Genetic Variation and Association Mapping in the F2 Population of the *Perilla* Crop (*Perilla Frutescens* L.) using new Developed *Perilla* SSR Markers

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Research Article

**Keywords:** *Perilla frutescens*, transcriptome, RNA-seq, SSR marker, F2 population, association analysis

**Posted Date:** April 5th, 2021

**DOI:** https://doi.org/10.21203/rs.3.rs-344394/v1

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Abstract

The transcriptome sequencing approach RNA-seq represents a powerful tool for transcriptional analysis and development of SSR markers for nonmodel crop. In the *Perilla* crop, analysis of the distribution of different repeat motifs showed that dinucleotide repeats were the most abundant (62.0%) type, followed by trinucleotide repeats (35.3%), with the two together comprising 97.3% of the eSSR repeats. In this study, we developed 39 new SSR primer sets by the transcriptome sequencing approach RNA-sEq. A total of 130 alleles were detected segregating in nine *Perilla* accessions with an average of 3.3 alleles per locus, ranging from 125 to 360 bp. The number of alleles per locus ranged from two to six. To detect SSR markers associated with morphological characteristics of *Perilla* crop, a total of 40 individuals from the *F*$_2$ population of *Perilla* were selected for association analysis based on their leaf- and plant-related characteristics. In an association analysis of 37 SSR markers and 9 leaf- and plant-related traits in the 40 individuals of the *F*$_2$ population, we identified 12 and 11 SSR markers associated with leaf-related traits and plant-related traits, respectively. Therefore, the new *Perilla* SSR primers described in this study could be helpful in identifying genetic diversity and genetic mapping, designating important genes/QTLs for *Perilla* crop breeding programs, and allowing *Perilla* breeders to improve leaf and plant quality through marker-assisted selection (MAS) breeding programs.

Introduction

*Perilla frutescens* Britt. is a self-fertilizing species of Labiatae. *Perilla* species is widely distributed and cultivated in East Asia, especially in South Korea, Japan, and China. There are two varieties (or two cultivated types) based on their uses and morphology: *P. frutescens* var. *frutescens* and var. *crispa*. Generally in East Asia, var. *frutescens* is used as an oil crop (dlggae in Korean, egoma in Japanese, and ren in Chinese), and var. *crispa* is used as a vegetable crop or herbal medicine crop (jaso in Korean, shiso in Japanese, and zisu in Chinese) (Lee and Ohnishi 2001, 2003; Lee et al. 2002; Nitta and Ohnishi 1999; Nitta et al. 2003). These two cultivated types of *Perilla* crop have a long history of cultivation and are very important as oil crop and vegetable or medicine crop in East Asia (Lee and Ohnishi 2001, 2003; Nitta et al. 2003). The two cultivated types of *Perilla* crop have different morphological characteristics, although var. *frutescens* and var. *crispa* can be crossed with each other by artificial pollination (Honda et al. 1990, 1994; Lim et al. 2019). For example, *Perilla frutescens* var. *frutescens* is taller, larger in seed size (above 2 mm), has either soft or hard seeds, has green leaves and stems, and has non-wrinkled leaves and a fragrance specific to the var. *frutescens*. Meanwhile, *Perilla frutescens* var. *crispa* is smaller in plant height and seed size (below 2 mm), has only hard seeds, has red or green coloration in the leaves and stem, and has wrinkly or non-wrinkly leaves and a fragrance specific to the var. *crispa* (Lee and Ohnishi 2001, 2003; Lee et al. 2002; Nitta et al. 2003).

Many taxonomic studies have been conducted to distinguish between the two cultivated types of *Perilla* crop using morphological characteristics and molecular markers. Their chromosome number is the same, 2n = 40 (Honda et al. 1994; Yamane 1950). A cross experiment between the two cultivated types by Nagai (1935), Honda et al. (1990, 1994) and Lim et al. (2019) showed that they are cross-fertile with each other by artificial pollination. Furthermore, studies by Koezuka et al. (1985, 1986), Honda et al. (1990), and Lee and Ohnishi (2001, 2003) failed to distinguish the two cultivated types of *Perilla* crop because of the existence of intermediate types. That is, essential oils, anthocyanins, and the color and hardness of the seeds cannot be used as key characteristics. It is still difficult to distinguish the two cultivated types of *Perilla* crop. In addition, weedy plants of *Perilla* crop were first reported in the two cultivated types of *P. frutescens* by Lee and Ohnishi (2001, 2003), Nitta and Ohnishi
(1999), and Nitta et al. (2003). They reported that analyses of random amplification of polymorphic DNA (RAPD) and amplified fragment length polymorphism (AFLP) markers showed that weedy plants could be grouped into two types: one belonging to the var. *frutescens* group and another belonging to the var. *crispa* group. However, the analysis does not provide concrete evidence of the origin of the weedy plants of the *Perilla* crop. Therefore, further research is needed to distinguish between cultivated and weedy types of *Perilla* crop.

Understanding the molecular genetic mechanism of phenotypic variation remains a great challenge because most complicated traits, such as yield, quality and resistance, are controlled by quantitative trait loci (QTLs) (i.e. quantitative and qualitative traits) (Mazzucato et al. 2008). In crop breeding studies, genetic mapping of functional loci for quantitative and qualitative related traits facilitates marker-assisted breeding. Two of the most commonly used tools for analyzing complex traits are linkage analysis and association analysis or association mapping. That is, linkage disequilibrium (LD)-based methods are used to detect associations between phenotypic variations and genetic polymorphisms. Thus, association mapping analysis complements traditional linkage analysis in understanding the genetic basis of complex traits and offers several advantages over linkage analysis, such as much higher mapping resolution, large number of alleles, wide reference population, and less research time in establishing an association (Buckler and Thornsberry 2002; Darvasi and Shifman 2005; Flint-Garcia et al. 2003; Mackay 2001). Furthermore, Shi et al. (2011) stated that using breeding populations may be a more practical approach for crop variety development, considering that markers linked to major QTLs can immediately be utilized in marker-assisted selection (MAS). Therefore, a thorough understanding of genetic diversity, population structure, and family relationships in a given panel is necessary for successful association analysis.

There are two methods to identify the genomic regions associated with important agricultural traits: 1) QTL mapping based on the results of crosses between two parents with linked and contrasting phenotypes and genotypes within segregated populations (Skot et al. 2005), and 2) association mapping using linkage disequilibrium (LD) between molecular markers and agronomic traits of interest (Pritchard et al. 2000, Zhang et al. 2012). Many population structures have been used for crop QTL detection and mapping. For example, backcross populations, F\textsubscript{2} populations, doubled haploid populations, recombinant inbred line populations, and near-isogenic line populations have all proven useful in identifying and confirming QTLs using various molecular marker systems (Choi et al. 2019; Farre et al. 2016; Park et al. 2013; Ramekar et al. 2018; Sa et al. 2015, Tanksley and Nelson 1996). However, for *Perilla* crop, QTL analysis using genetic maps of segregated populations is very difficult because various molecular markers, such as simple sequence repeat (SSR) markers for each chromosome, have not yet been developed for *Perilla* crop, in contrast with other major crops, such as rice, wheat and maize. In addition, more than 500 seeds of the F\textsubscript{2} population are produced in F\textsubscript{1} plant. For this reason, genetic analysis of the entire F\textsubscript{2} population is not possible (Lim et al. 2019). Therefore, in this study, some lines of the F\textsubscript{2} population were selected based on leaf characteristics such as leaf color (green or purple) and leaf shape (non-wrinkled or wrinkled) (see Fig. 1) to identify morphological differences between the two cultivated types of *Perilla* crop using molecular markers.

Meanwhile, molecular marker-based techniques in genetic research have been used in many crop species to identify the genomic regions associated with important agricultural traits. Among the various DNA molecular marker systems, SSRs are codominant markers, which are considered to be one of the most suitable means for assessing genetic diversity, genetic relationships, population structure, QTLs, and association mapping because of their reliability, reproducibility, and discrimination (Park et al. 2009; Pejic et al. 1998; Sa et al. 2018; Vathana et al. 2019). Thus, SSRs are widely used in many major crop species, such as rice (Pradhan et al. 2016), maize (Kim et
al. 2017), wheat (Crossa et al. 2007), and soybean (Qin et al. 2016), for assessing population structure and association analysis.

In *Perilla* crop, although SSR markers have recently been developed by several researchers, such as Kwon et al. (2005), Park et al. (2008), and Sa et al. (2018, 2019), the research is still insufficient. In our previous study, we sequenced and assembled one cultivated type (PF98095) of *P. frutescens* var. *frutescens* using transcriptome sequencing by RNA-seq (Tong et al. 2015), and we also obtained 15,991 SSR loci (Sa et al. 2018). The transcriptome sequencing approach RNA-seq represents a powerful tool for transcriptional analysis, novel gene discovery, and the development of SSR markers for nonmodel crops (Mutz et al. 2012, Wang et al. 2015, Yang et al. 2018). In particular, SSR markers developed from RNA-seq can enable marker-assisted selection for crop breeding programs because these regions are coding sequences and may be close to or within functionally transcribed genes (Mutz et al. 2012, Yang et al. 2018). In this study, we successfully developed SSR primers from *Perilla* crop, and these novel additional SSR markers can be used to analyze the genetic diversity, genetic relationships, population structure, QTL mapping, and association analysis in *Perilla* species.

Our study reports the results of the new *Perilla* SSR primers developed by the transcriptome sequencing approach RNA-seq and the results of association analysis in the F$_2$ population using these novel additional SSR primers and previously developed SSR primers. The results of this study are expected to provide useful information for future Perilla crop breeding programs.

**Materials And Methods**

**Plant materials and morphological characteristics of the F$_2$ population**

A total of 40 individuals from the F$_2$ population of the *Perilla* crop were selected from a cross between PF13-110 (female parent) and PF13-160 (male parent) (Fig. 1, Table 1). The female parent, PF13-110, is a var. *frutescens* that has non-wrinkled leaves with a green leaf surface and reverse side, green stem, and a fragrance specific to the var. *frutescens* (Fig. 1a, Table 1). The male parent, PF13-160, is a var. *crispa* that has wrinkled leaves with a purple leaf surface and reverse side, purple stem, and a fragrance specific to the var. *crispa* (Fig. 1b, Table 1).
Table 1
Morphological characteristics of the two parental lines used for crossing between *Perilla frutescens* var. *frutescens* (PF13-110, ♂, Parent A) and var. *crispa* (PF13-160, ♀, Parent B)

| Morphological trait          | Parent A | Parent A |
|------------------------------|----------|----------|
| QL1 (leaf size)              | Large    | Small    |
| QL2 (color of leaf surface)  | Green    | Purple   |
| QL3 (color of reverse side leaf) | Green | Purple   |
| QL4 (degree of pubescence)   | Heavily pubescent | Slightly pubescent |
| QL5 (number of leaf teeth)   | Many     | Few      |
| QL6 (flower color)           | White    | Purple   |
| QL7 (Stem color)             | Green    | Purple   |
| QL8 (leaf shape)             | Non-wrinkled | Wrinkled |
| QL9 (plant fragrance)        | Var. *frutescens* | Var. *crispa* |

To detect SSR markers associated with morphological characteristics, 40 individuals of the F\(_2\) population were selected based on leaf characteristics such as leaf color, leaf size, and leaf shape and plant characteristics such as pubescence degree, plant fragrance, and flower and stem colors (Tables 1, 2). Of the total individuals of the F\(_2\) population used in this study, approximately 20 individuals generally showed typical *frutescens* characteristics in leaf color and leaf shape, and the other 20 individuals showed a typical shape of *crispa* (Fig. 1, Table 2). To evaluate the morphological variation among the 40 individuals of the F\(_2\) population and two parental lines of *Perilla* crop, each line was grown in a greenhouse at the College of Agriculture and Life Sciences, Kangwon National University, Chuncheon, Gangwon-do, Korea. We examined nine leaf- and plant-related traits, as shown in Tables 1 and 2, that were selected based on a previous report by Lee and Ohnishi (2001).
Table 2
Morphological characteristics in two parental lines and 40 individuals of the F₂ population developed from cross between *Perilla frutescens* var. *frutescens* and var. *crispa*

| F₂ population line | QL1 | QL2 | QL3 | QL4 | QL5 | QL6 | QL7 | QL8 | QL9 |
|--------------------|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| 1                  | Large | Green | Green | S/P | Few | White | Green | None | Var. *frutescens* |
| 2                  | Small | Green | Green | S/P | Medium | Pink | Green | Wrinkle | Var. *crispa* |
| 3                  | Medium | Green | Green | S/P | Few | White | G/P | Wrinkle | Var. *frutescens* |
| 4                  | Small | Purple | Purple | Medium | Medium | Pink | Purple | Medium | Var. *frutescens* |
| 5                  | Large | Green | G/P | H/P | Medium | White | Green | Medium | Var. *crispa* |
| 6                  | Large | G/P | G/P | Medium | Medium | Pink | G/P | Medium | Var. *crispa* |
| 7                  | Large | G/P | G/P | Medium | Few | Pink | G/P | Medium | Var. *crispa* |
| 8                  | Medium | Green | Green | S/P | Medium | Pink | G/P | Wrinkle | Var. *crispa* |
| 9                  | Medium | Green | G/P | S/P | Medium | White | Purple | Medium | Var. *crispa* |
| 10                 | Small | Green | Green | S/P | Few | Pink | Green | None | Var. *frutescens* |
| 11                 | Large | Green | Green | S/P | Few | Purple | Purple | Medium | Var. *crispa* |
| 12                 | Medium | G/P | G/P | Medium | Few | Purple | Purple | None | Var. *crispa* |
| 13                 | Small | Green | Green | S/P | Medium | White | Purple | Wrinkle | Var. *crispa* |
| 14                 | Medium | Green | Green | Medium | Medium | White | Green | None | Var. *crispa* |
| 15                 | Medium | G/P | G/P | S/P | Few | Pink | G/P | None | Var. *frutescens* |
| 16                 | Small | Green | G/P | Medium | Few | White | Purple | None | Var. *frutescens* |
| 17                 | Small | G/P | G/P | Medium | Medium | Pink | Purple | None | Medium |
| 18                 | Medium | G/P | G/P | Medium | Medium | Purple | G/P | Medium | Var. *crispa* |
| 19                 | Small | Green | Green | H/P | Few | White | G/P | None | Var. *crispa* |
| 20                 | Medium | Purple | Purple | S/P | Medium | Pink | Purple | Medium | Var. *crispa* |
| 21                 | Medium | Green | Green | H/P | Many | Pink | Purple | Wrinkle | Var. *crispa* |
| 22                 | Medium | Green | G/P | S/P | Medium | White | Purple | Medium | Var. *frutescens* |

* Leaf color: G/P (Green/Purple)

** Pubescence: S/P (Slightly Pubescent), H/P (Heavily Pubescent)
| F₂ population line | QL1     | QL2     | QL3     | QL4   | QL5   | QL6   | QL7   | QL8   | QL9            |
|-------------------|---------|---------|---------|-------|-------|-------|-------|-------|----------------|
| 23                | Small   | Green   | Green   | H/P   | Many  | White | G/P   | Medium| Var. frutescens|
| 24                | Medium  | Green   | Green   | H/P   | Medium| White | G/P   | Wrinkle | None          |
| 25                | Small   | Green   | Green   | S/P   | Many  | White | Green | Medium| Var. frutescens|
| 26                | Large   | Purple  | G/P     | S/P   | Many  | Purple| G/P   | Medium| Var. crispa    |
| 27                | Large   | Green   | Green   | S/P   | Many  | Pink  | G/P   | Wrinkle| Var. crispa    |
| 28                | Large   | Green   | Green   | Medium| Many  | White | G/P   | Wrinkle| Var. crispa    |
| 29                | Medium  | Green   | Green   | S/P   | Many  | White | G/P   | Wrinkle| None          |
| 30                | Small   | Green   | G/P     | Medium| Many  | Pink  | G/P   | Wrinkle| None          |
| 31                | Large   | Green   | Green   | S/P   | Many  | White | G/P   | Wrinkle| Var. frutescens|
| 32                | Large   | Purple  | G/P     | H/P   | Medium| White | G/P   | Wrinkle| Var. crispa    |
| 33                | Medium  | Green   | Green   | H/P   | Medium| White | G/P   | None  | None          |
| 34                | Small   | Green   | Green   | H/P   | Many  | White | Green | Wrinkle| Var. crispa    |
| 35                | Small   | Purple  | Purple  | Medium| Many  | Pink  | Purple| Medium| None          |
| 36                | Small   | Green   | Green   | Medium| Many  | White | Green | Medium| None          |
| 37                | Large   | Purple  | Purple  | S/P   | Medium| Pink  | G/P   | None  | Var. crispa    |
| 38                | Large   | Green   | Green   | Medium| Medium| White | Green | Medium| Var. frutescens|
| 39                | Small   | Green   | G/P     | H/P   | Medium| Pink  | G/P   | Medium| Var. frutescens|
| 40                | Medium  | Green   | G/P     | S/P   | Many  | White | Purple| Wrinkle| Var. crispa    |

* Leaf color: G/P (Green/Purple)

** Pubescence: S/P (Slightly Pubescent), H/P (Heavily Pubescent)

** DNA extraction and SSR primer development **

Total DNA was extracted from young leaf tissues following the Plant DNAzol reagent protocols (GibcoBRL Inc., Grand Island, NY, USA). For construction of the transcriptome reference set in a previous study (Tong et al. 2015), *de novo* assembly of the PF98095 RNA-seq data was performed using Trinity software (http://TrinityRNASeq.sourceforge.net). The raw reads from next-generation sequencing (NGS) with a Phred quality score of at least 20 and a read length of at least 50 bp of HiSeq 2000 data were filtered before assembly. A Perl script MISA tool (http://pgrc.ipk-gatersleben.de/misa) was used to search microsatellite sites in the assembled transcriptome sequences of PF98095. The SSRs with dinucleotide, trinucleotide, and tetranucleotide...
repeat units were identified. Based on the SSR flanking sequences, PRIMER 3 software was employed to design the primer pairs. Then, we searched all unigenes in the cultivated type of var. frutescens (PF98095). In a previous study, Sa et al. (2018) detected a total of 15,991 SSR loci. All SSRs can be classified according to the number of repeating units: dinucleotide SSRs had the highest number of SSRs (9,910), followed by trinucleotide SSRs (5,652) and tetranucleotide SSRs (429). In this study, we designed 148 SSR primer sets based on the di-, tri-, and tetranucleotide types and the number of repeat units.

**SSR analysis and silver-staining**

SSR amplifications were conducted in a total volume of 20 µl consisting of 20 ng genomic DNA, 1× PCR buffer, 0.5 µM of forward and reverse primers, 0.2 mM dNTPs, and 1 unit of Taq polymerase (Biotools, Madrid, Spain). The PCR profile consisted of an initial denaturation at 95 ºC for 3 minutes followed by 36 cycles of 95 ºC for 30 seconds, 55 ºC for 30 seconds, and 72 ºC for 1 minute 30 seconds with a final extension step of 5 minutes at 72 ºC. After PCR, 5 µl of the final product was mixed with 10 µl of electrophoresis loading buffer (98% formamide, 0.02% BPH, 0.02% xylene C, and 5 mM NaOH). After denaturing and immediate cooling, 2 µl from each sample was loaded onto a 6% denaturing (7.5 M urea) acrylamide-bis acrylamide gel (19:1) in 1× Tris-borate-EDTA (TBE) buffer and then electrophoresed at 1,800 V and 60 W for 130 minutes. The separated fragments were then visualized using a silver staining kit (Promega, Madison, WI, USA).

**Data analysis**

TASSEL 3.0 (Bradbury et al. 2007) was used to evaluate marker-trait associations using a general linear model (GLM). The number of permutation runs was set to 10,000 to obtain a marker significance value of P ≤ 0.05. Power Marker version 3.25 (Liu and Muse 2005) was applied to obtain information on the number of alleles, major allele frequency (MAF), genetic diversity (GD), and polymorphic information content (PIC). GD was calculated for each pair of accessions using the Dice similarity index (Dice 1945). To examine the clustering pattern of all individuals of the F2 population by construction of a phylogenetic tree, Nei’s distance was calculated and used for the construction of unrooted phylogeny using a neighbor-joining (NJ) method, bootstrapping of the data (10,000 permutations) was implemented in POWERMARKER, and MEGA X was used to visualize the tree (Nei 1973, Kumar et al. 2018).

**Results**

**SSR identification and polymorphisms**

In our study, we surveyed the 148 newly developed SSR primer sets using nine Perilla accessions (3 cultivated var. frutescens; 3 weedy var. frutescens; 3 weedy var. crispa). The results showed that the 39 new SSR primer sets exhibited good amplification patterns and polymorphisms among the nine Perilla accessions (Supplement Tables 1, 2, Supplement Fig. 1). However, the remaining SSR primer pairs exhibited a monomorphic band (73) or ambiguous band pattern (24) and poor or no amplification (12). The 39 new SSR primer sets were used to measure polymorphisms in terms of the number of alleles, MAF, GD, and PIC among the nine Perilla accessions (Supplement Table 2). A total of 130 alleles were detected segregating in the nine Perilla accessions with an average of 3.3 alleles per locus, ranging from 125 to 360 bp. The number of alleles per locus ranged from two (KNUPE88, KNUPE90, KNUPE94, KNUPE100, KNUPE113, KNUPE114, KNUPE117, KNUPE118, KNUPE120, KNUPE122, KNUPE124, KNUPE125) to six (KNUPE105, KNUPE166). The GD ranged from 0.198 (KNUPE100, KNUPE113, KNUPE118, KNUPE124, KNUPE125) to 0.815 (KNUPE116), with an average of 0.552. The average PIC
value was 0.469, ranging from 0.178 (KNUPE100, KNUPE113, KNUPE118, KNUPE124, KNUPE125) to 0.788 (KNUPE116). The MAF per locus varied from 0.222 (KNUPE116) to 0.889 (KNUPE100, KNUPE113, KNUPE118, KNUPE124, KNUPE125), with an average of 0.590 (Supplement Table 2).

Meanwhile, we surveyed the 39 newly developed SSR primer sets between two parental lines of the F₂ population of the *Perilla* crop. From the results, among the 39 newly developed SSR primer sets, we selected 12 SSR primer sets that exhibited good amplification patterns and polymorphisms between the two parental lines of the F₂ population of the *Perilla* crop (Tables 3, 4). In addition, we surveyed a total of 105 SSR primer sets, which were developed in previous research, and selected 25 SSR primer sets that showed polymorphism between the two parental lines of the F₂ population (Table 4). Therefore, in our study, a total of 37 SSR primer sets, including 12 new SSR primer sets and 25 previous SSR primer sets, were used to measure polymorphisms in terms of GD, PIC, MAF, and separation patterns of allele bands (SPAB) among 40 individuals of the F₂ population of the *Perilla* crop (Table 4). In the results, the GD ranged from 0.185 (KNUPE95) to 0.708 (KNUPE102), with an average of 0.627. The average PIC value was 0.564, ranging from 0.177 (KNUPE95) to 0.650 (KNUPE102). The MAF per locus varied from 0.325 (KNUPE102) to 0.900 (KNUPE95), with an average of 0.481. In the case of SPAB, six SSR primer sets (KNUPF74, KNUPF79, KNUPF88, KNUPF89, KNUPF95, KNUPF103) were skewed towards AA (parent A), and four SSR primer sets (KNUPF5, KNUPF107, KNUPF108, KNUPF109) were skewed towards BB (parent B). Therefore, these SSR primer sets are considered to deviate from the expected Mendelian segregation ratio of 1:2:1 (or AA: AB: BB) in the F₂ population because more than 50% of the allele bands amplified in the F₂ population appeared to be biased towards one of the two parents (A or B). The remaining SSR primer sets showed a tendency towards a Mendelian segregation ratio of 1:2:1 (or AA: AB: BB) in 40 individuals of the F₂ population, although this does not represent the entire F₂ population. Meanwhile, among the SSR primer sets used in the analysis, 21 SSR primer sets showed a null band pattern in 40 individuals of the F₂ population (Table 4).
Table 3
Information of the 37 SSR primer sets used in the study

| SSR loci | Forward Primer(5'-3') | Reverse Primer(3'-5') | Repeat motif |
|----------|------------------------|-----------------------|--------------|
| KNUPF4   | TTTCAAAAATCTTACCAACGCT | TTCGTTTTTGCATCTAATTATTCA | (AT)$_{10}$ |
| KNUPF5   | TCCATCTCATCTCATTCAAACA | ATGGATCGGAAATCTAAACAA | (AT)$_{10}$ |
| KNUPF11  | TTTCAAAATCTTACCAACGCT | TGGAGGATTCACAATGTTGCT | (CA)$_{10}$ |
| KNUPF12  | AAATCCAATCTCGCCTCATATT | TTCTGAATCTTGAAGGTTGCT | (CA)$_{11}$ |
| KNUPF15  | CCACACGTAACCTCATAACC | TTATCTCAAAGAATCGGACA | (CT)$_{16}$ |
| KNUPF16  | CCTGTATCTTCCTCCTCGTAAAT | TGGATTTAATGCGTTGAGTTG | (CT)$_{22}$ |
| KNUPF21  | ATCTCATGAGTCTGGAGATTTC | AACCATGGCTATCCTACAGG | (TG)$_{10}$ |
| KNUPF23  | TTTCAAAATCTTACCAACGCT | CACTCCTTCCTCTCTTAAAT | (TG)$_{11}$ |
| KNUPF25  | GCTTAGTGTGAGGATAATCTAGGA | ACTCAGCATGCTTTGAATTCTC | (AAG)$_{12}$ |
| KNUPF29  | CTGGAAGTCTCCAGGAAAATG | GTCTAATCCGACGGAATCTG | (AGT)$_{10}$ |
| KNUPF30  | AACTGATATATATGGGCTGAAAAT | GACCTCTATCTCCACATCTATAA | (ATC)$_{10}$ |
| KNUPF39  | TCACCTTCCCTTCCATTAT | AGGATCGAACAGAACAATGT | (TCT)$_{13}$ |
| KNUPF49  | CTAGGTGTGCTGGGTATTTCAAT | AAACCGCTACCACCATTACC | (AG)$_{17}$ |
| KNUPF53  | GATTCATCATTTCAGCTCTCTCCC | ATGACCAATGGATTAAACAAGG | (CT)$_{17}$ |
| KNUPF54  | GCCATTGGAGATGGAATG | ATTCGGACAAAAAGCAACAAT | (GC)$_{5}$ |
| KNUPF59  | AATCTCGATGCCTAAACAGCT | TTTCTTGAATCCAGTTAAGG | (CAG)$_{7}$ |
| KNUPF60  | GCAATGGACATCTGTGAGAGTA | AATTGGTGTAATCATAGGGCAG | (CAG)$_{7}$ |
| KNUPF61  | GGGATACCCAAATTTCTACCAT | TCATGAAAAATCCAAAATTCA | (CAG)$_{7}$ |
| KNUPF72  | TAATTGAGGGATCCTCTCTCTCT | CGCCACCTTACTCTTACATC | (TCGA)$_{4}$ |
| KNUPF73  | CCATTCTTCAATTCGATCAACTA | TCTGCAAATCTCCAGTTAAAA | (CTAG)$_{4}$ |
| KNUPF74  | TTTCAAAATCTTACCAACGCT | GGTACACTCAACACTCTACAA | (AAAT)$_{6}$ |
| GBPFM111 | ATCATGGATGAATCGCACTT | CATTCTCCTCAATGGTACTCATTT | (ACACA)$_{8}$ |
| KWPE25   | ACGAAGGCGGCTTCAAATCTT | [[GT]$_{8}$] | |
| KNUPF79  | CCAACATCGACGACACTGATG | (AG)$_{26}$ |

* New SSR primer sets
| SSR loci  | Forward Primer(5’-3’)                      | Reverse Primer(3’-5’)          | Repeat motif |
|----------|------------------------------------------|-------------------------------|-------------|
| KNUPF81  | TTAAGCAACCAATTGCGAGGTA                   | GTTGTCAAAATTTGTTGATTT       | (AAT)$_7$   |
| KNUPF82  | AAACCAAGGAACCTGCACTCA                   | CGCTTCGTTTATTGTGTGA         | (AGA)$_7$   |
| *KNUPF88 | ATTCCTTTGAGTTGTTGGAGAA                  | CGTCTTTTGTGTGAGGCTAA        | (GTAC)$_4$  |
| *KNUPF89 | ATATCCCATTTCTGATGCAAG                | CTTTTTTCTGCTACTCATCA        | (AGT)$_6$   |
| *KNUPF95 | AGGATCAGGAGATCAGATGTGT                | GAATATTTAGATTTGTTATTTCCA   | (AT)$_9$    |
| *KNUPF102| AAGCTTATGCAACCTATGAGG                | ATGACTATCAGCAGATCTCGAA      | (AT)$_9$    |
| *KNUPF103| CAAAAGTACTTGTTCCTCCTC                | TGAAGCCCTCAAGCAACTATTA      | (CA)$_9$    |
| *KNUPF104| ATGCTTTTTGATTGTCTCTT                 | GCCAATCTTACATAATGTCCAAAA    | (TG)$_9$    |
| *KNUPF105| TCCAAAACCTGATTCTGAAC              | AATTTCGACTCATGGATCTTC       | (TG)$_9$    |
| *KNUPF107| TAAGGGTGCCGATTGTACTTTT            | ACAGAATGGCTCAAGAGATAG       | (AG)$_{25}$ |
| *KNUPF108| CCTCCTCTATAATACATGTTCTGC      | TCGAGCTTTCTGTATTTGAAA       | (AG)$_{22}$ |
| *KNUPF109| CTGAAAACCTTCTGAATGAGGG         | GCCCTATTTTGTAGCTGTTG       | (AG)$_{19}$ |
| *KNUPF111| CTAGGTGTGGGTGATTTCAAT               | AAATCCTACCACATTTCCC         | (AG)$_{17}$ |
| *KNUPF112| GACTTCTCCGAGCATATTCAC          | ATTGTCTTCTGGATAATGTGTG      | (CT)$_4$    |

* New SSR primer sets
Table 4
Estimates of GD, PIC, MAF, and separation pattern of allele bands (SPAB) of 37 SSR primer sets in 40 individuals of the F$_2$ population.

| Marker     | GD   | PIC  | MAF | Separation of the F2 population |
|------------|------|------|-----|---------------------------------|
|            |      |      |     | AA    | AB | BB | null |
| KNUPF4     | 0.666| 0.599| 0.425| 12  | 17 | 10 | 1    |
| KNUPF5     | 0.629| 0.565| 0.500| 2   | 13 | 20 | 5    |
| KNUPF11    | 0.636| 0.564| 0.475| 12  | 17 | 11 | 0    |
| KNUPF12    | 0.624| 0.553| 0.500| 9   | 20 | 11 | 0    |
| KNUPF15    | 0.694| 0.636| 0.400| 14  | 14 | 9  | 3    |
| KNUPF16    | 0.674| 0.622| 0.475| 5   | 20 | 11 | 4    |
| KNUPF21    | 0.645| 0.587| 0.500| 12  | 16 | 10 | 2    |
| KNUPF23    | 0.654| 0.580| 0.425| 11  | 17 | 11 | 1    |
| KNUPF25    | 0.661| 0.587| 0.375| 14  | 14 | 12 | 0    |
| KNUPF29    | 0.679| 0.618| 0.425| 10  | 17 | 11 | 2    |
| KNUPF30    | 0.634| 0.561| 0.475| 9   | 18 | 13 | 0    |
| KNUPF39    | 0.670| 0.610| 0.450| 9   | 19 | 11 | 1    |
| KNUPF49    | 0.596| 0.530| 0.550| 8   | 23 | 9  | 0    |
| KNUPF53    | 0.686| 0.625| 0.400| 11  | 16 | 12 | 1    |
| KNUPF54    | 0.629| 0.572| 0.525| 7   | 21 | 10 | 2    |
| KNUPF60    | 0.669| 0.608| 0.450| 11  | 18 | 10 | 0    |
| KNUPF61    | 0.651| 0.577| 0.425| 10  | 17 | 13 | 0    |
| KNUPF72    | 0.539| 0.481| 0.625| 8   | 25 | 7  | 0    |
| KNUPF73    | 0.646| 0.571| 0.425| 14  | 18 | 8  | 0    |
| KNUPF74    | 0.656| 0.596| 0.475| 21  | 9  | 8  | 2    |
| KNUPF79    | 0.601| 0.516| 0.450| 17  | 18 | 5  | 0    |
| KNUPF81    | 0.676| 0.613| 0.375| 15  | 15 | 8  | 2    |
| KNUPF82    | 0.690| 0.628| 0.350| 10  | 14 | 15 | 1    |
| KNUPF88    | 0.656| 0.587| 0.425| 17  | 14 | 8  | 1    |
| KNUPF89    | 0.681| 0.634| 0.475| 19  | 8  | 6  | 7    |
| KNUPF95    | 0.185| 0.177| 0.900| 36  | 2  | 2  | 0    |

*GD: Genetic Diversity, PIC: Polymorphism Information Content, MAF: Major Allele Frequency
In addition, a phylogenetic tree was constructed using a total of 37 selected SSR markers to elucidate the phylogenetic relationship between the 40 individuals of the F₂ population and their two parental lines (Fig. 2). The 40 individuals of the F₂ population and two parental lines were clustered into two major groups: Group I included 18 individuals of the F₂ population, including Parent B (PF13-160); Group II included 22 individuals of the F₂ population, including Parent A (PF13-110). In the results, with the exception of exceptional individuals, Group I mainly contained many individuals with similar characteristics to Parent B, and Group II included many individuals with similar characteristics to Parent A (Table 2, Fig. 2).

### Phenotypic variation and association analysis of leaf- and plant-related traits

Information on the distribution of leaf- and plant-related traits in 40 individuals of the F₂ population is shown in Table 5. In the case of QL1, among the 40 individuals of the F₂ population, 12 individuals showed large, 14 individuals showed medium, and 14 individuals showed small. For QL2, 28 individuals showed green, 6 individuals showed mixed green/purple, and 6 individuals showed purple. For QL3, 21 individuals showed green, 15 individuals showed mixed green/purple, and 4 individuals showed purple. For QL4, 9 individuals showed heavily pubescent, 13 individuals showed medium pubescent, and 18 individuals showed slightly pubescent. For QL5, 13 individuals showed many numbers, 18 individuals showed medium numbers, and 9 individuals showed few numbers. For QL6, 21 individuals were white, 15 individuals showed pink, and 4 individuals showed purple. For QL7, 9 individuals showed white, 19 individuals showed mixed green/purple, and 12 individuals showed purple. For QL8, 10 individuals showed non-wrinkles, 16 individuals showed medium wrinkles, and 14 individuals showed wrinkles. For QL9, 11 individuals showed specific to var. *frutescens*, 7 individuals showed medium, and 22 individuals showed specific to var. *crispa*. In addition, we performed a correlation analysis to detect the

| Marker      | GD    | PIC   | MAF   | Separation of the F₂ population |
|-------------|-------|-------|-------|---------------------------------|
|             |       |       |       | AA    | AB    | BB    | null  |
| KNUPF102    | 0.708 | 0.650 | 0.325 | 13    | 11    | 13    | 3     |
| KNUPF103    | 0.514 | 0.438 | 0.625 | 25    | 12    | 3     | 0     |
| KNUPF104    | 0.591 | 0.522 | 0.550 | 11    | 22    | 7     | 0     |
| KNUPF105    | 0.635 | 0.573 | 0.500 | 6     | 20    | 13    | 1     |
| KNUPF107    | 0.644 | 0.579 | 0.475 | 2     | 13    | 24    | 1     |
| KNUPF108    | 0.619 | 0.569 | 0.550 | 9     | 6     | 22    | 3     |
| KNUPF109    | 0.543 | 0.489 | 0.625 | 9     | 5     | 25    | 1     |
| KNUPF111    | 0.645 | 0.572 | 0.450 | 10    | 18    | 12    | 0     |
| KNUPF112    | 0.645 | 0.572 | 0.450 | 12    | 18    | 10    | 0     |
| GBPFM111    | 0.651 | 0.583 | 0.450 | 8     | 19    | 12    | 1     |
| KWPE25      | 0.586 | 0.506 | 0.525 | 14    | 20    | 6     | 0     |
| Mean        | 0.627 | 0.564 | 0.481 |       |       |       |       |

*GD: Genetic Diversity, PIC: Polymorphism Information Content, MAF: Major Allele Frequency*
relationship between leaf- and plant-related traits in the 40 individuals of the F\textsubscript{2} population (Table 6). The results showed significant variation among nine leaf- and plant-related traits (QL1, QL2, QL3, QL4, QL5, QL6, QL7, QL8, and QL9) in the 40 individuals of the F\textsubscript{2} population (Table 6). There were highly statistically significant positive correlations of QL2 with QL3 (0.682**) and QL6 (0.429**), of QL3 with QL6 (0.390*) and QL7 (0.462**), and of QL5 with QL8 (0.489**) and QL9 (0.313*) at P < 0.01 or P < 0.05.

Table 5
Distribution of morphological variation in 40 individuals of the F\textsubscript{2} population developed from cross between *Perilla frutescens* var. *frutescens* and var. *crispa*

| Morphological character      | Category                                      | F\textsubscript{2} population (N = 40) |
|------------------------------|-----------------------------------------------|---------------------------------------|
| QL1 (leaf size)              | Large (> 14cm\textsuperscript{2})-1, medium (= 14cm\textsuperscript{2})-2, small (< 14cm\textsuperscript{2})-3 | 1- (12)* 2- (14) 3- (14)             |
| QL2 (color of leaf surface)  | Green-1, green/purple-2, purple-3             | 1- (28) 2- (6) 3- (6)                 |
| QL3 (color of reverse side leaf) | Green-1, green/purple-2, purple-3             | 1- (21) 2- (15) 3- (4)                 |
| QL4 (degree of pubescence)   | Heavily pubescent-1, medium-2, slightly pubescent – 3 | 1- (9) 2- (13) 3- (18)               |
| QL5 (number of leaf teeth)   | Many (> 24 number)-1,medium-2, Few (< 17 number)-3 | 1- (13) 2- (18) 3- (9)               |
| QL6 (flower color)           | White-1, pink-2, purple-3                     | 1- (21) 2- (15) 3- (4)               |
| QL7 (stem color)             | Green-1, green/purple-2, purple-3             | 1- (9) 2- (19) 3- (12)               |
| QL8 (leaf wrinkle)           | Non-wrinkle-1, medium-2, wrinkle – 3          | 1- (10) 2- (16) 3- (14)               |
| QL9 (plant fragrance)        | Specific to var. *frutescens*-1, non-specific – 2, specific to var. *crispa*-3 | 1- (11) 2- (7) 3- (22)               |

* The number of individuals in the F\textsubscript{2} population for each trait type.
### Table 6
Pearson correlation coefficients among 9 qualitative traits in F<sub>2</sub> population

|     | QL2    | QL3    | QL4    | QL5    | QL6    | QL7    | QL8    | QL9    |
|-----|--------|--------|--------|--------|--------|--------|--------|--------|
| QL1 | -0.151 | -0.039 | -0.175 | -0.037 | -0.100 | 0.144  | -0.089 | 0.103  |
| QL2 | 0.682**| 0.047  | 0.126  | 0.429**| 0.269  | -0.165 | 0.191  |
| QL3 | 0.005  | -0.024 | 0.390* | 0.462**| -0.262 | 0.095  |
| QL4 |       | -0.114 | 0.182  | 0.129  | 0.045  | -0.243 |
| QL5 |       | -0.142 | -0.073 | 0.489**| 0.313* |
| QL6 |       |        | 0.244  | -0.161 | 0.130  |
| QL7 |       |        |        | -0.014 | 0.150  |
| QL8 |       |        |        |        |        |        | 0.167  |

** Significance at P < 0.01
* Significance at P < 0.05

Meanwhile, to select SSR markers associated with leaf- and plant-related characteristics in the 40 individuals of the F<sub>2</sub> population, genotypes of the 37 SSR markers and the nine leaf- and plant-related traits were used to confirm significant marker-trait associations (SMTAs) using TASSEL software (Table 7). In the results, we detected 24 SMTAs involving 21 SSR markers associated with nine leaf- and plant-related traits using GLM at a significance level of P ≤ 0.05 (Table 7). Among the 12 SSR markers for leaf-related traits, KWPF25 and KNUPF108 were associated with QL2, KNUPF11 was associated with QL3, KNUPF74 and KNUPF89 were associated with QL5, and seven SSR markers (KNUPF21, KNUPF49, KNUPF60, KNUPF81, KNUPF103, KNUPF109, GBPEM111) were associated with QL8. In the case of plant-related traits, 11 SSR markers were associated with plant-related traits. Among them, two SSR markers (KNUPF15 and KNUPF21) were associated with QL4, KNUPE11 was associated with QL6, four SSR markers (KNUPF11, KNUPF54, KNUPF82, and KNUPF88) were associated with QL7, and five SSR markers (KNUPF5, KNUPF29, KNUPF39, KNUPF61, and GBPEM111) were associated with QL9 (Table 7).

### Table 7. List of significant markers detected with the Q GLM model
| Trait                          | Marker   | $P$ value | Marker $R^2$ |
|-------------------------------|----------|-----------|--------------|
| **Leaf-related traits**       |          |           |              |
| QL2 (Color of leaf surface)   | KWPE25   | 0.02      | 0.19         |
|                               | KNUPF108 | 0.02      | 0.22         |
| QL3 (Color of reverse side leaf) | KNUPF11  | 0.02      | 0.20         |
| QL5 (Number of leaf teeth)    | KNUPF74  | 0.01      | 0.22         |
|                               | KNUPF89  | 0.01      | 0.26         |
| QL8 (Leaf wrinkle)            | KNUPF21  | 0.04      | 0.16         |
|                               | KNUPF49  | 0.05      | 0.15         |
|                               | KNUPF60  | 0.02      | 0.21         |
|                               | KNUPF81  | 0.03      | 0.18         |
|                               | KNUPF103 | 0.01      | 0.21         |
|                               | KNUPF109 | 0.04      | 0.17         |
|                               | GBPFM111 | 0.04      | 0.16         |
| **Plant-related traits**      |          |           |              |
| QL4 (Degree of pubescence)    | KNUPF15  | 0.03      | 0.19         |
|                               | KNUPF21  | 0.04      | 0.17         |
| QL6 (Flower color)            | KNUPF11  | 0.02      | 0.19         |
| QL7 (Stem color)              | KNUPF11  | 0.03      | 0.20         |
|                               | KNUPF54  | 0.04      | 0.17         |
|                               | KNUPF82  | 0.01      | 0.17         |
|                               | KNUPF88  | 0.02      | 0.21         |
| QL9 (Plant fragrance)         | KNUPF5   | 0.03      | 0.20         |
|                               | KNUPF29  | 0.03      | 0.19         |
|                               | KNUPF39  | 0.04      | 0.17         |
|                               | KNUPF61  | 0.00      | 0.33         |
|                               | GBPFM111 | 0.00      | 0.32         |

**Discussion**

Genetic variation between individuals within a population or between populations of crop species because of either or both genetic and environmental influences can be easily assessed using various molecular markers. Among various molecular markers, SSR markers in particular have many advantages compared with other marker
systems as follows: the first advantage is high reproducibility, which is most important in genetic analysis; the second advantage of the SSR marker system is the polymorphic genetic information contents, and the hypervariable nature of SSRs produces very high allelic variations even among very closely related varieties; and the third advantage relates to the codominant nature of SSR polymorphisms, as the codominant nature of SSRs is suitable for genetic analysis in segregation of F$_2$ populations or pedigree analysis in hybrids (Park et al. 2009; Pejic et al. 1998; Sa et al. 2018).

In plants, the presence of SSRs was first demonstrated by the hybridization of poly (G-T) and poly (A-G) oligonucleotide probes on phage libraries of tropical tree genomes (Condit and Hubbell 1991). A search of published DNA sequences revealed that SSRs are also abundant in diverse plant genomes (Morgante and Olivieri 1993; Wang et al. 1994). Therefore, SSRs have become the preferred molecular marker system for the analysis of plant genetics and ecology. However, SSR markers can only be applied to crop species with large amounts of EST or cDNA sequences that are freely accessible to the public. Recently, the transcriptome sequencing approach RNAseq has become a powerful tool for novel gene discovery and the development of molecular markers for nonmodel crop, such as Perilla crop. In the case of the Perilla crop, in a previous study, we sequenced and assembled one cultivated type (PF98095) of P. frutescens var. frutescens using transcriptome sequencing by RNA-seq (Tong et al. 2015). This information will help develop SSR primers for Perilla crop. As explained in the Introduction, Sa et al. (2018) detected a total of 15,991 SSR loci based on the SSR flanking sequences via de novo assembly of the PF98095 RNA-seq data (Tong et al. 2015), which were classified based on the number of repeating units. Dinucleotide SSRS had the highest number of SSRS (9,910), followed by trinucleotide SSRS (5,652) and tetranucleotide SSRS (429) (Sa et al. 2018). In the Perilla crop, analysis of the distribution of different repeat motifs showed that dinucleotide repeats were the most abundant (62.0%) type, followed by trinucleotide repeats (35.3%), with the two together comprising 97.3% of the eSSR repeats. This finding is important for developing a large number of effective eSSR primers in Perilla crop, as the markers associated with smaller SSR motif lengths (di- and tri-) are more variable than other motif lengths (Ellegren 2004). Our results also showed that dinucleotide repeats were more abundant than other repeat units in the Perilla crop. While an almost equal distribution of di- and trinucleotide repeats was reported in ramie (Liu et al. 2013), a high abundance of trinucleotide repeats was reported in flax (Cloutier 2009), jute (Saha et al. 2017), and kenaf ESTs (Zhang et al. 2015).

In this study, a total of 130 alleles with 39 SSRs were detected segregating in the 9 Perilla accessions, which yielded an average of 3.3 alleles per locus (Supplement Tables 1, 2). Depending on the number of repeat units in the 39 SSR primer sets, the dinucleotide SSRs such as KNUPF87, KNUPF97, KNUPF103, KNUPF105, KNUPF110, and KNUPF116 showed comparatively more allelic bands and genetic diversity values than the tri- and tetranucleotide SSRs in Perilla accessions. In our study, although we analyzed only nine Perilla accessions, the allele number, GD, PIC, and MAF values for 39 new Perilla EST-SSR markers were found to determine the unique genetic profiles of individual genotypes of the Perilla crop. Therefore, these new SSR primer sets may be useful in the future for studying GD for Perilla germplasm resources by association mapping and designating important genes/QTLs for future genetic and breeding programs in Perilla crop.

In addition, analysis of genetic diversity and relationships and association mapping of plant breeding materials will help develop new molecular markers associated with target morphological traits or useful varieties and lines in Perilla crop breeding programs. In our study, we developed an F$_2$ population from a cross between PF13-110 (female parent, cultivated var. frutescens) and PF13-160 (male parent, weedy var. crispa) to develop molecular
markers for traits showing morphological differences between two cultivated types of var. *frutescens* and var. *crispa* through 40 individuals of the F$_2$ population of the *Perilla* crop (Fig. 1, Tables 1, 2). According to our results, the 40 individuals of the F$_2$ population showed discontinuous variations within the range of morphological characteristics of their parental lines, as shown in Tables 2 and 5. That is, some morphological traits investigated in our study showed a Mendelian segregation ratio of 1:2:1 (or AA:AB:BB), while other traits showed different separation patterns in 40 individuals of the F$_2$ population. For example, in the case of QL1, of the 40 lines of the F$_2$ population, 12 lines were large, 14 lines were medium, and 14 lines were small; whereas in the case of QL2, 28 lines were green, 6 lines were mixed green/purple, and 6 lines were purple. However, unfortunately, as mentioned in the Introduction, the F$_2$ population lines used in this study did not represent the entire population, so it was not possible to predict the exact segregation ratio of the F$_2$ population.

In our study, we also performed a correlation analysis for nine leaf- and plant-related traits in 40 individuals of the F$_2$ population (Table 6). The results showed that there were highly statistically significant positive correlations of QL2 with QL3 (0.682**) and QL6 (0.429**), of QL3 with QL6 (0.390*) and QL7 (0.462**), of QL5 with QL8 (0.489**) and QL9 (0.313*) at P < 0.01 or P < 0.05. Among the morphological traits examined in our study, leaf and plant color-related traits, such as QL2, QL3, QL6, and QL7, and leaf shape-related traits, such as QL5 and QL8, were closely related to each other. In addition, to better understand the genetic relationships among 40 individuals of the F$_2$ population, we analyzed the phylogenetic relationships of the 40 individuals of the F$_2$ population and their two parental lines using 37 SSR markers. In the phylogenetic tree analysis, the 40 individuals of the F$_2$ population and two parental lines were clustered into two major groups. Except for exceptional individuals, Group I mainly contained many individuals with similar characteristics to Parent B, while Group II contained many individuals with similar characteristics to Parent A (Table 2, Fig. 2). Therefore, these separated individuals and leaf- and plant-related traits of the F$_2$ population are considered to be useful genetic materials for finding molecular markers associated with the leaf and plant morphological characteristics of *Perilla* crop through association mapping analysis.

Association mapping has been proposed as a selection method to identify loci involved in the inheritance of complex traits (Risch and Merikangas 1996). In this study, we developed an F$_2$ population for association mapping analysis to find molecular markers associated with leaf- and plant-related traits of *Perilla* crop. Additionally, in our study, we used SSR markers for association mapping analysis because SSR markers are highly polymorphic and generally codominant (Park et al. 2009; Powell et al. 1996; Sa et al. 2018, 2019). Therefore, the SSR marker is considered to be a useful analysis method for identifying separation patterns of allelic bands in the F$_2$ population.

To select the SSR markers associated with leaf- and plant-related traits of the F$_2$ population, we analyzed marker-trait associations (SMTAs) between 37 SSR markers and 9 leaf- and plant-related characteristics in 40 individuals of the F$_2$ population using TASSEL software. From the results, we identified 12 and 11 SSR markers associated with leaf-related traits and plant-related traits, respectively (Table 7). In the case of leaf-related traits, 12 SSR markers, namely, KWPF25, KNUPF11, KNUPF21, KNUPF49, KNUPF60, KNUPF74, KNUPF81, KNUPF89, KNUPF103, KNUPF108, KNUPF109, and GBPEM111, were associated with QL2, QL3, QL5, and QL8. In addition, in the case of plant-related traits, 11 SSR markers, namely, KNUPF5, KNUPF11, KNUPF15, KNUPF21, KNUPF29, KNUPF39, KNUPF54, KNUPF61, KNUPF82, KNUPF88, and GBPEM111, were associated with QL4, QL6, QL7, and QL9 (Table 7). Our results showed that with the exception of QL1, most leaf- and plant-related traits were associated
with 1 to 7 SSR markers, depending on the leaf- and plant-related characteristics. Additionally, some SSR markers (KNUPF11, KNUPF21, GBPEM111) were associated with leaf- or plant-related characteristics, such as QL3, QL4, QL6, QL7, and QL9. For example, KNUPE11 was found to be associated with QL3, QL6, and QL7 traits; GBPEM111 was associated with QL8 and QL9 traits; and KNUPE21 was associated with QL4 and QL8 traits (Table 7).

In a previous study by Lim et al. (2000), bulk segregant analysis (BSA) was performed to identify SSR markers linked to leaf- and seed-related traits in *Perilla* crop. They reported that four SSR markers (KNUPF15, KNUPF21, KNUPF29, and KNUPF60) were associated with the color of the leaf surface, four SSR markers (KNUPF11, KNUPF15, KNUPF21, and KNUPF60) were associated with the color of the leaf reverse side, and three SSR markers (KNUPF25, KNUPF61, and GBPEM111) were associated with the depth of leaf teeth. In addition, five SSR markers (KNUPF11, KNUPF12, KNUPF16, KNUPF29, and KNUPF42) were associated with seed-related traits. According to our results, among the SSR markers associated with the leaf- and plant-related traits, eight SSR markers (KNUPE11, KNUPF15, KNUPF21, KWPE25, KNUPF29, KNUPE60, KNUPE61, GBPEM111) appeared to overlap with leaf-related traits previously reported by Lim et al. (2000). Therefore, these SSR markers are thought to be useful molecular markers for selecting leaf- and plant-related traits in *Perilla* crop. Additionally, these SSR markers are thought to be useful molecular markers for distinguishing between two cultivated types of *Perilla* crop. However, in *Perilla* crop, it is difficult to compare genetic characteristics because genomic information on SSR markers associated with leaf- and plant-related characteristics is still lacking.

In the future, if studies such as genome analysis for each chromosome are actively conducted in *Perilla* crop, it will be possible to analyze genomic information of specific *Perilla* SSR markers related to leaf- and plant-related characteristics. As explained in the Introduction, many taxonomic studies have been conducted to distinguish between two cultivated and weedy types of *Perilla* crop using morphological characteristics and molecular markers, but they failed to distinguish the two cultivated types of *Perilla* crop because of the existence of intermediate types, such as weedy plants. Therefore, the SSR markers associated with leaf- and plant-related characteristics selected in this study are expected to provide useful information for distinguishing the two cultivated types of *Perilla* crop and their weedy types in the future. In addition, the new *Perilla* SSR primers described in this study could be helpful in identifying genetic diversity and genetic mapping, designating important genes/QTLs for *Perilla* crop breeding programs, and allowing *Perilla* breeders to improve leaf and plant quality through MAS breeding programs.

**Declarations**

**Acknowledgments**

This study was supported by the Basic Science Research Program through the National Research Foundation of Korea (NRF) funded by the Ministry of Education, Science, and Technology (#2016R1D1A1B01006461) and the Cooperative Research Program for Agriculture Science & Technology Development (project no. PJ014227032019 and PJ0142272019), Rural Development Administration, Republic of Korea.

**Funding** The Basic Science Research Program through the National Research Foundation of Korea (NRF) funded by the Ministry of Education, Science, and Technology (#2016R1D1A1B01006461) and the Cooperative Research Program for Agriculture Science & Technology Development (project no. PJ014227032019 and PJ0142272019), Rural Development Administration, Republic of Korea.
Author contributions  J.K. Lee wrote the manuscript and designed the experiments. J.Y. Kim and K.J. Sa performed the experiment and analyzed the data, and Y.J. Ha helped to draft the manuscript. All authors commented on previous versions of the manuscript and approved the final manuscript.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflicting interests.

Availability of data and material All data generated or analyzed during this study are included in this published article and its supplementary information files.

Ethical approval This article does not contain any studies with human subjects or animals performed by any of the above authors.

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Figures
Figure 1

Morphological features of two parental lines of the F2 population used for crossing between (a) *Perilla frutescens* var. *frutescens* (parent A: PF13-110, ♀) and (b) *P. frutescens* var. *crispa* (parent B: PF13-160, ♂).
Figure 2

Neighbor-joining tree of all 40 individuals of the F2 population and two parental lines inferred from 39 SSR markers; Group I (red) represents F2 individuals, including Parent B (PF13-160, ♂), and Group II (green) represents F2 individuals, including Parent A (PF13-110, ♀).

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