Integrated mRNA and microRNA transcriptome variations in the multi-tepal mutant provide insights into the floral patterning of the orchid *Cymbidium goeringii*  

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**Abstract**  
**Background:** *Cymbidium goeringii* is a very famous traditional orchid plant in China, which is well known for its spectacular and diverse flower morphology. In particular, the multi-tepal mutants have considerable ecological and cultural value. However, the current understanding of the molecular mechanisms of floral patterning and multi-tepal development is limited. In this study, we performed expression profiling of both microRNA (miRNA) and mRNA from wild-type and typical multi-tepal-mutant flowers of *C. goeringii* for the first time, to identify the genes and pathways regulating floral morphogenesis in *C. goeringii*.

**Results:** Total clean reads of 98,988,774 and 100,188,534 bp were obtained from the wild-type and mutant library, respectively, and *de novo* assembled into 98,446 unigenes, with an average length of 989 bp. Among them, 18,489 were identified as differentially expressed genes between the two libraries according to comparative transcript profiling. The majority of the gene ontology terms and Kyoto Encyclopedia of Genes and Genomes pathway enrichment responses were for membrane-building and ploidy-related processes, consistent with the excessive floral organs and altered cell size observed in the mutant. There were 29 MADS-box genes, as well as a large number of floral-related regulators and hormone-responsive genes, considered as candidates regulating floral patterning of *C. goeringii*. Small RNA sequencing revealed 132 conserved miRNA families expressed in flowers of *C. goeringii*, and 11 miRNAs corresponding to 455 putative target genes were considered to be responsible for multi-tepal development. Importantly, integrated analysis of mRNA and miRNA sequencing data showed two transcription factor/microRNA-based genetic pathways contributing to the multi-tepal trait: well-known floral-related miR156/SPL and miR167/ARF regulatory modes involved in reproductive organ development; and the miR319/TCP4–miR396/GRF regulatory cascade probably regulating cell proliferation of the multi-tepal development.

**Conclusions:** Integrated mRNA and miRNA profiling data provided comprehensive gene expression information on the wild-type and multi-tepal mutant at the transcriptional level that could facilitate our understanding of the molecular mechanisms of floral patterning of *C. goeringii*. These data could also be used as an important resource for investigating the genetics of floral morphogenesis and various biological mechanisms of orchid plants.

**Keywords:** *Cymbidium goeringii*, Multi-tepal mutant, Floral transcriptome, MicroRNA, Floral patterning, MiR396
Background

*Cymbidium goeringii* Rchb. f., which belongs to the subgenus *Jensoa* of the genus *Cymbidium*, is one of the most horticulturally important and popular ornamental plants in the orchid family (Orchidaceae) [1]. It blooms in the winter from January to March and is fragrant. In China, *C. goeringii* is called the spring orchid and is regarded as a Spring Festival flower, and holds a strong position in the traditional flower market [1, 2]. As a typical ornamental plant, *C. goeringii* is characterized by highly specialized reproductive strategies and extremely diversified flowers [3–5], and commercially plays a very important role in world flower markets, especially in Japan, Korea and Southeast Asia. Various mutations occur frequently in this orchid family, which greatly diversify the floral morphology and provide substantial commercial value. However, functional genomic studies and the gene discovery associated with floral pattern regulation remains greatly limited in *C. goeringii* [6].

Floral patterning, in terms of floral organ number, arrangement and initiation timing, has been well studied, especially in *Arabidopsis thaliana* and *Antirrhinum majus* [7–9]. Genes controlling floral organ identity have been identified through the genetic analysis of homeotic mutants, leading to the ABCDE model, in which five classes of regulatory genes (A, B, C, D and E) work in a combinatorial manner to confer the organ identities of the four whorls [10]. Among them, class A genes, such as *APETALA1* (*API*) in *Arabidopsis*, specify the outermost floral organs, the sepals. Class B genes, such as *APETALA3* (*AP3*) and *PISTILLATA* (*PI*), specify petals and male organs in concert with class A and C genes, respectively. Class C genes, such as *AGAMOUS* (*AG*), specify the inner-most floral organs, the carpels. The action of the class A, B and C genes is necessary for the specification of organ identity, and mutations in these genes produce homeotic transformations of one organ type into another. For example, the flowers of *Arabidopsis* *ag* mutants have no stamen or carpel, and lose the ability to terminate meristematic activity. As a result, the mutant generates multi-petal flowers, which have a repeated structure of sepal-petal-petal, including tens of petals [11, 12].

Although the ABCDE model is applicable to most flowering plants, in orchids it shows modifications in the expression domains of the class B *AP3* and *PI-like* genes and of the class C and E genes. The ‘orchid code’ model derived by Mondragon-Palomino and Theissen revealed that orchids typically have four *AP3/DEF-like* genes, representing the ancient gene Clades 1, 2, 3 and 4. With high levels of Clades 1 and 2, and low levels of Clades 3 and 4, gene expression specifies inner lateral tepals, whereas lip development requires low levels of Clades 1 and 2 expression and high levels of Clades 3 and 4 expression [13, 14]. Similarly, the orchid P-code model, which was derived primarily from *Oncidium* orchids, indicates that the higher-order heterotetrameric sepal/petal complex (OAP3-1/OAGL6-1/OAGL6-1/OPI) specifies sepal/petal formation, whereas the lip complex (OAP3-2/OAGL6-2/OAGL6-2/OPI) is exclusively required for lip formation [15]. However, beyond the hypothetical model of orchid code, we do not know exactly which genes are responsible for the subsequent floral organ specification or the mechanism underlying morphological diversity during orchid flower development. Compared with other plants, few reports have investigated the multi-tepal development of orchids [16]. Especially for the genus *Cymbidium*, available genomic resources are limited. This genetic data is insufficient for elucidating the molecular mechanism of floral regulation in *C. goeringii* [17–19].

Despite the genetic regulation of floral patterning, microRNAs (miRNAs)—which are small (18–25 nt) non-coding RNAs that control the epigenetic regulation of gene expressions—play crucial roles in different processes, from leaf and root morphogenesis to floral induction, organ formation, reproduction and stress response [20, 21]. They originate from single-stranded precursors (pre-miRNAs) able to self-pair and form hairpin structures. In plants, the Dicer1-like RNAse III enzyme excises the miRNA/miRNA* duplex from the pre-miRNA [22]. Subsequently, the guide strand of the duplex is recruited by an RNA-induced silencing complex, whose core component is the protein ARGONAUTE1. This complex permits the interaction between the miRNA and its target mRNA, thereby regulating its expression through mRNA cleavage, translational repression, or epigenetic modifications [22, 23]. In recent years, advances in next-generation sequencing techniques have prompted a plethora of studies on miRNAs, in both model and non-model species, and have led to the development of specific *in silico* analysis tools. Increasing numbers of studies on non-model plant species have highlighted the evolutionary conservation of a large number of miRNA families and the existence of taxon-specific ones. Recently, a few studies have also examined miRNAs in orchids, which are characterized by highly diversified floral structures and pollination strategies. Examples such as *Phalaenopsis aphrodite*, *Erycina pusilla* and *Dendrobium officinale* belonging to the sub-family *Epidendroideae* have been sequenced [24–27]. Both conserved and novel miRNAs have been identified in these species. By comparison, miRNA studies in the genus *Cymbidium* are very limited [28].

Here, we examined a typical multi-tepal mutant of *C. goeringii*, the cultivar ‘Yuhudie’, which continues to produce sepals and petals in the inner whorl. Compared to the wild-type plant, the gynostemium is replaced by a new emerged flower and both the lips and gynostemium
are misshapen in ‘Yuhudie’. From the computational analysis of the floral mRNA and miRNA expression levels based on deep sequencing data, we identified 98,446 unigenes and 226 conserved miRNAs that were expressed in flower. In total, 18,489 differentially expressed genes (DEGs) and 11 miRNA families corresponding to 455 putative target genes were probably related to the multi-tepal mutation. On this basis, we found a great number of floral-related and plant hormone-responsive genes contributing to multi-tepal development. Importantly, integrated analysis of mRNA and miRNA sequencing data showed that the transcription factor/microRNA-based genetic pathways—including floral-related miR156-SPL and miR167-ARF regulatory modes as well as the miR319/TCP4–miR396/GRF regulatory cascade—were involved in multi-tepal development. These results shed light on the regulatory mechanism of multi-tepal development of C. goeringii, and provide a base for future genome-wide orchid biology and biotechnology research.

Results

Morphology of the C. goeringii multi-tepal mutant

The wild-type flowers of C. goeringii have three sepals in the first whorl and three petals in the second whorl. Together these are called the tepals. Two of the petals are similar to each other and resemble unmodified sepals, while the third is highly evolved, with an ovate to triangular shape, and it is called the labellum (or lip). The male and female reproductive organs are highly fused to form a gynostemium, which evolved through the complete fusion of the style, stigma and staminal filament, and has four pollinia on a semi-circular viscidium (Fig. 1a & b). In comparison, the mutant has new flowers instead of a single gynostemium in the center, and continues to produce sepals and petals centripetally. Although they are morphologically similar to those of the wild-type, these newly emerged outgrowths are considerably smaller (Fig. 1c–e). The normal sepal and petal have diameters of ~1.23 and ~1.31 cm, respectively, and the total length of them ranges from ~2.78 to ~3.68 cm, whereas the newly emerged sepal and petal in the mutant are only 0.52–0.95 cm in diameter and the outgrowth’s lengths range from ~1.08 to ~3.38 cm (Table 1). On average, mutant petals are 28% shorter and almost 50% narrower than wild-type petals at anthesis. This difference in size is mainly due to a difference in cell size between them. The mutant shows about 30% smaller cell diameter than that of the wild-type (Additional file 1). Moreover, the lips are misshaped in the mutant, and the gynostemia are also mostly fused to the margin of C. goeringii, as illustrated in Fig. 1c–e.

![Fig. 1 Flower morphology of C. goeringii wild-type plant ‘Songmei’ and the multi-tepal mutant ‘Yuhudie’](image)
and 33,966 (34.50%) unigenes were aligned to a protein functional identifications. Among them, 46,342 (47.07%) more than half of the assembled sequences had putative one significant match with an existing gene model in public databases. A total of 78,175 unigenes had at least their similarities with known or putative annotations in unigenes is shown in Additional file 2.

The size distribution of the assembled transcripts and 1000 bp, which represented 19.1% of the total (Table 2). Of the putative unigenes, 23,137 were longer than 98,446 unigenes with a mean length of 989 bp, a N50 de novo were pooled together and unknown nucleotides (greater than 5%), the clean reads reads, reads containing adapters and reads containing exclusively. After filtering, and removing the low quality reads, reads containing adapters and reads containing unknown nucleotides (greater than 5%), the clean reads were pooled together and de novo assembled into 98,446 unigenes with a mean length of 989 bp, a N50 length of 1519 bp and a total length of 97,314,639 bp. Of the putative sequences, 23,137 were longer than 1000 bp, which represented 19.1% of the total (Table 2). The size distribution of the assembled transcripts and unigenes is shown in Additional file 2.

The entire set of unigenes was annotated on the basis of their similarities with known or putative annotations in public databases. A total of 78,175 unigenes had at least one significant match with an existing gene model in searches using the BLAST algorithm (Table 3). Thus, more than half of the assembled sequences had putative functional identifications. Among them, 46,342 (47.07%) and 33,966 (34.50%) unigenes were aligned to a protein present in the NCBI non-redundant protein (Nr) and SwissProt databases, respectively. There were 42,200 unigenes (42.87%) classified into 25 functional classifications in Clusters of Orthologous Groups (COG) classification. ‘General function prediction’ was dominant, followed by ‘Post-translational modification’ and ‘Signal transduction’. ‘RNA processing and modification,’ ‘Carbohydrate transport and metabolism,’ ‘Secondary metabolites biosynthesis’ categories and ‘Transcription’ also shared a relatively high percentage (Fig. 2), which is similar to results for the closely related C. ensifolium in our previous study [18].

A total of 24,563 unigenes (24.95%) were assigned to 29 functional groups using Gene Ontology (GO) assignments. The three major categories (biological process, cellular component and molecular function) were assigned 16,859, 16,255 and 18,122 GO terms, respectively. Within each of the three main categories of the GO classification scheme, the dominant subcategories were ‘ATP binding,’ ‘oxidation – reduction process’ and ‘mitochondrion,’ respectively. ‘Amino acid metabolic process,’ ‘protein phosphorylation,’ ‘plasma membrane, chloroplast cytoplasmic membrane-bounded vesicle’ were also well represented (Fig. 3). These genes were mainly correlated with ‘binding,’ ‘membrane-bounded organelles’ and ‘organelle parts,’ which correlated well with the genes identified in the stamen or pollen transcriptomes of other plants. Additionally, 11,875 unigenes were assigned to 331 Kyoto Encyclopedia of Genes and Genome (KEGG) pathways. Of these, the ‘metabolic pathways’ made up the majority, followed by ‘biosynthesis of secondary metabolites,’ ‘ribosome’ and ‘cell cycle’. Details of the pathway annotations for significant hits in unigene sets are provided in Additional file 3. We also aligned these unigenes to the Plant Transcription Factor Database (PTFDB) and identified 4451 unigenes as transcription factor sequences belonging to 56 putative transcription factor families (Fig. 4) using the BLASTX algorithm with a cut-off E-value below $10^{-5}$ (http://planttfdb.cbi.pku.edu.cn/). Thus, the ESTs generated during this study provide a valuable resource for gene discovery and future functional analyses of C. goeringii.

### Table 1 Morphological phenotype of the multi-tepal flowers.

Mature flowers are characterized and the data of each sample was the means ± SD from 20 plants. The experiments were repeated three times with similar results.

| Genotype | Sepal width (mm) | Sepal length (mm) | Petal width (mm) | Petal length (mm) |
|----------|------------------|------------------|------------------|------------------|
| Wild type | 1.23 ± 0.12      | 2.78 ± 0.18      | 1.31 ± 0.10      | 3.68 ± 0.22      |
| Mutant    | 0.52 ± 0.08      | 1.08 ± 0.08      | 0.95 ± 0.12      | 3.38 ± 0.16      |

The sepal and lack an organized four-pollinia structure (Fig. 1b & e).

### Table 2 Summary of C. goeringii floral transcriptome

| Type          | Wild type | Mutant | Total |
|---------------|-----------|--------|-------|
| Raw reads     | 98,988,774 | 100,188,534 |       |
| Clean reads   | 90,503,180 | 89,623,606  |       |
| Unigenes      | 79,888     | 94,004  | 98,446 |
| Residues      | 90,372,555 bp | 103,636,656 bp | 97,314,639 bp |
| Smallest unigene | 301 bp  | 301 bp  | 301 bp  |
| Largest unigene | 24,154 bp | 37,800 bp | 37,800 bp |
| Average length | 1131 bp  | 1102 bp | 989 bp  |

### Table 3 Summary of the blast hits against the known protein database

| Database     | Number | Percent |
|--------------|--------|---------|
| Total        | 78,175 | 79.40%  |
| Nr           | 46,342 | 47.07%  |
| Swiss prot   | 33,966 | 34.50%  |
| COG          | 42,200 | 42.87%  |
| GO           | 24,563 | 24.95%  |
| KEGG         | 11,875 | 12.06%  |
Comparison of transcriptomes between the wild-type and the multi-tepal mutant

We investigated the expression levels of unigenes in the wild-type and the mutant plant by comparing these two libraries using a Fragments Per Kilobase of transcript per Million mapped reads (FPKM) analysis, with a false discovery rate ≤ 0.05 and |log₂ ratio| ≥ 1. Subsequently, 18,489 unigenes were found to be differentially expressed, of which 9619 unigenes were up-regulated and 8871 unigenes were down-regulated in the mutant flower as compared with the wild-type. Among them, 412 unigenes (12.8%) showed over a 10-fold change in expression level (Additional file 4).

The set of 18,489 common DEGs between the wild-type and mutant was mapped in accordance with the GO biological process database and the KEGG pathway to identify the genes involved in important biological processes. After the GO term enrichment analysis, 8705 genes from the 18,489 DEGs were assigned to the three main categories: ‘biological process’ (1743), ‘cellular component’ (4682) and ‘molecular function’ (2280). The top five DEG-enriched GO terms were ‘mitochondrion’, ‘ATP binding’, ‘nucleus’, ‘plasma membrane’ and ‘membrane’ (Fig. 5). For the pathway enrichment analysis, we mapped those differentially expressed unigenes to terms in the KEGG database and searched for KEGG terms that were significantly enriched compared with the transcriptome background. In total, 4222 DEGs were assigned to 288 KEGG pathways. The pathways that were most represented by the unigenes were ‘RNA transport’ (122),
followed by ‘spliceosome’ (118), ‘biosynthesis of amino acids’ (102) ‘starch and sucrose metabolism’ (98), ‘protein processing in endoplasmic reticulum’ (95), ‘mRNA surveillance pathway’ (92) and ‘RNA degradation’ (89) (Additional file 5).

The morphological analysis indicated a significantly decreased cell size of the mutant flower, which was probably the primary cause of the small and narrower petals of the multi-tepal mutant. Considering that the expansion of plant cells necessarily requires a loosening of the cell wall, accompanied by ploidy increase through endoreduplication, with further remodeling of cell walls—including the production and trafficking of polysaccharides and membrane-associated proteins—it is reasonable that the unigenes responding to membrane-building and ploidy increase showed differential expression in the mutant flowers. This included ‘mitochondrion’, ‘plasma membrane’, ‘nucleus’ and ‘RNA transport’ recombination.
(Table S3). These GO terms and KEGG classifications serve as indications of significantly different biological processes occurring in the two different phenotypical flowers, which could offer clues for further studies to determine their functions in multi-tepal development.

**DEGs related to floral development**

The identities of flower organs are specified by the interactions of A, B, C, D and E class MADS-box genes. Additionally, the orchid plant has a specific P-code model that modulates floral development, in which the class B gene AP3/DEF-like determines the identity of the lateral petals and lip, while the class B gene PI/GLOBOSA-like, along with class A, C, D and E genes, retain their functions. Therefore, we focused on the changes in the floral-related MADS-box genes between the wild-type and mutant, aiming to reveal the candidates associated with multi-tepal development. Based on the reference annotation, we identified 110 putative MADS-box genes (Additional file 6). Among them, 29 unigenes, including AP1-related genes (2), AP3/DEF-like genes (16) and AGAMOUS-like classes (11), were differentially expressed, with 16 up- and 13 down-regulated in the mutant (Table 4). Eight of them were randomly chosen for further confirmation. Normalized expression values of these genes were created by calculating the value of Fragments Per Kilobase of transcript per Million mapped reads (FPKM) of each individual gene against the FPKM value of *Ubiquitin*, and the real-time RT-PCR expression data were also normalized to the expression of *Ubiquitin*. Results from the real-time RT-PCR assay followed similar trends to those of the read numbers (Fig. 6).

In addition to the typical ABCDE genes that related to floral development, a great number of other transcription factors were also differentially expressed, accounting for more than 25% of *C. goeringii* transcription factors. In total, 1118 of 4451 transcription factors showed significantly different expression levels in the mutant, indicating functions associated with the plant floral patterning. There were 504 and 614 transcription factors—including bZIP, FAR1 and Nin-like—specifically up- and down-regulated in the mutant compared to the wild-type, respectively (Fig. 7a). After normalization to the total number of family members, we found that most differences occurred in the transcription factor family of AP2, NF-YC, TALE and HSF (more than 40% of the members were differentially expressed) (Fig. 7b), suggesting a crucial role for them in *C. goeringii* floral development. For example, percent AP2 transcription factors act primarily in the regulation of developmental programs in *Arabidopsis*—AP2-13 (AP2), AP2-05 (AINTEGUMENT, ANT) and AP2-09 (ANT-LIKE1) genes regulate floral growth and ovule development [29, 30]. NF-Y interacts with CONSTANS in the photoperiod pathway and DELLA family genes in the gibberellin pathway. This CO/NF-Ys complex further regulates the transcription of *SOC1*, a major floral pathway integrator [31]. TALE genes are also required for meristem maintenance and proper patterning of organ initiation, and play a critical role in the diversity of flower and leaf form [32]. All of these new *de novo* transcriptome data sets provide a significant resource for the discovery of genes related to floral patterning and improve our understanding of the regulation of multi-petal mutants. However, the roles of these transcription factors in multi-tepal development and floral patterning require further investigation.

**DEGs related to hormone response involved in floral development**

Plant hormones play important roles in floral development, and their signaling pathways in plants are interconnected in a complex network to regulate flower organ primordia formation, organ specification and final organ size. Comparative analysis of transcriptomes between wild-type and mutant showed a significant enrichment of the plant hormone signal transduction pathway with a P-value of 8.6E-21, and with 70 unigenes involved (Fig. 8 and Additional file 7). Among them, auxin exerts its regulatory role, at least to an extent, by controlling the fundamental processes of cell division, expansion and differentiation. The ortholog of the gene encoding the F-box protein **TRANSPORT INHIBITOR RESPONSE1** (TIR1), an auxin receptor, showed altered expression levels with increases in transcript abundance. Another three well-known groups of early auxin-responsive genes [33] including the auxin/indole-3-acetic acid (Aux/IAA), the small auxin up RNA (SAUR) and the auxin-responsive Gretchen Hagen3 (GH3) gene families were generally repressed in the mutant. For example, the decrease of auxin-responsive protein IAA-like genes, ep123.comp37580/comp42487/comp50482/comp47687/comp42487, ranged from 6.8- to 2.2-fold, and the expressions of SAUR and GH3 family genes also decreased by 5.0- to 2.3-fold in the mutant (Additional file 7).

In addition to the auxin-associated genes, notable differential expressions were also observed among the cytokinin-responsive genes. For example, expressions of the hybrid histidine protein kinases known as cytokinin receptors [34], histidine phosphotransfer proteins (AHPs) [35] and nuclear response regulators (ARRs) that serve as transcriptional regulators in the cytokinin two-component signaling pathway [36] were also largely up-regulated, ranging from 3.8- to 5.4-fold, suggesting a positive regulation of cytokinin in multi-tepal development.

Genes involved in responses to other hormones, such as gibberellin, abscisic acid and ethylene, were also highly enriched. The homology levels of the DELLA family genes **GA INSENSITIVE** (GAI) and
**RGA-LIKE1 (RGL1) [37]** indicated 2.1- to 4.8-fold change in the mutant (Additional file 7). DELLA proteins inhibit growth first by altering the cell division rate during the proliferation phase, followed by altering the cell expansion rate during the expansion phase. A genetic analysis in *Arabidopsis* showed that the DELLA proteins RGA and RGL2 jointly repress petal, stamen and anther development in gibberellin-deficient plants, and that this function is enhanced by RGL1 activity [38, 39]. In our study, the expression of DELLA family genes and downstream responsive transcription factors significantly altered, which revealed a critical role of gibberellin in floral patterning regulation. Abscisic acid-related genes such as the PYR/PYL/RCAR family of receptors, 2C-type protein phosphatases, SnRK2-type kinases and downstream abscisic acid responsive element-binding factors [40, 41] were also greatly altered in the mutant. These results suggest important roles for the hormone pathway in floral patterning regulation, and all candidate genes provided important clues concerning multitepal development.

**Small RNA profiling in wild-type and mutant C. goeringii flowers**

Despite the numerous mRNAs involved in floral development, miRNAs, which regulate mRNA expressions at

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**Table 4** Differentially expressed MADS-box genes related to floral development. Fold change = log2(expression value of each gene of the mutant/wild-type), expression value = FPKM (total transcript fragments/mapped fragments (millions)) × transcript length (kb)

| Sequence ID                  | Fold change | Putative function                                           |
|------------------------------|-------------|-------------------------------------------------------------|
| **Up-regulated**             |             |                                                             |
| ep456.comp45403_c0_seq2      | 7.24        | MADS-box protein 3 [*Cymbidium ensifolium*]                 |
| ep456.comp45308_c0_seq2      | 6.83        | AP1-related protein [*Phalaenopsis amabilis*]               |
| ep456.comp59563_c0_seq4      | 6.51        | APETALA3-like protein [*Papaver somniferum*]                |
| ep456.comp58153_c0_seq7      | 5.79        | SOC1 [*Dendrobium hybrid cultivar*]                         |
| ep123.comp42801_c0_seq3      | 5.55        | SOC1 [*Dendrobium hybrid cultivar*]                         |
| CL11237Contig1               | 4.58        | MADS box transcription factor 1 [*Oncidium hybrid cultivar*] |
| ep456.comp52699_c0_seq6      | 3.89        | DEFICIENS-like MADS-box transcription factor [*Gongora galeata*] |
| ep456.comp50057_c0_seq1      | 2.90        | mads box protein, putative [*Ricinus communis*]             |
| ep456.comp50389_c0_seq1      | 2.67        | SOC1 [*Dendrobium hybrid cultivar*]                         |
| ep456.comp49009_c0_seq1      | 2.62        | AP3 [*Cymbidium goeringii*]                                 |
| ep456.comp58153_c0_seq3      | 2.45        | MADS box protein [*Phalaenopsis equestris*]                 |
| ep456.comp52699_c0_seq10     | 2.35        | DEFICIENS-like MADS-box transcription factor [*Gongora galeata*] |
| ep456.comp52699_c0_seq9      | 2.04        | DEFICIENS-like MADS-box transcription factor [*Gongora galeata*] |
| ep456.comp36865_c0_seq1      | 3.81        | putative MADS-box transcription factor family protein [*Zea mays*] |
| ep456.comp59687_c2_seq7      | 2.27        | MADS1 protein [*Euscholechia californica*]                  |
| ep456.comp59687_c2_seq1      | 2.26        | MADS1 protein [*Euscholechia californica*]                  |
| **Down-regulated**           |             |                                                             |
| ep123.comp50454_c0_seq6      | −2.12       | AGL66 protein [*Euscholechia californica*]                  |
| ep123.comp36586_c0_seq2      | −2.72       | AP3 [*Cymbidium goeringii*]                                 |
| ep123.comp44547_c1_seq1      | −2.98       | MADS box transcription factor [*Elaeis guineensis*]         |
| ep123.comp41679_c0_seq1      | −3.45       | DEFICIENS-like MADS-box transcription factor [*Gongora galeata*] |
| ep123.comp30197_c0_seq1      | −3.89       | C-class MADS-box-like protein [*Orchis italica*]            |
| ep123.comp40112_c1_seq1      | −6.16       | AGAMOUS-like transcription factor [*Dendrobium crumenatum*] |
| ep123.comp42801_c0_seq2      | −6.60       | SOC1 [*Dendrobium hybrid cultivar*]                         |
| ep123.comp35467_c0_seq1      | −6.63       | AP1-like protein [*Cymbidium faber*]                        |
| CL8650Contig1                | −6.92       | SOC1 [*Dendrobium hybrid cultivar*]                         |
| ep123.comp20001_c0_seq1      | −7.03       | Agamous-like MADS-box protein AGL62 [*Aegilops tauschii*]   |
| ep123.comp42801_c0_seq1      | −7.50       | SOC1 [*Dendrobium hybrid cultivar*]                         |
| ep123.comp41679_c0_seq4      | −7.91       | DEFICIENS-like MADS-box transcription factor [*Gongora galeata*] |
| ep123.comp42801_c0_seq4      | −8.59       | SOC1 [*Dendrobium hybrid cultivar*]                         |
the post-transcriptional level, also play essential roles in floral organ identity. To investigate the effects of miRNAs during multi-tepal development in *C. goeringii*, we constructed two small RNA libraries from the wild-type and the mutant flowers, independently. Deep sequencing of the small RNA libraries generated 14,263,624 and 15,042,951 raw reads, respectively, ranging from 17 to over 30 nucleotides in length, and the most represented lengths were 20–23 nt (Additional file 8). After removing adaptor, insert and poly(A) contaminations and low quality reads (smaller than 17 nt), 10,142,278 clean reads from wild-type and 10,400,447 from the mutant were obtained, among which 9,292,059 (91.6%) and 9,611,868 (92.4%), respectively, unique reads mapped to our transcriptome dataset. The mapped small RNA sequences were clustered into several RNA classes, including conserved miRNAs, transfer RNA (tRNA), ribosomal RNA (rRNA), small nuclear RNA (snRNA), small nucleolar RNA (snoRNA), Piwi-interacting RNA (piRNA) and an uncharacterized group (Table 5). Approximately 6.29% and 6.51% of the unique reads from wild-type and mutant, respectively, were annotated as miRNA.

To identify conserved miRNAs in *C. goeringii*, all of the mapped small RNA reads in the transcriptome were used as query against known miRNAs in the database. In total, 226 conserved miRNAs, representing 132 known miRNA families were obtained. Of these, 92 from wild-type and 94 from mutant were perfectly matched to known miRNA families of other plant species, respectively. The distribution of miRNAs in different miRNA families is shown in Table 6 and Additional file 9. The most abundant miRNA was the miR159 family, accounting for 38.6 and 21.6% of the total miRNA in the wild-type and mutant, respectively. Several other miRNA families, such as miR319, miR162 and miR166 families, also had high abundance of expression, consistent with the previous report of the floral miRNA expression profiling in *P. aphrodite* and *C. ensifolium*. Interestingly, we also found relatively high expressions of the miR396, miR6478 and miR2916 families both in the wild-type and the mutant flower, indicating their putative involvement in floral patterning of *C. goeringii*.

In addition to the conserved miRNAs, we identified novel miRNAs in *C. goeringii* flowers. In total, 28 new miRNA candidates from 25 predicted miRNA precursors were obtained, corresponding to 685 unique RNA sequences (Table 7). The miRNA* of these novel miRNAs were also predicted, providing more evidence that they were indeed novel miRNAs of *C. goeringii*. The precursor lengths of the new miRNAs ranged from 74 to 228 nt, with an average length of 128 nt. The average minimum free energy was $-38.3$ kcal mol$^{-1}$, with a range from $-81.9$ kcal mol$^{-1}$ to $-20.6$ kcal mol$^{-1}$. All of the predicted precursors could fold into a characteristic stem-loop structure with the mature miRNA on either the 5′ arm or the 3′ arm of the precursor.

**Target prediction and functional analysis of miRNA in *C. goeringii* flowers**

We used two approaches to identify miRNA targets in *C. goeringii* as described by Chao et al. (2013) [26] (see target prediction procedure section). The first approach, based on extensive complementarity between plant miRNAs and their targets, predicted 5004 miRNA target transcripts for 132 known miRNA families in *C. goeringii* without relying on known targets of other organisms,
which resulted in 3085 targets from wild-type and 4254 from the mutant. The second approach relies on the target homology of the previously reported targets of known miRNAs in Arabidopsis, and resulted in 3099 predicted target genes in 46 miRNA families (Additional file 10). Of the predicted miRNA targets, 1644 target genes from 12 conserved miRNA families were identified by both methods.

The truncated ESTs in our transcriptome limited the target prediction of the first approach because it relies on the direct alignment of miRNAs to EST sequences, while the second approach searched only for homologs, not for orchid-specific targets. Given that each of the approaches has its own strengths and limitations, using both homology-dependent and homology-independent approaches resulted in a more comprehensive target identification than using either one alone, especially for non-model organism lacking complete transcriptome sequence data.

The total of 6459 miRNA target unigenes were used as query against the Nr database, with 5089 hits that exceeded the E-value threshold. Among them, 64 were involved in floral development/flowering time (indicated by red in Additional file 11). Examples include the floral homeotic protein APETALA 2 (miR172), AP3/MADS5/DEFICIENS-like MADS-box transcription factor (miR156 and miR5179) and many other floral-related transcription factors, including TCP, MYB and SPL. Some are overlapping targets that resulted from miRNA sequence similarities. For example, miR156 and miR535 have overlapping predicted target sites in the SPL family of transcription factors. MiR159 and miR319 have five overlapping predicted targets, including two MYB genes and three kinase genes involved in post-translational modification. These results are consistent with observations in Arabidopsis and suggest that the expression of some mRNAs may be regulated by coordinated actions of multiple miRNAs in C. goeringii. Additionally, 29 targets involved in various signaling pathways, such as ARF (targeted by miR160 and miR167), NAC (targeted by 164), NF-YA (targeted by miR169), GRF (targeted by miR396) and HD-ZIP III transcription factors (targeted by miR165/166) were also

![Fig. 7](image_url) Transcription factors differentially expressed between the wild-type and the mutant. a: Number of up-regulated (blue) and down-regulated (red) transcripts were quantified in the mutant compared to the wild-type (based on the FPKM value with |log2(fold change)| > 1 and Q-value < 0.05). b: percentage of up- and down-regulated genes after normalization to the total number of family members.
considered to be associated with floral development (indicated by red in Additional file 11). However, a large number of targets were only annotated with vague information, such as 95 targets that were ‘unnamed protein’ and 93 that were not predicted.

Differentially expressed miRNAs in the multi-tepal mutant and their target genes
Normalized miRNA levels were compared between the wild-type and the mutant to identify the differential expression of miRNAs, and miRNAs that had a fold change log2 > 1 or < −1, and P < 0.05, were considered to be differentially expressed. Subsequently, 11 differentially expressed miRNAs were obtained in the mutant, with only two miRNAs being down-regulated, while nine miRNAs were up-regulated. 455 of their corresponding annotated targets were also obtained—some of them are listed in Table 8. Among them, the conserved miRNA families, miR319 and miR396, were more than two-fold greater in the mutant. Ten of their putative target mRNAs presented correlated expression changes, with seven showing a positive correlation and three showing a negative correlation. Examples included the target genes of miR396 and miR319, ep456.comp59964 (GRF) and ep123.comp31487 (TCP), respectively, which increased by more than eight- and four-fold, whereas their targeted mRNAs, ep456.comp55669 and CL10849Contig1, respectively, were down-regulated. Other floral-related miRNA families (miR156, miR166 and miR167) were approximately two-fold higher in the mutant. Of the targeted mRNAs, 13 were found to be correspondingly differentially expressed between the wild-type and the mutant. Among them, the putative target genes of miR156, SPL-like transcription factors ep456.comp55441, ep456.comp54065 and

| Table 5 Summary of small RNA sequencing data |
|-------------------------------------------|
| **Type** | **WT** | **Count %** | **MU** | **Count %** |
| Clean reads | 10,142,278 | 100% | 10,400,447 | 100% |
| Mapped reads | 9,292,059 | 91.6% | 9,611,868 | 92.4% |
| miRNA | 637,796 | 6.29% | 677,295 | 6.51% |
| tRNA | 232,530 | 2.29% | 257,376 | 2.47% |
| rRNA | 3,255,427 | 32.10% | 3,542,808 | 34.06% |
| snRNA | 79,518 | 0.78% | 88,029 | 0.85% |
| snoRNA | 209,573 | 2.07% | 280,213 | 2.69% |
| piRNA | 32,936 | 0.32% | 35,682 | 0.34% |
| Y_RNA_etc | 5,694,498 | 56.15% | 5,519,044 | 53.08% |

Fig. 8 DEG-enrichment of the plant hormone signal transduction pathway by KEGG annotation. The key regulatory components in multiple hormone response pathways are presented as their names (red, up-regulated; green, down-regulated). Gene IDs and fold changes in transcript abundance are indicated in Additional file 4.
Despite the conserved floral-related miRNAs, miR398 genes and in the multi-tepal mutant. Our previous studies mentioned above, we confirmed the expressions of miR396 and miR319, and their correlated targets among different floral organs in the wild-type and mutant. Our previous studies in Arabidopsis and tobacco indicated that miR396 can regulate expression of GRF genes, which are known to be involved in the control of cell proliferation during leaf and flower development. miR396-overexpressing plants produced flowers with various deformations: fused petals, extremely bent pistils, unfused carpels and single carpels [43, 44]. In this work, four of the GRF-like genes were predicted to be the targets of miR396 in C. goeringii. In silico RNA hybridization revealed a sequence complementary to miR396 in these sequences, and the target position was similar to that in Arabidopsis (Figs. 9 and 10). Since the conserved regulation of GRF genes by miR396 has been broadly reported in both monocotyledons and dicotyledons, our findings indicated that CgGRF-like genes were putative targets of miR396 in C. goeringii.

Expression validation of miRNAs and their targets involved in multi-tepal development

To validate the result obtained from small RNA sequencing and determine the potential roles of the miRNAs mentioned above, we confirmed the expressions of miR396 and miR319, and their candidate target genes were also correspondingly changed by 10- to 6.6-fold, and their candidate target genes were also correspondingly changed by 9.8 to 10.7-fold (Table 8), indicating their potential role in floral patterning.

### Table 6

| miRNA number of reads in clean data × 10⁶ | RPM (reads per million) |
|-----------------------------------------|-------------------------|
| miR159                                  | 6302.99                 |
| miR391                                  | 1307.90                 |
| miR396                                  | 559.20                  |
| miR612                                  | 716.16                  |
| miR6478                                 | 703.50                  |
| miR166                                  | 347.07                  |
| miR2916                                 | 465.28                  |
| miR172                                  | 249.73                  |
| miR171                                  | 204.25                  |
| miR902I                                 | 128.64                  |
| miR167                                  | 87.83                   |
| miR390                                  | 103.38                  |
| miR355                                  | 88.06                   |
| miR398                                  | 284.38                  |
| miR6300                                 | 33.59                   |
| miR528                                  | 3.84                    |
| miR164                                  | 29.08                   |
| miR168                                  | 28.56                   |
| miR156                                  | 27.50                   |
| miR5077                                 | 25.30                   |
| miR100                                  | 22.95                   |
| miR6173                                 | 17.42                   |
| miR9760                                 | 10.49                   |
| miR3630                                 | 10.38                   |
| miR5083                                 | 9.54                    |

ep456.comp52990 were changed by 1.9- to 3.9-fold. The target genes of miR166 and miR167, putative HD-ZIP transcription factor ep456.comp58119 and ARF-like unigenes, ep123.comp48918 and ep456.comp58489, respectively, also showed significant changes by 10- to 6.6-fold, indicating a regulatory relationship between the miRNA and the targets. Interestingly, we found that miR535 was notably enhanced in the mutant. MiR535 was expressed in a similar fashion to miR156 in the leaves and flowers of papaya and rice, and targeted the same SPBL genes [42]. Our result indicated a potentially redundant role for miR535 function in regulating SPL genes and in the multi-tepal mutant.

Despite the conserved floral-related miRNAs, miR398 and miR528, which are majorly involved in plant stress responses, were also altered in the mutant. The former was 2.7-fold lower and the latter was about three-fold higher. These results indicated a potential link between the stress response and the mutant phenotype. Moreover, we noted that miR6173, miR3630 and miR5083 exhibited different specificities with changes ranging from one- to 2.4-fold, and their candidate target genes were also correspondingly changed by –9.8 to 10.7-fold (Table 8), indicating their potential role in floral patterning.

Expression validation of miRNAs and their targets involved in multi-tepal development

To validate the result obtained from small RNA sequencing and determine the potential roles of the miRNAs mentioned above, we confirmed the expressions of miR396 and miR319, and their correlated targets among different floral organs in the wild-type and mutant. Our previous studies in Arabidopsis and tobacco indicated that miR396 can regulate expression of GRF genes, which are known to be involved in the control of cell proliferation during leaf and flower development. miR396-overexpressing plants produced flowers with various deformations: fused petals, extremely bent pistils, unfused carpels and single carpels [43, 44]. In this work, four of the GRF-like genes were predicted to be the targets of miR396 in C. goeringii. In silico RNA hybridization revealed a sequence complementary to miR396 in these sequences, and the target position was similar to that in Arabidopsis (Figs. 9 and 10). Since the conserved regulation of GRF genes by miR396 has been broadly reported in both monocotyledons and dicotyledons, our findings indicated that CgGRF-like genes were putative targets of miR396 in C. goeringii.
Comparatively, expression of miR319 was also broadly detected in different organs, with higher abundance in the petals and lips. Consistent with the transcriptome deep sequencing, the overall expression level of miR319 was higher in the mutant than in the wild-type (Fig. 12a). In addition, the mRNA amounts of two TCP-like unigenes that showed sequence complementary to miR319 (Additional file 12) were also determined. Real-time PCR showed large increases in TCP-like gene ep123.comp31487 in the central part of the mutant flower, with a positive correlation with that of miR319 (Fig. 12b). There were no significant correlations between the expression levels of ep123.comp27081 and miR319. This could be due to non-cleavage repression, feedback regulation and spatial or temporal exclusion of miRNAs and their targets. The expression of other miRNAs leading to different levels of target repression, or other levels of regulation, such as promoter methylation and translational repression also exist. Our results showed that the relative abundance of these putative targeted sequences followed similar trends to those of the read numbers, and the same differential expression patterns were also observed between the wild-type and the mutant in technical replicates. This indicated that the comprehensive profiling of C. goeringii miRNA provided a useful reference for further study. However, functional analysis is required to explore the regulatory mechanisms of various miRNAs and their targets contributing to the multi-tepal phenotype in the orchid.

### Discussion

Transcriptome sequencing and the sequence annotation of C. goeringii

C. goeringii is a very important potted plant in China, and the multi-tepal mutant is popular in the commercial orchid market. However, little is known about the mechanisms responsible for floral patterning and multi-tepal development. Currently, next-generation sequencing technologies make it convenient to study the transcriptome of a particular organism or tissue to gain insight into biological processes, and numbers of these studies have been reported in many orchid plants. In C. goeringii, however, genomic information is currently unavailable.

| miRNA   | Sequence                              | Read number | Chr-location      | Length (nt) | Potential_energy (kcal/mol) |
|---------|---------------------------------------|-------------|--------------------|-------------|-----------------------------|
| Cgo-m0001-3p | AGGCAGCCGUGCGCCGCGUUGG             | 11          | CL4439Contig1:400473:+ | 74          | −30.60                      |
| Cgo-m0001-5p | GAGCUUGUGGGCGGACAAAAG             | 32          | CL4073Contig1:867932:+ | 66          | −21.40                      |
| Cgo-m0002-3p | CGUGAGCCGACGGGCAUGA             | 10          | ep123.comp16229_c0_seq1:56148:+ | 93          | −24.70                      |
| Cgo-m0003-5p | CAUGCCUGUUCAGCGGCGUGC         | 76          | ep123.comp50928_c0_seq1:277367:+ | 91          | −30.20                      |
| Cgo-m0004-3p | AUCGAUUGUGUAAGUAGUGGCUGG       | 10          | ep123.comp41527_c0_seq1:10161092:+ | 77          | −24.20                      |
| Cgo-m0004-5p | CUGUGGCGCUUUCUCUGGCAGC          | 50          | ep123.comp41527_c0_seq1:311381220:+ | 83          | −42.00                      |
| Cgo-m0005-5p | CGGGAAUGUCAGGAAGUUGG          | 57          | ep123.comp41527_c0_seq1:313971476:+ | 80          | −37.30                      |
| Cgo-m0006-3p | CACCUUGACUAAUCAUGCGC         | 13          | ep123.comp41527_c0_seq1:81071091:+ | 75          | −24.20                      |
| Cgo-m0006-5p | UUCUGCGGAAGGCUUCAGU         | 18          | ep123.comp41527_c0_seq1:17971877:+ | 81          | −26.70                      |
| Cgo-m0007-3p | AUCGAUUGUGUAAGUAGUGGCUGG     | 10          | ep123.comp41527_c0_seq1:10171091:+ | 75          | −24.20                      |
| Cgo-m0008-3p | CUCUUCAAUCUACUGAUG             | 12          | ep123.comp45914_c0_seq1:13531426:+ | 74          | −21.40                      |
| Cgo-m0010-3p | CAGUUGGAGUUGUUGCGUGU          | 12          | ep123.comp29575_c0_seq1:267343:+ | 77          | −23.60                      |
| Cgo-m0010-5p | UAUUCGGCUUUCUGACUGUGC        | 20          | ep123.comp48991_c0_seq1:13381424:+ | 87          | −21.80                      |
| Cgo-m0011-3p | GGAGAAAGGUCAAGGAAGUUGG         | 15          | ep123.comp48451_c0_seq2:257343:+ | 87          | −30.80                      |
| Cgo-m0012-5p | AUUUUUGCCGCGAAGUUGACU        | 10          | ep123.comp48041_c0_seq1:15521628:+ | 77          | −20.60                      |
| Cgo-m0013-3p | GGAGAAAGGUCAAGGAAGUUGG         | 30          | ep123.comp48451_c0_seq2:258342:+ | 85          | −30.80                      |
| Cgo-m0013-5p | GAAUUGUGUGACGCAUUGCU        | 10          | ep123.comp53024_c0_seq1:65476616:+ | 70          | −24.60                      |
| Cgo-m0014-5p | SGCCCGAGUAGAGCUUGUUGUCUGG     | 26          | ep123.comp53024_c0_seq5:36193713:+ | 95          | −29.80                      |
| Cgo-m0015-5p | GGGCGGAAGUAAAGAAGUGGUGG        | 80          | ep123.comp53024_c0_seq5:54115491:+ | 81          | −37.60                      |
| Cgo-m0016-5p | UUGUGCGAGGUAUAGUGACU        | 21          | ep123.comp52937_c0_seq1:286375:+ | 90          | −25.30                      |
| Cgo-m0017-3p | CGGAAUUGGUGCGCAGGGCAGCGC       | 27          | ep123.comp53646_c0_seq1:117189:+ | 73          | −27.10                      |
| Cgo-m0017-5p | AUGCCUAUACUAGUGCUGUGGUGUGA     | 12          | ep123.comp55956_c0_seq1:16331732:+ | 100         | −20.80                      |
| Cgo-m0018-5p | AUGGCGUAGUAAUUGCAUUGG          | 30          | ep123.comp57349_c0_seq1:186273:+ | 88          | −29.30                      |
| Cgo-m0019-5p | AUGGCUGUAAUUGCAGUUGGUGG       | 40          | ep123.comp57349_c0_seq3:186273:+ | 88          | −29.30                      |
| Cgo-m0020-3p | UGGCGGCGAAGUUGGUGG          | 10          | ep123.comp58210_c0_seq1:41804271:+ | 92          | −26.40                      |
| miRNA | Fold change | Target | Target fold change | Nr annotation |
|-------|-------------|--------|-------------------|---------------|
| miR396 | 2.37 | ep123.comp47283_c0_seq1 | −2.11 | ceramidase, putative {Ricinus communis} |
|        |     | ep456.comp55669_c0_seq1 | −2.60 | LOC_Os02g47280.1|Oryza sativa_subsp._japonica|GRF|LOC_Os02g47280.1 |
|        |     | ep456.comp58258_c0_seq1 | 2.47 | ceramidase, putative {Ricinus communis} |
|        |     | ep123.comp17494_c0_seq1 | 4.00 | LOC_Os04g51190.3|Oryza sativa_subsp._japonica|GRF|LOC_Os04g51190.3 |
| CL10167 | 4.28 | ep123.comp17494_c0_seq1 | 4.00 | LOC_Os04g51190.3|Oryza sativa_subsp._japonica|GRF|LOC_Os04g51190.3 |
| miR319 | 2.04 | CL10849Contig1 | −1.11 | TPA: putative MYB DNA-binding domain superfamily protein {Zea mays} |
|        |     | ep123.comp27081_c0_seq1 | 1.81 | candidate developmental transcription factor TCP1 {Phalaenopsis hybrid cultivar} |
|        |     | ep123.comp44934_c0_seq2 | 3.19 | TCP family transcription factor {Dendrobium hybrid cultivar} |
|        |     | ep123.comp31487_c0_seq1 | 4.09 | candidate developmental transcription factor TCP1 {Phalaenopsis hybrid cultivar} |
| miR156 | 2.00 | ep456.comp55441_c0_seq1 | 1.91 | SQUAMOSA promoter-binding-like 3 {Erycinapusilla} |
|        |     | ep456.comp55441_c0_seq1 | 1.91 | SQUAMOSA promoter-binding-like 3 {Erycinapusilla} |
|        |     | ep456.comp55441_c0_seq1 | 1.91 | SQUAMOSA promoter-binding-like 3 {Erycinapusilla} |
| miR166 | 1.87 | ep123.comp46514_c0_seq3 | −4.20 | PREDICTED: receptor-like serine/threonine-protein kinase SD1-8-like {Vitis vinifera} |
|        |     | ep123.comp46514_c0_seq3 | −4.20 | PREDICTED: receptor-like serine/threonine-protein kinase SD1-8-like {Vitis vinifera} |
|        |     | ep123.comp46514_c0_seq3 | −4.20 | PREDICTED: receptor-like serine/threonine-protein kinase SD1-8-like {Vitis vinifera} |
| miR535 | 1.45 | ep456.comp49040_c0_seq1 | −5.89 | PREDICTED: ELMO domain-containing protein A-like {Glycine max} |
|        |     | ep456.comp49040_c0_seq1 | −5.89 | PREDICTED: ELMO domain-containing protein A-like {Glycine max} |
|        |     | ep456.comp49040_c0_seq1 | −5.89 | PREDICTED: ELMO domain-containing protein A-like {Glycine max} |
| miR398 | −2.67 | ep456.comp58639_c0_seq3 | −7.82 | hypothetical protein PRUE_ppa02063mg {Pruunpersica} |
|        |     | ep123.comp48546_c0_seq1 | −3.23 | PREDICTED: LOW QUALITY PROTEIN: DNA polymerase theta-like {Vitis vinifera} |
|        |     | ep123.comp47734_c0_seq2 | −2.29 | PREDICTED: uncharacterized protein LOC100245378 {Vitis vinifera} |
|        |     | ep456.comp58639_c0_seq4 | 2.00 | hypothetical protein PRUE_ppa02063mg {Pruunpersica} |
|        |     | ep456.comp58639_c0_seq4 | 2.00 | hypothetical protein PRUE_ppa02063mg {Pruunpersica} |
|        |     | ep456.comp58639_c0_seq4 | 2.00 | hypothetical protein PRUE_ppa02063mg {Pruunpersica} |
| miR528 | 2.19 | ep456.comp57151_c0_seq1 | 1.84 | polyphenol oxidase {Doritis pulcherima x Phalaenopsis hybrid cultivar} |
|        |     | ep456.comp53200_c0_seq2 | 2.87 | TPA: hypothetical protein ZEAMMB73_977642 {Zea mays} |
|        |     | ep456.comp52726_c0_seq1 | 5.34 | glutaredoxin family protein {Musa acuminata} |
|        |     | ep123.comp50029_c0_seq1 | 3.11 | Endoribonuclease Dicer-like protein 4 {Trincomaranta} |
| miR6173 | −1.11 | ep123.comp48621_c0_seq3 | −8.01 | predicted protein {Hordeum vulgare subsp. vulgare} |
|        |     | ep123.comp41527_c0_seq11 | −9.84 | Mitochondrial protein, putative {Medicago truncatula} |
|        |     | ep123.comp48621_c0_seq1 | −8.01 | TPA: hypothetical protein ZEAMMB73_631850 {Zea mays} |
| miR3630 | 1.96 | ep123.comp42566_c0_seq1 | 10.71 | SQUAMOSA promoter-binding-like 12 {Erycinapusilla} |
Our research primarily generated a large amount of cDNA sequences and miRNA data that will facilitate more detailed studies in *C. goeringii*, and will identify the genes controlling floral patterning in the multi-tepal mutant. In our study, a combination of wild-type and multi-tepal mutant libraries generated ~20 Gb of high quality reads that formed 98,446 unigenes, having an average sequence length of 989 bp. These unigenes were used in searches powered by the BLASTX algorithm and annotations against protein databases, like Nr, SwissProt, COG, GO and KEGG. In total, 78,175 sequences were identified through searches, and 20.6% of the unigenes had no homologs in the NCBI database.

When comparing with the orchid plants in other sub-families of the Orchidaceae, such as *P. equestris* and *D. officinale*, about 51% mapping rate was obtained, consistent with the similarity between *C. ensifolium* and *P. equestris* reported by Li et al. [45]. In contrast, comparison among three *Cymbidium* species, *C. ensifolium*, *C. sinense* and *C. goeringii*, showed almost 90% similarities. Among the aligned sequences, 74,377 (76%) unigenes had similarities with both *C. ensifolium* and *C. sinense*, whereas 10,628 (10%) isotigs had no similarity with either *C. ensifolium* or *C. sinense* (Fig. 13a). This may indicate that *C. goeringii* vegetative and reproductive growth contains many unique processes and pathways.

The GO analysis resulted in annotation of 4932 of the unigenes unique to *C. goeringii*, and the genes involved in metabolic processes, cellular processes and stimulus response were relatively more represented. For cellular components, the ‘cells’ and ‘cell part’ categories were most represented followed by ‘organelle’ (Fig. 13b); and for the molecular functions category, ‘binding’ and ‘catalytic activity’ were the most prevalent GO terms, followed by ‘structural molecular’. All of these unique unigenes belonging to different GO terms may be important for traits specific to *C. goeringii*, such as special flower formation, scent production and environment adaptation, which represent a valuable resource for exploring the genetic diversity of *C. goeringii*, and for comparative genomic studies among orchid species.

### Floral homeotic genes related to multi-tepal development of *C. goeringii*

According to the floral quartet model, BC gene products and the SEPALATTA protein assembled into quaternary complexes, which specify the stamen, while C gene and SEPALATTA control carpel identity. AG proteins can form a homodimer and exert CArG-box binding activity. Orchids have unique morphologies, and previous studies in the orchid plant indicated a modified model of patterning, although most of the gene orthologs involved in the regulation of organ identity were highly conserved in evolution [46, 47]. For example, orchids typically have four *AP3/DEF*-like genes, representing the ancient gene Clades 1, 2, 3 and 4. High gene expression levels of Clades 1 and 2, and low levels of Clades 3 and 4, specify inner lateral tepals, whereas labellum development requires low levels of Clade 1 and 2 expression and high levels of Clade 3 and 4 expression [13, 14, 48]. Findings of duplicated four-clade B-class genes with differential expression patterns in orchid floral organs and their divergent protein behaviors support the

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**Table 8** Differentially expressed miRNAs and their target genes between the wild type and the mutant. Fold change = log2(expression value of each gene of the mutant/wild-type) (Continued)

| miR   | Subject of expression | Target Gene | Fold Change |
|-------|-----------------------|-------------|-------------|
| miR396 | ep456.comp55669        | ep456.comp55669 | -0.27 kcal/mol |
| CL10167 | ep123.comp17494       | ep123.comp17494 | -0.33 kcal/mol |
| mfe    | -27.1 kcal/mol        |              |             |
| mfe    | -33.9 kcal/mol        |              |             |
| mfe    | -21.0 kcal/mol        |              |             |
| mfe    | -28.3 kcal/mol        |              |             |

**Fig. 9** Comparison of complementarity profiles for miR396 with GRF-like genes. Free energies of the duplex structures were calculated using RNA hybrid software (http://bibiserv.techfak.uni-bielefeld.de/mahybrid)
unique evolutionary routes of B-class genes associated with unique floral organ development. However, orchids remain under-represented in molecular studies of floral development compared with other species-rich plant families.

In this study, we primarily identified many floral homeotic genes differentially expressed between the wild-type and mutant, which were probably related to multi-tepal formation, including homologs of the A-function gene APETALA, B-function gene AP3/DEF, and E-function genes SEPELLATA/AGL. More importantly, we found four AG-like C-class genes that were notably repressed in the mutant (Table 3). According to the floral quartet model, C-class genes specify stamen and carpel development and function in meristem determination because C-class mutants are indeterminate in whorl 4 and form a new C-class mutant flower instead of carpels. In core eudicots, C lineage MADS-box genes have been separated into euAG and PLENA lineages. After duplication, the primary C functions were subfunctionalized. For example, in Antirrhinum majus, PLENA has a main role in C functions, including the termination of floral meristem and the establishment of stamen and carpel identities, but the euAG sublineage gene FARINERRI contributes only to male fertility [49]. Duplications of C-class genes in monocots have revealed that their functions may diversify and become partially redundant. In maize, C-class ZMM2 has a major function in organ identity, and ZAG1 is predominant in floral meristem determinacy [50]. In rice, OsMADS3 has a strong role in repressing lodicule development in whorl 2 and in specifying stamen identity, whereas OsMADS58 contributes more to conferring floral meristem determinacy and regulating carpel development [51]. Orchid C-class genes were identified and contained duplication events. Previous study in C. ensifolium indicated that the major C function provided by CeMADS1 and the paralogous CeMADS2 may have functional redundancy in meristem activity [16]. Here, we found four unigenes that act as C-class AG-like genes and were significantly decreased in the mutant. A further analysis will reveal the potential roles and regulatory mechanisms of these paralogous genes.

**Other floral regulators related to multi-tepal development of C. goeringii**

Along with the altered expression of ABCDE model genes, many other transcription factors that regulate class floral homeotic genes [52, 53], including the orthologs of LUG and EMF2, were up-regulated, whereas the expression of CLF orthologs were approximately five-fold lower in the mutant flower transcriptome. Other important categories of gene orthologs regulating patterning and symmetry also showed altered expression levels in the mutant floral transcriptome. For example, the putative orthologs of

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**Fig. 10** Sequence complementary to miR396 in four of the GRF-like genes. The comparison of complementary site of miR396 to GRF-like genes is indicated by the red frame.
TCP and NAC. For the TCP ortholog, comparatively higher expression levels of four TCP-like genes were observed in the mutant transcriptome compared with the wild-type. Additionally, 19 of the putative NAC orthologs were also altered in the mutant floral transcriptome (Additional file 13).

There were also homologs of meristem activity regulators, such as BAM and WUS, which enhance meristem proliferation [54], found to have higher expression levels in the mutant floral transcriptome. In parallel, GA1 and ANT, which are involved in decreasing meristem proliferation [55], also showed high levels in the mutant flower. Putative homologs of the genes ER, ERL1 and ERL2 encoding protein kinases that influence meristem cell fate and patterning in the inflorescence meristem [56], were also highly expressed. This \textit{in silico} expression analysis of genes related to flower development and cell fate determination provided new insights into multi-tepal development.

**Integrated actions of multiple hormones involved in the multi-tepal development**

During floral development, the floral organ identity genes specify the identity of different floral organs. However, to obtain their characteristic final size and shape, growth of the developing floral primordium needs to be tightly coordinated first through cell proliferation and then by cell expansion [9, 57]. Plant hormones play important roles in this progression, and the signaling pathways of phytohormones are interconnected in a complex network to regulate flower organ primordia formation, organ specification and final organ size [58].

In this study, transcriptome analyses showed that the plant hormone signal transduction pathway involving 70 unigenes was enriched with a \textit{P}-value of 8.6E-21. Among them, auxin represents a master player as it acts not only as a local morphogenetic trigger in flower organ primordia formation, but also in concert with other hormones during further growth, patterning and reproductive organ development.
Interestingly, three well-known groups of early auxin-responsive genes—the Aux/IAA, SAUR and GH3 gene families [33]—were generally repressed in the mutant. This repression of gene expression in the auxin signal pathway was well correlated with the retarded cell expansion and decreased cell size, which resulted in small and narrower tepals in the mutant.

Other hormones, such as cytokinin and gibberellin, are also critical for proper floral organ identity. Exogenous cytokinin applications and accumulation of endogenous cytokinin increase the flower number and induce an aberrant floral phenotype in several species. For example, an elevated level of endogenous cytokinin in Arabidopsis promotes cell division in the inflorescence meristem and the flower meristem, resulting in an increased number of flower organs [60]. In this study, the expressions of the ARR and AHP homologies in the cytokinin signaling pathway were also greatly up-regulated in the mutant, suggesting a positive regulation of cytokinin in the multi-tepal development. Gibberellin promotes the expression of floral homeotic genes, including AP3, PI and AG by antagonizing the effects of DELLA proteins—a family of nuclear growth repressors altering cell proliferation and expansion [37–39]. Our findings indicated that the expression of DELLA family genes and downstream responsive transcription factors were significantly altered, suggesting a critical role of gibberellin on floral patterning regulation. Moreover, gene homologies involved in abscisic acid, ethylene and brassinosteroid were also greatly altered in the mutant. These results suggest important roles for the hormone pathway in floral patterning regulation, and all of the candidate genes provide important clues to multi-tepal development.
MiRNA and transcription factor networks regulating multi-tepal development

MiRNAs are not only regulators of gene expression, but also play an essential role in the coordination of complex developmental processes through their extensive integration within genetic networks. In our study, 11 out of 132 miRNA families exhibited fold changes of log2 > 1 and P-value < 0.05 between the wild-type and the mutant, functioning as putative regulators for multi-tepal development. These included the well-known floral-related miR156, miR166/165, miR167, miR319 and miR396 (Fig. 14).

MiR156 and miR167 are well known to regulate floral, especially reproductive structure, development through interference with the auxin signaling pathway. For example, miR156-targeted SPL8 gene regulates gynoecium patterning in Arabidopsis, with down-regulation of miR156-targeted SPL genes resulting in a shortened style and an apically swollen ovary narrowing onto an elongated gynophore [61]. miR167 restricts ARF6 and ARF8 accumulation, resulting in failure to elongate filaments at anthesis (pre-anthesis filament elongation) and either lack of functional pollen grains or release of defective pollen grains before completion of filament elongation [62]. In this study, we found enhanced expression of miR156 and miR167, and corresponding alterations of their target transcripts in the C. goeringii multi-tepal mutant, which probably contributed to the significant deformation of its reproductive organs.

miR166/165 regulate HD-ZIP III transcripts through ARGONAUTE10 (AGO10) proteins to regulate shoot apical meristem formation and floral development by acting in parallel with the WUS–CLAVATA pathway [63]. In our work, homologs of these meristem activity regulators were induced in the mutant, and correlated well with the redundant floral structures of the multi-tepal mutant. However, further work is required to explore the targets and their regulatory mechanisms.

More importantly, miR396, another conserved miRNA, had the largest fold change difference between the wild-type and the mutant. The miR396 plays critical roles in plant leaf and flower development by modulating cell proliferation, at least to some extent. Seven of the nine members of the growth regulating factor family, GRF1–4 and GRF7–9, are the targets of miR396 in Arabidopsis. GRFs together with the family of putative transcriptional co-activators encoded by the GRF-INTERACTING FACTOR (GIF) genes regulate organ size by maintaining cell proliferation, with mutants forming smaller and narrower leaves, whereas overexpression leads to larger leaf size. Plants expressing high levels of miR396 produce narrow leaves, fused tepals and abnormal productive organs, such as extremely bent pistils, unfused carpels and single carpels [43, 44, 64].

Interestingly, the expression of miR396 is itself regulated by TCP4, a member of the TEOSINTE BRANCHED1/CYCLOIDEA/PCF (TCP) transcription factor family. TCP4 is a key regulator in petal development, to modulate cell division and cell differentiation, and is a target for miR319 [65]. Expression of a wild-type version of TCP4 under the AP3 promoter (AP3:TCP4) has very limited effect. By contrast, the AP3:mTCP4 lines with a miR319 resistant version exhibit a complete absence of petals and stamens. A point mutation in the miR319 target site of TCP4 reduces the interaction with miR319 and causes higher levels of miR396 and lower amounts of GRF transcripts [66]. In this study, we found higher expressions of both miR319 and miR396, and corresponding changes of their target transcripts in the multi-tepal mutant. In light of the regulation of miR396 by TCP4 in Arabidopsis, we hypothesize that a miR319/TCP4–miR396/GRF regulatory cascade also exists in C. goeringii to regulate cell proliferation. This probably contributed to the reduced cell size and narrow petals of the mutant.

Conclusions

Orchid plants with multiple tepals have considerable ecological and cultural value. In this work, we analyzed a typical multi-tepal mutant of C. goeringii, which continued to produce sepals and petals instead of a gynostemium in the center of the flower, with misshapen lips and gynostemia. Aiming to reveal the networks involved in regulatory mechanisms of multi-tepal development, we compared the mRNA and small RNA profiles between the wild-type and the mutant. Comparative transcriptome analysis showed a great number of floral-related genes, as well as plant hormone-responsive genes contributing...
to multi-tepal development. Combined with miRNA sequencing data, two transcription factor/microRNA-based genetic pathways were considered to regulate multi-tepal development: well-known floral-related miR156/SPL and miR167/ARF regulatory modes probably contributing to reproductive organs development, and the miR319/TCP4–miR396/GRF regulatory cascade involved in cell proliferation in the multi-tepal development. Moreover, the comprehensive profiling of the *C. goeringii* transcriptome and miRNAome provided a useful reference for further investigations of miRNA in Orchidaceae.

**Methods**

**Plant materials and growth conditions**

Wild-type plants of *C. goeringii* ‘Songmei’ and the multi-tepal mutant ‘Yuhudie’ (the most widely known commercial cultivars in China) used in this study were artificially cultivated and collected from the cultivation base of Environmental Horticulture Research Institute, Guangdong Academy of Agricultural Sciences, China. All plants were grown and maintained in pots in a greenhouse at day/night with temperatures of 26/23 °C and photoperiod of 16/8 h.

**Library construction and Illumina sequencing**

The primary aim of this study was to identify the genes responsible for the multi-tepal phenotype, including floral trait mutations such as multi-sepals and -petals, misshapen labella and gynostemium–petal fusions. We therefore respectively extracted the total RNA from the sepal, petal, labellum and gynostemium, rather than from the whole flower, to compare the unigene sets corresponding to individual floral organs. The cDNA and small RNA libraries were prepared from an equal mixture of RNAs isolated from different floral organs as outlined above. Each tissue sample consisted of individual floral organs isolated from 5–8 natural flowers. The mRNAs were purified from total RNA using the Oligotex mRNA Midi Kit (QIAGEN, Germany) and quantified using a Nano-Drop 2000 spectrophotometer according to the Illumina manufacturer’s instructions as described in our previous work [18]. Briefly, mRNAs were isolated from Total RNA and fragmented to approximately 200 bp. Subsequently, the collected mRNAs were subjected to first strand and second strand cDNA synthesis following by adapter ligation and enrichment with a low-cycle according to instructions of TruSeq® RNA HT Sample Prep Kit (Illumina, USA). The purified library products were evaluated using the Agilent 2200 TapeStation and diluted to 10 pM for cluster generation in situ on the HiSeq2500 single-end flow cell followed by sequencing (1 × 50 bp) on HiSeq 2500.

**Sequence assembly and annotation**

The resulting raw sequence reads with weak signal or low quality were screened and trimmed by GS FLX pyrosequencing software, to yield a final dataset comprising high-quality (HQ) (>99.5% accuracy of single-base reads) sequences. Prior to assembly, primer and adapter sequences were trimmed from the HQ dataset, and sequences shorter than 50 bp were removed. The remaining data were assembled into unique sequences (including contigs) using trinityrnaseq (http://trinityrnaseq.sourceforge.net/analysis/extract_proteins_from_trinity_transcripts.html).

For unigene annotations, the genes in which protein-coding sequences were found were searched against the Nr database (http://www.ncbi.nlm.nih.gov) and the SwissProt protein database (http://www.expasy.ch/sprot) using the BLASTP algorithm with an E-value cut-off of 10^-5. The remaining sequences were searched against public databases, using the BLASTX algorithm with an E-value cut-off of 10^-5.

GO classification was obtained by GO terms in the database (http://www.geneontology.org/). The unigene sequences were also aligned to the COG database to predict and classify functions. For identifying the transcription factors of *C. goeringii*, the protein sequence set of each transcription factor family from rice were downloaded from the PTFDB (http://planttfdb.cbi.pku.edu.cn/) and BLASTed against the sequences in our dataset with use of the TBLASTN programs. Sequence similarity was considered significant at E-value < 10^-5. Pathway assignments were made according to the KEGG mapping project (http://www.genome.jp/). Enzyme commission (EC) numbers were assigned to unique sequences that had BLASTX scores with an E-value cut-off of 10^-5 as determined by searching the KEGG database. The unique sequences were allocated to specific biochemical pathways according to the corresponding EC distribution in the KEGG database.
DEG analysis
The gene expression level was calculated by the FPKM value: FPKM = [total transcript fragments(mapped fragments (millions)] × transcript length (kb). The significance of difference in gene expression between the wild-type and mutant was determined using edgeR. False discovery rate (FDR) was applied to identify the threshold of the P-value in multiple tests; with FDR < 0.05 and |log2 ratio| > 1 (two-fold change) as the threshold for significant difference in gene expression.

The DEGs were annotated using GO and KEGG enrichment analyses according to a method similar to that described by Wang, which firstly mapped all DEGs to GO terms (or KEGG pathways) in the database (http://www.geneontology.org/, http://www.genome.ad.jp/) and calculated gene numbers for every term (or pathway). This was followed by a hypergeometric test to find the significantly enriched terms in DEGs compared with the genome background. We used the corrected P-value ≤ 0.05 or Q-value ≤ 0.05 as a threshold for the significantly enriched GO terms or KEGG pathways, respectively, in DEGs.

Bioinformatic analysis of small RNA deep-sequencing data
The small RNA data output from Illumina was filtered and the sequence mapping reads with poly-N, 5′-adapter or inset tags were removed. The clean data with read lengths in a specific size range of 17–30 nt were chosen for further analysis. These small RNA reads were mapped to Repeat Masker (www.repeatmasker.org), Rfam (rfam.janelia.org), UCSC (gtrnadb.ucsc.edu) and NONCODE (www.noncode.org/NONCODEv3/download.html), to distinguish miRNAs from the tags originating from protein-coding genes, repeat sequences, rRNA, tRNA, snRNA, snoRNA and piRNA.

Conserved miRNA alignment and its expression
All unique reads left from the previous screening were searched against the miRBase (version: 21.0) using BLASTN with the same parameters values as used in the miRBase Web-Blast server (−w 4 −r 5 −q −4). Unique reads that were identical to or had less than three mismatches to known miRNAs were regarded as potential conserved miRNAs in C. goeringii. The counts of unique reads were normalized to reads per million (RPM) by dividing the raw read count by the total number of reads in each library. The expression profiles for each miRNA family were calculated by summing all reads annotated to the same miRNA family in each library. MiRNAs that had change ratios of > 2 or < 0.5 (fold change log2 > 1 or < −1) and P < 0.05 were set as the threshold for significant differential expression by default. Statistical analysis was performed using the method described by Audic and Claverie.

Novel miRNA prediction
The miRNA precursor is characterized by its hairpin structure, which can be used to predict novel miRNA. Here, we used the software Mireap (https://sourceforge.net/projects/mireap/) to predict novel miRNA through exploring the secondary structure by identifying the Dicer cleavage site and the minimum free energy of the small RNA tags unannotated in the previous steps. The parameter value −f: (Flank sequence length of miRNA precursor) = 100 and the free energy < −20 kcal/mol.

Target gene prediction
Two computational approaches were applied to predict miRNA target transcripts in this study as described by Chao et al. (2014). In the first approach, miRNAs were searched against our EST dataset with -g (gapped alignment) F (false) appended for BLASTN to prevent the insertion of gaps in the middle of an alignment. Alignments containing positions 2–12 of the miRNA with a mapping length over 16 nt and ≤ 3 mismatches were considered to be miRNA target candidates. In the second procedure, genomic data of Arabidopsis were downloaded from the TAIR10 database (http://www.arabidopsis.org/index.jsp) and BLASTed against our EST dataset of C. goeringii using a cut-off value of 1e-30. An EST was considered a candidate target gene if its best BLAST match was a target of Arabidopsis miRNAs listed in the Arabidopsis small RNA project (ASRP) database (http://asrp.cgrb.oregonstate.edu/) [67].

Stem-loop RT-PCR of miRNAs and RT-qPCR of target genes
Individual floral organs of wild-type and multi-tepal mutant were used, independently, to extract RNA. The cDNA of the mature miRNA was prepared using the miRNA reverse transcription kit M-MLV (Takara, China), and the reverse-transcribed products were used as the template for RT-qPCR with gene-specific primers. The U6 snRNA was used for normalization. The miRNA specific stem-loop primers and gene-specific RT-qPCR primers were designed according to the rules described in Chen et al. [68]. For the RT-qPCR of target genes, total RNA extracted from different tissue types were reverse-transcribed by oligo(dT) primed cDNA synthesis protocol (Fermentas). The resulting cDNA was subjected to relative quantitative PCR using Bio-Rad CFX-96 RealTime PCR System (Bio-Rad, USA) in a final volume of 20 μl containing 2 μl of cDNA and 10 μl of SYBR premix Ex-taq™ (Takara, Japan). Ubiquitin was used as an internal control for normalization to make a comparison of gene expression level between the accessions. For each reported result at least three independent biological samples were subjected to a minimum of three technical replicates. The primers
designed with Primer 5.0 software are listed in Additional file 14.

Additional files

Additional file 1: Paradermal view of the epidermal cells of the petal both in the wild type and the mutant. (TIF 1951 kb)

Additional file 2: The length distribution of assembled unigenes. The x-axis represents the sequence length in base pairs. The y-axis represents the unigenes number. (TIF 478 kb)

Additional file 3: Details of the pathway annotations for the unigene sets by KEGG. (XLS 1399 kb)

Additional file 4: Gene ID and transcript fold changes of the DEGs (differentially expressed genes) between the wild type and mutant. (XLS 2888 kb)

Additional file 5: DEG-enrichment analysis by KEGG metabolic pathway classification. (XLS 619 kb)

Additional file 6: A total of 110 MADS-box genes in C. goeringii. (XLS 163 kb)

Additional file 7: Gene ID and fold changes of the unigenes involved in plant hormone signal transduction pathway annotated by KEGG pathway. (XLS 42 kb)

Additional file 8: Length distribution of small RNAs derived from Cymbidium goeringii 'Songmei' (WT) and the multi-tepal mutant 'Yuhudie'. (XLS 1182 kb)

Additional file 9: Distribution of the conserved miRNA families in Cymbidium goeringii 'Songmei' (WT) and the multi-tepal mutant 'Yuhudie'. (XLS 46 kb)

Additional file 10: putative target genes of the conserved miRNAs predicted in this study. (XLS 403 kb)

Additional file 11: Predicted miRNA target genes involved in floral development/flowering time. (XLS 79 kb)

Additional file 12: Phylogenetic analysis of Cymbidium goeringii TCP-like genes with their homologues in Arabidopsis and the sequence complementary to miR396 using RNA hybrid software (http://bibiserv.techfak.uni-bielefeld.de/mahybird). (TIF 496 kb)

Additional file 13: Gene ID and fold changes of the unigenes regulating floral homeotic genes and floral patterning. (XLS 43 kb)

Additional file 14: The primers of stem-loop RT-PCR and qRT-PCR. (DOC 37 kb)

Abbreviations

AGC: AGAMOUS; AHP: Histidine phosphotransfer protein; AP1: APETALA1; AP2/ERF: APETALA2/Ethylene Responsive Factor; AP3: APETALA3; ARF: Auxin response factor; ARR: Response regulator; Aux/IAA: Auxin/Indole-3-acetic acid; BAK1: Barely Any Meristem; BHLH: Basic helix-loop-helix; bZIP: Basic leucine zipper; CER: Cytokinin receptor; CLF: Curly Leaf; COG: Clusters of Orthologous Groups; DEF: DEFICIENS; DEGs: Differentially expressed genes; EMF2: Embryonic Flowers; FAR: Far-red impaired Response; GAI: GA insensitive; GH3: Auxin-responsive Gentiopicroside 2; GIF: GRF-interaction Factor; GO: Gene ontology; GRF: Growth-regulating factor; HD: Homeobox-leucine zipper factors; KEGG: Kyoto encyclopedia of genes and genomes; LUG: Leunig; MYB: Myeloblastosis transcription factors; NAC: NAM, ATAF, and CLC transcription factor; NF-Y: Nuclear transcription factor; NIN: Nodule inception; RGL: Repressor of GA-Like; RPM: Reads per million; SAM: Shoot apical meristem; SAUR: Small auxin up RNA; SPL: SQUAMOSA promoter binding protein-like; TALE: Three amino acid loop extension family; TCP: TEOSINTE BRACTHE1/CYCLOIDEA/PC transcription factors; WUS: Wuschel.

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Availability of data and materials

Uploaded as Additional files 3, 4, 5, 6 and 7, sequencing clean data were uploaded to the Sequence Read Archive (SRA) database of NCBI (Genbank accession number SRP091422).

Authors’ contributions

FX Y and GF Z designed the experiments, analyzed the data and wrote the manuscript with input from all authors. Z W, CY Z helped design the study and analyzed the data; FX Y, GF Z and Z W executed the experiments and assembled the figures. HL Q, XQ X and D H conducted the stem-loop qRT-PCR for miRNA and qRT-PCR for their putative targets. All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

Consent for publication

Not applicable.

Ethics approval and consent to participate

Not applicable.

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