Two complete chloroplast genome sequences of genus *Paulownia* (Paulowniaceae): *Paulownia coreana* and *P. tomentosa*

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**ABSTRACT**

The nucleotide sequence of the two chloroplast (cp) genomes from *Paulownia coreana* and *P. tomentosa* are the first to be completed in genus *Paulownia* of family Paulowniaceae. The structure of two Paulownia cp genomes shows similar characteristic with general cp genome of angiosperms. The lengths of two cp genomes are 154,545 bp and 154,540 bp, respectively. The cp genomes are divided into LSC region (85,241 bp and 85,236 bp) and SSC region (17,736 bp and 17,736 bp) by two IR regions (25,784 bp and 25,784 bp). Both of two cp genomes contain 113 genes (79 protein coding genes, 30 tRNA genes and 4 rRNA genes), eight protein-coding genes, seven tRNA genes and four rRNA genes duplicated in the IR regions. Similar to the general cp genome of angiosperms, 18 of the genes in the two cp genomes have one or two introns. The overall A-T contents of two genomes are 62.0% which is similar with general angiosperms. A-T content in the non-coding (64.6%) is higher than in the coding (60.1%) regions. Seventy-one and seventy simple sequence repeat (SSR) loci were identified in the *P. coreana* and *P. tomentosa* cp genomes, respectively. In phylogenetic analysis, genus *Paulownia* shows closed relationship with *Lindenbergia philippensis* of Orobanchaceae.

Genus *Paulownia* which included eight species is one of the four genera in the family Paulowniaceae (Nakai 1949; Olmstead et al. 2001; APG 2016). *Paulownia* is fast-growing plant which is used for ornamental tree, the materials of instruments and contributions in agroforestry system. We sequenced and analyzed chloroplast (cp) genomes of *Paulownia coreana* Uyeki and *P. tomentosa* Steud. *P. coreana* is controversial and unresolved species that there is no significant morphological difference which compared with *P. tomentosa*.

The plant materials of *P. coreana* and *P. tomentosa* were collected from a single individual that planted in the Korea University. Voucher specimens (KUS 2014-1539, KUS2014-1540) and DNA samples (PDBK 2014-1539, PDBK 2014-1540) were deposited in the Korea University Herbarium and Plant DNA Bank in Korea (PDBK), respectively. Chloroplast genome sequences were analyzed using Illumina MiSeq (San Diego, CA), and assembled by Geneious 8.1.7 (http://www.geneious.com, Kearse et al. 2012). The complete cp genome sequences were submitted into NCBI database with accession numbers of KP718622 and KP718624, respectively.

Length of complete cp genome sequence of *P. coreana* and *P. tomentosa* are 154,545 bp and 154,540 bp, respectively. The cp genome of *P. coreana* is composed of 85,241 bp of LSC region, 17,736 bp of SSC region and 25,784 bp of two IR regions, whereas the cp genome of *P. tomentosa* is composed of 85,236 bp of LSC region, 17,736 bp of SSC region and 25,784 bp of two IR regions. Both of two cp genomes are consist of 113 individual genes which included 79 protein-coding genes, 30 transfer RNA genes and four ribosomal RNA genes. Among them, eight protein-coding genes, seven tRNA genes and four rRNA genes are duplicated on the IR regions. Similar to the general cp genome of angiosperms such as Panax and Sesamum, 18 of the genes in the each cp genome have one or two introns. Of these, rps12, clpP and ycf3 have two introns (Shinozaki et al. 1986; Kim & Lee 2004; Yi & Kim 2012).

The major portion of the *P. coreana* and *P. tomentosa* cp genomes consist of gene-coding regions (57.4% and 57.4%) which consist of protein-coding region (51.1% and 51.2%) and RNA regions (6.2% and 6.2%), whereas the intergenic spacers (including 23 introns) of both cp genomes comprise 42.6%. The overall A-T contents of two genomes are 62.0% which is similar with general angiosperms and other cp genomes of Lamiaceae and some genus of Orobanchaceae (Shinozaki et al. 1986; Kim & Lee 2004; Yi & Kim 2012; Wicke et al. 2013; Zhu et al. 2014; Welch et al. 2016). In both of two genomes, the A-T content in the non-coding (64.6%) is higher than in the coding (60.1%) regions. The A-T contents of the IR region is 52.8% in two cp genomes and the A-T contents of LSC and SSC regions are 64.0% and 67.6%, respectively. Seventy-one and 70 SSR loci which repeated more than ten times identified in the *P. coreana* and *P. tomentosa* cp genomes, respectively.

For the phylogenetic analysis, we assembled the 54 complete cp DNA sequences from the Lamiales clade and two
Figure 1. Chloroplast phylogenetic tree of Lamiales. A maximum likelihood tree (−lnL = 458425.7176) inferred from analysis of alignment data containing 79 coding genes in 56 chloroplast genome sequences by use of the GTR + I + I model. The numbers above and below each node indicate the Bayesian support percentages and bootstrap value, respectively. Genbank accession numbers of used taxa are shown below, Ajuga reptans (NC 023102), Andrographis paniculata (NC 022451), Boulardia latissiquama (NC 025641), Cistanche deserticola (NC 021111), Cistanche phelypaea (NC 025642), Coffea arabica (NC 008535), Coffea canephora (NC 030053), Conopholis americana (NC 023131), Dorcoreras hygrometricum (NC 016468), Epifagus virginiana (NC 001568), Genlisea margareae (NC 025622), Haplostachys haplostachya (NC 029819), Hesperelaea palmeri (NC 025787), Jasminum nudiflorum (NC 029370), Lindenbergia philippensis (NC 022859), Olea europaea subsp. cuspidate (NC 015604), Olea europaea subsp. europea (NC 015401), Olea europaea subsp. maroccana (NC 015623), Olea woodiana subsp. woodiana (NC 015608), Orobanche californica (NC 025651), Orobanche crenata (NC 024845), Orobanche gracilis (NC 023461), Paulownia coreana (KP 718622), Paulownia tomentosa (KP 718624), Pedicularis ishidoyana (NC 029700), Phelipanche purpurea (NC 023132), Phelipanche ramose (NC 023465), Phyllostegia velutina (NC 029820), Pinguicula ehlersiae (NC 023463), Plantago maritima (NC 028519), Plantago media (NC 028520), Pogostemon stellate (KP 718620), Pogostemon yatateanus (KP 718618), Premna microphylla (NC 026291), Rosmarinus officinalis (NC 027259), Salvia miltiorrhiza (NC 020431), Schwalbea Americana (NC 023115), Scrophularia takesimensis (NC 026202), Scutellaria baicalensis (NC 027262), Scutellaria insignis (NC 028533), Sesamum indicum (NC 016433), Stachys byzantina (NC 029822), Stachys coccinea (NC 029823), Stachys sylvatica (NC 029824), Stenogyne bifida (NC 029818), Stenogyne haliakalae (NC 029817), Stenogyne kanehoana (NC 029821), Tanaecium tetragonolobum (NC 027955), Tectona grandis (NC 020098), Utricularia gibba (NC 021449), Utricularia macrorhiza (NC 025653), and Utricularia reniformis (NC 029719).
outgroup sequences from Rubiaceae in Gentianales. A total of 79 protein CDSs including rrn genes were aligned for the 56 analyzed taxa. The aligned data matrix consists of a total of 85,408 bp. An ML tree was obtained with an $-\ln L = 458,425.7176$ using the GTR + G + I base substitution model (Figure 1). Similar to APG system, genus Paulownia forms a monophyletic group which shows closed relationship with Orobanchaceae (Olmstead et al. 2001, 2009; Bremer et al. 2002; APG 2016).

Disclosure statement
The authors report no conflicts of interest. The authors alone are responsible for the content and writing of the paper.

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