

Genetic inhibition of flowering differs between juvenile and adult *Citrus* trees

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- **Background and Aims** In woody species, the juvenile period maintains the axillary meristems in a vegetative stage, unable to flower, for several years. However, in adult trees, some 1-year-old meristems flower whereas others remain vegetative to ensure a polycarpic growth habit. Both types of trees, therefore, have non-flowering meristems, and we hypothesize that the molecular mechanism regulating flower inhibition in juvenile trees is different from that in adult trees.
- **Methods** In adult *Citrus* trees, the main endogenous factor inhibiting flower induction is the growing fruit. Thus, we studied the expression of the main flowering time, identity and patterning genes of trees with heavy fruit load (not-flowering adult trees) compared to that of 6-month-old trees (not-flowering juvenile trees). Adult trees without fruits (flowering trees) were used as a control. Second, we studied the expression of the same genes in the meristems of 6-month, and 1-, 3-, 5- and 7-year-old juvenile trees compared to 10-year-old flowering trees.
- **Key Results** The axillary meristems of juvenile trees are unable to transcribe flowering time and patterning genes during the period of induction, although they are able to transcribe the *FLOWERING LOCUS T* citrus orthologue (*CiFT2*) in leaves. By contrast, meristems of not-flowering adult trees are able to transcribe the flowering network genes but fail to achieve the transcription threshold required to flower, due to *CiFT2* repression by the fruit. Juvenile meristems progressively achieve gene expression, with age-dependent differences from 6 months to 7 years, *FD-like* and *CsLFY* being the last genes to be expressed.
- **Conclusions** During the juvenile period the mechanism inhibiting flowering is determined in the immature bud, so that it progressively acquires flowering ability at the gene expression level of the flowering time programme, whereas in the adult tree it is determined in the leaf, where repression of *CiFT2* gene expression occurs.

Key words: Alternate bearing, *API*, *Citrus*, *FLC*, flowering, fruit, *FT*, *FD*, juvenility, *LFY*, *TFL1*.

INTRODUCTION

Flowering involves the transition of the meristem from the vegetative to the reproductive stage. In fruit trees, this process is impeded in at least two phases: (1) during the *juvenile period*, in which the meristem is unable to flower for several years even under conditions of floral bud induction (Albani and Coupland, 2010; Sgamma *et al.*, 2014), and (2) during the *adult period*, when fruit inhibits the ability of buds to differentiate into flowers (Martínez-Fuentes *et al.*, 2010). In both cases, the bud meristem remains dormant or grows only vegetatively. However, there is a paradox regarding the age of the bud meristem and its ability to flower. Juvenile trees grow and branch over several years from the apical and lateral meristems, respectively (Davies and Albrigo, 1994), giving rise to meristems one to several years in age that are unable to flower, i.e. in an *adult vegetative stage* (Bäurle and Dean, 2006). In adult trees, the inhibitory effect of fruit on flowering induction only occurs in one season, but subsequently, in the next one, the new emerging 1-year-old meristems are able to flower. Therefore, in juvenile trees none of the meristems are able to flower (regardless of age), whereas in adult trees

a proportion of the current 1-year-old meristems can develop into flowers, while other meristems remain in a resting state to ensure a polycarpic growth habit (Albani and Coupland, 2010).

In the model plant *Arabidopsis thaliana*, five genetic pathways explain the transition of meristems from the vegetative to the reproductive stage. They involve the interaction between a set of well-known key genes, such as the flowering time genes *FLOWERING LOCUS T* (*FT*), *FLOWERING LOCUS D* (*FD*) and *SUPPRESSOR OF THE OVEREXPRESSION OF CONSTANS 1* (*SOC1*), the meristem identity genes *LEAFY* (*LFY*) and *APETALLA1* (*API*), and the floral patterning genes *SEPALLATA* (*SEP*) and *FRUITFUL* (*FUL*) (see reviews by Albani and Coupland, 2010; Blümel *et al.*, 2015). In brief, flowering is mediated by the autonomous or induced up-regulation of *FT* gene expression in the leaf, and the resulting protein moves to the meristem where it modifies expression of the flower-identity genes through interaction with the *FD* transcription factor (Abe *et al.*, 2005). However, there are repressor genes that can inhibit flowering even under exogenous inductive conditions. For instance, *FT* transcription is repressed by *TEMPRANILLO1* (*TEMI*), a gene that regulates juvenility

in plants (Sgamma et al., 2014), or by *FLOWERING LOCUS C* (*FLC*), a gene that plays a central role in repressing flowering in the vernalization pathway (Seo et al., 2009). In the meristem, *TERMINAL FLOWER1* (*TFL1*) forms a heterodimer *TFL1/FD* which represses transcription of *FD* target genes (Sohn et al., 2007), delays flowering, and prevents upregulation of floral identity genes within the shoot apical meristem to maintain shoot indeterminacy (Hanano and Goto, 2011).

In *Citrus*, an evergreen tree species, flowering is triggered in adult trees either by low temperatures during the cold autumn–winter rest period (which varies dramatically from deciduous fruit tree species such as apple, pear, plum and peach) or by water stress, which upregulate expression of the citrus orthologues *CiFT2*, *CsLFY* and *CsAPI* genes (Nishikawa et al., 2007; Chica and Albrigo, 2013), and the orthologues of *SOC1*, *CsSL1* and *CsSL2* (Tan and Swain, 2007). Nevertheless, little is known about the endogenous mechanisms repressing them. The fruits produce an unknown signal(s) that inhibits floral bud induction, even under exogenous inductive conditions, from the time the fruit is close to complete growth late in summer. The process has been observed in *C. sinensis* (sweet orange) (Martínez-Fuentes et al., 2010) and *C. clementina* (mandarin) (Muñoz-Fambuena et al., 2011) under a Mediterranean climate, and in *C. paradisi* (grapefruit) under a tropical climate (Betancourt et al., 2014). The fruit represses expression of the *CiFT2* gene in leaves, and the meristem-identity genes *CsLFY* and *CsAPI* in the buds (Muñoz-Fambuena et al., 2011, 2012; Shalom et al., 2012). In addition, fruit removal induces *CiFT2* and *CsLFY* upregulation in citrus (Shalom et al., 2014) and apple (Haberman et al., 2016), and also *TFL1* downregulation in apple (Haberman et al., 2016).

In juvenile citrus trees, early studies investigated the constitutive expression of the *A. thaliana* *LFY* and *API* genes in transgenic seedlings, reducing flowering time from several years to 12–20 months (Peña et al., 2001). Similarly, the constitutive overexpression of *CiFT2* induced flowering within 3–22 months (Endo et al., 2005). Juvenile wild-type trees are not responsive to low temperatures in terms of *CiFT2* transcription in leaves and stems, and they accumulate a higher level of transcripts of *CsTFL1*, compared to adult flowering trees (Pillitteri et al., 2004; Nishikawa et al., 2007).

A comprehensive study including the expression of flowering time, meristem identity and flower patterning genes in woody crops is missing because it is difficult to obtain non-flowering tree mutants. To our knowledge, experiments conducted to study flowering repression in trees have compared (1) non-flowering seedlings and adult trees that are competent to flower (Pillitteri et al., 2004; Nishikawa et al., 2007; Castillo et al., 2013; Sgamma et al., 2014), and (2) ‘on’ trees (non-flowering phenotype) and ‘off’ trees (flowering phenotype) (Muñoz-Fambuena et al., 2011, 2012; Shalom et al., 2012; Haberman et al., 2016). Thus, we designed a different study which compares the time course of expression of flowering genes in non-flowering seedlings (juvenile trees) and adult non-flowering trees (‘on’ trees) during the period of floral bud induction. We hypothesized that transcription of flowering genes is hampered in the juvenile (immature) meristems, whereas the adult vegetative meristems are able to do so but fail to achieve the threshold level of flowering gene transcription required to flower (Blázquez et al., 1997), due to repression of *CiFT2* expression in the leaf. We

also used flowering adult trees (‘off’ trees) for comparison. The time course of expression was determined from *Citrus* orthologues of *A. thaliana* flowering time genes (*FT*, *FD* and *SOC1*), flowering identity genes (*LFY* and *API*), flower patterning genes (*SEP1*, *SEP3* and *FUL*) and flowering time inhibitors (*TFL1*, *TEM1* and *FLC*). The aim of this research was to study the mechanism which impedes the transition of the meristems from the vegetative to the reproductive stage in non-flowering citrus trees, both juvenile and adult.

MATERIAL AND METHODS

Plant material and tissue collection

In a first experiment, the time course of flowering gene expression during the period of floral bud induction was studied. The experiment involved 10-year-old ‘Moncada’ mandarin trees (adult) and 6-month-old ‘Cleopatra’ mandarin trees (juvenile). ‘Moncada’ is a parthenocarpic hybrid mandarin [‘Oroval’ (*Citrus clementina*) × ‘Kara’ (*C. unshiu* × *C. nobilis*)] that is known for its strictly biennial bearing (Muñoz-Fambuena et al., 2011). Adult trees flower profusely in spring and set a heavy fruit yield (‘on’ trees), do not flower in the following spring and develop only vegetatively (‘off’ trees). Five trees in each condition (‘on’ and ‘off’ trees) were selected for the experiments. In this study, ‘on’ trees are termed adult non-flowering trees (A-NFL) whereas ‘off’ trees are called adult flowering trees (A-FL). Trees were grafted onto ‘Carrizo’ citrange (*Citrus sinensis* × *Poncirus trifoliata*) rootstock, planted 5 × 5 m apart in a loamy clay soil, with drip irrigation in the IVIA Research Station (Moncada, Spain). ‘Cleopatra’ mandarin (*Citrus reshni*) is a self-compatible species that flowers profusely in the adult stage and produces seedy fruits. Fifty seeds were germinated indoors at 22 °C, and the 6-month-old potted plants (juvenile trees) (J-NFL) were transferred to the field in the IVIA Research Station under the same conditions as the adult trees.

Leaf and bud samples for study of flowering gene expression were collected from September to February. Note that although plants had different ages, all the buds studied in this experiment were 6–7 months old, i.e. buds produced in spring (April) were sampled in autumn (September). Five samples per date and plant were taken. In juvenile trees, all the buds (apical and axillary) and leaves of the trees were sampled. Samples were immediately ground and stored at –80 °C until analysis. Bud sprouting and flowering were evaluated in spring.

A second experiment to study the transition of the meristem from juvenile to adult stage was conducted using seedlings, juvenile and adult trees of ‘Carrizo’ citrange, which show a 7-year juvenile phase, on average (Spiegel-Roy and Goldschmidt, 1996). Six-month-old seedlings and 1-, 3-, 5-, 7- and 10-year-old trees were used in the experiment. Six biological replicates per tree were used. A sufficient quantity of buds were sampled in February, just before bud differentiation, for the gene expression analysis. To study the influence of age on the sensitivity to chilling as an inducer of flowering, six trees of each age were forced to flower by placing them, on November 10, in a culture chamber with controlled temperatures (15 °C/5 °C day/night), photoperiod (8 h/16 h day/night) and relative humidity (90 %). At 0, 15, 30 and 45 d of cold treatment, leaves from each tree

were sampled for gene expression analysis. In all cases, samples were immediately ground and stored at -80°C until analysis. Bud sprouting and flowering were recorded in spring.

RNA extraction and RT-PCR

Total RNA was isolated from frozen tissue using the RNeasy Plant Mini Kit (Qiagen, Hilden, Germany). RNA samples were treated with RNase-free DNase (Qiagen) through column purification following the manufacturer's instructions. RNA quality was tested based on the $\text{OD}_{260}/\text{OD}_{280}$ ratio and gel electrophoresis. RNA concentration was determined by fluorometric assays with the RiboGreen dye (Molecular Probes, Eugene, OR, USA) according to the manufacturer's instructions. Three fluorometric assays per RNA sample were performed. Quantitative real-time RT-PCR was performed with a LightCycler 2.0 Instrument (Roche Diagnostic, Basel, Switzerland) equipped with LightCycler Software version 4.0. One-step RT-PCR was carried out. Reactions contained 2.5 U of MultiScribe Reverse Transcriptase (Applied Biosystems, Carlsbad, CA, USA), 1 U of RNase Inhibitor (Applied Biosystems), 2 μL LC FastStart DNA MasterPLUS SYBR Green I (Roche Diagnostic), 25 ng total RNA and 250 nM of the specific forward and reverse primers of each gene in a total volume of 10 μL . Incubations were conducted at 48°C for 30 min, 95°C for 10 min followed by 45 cycles at 95°C for 2 s, 58°C for 8 s and 72°C for 8 s. Fluorescence intensity data were acquired during the 72°C extension step and transformed into relative mRNA values using a 10-fold dilution series of RNA sample as the standard curve. Relative mRNA levels were then normalized to total mRNA amounts and, in each case, an expression value of 1 was arbitrarily assigned to the sample which showed the lowest Ct value. *ACTIN* was used as the reference gene according to Mafra et al. (2012). Specificity of the amplification reactions was assessed by post-amplification dissociation curves and by sequencing the reaction product. Putative genes were identified through a homology search with related genes from an EST database of entry available in Phytozome v.12.1 (<https://phytozome.jgi.doe.gov>), the species selected being *C. clementina*. Synthetic oligonucleotides were designed to amplify the gene of the selected clones and, as stated before, sequenced for confirmation. Details about the forward and reverse primers are given in Supplementary Data Table S1. All the genes analysed were described in other studies (see references in Table S1) except the putative candidates for *FLC*, *FD* and *TEM1*. In the *C. clementina* genome, the sequence Ciclev10033420m (*FLC-like*) is quite similar to the *FLC* and *PEP1* genes from *A. thaliana*

and *Arabidopsis thaliana*, respectively, and was recently described by Hou et al. (2014). The sequence Ciclev10031846m (*TEM1-like*) shows high similarity to the *TEM1* gene from *A. thaliana* and, also, *Olea europaea* (olive tree), which was described by Sgamma et al. (2014). We also studied expression of the putative homologue to the *FD* gene in the *C. clementina* genome. The sequence Ciclev10003845m (*FD-like*) coded for a b-Zip transcription factor and showed similarity to the *FD* and *VEG2* (*FD* orthologue) genes from *A. thaliana* and *Pisum sativum*, respectively (Table 1).

Statistical analyses

Gene expression was statistically tested by analysis of variance (ANOVA), using the Student–Newman–Keuls test for separation of means. StatGraphics Plus software for Windows, version 5.1 (Statistical Graphics, Englewood Cliffs, NJ, USA), was used.

RESULTS

Tree flowering behaviour

In *Citrus*, adult trees with heavy fruit load in the previous year showed extremely low flowering in the following spring (0.2 flowers per 100 nodes, on average) (adult-not-flowering trees, A-NFL), whereas those without fruits flowered profusely (149 flowers per 100 nodes, on average) (adult flowering trees, A-FL). Juvenile trees (J-NFL) did not produce any flowers.

Time course of flowering gene expression during the period of floral bud induction

The time course of expression of *CiFT2* in the leaves of A-FL trees increased continuously in Spain's mild winter climate from September to February (Fig. 1A). At the time of floral bud induction (mid-November), coinciding with a decrease in average temperature (up to 12°C), relative expression *CiFT2* was significantly upregulated ($\times 1700$) and increased until the end of February ($\times 5000$). However, in the leaves of J-NFL trees, *CiFT2* expression was significantly lower in November ($\times 400$) compared to in A-FL trees (Fig. 1A and B). Finally, in the leaves of A-NFL trees, *CiFT2* expression was completely abolished during the period of floral bud induction (November), increasing ($\times 250$) in January only after fruit harvest (Fig. 1B).

TABLE 1. Sequence analysis of the putative *FD*, *FLC* and *TEM* homologue genes from *Citrus clementina*

Annotation	EST code	Homologous locus	BLASTP (E-value)	Query coverage (%)
<i>FD-like</i>	Ciclev10003845m	<i>FD</i> (<i>Arabidopsis lyrata</i>)	2×10^{-20}	92
		<i>VEG2</i> (<i>Pisum sativum</i>)	1×10^{-20}	92
<i>FLC-like</i>	Ciclev10033420m	<i>FLC-like</i> (<i>Citrus sinensis</i>)	4×10^{-93}	99
		K-box region MADS-box transcription factor family (<i>Arabidopsis thaliana</i>)	2×10^{-7}	83
<i>TEM1-like</i>	Ciclev10031846m	<i>RAV1</i> (<i>A. thaliana</i>)	2×10^{-150}	98
		<i>TEM1</i> (<i>Olea europaea</i>)	0.0	98

We also studied the expression of putative homologues to the *FLC* and *TEM1* genes, which are *FT* inhibitors. Expression of the *FLC-like* gene from *C. clementina* was significantly higher in the leaves of A-NFL (16.5) trees compared to the A-FL (3.8) and J-NFL (4.2) trees during the period of floral bud induction (November) (Fig. 1C). The *TEM1-like* gene showed significantly higher expression from September to November in the leaves of NFL trees (both juvenile and adult) compared to A-FL trees (Fig. 1D).

Transcription of flowering time, identity or patterning genes in the buds was strongly affected by the age of the plant. Thus, for most of the studied genes, no transcription was detected in the meristems of juvenile trees (Fig. 2A–D, H, I). *AtSOC1* orthologues, *CsSL1* and *CsSL2*, were upregulated in the meristems of adult trees, whereas they showed no transcription in those of juvenile trees. *CsSL1* was significantly upregulated ($\times 12$) in adult trees from September to February, irrespective of the presence of fruit, and *CsSL2* peaked in mid-November, much higher than for A-NLF trees ($\times 5$) (Fig. 2A, B). *FD-like* expression was also strongly affected by the age of the plant, as no transcription was detected in the meristems of juvenile trees. However, it was slightly downregulated from September to mid-November and upregulated from mid-November to February in both A-FL and A-NFL trees (Fig. 2C).

In the bud, relative expression of the meristem-identity citrus genes *CuFUL*, *CsAPI* and *CsLFY* differed significantly between flowering and not-flowering trees. As expected, *CsAPI* and *CsLFY* were upregulated from September to February in A-FL trees, with a significantly higher transcription level in February. The expression of these genes in A-NFL and J-NFL trees did not differ significantly; both had significantly lower expression in February compared to A-FL trees (Fig. 2E, F). A behaviour similar to *CsLFY* was observed for *CuFUL*, but surprisingly, it did not differ significantly between A-FL and A-NFL trees. Transcription of this gene in J-NFL trees was negligible (Fig. 2D).

Expression of the flowering inhibitor *CsTFL1* gene in November was significantly higher in the meristem of not-flowering trees than in that of A-FL trees (Fig. 2G). *CsTFL1*

showed the highest expression in juvenile buds in September (60 % higher than in adult trees), and diminished progressively, becoming not significantly different from that of adult trees, both A-FL and A-NFL, in February.

Finally, expression of the flower-patterning genes *CiSEP1* and *CiSEP3* was also strongly influenced by the age of the tree and by flowering ability. Both genes decreased their expression significantly from September to mid-November, followed by a significant upregulation ($\times 27$ and $\times 130$, respectively) from November to February in the A-FL meristem. Surprisingly, *CiSEP1* was also upregulated ($\times 15$) in the A-NFL meristem. By contrast, *CiSEP1* and *CiSEP3* transcription was hampered in the juvenile meristems (Fig. 2H, I).

Relationships between tree age and flowering gene relative expression

To determine when a juvenile tree transcribes flowering time, identity and patterning genes due to chilling, the relative expression of *CiFT2* was studied in leaves during the period of floral bud induction (November), and that of *FD-like*, *CsSL1*, *CuFUL*, *CsLFY*, *CsAPI*, *CiSEP1* and *CsTFL1* in buds just before the flower bud differentiation stage (February), in trees of different ages.

In our experiments, *CiFT2* gene expression increased over time, the magnitude of the expression depending on the tree age. Thus, for 6-month-old seedlings expression was upregulated five-fold at 30 d (480 chilling hours) of cold treatment (15 °C/5 °C, 8 h day/16 h night), and eight-fold at 45 d (720 chilling hours), whereas for 1-year-old and 3-year-old trees it upregulated to higher levels ($\times 16$ and $\times 36$, respectively), the latter being similar to those for 10-year-old trees (Fig. 3). Therefore, *CiFT2* gene expression in leaves increased with the age of J-NFL trees up to the 3 years old, with expression being similar to that of 10-year-old A-FL trees.

In the buds, a direct and significant relationship between tree age and floral bud induction gene expression was also found.

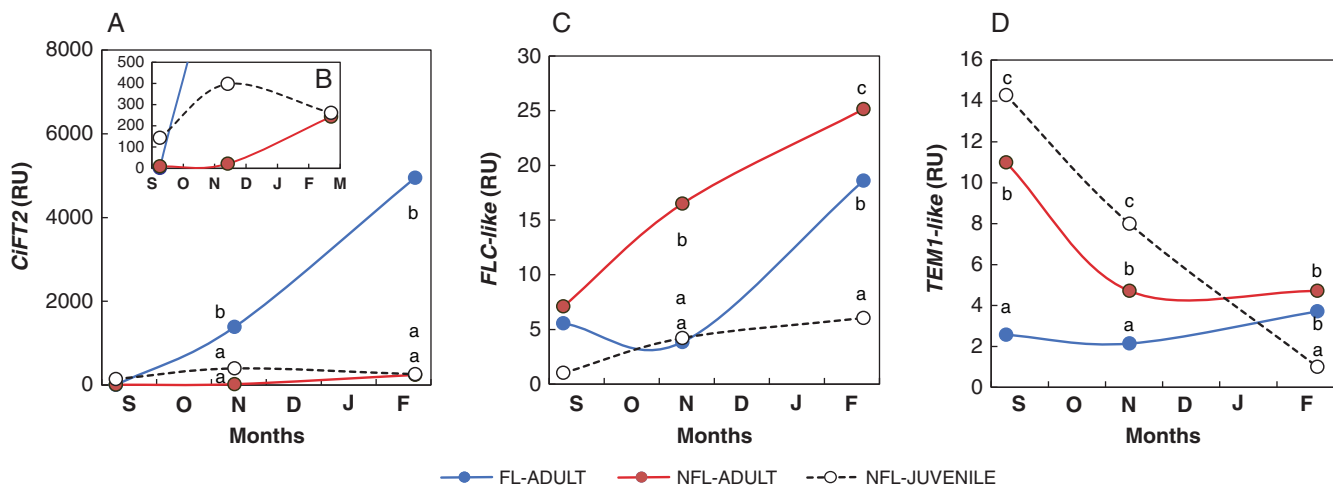


FIG. 1. Time course of *CiFT2* (A, B), *FLC-like* (C) and *TEM1-like* (D) gene expression in the leaves of juvenile (NFL-juvenile) and adult mandarin trees with heavy fruit load (on-trees, NFL-Adult) or without fruit (off-trees, FL-Adult). Data are mean of three independent replicates. In all cases, bars of s.e. are smaller than the symbol size. Different letters for a given month indicate significant differences ($P < 0.05$). RU, relative units.

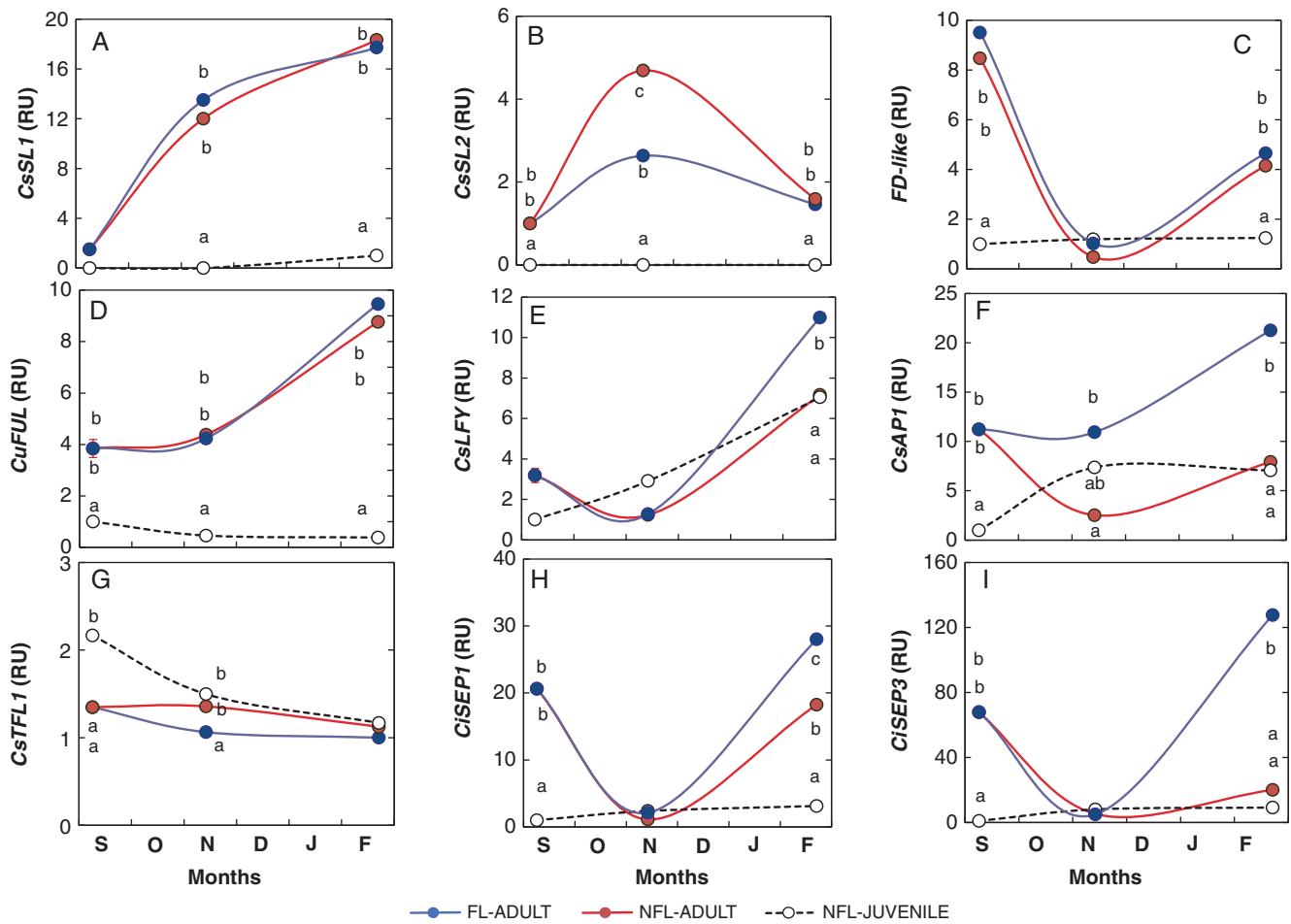


FIG. 2. Time course of *CsSL1* (A), *CsSL2* (B), *FD-like* (C), *CuFUL* (D), *CsLFY* (E), *CsAPI* (F), *CsTFL1* (G), *CiSEP1* (H) and *CiSEP3* (I) gene expression in the buds of juvenile (NFL-juvenile) and adult trees with heavy fruit load (on-trees, NFL-Adult) or without fruit (off-trees, FL-Adult). Data are mean of three independent replicates. In all cases, bars of s.e. are smaller than the symbol size. Different letters for a given month indicate significant differences ($P < 0.05$). RU, relative units.

A significant regression was fitted for *FD-like*, *CsLFY*, *CsAPI* and *CiSEP1* genes ($r^2 = 0.78$, $r^2 = 0.66$, $r^2 = 0.77$ and $r^2 = 0.81$, respectively), and for *CsSL1* and *CuFUL* genes ($r^2 = 0.86$ and $r^2 = 0.76$, respectively) (Fig. 4). *CsTFL1* expression in the buds showed a negative relationship with tree age.

Nevertheless, considering the population of J-NFL trees only, *FD-like* and *CsLFY* gene expression did not show a significant relationship with tree age (Fig. 3A and D), as their expression did not show significant differences between 6-month and 7-year-old trees. However, a significant and direct relationship with J-NFL tree age was found for *CsAPI*, *CsSL1*, *CuFUL*, *CiSEP1* and *FLC-like* genes ($r^2 = 0.61$, $r^2 = 0.80$, $r^2 = 0.66$, $r^2 = 0.98$ and $r^2 = 0.78$, respectively) (Fig. 3B, C, E, G), and an inverse relationship for the *CsTFL1* gene ($r^2 = 0.91$, Fig. 3I). The *TEM1-like* gene did not correlate with tree age.

DISCUSSION

Flower formation is conferred by four networking processes (Weigel, 1995; Blázquez et al., 2006): (1) the location of newly emerging primordia, (2) correct timing for the formation of flowers, (3) the floral identity of the primordia and (4) the

outgrowth of the flower with the correct patterning. Our results show that 6-month-old meristems from juvenile citrus trees (6-month-old seedlings) were unable to transcribe flowering time and flower patterning genes (*SL1*, *SL2*, *FD-like*, *CuFUL*, *CiSEP1* and *CiSEP3*), i.e. to fulfil networking processes 2 and 4, whereas the 6-month-old meristems from A-NFL trees were able to do so, although they failed to achieve the threshold level of transcription required to flower. This is similar to *Arabidopsis*, for which during the juvenile vegetative phase expression of flowering time genes in the meristems, under long-day induction, is very short (6–7 d), and 14–15 d after germination the plant achieves the level of transcription required to flower (Blázquez et al., 1997; Valentim et al., 2015). We also found that the juvenile meristem does not develop the four networking processes equally during maturation.

Long-term meristem maturation

In citrus, *FD-like* and *CsLFY* genes of 6-month to 7-year-old trees did not show any significant variation in their expression just before floral bud differentiation, but that for 10-year-old

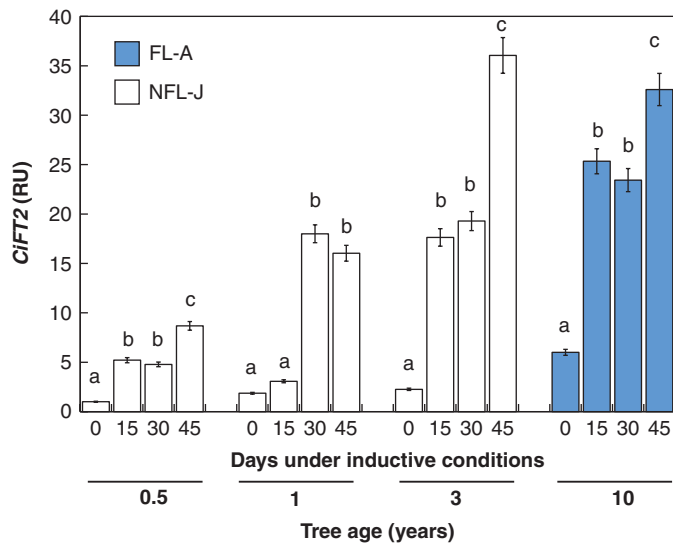


FIG. 3. Time course of *CiFT2* gene expression in leaves of 6-month, and 1- and 3-year-old juvenile (NFL-J) and 10-year-old adult 'Carrizo' citrange trees (FL-A). Data are mean of three independent replicates. Vertical bars indicate the standard error. Different letters for a given plant age indicate significant differences ($P < 0.05$). RU, relative units.

A-FL trees was upregulated eight- and 20-fold, respectively (Fig. 4). This result is similar to that obtained by Valentim *et al.* (2015) for *Arabidopsis* wild-type plants that exhibited an exponential trend for expression of these two genes during the juvenile to adult transition, taking into account the difference in the time scale of these two species, days for *Arabidopsis*, years for *Citrus*. However, the expression of other genes from the flowering programme, i.e. *CsSL1*, *CuFUL*, *CsAPI*, *CiSEPI* and *CsTFL1*, was progressively modified with the age of the tree. These results support the hypothesis that the four genetic programmes that confer flower formation (spatial, time, identity and patterning) might not necessarily occur sequentially (Blázquez *et al.*, 2006), and thus demonstrate that meristems in the juvenile tree progressively achieve a *mature vegetative stage*, with age-dependent differences in expression of flowering network genes. Specific flowering time is mainly uncoupled by a set of key genes (at least *FD-like* and *CsLFY* in our experiment), which play a fundamental role in the final decision to flower, while others maintain the vegetative stage. In monocarpic plants, the coordinated arrest of all meristems, a process called global proliferative arrest (GPA), which is phenotypically similar to the mature vegetative stage of our juvenile plants, is controlled by an age-dependent upregulation of the FUL transcription factor (Balanzá *et al.*, 2018). In our experiments, an age-dependent upregulation of *CuFUL* was also observed during the mature vegetative stage in juvenile trees (Fig. 4C).

Meristem and leaf role during the period of floral bud induction

The level of *API* and *LFY* transcription determines flower initiation in *Arabidopsis* (Blázquez *et al.*, 1997). In wild-type *Arabidopsis* plants, this threshold is achieved 2–3 d after the increase in flowering time gene expression (Valentim *et al.*, 2015).

Unexpectedly, our 6-month-old J-NFL trees were able to transcribe *CsLFY* and *CsAPI* genes in the meristem, from November to February, at least up to the same level (six- to seven-fold increase) as the 6-month-old meristems of the A-NFL trees. Neither the juvenile nor the adult trees flowered, and the only difference between them was the presence of fruits, whereas A-FL trees did flower. Therefore, it seems that transcription of *CsLFY* and *CsAPI* genes is dependent not only on juvenility but also on the presence of fruit, as previously shown by Muñoz-Fambuena *et al.* (2012).

The results suggest that the flowering-time programme prevails over the meristem identity programme. In citrus, *LFY* and *API* overexpression in transgenic 'Carrizo' citrange seedlings reduced flowering time to 12–20 months, but they needed the low-winter temperature signal to upregulate *CiFT2* expression and induce flowering (Peña *et al.*, 2001). Moreover, ectopic expression of 35S:*CiFT2* in citrus seedlings produced (1) extremely reduced flowering time (up to 16 weeks), and (2) continuous flowering regardless of exogenous induction conditions (Endo *et al.*, 2005), reinforcing the role of the flowering time programme in the control of juvenility in *Citrus*. In our experiments, the flowering time programme in the meristem significantly differed between trees. *FD-like*, *CsSL1* and *CuFUL* showed similar behaviours: (1) upregulation in the adult tree regardless of the flowering phenotype, and no transcription in the J-NFL tree (see Fig. 2A, C and D); and (2) a progressive upregulation of the *SL1* and *CuFUL* genes during the juvenile to adult meristem transition (Fig. 4B and C), which is in accordance with their implication in the age flowering pathway of *A. thaliana* (Wang *et al.*, 2009) and *Citrus* sp. (Castillo *et al.*, 2013). This latter result was not confirmed for the *FD-like* gene, at least in plants between 6 months to 7 years old, suggesting a key role for this gene for the vegetative-to-flowering meristem transition, as recently reported in pea (*P. sativum*) (Susmilch *et al.*, 2015). Moreover, in the adult stage, trees were able to express *FD-like* genes in the bud regardless of *CiFT2* transcription in the leaf (Fig. 1), as previously found by Muñoz-Fambuena *et al.* (2012). Phylogenetic analysis showed that the *C. clementina* *FD-like* gene was similar to the reported FD proteins (data not shown). The amino acid sequence alignment contains a bZIP domain, which is conserved in other FD proteins (Susmilch *et al.*, 2015). Therefore, it probably forms a florigen activation complex with *FT* genes, as reported previously (Susmilch *et al.*, 2015), although further studies are needed in this regard.

Pillitteri *et al.* (2004) compared *CsTFL1* gene expression in 4-month-old citrus seedlings and A-FL trees over 11 weeks, and found a correlation between *CsTFL1* gene expression and juvenility. Our results for 6-month-old trees agree with the latter, but further show that *CsTFL1* is not the cause of the lack of flowering in juvenile trees older than 1 year old, as it was downregulated in juvenile trees from 1 to 7 years old (Figure 2).

We also found that the flowering time programme was impeded in leaves of both juvenile and A-NFL trees, which were not able to transcribe the *CiFT2* gene up to the flowering level of the A-FL trees. Nishikawa *et al.* (2007) reported that 4-month-old citrus seedlings were not able to respond to low temperature (15 °C), compared to the adult flowering trees; however, we found a response to 15 °C/5 °C *CiFT2* upregulation in juvenile plants from 6 months to 3 years old (Figs 1 and 3). By contrast, no *CiFT2* expression was found up to fruit harvest in A-NFL

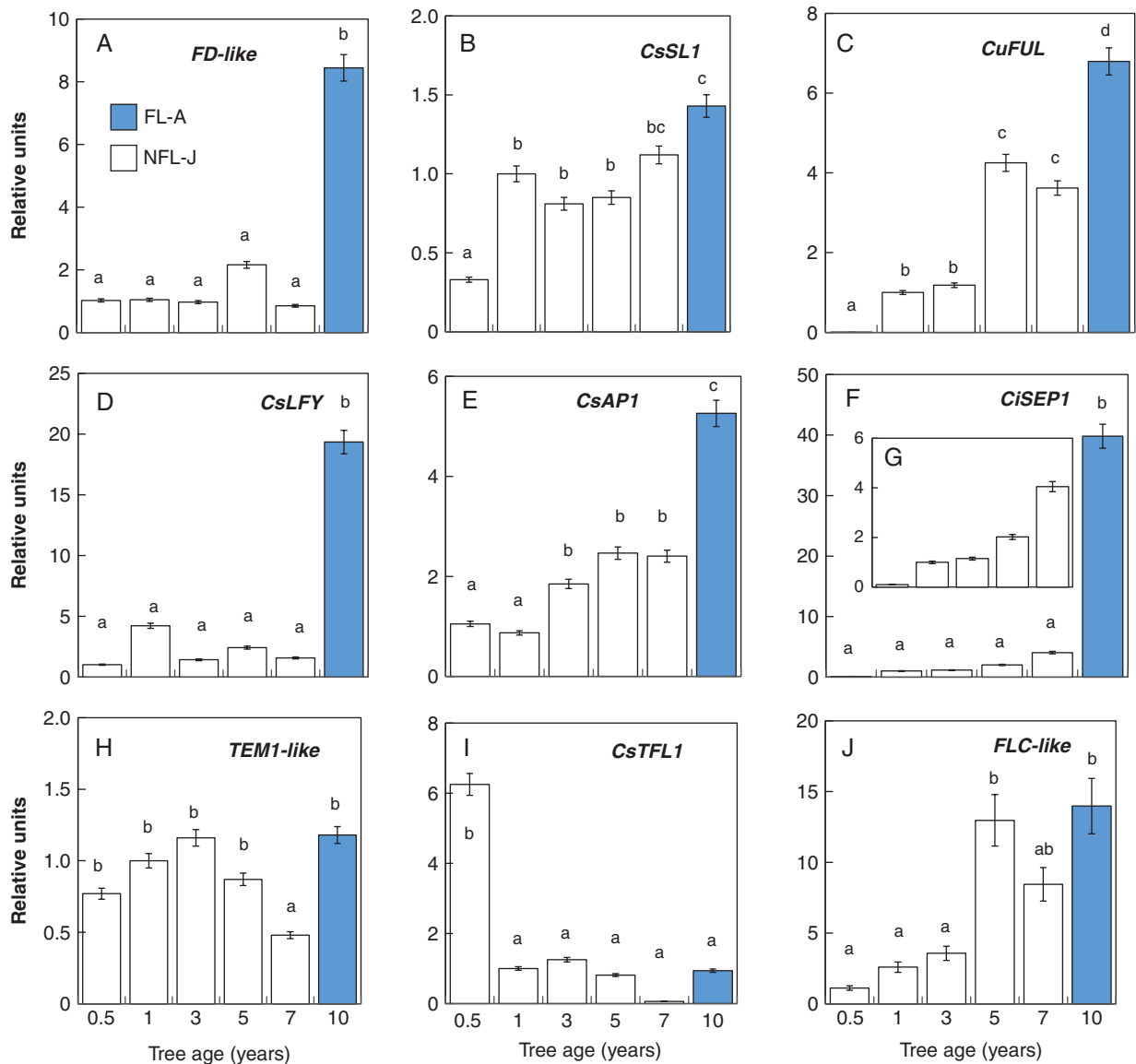


FIG. 4. Time course of *FD-like* (A), *CsSL1* (B), *CuFUL* (C), *CsLFY* (D), *CsAPI* (E), *CiSEP1* (F, G), *TEM1-like* (H), *CsTFL1* (I) and *FLC-like* (J) gene expression in the buds of 6-month, 1-, 3-, 5- and 7-year-old juvenile (NFL-J) and 10-year-old adult ‘Carrizo’ citrange trees (FL-A). Data are mean of three independent replicates. Vertical bars indicate the standard error. Different letters indicate significant differences ($P < 0.05$).

trees (Fig. 1), as previously shown by Muñoz-Fambuena *et al.* (2011). In *Arabidopsis* and *Antirrhinum* wild-type plants, *FT* is progressively transcribed during the adult vegetative phase, before the transition to the adult reproductive phase determined by *API* and *LFY* (Sgamma *et al.*, 2014; Valentim *et al.*, 2015). Our results suggest that leaves on the 6-month-old trees might be in transit to the adult vegetative stage, and that the reproductive stage is finally accomplished in the meristem. Supporting this hypothesis is the fact that early overexpression of *CiFT2* in transgenic citrus leads to the conversion of vegetative shoots into leafy inflorescences rather than the transition from the juvenile to adult phases (Endo *et al.*, 2005). Taken together, our results suggest that the inability of the meristem to differentiate may be the main constraint to flowering in the juvenile tree, whereas it is the fruit blocking expression of *CiFT2* in the

leaf, the main constraint to flowering in the adult tree. Further support for this hypothesis comes from the fact that *FLC-like* was barely expressed in leaves from young trees compared to adult citrus trees, a result also reported by Castillo *et al.* (2013), and its expression was highest in adult trees with fruits. The flowering inhibitor *FLC* is a transcription factor that acts in the vascular tissue to bind directly to the *FT* gene and to repress its transcription (Michaels and Amasino, 1999). However, a clear relationship between these two genes has yet to be established in fruit trees (Andrés and Coupland, 2012).

CONCLUSION

In *Citrus* species, axillary meristems of juvenile trees are unable to transcribe flowering time and flower patterning genes (*FD-like*,

CsSL1, *CuFUL*, *CiSEP1* and *CiSEP3*). Meristems progressively achieve the flowering time programme, and when they flower for the first time the derived meristems are always able to transcribe these genes, except when the adult tree produces a heavy crop load. In these cases, the axillary meristem is able to transcribe the flowering time programme genes, but fails to achieve the particular level of *CiFT2* transcription required to flower (Figure S1).

We propose that genetic inhibition of flowering time in juvenile trees is determined in the meristems and it is due to its immaturity, whereas in adult trees it is determined in the leaf, where repression of *CiFT2* gene expression occurs.

SUPPLEMENTARY DATA

Supplementary data are available online at <https://academic.oup.com/aob> and consist of the following. Table S1: List of primers used for quantitative real-time PCR. Figure S1: Diagrammatic representation of the repression of flowering time, meristem identity and flowering patterning genes in an adult fruiting *Citrus* tree and in a juvenile tree, compared to flowering promotion in an adult not-fruiting tree.

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