Morphological, Physiological, and Molecular Responses of Sweetly Fragrant Luculia Gratissima During the Floral Transition Stage Induced by Short-day Photoperiod

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Research article

Keywords: Luculia gratissima, Floral transition, Photoperiod, Flowering pathway, Phytohormone, Regulatory network

DOI: https://doi.org/10.21203/rs.3.rs-92695/v1

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Abstract

Background: Photoperiod-regulated floral transition is vital to the flowering plant. *Luculia gratissima* ‘Xiangfei’ is a flowering ornamental plant with high development potential and is a short-day woody perennial. However, the genetic regulation of short-day-induced floral transition in *L. gratissima* is unclear. To systematically research the responses of *L. gratissima* during this process, dynamic changes in morphology, physiology, and transcript levels were observed and identified in different developmental stages of long-day and short-day-treated shoot apexes.

Results: The results showed that floral transition in *L. gratissima* occurred 10 d after short-day induction, but flower bud differentiation did not occur under long-day conditions. A total of 1,226 differentially expressed genes were identified, of which 146 genes were associated with flowering pathways of sugar, phytohormones, photoperiod, ambient temperature, and aging signals, as well as floral integrator and meristem identity genes. The trehalose-6-phosphate signal positively modulated oral transition by interacting with SPL4 in the aging pathway. Endogenous gibberellin, abscisic acid, cytokinin, and jasmonic acid promoted floral transition, whereas strigolactone inhibited it. In the photoperiod pathway, FD, COL12, and NF-Ys positively controlled floral transition, whereas PRR7, FKF1, and LUX negatively regulated it. SPL4 and pEARLI1 positively affected floral transition. SOC1 and AGL24 integrated multiple flowering signals to modulate the expression of *FUL/AGL8, AP1, LFY, SEPs, SVP,* and *TFL1,* thereby regulating floral transition. Finally, we propose a regulatory network model for short-day-induced floral transition in *L. gratissima.*

Conclusions: Short-day photoperiod activated systemic responses of morphology, physiology, and transcript levels in *L. gratissima* and induced the generation of oral transition signals in the photoperiod pathway. Furthermore, multiple flowering signal pathways including phytohormone-, sugar-, temperature-, age-related genes synergistically control this process. This study improves our understanding of flowering time regulation in *L. gratissima* and provides knowledge for its production and commercialization.

Background

Floral transition (the switch from vegetative to reproductive development) is a critical stage in life history of flowering plants, particularly in horticultural ornamental plants [1, 2]. This process is regulated by both environmental and endogenous signals [1]. Recently, major breakthroughs have been made in research on the molecular regulatory networks of floral transition in *Arabidopsis thaliana* [3, 4]. In *A. thaliana,* different endogenous (autonomous, gibberellin, circadian rhythm, age, sugar signals) and environmental (vernalization, temperature, and photoperiod) signals congregate on some floral integrators, such as *SUPPRESSOR OF OVEREXPRESSION OF CONSTANS 1 (SOC1), FLOWERING LOCUS T (FT),* and *AGAMOUSLIKE24 (AGL24),* further activating floral meristem identity genes, such as *LEAFY (LFY)* and *APETALA1 (AP1),* which irreversibly convert vegetative meristem to floral meristem [5]. However, there is still much to learn regarding the regulation of floral transition in perennial woody plants. Perennial woody
plants do not die after flowering. Instead, they produce new flower buds and vegetative branches annually and have characteristics of long reproductive cycles and seasonal flowering [6]. Therefore, studies on annual plants cannot completely reveal the floral transition mechanisms in perennial woody plants. There are significant differences in the molecular mechanisms of floral transition in perennial woody plants compared with those of *A. thaliana*. For example, gibberellin (GA) promotes the transition from vegetative to reproductive development in *A. thaliana* but has inhibitory effects in some perennial woody plants [7–9]. Therefore, it is particularly important to study the molecular mechanisms of floral transition in perennial woody plants.

*Luculia gratissima* (Wall.) Sweet. (Rubiaceae) is a perennial evergreen shrub or small tree that is distributed in the south-eastern edge of the Tibetan plateau in southwest China and neighboring Nepal and Myanmar [10]. *L. gratissima* ‘Xiangfei’, a new cultivar cultivated by our research team for many years, has pink flowers, a strong fragrance, and a large and dense inflorescence (Fig. 1a, b); it is a woody horticultural flower with great ornamental value and development potential. However, this plant has not entered the large-scale commercial production stage because of imperfect flowering time regulation techniques. Previous studies showed that the cultivar ‘Xiangfei’ can only complete floral transition at short-day (SD) photoperiods [11], and thus controlling daylength to induce flowering is required to achieve year-round production. Therefore, understanding the mechanisms of short-day-induced floral transition in *L. gratissima* ‘Xiangfei’ has important significance for understanding and solving flowering-related problems.

In the present study, we investigated responses of *L. gratissima* during short-day-induced floral transition stage at the morphological, physiological, and transcriptome levels. The aims of this study were: (1) to observe shoot apexes of *L. gratissima* of short-day treatment during five developmental stages using morphological and histological methods to identify the time point of floral transition in *L. gratissima*; (2) to measure endogenous substance contents to study the soluble sugar and hormone effects in floral transition in *L. gratissima*; (3) to conduct an RNA sequencing (RNA-seq) analysis of the transcriptomes of *L. gratissima* shoot apexes at four different stages, 7, 10, 13, and 19 days after the initiation of long-day (LD) and short-day (SD) treatments, to study the molecular regulatory mechanism of short-day-induced floral transition in *L. gratissima*. The results presented in this research will aid in regulating *L. gratissima* flowering and achieving year-round production. Additionally, identification of important regulatory genes will provide important guidance for flowering-related molecular breeding in the future.

**Results**

**Morphological differentiation of shoot apexes during floral transition**

The morphological differentiation of *L. gratissima* shoot apexes was observed through paraffin sections. The results showed that 0 d to 7 d under the SD treatment (SD0 to SD7) was the vegetative growth stage (undifferentiated stage), in which the tip of the growth cone in the bud was narrow and pointed and surrounded by leaf primordia (Fig. 2a-c). At 10 d after the initiation of the SD treatment (SD10), the bract
primordial differentiation stage began in which the growth cone of the bud appeared hemispherical, which was an important marker of the transition from vegetative growth to reproductive growth. Subsequently, the two sides of the hemispherical growth cone elongated to form two protrusions, i.e., bract primordia (Fig. 2d-f). At 13 d after the initiation of the SD treatment (SD13), the inflorescence primordial differentiation stage began. At this stage, the growth cone in the bract primordia elongated to form three hemispherical protrusions, i.e., inflorescence primordia. Simultaneously, the lateral base of the bract primordia differentiated into lateral inflorescence primordia. Next, bilateral protrusions at each hemispherical inflorescence primordium differentiated into bract inflorescences (Fig. 2g-i). At 19 d after the initiation of the SD treatment (SD19), the floret primordial differentiation stage began and a single inflorescence primordium in the bract primordia gradually widened to become floret primordia at the tip of the bud (Fig. 2j-l). These results showed that the floral transition period began 10 d after the initiation of the SD treatment, and the selection of time points before and after this period could facilitate the physiological study of floral transition. Therefore, the LD treatment was used as a control in this study, and 7 d (SD7), 10 d (SD10), 13 d (SD13), and 19 d (SD19) in the SD treatment were selected to study the physiological and molecular regulation patterns of floral transition.

**Dynamic changes in endogenous substance content during floral transition**

Contents of soluble sugars and endogenous hormones (gibberellin [GA$_3$], indoleacetic acid [IAA], abscisic acid [ABA], and zeatin [ZT]) were measured at 0 d [SD0], 7 d [SD7], 10 d [SD10], 13 d [SD13], and 19 d [SD19] after the initiation of the SD treatment. The Kruskal-Wallis H test results showed that except for GA$_3$, which could not be detected because it was below the limit of quantitation (0.1 ng/ml), there were significant differences in the contents of the other substances among the five stages (adjusted $p < 0.05$; Fig. 3). Soluble sugar, ZT, and IAA reached their peaks at SD0, which were $28.86 \pm 0.67 \text{ mg g}^{-1} \text{ FW}$, $2.15 \pm 0.30 \text{ ng g}^{-1} \text{ FW}$, and $0.69 \pm 0.04 \text{ ng g}^{-1} \text{ FW}$, respectively. Additionally, soluble sugar and ZT decreased from SD0 to SD19. Interestingly, IAA showed an increase in SD13 before decreasing. Similarly, ABA initially increased from SD0 to SD13 and subsequently declined.

Additionally, post-hoc results showed that there were extremely significant differences in the pairwise comparisons between the five timepoints for ABA ($p < 0.001$). IAA only showed no significant differences between SD7 and SD13 ($p > 0.1$). Soluble sugar did not show any significant differences between SD0 and SD7 ($p > 0.1$). ZT did not show any significant differences between SD0 and SD10, between SD7 and SD10, or between SD13 and SD19 ($p > 0.1$). From these results, it can be seen that ABA levels changed rapidly and dynamically over these five stages, whereas ZT levels exhibited little change over this same time period. Changes in soluble sugar content mainly occurred in later periods (SD13 to SD19). In contrast to these substances, IAA changes were relatively constant (Fig. 3).

**RNA-seq and qRT-PCR identification of differentially expressed genes (DEGs)**

Transcriptomes were generated for three biological replicates from the SD and LD treatments at each of the time points corresponding to the four stages of bud differentiation in SD treatment plants, i.e., at 7 d,
10 d, 13 d, and 19 d (Fig. 2), yielding a total of 24 transcriptomes. A total of 1,236,426,670 raw sequencing reads were generated from 24 samples, $1.2 \times 10^9$ high-quality clean reads (181 Gb) were obtained after filtering, with mean Q20, Q30, and GC contents of 99.11%, 97.18%, and 43.53%, respectively (Additional file 1: Table S1). A total of 79,870 unigenes (≥ 200 b) were generated from de novo assembly, and the N50 length was 2,118 bp (Additional file 2: Table S2). Among these unigenes, 35,725 unigenes (44.73%) were successfully annotated to at least one database (Additional file 3: Fig. S1).

With RNA from the same 24 samples used for transcriptome generation, qRT-PCR was conducted for nine flowering-related unigenes identified through RNA-seq, including COP1 (Unigene0031506), ZTL (Unigene0041339), FKF1 (Unigene0038380), GI (Unigene0051409), ELF3 (Unigene0051761), PRR1 (Unigene0045946), PRR7 (Unigene0003564), PRR5 (Unigene0047475), and LHY (Unigene0035686). The results of qRT-PCR showed that except for PRR5 (Unigene0047475), the expression patterns of the other eight genes were generally consistent with the RNA-seq data (Additional file 4: Fig. S2), indicating that the transcriptome data generated in this study were reliable and valid.

A total of 113 (LD7-vs.-SD7), 420 (LD10-vs.-SD10), 483 (LD13-vs.-SD13), and 464 (LD19-vs.-SD19) differentially expressed genes (DEGs) were obtained by comparing the LD and SD treatments (Additional file 5: Fig. S3; Additional file 6: Table S3). A total of 1,226 DEGs were identified from these four comparisons, of which five DEGs were shared by four comparisons, and 250 DEGs were present in at least one comparison. There were 110, 302, 288, and 276 stage-specific DEGs in LD7-vs.-SD7, LD10-vs.-SD10, LD13-vs.-SD13, and LD19-vs.-SD19, respectively (Additional file 5: Fig. S3).

**Functional classifications of DEGs**

MapMan is an effective tool for systematic analysis of plant transcriptome metabolic pathways and other biological processes [12]. We employed MapMan to overview transcriptional changes in regulatory, metabolic, and cellular response-related genes (Additional file 7: Table S4). In “regulation overview”, more DEGs were detected in the other three comparisons contrasted with LD7-vs.-SD7, showing that the physiological and molecular characteristics after flower bud differentiation (SD10, SD13, and SD19) were significantly different from that before flower bud differentiation (SD7). In the IAA metabolic subclass, more DEGs were upregulated in LD19-vs.-SD19 compared with LD7-vs.-SD7, LD10-vs.-SD10, and LD13-vs.-SD13 (Additional file 8: Fig. S4). However, IAA content was the highest at SD0 and lowest on SD10 and continuously decreased from SD13 to SD19 (Fig. 3). Therefore, IAA was not a key factor mediating floral transition in *L. gratissima*. ABA metabolism-related DEGs were significantly upregulated in all four comparisons (Additional file 8: Fig. S4), and ABA levels were overall increasing in the process of floral transition (Fig. 3), demonstrating that ABA could promote floral transition in *L. gratissima*. In “minor CHO metabolism”, trehalose biosynthesis-related DEGs were only upregulated in LD7-vs.-SD7 (Additional file 9: Fig. S5). “Cellular response overview” showed that more development-related DEGs were upregulated in LD10-vs.-SD10 compared with the other three comparisons (Additional file 10: Fig. S6), indicating that these DEGs promoted floral transition in *L. gratissima*. 
Co-expression module analysis for DEGs

Weighted gene co-expression network analysis (WGCNA) is a systems biology method for analyzing the correlation relationships between genes in multiple samples [13]. In this study, the results of WGCNA showed that 1,226 DEGs in eight samples were clustered in 11 different co-expression modules (labelled with different colors) (Fig. 4a). It is noteworthy that four out of 11 co-expression modules significantly correlated with a single sample ($r > 0.9$, $p < 0.05$; Fig. 4b; Additional file 11: Table S5). For example, the largest module (black module) included 247 (20.15%) SD19-specific DEGs (Fig. 4b; Additional file 11: Table S5a).

We further conducted GO enrichment analysis on 11 co-expression modules, and only the greenyellow module was not significantly enriched for any GO terms (Additional file 12: Table S6). Some GO terms were specifically identified in only a single module. For example, 120 specific GO terms were identified in the black module, which mainly involved signal transduction and negative regulation of metabolic processes, and 34 module-specific GO terms were identified in the brown module, which was mainly associated with growth and development (Additional file 12: Table S6). However, several GO terms, including “response to organic substance” and “response to a stimulus”, appeared in multiple modules (Additional file 12: Table S6), indicating possible module-gene interactions. Overall, the extensively enriched GO terms showed that oral transition in *L. gratissima* activated functional activities at the systems level.

The 11 modules were divided into seven categories based on the correlations between modules (Fig. 4c). The heatmap showed that there was a high correlation between the blue, magenta, pink, and tan modules, in which the genes were highly expressed in SD7 and SD10 (Fig. 4b, c), and were significantly enriched in multiple GO terms involving secondary metabolite biosynthesis, signal transduction, and regulation of developmental processes (Additional file 12: Table S6).

Identification of DEG expression patterns associated with floral transition in *L. gratissima*

According to the above functional classifications and WGCNA of these DEGs, and flowering-related genes previously reported in model plants (such as *A. thaliana*) [5, 7], a total of 146 unigenes were identified as homologous genes related to floral transition in *L. gratissima*, involving several flowering pathways: sugar metabolism, hormone metabolism and signal transduction, photoperiod, ambient temperature, aging pathways, as well as floral integrator and floral meristem identity genes. Among these floral transition-related homologous genes, stage-specific DEGs, and common DEGs in LD7-vs.-SD7, LD10-vs.-SD10, LD13-vs.-SD13, and LD19-vs.-SD19 are listed in Additional file 13: Table S7.

The expression pattern of sugar signal-related homologs

The sugar signal pathway, which responds to the sugar budget in plants, is one of the important pathways mediating the transition from vegetative to floral meristems [5]. A total of 29 (19.86%) DEGs associated with sugar signal-related genes were identified, involving 23 sugar signal-related homologs.
These genes expressed differently in different development stages of *L. gratissima*. For example, *HEXOKINASE (HK)* homologs (Unigene0044869 and Unigene0044870) were significantly upregulated in LD7-vs.-SD7 and LD13-vs.-SD13, and a *BETA-GLUCOSIDASE 24 (BGLU24)* homolog (Unigene0013088) was significantly upregulated in LD10-vs.-SD10. Meanwhile, Unigene0009721 and Unigene0041893, homologs of *GALACTINOL SYNTHASE 2 (GOLS2)* and *RAFFINOSE SYNTHASE (RFS)* participating in raffinose synthesis, were upregulated in LD7-vs.-SD7. In addition, *TREHALOSE-6-PHOSPHATE SYNTHSE (TPS)* homologs (Unigene0019787, Unigene0024389, Unigene0013555, Unigene0054604, Unigene0004913, and Unigene0062998) were upregulated at various stages, and *SWEET16* homolog (Unigene0012661) was significantly upregulated in LD7-vs.-SD7 and LD10-vs.-SD10 (Fig. 5e; Additional file 14: Table S8). Hence, these genes may directly or indirectly participate in floral transition in *L. gratissima*.

The expression patterns of phytohormone metabolism and signal transduction homologs

Many studies have demonstrated that various phytohormones participate in the regulation of floral transition [7, 14-16]. A total of 20 (13.70%) DEGs associated with phytohormone metabolism were identified, and these involved 16 phytohormone metabolism homologous genes and were related to nine phytohormone metabolism pathways. Among these genes, *GIBBERELLIN 2-BETA-DIOXYGENASE 1 (GA2OX1)* homologs (Unigene0030732) and *GA2OX8* homologs (Unigene0073113), which are involved in GA metabolism, were significantly upregulated in LD10-vs.-SD10 and/or LD19-vs.-SD19. Meanwhile, Unigene0034382 (*CYP707A1* homolog) and Unigene0042754 and Unigene0042755 (*NCED1* homologs), respectively, encoding abscisic acid (ABA) 8'-hydroxylase 1 and nine-cis-epoxycarotenoid dioxygenase, were significantly upregulated in LD10-vs.-SD10. In addition, a homolog (Unigene0035296) of *YUC4*, encoding indole-3-pyruvate monoxygenase, which mediates auxin biosynthesis, was significantly upregulated in LD19-vs.-SD19. Additionally, genes encoding cytokinin (CK) dehydrogenase 7 (*CKX7*, Unigene0036599) and cytokinin dehydrogenase (*CYP735A1*, Unigene0029738) were significantly downregulated in LD19-vs.-SD19. *CYTOCHROME P450 745A1 (CYP745A1)* homolog (Unigene0036368), which participates in brassinolide (BR) biosynthesis, was upregulated in LD10-vs.-SD10 and LD19-vs.-SD19; the jasmonate (JA) metabolism-related *JASMONATE O-METHYLTRANSFERASE (JMT)* homolog (Unigene0020912) and the salicylic acid (SA) metabolism-related *UDP-GLYCOSYLTRANSFERASE 74F1 (UGT74F1)* homolog (Unigene0004033) were downregulated in LD19-vs.-SD19. A homolog of *CAROTENOID CLEAVAGE DIOXYGENASE 7 (CCD7*, Unigene0069349) involving in strigolactone (SL) biosynthesis, was also identified and showed significant downregulation in LD10-vs.-SD10 (Fig. 5c; Additional file 14: Table S8).

A total of 39 (25.85%) DEGs associated with phytohormone signal transduction were identified and involved 30 phytohormone signal transduction homologs that were associated with signal transduction for nine hormones. Among these DEGs, *GID1B* homologs (Unigene0032780, Unigene0032781, and Unigene0063035), encoding a gibberellin receptor, were upregulated in LD10-vs.-SD10, whereas an *RGL3* homolog (Unigene0071862), encoding a DELLA protein, was significantly downregulated in LD19-vs.-SD19. The ABA signal transduction-related *EID1-LIKE F-BOX PROTEIN 3 (EDL3)* homolog
(Unigene0018152) was upregulated in LD10-vs.-SD10, and SAUR71 homologs (Unigene0021953 and Unigene0025106), encoding auxin-responsive protein, were upregulated in LD13-vs.-SD13 and LD19-vs.-SD19. Moreover, in the CK signaling pathway, homologs of AHPs (Unigene0034629, Unigene0004315, and Unigene0034630), encoding histidine-containing phosphotransfer protein, were highly expressed in SD10, SD13, and SD19, and an ARR6 homolog (Unigene0049441), encoding a two-component response regulator, was upregulated in LD19-vs.-SD19. In addition, a BRI1 homolog (Unigene0024976) in the BR signaling pathway was significantly upregulated in LD7-vs.-SD7; homologs of MYC4 (Unigene0009399) and TIFYs (Unigene0022959 and Unigene0019294) in the JA signaling pathway were upregulated in LD10-vs.-SD10; a DWARF14 (D14) homolog (Unigene0028658), participating in SL signal transduction, was upregulated in LD7-vs.-SD7 but downregulated in LD19-vs.-SD19 (Fig. 5d; Additional file 14: Table S8).

Expression patterns of genes associated with photoperiod pathways

The photoperiod flowering pathways in plants include the photosensory pathway, the circadian clock, and the systemic effector [17]. A total of 10 (6.84%) photoperiod-related homologs were identified. Among these homologs, CHLOROPHYLL A-B BINDING PROTEIN (CAB40, Unigene0075619) was downregulated in LD19-vs.-SD19, whereas CONSTANS-LIKE 12 (COL12, Unigene0039617) and FD (Unigene0027311) were upregulated in LD13-vs.-SD13. Meanwhile, homologs of the FLAVIN-BINDING KELCH REPEAT F-BOX PROTEIN 1 (FKF1, Unigene0038380) and the PSEUDO-RESPONSE REGULATOR 7 (PRR7, Unigene0003564) were both downregulated in LD13-vs.-SD13, and a LUX homolog (Unigene0011585) was downregulated in LD10-vs.-SD10, whereas homologs of the NF-Ys (Unigene0025001, Unigene0002375, and Unigene0033157) were upregulated at one or more stages (Fig. 5a; Additional file 14: Table S8).

Expression patterns of genes associated with the ambient temperature pathway

Plant responses to photoperiod and temperature are coupled [18, 19]. The photoperiod-induced floral transition could also affect the expression of a series of ambient temperature-related genes in plants. We identified 28 (19.18%) ambient temperature-related DEGs involving 18 homologs, primarily including the HEAT STRESS TRANSCRIPTION FACTORS (HSFs), HEAT SHOCK PROTEIN/COGNATE (HSPs), and PEARL1, most of which were highly expressed at several stages under LD (Fig. 5b; Additional file 14: Table S8).

Expression patterns of aging pathway-related, floral integrator, and floral meristem identity genes

The aging pathway is an endogenous flowering pathway in plants [20]. SQUAMOSA PROMOTER-BINDING-LIKE PROTEIN 4 (SPL4) homologs (Unigene0024429 and Unigene0024430) in the aging pathway were upregulated in LD10-vs.-SD10, LD13-vs.-SD13, and LD19-vs.-SD19 (Fig. 5f; Additional file 14: Table S8).
Floral integrators combine environmental and endogenous signals to mediate flowering in plants [5]. The floral integrator gene SUPPRESSOR OF OVEREXPRESSION OF CONSTANS 1 (SOC1) homologs (Unigene0039572 and Unigene0039575) were upregulated in LD10-vs.-SD10, LD13-vs.-SD13, and LD19-vs.-SD19, whereas the AGAMOUSLIKE24 (AGL24) homolog (Unigene0049016) was downregulated in LD19-vs.-SD19 (Fig. 5g; Additional file 14: Table S8).

Genetic networks regulating floral transition in plants ultimately activated floral meristem identity genes, thereby causing the transformation from vegetative to floral meristems [21]. A total of 15 (10.27%) related DEGs were identified, involving nine floral meristem identity genes (Additional file 14: Table S8). Among these genes, homologs of AGL8/FUL (AGL8, also known as FUL; Unigene0019277, Unigene0004737, Unigene0042052, Unigene0042053, and Unigene0042058), APETALA 1 (API; Unigene0019278, Unigene0019279, and Unigene0031106), LEAFY (LFY; Unigene0030979 and Unigene0030980), and SEPALLATAs (SEPs; Unigene0000607, Unigene0034045, and Unigene0025130) were upregulated in one or more developmental stages, whereas homologs of SHORT VEGETATIVE PHASE (SVP, Unigene0049018) and TERMINAL FLOWER 1 (TFL1, Unigene0026727) were downregulated in LD19-vs.-SD19, LD10-vs.-SD10, and LD13-vs.-SD13 (Fig. 5h; Additional file 14: Table S8).

**Co-expression network of floral transition-related genes**

A co-expression network constructed using 126 floral transition-related DEGs with edge weights > 0.1 showed ten hub genes with great connectivity, including homologs of GLYCERALDEHYDE-3-PHOSPHATE DEHYDROGENASE (GAPDH, Unigene0005846), AKR1B1 (Unigene0076531), PKM (Unigene0073914), ENOLASE 1 (ENO1; Unigene0011083 and Unigene0011084), MED37E (Unigene0051600), L-LACTATE DEHYDROGENASE A CHAIN (LDHA, Unigene0009368), HSP83A (Unigene0031524), FUL (Unigene0042052), and SEP4 (Unigene0025130) (Additional file 15: Fig. S7). The genes with the highest network degree were GAPDH (Unigene0005846), AKR1B1 (Unigene0076531), and PKM (Unigene0073914), which participated in sucrose and starch catabolism (Additional file 14: Table S8).

**Discussion**

The timing of floral transition in plants is jointly regulated by internal and external environmental cues, of which photoperiod is one of the major environmental factors that affect floral transition in plants [5, 22]. *L. gratissima* is a horticultural ornamental plant with high development potential, and therefore elucidating the molecular mechanism of its SD photoperiod-induced floral transition is important to its year-round production for commercial purposes. In this study, we conducted transcriptome sequencing of *L. gratissima* shoot apexes at four stages under LD and SD treatments. A total of 79,870 unigenes were obtained by de novo assembly, of which 49.02% were successfully annotated. Currently, there is no report on *L. gratissima* transcriptome assembly and our assembled and annotated transcriptome of *L. gratissima* provides a valuable genetic resource for breeding this species.

**Sugar signal mediates floral transition in *L. gratissima***
Sugars are an important energy source and participate in oral transition in plants as important signaling molecules [23, 24]. Leaves are the primary organ of sugar synthesis in plants, and shoot apical meristems (SAMs) are the sites of sugar mobilization and consumption, both of which form an important source-sink unit [25]. Floral transition in plants is not only directly associated with sugar content from source and sink but is also regulated by sugar transport [26]. Previous studies have indicated that source-sink regulation could be achieved by the interaction between the bidirectional sugar transporter SWEET and the FT-like protein [27]. In this study, SWEET16 (Unigene0012661) was significantly upregulated in LD7-vs.-SD7 and LD10-vs.-SD10 (Fig. 5e; Additional file 14: Table S8), indicating that SWEET participated in sucrose transport during floral transition in L. gratissima. However, soluble sugars in SAMs decreased from SD0 to SD19 (Fig. 3), which is not consistent with the expression profile of genes associated with sucrose metabolism. We speculated that SAM only synthesized limited levels of soluble sugar but SWEET16 (Unigene0012661) expression in SAMs was only high at SD7 and SD10, and its expression level decreased as SD treatment duration increased (Fig. 5e; Additional file 14: Table S8), subsequently causing a decrease in the rate of the sucrose transport from leaves to SAMs; this suggests that sucrose only acts as an energy source in floral transition in L. gratissima.

Trehalose-6-phosphate (T6P) is a component of the plant sugar signaling system and has important effects on flowering and development [28]. In A. thaliana, the T6P pathway in leaves induced the expression of the florigen gene FT in the photoperiodic pathway to affect floral transition, whereas in SAMs, the expression of SPL in the aging pathway was controlled by the T6P pathway to directly affect the expression of floral transition-related genes [29]. Therefore, the T6P pathway is an important signal that coordinates flowering induction. In this study, except for the T6P synthase homolog TPS (Unigene0013555) that was downregulated in LD19-vs.-SD19, other TPSs were upregulated at one or more stages during floral transition in L. gratissima (Fig. 5e; Additional file 14: Table S8), showing that TPS homologs participate in floral transition in L. gratissima and the T6P signaling pathway is significantly enhanced during floral transition. SPL4 was also highly expressed at SD10, demonstrating that T6P in L. gratissima SAM promoted floral transition by regulating SPL4 expression. HK acts as a catalytic enzyme to catalyze hexose phosphorylation, as well as a glucose signal sensor mediating the interaction between the glucose signaling pathway and the ABA signaling pathway to regulate plant development [30, 31]. In this study, HK homologs (Unigene0044869 and Unigene0044870) were upregulated in LD7-vs.-SD7 and LD13-vs.-SD13 (Fig. 5e; Additional file 14: Table S8). We speculate that HK mainly catalyzed hexose phosphorylation to provide an energy source for initiating floral transition at SD7 and acted as a glucose signal sensor to participate in L. gratissima flower development at SD13.

In summary, the sugar metabolism-related genes TPS and HK entered the flowering regulatory network through the sugar signaling and hormone signaling pathways to regulate floral transition in L. gratissima. Phytohormones regulate floral transition in L. gratissima

Phytohormones play important regulatory roles in plant development and the mechanisms of their participation in floral transition in many plants are extensively studied [7, 14-16]. However, the complex
hormone regulatory network of floral transition in perennial woody plants remains unclear. We studied the regulatory patterns of hormones that participate in floral transition in *L. gratissima*.

As one of the most important phytohormones, the function of GA in regulating floral transition is mainly achieved through maintaining GA homeostasis and regulating the levels of DELLA, a growth inhibitor in the GA signaling pathway [7]. GA homeostasis in plants is maintained through coordinating the expression levels of the GA biosynthesis genes, such as *GA3OXs* and *GA20OXs*, and the catabolic enzyme genes *GA2OXs*, thereby regulating floral transition [7, 32]. In this study, homologs of *GA2OX1* (Unigene0030732) and *GA2OX8* (Unigene0073113) were both upregulated in LD10-vs.-SD10 (Fig. 5c; Additional file 14: Table S8). GA2OXs can catalyze the 2β-hydroxylation of bioactive GAs (such as GA$_1$, GA$_3$, GA$_4$, and GA$_9$), resulting in decreased levels of bioactive GAs [33]. This may be one of the reasons for low GA$_3$ content in shoot apexes of *L. gratissima*. The main components of GA signaling include the GA receptor GID1B and the growth inhibitors, DELLAs [7]. When GA concentrations increase, the DELLA protein forms a GA-GID1B-DELLA complex that undergoes degradation by the ubiquitination pathway, thereby activating the expression of downstream genes [7]. The GA signaling pathway mainly promotes floral transition by inducing the expression of *SOC1* and *LFY* [7, 34, 35]. In this study, *RGL3* (Unigene0071862) encoding DELLA had low expression in SD10, SD13, and SD19 (Fig. 5d; Additional file 14: Table S8). In contrast, *SOC1* (Unigene0039572 and Unigene0039575) and *LFY* (Unigene0030979) were highly expressed in SD10, SD13, and SD19 (Fig. 5g, h; Additional file 14: Table S8). This showed that low levels of the DELLA protein RGL$_3$ could induce the expression of *SOC1* and *LFY*. Additionally, the GA receptor genes *GID1Bs* (Unigene0032780, Unigene0032781, and Unigene0063035) were upregulated in LD10-vs.-SD10 (Fig. 5d; Additional file 14: Table S8), further demonstrating that GA promotes floral transition in *L. gratissima*. However, it may not be GA$_1$, GA$_3$, GA$_4$, or GA$_9$ but other active GAs that took effect. Previous studies indicated that GA has a promoting effect in floral transition in *A. thaliana* [7, 9], whereas GA was found to negatively regulate floral transition in woody plants [8]. GA regulation of floral transition in *L. gratissima* (a woody plant) is similar to herbaceous plants but not woody plants. This unique regulation pattern may be affected by many endogenous and environmental factors, which needs to be further studied in the future.

Other hormones also have some effects in regulating floral transition in *L. gratissima*. ABA is usually considered a stress-related hormone, but it also plays an important role in plant development [36]. However, there is still debate over the role of ABA in floral transition because both promoting and inhibitory effects were reported [16, 37]. In this study, the ABA synthase gene *NCED1* (Unigene0042754 and Unigene0042755) and the catabolic gene *CYP707A1* (Unigene0034382) were both upregulated in LD10-vs.-SD10 (Fig. 5c; Additional file 14: Table S8), and the ABA content in the SAMs was maintained at high levels that initially increased from SD0 to SD13 and subsequently declined, reaching its peak on SD13 (Fig. 3). ABF2 is a bZIP transcription factor that binds to ABA. It is also an important component of the glucose signaling pathway [38]. In this study, *ABF2* (Unigene0046988) was highly expressed in SD10, and likely participated in floral transition in *L. gratissima* by mediating the ABA and glucose signaling pathways. In the ABA core signaling pathway, the protein phosphatase PP2C (ABI1, ABI2, HAB1, and
PP2CA/AHG3) acts as a key negative regulatory factor, which has important regulatory effects on the activation of ABA signaling [39]. When ABA levels increase in plants, the ABA receptors PYR1/PYLs/RCARs bind and inhibit the phosphatase activity of PP2C, thereby activating the ABA signaling pathway [39]. In this study, PYL4 expression was high in SD13, whereas PP2C expression peaked on SD10 but was also high on SD13 (Fig. 5d; Additional file 14: Table S8), suggesting that the activation of the ABA signaling pathway mainly occurred on SD13 and that ABA promoted flower development in *L. gratissima* through the core signaling pathway. EDL3 is a positive regulator of the ABA signal cascade reactions, and it positively regulates the expression of the central component *CONSTANS* (*CO*) in the photoperiod pathway to regulate floral transition [40]. In this study, the expression of *EDL3* and *COL12* in the photoperiodic pathway peaked on SD10 (Fig. 5a, d; Additional file 14: Table S8), showing that ABA promoted floral transition in *L. gratissima* by interacting with EDL3 to induce *COL12* expression.

Plant growth depends on the continuous function of meristems, and CKs have positive effects on SAMs. In this study, the cytokinin synthase gene *LOGs* and the zeatin O-glucosyltransferase gene *ZOG1* were mainly upregulated in LD10-vs.-SD10 and LD13-vs.-SD13 (Fig. 5c, d; Additional file 14: Table S8). It is known that zeatin O-glucoside plays important roles in the transport and storage of CKs [41]. On the other hand, the *trans*-zeatin synthase gene *CYP735A1* and the cytokinin oxidase/dehydrogenase gene *CKX7* were downregulated in LD19-vs.-SD19 (Fig. 5c, d; Additional file 14: Table S8). Zeatin promotes cell division and has an important role in the early stages of flower bud development and cell division. This is likely the reason zeatin content gradually decreased from SD0 to SD19 (Fig. 3). The CK signaling pathway mainly cross talks with AGAMOUS (AG) to regulate SAM differentiation and maintenance [42]. RPN12A participates in ATP-dependent ubiquitinated protein degradation, which may inhibit the degradation of one or more factors in CK signaling and balance the proliferation rate of cells during bud development [43]. In this study, *AHPs*, which are key components in the cytokinin two-component signaling system [44], were highly expressed mainly at SD10, SD13, and SD19; *ARR6*, which is a CK responsive regulator [44], was significantly upregulated in LD19-vs.-SD19, and *RPN12A* was upregulated in LD13-vs.-SD13; moreover, *AGL8* was highly expressed in SD10, SD13, and SD19 (Fig. 5d, h; Additional file 14: Table S8), demonstrating that CK promotes floral transition and flower development in *L. gratissima* indirectly through the effects of *AGL8*.

In the JA signaling pathway, JAZ (jasmonate-ZIM domain, TIFY family) and *MYC2/3/4* regulate floral transition in plants [7]. In this study, *TIFYs* and *MYC4* were upregulated in LD10-vs.-SD10 (Fig. 5d; Additional file 14: Table S8), showing that the JA signaling pathway promotes floral transition in *L. gratissima*. In SL signaling pathway, *D14* negatively regulates SL signals as an SL receptor [45]. In this study, *D14* (Unigene0028658) expression was high at the early stage of SD treatment, and as treatment duration increased, its expression level decreased (Fig. 5d; Additional file 14: Table S8), which may have been caused by negative feedback regulation of SL signals by *D14*, thereby regulating SL changes during floral transition in *L. gratissima*. *CCD7* is a key enzyme in SL biosynthesis [7]. Compared with the LD treatment, *CCD7* (Unigene0069349) expression was lower in response to SD treatment and was significantly downregulated in LD10-vs.-SD10 (Fig. 5c; Additional file 14: Table S8), suggesting that SL may inhibit floral transition in *L. gratissima*. In contrast to the results of this study, recent studies have
shown that SL inhibits melatonin synthesis, thereby inducing floral transition in *A. thaliana* in an FLC-dependent manner [15]. As *L. gratissima* is a perennial woody plant, there may be differences in SL regulatory mechanisms in floral transition compared with *A. thaliana*, which requires further in-depth studies.

YUC-mediated auxin biosynthesis is vital for the formation of floral organs [46]. In this study, YUC4 was upregulated in LD19-vs.-SD19 (Fig. 5c; Additional file 14: Table S8), whereas IAA content was lowest in SD10 and continuously decreased from SD13 to SD19 (Fig. 3), whereas the auxin response gene *SAUR7* was upregulated in LD13-vs.-SD13 and LD19-vs.-SD19. These results showed that auxin does not participate in regulating floral transition in *L. gratissima* but instead has positive effects on the formation of floral organs.

These hormones interacted with other flowering regulation pathways to further ensure that *L. gratissima* rapidly responded to changes in environmental and endogenous signals to precisely regulate flowering time.

**Flowering pathways during floral transition in *L. gratissima***

The photoperiod pathway is involved in plant response to changes to day length and circadian rhythm, making it one of the most important flowering regulation pathways. In the photoperiod pathways of many plants, the bZIP transcription factor FD forms a transient complex in SAMs with the FT protein from leaves to induce the expression of floral meristem identity genes, thereby promoting floral transition [47]. In this study, *FD, AP1, FUL, and AGL8* were highly expressed in SD10 and SD13 (Fig. 5a, h), demonstrating that the FD protein directly or indirectly induced the expressions of *AP1, FUL,* and *AGL8,* thereby promoting floral transition in *L. gratissima*. CO is an important regulatory factor in the photoperiod pathway, and the expression of *CO* is regulated by a photoreceptor and circadian rhythm in *A. thaliana*, and when the expression rhythm of *CO* is consistent with the external photoperiod, expression of the downstream gene *SOC1* is activated [48]. In this study, *CO-like 12 (COL12)* was upregulated in LD13-vs.-SD13 (Fig. 5a; Additional file 14: Table S8), showing that the effects of COL12 in flower development in *L. gratissima* were similar to those of *CO* in *A. thaliana*.

The transcription factor LUX is one of the components of evening complex (EC) complex in circadian rhythm, and forms the HOS15-EC-HDA9 histone-modifying complex in *A. thaliana* to inhibit *GI* transcription, thereby inhibiting photoperiod-dependent flowering [49]. In this study, *LUX* was downregulated in LD10-vs.-SD10 (Fig. 5a; Additional file 14: Table S8), indicating that LUX had inhibitory effects on floral transition in *L. gratissima*. PRR7 positively regulates *CO* expression to promote floral transition in long-day plants, whereas the *PPR7/PRR3* genes delay floral transition by inhibiting *CO* expression in short-day plants [50]. In this study, *PPR7* was downregulated in LD13-vs.-SD13 (Fig. 5a; Additional file 14: Table S8), showing that PRR7 inhibits floral transition in *L. gratissima*, which was similar to the other short-day plants. In *A. thaliana*, FKF1 could degrade CDF1 (factor inhibiting *CO* transcription) to regulate *CO* expression and could directly bind to C0, or inhibit *COP1* to stabilize *CO* expression, thereby promoting flowering [51]. However, *FKF1* was downregulated in LD13-vs.-SD13 (Fig.
5a; Additional file 14: Table S8), which was not consistent with COL12 expression. This indicated that FKF1 inhibited floral transition in *L. gratissima* and does not interact with COL12, but other mechanisms may be present that require further study. NF-Ys interacts with CO in the photoperiod pathway to directly regulate SOC1 transcription [34]. In this study, NF-Ys, COL12, and SOC1 were highly expressed in SD10 and SD19 (Fig. 5a, h), showing that NF-Ys may interact with COL12 in the photoperiod pathway in *L. gratissima* to induce SOC1 expression, thereby positively regulating floral transition and flowering development in *L. gratissima*.

Previous studies showed that ambient temperature-associated EARLI1 regulated critical genes in the LD photoperiod pathway in *A. thaliana* to promote FLC expression and delayed flowering time [52]. In contrast, pEARLI1 was upregulated in LD13-vs.-SD13 and LD19-vs.-SD19 in this study (Fig. 5b; Additional file 14: Table S8), indicating that pEARLI1 promoted floral transition and flower development in *L. gratissima*.

In *A. thaliana*, age signals negatively regulate miR156 levels to promote SPL accumulation [20]. At SAMs, SPLs target FUL and SOC1 or directly regulate AP1 transcription to promote flowering [53]. In this study, SPL4 was upregulated in LD10-vs.-SD10, LD13-vs.-SD13, and LD19-vs.-SD19 (Fig. 5f; Additional file 14: Table S8), which was consistent with the expression patterns of SOC1, FUL, and AP1 (Fig. 5g, h), indicating that the aging pathway promoted floral transition and flower development in *L. gratissima* through SPL4-induced expression of FUL, SOC1, and AP1.

The floral integrators SOC1 and AGL24 integrate various flowering signals from photoperiod, temperature, hormone, and age-related signals to activate or inhibit downstream floral meristem identity genes, and ultimately lead to the transformation of vegetative to floral meristems in plants [5]. SOC1 can be indirectly activated by CO [54]. At SAMs, when SOC1 is activated, SOC1 and AGL24 form a heterodimer to directly activate LFY [55]. In this study, SOC1, AGL24, and LFY were highly expressed in SD10, showing that SOC1 and AGL24 can jointly promote LFY at this period to promote floral transition in *L. gratissima*. During early flower development, AP1 activates A function to inhibit SOC1 and AGL24 expression to prevent flowering reversion [54]. In SD19, AGL24 and SOC1 expression decreased and AP1 expression increased (Fig. 5g, h). These changes may prevent differentiated floral meristems from undergoing flowering reversion.

SEPs are important regulatory factors during flower development and form a heterodimer with AP1 to regulate genes during floral meristem development [56]. In this study, SEPs were highly expressed in SD10, SD13, and SD9, which was consistent with AP1 expression (Fig. 5h), showing that AP1 mediated positive regulation of floral transition and early flower development in *L. gratissima* by SEPs. In Arabidopsis, SVP is a flowering inhibitor and plays a role in floral transition by directly inhibiting SOC1 expression at SAMs and leaves [57]. In this study, SVP had low expressions in SD10, SD13, and SD19, whereas SOC1 expression was high (Fig. 5g, h), indicating that low levels of SVP induced SOC1 expression to promote floral transition and flower formation in *L. gratissima*. 
TFL1 is a key regulatory factor of oral transition and inflorescence meristem development in *A. thaliana*. TFL1 and FT have highly conserved amino acid sequences but opposite gene functions: FT promotes flowering, whereas TFL1 inhibits flowering [58]. Previous studies showed that TFL1 negatively regulated transcription of the target gene *FD*, thereby regulating the flowering time and inflorescence meristem development [59]. In this study, *TFL1* had low expression at SD10 and SD13, which is the opposite of *FD* expression (Fig. 5a, h), indicating that low levels of TFL1 promoted *FD* expression and, therefore, floral transition in *L. gratissima*.

Fig. 6 shows the hypothetical model of the regulatory network of SD photoperiod-induced oral transition in *L. gratissima*, involved in the regulation of multiple flowering signals in floral transition, including signals for photoperiod, phytohormones (GA, ABA, CK, JA, and SL), sugar, ambient temperature, age, and floral integrator and floral meristem identity genes.

**Conclusions**

Our study enables a comprehensive understanding of the gene expression patterns occurring during SD photoperiod-induced floral transition in *L. gratissima*. The histological, endogenous substance contents, and differential gene expression analyzes showed that short-day photoperiod activated systemic responses in *L. gratissima* and induced the generation of flowering signals in the photoperiod pathway. Furthermore, a complex regulatory network, including GA, ABA, CK, JA, and SL signals, sugar signals, and temperature and age signals was formed through the integration of SOC1 and AGL24. The outcomes of this study will aid in understanding flowering time regulation in *L. gratissima* at the molecular level, provide theoretical guidance for achieving year-round production, and further provide a reference for understanding the regulatory mechanisms of flowering time in other woody plants.

**Methods**

**Plant materials, growth conditions, and light treatments**

*L. gratissima* cultivar ‘Xiangfei’ cuttings were obtained from the central Yunnan Plateau experimental station of Research Institute of Resources Insects, Chinese Academy of Forestry (Yunnan, China; 25°13′N, 102°12′E, 1826 m a.s.l.). In mid-December 2016, cuttings with two stem nodes and shoot apexes were planted in a mixed matrix (peat and perlite at a 3:1 ratio) and grown in an 18–25°C greenhouse under natural lighting. Cuttings with roots were transplanted into pots and maintained in the same greenhouse under natural lighting. To prevent these plants from being induced by SD photoperiod, shoot apical meristems (SAM) were removed from all plants when 2–3 new stem nodes were formed, and high-pressure sodium lamps were used for additional lighting during 22:00–02:00 (night-break treatment) (Fig. 1c). In addition, considering the effects of individual developmental age on flowering time [60], some plants were placed in the natural environment as controls and the time when flower bud differentiation occurred in these plants was used as the start time for photoperiod treatments. On 10 August 2017 (when flower buds began to appear in some natural control plants), plants with the same number of branches
longer than 5 cm were selected from among the night-break treatment plants, and were then subjected to
either LD (night-break treatment as described above) or to SD (10 h light/14 h dark) (Fig. 1d) for a further
90 days. Shoot apexes of the main branches of SD and LD plants were sampled during 09:00–11:30
every 3–5 d after the initiation of the photoperiod treatments. For each stage, 10–20 shoot apexes were
packed together into each of the 10 biological replicates, of which one biological replicate was rapidly
immersed into FAA fixative (50% ethanol: acetic acid: formaldehyde, 18:1:1) for morphological analysis,
whereas the remaining nine biological replicates were snap-frozen in liquid nitrogen and then stored at −
80°C for measurements of soluble sugar and endogenous hormone contents, as well as RNA extraction.

Morphological anatomical observations

Ten FAA-fixed shoot apexes of SD and LD plants at each stage were made into sections with a thickness
of 8–10 μm using paraffin section method [61], and were stained with safranin O-fast green, and then
were mounted with neutral resin. Finally, the process of bud development was observed under a Carl
Zeiss Axio Scope A1 Microscope (Carl Zeiss Microscopy GmbH, Göttingen, Germany).

Measurements of soluble sugar and endogenous hormone contents

According to the anatomical observation results, samples from the SD treatment at five stages (0 d [SD0],
7 d [SD7], 10 d [SD10], 13 d [SD13], and 19 d [SD19]) close to flower bud differentiation (Fig. 2) were
selected for measurements of soluble sugar and endogenous hormone contents of three biological
replicates. For each of the three biological replicates from each stage, soluble sugar contents were
measured using sulfuric acid-anthrone colorimetric assays [62], and endogenous hormones (GA₃,
indoleacetic acid [IAA], ABA, and zeatin [ZT]) were quantified with high-performance liquid
chromatography-mass spectrometry (HPLC-MS; Aglient1290, Nanjing, China; AB 6500, Nanjing, China)
[63]. Before comparing changes in the soluble sugar and hormone contents among the five stages, the
Shapiro-Wilk test and Levene test were used to analyze the normality and homogeneity of variance of
each dataset. Because the four sets of data did not follow a normal distribution (% < 0.05), a Kruskal-
Wallis H test was employed for analysis of significant differences, and false discovery rate (FDR) was
used for the multiple testing correction of significant p-values. Additionally, the Tukey-Kramer method
was used for post-hoc testing of soluble sugar and hormone contents at the five stages. The above
analyzes were performed in the “car” and “stats” packages in R software and the data were expressed as
the mean ± SD.

Transcriptome sequencing and data analysis

Likewise, based on the anatomical observation results, samples from the SD and LD treatments at the
four stages (7 d [SD7 or LD7], 10 d [SD10 or LD10], 13 d [SD13 or LD13], and 19 d [SD19 or LD19]) close
to flower bud differentiation of SD plants (Fig. 2) were selected for RNA extraction. Total RNA extracted
from each of the three biological replicates was divided into two parts, of which one was used for RNA-
seq and the other was used for quantitative real-time PCR (qRT-PCR) validation. RNA was extracted with
the Plant total RNA Kit (Tiangen, Beijing, China) following manufacturer’s instructions. The cDNA library
construction and paired-end sequencing were conducted with an Illumina HiSeq™ 4000 (Illumina, San Diego, California, USA) at the Gene Denovo Biotechnology Company (Guangzhou, China). The generated raw reads were filtered by removing adapter sequences and ambiguous reads (N > 10%) and low-quality reads (more than 40% of bases with Q value ≤ 20) to obtain high-quality clean reads. Finally, the Trinity software [64] was used for de novo assembly of filtered clean reads to obtain unigenes.

All non-redundant unigenes were aligned with selected cut-offs of E-value ≤ 1e^-5 to six protein databases, including the NR (the NCBI non-redundant protein databases), KOG (EuKaryotic Orthologous Groups), KEGG (Kyoto Encyclopedia of Genes and Genomes), Swiss-Prot, eggNOG (evolutionary genealogy of genes: Non-supervised Orthologous Groups), and Pfam (Protein families database of alignments and hidden Markov models). Based on the NR annotation results, these unigenes were also annotated for GO (Gene Ontology) using the Blast2GO software [65], and then GO functional classification of unigenes was obtained by WEGO software [66].

**Quantitative real-time PCR analysis**

Quantitative real-time PCR (qRT-PCR) was conducted on nine flowering-related unigenes in this study, including COP1 (Unigene0031506), ZTL (Unigene0041339), FKF1 (Unigene0038380), GI (Unigene0051409), ELF3 (Unigene0051761), PRR1 (Unigene0045946), PRR7 (Unigene0003564), PRR5 (Unigene0047475), and LHY (Unigene0035686). To accurately measure gene expression levels, the ACT7/EF1-α combination obtained from the past screening was used as an internal reference gene for standardization and correction (Additional file 16: Supplementary Data). Primer3 software [67] was used to design specific primers for each gene (Additional file 17: Table S9). The KR106 FastQuantity RT Kit (with gDNase) (Tiangen, Beijing, China) was used for reverse transcription of 1 µg total RNA into cDNA according to the manufacturer’s instructions. The StepOnePlus™ Real-Time PCR System (Thermo Scientific, Wilmington, DE, USA) was used for qRT-PCR in a 20 µL reaction system, including 4 µL of 50 ng cDNA template, 10 µL of 2× qPCR Master Mix (Tiangen, Beijing, China), 0.4 µL each of 10 µM forward and reverse primers, and 5.2 µL ddH₂O. The qRT-PCR amplification conditions were as follows: pre-denaturation at 95°C for 90 s, followed by 40 cycles of denaturation at 95°C for 5 s, annealing at 60°C for 15 s, extension at 72°C for 20 s, followed by a final extension step at 72°C for 5 min. After amplification, a 65–95°C melting curve analysis was conducted to measure product specificity. The 2^-ΔΔCt method [68] was used to calculate the relative expression levels of the genes in the qRT-PCR experiment.

**Identification and functional enrichment of differentially expressed genes (DEGs)**

The Reads Per kb per Million reads (RPKM) method was used to evaluate unigene expression levels [69]. EdgeR package version 3.8 [70] was used for differential expression analysis of multiple comparisons, and the FDR method [71] was used for the correction of P-values. FDR < 0.05 and |log₂Ratio| > 1 were used as criteria to screen for DEGs between various groups (ratio refers to the RPKM ratios of SD7/LD7, SD10/LD10, SD13/LD13, and SD19/LD19).
The Mercator online tool (https://www.plabipd.de/portal/mercator-sequence-annotation) was employed for gene function predictions for the DEGs with a BLAST-CUTOFF of 50. The obtained mapping files were uploaded to MapMan version 3.6 [72] for the functional analysis of DEGs. Wilcoxon rank-sum test was used to analyze the Log₂Ratio of DEGs in each comparison before MapMan version 3.6 [72] was used for visualization of the results.

Co-expression network analysis

Weighted gene co-expression network analysis (WGCNA) [13] was employed to generate the co-expression network modules of DEGs. The parameter settings used were soft threshold = 20, minModuleSize = 30, TOMType = signed, mergeCutHeight = 0.25, and default values were used for the remaining parameters. The eigengene value of every module was calculated and the associations between every gene in eight samples were tested. KOBAS 3.0 [73] was used for GO enrichment analysis of genes in the clustering modules. Cytoscape version 3.7.1 [74] was used for visualization of the co-expression network.

List Of Abbreviations

ABA: abscisic acid; AG: AGAMOUS; AGL24: AGAMOUSLIKE24; AP1: APETALA1; BGLU24: BETA-GLUCOSIDASE 24; CAB40: CHLOROPHYLL A-B BINDING PROTEIN; CCD7: CAROTENOID CLEAVAGE DIOXYGENASE 7; CK: cytokinin; CO: CONSTANS; COL12: CONSTANS-LIKE 12; CYP734A1: CYTOCHROME P450 734A1; D14: DWARF14; DEG: differentially expressed gene; EC: evening complex; EDL3: EID1-LIKE F-BOX PROTEIN 3; FKF1: FLAVIN-BINDING KELCH REPEAT F-BOX PROTEIN 1; FT: FLOWERING LOCUS T; FUL: FRUITFULL; GA: gibberellin; GA2OX1: GIBBERELLIN 2-BETA-DIOXYGENASE 1; GOLS2: GALACTINOL SYNTHASE 2; HK: HEXOKINASE; HPLC-MS: high performance liquid chromatography-mass spectrometry; HSFs: HEAT STRESS TRANSCRIPTION FACTORS; HSPs: HEAT SHOCK PROTEIN/COGNATE; IAA: indole-3-acetic acid; JA: jasmonic acid; JMT: JASMONATE O-METHYLTRANSFERASE; LD: long day; LFY: LEAFY; NF-Y: nuclear factor Y; PRR7: PSEUDO-RESPONSE REGULATOR 7; RFS: RAFFINOSE SYNTHASE; SD: short day; SEP3: SEPALLATA3; SEPs: SEPALLATAs; SL: strigolactone; SOC1: SUPPRESSOR OF OVEREXPRESSION OF CONSTANS 1; SPL4: SQUAMOSA PROMOTER-BINDING-LIKE PROTEIN 4; SVP: SHORT VEGETATIVE PHASE; T6P: trehalose-6-phosphate; TFL1: TERMINAL FLOWER 1; TPS: TREHALOSE-6-PHOSPHATE SYNTHSE; UGT74F1: UDP-GLUCOSYLTRANSFERASE 74F1; WGCNA: weighted gene co-expression network analysis; ZT: zeatin

Declarations

Acknowledgments

The authors would like to thank Guangzhou Genedenovo Biotechnology Co., Ltd for assisting in RNA sequencing.

Author contributions
ZHL, YMW, and HM conceived and designed the study. XFL, YMW, JA, XJZ, YRC, and XXL conducted the experiment and collected the plant materials. XFL and YMW analyzed and interpreted the data. XFL and YMW wrote the manuscript. YMW, ZHL, and HM revised the manuscript. All authors read and approved the final version of the manuscript.

**Funding**

This work was funded by Key Research and Development Program of Yunnan Province (2018BB012 to Z.-H.L.), the Fundamental Research Funds for the Central Non-profit Research Institution of CAF (CAFYBB2017MB014 to Y.-M.W.), the Fundamental Research Funds for the Central Non-profit Research Institution of CAF (CAFYBB2019ZB007 to Z.-H.L.), and Ten Thousand Talent Program of Yunnan Province (grant to H.M.).

**Availability of data and materials**

The raw data of RNA sequencing from this study have been deposited into the NCBI Sequence Read Archive (SRA) database under the accession number SRP273656.

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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Figure 1

Features of Luculia gratissima ‘Xiangfei’ and the overview of greenhouses under two different photoperiods. a Whole plant of L. gratissima ‘Xiangfei’. b Flowers of L. gratissima ‘Xiangfei’. c Greenhouse under night-break treatment. d Greenhouse under short-day photoperiod.
Figure 2

Luculia gratissima morphological and histological characteristics, shoot apexes at five time points upon short-day treatment. a–c Vegetative buds in the undifferentiated stage (SD0 to SD7). d–f Bract primordial differentiation stage (SD10). a, b, d, g, h Histological images obtained from paraaffin-embedded sectioned samples (scale bar: 100 μm). e, j, k Histological images obtained from paraaffin-embedded sectioned samples (scale bar: 50 μm). c, f, i, l The external morphology of shoot apexes at different developmental
Luculia gratissima endogenous soluble sugar content and hormonal changes, shoot apexes at five stages upon short-day treatment. The y-axis shows soluble sugar and four hormones, and the x-axis
shows the average relative abundance of the endogenous soluble sugars and hormones. Colored columns represent different developmental stages. * 0.01 < p ≤ 0.05, ** 0.001 < p ≤ 0.01.

**Figure 4**

Weighted co-expression network analysis of 1,226 DEGs at four developmental stages of Luculia gratissima, short- or long-day treatments. a Hierarchical cluster tree showing the co-expression modules, with each tree leaf representing one gene. The major tree branches constitute 11 modules labelled by
different colors. b Heat map of gene relative expression of different modules (y-axis) in eight samples (x-axis). The normalized RPKM value for an individual gene at a given developmental stage is indicated in a green (low expression) to red (high expression) scale. c Eigengene network representing the relationships among the modules and the clinical trait weight. The hierarchical clustering dendrogram of the eigengenes shows the relationships among the modules, whereas the heat map shows the correlation between the different modules, with deeper red color representing a stronger correlation.

Figure 5

Expression profiles of genes associated with Lucilia gratissima oral transition at four developmental stages, short- or long-day treatments. Relative expression profile of (a) photoperiod pathway-related genes, (b) ambient temperature pathway-related genes, (c) phytohormone metabolism-related genes, (d) phytohormone signal transduction-related genes, (e) sugar signal-related genes, (f) aging pathway-related genes, (g) floral integrator-related genes, and (h) floral meristem identity genes. The normalized RPKM value for an individual gene at a given developmental stage is represented in a green (low expression) to red (high expression) scale.
Figure 6

Proposed regulatory network of short-day photoperiod-induced floral transition in Luculia gratissima. Colored fonts represent down-regulated (green) or up-regulated (red) genes.

Supplementary Files

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