-alpha 1 (EF1ALPHA) primers (Kirik et al., 2007) spanning an intron served as a control to ensure that there was no genomic DNA in the cDNA synthesis (Primers in Table S8).

qRT-PCR was performed using the QuantStudio 5 Real-Time PCR Systems (ThermoFisher) with POWER SYBR Green PCR-Master Mix (Applied Biosystems), the respective cDNA and gene-specific primers. UBQ10 (UBIQUITIN10) was used as a reference gene (Harari-Steinberg, Ohad & Chamovitz, 2001; Sun & Callis, 1997). Three biological replicates with three technical replicates each were performed. Calculations are
described in detail in Tables S3 and Table S5. All used primers are listed in Table S8. Most of these were described before (Grigorova et al., 2011; Hayama et al., 2017; Li et al., 2008; Maier et al., 2013; Nakamichi et al., 2007; Shin et al., 2017; Wang et al., 2014; Wang et al., 2011; Wenden et al., 2011; Yu et al., 2012; Zhang et al., 2015; Zou et al., 2013). For TTG1 endo, TTG1 both and TTG1 no LWD see Fig. 1G and Fig. S2.

Comparison of protein levels
Samples were homogenized under liquid nitrogen to compare YFP-tagged TTG1 in the overexpression lines OE01-03 and OE19-21. 150 µl of lysis buffer (Kirik et al., 2007) were added to the powder and incubated for 30 min at 4 °C (rotating). 100 µl of the supernatant following centrifugation were mixed with 100 µl 2x Laemmli, boiled for 10 min at 95 °C and centrifuged for 1 min at 10 600 g. Samples were separated on SDS-PAGE gels, subsequently blotted and immunodetected (α-GFP (IgG1K, Roche), α-mouse (Jackson ImmunoResearch, http://www.jacksonimmuno.com)). After GFP detection using the SuperSignal® West Femto Maximum Sensitivity Substrate (ThermoFisher) and a LAS-4000 Mini bioimager (GE Healthcare Life Sciences (formerly Fuji), http://www.gelifesciences.com), blots were stripped as suggested by Abcam (http://www.abcam.com/ps/pdf/protocols/stripping%20for%20reprobing.pdf) using mild stripping buffer (1L: 15 g glycine, 1 g SDS, 10 ml Tween20, pH to 2.2) and re-probed with α-histone H3 (ab1791, Abcam, http://www.abcam.com) and α-rabbit (A6154, Sigma-Aldrich, http://www.sigmaaldrich.com).

Y2H experiments and Y2H screening
The TTG1pAS2.1-attR construct (Pesch et al., 2015) was used as bait to screen an A. thaliana root cDNA library in yeast (Klopffleisch et al., 2011). Y2H screening was performed as described before using 5mM of 3-AT (Soellick & Uhrig, 2001). Y2H assays were done by co-transformation of pAS2.1-attR/pACT-attR (or TTG1-pcACT2 Pesch et al., 2015) vector combination as described previously (Gietz & Schiestl, 2007). GFPpAS2.1-attR and GFPpACT (Schrader et al., 2013) served as a negative control. At least three replicates were conducted for each Y2H co-transformation experiment and 6 or 8 individual colonies per transformation were resolved in water in 96-well plates and transferred to SD-LW or SD-LWH plates supplemented with different 3-AT concentrations (3, 5, 10, 15, 20, 30 mM) using a 96-well replica plater. Plates were scanned after one (only SD-LW controlling for successful double transformation and providing a relative comparison of transferred yeast amounts), three and seven days.

Co-localization, microscopy, phenotypic characterization of ttg1-21 and ttg1-22
Nicothiana benthamiana leaves were infiltrated as described before (Yang, Li & Qi, 2000) but using the Agrobacterium tumefaciens strain GV3101 pMP90RK harboring the respective constructs and Agrobacteria expressing the silencing suppressor TBSV19K (Voinnet, Pinto & Baulcombe, 1999). Infiltrated plants were analyzed three days post-infiltration. CFP-attB1-pBat-TL-B-p35s (this study), YFP-attB1-pBat-TL-B-p35s and RFP-HA-attB1-pBat-TL-B-p35s (Schrader et al., 2013) were used as controls for co-expression with single
Figure 1  TTG1 has an effect on flowering time regulation in the A. thaliana Col-0 ecotype at long-day conditions. (A–D) Flowering time of ttg1 mutants in Col-0 (continued on next page...)

Full-size DOI: 10.7717/peerj.8303/fig-1
Figure 1 (...continued)

background. In (A–F, H, I), flowering time was recorded as the number of days until bolting (A, C, E, H) or the number of leaves at the time-point of bolting (B, D, F, I). Plants were grown at long-day conditions (16 h light, 8 h darkness) at 21°C ("cold") or 23°C ("warm"). Black lines in the box plots represent the median, red dots are the mean for the shown representative experiment and crosses mark outliers. Asterisks indicate significant differences (*$P < 0.05$) between the mutants and the Col-0 wild type (orange) or between the two conditions (grey) in the shown representative experiment. Numbers in brackets indicate the number of experiments for which a significant difference was observed out of the total number of experiments. (G) Primer binding sites for the used primer pairs for qRT-PCR (arrow heads) relative to the TTG1 genomic, cDNA, CDS and overexpression construct sequence. The position of the point mutations in the ttg1-9 and ttg1-11 is indicated above. The T-DNA insertion for ttg1-21 and ttg1-22 are provided in Fig. S1 characterizing the mutants’ phenotypes. See Fig. S2 for an alignment of TTG1 with LWD1 and LWD2 in the primer binding region. The sense primers for the primer pairs “TTG both” and ”TTG1 no LWD” are overlapping and the sequence of the latter deviates at its 3′ end from the sequence for LWD1 and LWD2. The name “TTG1 both” indicates that the CDS as well as the construct are amplified by this primer pair. (H, I) Flowering time of TTG1 overexpression lines as described above. (J–M) Characterization of transcript and protein levels in lines overexpressing YFP-TTG1 in Col-0 (OE01-03) and cop1-4 (OE19-21) background driven by the 35S promoter. (J–L) Highest levels of TTG1 transcript were observed in OE01-03 (Col-0) and OE20 (cop1-4) while the endogenous TTG1 transcript levels were in general not affected by overexpression despite in the OE19 (cop1-4) line. Note the elevated TTG1 levels in cop1-4 mutants. Transcript levels were determined by qRT-PCR relative to UBQ10 using 8-day-old seedlings and are normalized with Col-0 wild type values. Data represent the mean of three independent experiments. Asterisks indicate significant differences (#, $P < 0.1$; *, $P < 0.05$) between overexpression lines and the backgrounds Col-0 (orange) and cop1-4 (magenta), respectively. Please note that the scale in (K, L) differs from the scale in (J). The solid line equals 1. The y-axis is in log$_{10}$-scaled. Error bars indicate the SD. (M) Western blot using 7d-old long-day grown seedlings from one of the repeats used in (J–L). In OE03 (Col-0), YFP-TTG1 levels varied between experiments never exceeding those in OE19 (Col-0). See also Fig. S3. Tables S2–S4 provide more details on the underlying data and statistics. OE01-03: Pro35s:YFP-TTG1 (Col-0), three independent insertion lines. OE19-21: Pro35s:YFP-TTG1 (cop1-4), three independent insertion lines.

fluorescent tag fusion protein. The experiment was conducted at least three times for each combination. Infiltrations for the co-expression of CFP and RFP-TTG1 or YFP-TTG1 and RFP, respectively, were included in all experiments. CLSM was performed using a Leica SP8 confocal microscope (Leica Microsystems). Z-stacks were acquired by sequential scanning starting with the laser with the higher wavelength. The LAS Application Suite X (Leica Microsystems) was used to extract and merge images for co-localization figures.

Stacks of small leaves acquired as described before (Failmezger et al., 2013) were merged using Combine ZP (by Alan Hadley, https://combinezp.software.informer.com/) for Fig. S1. Pictures of seeds, seedlings and older leaves (14d-old soil and LD grown) were acquired using a stereo microscope Leica stereomicroscope (MZ FLIII) with the MultiFocus and Montage option of the Leica Application Suite V3 (Leica Microsystems) The step-size was 20 μm (seeds, seedlings) and 50μm (leaves). Seedlings were sterilized as described above and grown on MS (4% sucrose) at constant light at 21 °C.

Data analysis and statistics

All statistics (Tables S2, S3 and S5), most data analysis, all box plots and plots for qRT-PCR results were generated using R version 3.4.1 (R Core Team, 2017) with the following packages: dplyr (Wickham et al., 2019), extrafont (Chang, 2014), ggplot2 (Wickham, 2016), plyr (Wickham, 2011), scales (Wickham, 2017), tidyr (Wickham & Henry, 2018). Schematics
for Fig. 1G, Figs. S1A and S2 were extracted from CLC DNA Workbench (CLC bio A/S, http://www.clcbio.com).

Relative protein amounts were determined using Fiji (imageJ 1.52 h, http://imagej.nih.gov/ij). ROIs of the same size for all bands analyzed within both images of one blot detected with both antibody combinations were measured for their mean grey value intensity. The background close to each band was subtracted, GFP values were set relative to the respective Histone H3 values and values obtained for one blot were normalized to OE01 and, in one case for which OE01 was not evaluated, OE20 was used (for OE19-OE21 analysis). Results are shown in Fig. S3.

RESULTS

**TTG1 has an effect on flowering time**

To date, TTG1 has been analyzed in detail for early traits while little is known about its role in the regulation of late developmental traits. When growing *ttg1* mutants, we observed that these flowered slightly earlier than the wild type. Therefore, we selected flowering time as a key late developmental trait. We analyzed the classical *ttg1-9* and *ttg1-11* mutants at the same “warm” long-day conditions (chamber set to 22 °C, on average 23.7 °C at the plant’s level, Table S1). We also analyzed flowering time at an about 2 °C reduced temperature which we named “cold” long-day conditions (chamber set to 20 °C, on average 21.4 °C at the plant’s level, Table S1). Both mutants flowered significantly earlier at both conditions (Figs. 1A–1B, Table S2). *ttg1-9* exhibited the strongest flowering time phenotype and was only slightly responsive to the difference in temperature in contrast to *ttg1-11* and the wild type. Here, the “cold” condition proved to beneficial in order to reveal differences in flowering time regulation among the *ttg1* mutants.

In both used EMS mutants, an amino acid in the TTG1 protein is changed. These mutants are known to be no null-mutants at least with respect to their effect on trichome patterning. Therefore, we obtained the recently described additional mutants *ttg1-21* and *ttg1-22* in Col-0 background to extend the flowering time analysis (Appelhagen et al., 2014; Rosso et al., 2003). In both mutants, a T-DNA insertion causes a premature stop codon within the inserted T-DNA. The insertion in *ttg1-21* is close to the start and in front of the WD40 domain (Fig. S1A). Therefore, it can be expected that this mutant is a null mutant or at least a comparably strong mutant. In *ttg1-22*, the T-DNA inserts in proximity to the stop codon. This might also cause a strong phenotype as seen for the premature stop codon mutant *ttg1-1* in Landsberg erecta (Ler) background. We tested these mutants for some of the early TTG1-dependent developmental traits. Compared to the wild type, all used Col-0 mutants showed the analyzed aspects of the *ttg1* syndrome (Koornneef, 1981) (Fig. S1). Moreover, we wondered, why the flowering time phenotype of *ttg1* mutants was not reported before. Therefore, we added an often used mutant in Ler background, *ttg1-1*, a point mutant with a premature stop codon close to the end of TTG1 (Larkin et al., 1994; Walker et al., 1999), and a mutant in Wassilewskija (Ws) background, *ttg1-10*, carrying a point mutation in the TTG1 promoter (Larkin et al., 1994; Larkin et al., 1999).

The different mutants and variants showed different patterns of flowering time phenotypes relative to the respective backgrounds (Figs. 1C–1F). In Col-0 background,
similar to ttg1-9 and ttg1-11, ttg1-21 and ttg1-22 flowered significantly earlier in terms of time and leave number at both tested conditions with one exception. For ttg1-21 grown in the warm condition, the number of days only deviated significantly from the wild type in one out of three repeats. Therefore, early flowering cannot be concluded in this case. For ttg1-21 and ttg1-22 it can be summarized that the mutants responded to temperature in the same way as the wild type in regard to time (days) and number of leaves produced until flowering. Again, there was one exception, only in one of three repeats for ttg1-21, a significantly different number of leaves was recorded when comparing between the warm and cold condition at flowering time. This suggests that these mutants are less responsive to temperature affecting its number of leaves at flowering time than the wild type and ttg1-22.

As compared to the Ler wild type, flowering time was not significantly reduced in ttg1-1 at both conditions. Therefore, it is not surprising that the flowering time phenotype was not reported before for this heavily used ttg1 mutant. In contrast to the other analyzed mutants, ttg1-10 carries its mutation as a point mutation in the TTG1 promoter. Interestingly, ttg1-10 mutants flowered significantly later at both conditions for both recorded flowering phenotypes as compared to its wild type. Ler and ttg1-1 did not respond to the difference in temperature, when it comes to the number of leaves produced at flowering time. ttg1-1 and Ws responded only in one out of two experiments to the difference in temperature for the number of days (ttg1-1) and leaves (Ws), respectively. In these cases, a reduced response to temperature cannot be concluded. In all other cases, the reduced temperature caused a delay in flowering time as also observed for Col-0 in the other experiments.

TTG1-9 and TTG1-11 encode TTG1 protein variants. Therefore, the observed early flowering might be due to a gain- or loss-of-function of the TTG1 variant or TTG1 gene in the respective mutants. To specify this, we generated overexpression lines of TTG1 (Col-0) driven by the constitutively active 35s promoter in Col-0 background (Pro35S::YFP-TTG1 (Col-0), the three lines are subsequently named OE01-OE03, Fig. 1G). As a regulatory hub in light signaling, COP1 is known to regulate protein stability of relevant flowering time regulators (Jang et al., 2008; Liu et al., 2008; Yu et al., 2008) and COP1 can interact with a TTG1 gene regulatory network component at the protein level (Maier et al., 2013). Therefore, we included cop1-4 as a background in the flowering time analysis (Pro35S::YFP-TTG1 (cop1-4), the three lines are subsequently named OE19-OE21). All overexpression lines in wild-type background flowered later than the wild type at both temperatures and in respect to time and number of leaves (Figs. 1H–1I). This suggests a loss-of-function in the mutant scenario. Compared to Col-0, cop1-4 produced significantly less leaves at the time point of flowering. An increase in number of leaves and days in this background was only observed for the overexpression line OE20 as compared to its background (Figs. 1H–1I). This might be due to different transcript or protein levels in the overexpressors.

The overexpression constructs did not affect the endogenous TTG1 transcript levels despite for OE19 in cop1-4 background. In this line, a significant reduction of endogenous TTG1 transcript was observed (Fig. 1J, Table S3, see Fig. 1G and Fig. S2 for the selective primer design). Interestingly, in cop1-4 mutants, TTG1 CDS transcript levels were significantly increased (3-4-fold) as compared to Col-0 (Figs. 1J–1L). All overexpression
lines showed a significant overexpression of the construct (Figs. 1K–1L). Highest expression and protein levels were reached by line OE01 in Col-0 background and by line OE20 in cop1-4 background (Figs. 1K–1M, Fig. S3). Both, expression and protein levels, were comparable for line OE02 (Col-0) and line OE20 (cop1-4). For OE19 and OE21 (cop1-4), in different repeats, YFP-TTG1 levels were either comparable (Fig. 1M) or close to or even below the detection limit (Fig. S3). In Col-0 background, line OE03 exhibited the highest variance of YFP-TTG1 levels compared to other lines in between the repeats. This is in agreement with the observation at the fluorescence stereo microscope using older plants. Here, in several OE03 plants YFP-fluorescence was absent in areas of the leaves or in the center of the rosette. This patchiness of YFP fluorescence was also observed in OE21 (cop1-4) and sometimes in OE01 (Col-0) plants but only in one out of 50 OE02 (Col-0) plants and in none of the OE20 (cop1-4) plants (Table S4). Therefore, and due to the similar transcript and protein level, OE02 (Col-0) and OE20 (cop1-4) were chosen for subsequent quantitative RT-PCR (qRT-PCR) experiments.

Together, we revealed that TTG1 has an effect on flowering time. Subsequently, we used q-RT-PCR experiments for an initial embedding of TTG1 in the transcriptional regulation of the flowering time pathway.

**TTG1 can reduce *FT* and *SOC1* transcript levels**

*TTG1* acts early in cell fate determination and is a pleiotropic regulator of transcription. The known mechanisms of TTG1 molecular activity are at the protein level at which it acts in differing complex composition with transcription factors that act as the direct modulators of transcription (e.g., Zhang & Schrader, 2017). Therefore, overexpression lines are most informative to initially reveal if the TTG1 protein can have an impact on the transcriptional regulation of specific targets within the individual branches of the flowering time regulatory pathway. Moreover, on the one hand, by using the *cop1-4* mutant, we added a sensitized background with a significant modulation in protein composition. On the other hand, COP1 interacts with at least one TTG1-complex component (PAP2), therefore, this background also allows for a conclusion if TTG1 activity requires a functional COP1 protein. Results in this background provide insides if TTG1 at elevated protein levels is able to even overwrite the transcriptional scenario in the light-signaling and LD flowering time mutant *cop1-4*. This would underline even more than in the wild-type scenario the potential adaptive value of TTG1 and relevance as a valuable target for flowering time modulation in various environmental settings. All selected targets were analyzed with the most suitable overexpression line as characterized and described above. In addition, seed material of the two EMS mutants *ttg1-9* and *ttg1-11* was available.

First, we assessed the circadian expression profile of endogenous and overexpressed *TTG1* in OE02. As for all circadian qRT-PCR experiments in this study, we used 8-day-old LD grown seedlings harvested first at ZT0 and thereafter in intervals of 4 h with the last sample at ZT20. A similar overexpression level of *TTG1* was seen throughout the day. The endogenous expression was not affected (see Fig. S4, Table S5).

Due to the strong late flowering phenotype of the overexpression lines, we expected that the transcript levels of the floral integrators (e.g., Blumel, Dally & Jung, 2015) differed from...
the respective backgrounds. Towards this end, we continued with analyzing the CO-FT module’s and SOC1 transcript levels. In line with the flowering time phenotype, for CO, an activator of FT (Suarez-Lopez et al., 2001), a slight tendency to lower transcript levels in the overexpression lines was observed. Nevertheless, this was neither significant nor sufficient to explain the strong phenotype especially in the overexpression in wild-type background (Fig. 2, Table S5). GI is an FT regulator that can increase CO transcript levels but can also activate FT in a CO-independent way. It can directly bind to the FT promoter and interacts with FT suppressors (Sawa & Kay, 2011; Sawa et al., 2007). Also, GI transcript levels in OE02 were not significantly changed (Fig. S5). In both overexpression lines, for FT, the transcript levels almost dropped to the detection limit and exhibited a general reduction at all timepoints which was significant at ZT12 and 16 in OE02. The overexpressors’ SOC1 transcript profiles were very similar in their circadian pattern and both exhibited reductions throughout the day, which was significant at ZT0, 4, 8 and 16 for OE20. In the mutant scenario, we expected to find more subtle effects. We found only a trend to elevated SOC1 levels in ttg1-9 mutants which might explain the early flowering time phenotype but was not significant. In summary, we found that TTG1 can reduce FT and SOC1 transcript levels.

**Early and late effects on transcript levels of AP2-domain containing factors in ttg1 mutants**

We moved our focus on factors that can suppress FT transcript levels, the AP2 domain containing factors of the flowering time regulatory pathway: TEM1, TEM2 (RAV transcription factors with AP2/ERF and B3 DNA-binding domain) and AP2, SMZ, SNZ, TOE1-3 etc., (Song, Ito & Imaizumi, 2013; Wang, 2014). In addition, we tested the transcript level of SVP which acts as an activator of the aforementioned AP2-like factors and as a suppressor of FT (Lee et al., 2007; Tao et al., 2012). Only at night, a significant increase of the TEM2 transcript level was observed for OE02 (Fig. 3, Table S5). No significant change was found for OE20 suggesting that elevated TTG1 levels do not have a strong impact on these genes. However, in the mutants’ case a trend for reduced transcript levels was observed for TEM2 (ttg1-9), AP2, TOE1 and TOE3 (all ttg1-11) and a slight reduction for SNZ (ttg1-11, at night also for ttg1-9) and SMZ (both mutants). Significance analysis ($P < 0.05$) supported the reduction of TOE1 transcript levels at ZT4 and ZT16 for ttg1-11 and SNZ transcript levels at ZT20 for ttg1-9. Interestingly, although not supported by the significance analysis, the trend of reduced transcript levels for SVP seemed to be opposed by elevated SVP levels at ZT0 and ZT20 for OE02 pointing to a possibly flattened circadian amplitude of SVP transcript levels upon TTG1 overexpression. Therefore, we had a closer look on circadian clock components for OE02.

**TTG1 can regulate circadian clock components**

LWD1 and LWD2, the closest homologs of TTG1, regulate flowering through transcriptional modulations within the circadian clock (Wang et al., 2011; Wu et al., 2016; Wu, Wang & Wu, 2008). LWD1 was shown to bind to the promoter of PRR5, PRR9 and PRR1/TOC1 (Wang et al., 2011). Therefore, we analyzed the transcript levels of the core
TTG1 overexpression suppresses FT and SOC1 transcript levels which cannot be explained by CO transcript levels. (A–J) Eight-day-old seedlings grown under LD conditions (light: ZT0-16, dark: ZT16-0) at 21 °C were harvested in 4h-intervals starting at ZT0. The analyzed genotypes were $ttg1$ mutants in Col-0 background, TTG1 overexpression lines in Col-0 (OE02) and in $cop1-4$ (OE20) background with their respective backgrounds. CO, FT and SOC1 transcript levels were analyzed. Transcript levels are presented relative to the $UBQ10$ transcript levels and normalized with the maximal mean per target within a genotype set. (K–M) Same growth conditions as used for A–J. The seedlings were harvested at ZT = 11 and ZT = 13 from OE01-03 (Pro35s:YFP-TTG1 (Col-0), three independent insertion lines) and OE19-21 (Pro35s:YFP-TTG1 (cop1-4), three independent insertion lines), Col-0 and cop1-4. Data are means from three biological replicates originating from three independent seed batches (A–J) or from one seed batch of parallel grown parental plants (K–M). Error bars are SD. Asterisks indicate significant differences (# $P < 0.1$, *$P < 0.05$) between the mutants (blue: $ttg1$-9, cyan: $ttg1$-11) and their background or the overexpression lines and their respective backgrounds Col-0 (orange) and cop1-4 (magenta). Dashed line: averaged lower threshold in the set of experiments for the respective target ($Ct = 35$) relative to the respective $UBQ10$ levels. Dotted line: average lower threshold + SD. The solid line equals 1. The y-axis is log$_{10}$-scaled. See Table S3 for more details on the underlying data and statistics.

Figure 2  TTG1 overexpression suppresses FT and SOC1 transcript levels which cannot be explained by CO transcript levels. (A–J) Eight-day-old seedlings grown under LD conditions (light: ZT0-16, dark: ZT16-0) at 21 °C were harvested in 4h-intervals starting at ZT0. The analyzed genotypes were $ttg1$ mutants in Col-0 background, TTG1 overexpression lines in Col-0 (OE02) and in $cop1-4$ (OE20) background with their respective backgrounds. CO, FT and SOC1 transcript levels were analyzed. Transcript levels are presented relative to the $UBQ10$ transcript levels and normalized with the maximal mean per target within a genotype set. (K–M) Same growth conditions as used for A–J. The seedlings were harvested at ZT = 11 and ZT = 13 from OE01-03 (Pro35s:YFP-TTG1 (Col-0), three independent insertion lines) and OE19-21 (Pro35s:YFP-TTG1 (cop1-4), three independent insertion lines), Col-0 and cop1-4. Data are means from three biological replicates originating from three independent seed batches (A–J) or from one seed batch of parallel grown parental plants (K–M). Error bars are SD. Asterisks indicate significant differences (# $P < 0.1$, *$P < 0.05$) between the mutants (blue: $ttg1$-9, cyan: $ttg1$-11) and their background or the overexpression lines and their respective backgrounds Col-0 (orange) and cop1-4 (magenta). Dashed line: averaged lower threshold in the set of experiments for the respective target ($Ct = 35$) relative to the respective $UBQ10$ levels. Dotted line: average lower threshold + SD. The solid line equals 1. The y-axis is log$_{10}$-scaled. See Table S3 for more details on the underlying data and statistics.

clock components LHY, CCA1 and TOC1/PRR1 and also those of its feed-back regulators PRR5, PRR7 and PRR9 (Shim, Kubota & Imaizumi, 2017) in the TTG1 overexpression line OE02. Figure 4 shows the results sorted by the time point of the maximal peak in the wild
Figure 3  AP2-domain containing factors: reduction of TEM2, SNZ, TOE1 and TOE3 transcript levels in *ttg1* mutants occurs early and late under long-day conditions. In the mutants and overexpressors, we analyzed the transcript levels of transcriptional FT suppressors: AP2 family genes AP2, SMZ, SNZ, TOE1, TOE2, TOE3, the two RAV factors TEM1 and TEM2 as well as SVP, an activator of AP2 family genes and suppressor of FT. The same samples as in Fig. 2 were used. Data are means from three biological replicates. Error bars are SD. Dotted line: average lower threshold in the set of experiments for the respective target (C<sub>T</sub> = 35) relative to the respective UBQ10 levels + SD. The solid line equals 1. The y-axis is in log<sub>10</sub>-scaled. For details on the genotypes and data presentation please refer to Fig. 2 and for details on the underlying data and statistics to Table S5.
TTG1 can modulate transcript levels of circadian clock components, flattens their circadian amplitude and TTG1 overexpression increases FLC transcript levels. To further explore an additional relevant part of the flowering time regulatory pathway, we used the overexpression line in Col-0 background to analyze eight-day-old seedlings grown under LD conditions (light: ZT0-16, dark: ZT16-0) at 21 °C which were harvested in 4 h-intervals starting at ZT0. (A–F) The analyzed clock components are sorted according to their peak during the day. (G) The selected overexpression line in Col-0 background (OE02) was also employed to further explore another relevant FT-suppressive branch of the flowering time regulatory pathway: FLC transcript levels. All transcript levels are relative to that of UBQ10 and normalized with the maximal mean per target. Data are means from three biological replicates. Error bars are SD. Asterisks indicate significant differences (*P < 0.05) between the overexpression line and the Col-0 wild type. Dashed line: averaged lower threshold in the set of experiments for the respective target (Ct = 35) relative to the respective UBQ10 levels. Dotted line: average lower threshold + SD. The solid line equals 1. The y-axis is in log_{10}-scaled. See Table S5 for more details on the underlying data and statistics.

In general, a flattened circadian amplitude is to be seen despite for TOC1 which is not affected by TTG1 overexpression. For LHY, the minimum seems to be shifted to ZT8 instead of ZT12 in the wild type. A significant increase (P < 0.5) in transcript levels was observed for CCA1 at ZT12 and ZT16, for PRR9 at ZT20, for PRR7 at ZT20 and ZT0 and for PRR5 at ZT0 and ZT4. Thus, we found that TTG1 can regulate the transcript levels of circadian clock components and modulates their transcriptional profiles mainly through flattening the amplitude as a consequence of elevated transcript levels at distinct timepoints on the day under investigation. Similar as its homologs LWD1 and LWD2, TTG1 can change the transcript levels of PRRs but did not modulate TOC1 transcript levels.

The transcript levels of the so far analyzed branches of the flowering time regulatory pathway did not provide a convincing explanation for the strong suppression of FT and SOC1 transcript levels in the overexpression lines. FLC represents and integrates additional branches of this pathway. The FLC protein can bind directly to the promoters of FT.
and SOC1 and suppresses the respective transcript levels (Helliwell et al., 2006; Searle et al., 2006). Therefore, we decided to complete our initial embedding of TTG1 in the transcriptional control of the flowering time pathway by analyzing the transcript levels of FLC. We found that overexpression of TTG1 resulted in elevated FLC transcript levels throughout the day (Fig. 4). These were significantly and more than 2-fold increased as compared to the wild type at ZT8. We conclude that TTG1 can act as an activator of FLC.

One well selected and characterized overexpression line was analyzed in three biological replicates based on three independent seed batches for this initial embedding of TTG1 in the flowering time regulatory pathway. The different and specific time points of transcript modulation spread throughout the day based on the overexpression line OE02 suggest the general ability of TTG1 to elevate transcript levels of clock components and FLC and identify both branches as targets for specific and more detailed follow up studies.

Together, we found that TTG1 acts as a transcriptional regulator in various parts of the flowering time regulatory pathway.

PRR5 recruits TTG1 to subnuclear foci and bHLH92 nuclear enrichment is counteracted by TTG1

In its role as a regulator of early developmental traits, TTG1 acts through differential complex composition, it acts in competitive scenarios, trapping to the nucleus and mutual localization with respective interactors (Balkunde, Bouyer & Hulskamp, 2011; Bouyer et al., 2008; Pesch et al., 2015; Wester et al., 2009; Zhang et al., 2019; Zhang & Schrader, 2017; Zhao et al., 2008). All these scenarios occur at the protein level in dependence of its interactors. bHLH factors like GL3, EGL3, TT8 or MYC1 and R2R3-MYB factors form R2R3MYB-bHLH-WD40 (MBW) complexes with TTG1 to regulate the early TTG1-dependent traits (Balkunde, Pesch & Hulskamp, 2010; Brown, 2005; Koornneef, 1981; Lepiniec et al., 2006; Miller, Chezem & Clay, 2015; Ramsay & Glover, 2005; Tominaga-Wada, Ishida & Wada, 2011; Walker et al., 1999; Zhang et al., 2003). We wondered, which interactors might be relevant for TTG1 function towards flowering time regulation. As the classical bHLH interactors did not modulate flowering time in a similar way as TTG1 (Fig. S6), we conducted a Y2H screening to identify candidates which are related to the flowering time regulatory pathway. Among the results of this screen was one of the classical bHLH interactors, EGL3 (Table S6) (Zhang et al., 2003). In addition, and related to the flowering time regulatory pathway, we identified PRR5 and bHLH92, a bHLH factor shown to be expressed in a circadian pattern (Hanano et al., 2008).

Both selected candidates could be verified as interactors of TTG1 in Y2H experiments in which they were fused to the GAL4-binding as well as to the GAL4-activation domain (Figs. S7A–S7B). We also tested PRR7, PRR9 and TOC1 in the GAL4 system (Figs. S7A–S7B) with different 3-AT (3-amino-1,2,3triazole) concentrations and an adjusted optical density of the samples (Figs. S7A–S7B). All three PRRs interacted with TTG1 when fused to the activation domain of GAL4. TOC1 exhibited the weakest interaction, as indicated by growth of yeast on the respective plates, followed by PRR9, PRR7 and PRR5. A very weak possible interaction of LWD1 and LWD2 was seen with PRR5 in yeast (Figs. S7B–S7C). However, among the homologs, the interaction with all PRRs is TTG1-specific. bHLH92
Figure 5  TTG1 interacts with PRR5 and bHLH92 in yeast with suggested functional relevance due to protein re-localizations in planta.  

(A–B) Yeast two-hybrid assay with TTG1 as a bait (A). TTG1 was tested for interaction with the PRRs (TOC1/PRR1, PRR5, PRR7, PRR9) and bHLH92. GFP serves as a negative control (B). Yeast colonies were transferred to interaction plates (SD-LWH) and plates to test for successful co-transformation (SD-LW). Interaction plates were supplemented with different 3-AT concentrations: 3, 5, 10, 15, 20, 30 mM to qualitatively assess differences in interaction strength. See Fig. S7 for additional results discussed in the text. AD = GAL4-activation domain; BD = GAL4-DNA-binding domain (used for bait constructs); SD = synthetic defined medium. (C–U) Representative sequentially scanned, merged confocal stacks of *N. benthamiana* leaf epidermal cells co-expressing RFP- and YFP- (C–K) or CFP- tagged (L–U) proteins. (C-U) YFP-TTG1 is recruited by RFP-PRR5 to subnuclear foci. Constructs were co-infiltrated to co-express YFP-TTG1 and RFP (C, F, I), YFP and RFP-PRR5 (D, G, J) or YFP-TTG1 and RFP-PRR5 (E, H, K) as also indicated above each column. The indicated channels (left of the respective row) were subsequently merged (I–K) and no brightness-contrast correction was applied. Please note that we did neither differentially adjust the detection nor the pictures for YFP among the combinations to improve visualization of the faint YFP-TTG1 signal in order to visualize the strong effect of RFP-PRR5 on YFP-TTG1 protein localization and abundance within the nucleus (E). (L–U) Enrichment of CFP-bHLH92 in the nucleus is reduced when RFP-TTG1 is co-expressed. Constructs were co-infiltrated to co-express CFP and RFP-TTG1 (M, P, S), CFP-bHLH92 and RFP (N, Q, T) and CFP-bHLH92 and RFP-TTG1 (L, O, R, U) as also indicated above each column. L–O show infiltrated leaf areas of the indicated tagged protein in the respective channel. The same post-acquisition brightness and contrast adjustment was applied to all pictures marked with “adj”. White arrows point to representative nuclei with differing, relative CFP-bHLH92 enrichment as compared to the cytoplasm and in dependence on the presence of RFP-TTG1. All pictures within C-U were acquired with the same settings for RFP/YFP and RFP/CFP, respectively, despite for a reduced laser intensity for the RFP detection of the CFP-bHLH92/RFP combination (T) and a smaller image size for the CFP/RFP-TTG1 combination (P, S) (512 × 512 as compared to 2,058 × 2,058 px²). In Fig. S9 we provide adjusted and non-adjusted pictures for RFP-TTG1 to visualize the protein’s localization. See also Fig. 6 for CFP-PRR/RFP-TTG1 combinations. Bars equal 100 µm in the leaf area pictures and 50 µm in all other pictures. Additional confocal images are shown in Figs. S8–S9. All experiments in the figure were at least conducted three times independently with the same results.

Full-size DOI: 10.7717/peerj.8303/fig-5
did neither interact with LWD1, LWD2 nor with the PRRs in this assay (Figs. S7B–S7C). This suggests that the mechanisms of LWDs and of the TTG1 protein in flowering time regulation including the transcriptional modification of the circadian clock differ.

The TTG1 protein is cell-to-cell mobile (Bouyer et al., 2008). Knowing that nuclear trapping by GL3 is a relevant mechanism for TTG1 function in trichome patterning (Balkunde, Bouyer & Hulskamp, 2011; Bouyer et al., 2008), we analyzed the localization of TTG1 in presence and absence of PRR5 and bHLH92 and vice versa (Figs. 5C–5U, Figs. S8–S9). As reported before, tagged TTG1 is localized in the cytoplasm and in nucleus in epidermal cells of infiltrated tobacco leaves (e.g., Bouyer et al., 2008). PRR5 is only localized to the nucleus where it forms nuclear foci. When co-expressing RFP-tagged TTG1 and YFP-tagged PRR5, RFP-TTG1 localized predominantly in the nucleus where it co-localized with YFP-PRR5 (Figs. 5C–5K, Figs. S8–S9). In case of bHLH92, we obtained a different result. YFP-bHLH92 is enriched in the nucleus but also localizes to the cytoplasm when co-expressed with RFP alone. When co-expressed with RFP-TTG1, TTG1 localization did not change but the nuclear enrichment of bHLH92 did not occur (Figs. 5L–5U, Figs. S8–S9). We repeated the experiment with each of the PRRs (PRR5, PRR7, PRR9 and TOC1/PRR1) being fused to CFP in combination with RFP alone or RFP-TTG1 (Fig. 6, Figs. S8–S9). The PRRs did not only recruit TTG1 to the nucleus, TTG1 also changed the subnuclear localization for PRR9 and PRR7.

DISCUSSION

TTG1 is a pleiotropic regulator of early developmental traits in A. thaliana. Here we show, that this view has to be extended as TTG1 also acts later in the plant’s life cycle as a regulator of flowering time in A. thaliana.

The four used ttgl mutants in Col-0 background flowered earlier and overexpressors in the same background flowered later at long-day conditions than the respective wild type. This is consistent with an observed suppression of FT and SOC1 transcript levels upon TTG1 overexpression. In our study, we followed the hypothesis that TTG1 acts upstream of FT. In this line, if a suppressor of FT is regulated by TTG1, its transcript level should be suppressed in ttgl mutants and increased in the TTG1 overexpressor line. For the mutants and overexpressors, we find a mixed bag of transcript profiles suggesting TTG1 to act in multilayered regulatory mechanisms.

TTG1 variants, protein levels and the background

The Ler wild type exhibited the earliest flowering wild type in our experiment which could impair an early flowering of ttgl-1 relative to its wildtype with respect to time and number of leaves produced. Due to these results, it is not surprising that the flowering phenotype was not reported earlier as the ttgl-1 has been heavily used in previous studies since the 1980s. The late flowering phenotype of ttgl-10 in Ws background might be initially surprising but can be explained by the localization of the mutation within the TTG1 gene. Floral buds did not express the TTG1 transcript in this mutant (Larkin et al., 1999). This might deviate at different developmental stages and tissues as ttgl-10 is an EMS mutant with a point mutation in the TTG1 promoter. The mutation might change the expression
Figure 6  PRRs can re-localize TTG1 to different distinct (sub-)nuclear localizations. Representative, sequentially scanned and merged confocal stacks of *N. benthamiana* leaf epidermal cells. Combinations of RFP-TTG1 or RFP with CFP-PRR and CFP given above each column were co-expressed: CFP/RFP-TTG1 (A,B,G,H,M,N,W,X), CFP-PRR9/RFP (C,I,O), CFP-PRR9/RFP-TTG1 (D,J,P,CC), CFP-PRR7/RFP (E,K,Q), CFP-PRR7/RFP-TTG1 (F,L,R,DD), CFP-PRR5/RFP (S,Y,EE) CFP-PRR5/RFP-TTG1 (T,Z,FF,II), CFP-TOC1/RFP (U,AA,GG) and CFP-TOC1/RFP-TTG1 (V,BB,HH,JJ). The CFP- (A-F, S-V) and RFP-channel (G-L, W-DD, II, JJ) were merged (M-R, EE-HH) and a brightness-contrast correction was only applied for the CFP/RFP-TTG1 combination in B, H and N to visualize RFP-TTG1 presence and localization. Insets show representative observed nuclear localization. Please note that for CFP-PRR7/RFP (and to a minor extend for CFP-TOC1/RFP-TTG1) different subnuclear localizations were seen in all three experiments conducted and that the subnuclear localization of PRR9 and PRR7 was also modified by RFP-TTG1. White arrows for CFP-PRR7/RFP (E,Q) point to week CFP fluorescing nuclei without intense subnuclear foci. These were not dominant but present. For the CFP-TOC1/RFP-TTG1 combination (BB), arrowheads in the left inset of a characteristic nucleus point to faint subnuclear foci into which RFP-TTG1 is recruited. Pictures of all nuclei, including bars for those which are shown as insets, are shown in Fig. S8. Orange box: Non-adjusted pictures of the RFP channel detection for co-infiltrations of RFP-TTG1 with CFP or CFP-tagged PRRs showing an area of the respective leave. See also Fig. S9 for pictures with increased brightness and contrast. Bars equal 50 µm for representative cells and 100 µm for the leaf areas.
pattern of TTG1, which in turn can suppress flowering. Also, a manifested second site mutation cannot be excluded as well as an effect of the Ws background.

Interestingly, the *ttg1-21* mutant, the presumably strongest mutant analyzed in this study, showed the mildest effect on flowering time as compared to the wildtype. At the warm condition, a significant deviation from the wild type could not be concluded for this mutant in terms of time. This suggests that the properties of the TTG1 protein might have a stronger effect on flowering time regulation than the reduced level or absence of TTG1. Presence of the TTG1 protein is not required for the plant to flower but elevated levels and variants can modulate flowering time in *A. thaliana*. Therefore, an overexpression line seems to be more beneficial to initially identify targets and interaction partners for TTG1-dependent regulation within the flowering time regulatory pathway. Thereby, young plant material can be used to provide the basis for a broad embedding of TTG1 in this pathway. The phenotype of the mutants is in agreement with the phenotype of the overexpression lines in the same background. However, TTG1-dependent regulation through protein level and type of protein variant might follow overlapping as well as differing regulatory mechanisms which might be dependent on the respective background and variant-specific interaction patterns to be dissected in the future.

**The developmental stage and age pathway**

Our results acquired along the chronological and developmental axis point to an involvement of TTG1 in modulating the plastochron. Detailed meristem analysis at and around the time point of flowering are required in the future. The developmental stage and a possible tissue-specificity might also explain the *FT* and *SOC1* transcript levels which were not significantly increased in *ttg1* mutants. As TTG1 is a factor required for cell fate determination (*Galway et al., 1994*), developmental stage, tissue and cell specific effects might occur. Therefore, older plants than the used seedlings and the analysis of tissue specific expression might be required in a more detailed future analysis. This would be of particular relevance for the age pathway and GA signaling which were not covered in this study.

Overlaps with the age pathway might occur at the level of the *SQUAMOSA BINDING PROMOTER BINDING PROTEIN-LIKE* (SPL)s which are suppressed by microRNA156. SPLs are involved in regulating trichome density at later stages e.g., at the stem (*Yu et al., 2010*), SPL9 activates *TRIPTYCHON* and *TRICHOMELESS1* and is thought to modulate trichome density thereby (*Yu et al., 2010*). Both R3-MYBs belong to the so-called inhibitors which compete in the MBW complex scenarios with R2R3-MYBs for bHLH factor binding (*Balkunde, Pesch & Hulskamp, 2010; Esch et al., 2003; Wang et al., 2008; Wester et al., 2009*). TTG1 itself was reported to interact with SPL4 and SPL5 in yeast (*Ioannidi et al., 2016*) and the mutant of another inhibitor, *enhancer of try and cpc 3* (*etc3*), exhibits a differential *FT* and *SOC1* regulation in 21d-old LD-grown plants as compared to the wild type which equals the time of its early flowering time phenotype (*Wada & Tominaga-Wada, 2015*).

**WERWOLF (WER) and FT mRNA stability**

If TTG1 acts at the *FT* gene, TTG1 can either act on the promoter or on other regulatory regions of *FT* (and *SOC1*) or affect the *FT* mRNA stability as reported for WER, a TTG1
network component (Seo et al., 2011). Seo and co-workers found that the mutant of WER flowers late and the R2R3-MYB factor WER is required for FT mRNA stability. The role of WER towards flowering time regulation was found to be independent of CO and FLC. Together with our results, the role of TTG1 and WER would be opposing which is not in line with a joined regulation following the classical MBW complexes regulatory mechanisms. However, we can not exclude that FT mRNA stability is changed in dependence of TTG1 complexes. By forming an MBW complex with WER, TTG1 could prevent WER from its function towards FT mRNA stability which would add to the late flowering time phenotype observed in TTG1 overexpressors.

**AP2-domain containing factors and the GA signaling branch**

Several AP2-domain containing factors exhibit the expected tendencies to reduced transcript levels in ttg1 mutants. However, the overexpressor lines did not show the respective opposing effect. Hence, an intact TTG1 seems to be required for normal circadian transcript profiles of these AP2-domain containing factors.

At the transcriptional level, TEM1 and TEM2 act as repressors of the TTG1-MBW complex components GL1, an R2R3-MYB factor, and the bHLH factors GL3 and EGL3 while TTG1 transcript levels are not affected (Matias-Hernandez et al., 2016). On the one hand, TEM1 and TEM2 act in a cell type-dependent manner (Matiaz-Hernandez, 2016). Therefore, an effect of deviating cell fate and differentiation in the ttg1 mutants can have additional effects. On the other hand, TEMs control GA accumulation and distribution in the leaf mesophyll. They also integrate the photoperiod and GA signaling pathway in LD and SD conditions (Castillejo & Pelaz, 2008; Matiaz-Hernandez, 2016; Osnato et al., 2012). At the molecular level, reduced TEM2 transcript levels might circumvent FT and SOCI transcript levels and cause an early flowering of ttg1-9. Reduced TEM2 levels lead to elevated GA levels which promote flowering time. The bHLH factors GL3, EGL3 and the R2R3-MYB factor GL1, can interact with the DELLA proteins RGA (REPRESSOR OF GA REQUIRING 1) 1 and RGA2 which both repress the transcripational activation properties of the MBW complex. This suppression is derepressed by GA through GA-induced degradation of the DELLA proteins (Qi et al., 2014). Therefore, different binding properties of the mutant protein variants might cause additional regulatory loops through the GA signaling pathway to play a role.

**Competitive scenarios modulating CO protein levels**

AP2-domain containing factors can bind directly to the FT promoter (Castillejo & Pelaz, 2008; Mathieu et al., 2009; Zhang et al., 2015). In addition, at the protein level, TOEs can interact with CO and thereby prevent CO from activating FT transcription (Zhang et al., 2015). In contrast, PRRs can stabilize the CO protein (Hayama et al., 2017). Although a reduction of CO transcript levels upon TTG1 overexpression at ZT12 is observed with similarities to the patterns of PRR overexpressors (Hayama et al., 2017), a leading role of CO in the TTG1-dependent regulation of FT transcript levels cannot be concluded based on the CO transcript levels. Nevertheless, at the protein level, TTG1 might either decrease CO protein levels or inactivate the CO protein and, thereby, reduce a CO-mediated FT
activation. The interaction of TTG1 with PRRs and re-localization of PRRs, as suggested by our results, could have such an effect.

CO protein levels are elevated in cop1-4 background (Jang et al., 2008). Overexpression of TTG1 can also delay flowering time in cop1-4 background. This indicates that either the TTG1 protein levels were sufficient to counteract the CO protein function at the protein level or that the TTG1 function can be or is mainly independent of CO. However, the effects on flowering time are difficult to compare among wild type and cop1-4 background as we found increased TTG1 transcript levels in cop1-4.

**LWDs, TTG1 and the clock**

Further upstream in the photoperiodic pathway, LWD proteins act as activators within the loops of the circadian clock (Shim, Kubota & Imaizumi, 2017). It is conceivable that a partial overlap in function exists as these are the two closest homologs of TTG1 in A. thaliana but also differences are expected. Towards this end, a detailed evolutionarily focused analysis will be of interest.

While LWD1 transcript levels show a strong circadian response with highest levels late at night and in the long-day morning, LWD2 and, in our results, TTG1 do not show this pattern and remain at a similar level during the day and night (Wu, Wang & Wu, 2008). Interestingly, promoter-luciferase constructs showed rhythmic activity of both LWD promoters (Wang et al., 2011). With a focus on LWD1, its binding in a time dependent manner at the promoters of PRR5, PRR7, PRR9 and CCA1 was revealed (Wang et al., 2011). In lwd1lwd2 double mutants, CCA1 and LHY transcript levels are reduced, the period is shortened and shifted forward (Wu, Wang & Wu, 2008). In the late afternoon to early morning at LD condition, TTG1 overexpression increases the transcript levels of PRR5, PRR7, PRR9, CCA1 and LHY and potentially reduces the respective transcriptional amplitudes.

CCA1 expression can be activated by LWD1 which acts together with interacting TCP transcription factors (Wang et al., 2011; Wu et al., 2016). TTG1 might act similarly but, also, a mechanism as suggested for CO is possible for CCA1 and LHY. Through binding of TTG1 to PRRs, these might no longer be able to form a complex with the transcriptional repressors TOPOLESS/TOPOLESS RELATED PROTEINS at the CCA1 and LHY promoters (Wang, Kim & Somers, 2013). Consequently, CCA1 and LHY transcript levels would increase and the circadian period would be lengthened (Wang, Kim & Somers, 2013). We observed respective transcript level characteristics upon TTG1 overexpression apart of the period which was not tested.

Differential complex composition of TTG1 with interacting transcriptional activators and repressors might act in comparable regulatory mechanisms mediating target specific DNA-interaction. PRRs and bHLH92 described in this study are excellent candidates. As we did not find strong indications for a clear interaction with LWDs, mechanisms related to PRRs and bHLH92 at the protein level are likely to differ between the TTG1 and the LWDs. Being a bHLH factor, bHLH92 might fit well into the regulatory scheme known for TTG1. This would require the identification of an R2R3-MYB interacting with bHLH92.
TTG1 might act through TTG1-PRR and TTG1-FLC-FT modules (Fig. 7, Fig. S10)

The subnuclear localization patterns upon co-expression of PRRs and bHLH92 with TTG1 can provide a regulatory mechanism. Does this have an influence on transcriptional activation? What is the identity of the subnuclear foci? Are the interaction partners binding in concert to specific loci, stabilize each other, are they stored or deactivated within these foci or are competitive scenarios at work? These are pressing questions to be answered.

Through such mechanisms, known trait networks of the clock, like those of the PRRs, can be linked with the TTG1 trait network. For PRR5, target promoters were identified which comprise different transcription factors involved e.g., in auxin production, hypocotyl growth and cold-stress response. This might be intermingled with traits and temperature responsiveness in TTG1-dependence. TTG1 might suppress PRR target modulation on relevant downstream targets of both factors at a respective developmental stage. FT might be such a target as it was shown that a PRR-CDF-FT module can exists. PRR5 can directly suppress CDF expression (Nakamichi et al., 2012) and CDFs can suppress FT at its promoter (Song et al., 2012). A TTG1-PRR-CDF-FT module could bypass GI and CO and could link TTG1 effects on clock gene modulation with FT suppression in a competitive scenario with PRR-CDF-FT modules.

Our results suggest a possible, parallel TTG1-FLC-FT module. Elevated FLC transcript levels lead to an increase in FLC-mediated FT and SOC1 suppression and consequently to late flowering in the overexpression line. In line with this role of TTG1, the weak allele of FLC in Ler background could explain the absence of a flowering time phenotype in the strong and heavily used ttg1-1 mutant. Therefore, when analyzing this mutant, a role of
TTG1 towards flowering time regulation was not detectable before. All used accessions in this study are rapid cycling accessions lacking a functional FRI allele and therefore immediately exposing modulations at \( FLC \) to potential phenotypic detection.

**Overlapping regulatory network**

The annual plant \( A.\ thaliana \) completes its life cycle with the production and ripening of seeds and enters the new life cycle following seed dormancy with seed germination. The reproductive success depends therefore on the appropriate timing of flowering and seed ripening as well as germination thereafter. Therefore, it is not surprising that a pleiotropic regulator like TTG1 which is strongly involved in the regulation of various relevant seed traits, is also involved in the regulation of flowering time. Here, it is noteworthy that TTG1-dependent gene regulatory network components including TTG1 have the potential to intervene in several sub-pathways of flowering time regulation. We found that TTG1 can even overwrite the transcriptional scenario in \( \textit{cop1-4} \) in regard to the floral integrators \( \textit{FT} \) and \( \textit{SOC1} \). Moreover, TTG1 variants are likely to be of relevance in adaptation to temperature seasonality, minimum temperature and daylength (Hancock et al., 2011). This strongly suggests an adaptive value of the TTG1-dependent trait network which is strengthened by the overlapping gene regulatory networks of TTG1 and flowering time regulation as substantiated with this study.

**CONCLUSIONS**

Plants can respond to endogenous and exogenous cues through concerted regulation of specific trait networks. Pleiotropic regulators can aid to reveal such trait networks of adaptive value. The pleiotropic regulator \( \textit{TRANSPARENT TESTA GLABRA 1} \) is known as head of a conserved gene regulatory network regulating early developmental traits. Surprisingly little has been known about its involvement in late developmental trait regulation. We reveal that TTG1 is a flowering time regulator in \( \textit{Arabidopsis thaliana} \) and initially embedded TTG1 in the flowering time regulatory pathway. TTG1 modulates transcript levels of key elements within this pathway—the floral integrators \( \textit{FT} \) and \( \textit{SOC1} \). We show that TTG1 might act upstream of \( \textit{FLC} \) and the circadian clock. At the protein level we found differential interdependencies with regard to the subcellular and subnuclear localization of clock proteins and TTG1 \textit{in planta}. In summary, our results provide an initial embedding of TTG1 in the flowering time regulatory pathway.

**ACKNOWLEDGEMENTS**

The authors thank Martin Hülskamp (University of Cologne) for hosting A.S. We thank Alexander Schurz for laboratory and plant work assistance, Iqra Javed for a tobacco infiltration and support with yeast work, Julian Schiffner for “TTG1 both” qRT-PCR primers, Sabine Lohmer for maintenance of the yeast strain and Bastian Welter for technical assistance while cloning CFPattB1-pBat-TL-p35s. We are grateful to the following persons for providing seeds and plasmids: Martin Hülskamp providing access to his lab stock, Martina Pesch for bHLH vectors used for cloning, George Coupland (Liron Sarid-Krebs/Brigitte Koop) (Max Planck Institute for Plant Breeding Research, Germany) for...
35S::PRR7:CFP and 35S::PRR9:CFP and Martin Sagasser/Bernd Weisshaar (Center for Biotechnology, University of Bielefeld, Germany) for ttg1-21 and ttg1-22. We wish to thank Klaus Menrath (University of Cologne) and his team for perfect plant care and providing tobacco plants. Last but not least we wish to thank for all constructive contributions within the reviewing process.

**ADDITIONAL INFORMATION AND DECLARATIONS**

**Funding**
Martin Hülskamp (University of Cologne) financed Andrea Schrader and this work. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**Grant Disclosures**
The following grant information was disclosed by the authors:
University of Cologne.

**Competing Interests**
The authors declare there are no competing interests.

**Author Contributions**
- Barbara A.M. Paffendorf performed the experiments, analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, approved the final draft.
- Rawan Qassrawi, Andrea M. Meys and Laura Trimborn performed the experiments, prepared figures and/or tables, approved the final draft.
- Andrea Schrader conceived and designed the experiments, performed the experiments, analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.

**Data Availability**
The following information was supplied regarding data availability:
Raw data are available in the Supplemental Files.

**Supplemental Information**
Supplemental information for this article can be found online at http://dx.doi.org/10.7717/peerj.8303#supplemental-information.

**REFERENCES**
Alonso JM, Stepanova AN, Leisse TJ, Kim CJ, Chen H, Shinn P, Stevenson DK, Zimmermann J, Barajas P, Cheuk R, Gadrinab C, Heller C, Jeske A, Koesema E, Meyers CC, Parker H, Prednis L, Ansari Y, Choy N, Deen H, Geralt M, Hazari N, Hom E, Karnes M, Mulholland C, Ndubaku R, Schmidt I, Guzman P, Aguilar-Henonin L, Schmid M, Weigel D, Carter DE, Marchand T, Risseeuw E, Brogden D, Zeko A,
Crosby WL, Berry CC, Ecker JR. 2003. Genome-wide insertional mutagenesis of Arabidopsis thaliana. Science 301:653–657 DOI 10.1126/science.1086391.

An H, Roussot C, Suarez-Lopez P, Corbesier L, Vincent C, Pineiro M, Hepworth S, Mouradov A, Justin S, Turnbull C, Coupland G. 2004. CONSTANS acts in the phloem to regulate a systemic signal that induces photoperiodic flowering of Arabidopsis. Development 131:3615–3626 DOI 10.1242/dev.01231.

Andres F, Coupland G. 2012. The genetic basis of flowering responses to seasonal cues. Nature Reviews Genetics 13:627–639 DOI 10.1038/nrg3291.

Appelhagen I, Jahns O, Bartelniewoehner L, Sagasser M, Weisshaar B, Stracke R. 2011. Leucoanthocyanidin Dioxygenase in Arabidopsis thaliana: characterization of mutant alleles and regulation by MYB-BHLH-TTG1 transcription factor complexes. Gene 484:61–68 DOI 10.1016/j.gene.2011.05.031.

Appelhagen I, Thiedig K, Nordholt N, Schmidt N, Hupel G, Sagasser M, Weisshaar B. 2014. Update on transparent testa mutants from Arabidopsis thaliana: characterisation of new alleles from an isogenic collection. Planta 240:955–970 DOI 10.1007/s00425-014-2088-0.

Balasubramanian S, Sureshkumar S, Lem J, Weigel D. 2006. Potent induction of Arabidopsis thaliana flowering by elevated growth temperature. PLOS Genetics 2:e106 DOI 10.1371/journal.pgen.0020106.

Balkunde R, Bouyer D, Hulskamp M. 2011. Nuclear trapping by GL3 controls intercellular transport and redistribution of TTG1 protein in Arabidopsis. Development 138:5039–5048 DOI 10.1242/dev.072454.

Balkunde R, Pesch M, Hulskamp M. 2010. Trichome patterning in Arabidopsis thaliana from genetic to molecular models. Current Topics in Developmental Biology 91:299–321 DOI 10.1016/S0070-2153(10)91010-7.

Blazquez MA, Ahn JH, Weigel D. 2003. A thermosensory pathway controlling flowering time in Arabidopsis thaliana. Nature Genetics 33:168–171 DOI 10.1038/ng1085.

Blumel M, Dally N, Jung C. 2015. Flowering time regulation in crops—what did we learn from Arabidopsis? Current Opinion in Biotechnology 32:121–129 DOI 10.1016/j.copbio.2014.11.023.

Bouyer D, Geier F, Kragler F, Schnittger A, Pesch M, Wester K, Balkunde R, Timmer J, Fleck C, Hulskamp M. 2008. Two-dimensional patterning by a trapping/depletion mechanism: the role of TTG1 and GL3 in Arabidopsis trichome formation. PLOS Biology 6:e141 DOI 10.1371/journal.pbio.0060141.

Broun P. 2005. Transcriptional control of flavonoid biosynthesis: a complex network of conserved regulators involved in multiple aspects of differentiation in Arabidopsis. Current Opinion in Plant Biology 8:272–279 DOI 10.1016/j.pbi.2005.03.006.

Castillejo C, Pelaz S. 2008. The balance between CONSTANS and TEMPRANILLO activities determines FT expression to trigger flowering. Current Biology 18:1338–1343 DOI 10.1016/j.cub.2008.07.075.

Chang W. 2014. extrafont: tools for using fonts. R package version 0.17. Available at https://CRAN.R-project.org/package=extrafont.
Chen M, Zhang B, Li C, Kulaveerasingam H, Chew FT, Yu H. 2015. TRANSPARENT TESTA GLABRA1 regulates the accumulation of seed storage reserves in Arabidopsis. Plant Physiology 169:391–402 DOI 10.1104/pp.15.00943.

Clough SJ, Bent AF. 1998. Floral dip: a simplified method for Agrobacterium-mediated transformation of Arabidopsis thaliana. The Plant Journal 16:735–743 DOI 10.1046/j.1365-313x.1998.00343.x.

Deng X, Qiu Q, He K, Cao X. 2018. The seekers: how epigenetic modifying enzymes find their hidden genomic targets in Arabidopsis. Current Opinion in Plant Biology 45:75–81 DOI 10.1016/j.pbi.2018.05.006.

Dodd AN, Salathia N, Hall A, Kevei E, Toth R, Nagy F, Hibberd JM, Millar AJ, Webb AA. 2005. Plant circadian clocks increase photosynthesis, growth, survival, and competitive advantage. Science 309:630–633 DOI 10.1126/science.1115581.

Esch JJ, Chen M, Sanders M, Hillestad M, Ndkium S, Idelkope B, Neizer J, Marks MD. 2003. A contradictory GLABRA3 allele helps define gene interactions controlling trichome development in Arabidopsis. Development 130:5885–5894 DOI 10.1242/dev.00812.

Failmezger H, Jaegle B, Schrader A, Hulskamp M, Tresch A. 2013. Semi-automated 3D leaf reconstruction and analysis of trichome patterning from light microscopic images. PLOS Computational Biology 9:e1003029 DOI 10.1371/journal.pcbi.1003029.

Feys BJ, Wiermer M, Bhat RA, Moisan LJ, Medina-Escobar N, Neu C, Cabral A, Parker JE. 2005. Arabidopsis SENESCENCE-ASSOCIATED GENE101 stabilizes and signals within an ENHANCED DISEASE SUSCEPTIBILITY1 complex in plant innate immunity. The Plant Cell 17:2601–2613 DOI 10.1105/tpc.105.033910.

Galway ME, Masucci JD, Lloyd AM, Walbot V, Davis RW, Schiefelbein JW. 1994. The TTG gene is required to specify epidermal cell fate and cell patterning in the Arabidopsis root. Developmental Biology 166:740–754 DOI 10.1006/dbio.1994.1352.

Gietz RD, Schiestl RH. 2007. High-efficiency yeast transformation using the LiAc/SS carrier DNA/PEG method. Nature Protocols 2:31–34 DOI 10.1038/nprot.2007.13.

Grigorova B, Mara C, Hollender C, Sijacic P, Chen X, Liu Z. 2011. LEUNIG and SEUSS co-repressors regulate miR172 expression in Arabidopsis flowers. Development 138:2451–2456 DOI 10.1242/dev.058362.

Hanano S, Stracke R, Jakoby M, Merkle T, Domagalska MA, Weisshaar B, Davis SJ. 2008. A systematic survey in Arabidopsis thaliana of transcription factors that modulate circadian parameters. BMC Genomics 9:182 DOI 10.1186/1471-2164-9-182.

Hancock AM, Brachi B, Faure N, Horton MW, Jarymowycz LB, Sperone FG, Toomajian C, Roux F, Bergelson J. 2011. Adaptation to climate across the Arabidopsis thaliana genome. Science 334:83–86 DOI 10.1126/science.1209244.

Harari-Steinberg O, Ohad I, Chamovitz DA. 2001. Dissection of the light signal transduction pathways regulating the two early light-induced protein genes in Arabidopsis. Plant Physiology 127:986–997 DOI 10.1104/pp.010270.
Hayama R, Sarid-Krebs L, Richter R, Fernandez V, Jang S, Coupland G. 2017. PSEUDO RESPONSE REGULATORs stabilize CONSTANS protein to promote flowering in response to day length. EMBO Journal 36:904–918 DOI 10.15252/embj.201693907.

Hellwell CA, Wood CC, Robertson M, James Peacock W, Dennis ES. 2006. The Arabidopsis FLC protein interacts directly in vivo with SOC1 and FT chromatin and is part of a high-molecular-weight protein complex. The Plant Journal 46:183–192 DOI 10.1111/j.1365-313X.2006.02686.

Hepworth J, Dean C. 2015. Flowering Locus C’s lessons: conserved chromatin switches underpinning developmental timing and adaptation. Plant Physiology 168:1237–1245 DOI 10.1104/pp.15.00496.

Huang H, Nusinow DA. 2016. Into the evening: complex interactions in the Arabidopsis Circadian clock. Trends in Genetics 32:674–686 DOI 10.1016/j.tig.2016.08.002.

Ioannidi E, Rigas S, Tsitsekiad D, Daras G, Alatzas A, Makris A, Tanou G, Argiriou A, Alexandrou D, Poethig S, Hatzopoulos P, Kanellis AK. 2016. Trichome patterning control involves TTG1 interaction with SPL transcription factors. Plant Molecular Biology 92:675–687 DOI 10.1007/s11103-016-0538-8.

Jaegle B, Uroic MK, Holtkotte X, Lucas C, Termath AO, Schmalz HG, Bucher M, Hoecker U, Hulskamp M, Schrader A. 2016. A fast and simple LC-MS-based characterization of the flavonoid biosynthesis pathway for few seed(ling)s. BMC Plant Biology 16:190 DOI 10.1186/s12870-016-0880-7.

Jakoby MJ, Falkenhan D, Mader MT, Brininstool G, Wischnitzki E, Platz N, Hudson A, Hulskamp M, Larkin J, Schnittger A. 2008. Transcriptional profiling of mature Arabidopsis trichomes reveals that NOECK encodes the MIXTA-like transcriptional regulator MYB106. Plant Physiology 148:1583–1602 DOI 10.1104/pp.108.126979.

Jang S, Marchal V, Panigrahi KC, Wenkel S, Soppe W, Deng XW, Valverde F, Coupland G. 2008. Arabidopsis COP1 shapes the temporal pattern of CO accumulation conferring a photoperiodic flowering response. EMBO Journal 27:1277–1288 DOI 10.1038/emboj.2008.68.

Kirik V, Schrader A, Uhrig JF, Hulskamp M. 2007. MIDGET unravels functions of the Arabidopsis topoisomerase VI complex in DNA endoreduplication, chromatin condensation, and transcriptional silencing. The Plant Cell 19:3100–3110 DOI 10.1105/tpc.107.054361.

Klopfleisch K, Phan N, Augustin K, Bayne RS, Booker KS, Botella JR, Carpita NC, Carr T, Chen JG, Cooke TR, Frick-Cheng A, Friedman EJ, Fulk B, Hahn MG, Jiang K, Jorda L, Kruppe L, Liu C, Lorek J, McCann MC, Molina A, Moriyma EN, Mukhtar MS, Mudgil Y, Pattathil S, Schwarz J, Seta S, Tan M, Temp U, Trusov Y, Urano D, Welter B, Yang J, Panstruga R, Uhrig JF, Jones AM. 2011. Arabidopsis G-protein interactome reveals connections to cell wall carbohydrates and morphogenesis. Molecular Systems Biology 7:Article 532 DOI 10.1038/msb.2011.66.

Koornneef M. 1981. The complex syndrome of ttg mutants. Arabidopsis Information Service 18:45–51.

Koornneef M, Meinke D. 2010. The development of Arabidopsis as a model plant. The Plant Journal 61:909–921 DOI 10.1111/j.1365-313X.2009.04086.x.
Larkin JC, Oppenheimer DG, Lloyd AM, Paparozzi ET, Marks MD. 1994. Roles of the GLABROUS1 and TRANSPARENT TESTA GLABRA genes in Arabidopsis Trichome development. *The Plant Cell* 6:1065–1076 DOI 10.1105/tpc.6.8.1065.

Larkin JC, Walker JD, Bolognesi-Winfield AC, Gray JC, Walker AR. 1999. Allele-specific interactions between ttg and gl1 during trichome development in *Arabidopsis thaliana*. *Genetics* 151:1591–1604.

Laubinger S, Marchal V, Gourrierec JLe, Wenkel S, Adrian J, Jang S, Kulajta C, Braun H, Coupland G, Hoecker U. 2006. *Arabidopsis* SPA proteins regulate photoperiodic flowering and interact with the floral inducer CONSTANS to regulate its stability. *Development* 133:3213–3222 DOI 10.1242/dev.02481.

Lee J, Lee I. 2010. Regulation and function of SOC1, a flowering pathway integrator. *Journal of Experimental Botany* 61:2247–2254 DOI 10.1093/jxb/erq098.

Lee JH, Yoo SJ, Park SH, Hwang I, Lee JS, Ahn JH. 2007. Role of SVP in the control of flowering time by ambient temperature in *Arabidopsis*. *Genes and Development* 21:397–402 DOI 10.1101/gad.1518407.

Lepiniec L, Debeaujon I, Routaboul JM, Baudry A, Pourcel L, Nesi N, Caboche M. 2006. Genetics and biochemistry of seed flavonoids. *Annual Review of Plant Biology* 57:405–430 DOI 10.1146/annurev.arplant.57.032905.105252.

Li D, Liu C, Shen L, Wu Y, Chen H, Robertson M, Hellwell CA, Ito T, Meyerowitz E, Yu H. 2008. A repressor complex governs the integration of flowering signals in *Arabidopsis*. *Developmental Cell* 15:110–120 DOI 10.1016/j.devcel.2008.05.002.

Li C, Zhang B, Chen B, Ji I, Yu H. 2018. Site-specific phosphorylation of TRANSPARENT TESTA GLABRA1 mediates carbon partitioning in *Arabidopsis* seeds. *The Plant Journal* 74:638–651 DOI 10.1111/tpj.12153.

Locke JC, Kozma-Bognar L, Gould PD, Feher B, Kevei E, Nagy F, Turner MS, Hall A, Millar AJ. 2006. Experimental validation of a predicted feedback loop in the multi-oscillator clock of *Arabidopsis thaliana*. *Molecular Systems Biology* 2:Article 59 DOI 10.1038/msb4100102.

Maier A, Schrader A, Kokkelink L, Falke C, Welter B, Iniesto E, Rubio V, Uhrig JF, Hulskamp M, Hoecker U. 2013. Light and the E3 ubiquitin ligase COP1/SPA control the protein stability of the MYB transcription factors PAP1 and PAP2 involved in anthocyanin accumulation in Arabidopsis. *The Plant Journal* 74:638–651 DOI 10.1111/tpj.12153.

Makino S, Kiba T, Imamura A, Hanaki N, Nakamura A, Suzuki T, Taniguchi M, Ueguchi C, Sugiyama T, Mizuno T. 2000. Genes encoding pseudo-response regulators: insight into His-to-Asp phosphorelay and circadian rhythm in *Arabidopsis thaliana*. *Plant and Cell Physiology* 41:791–803 DOI 10.1093/pcp/41.6.791.
Martel F, Grundemann D, Schomig E. 2002. A simple method for elimination of false positive results in RT-PCR. *Journal of Biochemistry and Molecular Biology* 35:248–250.

Mathieu J, Yant LJ, Murdter F, Kuttner F, Schmid M. 2009. Repression of flowering by the miR172 target SMZ. *PLOS Biology* 7:e1000148 DOI 10.1371/journal.pbio.1000148.

Matias-Hernandez L, Aguilar-Jaramillo AE, Osnato M, Weinstain R, Shani E, Suarez-Lopez P, Pelaz S. 2016. TEMPRANILLO reveals the mesophyll as crucial for epidermal trichome formation. *Plant Physiology* 170:1624–1639 DOI 10.1104/pp.15.01309.

Matsushika A, Makino S, Kojima M, Mizuno T. 2000. Circadian waves of expression of the APRR1/TOC1 family of pseudo-response regulators in *Arabidopsis thaliana*: insight into the plant circadian clock. *Plant and Cell Physiology* 41:1002–1012 DOI 10.1093/pcp/pcd043.

McNellis TW, Von Arnim AG, Araki T, Komeda Y, Misera S, Deng XW. 1994. Genetic and molecular analysis of an allelic series of cop1 mutants suggests functional roles for the multiple protein domains. *The Plant Cell* 6:487–500 DOI 10.1105/tpc.6.4.487.

Michael TP, Salome PA, Yu HJ, Spencer TR, Sharp EL, McPeek MA, Alonso JM, Ecker JR, McClung CR. 2003. Enhanced fitness conferred by naturally occurring variation in the circadian clock. *Science* 302:1049–1053 DOI 10.1126/science.1082971.

Michaels SD, He Y, Scortecci KC, Amasino RM. 2003. Attenuation of FLOWERING LOCUS C activity as a mechanism for the evolution of summer-annual flowering behavior in Arabidopsis. *Proceedings of the National Academy of Sciences of the United States of America* 100:10102–10107 DOI 10.1073/pnas.1531467100.

Miller JC, Chezem WR, Clay NK. 2015. Ternary WD40 repeat-containing protein complexes: evolution, composition and roles in plant immunity. *Frontiers in Plant Science* 6:Article 1108 DOI 10.3389/fpls.2015.01108.

Nakamichi N, Kiba T, Henriques R, Mizuno T, Chua NH, Sakakibara H. 2010. PSEUDO-RESPONSE REGULATORS 9, 7, and 5 are transcriptional repressors in the Arabidopsis circadian clock. *The Plant Cell* 22:594–605 DOI 10.1105/tpc.109.0972892.

Nakamichi N, Kita M, Kamioka M, Suzuki T, Yamashino T, Higashiyama T, Sakakibara H, Mizuno T. 2012. Transcriptional repressor PRR5 directly regulates clock-output pathways. *Proceedings of the National Academy of Sciences of the United States of America* 109:17123–17128 DOI 10.1073/pnas.1205156109.

Nakamichi N, Kita M, Niinuma K, Ito S, Yamashino T, Mizoguchi T, Mizuno T. 2007. Arabidopsis clock-associated pseudo-response regulators PRR9, PRR7 and PRR5 coordinately and positively regulate flowering time through the canonical CONSTANS-dependent photoperiodic pathway. *Plant and Cell Physiology* 48:822–832 DOI 10.1093/pcp/pcm056.

Neff MM, Turk E, Kalishman M. 2002. Web-based primer design for single nucleotide polymorphism analysis. *Trends in Genetics* 18:613–615 DOI 10.1016/S0168-9525(02)02820-2.
Oakenfull RJ, Davis SJ. 2017. Shining a light on the Arabidopsis circadian clock. *Plant, Cell and Environment* **40**:2571–2585 DOI 10.1111/pce.13033.

Osnato M, Castillejo C, Matias-Hernandez I, Pelaz S. 2012. TEMPRANILLO genes link photoperiod and gibberellin pathways to control flowering in Arabidopsis. *Nature Communications* **3**:Article 808 DOI 10.1038/ncomms1810.

Pesch M, Schultheiss I, Digiuni S, Uhrig JF, Hulskamp M. 2013. Mutual control of intracellular localisation of the patterning proteins AtMYC1, GL1 and TRY/CPC in Arabidopsis. *Development* **140**:3456–3467 DOI 10.1242/dev.094698.

Pesch M, Schultheiss I, Klopffleisch K, Uhrig JF, Koegl M, Clemen CS, Simon R, Weidtkamp-Peters S, Hulskamp M. 2015. TRANSPARENT TESTA GLABRA1 and GLABRA1 Compete for Binding to GLABRA3 in Arabidopsis. *Plant Physiology* **168**:584–597 DOI 10.1104/pp.15.00328.

Qi T, Huang H, Wu D, Yan J, Qi Y, Song S, Xie D. 2014. Arabidopsis DELLA and JAZ proteins bind the WD-repeat/bHLH/MYB complex to modulate gibberellin and jasmonate signaling synergy. *The Plant Cell* **26**:1118–1133 DOI 10.1105/tpc.113.121731.

R Core Team. 2017. *R: a language and environment for statistical computing*. Vienna: R Foundation for Statistical Computing.

Ramsay NA, Glover BJ. 2005. MYB-bHLH-WD40 protein complex and the evolution of cellular diversity. *Trends in Plant Science* **10**:63–70 DOI 10.1016/j.tplants.2004.12.011.

Rosso MG, Li Y, Strizhov N, Reiss B, Dekker K, Weisshaar B. 2003. An *Arabidopsis thaliana* T-DNA mutagenized population (GABI-Kat) for flanking sequence tag-based reverse genetics. *Plant Molecular Biology* **53**:247–259 DOI 10.1023/B:PLAN.000009297.37235.4a.

Sawa M, Kay SA. 2011. GIGANTEA directly activates flowering locus T in *Arabidopsis thaliana*. *Proceedings of the National Academy of Sciences of the United States of America* **108**:11698–11703 DOI 10.1073/pnas.1106771108.

Sawa M, Nusinow DA, Kay SA, Imaizumi T. 2007. FKF1 and GIGANTEA complex formation is required for day-length measurement in Arabidopsis. *Science* **318**:261–265 DOI 10.1126/science.1146994.

Schrader A, Welter B, Hulskamp M, Hoecker U, Uhrig JF. 2013. MIDGET connects COP1-dependent development with endoreduplication in *Arabidopsis thaliana*. *The Plant Journal* **75**:67–79 DOI 10.1111/tpj.12199.

Searle I, He Y, Turck F, Vincent C, Fornara F, Krober S, Amasino RA, Coupland G. 2006. The transcription factor FLC confers a flowering response to vernalization by repressing meristem competence and systemic signaling in Arabidopsis. *Genes and Development* **20**:898–912 DOI 10.1101/gad.373506.

Seo E, Yu J, Ryu KH, Lee MM, Lee I. 2011. WEREWOLF, a regulator of root hair pattern formation, controls flowering time through the regulation of FT mRNA stability. *Plant Physiology* **156**:1867–1877 DOI 10.1104/pp.111.176685.
Shim JS, Kubota A, Imaizumi T. 2017. Circadian clock and photoperiodic flowering in Arabidopsis: CONSTANS is a hub for signal integration. *Plant Physiology* **173**:5–15 DOI 10.1104/pp.16.01327.

Shin J, Sanchez-Villarreal A, Davis AM, Du SX, Berendzen KW, Koncz C, Ding Z, Li C, Davis SJ. 2017. The metabolic sensor AKIN10 modulates the Arabidopsis circadian clock in a light-dependent manner. *Plant, Cell and Environment* **40**:997–1008 DOI 10.1111/pce.12903.

Simpson GG, Dean C. 2002. Arabidopsis, the Rosetta stone of flowering time? *Science* **296**:285–289 DOI 10.1126/science.296.5566.285.

Soellick TR, Uhrig JF. 2001. Development of an optimized interaction-mating protocol for large-scale yeast two-hybrid analyses. *Genome Biology* **2**:Article research0052.1.

Song YH, Ito S, Imaizumi T. 2013. Flowering time regulation: photoperiod- and temperature-sensing in leaves. *Trends in Plant Science* **18**:575–583 DOI 10.1016/j.tplants.2013.05.003.

Song YH, Shim JS, Kinmonth-Schultz HA, Imaizumi T. 2015. Photoperiodic flowering: time measurement mechanisms in leaves. *Annual Review of Plant Biology* **66**:441–464 DOI 10.1146/annurev-arplant-043014-115555.

Song YH, Smith RW, To BJ, Millar AJ, Imaizumi T. 2012. FKF1 conveys timing information for CONSTANS stabilization in photoperiodic flowering. *Science* **336**:1045–1049 DOI 10.1126/science.1219644.

Strayer C, Oyama T, Schultz TF, Raman R, Somers DE, Mas P, Panda S, Kreps JA, Kay SA. 2000. Cloning of the Arabidopsis clock gene TOC1, an autoregulatory response regulator homolog. *Science* **289**:768–771 DOI 10.1126/science.289.5480.768.

Suarez-Lopez P, Wheatley K, Robson F, Onouchi H, Valverde F, Coupland G. 2001. CONSTANS mediates between the circadian clock and the control of flowering in Arabidopsis. *Nature* **410**:1116–1120 DOI 10.1038/35074138.

Sun CW, Callis J. 1997. Independent modulation of *Arabidopsis thaliana* polyubiquitin mRNAs in different organs and in response to environmental changes. *The Plant Journal* **11**:1017–1027 DOI 10.1046/j.1365-313X.1997.11051017.x.

Tao Z, Shen L, Liu C, Liu L, Yan Y, Yu H. 2012. Genome-wide identification of SOC1 and SVP targets during the floral transition in Arabidopsis. *The Plant Journal* **70**:549–561 DOI 10.1111/j.1365-313X.2012.04919.x.

Tominaga-Wada R, Ishida T, Wada T. 2011. New insights into the mechanism of development of Arabidopsis root hairs and trichomes. *International Review of Cell and Molecular Biology* **286**:66–106 DOI 10.1016/B978-0-12-385859-7.00002-1.

Turck F, Fornara F, Coupland G. 2008. Regulation and identity of florigen: FLOWERING LOCUS T moves center stage. *Annual Review of Plant Biology* **59**:573–594 DOI 10.1146/annurev.arplant.59.032607.092755.

Valverde F, Mouradov A, Soppe W, Ravenscroft D, Samach A, Coupland G. 2004. Photoreceptor regulation of CONSTANS protein in photoperiodic flowering. *Science* **303**:1003–1006 DOI 10.1126/science.1091761.

Voinnet O, Pinto YM, Baulcombe DC. 1999. Suppression of gene silencing: a general strategy used by diverse DNA and RNA viruses of plants. *Proceedings of the
Wada T, Tominaga-Wada R. 2015. CAPRICE family genes control flowering time through both promoting and repressing CONSTANS and FLOWERING LOCUS T expression. Plant Science 241:260–265 DOI 10.1016/j.plantsci.2015.10.015.

Walker AR, Davison PA, Bolognesi-Winfield AC, James CM, Srinivasan N, Blundell TL, Esch JJ, Marks MD, Gray JC. 1999. The TRANSPARENT TESTA GLABRA1 locus, which regulates trichome differentiation and anthocyanin biosynthesis in Arabidopsis, encodes a WD40 repeat protein. The Plant Cell 11:1337–1350 DOI 10.1105/tpc.11.7.1337.

Wang JW. 2014. Regulation of flowering time by the miR156-mediated age pathway. Journal of Experimental Botany 65:4723–4730 DOI 10.1093/jxb/eru246.

Wang CQ, Guthrie C, Sarma MK, Dehesh K. 2014. BBX19 interacts with CONSTANS to repress FLOWERING LOCUS T transcription, defining a flowering time checkpoint in Arabidopsis. The Plant Cell 26:3589–3602 DOI 10.1105/tpc.11.130252.

Wang S, Hubbard L, Chang Y, Guo J, Schieldbein J, Chen JG. 2014. Comprehensive analysis of single-repeat R3 MYB proteins in epidermal cell patterning and their transcriptional regulation in Arabidopsis. BMC Plant Biology 8:81 DOI 10.1186/1471-2229-8-81.

Wang L, Kim J, Somers DE. 2013. Transcriptional corepressor TPOLESS complexes with pseudoresponse regulator proteins and histone deacetylases to regulate circadian transcription. Proceedings of the National Academy of Sciences of the United States of America 110:761–766 DOI 10.1073/pnas.1215010110.

Wang Y, Wu JF, Nakamichi N, Sakakibara H, Nam HG, Wu SH. 2011. LIGHT-REGULATED WD1 and PSEUDO-RESPONSE REGULATOR9 form a positive feedback regulatory loop in the Arabidopsis circadian clock. The Plant Cell 23:486–498 DOI 10.1105/tpc.110.081661.

Wenden B, Kozma-Bognar L, Edwards KD, Hall AJ, Locke JC, Millar AJ. 2011. Light inputs shape the Arabidopsis circadian system. The Plant Journal 66:480–491 DOI 10.1111/j.1365-313X.2011.04505.x.

Wester K, Digiuni S, Geier F, Timmer J, Fleck C, Hulskamp M. 2009. Functional diversity of R3 single-repeat genes in trichome development. Development 136:1487–1496 DOI 10.1242/dev.021733.

Wickham H. 2011. The split-apply-combine strategy for data analysis. Journal of Statistical Software 40:1–29.

Wickham H. 2016. ggplot2: elegant graphics for data analysis. Springer-Verlag New York. Available at https://ggplot2.tidyverse.org.

Wickham H. 2017. scales: scale functions for visualization. R package version 0.5.0. Available at https://CRAN.R-project.org/package=scales.

Wickham H, Francois R, Henry L, Müller K. 2019. dplyr: a grammar of data manipulation. R package version 0.8.0.1. Available at https://CRAN.R-project.org/package=dplyr.
Wickham H, Henry L. 2018. tidyr: easily tidy data with 'spread()' and 'gather()' functions. R package version 0.8.0. Available at https://CRAN.R-project.org/package=tidyr.

Wu JF, Tsai HL, Joanito I, Wu YC, Chang CW, Li YH, Wang Y, Hong JC, Chu JW, Hsu CP, Wu SH. 2016. LWD-TCP complex activates the morning gene CCA1 in Arabidopsis. Nature Communications 7:Article 13181 DOI 10.1038/ncomms13181.

Wu JF, Wang Y, Wu SH. 2008. Two new clock proteins, LWD1 and LWD2, regulate Arabidopsis photoperiodic flowering. Plant Physiology 148:948–959 DOI 10.1104/pp.108.124917.

Yang Y, Li R, Qi M. 2000. In vivo analysis of plant promoters and transcription factors by agroinfiltration of tobacco leaves. The Plant Journal 22:543–551 DOI 10.1046/j.1365-313x.2000.00760.x.

Yant L, Mathieu J, Schmid M. 2009. Just say no: floral repressors help Arabidopsis bide the time. Current Opinion in Plant Biology 12:580–586 DOI 10.1016/j.pbi.2009.07.006.

Yu N, Cai WJ, Wang S, Shan CM, Wang LJ, Chen XY. 2010. Temporal control of trichome distribution by microRNA156-targeted SPL genes in Arabidopsis thaliana. The Plant Cell 22:2322–2335 DOI 10.1105/tpc.109.072579.

Yu S, Galvao VC, Zhang YC, Horrer D, Zhang TQ, Hao YH, Feng YQ, Wang S, Schmid M, Wang JW. 2012. Gibberellin regulates the Arabidopsis floral transition through miR156-targeted SQUAMOSA promoter binding-like transcription factors. The Plant Cell 24:3320–3332 DOI 10.1105/tpc.112.101014.

Yu JW, Rubio V, Lee NY, Bai S, Lee SY, Kim SS, Liu L, Zhang Y, Irigoyen ML, Sullivan JA, Zhang Y, Lee I, Xie Q, Paek NC, Deng XW. 2008. COP1 and ELF3 control circadian function and photoperiodic flowering by regulating GI stability. Molecular Cell 32:617–630 DOI 10.1016/j.molcel.2008.09.026.

Zhang B, Chopra D, Schrader A, Hülskamp M. 2019. Evolutionary comparison of competitive protein-complex formation of MYB, bHLH, and WDR proteins in plants. Journal of Experimental Botany 70:3197–3209 DOI 10.1093/jxb/erz155.

Zhang F, Gonzalez A, Zhao M, Payne CT, Lloyd A. 2003. A network of redundant bHLH proteins functions in all TTG1-dependent pathways of Arabidopsis. Development 130:4859–4869 DOI 10.1242/dev.00681.

Zhang B, Schrader A. 2017. TRANSPARENT TESTA GLABRA 1-dependent regulation of flavonoid biosynthesis. Plants 6(4):Article 65 DOI 10.3390/plants6040065.

Zhang B, Wang L, Zeng L, Zhang C, Ma H. 2015. Arabidopsis TOE proteins convey a photoperiodic signal to antagonize CONSTANS and regulate flowering time. Genes and Development 29:975–987 DOI 10.1101/gad.251520.114.

Zhao M, Morohashi K, Hatlestad G, Grotewold E, Lloyd A. 2008. The TTG1-bHLH-MYB complex controls trichome cell fate and patterning through direct targeting of regulatory loci. Development 135:1991–1999 DOI 10.1242/dev.016873.

Zou Y, Wang Y, Wang L, Yang L, Wang R, Li X. 2013. miR172b controls the transition to autotrophic development inhibited by ABA in Arabidopsis. PLOS ONE 8:e64770 DOI 10.1371/journal.pone.0064770.