The complete chloroplast genome sequence of the *Alstroemeria* hybrid variety ‘Hanhera’

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**ABSTRACT**

*Alstroemeria,* a member of the Alstroemeriacae family, is a species from South America. The chloroplast genome of *Alstroemeria* spp. was completed by de novo assembly using a small amount of whole genome sequencing data. The chloroplast genome of *Alstroemeria* spp. was 155,672 bp in length consisting of 84,379 bp of large single copy, 17,815 bp of small single copy, and 26,739 bp of a pair of inverted repeat regions. A total of 157 genes were annotated including 103 protein-coding genes (PCGs), 46 tRNA genes, and eight rRNA genes. Maximum likelihood phylogenetic analysis with seven species belonging to the Alstroemeriaeae or Liliaceae family revealed that *Alstroemeria* spp. is grouped with the species in the Alstroemeriaeae family.

*Alstroemeria* belonging to the Alstroemeriaeae family is a popular cut flower plant because of the wide variety of flower colors and includes approximately 60 species (Aker and Healy 1990). The native species are found with many different shades of reds and yellow colors in numerous habitats and the wide color range in the interspecific hybrids arises from this diverse color base found in the species. The white and yellow types of color originated from one interspecific cross and the red and orange types of color originated from several different interspecific crosses (Healy and Wilkins 1986). Interspecific breeding is considered as the most important source of genetic variation in ornamental crops including *Alstroemeria.* However, albinogenes and variegated-color phenotypes caused by plastome-genome incompatibility appear during interspecific breeding in diverse plants, such as the genus, *Pelargonium* (Metzlaff et al. 1982), *Trifolium* (Pandey et al. 1987), *Zantedeschia* (Yao et al. 1994, 1995), etc. Therefore, the information of chloroplast genome of the hybrid ‘Hanhera’ (*Alstroemeria* spp.) obtained in this study will provide an insight of plastid inheritance, investigate more detailed evolutionary aspect, and facilitate breeding of colored *Alstroemeria* efficiently.

The *Alstroemeria* variety ‘Hanhera’ was provided by Prof. Tae-Ho Han, Department of Horticulture, Chonnam National University, South Korea (35.2° N, 126.9° E). It is an *Alstroemeria* hybrid (*Alstroemeria* spp.) derived from a cross between other *Alstroemeria* spp. hybrids ‘Leon’ and ‘Everest.’ Their genetic backgrounds are complex because they are originally derived from interspecific crosses between different species belonging to the *Alstroemeria* genus. The variety ‘Handera’ has white-colored flowers with very faint pink and yellow petal. Its flower size is bigger and plant height is taller than other *Alstroemeria* varieties. The completion of the chloroplast genome sequence was performed at Phyzen bioinformatics pipeline (Kim et al. 2015). According to the PE standard protocol (Illumina, San Diego, CA), an Illumina paired-end (PE) genomic library was constructed with a total genomic DNA and they were sequenced using an Illumina HiSeq2000 platform at Macrogen (http://www.macrogen.com/kor/). The chloroplast genome assembly was performed with approximately 1.9 Gbp of trimmed high-quality reads obtained by a CLC assembly cell package version 4.2.1 (CLC Inc., Aarhus, Denmark). The principal contigs representing the chloroplast genome were retrieved using Nucmer (Kurtz et al. 2004) with the chloroplast genome sequence of *Alstroemeria aurea* (KC968976, Do et al. 2013; Kim and Kim 2013) as reference sequence. The representative chloroplast contigs were arranged in order based on BLASTZ analysis (Schwartz et al. 2003) and connected to a single draft sequence by joining overlapping terminal sequences and manual editing through a comparison with the reference chloroplast genome sequence of *Alstroemeria aurea* (KC968976) as described previously (Cho et al. 2015; Cho and Park 2016). GeSeq program (Tillich et al. 2017) and manual curation based on the results of BLAST searches were used for gene annotation.

The complete chloroplast genome of the *Alstroemeria* hybrid variety ‘Hanhera’ (GenBank accession no. MK327552) was 155,672 bp in length including 26,739 bp inverted repeats (IRa and IRb) regions separated by small single copy (SSC) region of 17,815 bp and large single copy (LSC) region of 84,379 bp. When the chloroplast genome sequence was compared with those of other Alstroemeriaeae species, such as *A. aurea* (KC968976, Do et al. 2013; Kim and Kim 2013) and *Bomarea edulis* (KM233641, Kim et al. 2016), it was slightly longer and the sequence conservation between them was expectedly very
high with a sequence identity of 99 and 96%, respectively. The structure was quadripartite that is typical in most plastids, and the structure and gene features were typically identical to those of higher plants. A total of 157 genes with an average size of 588.2 bp were annotated including 103 protein-coding genes (PCGs) with an average size of 780.9 bp, 46 tRNA genes, and eight rRNA genes with an average size of 220.7 bp. An overall GC content was 38.05%.

The results from the phylogenetic analysis performed using chloroplast coding sequences of the Alstroemeria hybrid ‘Hanhera’ and three and four published species in the Alstroemeriaceae and Liliaceae family, respectively, by a maximum likelihood method in MEGA version 6.0 (Tamura et al. 2013) revealed that the Alstroemeria hybrid ‘Hanhera’ was expectedly grouped with A. aurea followed by Bomarea edulis and Luzuriaga radicans which all belonged to the Alstroemeriaeae family (Figure 1).

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 MK327552.

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Figure 1. Maximum likelihood phylogenetic tree of the Alstroemeria hybrid ‘Hanhera’ with seven species belonging to the Alstroemeriaceae or Liliaceae based on chloroplast protein-coding sequences. Numbers in the nodes are the bootstrap values from 1000 replicates.