The complete chloroplast genome of Cryptomeria japonica var. sinensis (Cupressaceae)

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ABSTRACT
The complete chloroplast genome of Cryptomeria japonica var. sinensis Miquel was assembled and analyzed. The chloroplast genome of C. japonica var. sinensis Miquel did not have a typical quadripartite structure with the inverted repeats (IR) absent, and the size of C. japonica var. sinensis is 131,412 bp. The overall GC content was 35.4%. The genome encoded a set of 119 genes, containing 83 protein-coding genes, 32 tRNA genes, and 4 rRNA genes. Phylogenomic analysis indicated that C. japonica var. sinensis is sister to C. japonica (Thunb. ex L. f.) D. Don.

Cupressaceae is a family of coniferous plants, and Cryptomeria japonica (Thunb. ex L. f.) D. Don is a modeling species of Cupressaceae, which is not only beautiful in tree figure, but also capable of purifying the air and improving the environment. It is mainly distributed in the high altitude regions of China and is an important ornamental tree species (Dou et al. 2010). Currently, its natural forests are mainly distributed in Wuyi Mountain in Fujian and Tianmu Mountain in Zhejiang across southeast China, and broadly introduced at different provinces in China. Consequently, it has great prospects and the genetic resources are worth preservation (Wang et al. 2007). According to the Flora of China, there are obvious morphological differences between C. japonica var. sinensis Miquel and C. japonica (Thunb. ex L. f.) D. Don. However, whether their chloroplast genomes differed remains unclear. In this study, the completed chloroplast genome sequence of C. japonica var. sinensis Miquel is reported contributing for better understanding its evolution and population genetics, and also providing significant information for the phylogeny of Cupressaceae.

The fresh leaves of C. japonica var. sinensis Miquel were collected from Fujian province, China (Xiapu, Ningde: 119°56′37″E, 26°52′19″N). Voucher specimen (specimen code FJFC8003) was deposited in the Key Laboratory of Integrated Pest Management in Ecological Forests, Fujian Province University, Fujian Agriculture and Forestry University. Total genomic DNA was extracted from 100 mg fresh leaves using a modified CTAB method (Murray and Thompson 1980). Libraries with an average length of 350 bp were constructed using the NexteraXT DNA Library Preparation Kit. Sequencing was performed on the Illumina NovaSeq 6000 platform (Total Genomics Solution Limited, SZHT), and the average length of the generated reads was 150 bp. The Illumina raw sequence reads were edited using the NGS QC Tool Kit version 2.3.3 (National Institute of Plant Genome Research (NIPGR), New Delhi, India). High-quality reads were assembled into contigs using the de novo assembler SPAdes version 3.11.0 (Bankevich et al. 2012) and finally annotated by PGA software (Qu et al. 2019).

Like some gymnosperms chloroplasts, the chloroplast genome of C. japonica var. Sinensis Miquel (MT554702) did not have a typical quadripartite structure with the inverted repeats (IR) region absent, and the size of C. japonica var. sinensis Miquel is 131,412 bp. The overall GC content was 35.4%. The genome encoded a set of 119 genes, containing 83 protein-coding genes, 32 tRNA genes, and 4 rRNA genes. To confirm the phylogeny of C. japonica var. Sinensis Miquel, 12 complete genomes were obtained from GenBank, and were aligned using HomBlocks software (Bi et al. 2018). A maximum likelihood (ML) tree was inferred using RAXML version 8.2.9 (Stamatakis 2014), with the combined rapid bootstrap (1000 replicates) and search for ML tree (the ‘-f a’ option). The result of ML phylogenetic tree showed that C. japonica var. sinensis Miquel is closely related to C. japonica (Thunb. ex L. f.) D. Don.

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The chloroplast genome of *C. japonica* var. *sinensis* Miquel will provide useful genetic information for further study on genetic diversity and conservation of Cupressaceae species.

**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 at [https://www.ncbi.nlm.nih.gov/nuccore/MT554702/](https://www.ncbi.nlm.nih.gov/nuccore/MT554702/), reference number MT554702.

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