

Chloroplast genome of white wild chrysanthemum, *Dendranthema* sp. K247003, as genetic barcode

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Dendranthema boreale and *D. indicum* are easily distinguished from other Korean *Dendranthema* spp. by having yellow flowers. We have found a putative new taxon of *Dendranthema* having white flowers, except for sharing most characters with *Dendranthema boreale*. The chloroplast (cp) genome of the putative new taxon of *Dendranthema*, *Dendranthema* sp. K247003, registered in National Agro-Biodiversity Center (ABC), was completely characterized as a genetic barcode. The cp-genome of *Dendranthema* sp. K247003 was 151,175-bp in size: LSC was 82,886-bp, IR 24,971-bp, SSC 18,347-bp. The cp-genome of *Dendranthema* sp. K247003 contains 113 genes and 21 introns consisted of 79 protein coding genes, 4 RNA genes, and 30 tRNA genes, with 20 group II introns and one group I intron. Some of the genes and there introns were duplicated in IR. The cp-DNA of *Dendranthema* sp. K247003 is distinguished from that of *D. boreale* IT121002 by 67 SNPs in genic regions of 24 protein coding genes and by a 9-bp INDEL in *ycf1*. Further cp-DNA study will give us better information on genetic markers of *Dendranthema* species.

Keywords: Asteraceae, chloroplast genome, Compositae, *Dendranthema*, INDEL, SNP

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INTRODUCTION

Dendranthema, commonly called as chrysanthemum, one of the most popular and economically important ornamental plants due to its huge diversity in growing habits and wide range of colors. Five native *Dendranthema* species are reported from Korea (Park, 2007). Three species, *D. zawadskii*, *D. sichotense* and *D. coreanum*, are known to have ray flowers from white, pale pink to pale violet, while two species, *D. boreale* and *D. indicum* have yellow ray flowers. *Dendranthema boreale* is characterized by having smaller floral heads, 1.2~2 cm in diameter, and globose involucre. During the last several years, we have collected wild *D. boreale* individuals from Korean peninsula and Jeju Island. Their chromosome numbers varied among the populations, i.e. $2n = 18$, $18 + 2$ (Hwang *et al.*, 2014). During the field work, we have

found individuals with white floral heads in the middle of yellow headed *D. boreale* population. These white flowered individuals shared other characteristics with the rest of the yellow-flowered population. We registered two white individuals of *D. boreale* (K247002 and K247003) from different localities to National Agro-Biodiversity Center, Rural Agricultural Administration (RDA), as genetic resources for vegetative clones, because chrysanthemum can easily be propagated by cutting.

As the National Agricultural Genome Center Project, Rural Agricultural Administration (RDA), the genome of *D. boreale* IT121002 ($2n = 18$) has been characterized as a reference genome for chrysanthemum molecular breeding, Marker Assisted Selection (MAS). Organellar genomes having maternal inheritance have been characterized in more than 16 plants of *Dendranthema* including all the five Korean species. Currently, chloroplast genomes from

more than six Asteraceae genera were reported (Dempe-wolf *et al.*, 2010; Doorduyn *et al.*, 2011; Nie *et al.*, 2012; Liu *et al.*, 2013; Walker *et al.*, 2014). Among them, 47 cp-genomes were reported only from agriculturally important *Helianthus* (Shaw *et al.*, 2007; Timme *et al.*, 2007; Bock *et al.*, 2014), Asteraceae. For *Dendranthema*, while published as *Chrysanthemum*, some IGS regions of chloroplast were used for genetic diversity study (Liu *et al.*, 2012). For MAS, genetic information on genic regions is valued than that of IGS regions. Thanks to the dramatic development of Next Generation Sequencing method (NGS) in recent years, it has become possible to complete chloroplast genome sequencing at low cost. Complete cp-genomic sequences have become more useful as genetic barcode of plants (Nock *et al.*, 2011; Li *et al.*, 2015). Here, we report the complete genome of *Dendranthema* sp. K247003 as genetic barcode.

MATERIALS AND METHODS

Chloroplast DNA extraction, genome sequencing, assembly, and PCR-based validation

White flowered chrysanthemum, *Dendranthema* sp. K247003, was collected at the population of *D. boreale* in Geounri, Yeongwol of Gangwon province (N: 37°15' 18.9" E: 128°31'39.3"). The plant was registered at Agro-Biodiversity Center (ABC), Rural Agricultural Administration (RDA), as genetic resources for vegetative clones (IT number: K247003). The plant was propagated in Floriculture Research Division, National Institute of Horticultural and Herbal Science (NIHHS), RDA. Fresh leaves of *Dendranthema* sp. K247003 were collected from the Floriculture Research Institute, NIHHS in Rural Development Administration (RDA), Jeonju, and stored in liquid nitrogen until usage. Total DNA was extracted using the Qiagen DNeasy Plant Mini Kit (Qiagen, Germany), and DNA concentration and quality were determined using a Scandrop Nano-volume spectrophotometer (Analytik Jena, Germany). High quality DNA (concentration = 300 ng/μL, A260/280 ratio = 1.8-2.0, and A260/230 ratio = 1.7) was used for PCR and sequencing.

For NGS data production, purified DNA was fragmented and used to construct short-insert libraries (insert size, 200-bp), according to the manufacturer's instructions (Illumina, USA). The short fragments were paired-end sequenced using an Illumina Hi-Seq 2500 sequencing system at NICEM of Seoul National University. NGS data (7.63 Gb of 82.97 M reads) were analyzed using CLC Genomic Workbench ver. 7.5.1 (Qiagen, Hilden, Germany), as described by Jeong *et al.* (2014). For Sanger sequencing, the whole cp-genome of *Dendranthema* sp. K247003 was PCR-amplified in ~1-2 kb fragments,

and cp-genome structure was verified using Long PCR, with ~5-28 kb fragments, as described by Lee and Manhart (2002a; 2002b). Only PCR products ranging from ~1-2 kb were sequenced using Bigdye (ver. 3.1) and ABI3730 at NICEM of Seoul National University. Assembled cp-sequences were verified using Sequencher ver. 5.0 (Gencode, USA) by combining Sanger data and the assembled NGS sequence.

Genome annotation, genome comparison and sequence analysis

Protein coding and ribosomal RNA genes were annotated using DOGMA (<http://dogma.cccb.utexas.edu/>; Wyman *et al.*, 2004). The boundaries of each annotated gene were manually determined by comparison with orthologous genes from other known cp-genomes. Genes encoding tRNAs were first predicted using tRNAscan (<http://lowelab.ucsc.edu/tRNAscan-SE>; Lowe and Eddy, 1997) and ARAGORN, version 1.2 (<http://130.235.46.10/ARAGORN/>; Laslett and Canback, 2004), and were manually verified by predicting the tRNA secondary structure. Circular genome maps were drawn using GenomeVx (Conant and Wolfe, 2008), followed by manual modification. The sequencing data and gene annotation were submitted to National Agricultural Biotechnology Information Center (NABIC), Jeonju, with accession number NG-0482-000001. The mVISTA program in ShuffleLAGAN mode (Fraser *et al.*, 2004) was used to compare the cp-genome of *Dendranthema* sp. K247003 with that of *D. boreale* IT121002 (NABIC: NG-0478-000001; unpublished).

RESULTS AND DISCUSSION

The cp-genome of *Dendranthema* sp. K247003, was determined (Fig. 1) and found to be 151,175 bp in length. It includes small and large single copy (SSC, LSC) regions of 18,347 bp and 82,886 bp, respectively, separated by a pair of 24,971 bp Inverted Repeats (IRs). A total of 113 genes were detected, including 79 protein coding genes, 30 tRNA genes, and four rRNA genes (Table 1). This cp-genome was also found to contain 20 different introns, including 19 group II introns and a group I intron with a cyanobacterial origin (Besendahl *et al.*, 2000) found within the *trnL_uua* gene. Three protein coding genes, including *clpP*, *rps12*, and *ycf3*, contain two group II introns (*clpP.i1*, *clpP.i2*, *rps12.i1*, *rps12.i2*, *ycf3.i1* and *ycf3.i2*), and 14 genes contain a single group II intron: *rpoC1.i*, *rpl2.i*, *rpl16.i*, *rps16.i*, *atpF.i*, *petB.i*, *petD.i*, *ndhA.i*, *ndhB.i*, *trnA_ugc.i*, *trnG_ucc.i*, *trnL_gau.i*, *trnK_uuu.i*, and *trnV_uac.i*. Among the 20 group II introns, the intron in *rps12*, between exons 1 and 2, is trans-splicing, while the other 19 group II introns are cis-splicing.

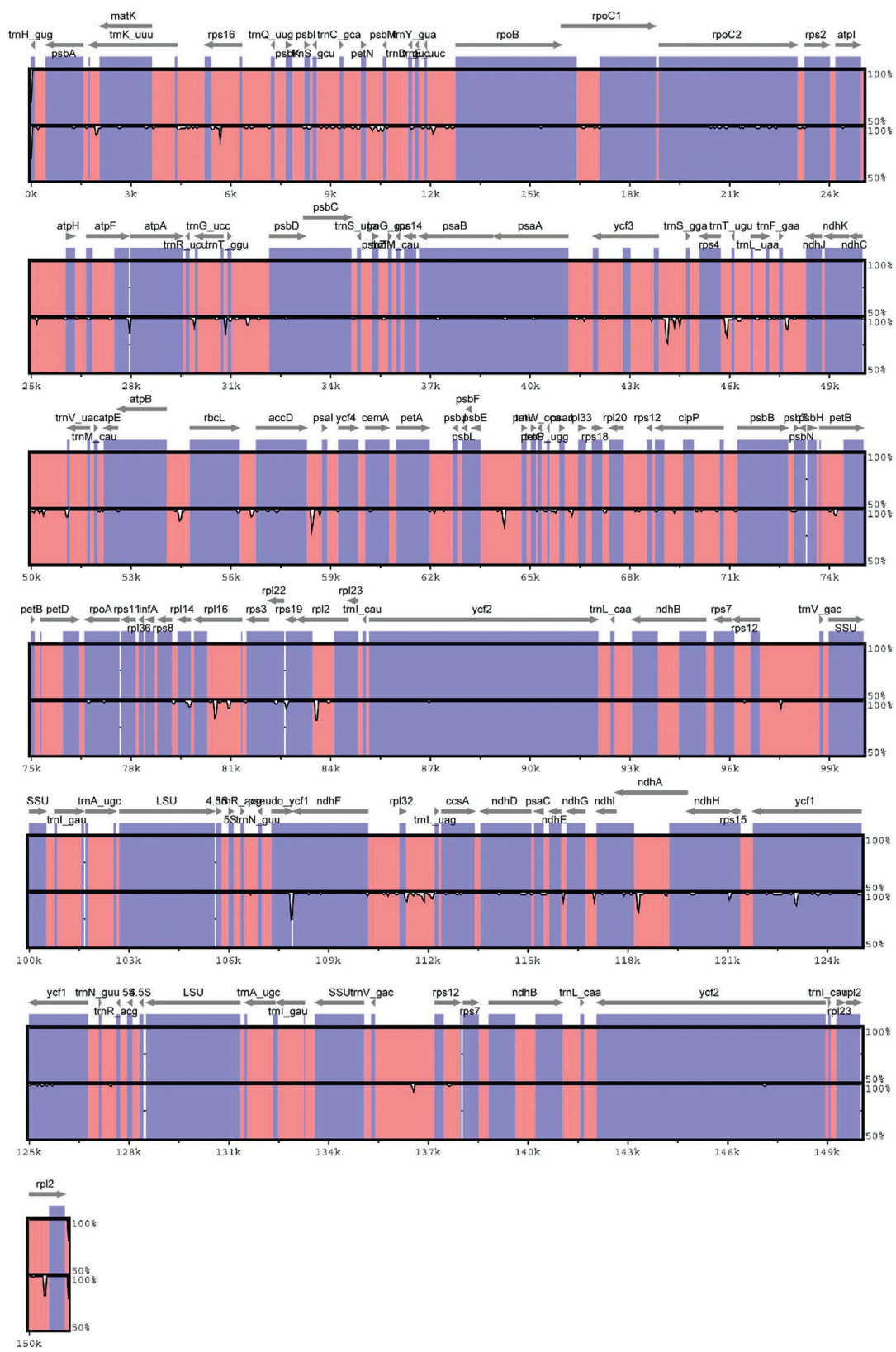


Fig. 2. Comparison of chloroplast genomes of *Dendranthema* sp. K247003 and *D. boreale* IT121002 using mVISTA program. Grey arrows and thick black lines above the alignment indicate genes with their orientation and the position of the IRs, respectively. The Y-scale represents the percent identity between 50-100%. Genome regions are color-coded: Coding regions in blue; noncoding sequences (CNS) in red.

Table 2. Variation in cp-genic regions between *Dendranthema* sp. K247003 and *D. boreale* IT121002. SNP: Single Nucleotide Polymorphism.

	Gene name	Size (<i>D. sp</i> / <i>D. boreale</i>)	No of SNP	INDEL		Gene name	Size (<i>D. sp</i> / <i>D. boreale</i>)	No of SNP	INDEL
1	<i>psbA</i>	1062	1	-	13	<i>accD</i>	1503	4	-
2	<i>matK</i>	1518	2	-	14	<i>cemA</i>	690	1	-
3	<i>rpoB</i>	3183	1	-	15	<i>rpoA</i>	1008	2	-
4	<i>rpoC2</i>	4152	10	-	16	<i>rpl22</i>	468	2	-
5	<i>atpI</i>	744	1	-	17	<i>ycf2</i>	6849	1	-
6	<i>atpF</i>	555	2	-	18	<i>ndhF</i>	2226	1	-
7	<i>psbD</i>	1062	1	-	19	<i>ccsA</i>	975	3	-
8	<i>psaB</i>	2205	1	-	20	<i>ndhD</i>	1503	3	-
9	<i>psaA</i>	2253	2	-	21	<i>ndhI</i>	561	1	-
10	<i>ndhJ</i>	477	1	-	22	<i>ndhA</i>	1092	3	-
11	<i>atpB</i>	1479	1	-	23	<i>ndhH</i>	1236	2	-
12	<i>rbcL</i>	1479	1	-	24	<i>ycf1</i>	5016 / 5007	19	9-bp INDEL

genes. As the genetic system genes, two conserved open reading frames (*ycf1* and *ycf2*), maturase K gene (*matK*), 3 RNA polymerase genes (*rpoA*, *rpoB* and *rpoC2*) and ribosomal protein large subunit gene (*rpl22*) are included. Among photosynthesis genes, six genes of NADH dehydrogenase (*ndhA*, *ndhD*, *ndhF*, *ndhG*, *ndhH*, *ndhI* and *ndhJ*), three genes of ATP synthase (*atpB*, *atpF* and *atpI*), two photosystem I genes (*psaA* and *psaB*), two photosystem II genes (*psbA* and *psbD*), a rubisco gene (*rbcL*), a membrane protein gene (*cemA*), cytochrome C biogenesis gene (*ccsA*), and a Acetyl-CoA carboxylase gene (*accD*) were variable. In addition to SNP, 9-bp INDEL was found in *ycf1* gene containing 19 SNPs.

CONCLUSION

This is the first report of chloroplast genome in *Dendranthema*, Asteraceae. As genetic barcode of *Dendranthema* sp. K247003, a possible new species, 151,175-bp of chloroplast genomic sequence was registered to NABIC (NG-0482-000001). The chloroplast genome is distinguished from that of *D. boreale* IT121002, by 67 SNPs and an INDEL in coding regions, in addition to 97 variable IGS sites. As suggested by Dong *et al.* (2015) in land plants, *ycf1* would be useful for plant identification as having 19 SNPs and an INDEL in the comparison of *Dendranthema* sp. K247003 and *D. boreale* IT121002. In addition, as suggested by Li *et al.* (2015), we show that chloroplast genomic information is useful for genetic barcode in *Dendranthema*. Further characterization of organellar genomes using NGS data would facilitate our phylogenomic study and molecular marker developments in *Dendranthema* at low cost. Finally, further morphological and cytological studies on *Dendranthema* sp. K247003 are remained for the taxonomic treatment of this taxon.

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