miR390 family of Cymbidium goeringii is involved in the development of reproductive organs in transgenic Arabidopsis

Zihan Xu1, Qian Liu1, Yue Chen2, Yuanhao He1 and Fengrong Hu1*

Abstract
Background: miR390s is an ancient family with a high level of conservation among plant miRNAs. Through the auxin signal transduction pathway, miR390 participates in diverse biological processes of plant growth and development. As an important Chinese traditional orchid, Cymbidium goeringii has unique flower shape and elegant fragrance. But its development has been greatly restricted because of the low flower bud differentiation and the difficult reproduction. This study aims to provide guidance for the role of cgo-miR390 in reproductive organ development to enhance the ornamental and economic value of Cymbidium.

Results: MIR390a, MIR390b and MIR390c of C. goeringii were cloned, and their length ranged from 130 to 150 nt. Each precursor sequence of cgo-miR390 contains 2 to 3 mature miRNAs. Three kinds of cgo-miR390s displayed distinct temporal and spatial expression patterns during floral development in C. goeringii. The overexpression of MIR390s alters morphology and function of stamens and pistils in Arabidopsis, such as enlargement of anther aspect ratio and separation of stylar and stigmas, which affects the development of fruits and seeds. In particular, the pollen amount decreased and the seed abortion rate increased in cgo-MIR390c-overexpressed plants.

Conclusions: cgo-miR390 family affected the development of reproductive organs in transgenic Arabidopsis. The study provides references for the genetic improvement for orchid with potentially great economic benefit.

Keywords: Cymbidium goeringii, miR390, Reproductive organ, Flower development, Arabidopsis

Background
MicroRNAs (miRNAs) are a class of non-coding endogenous small RNAs with 20–24 nucleotide (nt) in length [1], which have been identified as new regulators of gene expression at transcriptional, post-transcriptional and post-translational levels [2]. The ancient and highly conserved miRNA, miR390, plays an important role in diverse processes of plant growth and development, including apical dominance, leaf senescence, root formation and abiotic and biotic stress responses, through the auxin signal transduction pathway. According to the miR-base database, miR390 family comprises no more than 4 members in most plants, such as only two members in Arabidopsis thaliana [3]. It has shown that some Auxin response factors (ARFs) are the target genes of miR390. Notably, miR390 triggers the production of trans-acting (ta) siRNAs from TAS3 transcripts, which regulated ARFs in trans [4, 5].

miR390-TAS3-ARF regulatory network confers sensitivity and stability to auxin responses in various tissues and developmental stages. During root development, an auxin-response element (AuxRE) in the promoters of miR390 was bound by ARFS/MONOPTEROS (MP), which affects the expression of miR390 in the transit-amplifying compartment of the root meristem.
then modulated the abundance of ARF2/3/4 by TAS3 [6]. miR390 was also specifically expressed at the sites of lateral root (LR) initiation where it triggers the biogenesis of ta-siRNAs to inhibit ARF2/3/4, and these ARFs affect auxin-mediated 390 accumulation as well [7, 8]. Therefore, an ARF-miR390-TAS3 tasiRNA-ARF regulatory network was established due to the positive and negative feedback regulation of miR390 by ARF2/3/4. In leaf development, it is likely that AGO associates with miR390 and cleaves TAS3 precursor RNAs as ta-siRNAs. After DCL4-mediated processing, this complex cleaves their targets, including ARF3 and ARF4, to regulate leaf morphology development in Arabidopsis [9–11]. Dissimilarly, the expression of miR390 in leaf polarity of maize is established and maintained independently of the ta-siRNA pathway [12]. In addition, miR390a has been previously reported to be involved in dark-induced leaf senescence, while ARF2, a target gene of miR390, positively regulated auxin-mediated leaf longevity in Arabidopsis [13–15].

Results

Sequence analysis of miR390s in Cymbidium goeringii

MIR390a, MIR390b and MIR390c sequences were identified based on the transcriptome sequence data. Then the three sequences were cloned from the leaf of C. goeringii ‘Songmei’, and their length ranged from 130 to 150 nt (Table S1). Sequencing results showed that these three cgo-MIR390s produced a total of 8 mature miRNAs, in which the sequence of 3 mature miRNAs each detected on MIR390a and MIR390b were identical (Table 1; Fig. 1A). Predicted stem-loop structures of three precursor sequences were shown in Fig. 1, all of them had 2 nt overhang structure at the distal end of their arms, which resulted from Dicer cleavage [19]. According to reference [20], the minimal folding free energy indexes (MFEIs) of cgo-MIR390a/b/c were 1.00, 1.35 and 1.18, respectively. These calculation results indicated that the sequences

Table 1  List of gene information

| Gene type       | Gene name     | Inclusion relation | Sequence length |
|-----------------|---------------|--------------------|-----------------|
| Precursor sequences | cgo-MIR390a   | ●■▲               | 145bp           |
|                 | cgo-MIR390b   | ●■▲               | 133bp           |
|                 | cgo-MIR390c   | ●■                | 150bp           |
| Mature miRNAs   | cgo-miR390a-5p| ●                 | 21nt            |
|                 | cgo-miR390b-5p| ●                 | 21nt            |
|                 | cgo-miR390c-5p| ●                 | 21nt            |
|                 | cgo-miR390a-3p.1 | ▲          | 21nt            |
|                 | cgo-miR390b-3p.1 | ▲          | 21nt            |
|                 | cgo-miR390c-3p  | ▲                | 21nt            |
|                 | cgo-miR390a-3p.2 | ▲          | 21nt            |
|                 | cgo-miR390b-3p.2 | ▲          | 21nt            |
Fig. 1  Sequence analysis of cgo-MIR390s. A The secondary structures of cgo-MIR390s in Cymbidium goeringii. B The phylogenetic relationship of cgo-MIR390s with other homologous sequences. The triangles represent C. goeringii, the green dots represent monocotyledons, and the yellow dots represent dicotyledons.
were all most likely to be miRNA due to the MFEIs were more than 0.85. In addition, no secondary stems or large loops interrupted the miRNA:miRNA* duplex, and there were fewer than 5 mismatched positions in duplex. These features are consistent with the universal characteristics of pre-miRNAs [21].

Analysis of homologous sequences found that three MIR390 genes of C. goeringii showed strong homology, and all clustered together within the same branch of the phylogenetic tree (Fig. 1B). Furthermore, MIR390s in monocot plants display a higher sequence identity with cgo-MIR390s than those in dicots. It is obvious that this kind of pre-miRNA in monocot and dicot plants underwent evolutionary diversification. For analysis of mature miRNAs, cgo-miR390a-5p (cgo-miR390b-5p, cgo-miR390c-5p) was the most conserved one, followed by miR390a-3p.1 (miR390b-3p.1, miR390c-3p) and miR390a-3p.2 (miR390b-3p.2) (Fig. 2). This may be related to the evolution of miR390 family in C. goeringii as well.

Expression analysis of cgo-miR390s in Cymbidium goeringii

All three miR390s were expressed in each floral organ of fully open flowers. The expression of cgo-miR390a-5p was specific to the ovary (Fig. 3 A), whereas cgo-miR390a-3p.1 was mainly expressed in lateral petal and stamen. By contrast, the expression of cgo-miR390a-3p.2 was extremely low in all floral organs. During flower bud development, cgo-miR390a-5p and cgo-miR390a-3p.1 had the highest expression when the flower bud length was 1-3 cm (Fig. 3B), while cgo-miR390a-3p.2 has the lowest at that time. When the length of flower buds was 3-5 cm, these three miR390s were all expressed at about the same level. These three kinds of cgo-miR390s showed overall non-overlapping expression patterns both temporally and spatially, which suggested they probably have functional divergence in regulating Cymbidium flower development (Fig. 3).

Effect of cgo-miR390s on stamens and pistils development in Arabidopsis

To investigate the function of cgo-miR390 family in flower development, we overexpressed cgo-MIR390s in Arabidopsis. The flowering time of wild-type and transgenic plants was almost consistent (Fig. 4A). By observing the external morphology of fully open flowers, compared with wild-type (WT) plants, the pollen attached to the anthers of the three transgenic plants was less, which was most obvious in cgo-MIR390c-overexpressing Arabidopsis (Fig. 4B). We speculate that this might be due to anthers’ early maturation, early dehiscence and insufficient pollen production.

Alexander staining was performed on the stamens of unopened flowers to further confirm this phenomenon. Figure 5 A showed that the anther aspect ratio (anther width/anther length) of three cgo-MIR390s-overexpressed plants was all larger than that of wild-type, suggesting that these three genes all regulated the morphological development of stamens. At this time, the pollen of wild-type did not reach full maturity. In contrast, pollen cells of cgo-MIR390a and MIR390b over-expressing plants were fully mature and kept inside the locules of anther, while the anther of cgo-MIR390c-overexpressing plants was prematurely dehisced and pollen grains were released (Fig. 5 A). Figure 5B-D also illustrated that the pollen amount from fully open flowers of

![Fig. 2 Nucleotide sequence alignment of cgo-miR390s with other homologous sequences](image-url)
**Effect of cgo-miR390s on fruits and seeds development in Arabidopsis**

Although all transgenic plants flowered at a similar time as the wild-type, their senescence time was earlier than WT, especially in the reduced flowering (Fig. 6) and fruit ripening time (Fig. 7 A). By observing under a stereomicroscope, the fruit pods of all transgenic plants were thicker than those of the wild-type, with those of cgo-MIR390a and MIR390b overexpressed plants longer than MIR390c and WT (Fig. 7 A and B). The anatomical observation of fruits was further performed, which showed that the three genes overexpressed plants had different degrees of ovule abortion, and the seed volume of transgenic plants was larger than that of WT (Fig. 7 C). These results further indicated that cgo-miR390s were involved in fruit development by regulating the development of floral organs.

**Discussion**

miR390s in plants are the ancient family of miRNAs with a high level of conservation among species [22]. Mature sequence analysis indicated that the three kinds of miR390s in *Cymbidium goeringii* were conserved. Secondary structure prediction showed that the mature miRNAs were all clustered in the same precursor. Studies of miRNA clusters in animals showed that evolutionarily conserved miRNAs are significantly enriched in clusters. The miRNAs in the same cluster have cooperative effects, which is called “functional co-adaptation” model [23]. We then analyzed the expression patterns by Quantitative real-time PCR (qRT-PCR), observing that the expression levels of cgo-miR390a-5p (miR390b-5p, miR390c-5p) and cgo-miR390a-3p.1 (miR390b-3p.1, miR390c-3p) were opposite in ovary but the expression trends of them in other floral organs were similar. During floral buds development, the highest expression of cgo-miR390a-5p and cgo-miR390a-3p.1 was obtained when the floral buds grew to 1-3 cm and significantly decreased after this stage. In particular, the expression pattern of miR390a-3p.2 (miR390b-3p.2) was different from the other two kinds of cgo-miR390s not only in floral organs but also in floral bud development. The above results indicated that there may be synergy between cgo-miR390a-5p and

![Fig. 3](image-url)

**Fig. 3** Expression patterns of cgo-miR390s in *Cymbidium goeringii*. A Expression level of miR390s in various floral organs. Se, sepal; Li, lip; La, lateral petal; St, stamen; Ov, ovary. B Expression level of miR390s in developing stages of floral buds. The mean lengths of flower buds at different stage were expressed in cm. The different letters indicate significant differences among 5 samples within a single group (P < 0.05).
cgo-miR390a-3p.1, while miR390a-3p.2 were inclined to play roles independently. These inferences seemed to be further substantiated by the functional verification in *Arabidopsis*.

The overexpressed plants of cgo-miR390s cluster showed similar phenotypes in some of the organs, such as pod thickening, anther aspect ratio enlargement, and seed abortion. Moreover, the similarity of functions between cgo-MIR390a and cgo-MIR390b was more clear, while the phenotypes of cgo-MIR390c-overexpressed plants were different. This might be caused by the lack of a mature miRNA in cgo-MIR390c, which needs to be validated by further experiments. Of course, there are some differences between cgo-MIR390a and cgo-MIR390b as well, such as the effect on pods morphology. Although both cgo-MIR390a and MIR390b produce mature miRNAs, the stem-loop structure and precursor sequences between them were all different. In addition, the expression of mature miRNAs was not linearly associated with that of precursor miRNAs, which might be regulated at several levels [24]. These might be the reasons for their functional difference.

Next, how cgo-miR390s is involved in the regulation of reproductive organs development remains an open

![Figure 4](image-url)  
**Fig. 4** A Flowering time observation of wild type and transgenic *Arabidopsis* plants. B Flower characterization of wild type and transgenic *Arabidopsis* plants. The first line showed the top view of flowers and the second line showed the side view of flowers under a stereomicroscope.

![Figure 5](image-url)  
**Fig. 5** Phenotype changes in floral organs of wild type and transgenic *Arabidopsis* plants. A The stamens staining with Alexander solution under an optical microscope. B The pollen grains staining with Alexander solution under an optical microscope. C-D The statistical results of the pollen stains' number and staining rate under an optical microscope with 10x magnification. E In vitro pollen germination and pollen-tube growth under an optical microscope with 20x magnification. F-G The statistical results of pollen germination rate and pollen tubes' length. H The stigma of pistils under an electron microscope. The different letters indicate significant differences (P<0.05).
Fig. 5 (See legend on previous page.)
question, but we offer some possibilities. Existing studies indicated that miR390s play a potential role in the anther development and regulation of male-sterility in tomato, which was similar to the results in our study. The researchers speculated that it might be related to Ta siARFs, even though their regulatory mechanisms are poorly understood [25]. In Arabidopsis, the miR390-TAS3-ARF2/3/4 regulatory network is an integral part of the regulatory networks mediating auxin response [6, 26]. Flower phenotypes of arf2 mutants showed infertility of the early-produced flowers, which was probably caused by the early elongation of gynoecium [27]. TEX1- and TAS3-mediated restriction of ARF3 expression limits excessive megaspore mother cell formation [28]. The overexpressed plant of cgo-MIR390c showed a tendency towards these phenotypes. Moreover, the ETT/ARF3 mutant exhibited some developmental abnormalities such as an expansion of the stylar and stigmatic regions [29]. Seed size and weight were dramatically increased in an AtARF2 mutant [30]. These were very similar to all cgo-miR390s-overexpressed plants. Notably, three cgo-MIR390s overexpressed plants exhibited different degrees of ovule abortion, but Alexander staining and in vitro pollen germination assays showed that the pollen collected from transgenic Arabidopsis had normal viability. Therefore, we speculated that the ovule abortion of transgenic plants could result from aberrant development of female reproduction, such as stigma defect shown in Fig. 5 H, transmission tract defect and/or female gametophyte defects [31]. Of these, failed fertilization in cgo-MIR390c-overexpressed plants may also be caused by premature release of pollen grains. In addition, the miR390-TAS3-ARF pathway was also involved in other biological processes of non-model plants like the response to salt stress [32]. But this functionally important and archetypal regulatory pathway in land plants occurred significant variation [33]. All of these have brought more light to the verification experiments of target genes and the regulatory mechanism of cgo-miR390s in reproductive organs.

Conclusions
In conclusion, cgo-miR390 family plays important roles in the reproductive organs of plants. Three kinds of cgo-miR390s displayed distinct temporal and spatial expression patterns during floral development in Cymbidium goeringii. The results also demonstrate that the overexpression of MIR390s alters the morphology and function of stamens and pistils in Arabidopsis, which affects the development of fruits and seeds. These findings preliminarily reveal the functions of cgo-miR390s and suggest that they can potentially be used for genetic improvement for orchid.

Methods
Plant materials and growth conditions
The Songmei cultivar of C. cymbidium was cultivated in a naturally lit glasshouse in the Institute of Horticulture, Zhejiang Academy of Agricultural Sciences (Hangzhou, Zhejiang, China). The Arabidopsis variety Columbia (Col-0) was used for gene overexpression and was grown at (22±1) °C, 16 h day/8 h night in an artificial climate chamber.

Extraction of RNA and qRT-PCR analysis
Total small RNA was extracted from the plants using MiniBEST Universal RNA Extraction Kit (Takara, Dalian, China). The total small RNA was then reverse transcribed into cDNA using miRNA 1st Strand cDNA Synthesis
Kit (Vazyme, Nanjing, China) with stem-loop primer. qRT-PCR reaction solution was configured according to the instruction of ChamQ Universal SYBR qPCR Master Mix (Vazyme, Nanjing, China), and the action was performed using a StepOnePlus real-time PCR system (Thermofisher, Waltham, USA) under the following conditions: polymerase activation at 95°C for 5 min, followed by 40 cycles at 95°C for 15s and 60°C for 1 min. We used 18 S rRNA and U6 as internal reference genes for *C. goeringii* and *Arabidopsis*, respectively. The threshold cycle (Ct) values of the PCR were averaged, and the relative transcript levels were quantified using the $2^{-\Delta\Delta Ct}$ method. The sequences of all the primers for this experiment are shown in Table 2.

**Cloning of cgo-miR390 family and bioinformatic analysis**

The putative precursors of cgo-miR390 family, *MIR390a* (145 bp), *MIR390b* (133 bp), and *MIR390c*...
(150 bp), were amplified with the primer pairs listed in Table 2, and then cloned into pBI121 vector at double restriction sites (XbaI and SmaI) to construct into recombinant vector. Gene amplification was performed by a standard PCR procedure, and the annealing temperatures were all 60 °C.

Secondary structure and minimum free energy (MFE) prediction were performed with the online tool RNAfold (http://rna.tbi.univie.ac.at/). Homologous sequence of precursors and matures were download from miRbase (http://www.mirbase.org/index.shtml), and then aligned in DNAMAN 8.0 software. The phylogenetic tree was constructed using MEGA 6.0 with the neighbor joining (NJ) method.

Transformation into Arabidopsis
pBI121-35 S::MIR390a, pBI121-35 S::MIR390b and pBI121-35 S::MIR390c were transformed into Arabidopsis Col-0 by the Agrobacterium tumefaciens-mediated floral dipping method [34]. T0 seeds were screened on MS media containing 50 mg L⁻¹ kanamycin to obtain T1 plants, and the gene insertions were confirmed by detection of genomic DNA PCR (Fig. S1). T2 plants were obtained through phenotypic observation and gene expression measurement. Screening was performed until homozygous lines of T3 generation were obtained.

In vitro pollen germination and pollen-tube growth
Pollen was collected from fully open flowers and germinated on Arabidopsis thaliana pollen medium according to the procedure of Fan et al. [35] with minor modifications. This medium consisted of 1mM KCl, 10mM CaCl2, 0.8mM MgSO4, 1.5mM boric acid, 10 mg/L myo-inositol, 5mM Mes, 18% (w/v) sucrose and 1.0% (w/v) agarose. And the pH was adjusted to 5.8 using Tris-hydroxymethyl aminomethane (Tris) base. Pollen germination was determined by microscopy after 10 h incubation in a chamber at 25 °C and 100% relative humidity. The pollen grains were recorded as germinated when the pollen tube length was equal to or greater than the pollen grain diameter, as described by Boavida and McCormick [36].

Alexander staining of stamens and pollen
For examination of stamen samples, the stamens were collected from flowers that were about to bloom, when the anthers had not yet split but the pollens were mature. The stamens were fixed in Alexander stain solution (Jiskang, Qingdao, China) at 55°C for 3 h, and observed under the optical microscope.

For examination of pollen samples, 30 fully open flowers were randomly collected from the plants and mixed with ddH2O in a 1.5 mL centrifuge tube by vortex for 2 min, then took the flowers out. After centrifugation at 11,000 rpm for 1 min (at room temperature), the

Table 2 List of primers sequences used for the experiments

| Sequence type       | Gene name   | Primer sequences                                                                 |
|---------------------|-------------|----------------------------------------------------------------------------------|
| Precursor cloning   | cgo-MIR390a | F: gagaacacggggactctagaaaagcagggagaaaattcttaag                  |
|                     | cgo-MIR390b | R: ataaagggactcaggacccgggagacaggagagaaaccatatggaact | |
|                     | cgo-MIR390c | R: ataaagggactcaggacccgggagacaggagagaaaccatatggaact | |
| Reverse transcription| cgo-miR390a-5p| GTGTATCCAGTCAGGGTTCCAGGGTTTCCAGCTTGCACCTGAGCACAGCCTCA |
|                     | cgo-miR390a-3p.1 | GTGTATCCAGTCAGGGTTCCAGGGTTTCCAGCTTGCACCTGAGCACAGCCTCA |
|                     | cgo-miR390a-3p.2 | GTGTATCCAGTCAGGGTTCCAGGGTTTCCAGCTTGCACCTGAGCACAGCCTCA |
| Fluorescence quantification | cgo-miR390a-5p | F: CGCGAGAGCTCGAGGGAGATA |
|                     | cgo-miR390a-3p.1 | R: AGTGCAGGGTCCAGGGTTTCCAGCTTGCACCTGAGCACAGCCTCA |
|                     | cgo-miR390a-3p.2 | R: AGTGCAGGGTCCAGGGTTTCCAGCTTGCACCTGAGCACAGCCTCA |
|                     | 18S         | F: GTCTCTATTGTTGGCCT |
|                     | U6          | R: TCGCAGTGGTTGCCTTGCCT |
|                     |             | F: GTGTCATAGAGAGGGAGAAT |
|                     |             | R: CTCCTTTCTTTGTGAACCTCA |
supernatant was discarded. The pollen precipitation was fixed in Alexander stain solution at 55°C for 3 h. Then the morphology of pollen was observed under a microscope, and the number, coloration rate and aberration rate of pollen were counted.

### Scanning electron microscopy (SEM)

The pistils collected from fully open flowers were infiltrated in 4% glutaraldehyde (Electron Microscopy China, Beijing, China) overnight. Samples were dehydrated in an ethanol series, dried by EMITECH-K850 (Quorum, Hertfordshire, England) and sputter coated with Platinum. Images were taken with a Quanta 200 Scanning Electron Microscope (FEI, Hillsboro, USA) at an accelerating voltage of 25 kV.

### Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s12870-022-03539-3.

**Additional file 1.**
**Additional file 2.**
**Additional file 3.**
**Additional file 4.**
**Additional file 5.**
**Additional file 6.**
**Additional file 7.**
**Additional file 8.**

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Not application.

**Plant ethics**

Experimental research and field studies on plants (either cultivated or wild), including the collection of plant material are conducted in compliance with relevant institutional, national, and international guidelines and legislation.

**Authors’ contributions**

ZX and QL conducted the experiments, analyzed the data and wrote the manuscript. ZX and YH edited the manuscript. YC and FH designed the experiment and edited the manuscript. All authors read and approved the manuscript.

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### Availability of data and materials

All data generated or analyzed during this study are included in this published article and its supplementary information files. The precursor sequences and mature miRNAs of cgo-miR390s described here are available in the Supplemental Files and at Genbank: OM824433–OM824435.

### Declarations

**Ethics approval and consent to participate**

There are no ethical issues involved.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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