Characterization of the complete chloroplast genome of *Coelogyne fimbriata* (Orchidaceae)

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**ABSTRACT**

*Coelogyne fimbriata* has been classified as a national second-class protected orchid species in China. In this study, we report and characterize the complete chloroplast (cp) genome sequence of *C. fimbriata* in an effort to provide genomic resources useful for promoting its conservation and systematic research. The complete genome is 159,010 bp in length and the overall GC content is 43.3%. The cp genome sequence has a typical quadripartite structure, comprising two inverted repeats (IRA and IRB) regions, which are separated by a small single-copy (SSC) region and a large single-copy (LSC) region. Moreover, a total of 135 functional genes were annotated, including 89 protein-coding genes, 38 tRNA genes, and 8 rRNA genes. The phylogenetic analysis recovered a close relationship between *C. fimbriata* and *Pleione formosana*, and both species are placed within the tribe Arethuseae (Orchidaceae).

*Coelogyne fimbriata* Lindl. (Orchidaceae) is native to a wide range of countries from Nepal, Bhutan, southern Fujian, Guangdong, Guangxi, Guizhou, Hainan, Hong Kong, Jiangxi, Sichuan, southeastern Xizang, Yunnan of China, northeastern India, Myanmar, Thailand, Vietnam, Cambodia, Laos to Peninsular Malaysia. This orchid grows in more humid areas on trees and rock walls, which have ovoid to ellipsoid pseudobulbs. It has yellowish-green flowers with reddish-brown markings. The epithet refers to the fimbriate margins of the lip of the flower (Clayton 2002).

*Coelogyne fimbriata* has been classified as a national second-class protected orchid species in the Information System of Chinese Rare and Endangered Plants (ISCRPE; http://www.iplant.cn/rep/prot/Coelogyne). It is urgent to design effective conservation strategies for Chinese *C. fimbriata* populations because southern China is the northernmost edge of its distribution region where the populations are subject to ecological marginality and vulnerable to disturbance (Channell and Lomolino 2000; Swarts and Dixon 2009). Furthermore, given that *C. fimbriata* is so variable in morphology that scientists have defined many different species (Gravendeel et al. 2001), it is more necessary to protect the germplasm resources of this species.

The samples of *C. fimbriata* were collected from Nanling National Nature Reserve, Northwest Guangdong, China (112.35E, 24.40N) and cultivated in the greenhouse of South China Botanical Garden, Guangzhou, China (113.36E, 23.18N). The voucher specimen (WuQiu018) was deposited in the herbarium of South China Botanical Garden (IBSC). Total genome DNA was extracted with the Trelief plant genomic DNA kit (TsingKe Biotechnological Technology, Beijing, China) and tested for gel electrophoresis and concentration detection.

The high-throughput sequencing of the plastid genomes was performed using the Illumina HiSeq 2000 at the Beijing Genomics Institute (Beijing, China). A total of 1.73GB clean data were used for assembling the sequence as described by Jin et al. (2018). The plastid genome sequences of *Pleione bulbocodioides* (KY849819.1), *Pleione formosana* (MK361027.1), *Bletilla ochracea* (KT695602.1), and *Bletilla striata* (KT588924.1) were used as references to annotate the plastome using the PGA (Qu et al. 2019). Geneious R11 (Kearse et al. 2012) was used to check the accuracy of the assembly, and to adjust the start/stop codons and intron/exon boundaries of the annotation. The raw sequencing reads used in this study has been deposited in SRA with the accession number SRX12560166 and the annotated chloroplast (cp) genome sequence has been deposited into the GenBank with the accession number MT705722.

The complete cp genome sequence of *C. fimbriata* was found to be 159,010 bp in length. A total of 135 genes were annotated, including 89 protein-coding genes, 38 tRNA genes, and 8 rRNA genes. It has a typical quadripartite structure containing two short inverted repeat (IRA and IRB).
regions (26,350 bp, each), separated by a small single-copy (SSC) region (18,835 bp) and a large single-copy (LSC) region (87,475 bp). The overall GC content is 43.3%, and the A, T, C, G base composition of the genome is 30.9, 31.7, 19, and 18.4%, respectively.

To investigate the phylogenetic position of *C. fimbriata*, a maximum-likelihood (ML) phylogenetic tree was constructed in IQ-TREE 1.6.12 with default parameters (Nguyen et al. 2015), based on 19 complete plastid genome sequences downloaded from the GenBank (Figure 1). We conducted sequences alignment using the MAFFT v.7.0.17 (Katoh and Standley 2013) plugin in Geneious, using *Goodyera fumata* (GenBank: NC_026773) and *Anoectochilus emeiensis* (GenBank: NC_033895) as outgroups. Our phylogenetic analyses indicate that *C. fimbriata* and *P. formosana* form a well-supported sister lineage, with *Bletilla formosana* as a successive closer taxon within the tribe Arethuseae. The complete cp genome of *C. fimbriata* will contribute to the development of conservation strategy for this rare orchid, as well as for further phylogenetic studies.

**Disclosure statement**

No potential conflict of interest was reported by the author(s).

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**Data availability statement**

The data that support the findings of this study are openly available in GenBank of NCBI at https://www.ncbi.nlm.nih.gov/, reference number MT705722.
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