Cladrastis yunchunii X. W. Li et G. S is a plant species belonged to the family Papilionaceae. Cladrastis yunchunii is currently found in broad-leaved forests in the limestone area of Luxi County, Yunnan Province. It is suitable for afforestation and urban greenin in limestone areas. In this study, for the first time, we report the complete chloroplast genome sequence of C. yunchunii. We sequenced and assembled the entire chloroplast genome of C. yunchunii. The chloroplast genome was determined to be 158,250 bp in length. It contained large single-copy (LSC) and small single-copy (SSC) regions of 84,930 bp and 12,664 bp, respectively, which were separated by a pair of inverted repeats (IR) regions of 30,328 bp. The genome contained 132 genes, including 87 protein-coding genes, 8 rRNA genes, and 37 tRNA genes. The overall GC content of the whole genome is 38.1%, and the corresponding values of the LSC, SSC, and IR regions were 36.4%, 33.6%, and 41.3%, respectively. Phylogenetic analysis suggested that C. yunchunii is closely related to the genus Ormosia in the Papilionaceae.

Until now, the genome research on C. yunchunii has not been revealed, and the complete genome sequence of its chloroplast has not been reported. Complete chloroplast genomes are useful tools for providing the information of species identification, phylogenetic relationships, and germplasm diversity (Wang et al. 2019). In this study, we reported the complete genomic sequence of C. yunchunii. And the chloroplast genome will provide a useful resource for C. yunchunii conservation as well as for the phylogenetic studies of Papilionaceae.

The chloroplast genome is a circular DNA molecule (Tonti-Filippini et al. 2017). It is highly conserved in plants and an ideal for ecological, evolutionary, and diversity studies (Wicke et al. 2011). So far, the complete chloroplast genomes of some species in the family Papilionaceae have been studied and saved deposited in the GenBank database, e.g. Ormosia hosiei (Zhai et al. 2019) and Pterocarpus macrocarpus (Zhang et al. 2020). But there is little known about the genome information of C. yunchunii. Therefore, we studied the complete sequence of the chloroplast genome of C. yunchunii and compared it with other known chloroplast genomes to determine phylogenetic relationships among in family Papilionaceae.

The fresh young and healthy leaves of C. yunchunii were collected from Southwest Forestry University Kunming, China (Yunnan, China; geospatial coordinates: 25°04’00”N, 102°45’41”E). The voucher specimens of C. yunchunii were preserved in the herbarium of Southwest Forestry University (Voucher code: SWFU-XW919028). Total genomic DNA was extracted using the Magnetic beads plant genomic DNA prep Kit (TSINGKE Biological Technology, Beijing, China). Then DNA were sequenced using the Illumina HiSeq 2000 platform and the compiled chloroplast genome were assembled using Velvet and Getorganelle software (Zerbinio and Birney 2008; Jin et al. 2020). Genome annotation was performed with the online annotation tool DOGMA (Dual Organellar GenoMe Annotator, http://dogma.ccbj. utexas.edu/) and program Geneious R8 (Biomatters Ltd, Auckland, New Zealand) (Kearse et al. 2012). Finally, the annotated genomic sequence was submitted to GenBank under accession number MW410239.

The complete chloroplast genome of C. yunchunii was 158,250 bp in length, which was contained large single-copy
(LSC) and small single-copy (SSC) regions of 84,930 bp and 12,664 bp, respectively, and a pair of inverted repeats (IR) regions of 30,328 bp. The overall GC content of the whole genome was 38.1%, and the corresponding values of the LSC, SSC, and IR regions were 36.4%, 33.6%, and 41.3%, respectively. A total of 132 functional genes were contained in the chloroplast genome, including 87 protein-coding genes, 8 ribosomal RNA genes, and 37 transfer RNA genes.

To investigate its taxonomic status, 13 Papilionaceae species plastomes were obtained from the Genbank for phylogenetic analysis, and *Maackia floribunda* (Papilionaceae) was used as out-group. The 13 complete chloroplast sequences were aligned by the MAFFT version 7 software (Katoh and Standley 2013). A maximum-likelihood analysis based on the GTR+F+R2 model was performed with iqtree version 1.6.7 using 1000 bootstrap replicates (Nguyen et al. 2015). The results revealed that *C. yunchunii* was closest to *Ormosia* (Figure 1). This information was crucial for the correct identification of *C. yunchunii* and provides valuable genetic resources for the future development of chloroplast derived molecular markers. In conclusion, the complete chloroplast genome of *C. yunchunii* was decoded for the first time, which will facilitate the species identification, molecular biology, and phylogenetic studies of *C. yunchunii*. The chloroplast genome will contribute to the research and conservation of *C. yunchunii*.

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

**Funding**
The study was supported by the National Natural Science Foundation of China [Grant Number: 31860203 & 31760204] and the Youth Top Talent Program of the Ten Thousand Talents Program of Yunnan Province.

**Data availability statement**
The genome sequence data that support the findings of this study are openly available in GenBank of NCBI at [https://www.ncbi.nlm.nih.gov] under the accession no.MW410239. The associated BioProject, SRA, and Bio-Sample numbers are PRJNA687643, SUB8793795, and SAMN17151148 respectively.

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**Figure 1.** Phylogenetic relationships among 13 complete chloroplast genomes (*C. yunchunii* in this study and 12 previously reported species). Bootstrap values based on 1000 replicates were provided near branches. The ML phylogenetic tree for *C. yunchunii* is based on the other 12 species (six in *Sophora*, five in *Ormosia*, and one in *Maackia*) chloroplast genomes.
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