The complete plastome sequences of nine diploid potato clones

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\textbf{ABSTRACT}

Potato (\textit{Solanum tuberosum} L.) is the world’s fourth most important food crop and essential for global food security. The potato chloroplast genomes, the plastomes, are highly conserved and are largely studied for their maternal lineages. In this study, we assembled the complete circular plastome sequences of nine diploid potato clones, with sizes ranging between 155,296 bp and 155,564 bp. Annotation of these plastomes reveals that they each have 141 genes in a similar order. The computational chloroplast DNA typing reveals three plastid DNA types among the nine plastomes and they are grouped according to these types in the phylogeny.

The plastid DNA sequences (plastomes) of land plants are highly conserved compared to nuclear and mitochondrial DNA, and are extensively studied for their maternal phylogenetic relationships (Hosaka and Hanneman 1988). However, previous studies have shown considerable differences in plastomes and analysis of their complete sequences is important in understanding their diversity and evolutionary links (Hosaka 1986; Hosaka and Hanneman 1988; Chung et al. 2006; Achakkagari et al. 2020). In this study, a panel of nine diploid potato clones were selected for their unique characteristics such as tuber size, shape and disease resistance. Disease-free seed tubers from the following nine diploid \textit{Solanum tuberosum} clones were obtained from the Benton Ridge Substation of Agriculture and Agri-Food Canada in Benton, New Brunswick (https://www.agr.gc.ca/eng/scientific-collaboration-and-research-in-agriculture/agriculture-and-agri-food-research-centres-and-collections/atlantic-provinces/fred-ericton-research-and-development-centre/?id=1180622499704) (Latitude: 45.923397 Longitude: 66.606842): \textit{S. tuberosum} cv. W5281-2, \textit{S. tuberosum} cv. 08675-21, \textit{S. tuberosum} cv. 07506-01, \textit{S. tuberosum} cv. 12625-02, \textit{S. tuberosum} cv. 12120-03, \textit{S. tuberosum} cv. 11379-03, and \textit{S. tuberosum} cv. 10908-06. Potato plants were propagated in the greenhouse at the Fredericton Research and Development Centre, Fredericton NB, Canada, in six-inch clay pots with potting soil, under conditions of 16 h of light. Leaf tissue was ground and used for DNA extraction using the DNeasy Plant Mini Kit (Qiagen) according to the manufacturer’s instructions for plant tissue. Genomic DNA libraries were prepared using 10X Genomics GemCode technology (https://www.10xgenomics.com/) following the manufacturer’s instructions and whole genome de novo sequencing was carried out using the Novaseq 6000 DNA sequencer at the Génome Québec/Centre d’Expertise et de services Génome Québec (Montréal, Québec) at 89× coverage.

The raw reads obtained from the 10× Genomics GemCode technology were run through LongRanger Basic (https://www.10xgenomics.com/) to perform read trimming and barcode error correction. The filtered reads of each plant clone were used to assemble complete plastomes using the NOVOPlasty: de novo organellar genome assembler (Dierckxens et al. 2016), which can separate plastome from nuclear reads. A seed sequence of 1000 bp was randomly selected from a \textit{S. stenotomum} subsp. \textit{goniocalyx} Juz. & Bukasov plastome (MT120855, CIP 702472) (Achakkagari et al. 2020). The expected genome range was set to 145–165 kbp, and other parameters set to default. The assembled sequences were annotated using GeSeq (Tillich et al. 2017) and PGA (Qu et al. 2019) with \textit{S. stenotomum} subsp. \textit{goniocalyx} (MT120855), \textit{S. phureja} Juz. (MT120858, CIP 703654), \textit{S. curtibum} Juz. and Bukasov (MT120866, CIP 702937), and \textit{S. tuberosum} subsp. \textit{tuberosum} (L.) Hawkes (MT120865, CIP 705053) species as references (Achakkagari et al. 2020). The annotations were manually examined to adjust start and stop codons and to remove redundant annotations. The circular maps of these plastomes were drawn using OGDRAW v1.3.1 (Greiner et al. 2019). To understand their phylogenetic position, a maximum-likelihood phylogenetic tree was constructed using MEGA X v10.0.5 with Kimura 2-parameter substitution model (Kumar et al. 2018). The complete plastomes were used in constructing the phylogenetic tree. The chloroplast DNA types were identified by looking for specific polymorphisms that were previously identified in a panel of plastomes from a range of potato taxa (Achakkagari et al. 2020).
The assembled sizes range from 155,296 bp to 155,564 bp with the typical quadripartite structure of potato plastomes. All the nine of them have 25,593 bp of inverted repeats. The size of SSC and LSC ranges from 18,373 bp to 18,376 bp and 85,737 bp to 86,005 bp respectively. The plastomes of the W5281.2, 12625-02, 12120-03, and 11379-03 clones have the
same size of 155,492 bp. Similarly, the plastomes of clones H412-1, DW84-1457, 07506-01, and 08675-21 have the same size of 155,296 bp. Overall, 141 genes are present in each plastome, with similar gene content and gene order (Figure 1). Of these, 88 are protein-coding genes, 45 are tRNAs and 8 are rRNAs. There are five basic plastid DNA types in potato that were derived through point mutations (Hosaka 1986). The DNA typing reveals three types among the nine genomes. The W5281.2, 12625-02, 12120-03, and 11379-03 genomes have the S-type, while the H412-1, DW84-1457, 07506-01, and 08675-21 genomes have the T-type plastid DNA and the 10908-06 has the W-type. The complete plastomes of these nine potato clones and the four reference plastomes used for annotation are included in the phylogeny to determine their phylogenetic relationships. It was observed that all thirteen genomes are grouped together according to their plastid DNA type.

Future studies focusing on the agronomic traits associated with the plastome types in this study can greatly facilitate breeding programs in potato. Also, determining parental origin is trivial to deal with more and more fertility problems in diploid potato breeding. The phylogeny in this study is useful in understanding the evolutionary relationships, hybridization and introgression between these species, when combined with a nuclear phylogeny.

### 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/nuccore/](https://www.ncbi.nlm.nih.gov/nuccore/) under the accession numbers MT511702-MT511710. The associated BioProject, SRA, and Bio-Sample numbers are PRJNA684565, SRR13321685-SRR13321693, and SAMN17059154-SAMN17059162, respectively.

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### Disclosure statement

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

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