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Expression of coffee florigen *CaFT1* reveals a sustained floral induction window associated with
 asynchronous flowering in tropical perennials

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23

24 **Running title**

25 *CaFT1* is the *FLOWERING LOCUS T* ortholog in *Coffea* sp. L.

26

27 Highlight

Coffee florigen CaFT1 and related regulators revealed an extended floral induction window clarifying the asynchronicity and influence of environment for flowering in tropical perennial crops, providing perspectives to its control.

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32 Abstract

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The behavior of florigen(s) and environment-influenced regulatory pathways that control 34 flowering in tropical perennials with complex phenological cycles is poorly understood. 35 Understanding the mechanisms underlying this process is important for food production in the 36 face of climate change. To explore this, homologs of Arabidopsis florigen FLOWERING LOCUS 37 T (CaFT1) and environment-related regulators CONSTANS (CO), PHYTOCHROME 38 INTERACTING FACTOR 4 (PIF4) and FLOWERING LOCUS C (FLC) were isolated from 39 Coffea sp. L. (Rubiaceae). Overexpression of CaFT1 in Arabidopsis showed typical early-40 flowering and yeast two hybrid studies indicated CaFT1 binding to bZIP floral regulator, FD, 41 demonstrates that CaFT1 is a coffee orthologue of florigen. Expression of CaFT1 and floral 42 regulators were evaluated over one year using three contrasting genotypes: two C. arabica and 43 one C. canephora. All genotypes showed active CaFT1 transcription from February until 44 October, indicating a potential window for floral induction. CaCO expression, as expected, 45 varied over the day period and monthly with day length, whereas expression of temperature-46 47 responsive homologs, *CaFLC* and *CaPIF4*, did not correlate with temperature changes. Using coffee as a model, we suggest a continuum of floral induction that allows different starting points 48 49 for floral activation, which explains developmental asynchronicity and prolonged anthesis events in tropical perennial species. 50

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Keywords - Floral induction and development; *FLOWERING LOCUS T (FT)*; Asynchronous
flowering; *Coffea* sp. L. (Rubiaceae); Tropical perennial crops; Environment.

54

55 Abbreviations

- 56
- 57 BFT BROTHER OF FT
- 58 CO CONSTANS
- 59 FLC FLOWERING LOCUS C
- 60 FM Floral meristem
- 61 FT FLOWERING LOCUS T
- 62 GA Gibberellin

- 63 LD Long day
- 64 MFT MOTHER OF FT
- 65 PEBP Phosphatidylethanolamine-binding protein
- 66 PIF4 PHYTOCHROME INTERACTING FACTOR 4
- 67 SAM Shoot apical meristem
- 68 SFT TWEEN SISTER OF FT
- 69 SD Short day
- 70 TFL1 TERMINAL FLOWER 1
- 71 VM Vegetative meristem
- 72 73

INTRODUCTION

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In flowering plants, the transition of vegetative meristems (VMs) to a reproductive state 75 is mainly controlled by members of the phosphatidylethanolamine-binding protein (PEBP) 76 family that act as transcriptional coactivators (Cao et al., 2016). The Arabidopsis FLOWERING 77 78 LOCUS T (FT) gene is the first PEBP member identified to encode a florigen; that is, FT translocation via phloem from source leaves to VMs induces floral meristem (FM) production 79 (Wigge et al., 2005; Corbesier et al., 2007). To ensure proper reproductive timing, the FT 80 pathway integrates inductive exogenous stimuli, such as photoperiod, temperature, gibberellins 81 82 (GA) and vernalization, together the endogenous autonomous pathway (Amasino, 2010; Song et al., 2008; Kim et al., 2009; Andrés and Coupland, 2012). The flowering process is very well 83 84 explored in model plants such as Arabidopsis, however, it is poorly understood in perennial plants that have more complex reproductive cycles, such as *Coffea* sp. (Rubiaceae). 85

86 An example of this complexity is that perennial plants must maintain simultaneously in a branch the shoot apical meristem (SAM) in a vegetative state to guarantee the continuity of plant 87 vegetative growth and reproduction in subsequent years, whereas older meristems at the base of 88 branches form FMs and then flourish (Amasino, 2009; Albani and Coupland, 2010). Thus, this 89 mode of reproduction is intrinsically connected with plant architecture indicating a precise 90 molecular control to determine the different forms of meristems (Wang et al., 2009; Albani and 91 Coupland, 2010). The coffee phenological cycle is biennial, which means that it takes two years 92 to produce fruits (Teketay, 1999; Camargo and Camargo, 2001). The first year is characterized 93

by the floral induction and floral development until anthesis (or bloom) which, in *Brazil*, occurs
from January to March and in September, respectively (Camargo and Camargo, 2001;
Majerowicz and Söndahl, 2005; Morais *et al.*, 2008). Importantly, the floral induction period is
based on morphological observations, when vegetative and floral buds are distinguishable (de
Oliveira *et al.*, 2014), but without precise molecular characterization of the inductive floral
pathway (reviewed by López *et al.*, 2021 in press).

Organogenesis during coffee floral development was previously characterized 100 (Majerowicz and Söndahl, 2005; de Oliveira et al., 2014) and shown to be affected by 101 environmental conditions that result in uneven development and sequential flowering (Camargo 102 and Camargo, 2001; DaMatta and Ramalho, 2006; DaMatta et al., 2007). For example, in the G4 103 stage coffee buds enter a latent state during the cold and dry period of Brazilian winter in June to 104 August (Wormer and Gituanja, 1970; Camargo and Camargo, 2001; Morais et al., 2008). After 105 this period, bud growth resumes in a quick passage from the G4 to G6 following the anthesis that 106 coincides with the rain period around September. Thus, this suggests that coffee flowering is 107 controlled by different factors, such as low temperatures and plant water potential (Crisosto et 108 109 al., 1992; Majerowicz and Söndahl, 2005; DaMatta and Ramalho, 2006), in addition to photoperiod, as commonly observed in angiosperms. Nevertheless, from the molecular point of 110 view, the mechanism involved in the perception of these environmental stimuli and its 111 relationship with the FT-dependent inductive pathway has never been explored in coffee. 112

113 The PEBP family of proteins can be divided into five subgroups defined by Arabidopsis family members: FLOWERING LOCUS T (FT), TWIN SISTER OF FT (TSF), TERMINAL 114 FLOWER 1 (TFL1), BROTHER OF FT AND TFL (BFT) and MOTHER OF FT AND TFL 115 (MFT). These phosphatidyl ethanolamine-binding protein (PEBP) genes have important roles in 116 117 plant development, most notably floral induction and inflorescence architecture (Karlgren et al., 2011; Zhu et al., 2021). Accordingly, PEBP members are highly conserved between flowering 118 species, especially FT and TFL1 sequences (Turck et al., 2008; Wickland and Hanzawa, 2015). 119 Interestingly, FT and TFL1 proteins are highly similar, yet they have opposite functions. That is, 120 FT promotes floral meristem formation whereas TFL1 represses flowering and maintains the 121 inflorescence in an indeterminate state (Kaneko-Suzuki et al., 2018; Nakamura et al., 2019). 122 These antagonistic roles are due FT and TFL1 competition for the bZIP transcription factor FD 123 124 and 14-3-3 proteins to create a floral activation complex (FAC) or a floral repression complex

(FRC), respectively (McGarry and Ayre, 2012; Kaneko-Suzuki *et al.*, 2018). Another difference
is that FT protein accumulates in leaves and then moves long distances to the SAM (Yoo *et al.*,
2013), whereas *TFL1* is expressed in a subdomain of the SAM (Bradley *et al.*, 1997).

In Arabidopsis, long-day (LD) induction of FT expression is largely dependent on the B-128 box transcription factor CONSTANS (CO), which is modulated by the circadian clock and day 129 length (Amasino, 2010; Andrés and Coupland, 2012; Liu *et al.*, 2014). Under LD conditions, CO 130 protein accumulates to a threshold sufficient to activate FT expression, whereas in short days 131 (SD) CO is degraded via the 26S proteasome complex and does not activate FT expression 132 (Suárez-López et al., 2001; Valverde et al., 2004; Zuo et al., 2011). Other key transcription 133 factors (TFs) regulating FT are PIF4 and FLC, both responsive to temperatures changes and 134 activity depending on epigenetic changes in chromatin (Helliwell et al., 2006; Kumar et al., 135 2012; Madrid *et al.*, 2021). PIF4 is responsive to warmer temperatures and positively regulates 136 FT expression, whereas FLC is a repressor that is inactivated by prolonged cold periods 137 (vernalization) allowing FT expression and flowering initiation (Michaels et al., 2005; Amasino, 138 2010). FLC homologs together with other MADS-box genes were described in C. arabica (de 139 Oliveira et al., 2010; 2014) and related to drought (Barreto et al., 2012) that occurs in the cold 140 period of the Brazilian winter. However, the interplay between the FT pathway and 141 environmental signals through the identification and analysis of CO, PIF4 and FLC and their 142 roles in modulating floral induction is poorly understood in perennials and has never been 143 144 explored in coffee.

In addition to environmental effects, another regulatory factor is sugar signaling, which 145 146 plays an important role in plant development, including flowering regulation (Corbesier *et al.*, 1998; Lastdrager et al., 2014; Sheen, 2014). Day length directly affects sucrose and starch 147 148 accumulation as a consequence of photosynthesis, which has been correlated with reproductive induction (Corbesier et al., 1998). Accordingly, varying day length of different seasons is 149 150 directly associated with the transition from vegetative to reproductive time in plants (reviewed by Andrés and Coupland, 2012). A possible role for sucrose as a signal for flowering is 151 152 implicated by its influence on FT expression (Moghaddam and Van den Ende, 2013). Sucrose has been shown to act in leaves, downstream of CO and upstream to FT, to affect flowering 153 (Corbesier et al., 1998; Ohto et al., 2001). Moreover, transcriptomic studies of maize floral 154 regulator indeterminate1 (id1) shows a correlation between sugar and starch metabolism and 155

floral induction in this autonomously flowering plant (Coneva *et al.*, 2007, 2012; Minow *et al.*,
2018).

158 Florigen and FT orthologs in diverse plant species are key integrators of reproductive meristem specification via interconnecting molecular pathways that determine the proper floral 159 induction window (Amasino, 2009; He et al., 2020). Despite conservation across plant species, 160 variable expression patterns of FT homologs have been reported (Coelho et al., 2014; Cao et al., 161 162 2016; Wolabu et al., 2016; Štorchová et al., 2019). This variable expression likely reflects the variety of flowering evolutionary strategies required to tailor flowering time and floral 163 architecture diversification to particular geographical locations (Pin and Nilsson, 2012; Wickland 164 and Hanzawa, 2015; Jin et al., 2020). Very little research on the mechanisms underlying floral 165 induction and its interplay with environmental-dependent pathways has been carried out in 166 *Coffea* sp, a tropical perennial crop of considerable socio-economic importance (IOC, 2021). 167

Here we report the first characterization of homologs of the key floral regulators FT, CO, 168 PIF4, and FLC from C. arabica, C. canephora and C. eugenioides. Through transgenic analysis 169 and protein interaction assays we demonstrate that *CaFT1* from coffee is a functional floral 170 171 inducer (an FT ortholog). To determine the precise floral induction window and establish a possible correlation with environmental signals, expression of CaFT1 was determined during a 172 year at two daily time points together with FT regulators CaCO, CaPIF4, and CaFLC. 173 Expression was analysed in three coffee genotypes with contrasting flowering patterns, two C. 174 arabica cvs. Acauã and IAPAR59 and C. canephora cv. Conilon, to examine related intra- and 175 interspecific variation. In addition, to investigate a possible connection between sugar levels, 176 177 gene expression and floral development, carbohydrate content was determined at the same time points for all genotypes. Based on these results we propose that an extended floral induction 178 179 window for coffee could explain the asynchronous development and sequential flowering in this perennial species. Whereas *CaFT1* expression was correlated to short day photoperiod and cold 180 (Brazilian winter), a similar pattern was not detected in the expression profiles of CaCO, 181 *CaPIF4*, and *CaFLC*, indicating a more complex process regulating the flowering process. Thus, 182 183 this work contributes to our understanding of the floral transition in perennial species and could support future strategies aimed at mitigating asynchronous flowering. 184

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MATERIAL AND METHODS

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188 Plant material

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Plant material used for RNA extraction, cloning, gene expression, carbohydrate content 190 (total soluble sugar, sucrose, reducer sugar, and starch) was obtained from 4-year-old coffee 191 plants of three different cultivars established at the National Institute of Science and Technology 192 (INCT-Cafe) experimental field at the Federal University of Lavras, Brazil (21°23'S, 44°97'W): 193 C. arabica genotypes IAPAR 59 and Acauã, and C. canephora Conilon. All plants were 194 cultivated under nutritional and pest control conditions recommended for coffee (Vieira, 2008). 195 Three biological repetitions of each coffee cultivar were distributed randomly in the field. Each 196 sampling consisted of a mix of three completely expanded leaves collected and immediately 197 immersed in liquid nitrogen and stored at -80 °C until RNA extraction. Samples were collected 198 at five-time points (December 2016, February, April, June, and October 2017) over a two-day 199 period, at 6:00 am and 5:00 pm, considered the start and end of the day, totaling 90 samples. 200 Arabidopsis thaliana var. Landsberg erecta (Ler) was used for CaFT1 heterologous expression 201 studies following Coelho et al (2014) plant growth conditions. Plants were grown in growth 202 chambers under 16 hrs light / 8 hrs dark at 22 °C and 60 % humidity. 203

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205 In silico and phylogenetic analysis

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Searches for coffee homologs of FLOWERING LOCUS T (FT), CONSTANS (CO), and 207 PHYTOCHROME INTERACTING FACTOR 4 (PIF4) were performed by sequence comparisons 208 using the BLAST tool (Johnson et al., 2008). First, the described genes FT (At1G65480), CO 209 210 (At5G15840), and PIF4 (At2G43010), and FLC (At5g10140) from A. thaliana were used as queries (Lamesch et al., 2012) against the C. arabica (NCBI:txid13443) annotated genome 211 212 deposited at the National Center for Biotechnology Information (NCBI) and the Coffee Genome Hub (Denoeud *et al.*, 2014). To enrich searches for new coffee sequences phylogenetic analysis 213 214 also was performed with different homologs from various species, including Arabidopsis thaliana, Solanum lycopersicum, Brassica napus, Jatropha curcas, Nicotiana tabacum, Glycine 215 max, Oryza sativa, Zea mays, Sorghum bicolor, Solanum tuberosum and Populus nigra. The 216 sequence for coffee FLC, CaFLC, was retrieved from the previously reported sequence (de 217

218 Oliveira *et al.*, 2010).

Putative homologous sequences (>80% similarity and e-Value < 0.005) were aligned by 219 220 ClustalX2 (Larkin et al., 2007), analyzed by Genedoc software (Nicholas and Nicholas, 1997), and with the translated nucleotide sequence to protein, phylogeny was inferred with the nearest 221 neighbour joining method in MEGA-X (Kumar et al., 2018). Duplicate sequences were deleted 222 from the first phylogenetic tree and only considered A. thaliana and S. lycopersicum sequences 223 for the final tree (Fig. S1). The IDs of the used sequences are presented in the legend of Fig. S1. 224 To provide further evidence of the putative coffee FT as a floral inducer, key amino acids were 225 identified in alignment with CETS members from A. thaliana and S. lycopersicum (Fig. S2). The 226 conserved amino acids that distinguish the related proteins FT and TFL1, a floral inducer and a 227 repressor respectively (Ahn et al., 2006; Wickland and Hanzawa, 2015; Jung et al., 2016), are 228 shown in Fig. S2. Similar analyses were made to show the conserved amino acids between 229 coffee proteins and homologs of CO and PIF4 (Fig. S3 and S4). 230

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232 Gene cloning

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Total RNA from coffee leaves was isolated using ConcertTM Plant RNA Reagent 234 (Invitrogen) following the manufacture's recommendation. RNA concentration and purity were 235 measured by spectrophotometric analysis (GE NanoVueTM Spectrophotometer). All samples 236 237 were treated with Ambion DNase I (RNase-free) kit (Thermo Fisher) and cDNA synthesized using High-Capacity cDNA Reverse Transcriptase Kit (Thermo Fisher) following the 238 manufacturer's recommendation. Primers were designed for gene isolation with the putative 239 *CaFT1* sequence previously identified by *in silico* analysis and including the restriction sites 240 241 5° Rv = TTATCGTCTTCTGCCTC). The Polymerase Chain Reaction (PCR) was carried out 242 using *iProof* High-Fidelity DNA Polymerase (Bio-Rad) following the manufacture's protocol. 243 PCR fragments were isolated from 1 % agarose gel after electrophoresis and purified by 244 GeneJET Gel Extraction Kit (Thermo Fisher). The fragment generated was inserted into the 245 PJET1.2/blunt cloning vector (Thermo Fisher) and transferred to pK2WG7 plasmid (Thermo 246 Fisher), both previously digested using restriction enzymes, and then ligated with T4 DNA ligase 247 (New England Biolabs – NEB). All procedures followed the manufacturer's instructions. 248

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250 Plant transformation

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Arabidopsis var. Landsberg erecta (Ler) wild type plants (WT) and ft loss-of-function 252 mutants were transformed to overexpress CaFT1 using the Agrobacterium tumefaciens strain 253 GV3101::pMP90 and floral dip protocol (Clough and Bent, 1998), using the isolated and cloned 254 255 CaFT1 fragment under the control of the CaMV 35S promoter (Gene cloning above section). Seeds from different transformation events were harvested separately for each background 256 genotype. Positive transformed seeds were screened in selective growth media with Kanamycin 257 30 ug/mL and, after 2 days of incubation in darkness and at 4 °C, they were maintained in a 258 growth chamber under continuous light for one week. Twenty positive T1 plants were transferred 259 to soil and after two weeks, DNA was extracted from leaves and PCR analysis was conducted to 260 confirm insertions. Overexpression of CaFT1 driven by 35S CaMV was analyzed in 9 261 independent lines from 24 in T2 both Ler wild type (Ler-WT) plants as well as ft mutants. 262

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264 Yeast two Hybrid assay

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Protein-protein interaction analysis by Yeast Two Hybrid assay with CaFT against AtFD 266 and At14-3-3 was performed with the Matchmaker Gold Yeast Two-Hybrid System (Chien et 267 268 al., 1991). The CaFT1 sequence was amplified by *iProof* High-Fidelity DNA Polymerase (Bio-Rad) with restriction enzyme site tag insertion (Fw-EcoRI and Rv-BamHI), inserted to pBridge 269 plasmid by T4 DNA ligase and transformed in DH5a E. coli competent cells. Gold Yeast was 270 transformed with CaFT1-pBridge (Bait) and AtFD-pGADT7-RecAB (Prey); and CaFT1-pBridge 271 272 (Bait) and At14-3-3-pGADT7-RecAB (Pry). AtFD and At14-3-3 were screened from Arabidopsis GoldY2H library. 273

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275 Gene expression analysis (RT-qPCR)

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The primers for Reverse Transcription quantitative Polymerase Chain Reaction (RTqPCR), shown in Table S1, were designed using as template the *CaFT1*, *CaCO* and *CaPIF4* sequences identified in this work by *in-silico* analysis, whereas for *CaFLC*, the sequence was

280 previously characterized by de Oliveira et al. (2010). Primers were designed following specifications for standard qPCR such as amplicon length (between 80 and 150 bases), having no 281 282 sequences at the conserved domain, 40% to 60% GC content, and others as suggested by MIQE (Bustin et al., 2009) and examined for hairpins and dimers with the Oligoanalyzer Tool IDT 283 (https://www.idtdna.com/pages/tools/oligoanalyzer). The genes CaUBO2 and CaMDH and their 284 described primers were used as reference genes (Martins et al., 2017). RNA extraction, DNAse 285 treatment, and reverse transcription reaction were performed as described above (Gene cloning 286 session). RT-qPCR analysis was performed using 15 ng of cDNA to a final volume of 15 µL 287 reaction with Rotor-Gene SYBR® Green PCR Kit (Qiagen), in a Rotor Gene-Q (R) 288 thermocycler (Venlo, Netherlands). Mix reagents: 7.5 µL of SYBR-green (QuantiFast SYBR 289 Green PCR Kit - Qiagen), 3.0 µL of forward and reverse gene-specific primers, 1.5 µL of cDNA 290 291 at 10 ng/µL, and 3 µL of RNase-DNase-free water, resulting in at 15 µL final volume. Three biological repetitions were used, run in duplicate, and amplification was performed following 292 293 manufacturer's instructions. Relative expression differences were calculated by log2 of fold change and statistical analysis by Linear Mixed Model as described by Steibel *et al.* (2009). 294

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296 Carbohydrate Content Analysis

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Carbohydrate quantification was carried out following methods adapted by Meireles da 298 299 Silva et al, (2014) with modifications: 100 mg of fresh frozen tissue previously powdered with liquid nitrogen were homogenized in 5 mL potassium buffer (100 mM, pH 7.0) followed by 30 300 min incubation in a water bath at 40 °C. Supernatant was collected after 10 min centrifugation at 301 10,000 xG and stored at -20 °C. Pellets were used for starch extraction by 5 mL potassium 302 303 acetate buffer (200 mM, pH 4.8) resuspended with 16 units of amyloglucosidase enzyme, 304 followed by two hours at 40 °C and 20 min of centrifuge at 10,000 xG for supernatant separation and starch quantification. Starch, sucrose, and total soluble sugars were quantified according to 305 Dische (1962) and, reducing sugar levels according to Miller (1959). 306

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308 Statistical analysis

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A Linear Mixed Model was used for statistical analysis of biochemistry quantification by

the lme4 R package (Bates et al., 2015), and adjustment parameters following Oliveira et al,

312 (2020) statistical methods. RT-qPCR statistical analysis was carried out by log2 of fold change

by Linear Mixed Model as described by Steibel *et al.* (2009).

314 315

RESULTS

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317 Identification of *FT*, *CO* and *PIF4* homologs from *Coffea* sp.

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To identify FT sequences and other genes related to floral regulatory pathways, such as 319 CO and PIF4, we searched for homologous genes in C. arabica (NCBI:txid13443) and C. 320 canephora (https://coffee-genome.org/) genome databases. Through phylogenetic analysis, 321 322 sequences were compared to their homologs in other species to identify the most similar counterparts as putative orthologs (Fig. S1). Genome analysis of homologous sequences 323 324 identified seven FT-like genes, four CO-like genes and four PIF4-like genes. The PEBP family members, which include FT-like (Fig. S1-A), were separated into three subgroups based on 325 326 previous reports: FT-LIKE (FT and TSF), TFL1-like (TFL1 and BFT) and MFT-like (MFT) (Karlgren et al., 2011; Nasim et al., 2017). 327

Sequence alignments of PEBP homologs, including those from Arabidopsis thaliana and 328 Solanum lycopersicum (Fig. S2), revealed conserved amino acids in Coffea sequences related to 329 330 FT-like floral inducers. For example, the conserved Y85 amino acid residue reported to be a hallmark of FT orthologs supports the identified coffee genes as a possible florigen (Hanzawa et 331 al., 2005). These FT-like sequences used in further functional characterization were identical in 332 C. arabica and C. canephora and here both are referred to as CaFT1. Similar genomic analysis 333 was undertaken for CO-like (Fig. S3) and PIF4-like genes (Fig. S4), which showed conserved 334 335 motifs indicating they are putative *Coffea* sp. orthologs.

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Overexpression of *CaFT1* in *Arabidopsis* and protein interaction analysis suggest that it is a *Coffea* sp. *FT* ortholog

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We explored *CaFT1* function by over-expressing this gene in *Arabidopsis thaliana* Landsberg *erecta* (L*er*) to examine the effects on flowering. In addition, yeast two hybrid (Y2H)

assays were performed to test conserved protein-protein interactions between CaFT1 and 342 components of the Arabidopsis floral regulatory complexes. In 24 independent lines from T2 343 344 generations of *CaFT1* overexpression driven by the 35S CaMV promoter, nine independent lines caused a strong early flowering phenotype in both Ler wild type (Ler-WT) plants as well as ft 345 mutants. Transgenic plants flowered 12 days after germination (DAG) in contrast to Ler-WT 346 plants, which flowered at 28 days (Fig. 1A). Accelerated transition from vegetative to 347 reproductive development was also indicated by the reduced number of rosette leaves initiated in 348 both Ler-WT and ft mutant backgrounds (Fig. 1B). 349

Known conserved components of the Arabidopsis Floral Activation Complex (FAC) 350 include FT and FD together with a 14-3-3 protein (Kaneko-Suzuki et al., 2018). Y2H assays with 351 CaFT1 were carried out to test whether it interacts with AtFD and a 14-3-3 protein. CaFT1 352 protein was shown to interact with AtFD (Fig. 1C) as expected, since both proteins are reported 353 as partners to form a protein complex (Abe et al., 2005; Wigge et al., 2005; Corbesier et al., 354 2007; Jung et al., 2016). However, despite the 14-3-3 protein being a component of the FAC 355 with FT and FD proteins, no interaction was detected between CaFT1 and At14-3-3, suggesting 356 diversification on 14-3-3 homologs and/or components of the FAC in Coffea sp. Together these 357 results strongly support a role for CaFT1 as a floral inducer and that the selected CaFT1 gene is a 358 functional FT ortholog in Coffea sp. 359

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361 *CaFT1, CaFLC, CaCO, CaPIF4* show variable expression patterns over the course of a year

Cultivars of the same perennial species can show differential floral development patterns, 362 363 especially flowering time (reviewed by Lopez et al., 2021 in press). Such variation between cultivars is important for breeding programs to derive different cultivar genotypes with 364 365 contrasting floral development times (Carvalho, 2008). Relevant to this study, variable flowering patterns were observed for the C. arabica cvs. Iapar 59, Acauã and C. canephora cv. Conilon 366 cultivars (Fig. S5). To determine the floral induction window and explore a possible intra- and 367 inter-specific diversification of *CaFT1* related to the contrasting flowering patterns of coffee 368 369 cultivars, the expression of CaFT1 in leaves of these genotypes were examined by RT-qPCR over the course of a year (Fig. 2). 370

Despite quantitative differences, all coffee genotypes showed a clear "bell shaped" pattern of *CaFT1* expression during the year for both morning and afternoon samples. *CaFT1*

expression was observed in leaf samples from February, and increased progressively to 373 maximum expression in June, followed by drastically reduced transcript levels in October and 374 375 December (Fig. 2A). This result was correlated with the precise floral induction window, extending and encompassing the described emergence of floral buds (Majerowicz and Söndahl, 376 2005; de Oliveira et al., 2014). Comparing genotypes, C. arabica cv. Iapar 59 and C. canephora 377 showed similar values for *CaFT1* expression throughout the year and independent of the time 378 period, whereas C. arabica cv. Acauã showed lowest expression levels in February and highest 379 in April and June (Fig. 2A). This result can be associated with phenotypic differences between 380 early- and late flowering coffee cultivars (Fig. S5). 381

Moreover, FT and florigen genes from diverse plant species have been shown to act as a 382 hub for floral meristem activation by integrating different interconnected pathways responsive to 383 environmental signals (Amasino, 2010; Andrés and Coupland, 2012). Thus, to gain insights into 384 the regulation of CaFT1, the co-expression expression of photoperiod and thermosensitive FT 385 regulators, as CaCO, CaFLC and CaPIF4 (Figs. S1 to S4) were also determined at the same 386 time points (Fig. 2B, 2C and 2D). Expression analysis showed that *CaFT1* transcription is 387 detected first in leaves collected in February and increases gradually until it reaches a maximum 388 in June. This period coincides with the shorter day length photoperiod and colder temperatures, 389 typical of the winter in coffee-growing regions of Brazil (Fig. 4A and S6). 390

For all coffee genotypes, expression of *CaCO* was higher in the morning compared to 391 392 afternoon (Fig. 2B), in agreement with the photoperiod-dependent regulation of CO orthologs reported in other plants (Suárez-López et al., 2001; Zuo et al., 2011). In the morning, CaCO 393 394 expression presents a "smile shape" (or inverted bell shape) pattern throughout the year and was quite similar among genotypes, which means higher values in December and October and lower 395 396 in the middle timepoints. This result correlates with the photoperiod of longer day length (Fig. 4A and S6-A). In afternoon samples, CaCO expression was lower and stable with few 397 differences observed for genotypes, except in April and October. Comparing genotypes, in the 398 morning *CaCO* was expressed at lower levels in cv. Conilon than *C. arabica* both cvs. for all 399 400 time points throughout the year.

Regarding thermosensitive FT regulators, expression of *CaFLC* and *CaPIF4* was detected in all months for all three coffee genotypes, with few differences during the year in the morning, but varying in the afternoon (**Fig. 2C and 2D**). In the afternoon, *CaFLC* expression 404 presents different patterns and maximum levels depending on genotype, for *C. arabica* cv. Iapar 405 59 it is in December and cv. Acauã in April and *C. canephora* cv. Conilon June. In the morning, 406 no significant differences were found for *CaFLC* expression between *C. arabica* genotypes 407 while cv. Conilon were always expressed less. Similar to *CaFLC*, *CaPIF4* expression in leaves 408 showed little variation throughout the year, and no expression differences between coffee 409 genotypes were observed during the day or throughout the year, except for cv. Acauã in October 410 (**Fig. 2D**).

The CaFLC and CaPIF4 expression did not vary with the temperature changes in Brazil 411 (Fig. 4A and S6-B), with the lowest values in June and highest in October and December, 412 except for CaFLC in June and afternoon for cv. Conilon. Moreover, and importantly, CaFLC and 413 *CaPIF4* expression did not correlate with the expression pattern of *CaFT1* since it is expressed at 414 higher levels in June when the putative negative regulator CaFLC is expressed and CaPIF4 did 415 not change. In addition, no differences or expression correlations were found for precipitation 416 and humidity (Fig. S6-C and D). Thus, these results suggest that the floral inductive CaFT1 417 pathway is not influenced, or at least to a lesser extent, by these thermosensitive floral FT 418 419 regulators.

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421 Analysis of carbohydrate content in coffee leaves shows correlation with *CaFT1* expression 422

Sugars provide energy to all metabolic processes in plants and, with respect to the preparation for reproductive development, possibly expression of florigens such as *FT* as well (Corbesier *et al.*, 1998; Ohto *et al.*, 2001; Cho *et al.*, 2018). To examine a possible correlation with the *CaFT1* expression pattern and the energy status of coffee leaves, the carbohydrate content (total soluble sugar, sucrose, reducing sugar and starch) in adult plants was quantified at two daily timepoints over the course of a year.

Sucrose, starch, total soluble sugar and reducing sugars were quantified from the same samples used for expression analysis (**Fig. 2**). Results show patterns of distribution for the different coffee genotypes (**Fig. 3**). In general, all genotypes showed higher values for Total Soluble sugars (TS) and Sucrose (SC) in December and April (**Figs. 3A and B**), the summer period in Brazil, which is hotter and has more hours of daylight (**Figs. 4A**, **S6-A and B**). Similar results were observed for sucrose, but it was noted that cv. Iapar 59 had higher levels in all

435 months (Fig. 3B). For reducing sugars (Fig. 3C), all genotypes had similar levels throughout the 436 year, with *C. arabica* cv. Acauã showing higher values at all timepoints, except April. Finally, 437 the quantification of starch levels showed a contrasting pattern between coffee genotypes. For 438 example, the cv. Iapar 59 and cv. Conilon showed a clear pattern of increasing values from 439 December to October in the next year, differing only at the highest timepoints, June and April, 440 respectively. Whereas the cv. Acauã showed lower starch levels overall compared to other 441 genotypes and the highest levels in October (Fig. 3D).

These findings showed different patterns among the three coffee genotypes that could be 442 related to differences observed for *CaFT1* expression (Fig. 2A) and, consequently, flowering 443 time (Fig. S5). Thus, sucrose and other soluble sugars were the only types analysed whose levels 444 correlated with the period of higher CaFT1 expression, in April and June (Fig. 2A). Since 445 sucrose is the main transport sugar (Lemoine, 2000) it is possible that CaFT1 expression could 446 be responsive to the sugar state. Similarly, a correlation between the energy status and flowering, 447 as well as FT expression with trehalose-6-phosphate was established in Arabidopsis (Wahl et al., 448 2013; Fichtner et al., 2021). 449

450 451

DISCUSSION

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453 *CaFT1* function and expression revealed the floral induction window in coffee that could be 454 associated with the asynchronous flowering in perennial species

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Previous studies focused on understanding the transition from juvenile to reproductive 456 phase in perennial species have demonstrated that FT-related genes are involved in floral 457 458 induction and dormancy time in response to seasonal stimuli in trees (Böhlenius et al., 2006). However, the link between regulatory pathways of floral activation and asynchronous flowering 459 460 is still poorly understood. First, in this work we demonstrate that overexpressing *CaFT1* in Arabidopsis, wild type and ft-1 mutants, causes the typical early flowering phenotype and that 461 462 CaFT1 interacts with AtFD (Fig. 1), a component of the floral activation complex (Kaneko-Suzuki et al., 2018). Similar results were found for other annual or perennial plants when FT 463 homologs were overexpressed in Arabidopsis and/or also in their respective species; for 464 465 example, Poplar and apple (Kotoda et al., 2010; Tränkner et al., 2010), tomato (Cao et al., 2016),

Eucalyptus (Klocko *et al.*, 2016), Medicago (Laurie *et al.*, 2011) and blueberry (Gao *et al.*, 2016). Thus, CaFT1 can act as a florigen in *Arabidopsis* strongly suggesting same function in coffee plants as observed in other perennials. Further studies demonstrating that CaFT1 can act as a florigen in *Coffea* sp. will need to overcome the difficulty of transforming this species (Ribas *et al.*, 2011).

In coffee, the meristem floral transition window is reported to occur from January to 471 March (Camargo and Camargo, 2001; Majerowicz and Söndahl, 2005). To understand the 472 molecular events occurring at this transition, here we characterize an FT homolog from coffee 473 that acts as a florigen and show that it is transcriptionally active for a longer period than the 474 previously described window (Fig. 2A). CaFT1 expression extends from February to October, 475 reaching a maximum in June, which is a more precise characterization to indicate the potential 476 floral induction window. Such a window overlaps with the entire period of floral bud 477 development, which was interpreted as a continuum of induction (Fig. 4B). This extended 478 window explains the formation of floral buds at the base of branches (older buds) together with 479 induction of newly formed buds from the SAM at the distal end of branches (Fig. 4C). Floral bud 480 481 development starts at different times until anthesis in September, when CaFT1 expression decreases rapidly, followed by vegetative growth of new branches, restarting the cycle (Fig. 4B 482 and C). Moreover, our finding suggests that the developmental time is also different, or, in other 483 words, floral meristems induced early (at the base of the branch) develop slower than later ones 484 485 (those closer to SAM). Thus, the *CaFT1* expression patterns observed in this study suggest that the prolonged expression window is associated with the asynchronous flowering behavior of 486 coffee (Fig. 4B and C). Moreover, due to the conserved nature of flowering mechanisms, we 487 speculate that this occurs in other tropical perennials plants. 488

489 In addition, the *CaFT1* expression pattern in coffee supports evidence that floral induction occurs only in axillary meristems because the FT from the leaves is distributed along 490 491 the branch controlling floral induction to proximal meristems (McGarry and Ayre 2012). This also may explain the asynchronous flowering behavior of Coffea sp L., in which floral bud 492 development occurs at axillary meristems from branches formed at different times and does not 493 produce terminal flowers from the SAM (Fig. 4C). Alternatively, from the expression results 494 (Fig. 2 and 4B), it is possible that *CaFT1* other functions in addition to floral induction, since it 495 496 is reported to have a role in a wide range of developmental processes such as fruit set, vegetative

growth, stomatal control and tuberization (Pin and Nilsson, 2012). Further study is required toexplore this possibility.

499

500 **Coffee floral transition and photoperiodic stimulus** 501

Despite the extended window of *CaFT1* expression, it is clear from the analysis with 502 three genotypes that there is a conservation of the maximum level in June (Fig. 2), raising the 503 question of whether there is a correlation with environmental cues during the period (Fig. 4). The 504 505 florigen encoded by FT and its orthologs from numerous plants has been shown to induce the floral transition in response to temperature and photoperiod (Tsuji, 2017). With the isolation of 506 an FT orthologue in coffee, this analysis can be extended to floral induction in tropical species 507 that are subjected to high temperatures throughout the year. The expression profile of CaFT1 508 (Fig. 2A) may show less variation with respect to photoperiod effects, possibly because there is 509 less daylight variation in equatorial regions compared to temperate zones. 510

Examination of flowering control in trees provides evidence that CONSTANS (CO) 511 protein is responsible for FT ortholog expression induction by its accumulation in long days 512 (LD) (Böhlenius et al., 2006). However, despite CaCO showing expression variation in response 513 to day length (Fig 2B), its expression over the course of the year showed an inverse relationship. 514 515 The *CaCO* showed a variable expression pattern for all three coffee genotypes with higher values reached in the morning, according to the circadian clock that is mainly responsive to photoperiod 516 (Valverde et al., 2004). Comparing expression over the year, CaCO was higher in December and 517 October (Fig. 2B), coinciding with summer in the Southern hemisphere and the period of 518 519 greatest solar light incidence (Fig. 4 and S6-A). This was opposite to the pattern since CaFT1 expression is higher during the Brazilian winter (Fig. 2A), with the shortest day length (Fig. 4A 520 and S6-A) and more highly expressed in summer in December and October (Fig. 2A). Other 521 factors might be involved in the CO regulatory pathway of CaFT1 expression; for example, 522 523 Wenkel, et al, (2006) showed that the protein HEME ACTIVATOR PROTEIN2 (HAP2) or HAP3 can create a complex with CO that causes a reduction in the expression of FT. 524

525

526 Expression of *CaFT1* is not correlated with *CaFLC* and *CaPIF4* expression, suggesting 527 alternative thermoregulatory pathways that are not influenced by temperature

528

In regions where temperatures vary during the year, floral initiation and dormancy for 529 some species can be controlled by cold exposure or vernalization (Michaels et al., 2005; Kim et 530 al., 2009; Madrid et al., 2021). In the Arabidopsis vernalization pathway, FLC expression is 531 repressed after prolonged exposure to colder temperatures, thus activating FT expression and 532 allowing floral meristem induction (Michaels and Amasino, 1999; Helliwell et al., 2006; Aikawa 533 et al., 2010). On the other hand, PIF4 is responsive to warmer temperatures and positively 534 regulates FT expression (Kumar et al., 2012). Thus, we hypothesized that FLC and PIF4 535 homologs in coffee could be related to thermal-sensitive pathways, possibly regulating CaFT1 536 expression. Here we use them as a probe to evaluate co-expression patterns that are integrated 537 538 with temperature changes during the growing season.

In this work, FLC homologs were found in coffee, which is interesting given that it is a 539 tropical plant that does not normally experience freezing temperatures. In line with this, CaFLC 540 expression was not associated with temperature variation, as might be expected. A similar result 541 542 was found for CaPIF4 expression (Fig. 2C and D, Fig. 4A and S6-A). Over the course of the experiment, when the lowest temperatures were registered in June and the highest occurred in 543 December, CaFLC and CaPIF4 showed stable expression patterns or changes in different 544 periods not correlated with these temperature changes. Moreover, comparing *CaFT1* and *CaFLC* 545 expression patterns, there was no inverse co-expression correlation as would be expected if FLC 546 repressed FT activity, nor was there a positive correlation with CaPIF4, suggesting it is not an 547 FT inducer. Overall, expression of *CaFT1* was highest in June, during the Brazilian winter (Fig. 548 2A, Fig. 4A and S6-A), suggesting positive regulation by cold or even drought, typical 549 characteristics of the Brazilian winter. Further analysis under controlled conditions will 550 demonstrate whether CaFT1 is responsive to cold and/or drought. In addition, CaFLC expression 551 552 varied between coffee genotypes, with Iapar 59 and Acauã showing similar expression, whereas 553 Conilon showed lower expression levels throughout the growth period and higher in afternoon samples of June (Fig. 2C). These differential expression patterns suggest intra- and inter-specific 554 transcriptional differences, which might coincide with contrasting flowering behavior (DaMatta 555 and Ramalho, 2006), or it could be associated with greater phenotypic homeostasis of the 556 557 allotetraploid C. arabica than its diploid parents in response to different temperature conditions (Bertrand *et al.*, 2015). 558

559 These results show no evidence that the FT-dependent floral transition pathway is regulated by coffee FLC or PIF4 homologs, thus the potential roles of CaFLC and CaPIF4 in 560 561 flowering of tropical perennials awaits further study. Alternatively, despite tropical species such Coffea sp. having sequences that are homologous to Arabidopsis FLC and PIF4, they may have 562 functions not necessarily related to floral control in response to temperature variation. The 563 finding that *CaFLC* expression varied widely in different coffee tissues such as, roots, leaf, SAM 564 at all floral and fruit developmental stages (de Oliveira et al., 2014) suggests functions other than 565 flowering repression, such as coordinating organogenesis together with SOC1 as reported in 566 Arabidopsis (Deng et al., 2011). In support of a possible role in organogenesis, CaFLC 567 expression is upregulated in response to drought (Barreto et al., 2012), which coincides with the 568 growth latency in the G4 stage interpreted as a dormant stage (Wormer and Gituanja 1970; 569 Majerowicz and Söndahl, 2005). At present, however, no direct mechanism for this has been 570 described yet (reviewed by Lopez et al., 2021 in press). 571

572 Future research will decipher whether CaFLC and CaPIF4 are involved in flowering, and 573 their relationship with environmental signals and the FT-regulatory pathways. In addition, both 574 genes are regulated by chromatin epigenetic changes in Arabidopsis (Helliwell *et al.*, 2006; 575 Kumar *et al.*, 2012; Madrid *et al.*, 2021), a very little explored field of research in *Coffea* sp. and 576 crop perennials in general.

577

578 The role of carbohydrates in coffee floral induction as a model for perennials

579

Sugars have been shown to be important chemical signals that affect flowering, as 580 strongly supported in model plants like Arabidopsis (Wahl et al., 2013; Cho et al., 2018; 581 582 Fichtner et al., 2021). Since no correlation was found between CaFT1 expression and possible environment-related regulators, we examined sugar levels to assess their role as possible 583 584 regulators in tropical perennial species. Gene expression analysis showed association with Total Soluble Sugar and Sucrose levels (Fig. 3A and B) in relation to temperatures during the five 585 586 analyzed months, showing an association with warmer periods of the year (Fig. 4A and S6-A). This association is related to higher levels of TS and SC and coincided with CaFT1 expression 587 levels (Fig. 2A). As previously described by Cho et al. (2018) and, in accordance with our 588 results (Fig. 4), sugar levels, a product of photosynthesis, can change according to seasons in 589

response to more hours of light and higher temperature.

Sugar variation is an important indicator of reproductive phase initiation, with evidence 591 592 that it affects the expression of floral integrators, such as FT (Rolland et al., 2006; Moghaddam and Van den Ende, 2013). In our study, sugar accumulation was higher in December and April, 593 the summer period in Brazil that is hotter and has more hours of daylight (Fig. S6-A). 594 Interestingly, these results coincide with two relevant periods associated with *CaFT1* expression 595 - in December, before the increase in CaFT1 expression in February, and in April which is a 596 period before *CaFT1* expression peaks in June. Whether this pattern suggests that sugars act as a 597 stimulus for *CaFT1* expression requires further evidence (Fig. 1A). Association of *CaFT1* with 598 TS and sucrose levels suggest that its expression is responsive to carbohydrate signals, showing a 599 possible connection between sugars and floral induction in perennial species. Thus, it will be 600 interesting to determine whether energy status is a key regulator of CaFT1 expression and 601 controls coffee floral development, further suggesting that environmental factors are indirectly 602 involved since photo-assimilate production is affected by photoperiod and temperature (Pego et 603 al., 2000; Lastdrager et al., 2014). Accordingly, we demonstrate that sugar content in coffee 604 605 plants changes in response to temperature regimens (de Oliveira et al., 2020). These findings support previous studies demonstrating a correlation between FT expression pattern and 606 carbohydrate content (Corbesier et al., 1998). 607

608 609

CONCLUSIONS

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Floral induction in perennial plants is a poorly understood process that integrates 611 endogenous and environmental factors. Elucidating the molecular components of this floral 612 613 regulatory process is a necessity in the face of imminent climate change to secure food production (Howden et al., 2007; Lobell et al., 2011; Zhao et al., 2017). In this work we 614 615 identified the coffee FT ortholog and determined its expression profile to describe the precise and potential floral induction window. This analysis includes the analysis of key FT regulators 616 617 responsive to environmental signals, evaluation of climatic parameters and sugar content over the period of one year. Together, our results indicate a continuum of florigen transcription, 618 conserved between contrasting coffee genotypes, that could underlie asynchronous floral 619 development and flowering. The environment-related floral regulators, CaCO, CaFLC and 620

CaPIF4, were not co-expressed as expected with *CaFT1*, whereas a correlation was found with sugar content, which is affected by environmental changes over the course of a year. This suggests that *CaFT1* is not directly regulated by these genes, but that there may be a connection between sugar metabolism and florigen function in coffee. Thus, the present work contributes to comprehending asynchronous flowering in tropical perennials plants and provides a basis for targeting molecular components in crop breeding programs.

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628 **SUPPLEMENTARY DATA** 629

Table S1 shows all the primers used in this work; figures S1 to S4 show the phylogenetic analyses and amino acids alignments for the gene families *FT*, *CONSTANS*, *FLC* and *PIF4*; Figure S5 shows a photo panel of the plagiotropic branches with floral meristems at different stage of development comparing *C. arabica* cvs. Iapar 59 and Acauã and *C. canephora* cv. Conilon in three different timepoints; Figure S6 shows the variation of photoperiod, temperature, precipitation and relative humidity during the experiments at the experimental field of Federal University of Lavras (UFLA, MG/Brazil).

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650

651 AUTHORS' CONTRIBUTIONS

652

653 C.H.C., R.R.O. and A.C-J. conceptualized the project. C.H.C., conducted all experiments and 654 data analyses. L.P.P. participated in the collection of plant materials, RNA extractions and the 655 RT-qPCR execution. V.L. and C.F. participated in the yeast two hybrid and heterologous 656 expression analyses. T.H.C.R. assisted bioinformatic and statistical analyses. R.R.O., J.C. and 657 A.C-J. conceived of and supervised the experiments and data analyses. C.H.C. wrote the 658 manuscript. R.R.O., J.C and A.C-J revised the manuscript and contributed to writing.

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661 CONFLICT OF INTEREST

The authors declare that there is no conflict of interest.

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DATA AVALIABILTY

Main data supporting the findings of this study are available within the paper and within its
supplementary materials published online. The raw data used for analyses and figures are
available from the corresponding author, Antonio Chalfun-Júnior, upon request.

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930 FIGURE LEGENDS

931

Figure 1 – Functional analysis of CaFT1. A – Heterologous overexpression of CaFT1 in 932 Arabidopsis thaliana Ler ecotype causes early flowering. From left to right: Wild-type (Ler-933 934 WT); 35S::CaFT1 construct in Ler-WT background; FT loss-of-function (Ler-ft); 35S::CaFT1 construction with Ler-ft background. The overexpression of CaFT1 driven by 35S CaMV was 935 analyzed in 9 independent lines from 24 in T2 both Ler wild type (Ler-WT) plants as well as ft 936 mutants. **B** – Rosette leaf number compared between Ler-WT / 35S::CaFT1, Ler-ft / 937 35S:: CaFT1, Ler-WT, and Ler-ft. C - Yeast Two Hybrid Protein-protein interaction assay: Yeast 938 transformed with Empty plasmid, AtFD, and At14-3-3 inserted into pGADT7-RecAB plasmid 939 (Prey), against CaFT1 inserted into pBridge plasmid (Bait). Transformed yeast grown in SD 940 medium as negative control and SD/-LTH/X-α-Gal as select medium and, as positive control 941 Arabidopsis FT interacted with Arabidopsis FD in SD/-LTH/X-α-Gal select medium. 942

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Figure 2 - RT-qPCR expression analysis of putative floral regulators genes in leaves 944 collected over one year from three different coffee genotypes: C. arabica cvs. Iapar 59 and 945 Acauã and C. canephora cv. Conilon. Expression analysis were performed for the genes 946 previously characterized (Fig. 1 and S1 to S4), CaFT1 (A), CaCO (B), CaFLC (C) and CaPIF4 947 (D). Leaf samples of different coffee genotypes were used. C. arabica cvs. IAPAR59 (black 948 bars) and Acauã (white bars) and C. canephora cv. Conilon (gray bars). Samples collected at the 949 950 first hours of the day (morning) and at the last hours of the day (Afternoon) in December of 2016 (DEC) and February (FEB), April (APR), June (JUN) and October (OCT) of 2017. Relative 951 expression differences were calculated as log2 fold Change by Linear Mix Model as described 952 by Steibel, et al. (2009). Letters at the top of the bars indicate the statistical differences, in which 953 954 lower case letters represent comparisons between genotypes in each month, and capital letters 955 compare the same genotype at different time points during the year.

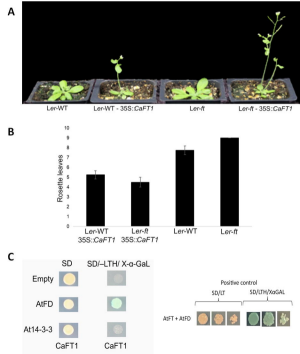
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Figure 3 – Carbohydrate content quantification from coffee leaves in microgram per gram
of fresh weight. Samples collected from two *Coffea arabica* cultivars, Iapar 59 (red line), Acauã
(blue line), and one *Coffea canephora* cultivar, Conilon (purple line), collected along five
months of a year (DECEMBER, FEBRUARY, APRIL, JUNE, and OCTOBER). A – Total

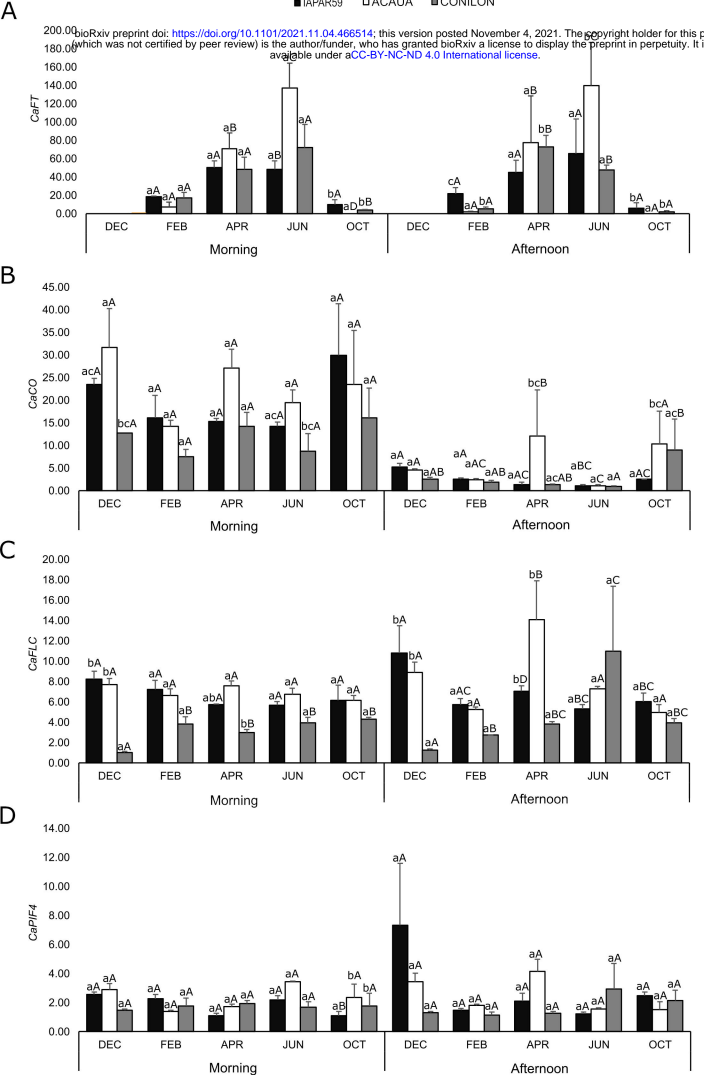
soluble sugar; \mathbf{B} – Sucrose; \mathbf{C} – Reducing sugar; \mathbf{D} – Starch. Each sampling data set is represented by a point in the line with the standard deviation. All cultivars in each timepoint were represented by 3 biological samples and two technical replications.

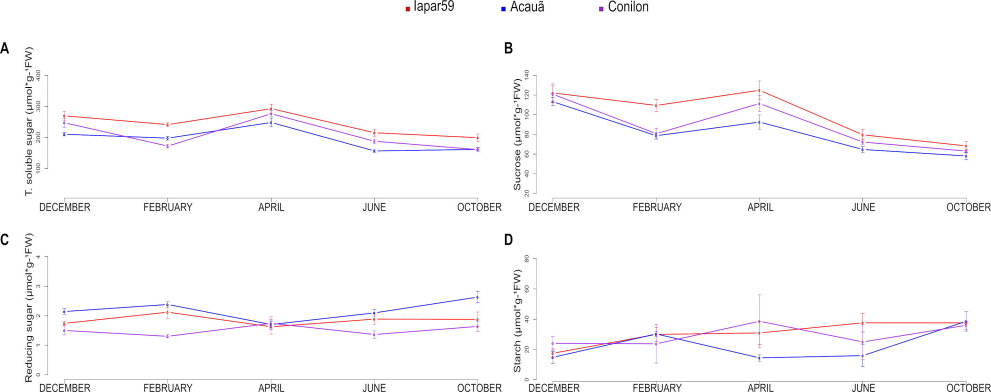
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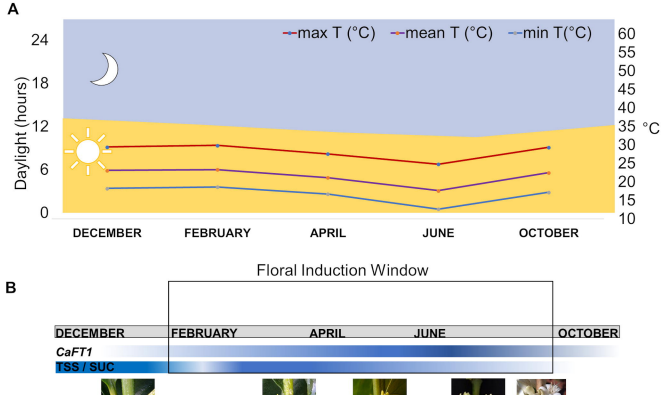
Figure 4 – The floral induction window in coffee trees as a model for perennial species. 966 Figure A, left axis of the graph represents the daylight hours variation divided by 24 hours which 967 is separated in yellow color (day) and light blue color (night) in the graphic area. Right axis 968 represents degree values (Celsius) from minimum (blue line), mean (purple line), and max (red 969 line) temperature variation over the year; **B**, proposed floral induction window from February 970 until October based on variation of *CaFT1* expression levels, total soluble sugars (TSS) and 971 sucrose (SUC) content over the year. Gradients of color intensity (dark blue indicates higher 972 level, light blue lower levels and white not detectable) represents the content level in terms of 973 gene expression, total soluble sugar, and sucrose variation. The floral development stages 974 observed in plants when the leaves were collected for the five time points (months) are also 975 976 indicated in the figure; C, shows a plagiotropic branch with floral buds in the same development in both parts, the early formed branch (brown and lignified stem) and the late formed branch 977 978 (green stem). Because the late formed branch originates from early nodes, the continuous vegetative activity of the shoot apical meristem (SAM) along the year (details in Figs. S5 e S6), 979 980 in this figure suggests a need for extended florigen activity to induce axillary meristems formation at different times. Interestingly, floral meristems induced at different times reach 981 982 flower buds at the same time.













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Coffea arabica

