Rapid Identification of Ginseng Cultivars (*Panax ginseng* Meyer) Using Novel SNP-Based Probes

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In order to develop a novel system for the discrimination of five ginseng cultivars (*Panax ginseng* Meyer), single nucleotide polymorphism (SNP) genotyping assays with real-time polymerase chain reaction were conducted. Nucleotide substitution in gDNA library clones of *P. ginseng* cv. Yunpoong was targeted for the SNP genotyping assay. From these SNP sites, a set of modified SNP specific fluorescence probes (PGP74, PGP110, and PGP130) and novel primer sets have been developed to distinguish among five ginseng cultivars. The combination of the SNP type of the five cultivars, Chungpoong, Yunpoong, Gopoong, Kumpoong, and Sunpoong, was identified as ‘ATA’, ‘GCC’, ‘GTA’, ‘GCA’, and ‘ACC’, respectively. This study represents the first report of the identification of ginseng cultivars by fluorescence probes. An SNP genotyping assay using fluorescence probes could prove useful for the identification of ginseng cultivars and ginseng seed management systems and guarantee the purity of ginseng seed.

**Keywords:** *Panax ginseng*, Nucleotide substitution, Fluorescence probes, Single nucleotide polymorphism genotyping

**INTRODUCTION**

Ginseng (*Panax ginseng* Meyer), which belongs to the genus *Panax* in the family *Araliaceae*, is a herbaceous perennial plant native to Korea and China; it has been used for over 2,000 years as a medicine in the Oriental countries. The major biologically active components of ginseng include ginsenosides, polysaccharides, peptides, polyacetylenic alcohols, phenolic compounds, and fatty acids [1]. *P. ginseng* evidences a variety of beneficial biological actions, including anti-stress [2], anti-carcinogenic [3,4], anti-hyperlipidemic [5], anti-aging, anti-amnestic [6], anti-diabetic [7], cardiovascular protection, and neuroprotection activities [8]. For these reasons, *P. ginseng* has become one of the most sought-after herbal remedies in Korea. The ginseng (*P. ginseng*) has three variants in Korea: the Jakyung, Chungkyung, and Hwangsook varieties.

Until now, a total of nine cultivars have been bred via pure line selection [9] and were registered in Korea Seed & Variety Service (Anyang, Korea). Among these nine cultivars, the Yunpoong [10], Gopoong [11], Sunpoong, Sunwon, Sunwoon, and Sunhyang cultivars were selected from Jakyung, whereas the Chunpoong, Chungsun and Kumpoong cultivars were selected from the Chungkyung and Hwangsook variants, respectively. Recently,
ginseng cultivars are frequently mixed-cultivated on local farms, and the seeds of ginseng cultivars are sold in the market by home seed producers. These illegal practices have caused serious social problems, including breeders’ intellectual property rights infringement, deteriorations in ginseng product quality and monetary damage to farmers caused by the use of mixed seeds.

Therefore, an effective management system should be established to maintain the high quality of ginseng cultivars. DNA marker techniques have been successfully applied for the classification of families and the estimation of genetic divergence among and between tested families. DNA-based markers are less profoundly affected by age, the physiological condition of samples, and environmental factors [12]. Thus, several researchers have developed molecular markers for the authentication of ginseng cultivars, including inter-simple sequence repeat markers [13], randomly amplified polymorphic DNA markers [14], restriction fragment length polymorphism markers [15], simple sequence repeat markers [16-18], single nucleotide polymorphism (SNP) markers [19,20], sequence characterized amplified region markers [21], and expressed sequence tag polymerase chain reaction (PCR) markers [22].

However, these markers failed to develop a robust molecular marker for the authentication of ginseng cultivars, owing to low levels of polymorphism and reproducibility. Additionally, several markers have proven effective only on a limited range of cultivars. In an effort to overcome these limitations, we developed sequence-tagged site (STS) markers containing a cleaved amplified polymorphic sequence (CAPS) system for the authentication of ginseng cultivars at the DNA level [23].

However, STS markers containing CAPS systems require complicated steps and electrophoresis and restriction fragment length polymorphism analysis after PCR amplifications are quite time-intensive.

These disadvantages can be complemented using a SNP genotyping assay with real-time PCR. Real-time PCR has advantages over competitive PCR, in that it is faster, more sensitive, and more robust against contamination due to minimal sample manipulation in closed-tube assays [24]. SNP genotyping assays are used routinely for the quantification of genetically modified organisms in food [25]. Moreover, a recently conducted genotyping assay by real-time PCR has also been employed for the identification of species such as beef [26], peanut [27], and fish [28-30].

The aim of the present study was to develop a novel method using fluorescence probes with a real-time PCR machine for the rapid identification of major ginseng cultivars, in order to protect intellectual property rights, to establish ginseng seed management systems and to guarantee the purity of ginseng seed.

**MATERIALS AND METHODS**

**Plant materials**

Fresh leaves of 3-years-old plants from five ginseng cultivars were cut and quickly frozen in a deep freezer prior to use. The five ginseng cultivars were used as follows; Chunpoong, Yunpoong, Gopoong, Kumpoong, and Sunpoong Sunpoong. These plant materials were preserved and cultivated at an experimental field of the National Institute of Horticultural and Herbal Science (NIHHS) of the Rural Development Administration, Chungbuk Province, Korea, and voucher samples were deposited at the Korean medicinal herbarium at NIHHS (Table 1).

**DNA extraction**

The samples were frozen in liquid nitrogen and immediately ground to a fine powder in a 1.5 mL micro-centrifuge tube with a micro-pestle. Total genomic DNAs were extracted from fresh leaves of the five cultivars of ginseng (3-year-old) using Dneasy Plant Mini Kit (Qiagen, Hilden, Germany) according to manufacturer’s protocol and DNA was eluted in 100 μL elution solution.

**Polymerase chain reaction analysis by sequence-tagged site markers**

Table 1. Ginseng cultivars used in this study

| No. | Cultivars               | Voucher no. | Collection area              |
|-----|-------------------------|-------------|------------------------------|
| 1   | Panax ginseng cv. Chunpoong | MPS002375   | Eumsung, Chungbuk, Korea     |
| 2   | P. ginseng cv. Yunpoong  | MPS002380   | Eumsung, Chungbuk, Korea     |
| 3   | P. ginseng cv. Gopoong  | MPS002385   | Eumsung, Chungbuk, Korea     |
| 4   | P. ginseng cv. Kumpoong | MPS002390   | Eumsung, Chungbuk, Korea     |
| 5   | P. ginseng cv. Sunpoong | MPS002395   | Eumsung, Chungbuk, Korea     |

**Table 1. Ginseng cultivars used in this study**
Gp130A) used in this study were developed from the gDNA library clones of *P. ginseng* cv. Yunpoong [23]. By using the methylation filtering technique, a genomic library was constructed, in which clone inserts were derived from the hypomethylated regions of ginseng genome. We collected more than 3000 white colonies from the methylation filtered library. Colony PCR was carried out on randomly selected 1099 from the methylation filtered library, to select clones between 0.8 and 1.5 kb as their estimated inserts size, and those clones were then subjected for sequence analysis. STS primer sets UFGp74, MFGp110A and MFGp130A were designed using Primer3 based on sequence information from gDNA library clones of *P. ginseng* cv. Yunpoong [31].

The STS primers synthesized by Bioneer (Daejeon, Korea) and the sequences of the STS primers are listed in Table 2. PCR amplification was performed using the following mixture: 50 ng of genomic DNA, 20 pmole of each primer, 200 uM dNTPs, 0.5 U DNA polymerase (5 U/L uL), 1X PCR buffer (Solgent, Daejeon, Korea); giving a 25 uL reaction mixture according to the manufacturer’s protocol. Amplification reactions were carried out on a Thermal-cycler machine (TProfessional 96; Biometra, Göttingen, Germany); the procedure used was an initial 5 min at 94°C followed by 40 cycles of 1 min at 94°C, 1 min at 60°C, 1 min at 72°C, and a final 7 min at 72°C. Amplification products were analyzed by electrophoresis on 1.5 % agarose gel in TBE buffer (45 mM Tris-HCl, pH 8.0, 45 mM Boric acid, 1 mM EDTA).

### Cloning and sequencing

The PCR products from five samples per cultivar were amplified and purified with a PCR purification kit (Qiagen) in accordance with the manufacturer’s instructions. Purified PCR products were cloned into the pGEM-T Easy vector System (Promega, Madison, WI, USA) and transformed into competent *Escherichia coli* KL1-Blue cells. The plasmids were purified with a plasmid mini kit (Qiagen). Sequencing was performed by Sanger’s method using ABI3730 automatic sequencer and the sequences were edited with the BioEdit program [32].

### Probes and primer design

The primers and fluorogenic probes were designed on the basis of the sequences of the PCR-amplified DNA fragments by STS markers (UFGp74, MFGp110A, and MFGp130A). The probes were labelled with the reporter molecules 5′-VIC™ (emission wavelength, 552 nm) or FAM™ (emission wavelength, 518 nm) and the 3′ends were labelled with an minor groove binder (MGB) molecule and a nonfluorescent quencher. The primers and fluorogenic probes were synthesized by Applied Biosystems (Foster city, CA, USA).

### Real-time polymerase chain reaction

Real-time PCR was performed by amplification using the ABI Step One Plus system (Applied Biosystems). Each reaction was carried out by duplicate in a Micro Amp Optical 96-well reaction plate. The 10 μL reaction mixtures contained 5 μL TaqMan® Universal Master Mix (which includes the heat-activated Ampli-Taq Gold Enzyme), 300 nM of each specific oligonucleotide primer, MGB probe 200 nM, and 1 μL of different dilution of

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**Table 2. STS primer sets discriminating ginseng cultivars**

| gDNA library | Primer name | PCR information          |
|--------------|-------------|--------------------------|
| ID           | Insert size (bp) | Gene bank no. | Primer sequence| PCR product size (bp) | Anneal temp (°C) |
| UFG0074      | 717         | HN339415                | UFGp74          | 920                 | 65             |
| MFG0110      | 1129        | HN339416                | MFGp110A        | 322                 | 65             |
| MFG0130      | 1124        | HN339417                | MFGp130A        | 930                 | 65             |

1. gDNA library clones of *Panax ginseng* cv. Yunpoong.
2. Primers were converted from gDNA library clones of *P. ginseng* cv. Yunpoong and sequence-tagged site (STS) primers were designed by Lee [23].
3. The polymerase chain reaction (PCR) product size by using the recommended annealing temperature (anneal temp) and the genomic DNA of *P. ginseng* cv. Yunpoong.
the DNA mixture, corresponding to 10 to 100 ng of total DNA. The reaction conditions were as follows: 60°C for 30 s, 95°C for 10 min followed by 40 cycles of 95°C for 15 s and 60°C for 1 min.

Genotyping

For the genotyping assay of ginseng cultivars, six TaqMan® MGB probes and three pair of oligonucleotide primers were employed. The one specific for the wild-type allele was labeled with a VIC™ dye (green fluorophore), and the other one specific for the mutant allele was labeled with a FAM™ dye (blue fluorophore). The generation of green fluorescence during amplification indicates homozygous wild-type, blue fluorescence indicates homozygous mutants, and both green and blue fluorescence indicates heterozygotes, respectively. For scatter plot analysis, the automatically generated threshold cycle values from each sample were plotted at coordinates that correspond to the signal of either FAM™ or VIC™.

RESULTS

Polymerase chain reaction analysis by sequence-tagged site markers

To identify useful variations for the development of a cultivar identification system, UFGp74, MFGp110A, and MFGp130A were tested to detect polymorphisms among ginseng cultivars (Korea). Polymorphic fragments were detected by MFGp130A ranging between 300 and 400 base pairs. MFGp130A generated two alleles among the ginseng cultivars. Chunpoong, Gopoong, and Kumpoong shared the same allele of 375 bp band and Yunpoong and Sunpoong shared another identical allele of the 322 bp band. UFGp74 and MFGp110A amplified uni-bands of 639 bp and 930 bp, respectively (Fig. 1).

Identification of single nucleotide polymorphisms based on sequences

According to the sequencing results, the amplified DNA with MFGp130A (gene bank accession number: HN339417) observed in one site base substitutions at 212 bp position and an insertion/deletion variation of 53 bp nucleotides at between 1042 bp and 1094 bp position (Fig. 2C). DNA with MFGp110A (gene bank accession number: HN339416) was observed in base substitutions of 6 sites. Chunpoong and Gopoong contain C at the 154 bp nucleotide position, but Yunpoong, Kumpoong, and Sunpoong were replaced with T in the same region. Yunpoong, Kumpoong, and Sunpoong contain C at nucleotide positions of 158 bp, 254 bp, and 630 bp, but Chunpoong and Gopoong were replaced with T at the same region. Chunpoong and Gopoong harbor A at the nucleotide position of 602 bp, but Yunpoong, Kumpoong, and Sunpoong were replaced with G at the same region. Yunpoong, Kumpoong, and Sunpoong harbor G at the nucleotide position of 859 bp, but Chunpoong and Gopoong were replaced with T at the same region. At the nucleotide positions 174 bp and 202 bp in the amplified DNA with UFGp74 (gene bank accession number: HN339415), Yunpoong, Gopoong, and Kumpoong harbor nucleotide G, but Chunpoong and Sunpoong contained C (Fig. 2A).
Probes and primers design

In the sequence alignment, nine SNP sites were detected. Based on the SNP site detected in the amplified region by UFGp74 (A), MFGp110A (B) and MFGp 130A (C), the primer and probe set were designed for identification of the ginseng cultivars (Korea). The wild-type probes were labeled on the 5' end with the fluorescent VIC™ dye, whereas the mutant type probes were labeled with the fluorescent FAM™ dye.
quencher and a MGB. To obtain a specific and reliable PCR amplification reaction, each fluorescent probe was placed to target SNP sequence between upstream and downstream primer (Fig. 2). The primers that flank each target were located close to the probe by means of forcing the Primer Express program. The primers and probes used in the present study are shown in Table 3.

Real-time polymerase chain reaction

The molecular authentication of major ginseng cultivars (Chunpoong, Yunpoong, Gopoong, Kumpoong, and Sunpoong) was conducted via an SNP genotyping assay with real-time PCR using cultivar-specific probe and primer sets. According to the results of real-time PCR using PGP74, Chunpoong and Sunpoong generated a FAM™

**Fig. 3.** Single nucleotide polymorphism (SNP) detection of five ginseng cultivars (Korea) by using PGP74. Each diagram shows two real-time polymerase chain reaction fluorescence curve with either SNP-A type or SNP-G type.

**Fig. 4.** Single nucleotide polymorphism (SNP) detection of five ginseng cultivars (Korea) by using PGP110. Each diagram shows two real-time polymerase chain reaction fluorescence curve with either SNP-T type or SNP-C type.
fluorescent (SNP-A type) signal, whereas Yunpoong, Gopoong, and Kumpoong generated a VIC\textsuperscript{TM} fluorescent (SNP-G type) signal (Fig. 3). In the case of real-time PCR using PGP110, Chunpoong and Gopoong generated a FAM\textsuperscript{TM} fluorescent (SNP-T type) signal, whereas Yunpoong, Kumpoong, and Sunpoong generated a VIC\textsuperscript{TM} fluorescent (SNP-C type) signal (Fig. 4). Finally, the results of real-time PCR using PGP130, Chunpoong, Gopoong, and Kumpoong generated a FAM\textsuperscript{TM} fluorescent (SNP-A type) signal, whereas Yunpoong and Sunpoong generated a VIC\textsuperscript{TM} fluorescent (SNP-C type) signal (Fig. 5).

**Genotyping**

The genotypes of ginseng cultivars were determined at the end of the PCR when the accumulated fluorescence was determined as predominantly a VIC\textsuperscript{TM} signal, a FAM\textsuperscript{TM} signal, or both VIC\textsuperscript{TM} and FAM\textsuperscript{TM} signals. The results were displayed on a scatter plot in which each axis corresponds to a reporter signal. Fig. 6 shows the results of genotyping of the structural variants of PGP74, PGP110, and PGP130, respectively. Two genotypes (SNP-A or SNP-G) were detected in PGP74. Chunpoong and Sunpoong shared the same SNP-A, and Yunpoong, Gopoong, and Kumpoong shared another identical SNP-G. In the case of PGP110, two genotypes (SNP-T or SNP-C) were detected in the five ginseng cultivars (Korea). Chunpoong and Gopoong shared the same SNP-T, whereas Yunpoong, Kumpoong, and Sunpoong shared another identical SNP-C. Finally, two genotypes (SNP-A or SNP-C) of PGP130 in the five cultivars of ginseng have been identified. Chunpoong, Gopoong, and Kumpoong shared the same SNP-A, whereas Yunpoong and Sunpoong shared another SNP allele, allele-C. The allele combination of the SNP type of the five cultivars, Chunpoong, Yunpoong, Gopoong, Kumpoong, and Sunpoong, was identified as ‘ATA’, ‘GCC’, ‘GTA’, ‘GCA’, and ‘ACC’, respectively (Table 4).

**DISCUSSION**

Ginseng (*P. ginseng*) is a Korea’s national heritage and Korea’s most representative agricultural crop. Its excellent quality has been recognized worldwide, and it has been exported to neighboring Oriental countries, including China. Despite the importance of ginseng, there are no any system and regulation to manage the protection of ginseng cultivars in Korea. In recent years, several efforts have focused principally on the development of various molecular markers to prevent the contamination of cultivars and to authenticate ginseng cultivars (Korea). Among them, the most recently developed STS system containing CAPS markers has proven the most useful in terms of its reproducibility and reliability [23].

STS containing CAPS markers are powerful tools for identification of the ginseng cultivars (Korea). However,
these methods require either restriction enzyme digestion or an agarose gel electrophoresis step. By way of contrast, the SNP genotyping assay using real-time PCR does not require additional post-PCR manipulations such as restriction enzyme digestion or agarose gel electrophoresis.

The authentication of ginseng cultivars (Korea) via SNP genotyping assays is based on allele-specific amplification via real-time PCR. The allelic discrimination of each probe depends on duplex allele-specific probes, labeled with a probe-specific fluorescent dye and a generic quencher that reduces fluorescence in the intact probes. During the amplification of the sequence surrounding the SNP, probes complementary to the DNA target are cleaved by the 5’ exonuclease activity of Taq polymerase [33].

The SNP genotyping assay used for identification of the ginseng cultivars (Korea) has several advantages. First, large numbers of samples can be processed within a short time. The conventional PCR method using gel electrophoresis takes approximately 5 h. But, the TaqMan-MGB probe system can analyze 96 samples in less than 2 h. Thus, the TaqMan-MGB probe system has a high-throughput capacity, because it requires no post-PCR manipulations. Second, this system can be used with a minimum amount of waste; because this system does not require electrophoresis, toxic substances, such as ethidium bromide and silver staining reagents and the discharge of plastic tubes is avoided. Third, conjugated MGB can increase the melting temperature of probes, thereby increasing probe specificity. Additionally, it allows for the use of shorter probes (usually 13 to 18 nucleotides), which can facilitate probe design, particularly in the AT-rich region. Finally, TaqMan-MGB probes can be employed without restriction enzymes recognizing specific regions. That is, almost all detected DNA sequence variation positions can be employed for probe design [34]. Thus, it is considered a powerful tool for use in SNP genotyping assays.

In this study, the real-time PCR using TaqMan-MGB probes was conducted for the application of SNP genotyping assays to discrimination among the ginseng cultivars (Korea). Consequently, fluorescence probes (PGP74, PGP110, and PGP130) and novel primer sets can be used to distinguish among ginseng cultivars (Chunpoong, Yunpoong, Gopoong, Kumpoong, and Sunpoong). The combination of the SNP type of the five cultivars, Chunpoong, Yunpoong, Gopoong, Kumpoong, and Sunpoong, was identified as ‘ATA’, ‘GCC’, ‘GTA’, ‘GCA’, and ‘ACC’, respectively (Table 4). However, to distinguish the nine cultivars used in the study by using only three pairs of probes would be too limited, as the SNP

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**Table 4. SNP combinations of ginseng cultivars by SNP genotyping**

| Probe | Cultivars/allele type |
|-------|-----------------------|
|       | Chunpoong | Yunpoong | Gopoong | Kumpoong | Sunpoong |
| PGP74 | A         | G        | G       | G        | A        |
| PGP110| T         | C        | T       | C        | C        |
| PGP130| A         | C        | A       | A        | C        |
| SNP combination | ATA  | GCC      | GTA     | GCA      | ACC      |

SNP, single nucleotide polymorphism.
variations are not sufficient enough. Accordingly, it will be required to build additional probes to distinguish five cultivars out of nine. By discovering additional SNP site, we will try to add the number of probes.

The development of a rapid authentication system of the ginseng cultivars (Korea) using TaqMan-MGB probes has proven helpful in protecting the unique national resources, in establishing a distribution system for Korean herbal markets, in the quality control of the ginseng seed production process, and in the protection of intellectual property.

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