High-Throughput Development of Polymorphic Simple Sequence Repeat Markers Using Two Whole Genome Sequence Data in *Peucedanum japonicum*

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ABSTRACT Resource plants are important and have strong potential for a variety of utilities as crops or pharmaceutical materials. However, most resource plants remain wild and thus their utility for breeding and biotechnology is limited. Molecular markers are useful to initiate genetic study and molecular breeding for these understudied resource plants. We collected various wild collections of *Peucedanum japonicum* which is indigenous resource plants utilized as oriental medicine and leafy vegetables in Korea. In this study, we produced two independent whole genome sequences (WGSs) from two collections and identified large scale polymorphic simple sequence repeat (pSSR) based on our pipeline to develop SSR markers based on comparison of two WGSs. We identified a total of 452 candidate pSSR contigs. To confirm the accuracy and utility of pSSR, we designed ten SSR primer pairs and successfully applied those to seven collections of *P. japonicum*. The WGS and pSSR candidates identified in this study will be useful resource for genetic research and breeding purpose for the valuable resource plant, *P. japonicum*.

Keywords Molecular markers, Polymorphisms, Simple Sequence Repeats, Whole Genome Sequence, *Peucedanum japonicum*

INTRODUCTION

Indigenous resource plants have important values for a variety of utilities such as crops or pharmaceutical materials although there is limited research. Molecular markers are actively applied in many major crops, but the development of marker on the minor crops and resource plants is still limited. A molecular marker is a particular DNA fragment which includes specific genetic information with genome-level polymorphisms (Agarwal et al. 2008). Molecular markers have been widely used as valuable tools to measure genetic variation and thereby highly enhanced the genetic analysis on plants (Varshney et al. 2005).

Simple sequence repeats (SSRs, otherwise called as microsatellites) (Hiroe et al. 1958; Jones et al. 2010) which consist of sequential repeat of one to six or more nucleotide motifs in a head-to-tail structure (Kelkar et al. 2010) have been used often in genetic studies. SSR markers are further universally implemented for intraspecies variation and population genetic studies such as parentage analysis, fingerprinting, genetic structure analysis and genetic mapping (Mittal et al. 2009; Meng et al. 2014; Grover et al. 2016) due to variable polymorphic features and high...
reproducibility. Because of the utility and advantage of SSR markers, various programs have been developed to identify SSR motifs and primer design such as MISA (http://pgrc.ipk-gatersleben.de/misa/), SSR Locator, Primer3, and FullSSR (Rozen et al. 2000; da Maia et al. 2008; Abdelkrim et al. 2009; Untergasser et al. 2012; Metz et al. 2016). However, conventional SSR marker development and experimental verification still require long guided sequence information and a labor-intensive trial and error procedure by repetitive wet experiments to find polymorphic SSR markers (Ma et al. 2009).

*Peucedanum japonicum* is a perennial herb that belong to the Apiaceae family and grows on the cliff or between rocks in coastal regions of Taiwan, Japan, China and Korea (Buwalda 1949; Hiroe and Constance 1958; She 1992). The aerial part of *P. japonicum* such as young buds, leaves and berries are used as foodstuff whereas the underground roots have been used as oriental herbal medicine in the treatment of headache, anemia, apoplexy, paralysis, common cold and respiratory illnesses (Amano 1987; Hisamoto et al. 2003; Kang et al. 2013). *P. japonicum* has a variety of utilities. It is commonly used as a leafy vegetable and an herbal medicine, and a growing interest in this plant has increased farmer's cultivation in Korea. However, despite this growing interest, genomic information to advance breeding efforts on *P. japonicum* is limited (Seo et al. 2001; Han et al. 2017).

Next-generation sequencing (NGS) technology is widely used because of its high-throughput productivity (Varshney et al. 2009; Choi et al. 2011; Kim 2012). Recently, we developed a high-throughput method to assemble complete sequences of chloroplast genome and nuclear ribosomal DNA (nrDNA) simultaneously using low coverage WGS data, coined as *de novo* assembly using low coverage WGS (dnaLCW) (Kim et al. 2015b). We applied the dnaLCW method very efficiently to finding inter- or intra-species polymorphism and developing markers to differentiate species or cultivars based on the completely assembled chloroplast and nrDNA sequences in diverse plant species (Kim et al. 2015a). The dnaLCW method was successfully applied to develop markers for genetic diversity for resource plants. However, the information is limited to the chloroplast genome and nrDNA which are highly conserved genomic regions. Therefore, large-scale SSR markers derived from nuclear genomes are necessary for further genetics and genomics approach for breeding. Here, we would like to expand our approach to find large-scale polymorphic SSR markers using WGS data and provide useful genome information and genetic pSSR markers that can be utilized for genetics and breeding of the valuable resource plant, *P. japonicum*.

**MATERIALS AND METHODS**

**Plant materials, genomic DNA extraction and NGS sequencing**

We collected various wild *P. japonicum* collections in the Southern coastal area in Korea. We maintain these collections in Seoul National University farm, Suwon, in Korea. Among them, seven *P. japonicum* collections were used in this study. Three wild individuals were collected from Jeju Island (hereafter PJ #1), Geumo Island (hereafter PJ #2) and Wan Island (hereafter PJ#3) in Korea. The other two individuals (hereafter PJ#4 and PJ#5) were provided from Hantaek botanical garden (http://www.hantaek.co.kr, South Korea) and other two individuals (hereafter PJ#6 and PJ#7) were provided from Medicinal herb garden of college of Pharmacy in Seoul National University (http://www.snuherb.ac.kr). Hereafter we will refer all *P. japonicum* sample to their collection number. The fresh leaves of seven *P. japonicum* individuals were collected for DNA preparation. Each leaf sample was grinded separately using liquid nitrogen. The genomic DNAs were extracted using a modified cetyltrimethylammonium bromide (CTAB) method (Allen et al. 2006). Among seven plants, PJ#1 and PJ#2 were used to construct genomic libraries with insert size of 500 bp, according to Illumina paired-end (PE) standard protocol (http://www.illumina.com) and sequenced using Illumina MiSeq genome analyzer from LabGenomics (www.labgenomics.co.kr). The raw sequences of PJ#1 and PJ#2 have been uploaded at NABIC database and registered as NN-2130-000001 and NN-2132-000001, respectively (http://nabic.rda.go.kr/) (Seol et al. 2016).
Sequence preparation (quality control and paired reads joining), identification of SSR motif and clustering

WGS reads of PJ#1 and PJ#2 were trimmed off using Trimmomatic (ver. 0.33) based on quality score and sequence length; condition is set to the minimum quality score of $\geq 20$ and read length of $\geq 70$ bp (Bolger et al. 2014). Trimmed PE reads of PJ#1 were further assembled for joining of both forward and reverse PE sequences for each read by clc_overlap_reads (ver. 4.21.104315, CLC Inc, Aarhus, Denmark). We assembled forward and reverse sequences for each PE reads and made one PE-joined contigs (contigs) by combining of both forward and reverse reads based on the overlap. In the contig sets, contigs greater than 250 bp including SSR motifs were identified using microsatellite search module (MISA, http://pgrc.ipk-gatersleben.de/misa/) with the minimum repeat number of 6, 5, 5, 5, 5, and 4 for di-, tri-, tetra-, penta-, hexa-, and hepta-nucleotides, respectively. The contigs of reference species (PJ#1) were clustered based on homology using BLASTClust (https://www.ncbi.nlm.nih.gov/Web/Newsltr/Spring04/blastlab.html, Altschul et al. 1997) and the clusters with single contigs were selected for further analysis because we considered these sequence as derived from non-repeat genomic regions.

Discovery of pSSRs between two WGS reads

The WGS reads of PJ#2 were aligned with the contigs of PJ#1 in previous step by using clc_mapper (ver. 4.21.104315, CLC Inc, Aarhus, Denmark). Afterward, we have located the candidates of pSSR through comparison between contigs of PJ#1 and WGS reads of PJ#2 using clc_find_variation (ver. 4.21.104315, CLC Inc, Aarhus, Denmark). The information of the sequence variation file and the SSR motif file from MISA were combined to estimate the polymorphic regions including microsatellite sites using in-house python program.

PCR validation of SSR markers

Using sequences information with the candidate pSSR sites, SSR markers were designed by Primer3 (Rozen et al. 2000; Koressaar and Remm 2007; Untergasser et al. 2012). Genomic DNAs of seven P. japonicum collections were used as templates for PCR validation of SSR markers. PCR amplification was carried out in a 25-μL reaction volume containing the following components: 20 ng of DNA template, 10 μM of primer set, 5 mM of dNTP, and one unit Taq DNA polymerase (Vivagen, Seongnam, Korea). For six primer sets, PjSSR01 to PjSSR06, the amplification condition was as follows: 5 minutes at 94°C, 35 cycles of 94°C 20 seconds, 58°C 20 seconds, and 72°C 20 seconds and then 72°C for 7 minutes. For other primer sets, PjSSR07 to PjSSR10, the amplification condition was the same as above except annealing condition of 54°C for 20 seconds. PCR products were then separated by 9% polyacrylamide gel electrophoresis for identification of polymorphisms. Gel was stained with ethidium bromide and visualized under UV lamps for manual genotyping.

RESULTS

Identification of pSSR using two WGS in P. japonicum

We collected diverse germplasms for P. japonicum which is an indigenous plant in Korea. Among the

|                  | P. japonicum acc. #1 (PJ#1) | P. japonicum acc. #2 (PJ#2) |
|------------------|------------------------------|------------------------------|
| Raw data         | 3,550,678                    | 3,937,550                    |
| Trimming data    | 3,438,470                    | 3,817,637                    |
| PE joined contig (contigs) | 1,646,551                    | 595,380,248                  |
| SSR containing contigs | 25,814                      | 9,891,314                    |
| Primary filtered contigs by clustering | 12,206                       | 4,565,371                    |
| Secondary filtered contigs by mapping | 4,698                        | 1,878,413                    |
| pSSR containing contigs | 452                         | 179,609                      |
collections, we carried out WGS sequencing for two individuals, PJ#1 and PJ#2, which were collected from Jeju Island and Geumo Island, respectively. Raw WGS of PJ#1 and PJ#2 were approximately 1.1 Gbp and 1.2 Gbp, respectively. After sequence quality filtering, a total amount of 870 Mb and 980 Mb of WGS data remained for PJ#1 and PJ#2, respectively (Table 1). MiSeq sequences usually provide average 300 bp for forward and reverse sequence of each PE read. Our PE library was constructed from 500 bp insert size DNA fragments. To obtain relatively long reads and reduce the redundant finding from same sequences derived from forward and reverse sequence of same PE read, we assembled forward and reverse sequences for each PE reads to make PE-joined contigs (contigs) based on the overlapped sequences. The high quality 3.4 million reads were reduced to 1.6 M contigs by pair joining. Using the contigs, we conducted following steps for in silico identification of pSSRs. First, we selected SSR motif from PJ#1 through MISA program. Second, considerably non-redundant contigs were chosen as candidates by sequence clustering because high copy reads might be derived from repetitive sequence regions. SSR motifs were identified from 25,814 contigs, of which 12,206 were chosen from the solitary clustered contigs (Table 1). To identify pSSR motifs between PJ#1 and PJ#2, WGS reads of PJ#2 were aligned onto 12,206 contigs of PJ#1. During alignment, the 1,376 contigs with high mapping depth (≥10) and 6,132 contigs without any counterpart mapped reads were excluded. The remaining 4,698 contigs were then compared with SSR motif of PJ#2 and only 452 contigs contains pSSR motif were selected for further analysis. Among them, 371, 60, 21, and 7 contigs have di-, tri-, tetra- and more than penta-nucleotide SSR motifs, respectively (Fig. 1). Among the 452 pSSR contigs, five contigs had more than 2 SSR motifs (Supplementary Table S1).

Validation of pSSR and application for P. japonicum germplasm

To validate the pSSRs, we randomly chose ten contigs and designed primer pairs to amplify the SSR regions using Primer3 (Table 2). We included five more P. japonicum plants (PJ#3-7) for genotyping each collections using the primer pairs. All ten primer sets (PjSSR01 ~ 10) successfully amplified each of P. japonicum collections. Nine of ten produced the estimated polymorphic SSR

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**Fig. 1.** Classification of pSSR candidates based on the SSR motif in *P. japonicum*. The horizontal axis represents the type of SSR motif and vertical axis represents the number of SSR motifs.
Table 2. Primer information of ten SSR markers.

| Marker ID | SSR motif          | Contig Length (bp) | Estimated PCR product (bp) | Primers                          | Description based on BLASTN (E-value)          |
|-----------|--------------------|--------------------|----------------------------|----------------------------------|-----------------------------------------------|
| PjSSR01   | (GTAGATG)₄         | 543                | 203 196                    | F AGGTTGTGGGTTCAATTCCCA          |                                                |
|           | (GTAGATG)₃         |                    |                            | R GTCAGAAGTTGCGGTCAGA            |                                                |
| PjSSR02   | (TTG)₇ (TTG)₅      | 521                | 246 240                    | F GTGTTATTCACCTCACAAGGAAGG       |                                                |
|           |                    |                    |                            | R TCCCGTCCAAACGAACCTTA           |                                                |
| PjSSR03   | (TC)₁₇ (TC)₁₀      | 520                | 102 88                     | F TGCCGTTACACTGATTACCT           | XM_017381863.1 Daucus carota serine/threonine-protein kinase, mRNA (2e-14) |
|           |                    |                    |                            | R CTATGCTTTGCTGTCGAGCT           | XM_014765223.1 Glycine max receptor-like protein kinase 2-like, transcript variant X1, mRNA (5e-10) |
| PjSSR04   | (GAA)₈ (GAA)₅      | 519                | 213 204                    | F GAGAAATGACGGGAGAAATGG         | XM_017363412.1 Daucus carota cytochrome P450 8α1-like, mRNA (1e-16) |
|           |                    |                    |                            | R CTGCTCTGCAGTTGCGCTATT          |                                                |
| PjSSR05   | (AGAAG)₅ (AGAAG)₄ | 515                | 220 215                    | F TGTTGAACGACGGGAAGAGAGTG        |                                                |
|           |                    |                    |                            | R CTATGCTACATGACGGGTTT          |                                                |
| PjSSR06   | (GGGA)₅ (GGGA)₄    | 514                | 137 133                    | F AGATAGATGGCCACAGC             |                                                |
|           |                    |                    |                            | R AAACGCCCGCGCCGACCT            |                                                |
| PjSSR07   | (GTAA)₈ (GTAA)₇   | 509                | 245 241                    | F TGCTATTTGTCAGCTTTT           |                                                |
| PjSSR08   | (TAA)₆ (TAA)₈     | 541                | 257 263                    | F TGCCGTTACATGACAAGCAACT        | XM_010656117.2 Vitis vinifera uncharacterized, transcript variant X2, mRNA (4e-04) |
|           |                    |                    |                            | R TCAGACTGTTCAGGAAATAGA         |                                                |
| PjSSR09   | (TA)₁₁ (TA)₇      | 525                | 244 236                    | F ACACAATATAGATAGACACGGCTG      |                                                |
|           |                    |                    |                            | R CGGAGCTCTTCAGGAGCT           |                                                |
| PjSSR10   | (AGA)₅ (AGA)₃     | 517                | 290 284                    | F AGTGATGCGGAGAAGAAACAG         |                                                |
|           |                    |                    |                            | R TCAGCTGGAACCTTTGGAACCAAT      |                                                |
genotypes derived from the copy numbers of SSR units among *P. japonicum* collections (Fig. 2). Meanwhile, one primer, PjSSR08, did not show any polymorphism among genotypes which differ from the expectation (Table 3). The genotypes of the *P. japonicum* collections for each marker are represented in Table 3. The mean allele number of developed markers was around 2.7, whereas the mean genetic diversity and polymorphic information contents

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**Fig. 2.** *In silico* prediction and PCR validation of PjSSR01 and PjSSR06 primers. (A, C) Sequence alignment between contigs of PJ#1 and the counterpart WGS reads of PJ#2 for the primer regions, PjSSR01 and PjSSR06. Blue arrows indicate the forward and reverse primers and the colored triangle boxes in red and yellow represent the SSR motifs for PJ#1 and PJ#2, respectively. (B, D) Gel electrophoresis results of PCR products for PjSSR01 and PjSSR06. The number 1 to 7 indicate seven *P. japonicum* collections, PJ#1 to PJ#7, respectively. M indicates 100-bp ladder. Arrows with alphabets (a,b) indicate different alleles among seven collections.

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**Table 3.** Genotypes and allele diversity of 10 markers among seven collections of *P. japonicum.*

| Primers   | PJ#1 | PJ#2 | PJ#3 | PJ#4 | PJ#5 | PJ#6 | PJ#7 | No. of alleles | Genetic diversity\(^3\) | PIC\(^3\) |
|-----------|------|------|------|------|------|------|------|---------------|----------------|---------|
| PjSSR01 a | a    | b    | a    | a    | a    | a    | b    | 2             | 0.408          | 0.325   |
| PjSSR02 a | b    | b    | b    | b    | b    | b    | c    | 3             | 0.449          | 0.406   |
| PjSSR03 a | c    | c    | c    | c    | c    | c    | c    | 3             | 0.449          | 0.406   |
| PjSSR04 b | a    | d    | a    | a    | a    | c    | c    | 3             | 0.694          | 0.641   |
| PjSSR05 a | b    | a    | c    | c    | a    | b    | a    | 3             | 0.653          | 0.580   |
| PjSSR06 a | b    | b    | b    | b    | a    | a    | a    | 2             | 0.490          | 0.370   |
| PjSSR07 b | c    | d    | c    | c    | c    | c    | a    | 4             | 0.612          | 0.570   |
| PjSSR08 a | a    | a    | a    | a    | a    | a    | a    | 1             | 0.000          | 0.000   |
| PjSSR09 a | b    | b    | b    | b    | b    | b    | b    | 2             | 0.245          | 0.215   |
| PjSSR10 a | a    | b    | c    | b    | c    | d    | c    | 4             | 0.735          | 0.685   |
| Mean      |      |      |      |      |      |      |      | 2.7           | 0.473          | 0.420   |

\(^3\)Genetic diversity is the probability to show difference between two randomly chosen alleles from the population. 
\(^3\)PIC: polymorphism information content. Genetic diversity and PIC values were calculated based on genotype of individuals by PowerMarker ver. 3.0 (Liu *et al.* 2005).
(PIC) were around 0.47 and 0.42, respectively. We believe that the pSSR targets will be valuable resources and the ten SSR markers will be beneficial to evaluate and classify *P. japonicum* genotype for genetic study. We carried out blastn analysis (https://blast.ncbi.nlm.nih.gov/Blast.cgi) against non-redundant (nr) nucleotide sequence database and identified that four of ten *P. japonicum* primer regions were similar to genic regions of other plant species (Table 2).

**DISCUSSION**

Resource plants have a variety of useful potentials and some resource plants can be developed as minor food crops or pharmaceutical materials. Minor crops are crops with high value but not widely studied, hence it is also known as an ‘orphan crops’ or ‘underutilized crops’. In addition, genomic or genetic information of minor crops is rarely reported. SSR marker is one of the most applicable markers used the genetic evaluation and utilization of crops due to its high-reproducibility, hyper-variable and co-dominant nature (Park *et al.* 2009). Therefore, many efforts were conducted to develop SSR markers by construction and sequencing of SSR-rich genomic libraries for the minor crops and by utilization of EST sequence (Choi *et al.* 2011; Kim 2012; Izzah *et al.* 2014). However, previous researches required the time consuming wet experiments to find polymorphic SSR markers due to low polymorphism rate (with less than 5% of success rate) from the SSR candidates because they utilized sequence harboring the SSR motif from one genotype (Choi *et al.* 2011; Kim 2012; Izzah *et al.* 2014). In this study, we explored two WGSs to discover polymorphic SSR markers for the valuable resource plants *P. japonicum* as leafy vegetable and functional foods. Moreover, we have also confirmed that our designed SSR primer sets could be successfully applied in other collections. Ten SSR primers (PjSSR01~10) were successfully amplified, with high polymorphism in all collections (Table 2 and 3). Some primers showed different results from our expectation. The PCR products of both PjSSR04 and PjSSR10 were smaller than expected sizes and PjSSR08 did not show any polymorphism. These unexpected results might be caused by paralogous sequences in *P. japonicum*. Despite the unexpected results, however, our SSR markers displayed polymorphic amplicons with high success rate (9 out of 10). The 452 pSSR candidates sequence will provide fundamental information on evaluation of useful and fundamental genetic resources for breeding in *P. japonicum* (Table 2 and Supplementary Table S1).

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