First microsatellite markers for *Scaligeria lazica* Boiss. (Apiaceae) by next-generation sequencing: population structure and genetic diversity analysis

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The Apiaceae family includes a few agronomic and medicinal species, one of which is *Scaligeria lazica* Boiss. In this study, the genetic diversity of *S. lazica* was analyzed based on novel simple sequence repeat (SSR) markers using next-generation sequencing (NGS). A total of 15.17G clean Illumina data set was obtained and dinucleotide repeats were the most abundant repeats in *S. lazica*. Of the tested 150 SSR primer pairs, 139 ones produced amplification and 84 ones were polymorphic. Forty polymorphic SSR loci were used in genetic diversity analysis of 40 *S. lazica* accessions from four locations. A total of 264 alleles were amplified with an average of 6.6 alleles per locus. The polymorphism information content (PIC) was 0.60, while the observed homozygosity (Ho) and expected heterozygosity (He) values were 0.47 and 0.66, respectively. According to cluster and structure analysis, all accessions were grouped into four different clusters according to their collection sites. The SSR markers developed in this study can be tested for other *Scaligeria* species due to their high transferability and can be used for genetic studies in genus *Scaligeria* DC.

**ABSTRACT**

The Apiaceae family includes a few agronomic and medicinal species, one of which is *Scaligeria lazica* Boiss. In this study, the genetic diversity of *S. lazica* was analyzed based on novel simple sequence repeat (SSR) markers using next-generation sequencing (NGS). A total of 15.17G clean Illumina data set was obtained and dinucleotide repeats were the most abundant repeats in *S. lazica*. Of the tested 150 SSR primer pairs, 139 ones produced amplification and 84 ones were polymorphic. Forty polymorphic SSR loci were used in genetic diversity analysis of 40 *S. lazica* accessions from four locations. A total of 264 alleles were amplified with an average of 6.6 alleles per locus. The polymorphism information content (PIC) was 0.60, while the observed homozygosity (Ho) and expected heterozygosity (He) values were 0.47 and 0.66, respectively. According to cluster and structure analysis, all accessions were grouped into four different clusters according to their collection sites. The SSR markers developed in this study can be tested for other *Scaligeria* species due to their high transferability and can be used for genetic studies in genus *Scaligeria* DC.

**Introduction**

There is an overwhelming plant biodiversity on earth and efforts have been made to categorize plants based on their size, forms, habitat, structure, anatomy, biochemical and molecular features to interpret the relationships among them [1–4].

Apiaceae Lindl. (Umbelliferae) includes 300–455 genera with 3000–3750 species and is one of the best known families of flowering plants. Members of this family are of great economic significance due to their culinary use (vegetables, condiments) and for their application in medicine [5,6]. The genus *Scaligeria* DC. belongs to Apiaceae and is represented by seven species, of which two are endemic in Turkey [7,8]. The naturally growing *Scaligeria* species in Turkey are as follows: *S. napiformis* (Wildl.ex Spreng) Grande, *S. tripartita* (Kalen.) Schischkin, *S. lazica* Boiss. (endemic), *S. meifolia* (Fenzl) Boiss., *S. glaucescens* (DC.) Boiss., *S. hermonis* Post and *S. capillifolia* Post. (endemic). *S. lazica* has a strong characteristic smell of anise due to the presence of trans-anethol (1-methoxy-4-(1-propenyl) benzene) in its oil, which is known as ‘laz anasonu’ in the Black Sea Region of Turkey [8,9]. The economically important trans-anethole-containing oils are used widely in the food and liquor industries [10]. trans-Anethole originates from anise in raki and is also used for flavour in many alcoholic drinks such as Middle Eastern arak, Colombian aguardiente, French spirits absinthe, anisette and pastis, Greek ouzo, mastika in Bulgaria and the former Yugoslav Republic of Macedonia. [11]. In addition, anethole shows estrogenic effects and is useful in providing relief from rheumatism and lower back pain [12,13]. Interestingly, some phytochemical markers of *Pimpinella* spp. are also detected in *S. tripartita* and *S. lazica* species [9,14–16]. Studies on *Scaligeria* spp. are still very limited in the literature to clarify the taxonomy of the genus. A promising approach to resolving these problems in the taxonomy of *Scaligeria* species are molecular studies.

Simple sequence repeats (SSRs) are a very powerful technique due to their high rate of polymorphism and repeatability. The potential use of SSR molecular markers in the family Apiaceae has been demonstrated by different studies. Novel SSR markers were developed in Apiaceae and they have been used to study the genetic diversity, phylogenetic relationships and germplasm characteristics. The potential use of SSRs in Apiaceae has been demonstrated by different studies [17–21].
Next-generation sequencing is a very advantageous technology as regards the millions of sequences that can be generated at once with a lower unit cost. It is possible to have thousands of microsatellite loci sequenced very rapidly and cost-effectively in plants [22,23]. SSR markers in plants are used as a prominent genomic resource in the botanical sciences. SSRs are highly repeatable and transferable across closely related species [24–27]. An important feature of SSR markers is that multiple alleles per locus can be amplified, e.g. 10 alleles in a population. This variability makes SSR markers highly polymorphic and useful for genetic diversity analysis [27,28].

To date, there are no reports on SSR development and genetic diversity studies in _Scaligeria lazica_. Therefore, in this study, we aimed to develop novel SSR markers by next-generation sequencing and to study the population structure in _S. lazica_. Due to their high transferability, the markers from this study can also be useful as molecular tools for other species in genus _Scaligeria_.

**Materials and methods**

**Plant materials and DNA extraction**

Stems of _Scaligeria lazica_ [7] were collected from four locations: Camlıhemsin (RCH) and Cayeli (RCE) from Rize province and Altindere (TAD) and Sumela (TSM) from Trabzon province of Turkey. Ten accessions were collected from each location. Figure 1 and Table 1 represent

| No. | Accession name | Acronym | Location          | Latitude (N) | Longitude (W) |
|-----|----------------|---------|-------------------|--------------|---------------|
| 1   | Camlıhemsin    | RCH     | Rize Camlıhemsin  | 41° 34'      | 41° 10'       |
| 2   | Cayeli         | RCE     | Rize Cayeli       | 40° 53'      | 40° 44'       |
| 3   | Altindere      | TAD     | Trabzon Altindere | 40° 43'      | 39° 38'       |
| 4   | Sumela         | TSM     | Trabzon Sumela    | 40° 41'      | 39° 39'       |

Figure 1. Sampling sites of _S. lazica_ accessions from four different locations in Turkey.

**SSR loci development by illumina sequencing and primer design**

A DNA sample of one _S. lazica_ accession (RCH-5) from the Rize–Camlıhemsin location was used for next-
PCR reactions were performed in a total volume of 12.5 μL containing 20 ng DNA, 75 mmol/L Tris–HCl (pH: 8.8), 20 mmol/L (NH₄)₂SO₄, 2.0 mmol/L MgCl₂, 0.01% Tween 20, 200 μmol/L of each deoxyribonucleoside triphosphate (dNTP), 10 nmol/L M13 tailed forward primer at the 5’ end, 200 nmol/L reverse primer, 200 nmol/L universal M13 tail primer (5’-TGTTAAAAACGACGCCAGT-3’) labelled with one of the FAM, VIC, NED or PET dyes, and 0.6 U of Hot Start Taq DNA polymerase.

Amplification was performed in a Veriti thermal cycler (Applied Biosystems, Massachusetts, USA) in two steps as follows: initial denaturation at 94 °C for 3 min, followed by 10 cycles at 94 °C for 30 s, 56–60 °C for 45 s and 72 °C for 60 s. Then, a second step included 30 cycles at 94 °C for 30 s, 58 °C for 45 s and 72 °C for 60 s. A final extension at 72 °C for 10 min was included in the program. When the PCRs were completed, the reactions were subjected to denaturation for capillary electrophoresis in ABI 3130xl genetic analyzer [Applied Biosystems Inc., Foster City, CA, USA (ABI)] using 36 cm capillary array with POP7 as the matrix (ABI). Denaturation of the samples was completed by mixing 0.5 μL of the amplified product, 0.2 μL of the size standard and 9.8 μL of formamide. The fragments were resolved using ABI data collection software 3.0. and SSR fragment analysis was performed with GeneScan Analysis Software 4.0 (ABI).

Data analysis

After determining the alleles of each SSR locus by capillary electrophoresis (Figure 2), the number of alleles (Na), the number of effective alleles (Ne), observed (Ho), expected (He) heterozygosity and the number of private alleles were calculated using GenAlEx V6.5 software [34]. The polymorphism information content (PIC) was also calculated using PowerMarker ver 3.25 software [35]. A dendrogram was obtained using Jaccard coefficient in NTSYSpc v2.21c [36] software by unweighted pair-group method with arithmetic averages (UPGMA).

Population structure and identification of admixed individuals was performed using the model-based software program, STRUCTURE 2.3.4 [37]. In this model, a number of populations (K) are assumed to be present, each of which is characterized by a set of allele frequencies at each locus. Individuals in the sample are assigned to populations (clusters), or jointly to more populations if their genotypes indicate that they are admixed. The posterior probabilities were estimated using the Markov Chain Monte Carlo (MCMC) method. The MCMC chains were run with a 50,000 burn-in period, followed by 500,000 iterations, using a model allowing for admixture and correlated allele frequencies. Run of STRUCTURE was performed by setting K from 1 to 5.

Results and discussion

Development of novel SSR loci and genetic diversity analysis in S. lazica

A total of 15.17G clean Illumina data were generated to search SSR loci and 1982 SSR primer pairs were successfully designed. A total of 1982 SSRs dinucleotide repeats were the most abundant (70.63%), followed by trinucleotide (11.5%) and hexanucleotide repeats (11.3%), while tetranucleotide (4.3%) and pentanucleotide (2.1%) repeats were the least frequent (Figure 3). The most abundant dinucleotide and trinucleotide repeat motif types were TC/GA and TTC/GAA, respectively.
Of the tested 150 SSR primer pairs by gradient PCR analysis, 139 primer pairs produced successful amplification. Fifty-five of 139 primer pairs were monomorphic in the screening of the SSRs in the eight accessions. Eighty-four SSR loci (60.4%) generated polymorphic fragments (Table S1 in the Online Supplement) and 40 were used for further analysis to fingerprint 40 Scaligeria accessions. The Genebank accession numbers, the sequences of forward and reverse primers, repeat motifs, annealing temperatures, product sizes, $Na$, $Ne$, $Ho$, $He$ and PIC values of the 40 SSR loci are presented in Table 2.

Forty microsatellite loci were used to fingerprint 40 S. lazica accessions belonging to four populations and a total of 264 alleles were obtained, ranging from 2 to 13 per locus (Table 2). The loci SclgSSR1636 (13), SclgSSR285 (12), SclgSSR762 (12), SclgSSR1614 (11) and SclgSSR2725 (11) produced the highest number of alleles. The effective number of alleles ranged from 1.35 (SclgSSR323) to 7.04 (SclgSSR285). The $Ho$ was between 0.10 (SclgSSR179) and 1.00 (SclgSSR1645 and SclgSSR1544). The $He$ values changed from 0.25 (SclgSSR323) to 0.86 (SclgSSR285) with an average of 0.66. The PIC values ranged from 0.24 to 0.84 with an average of 0.60. The locus SclgSSR285 had the highest PIC value (Table 2).

The average values of population genetic parameters in this study are summarized in Table 3. The highest degree of genetic diversity values occurred in the RCE location, while the TSM location had the lowest values and RCH and TAD locations had similar degrees of genetic diversity. The number of private alleles in the RCH location was quite higher than that in the other three locations. For example, it was four times more than in TAD and 16 times more than in TSM. The polymorphism rate of 264 alleles was the highest (97.3%) in the RCE location, whereas the TSM (91.3%) and RCH (92.4%) locations had the lowest polymorphisms.
Table 2. Repeat motifs, product sizes, number of alleles (Na), number of effective alleles (Ne), observed heterozygosity (Ho), expected heterozygosity (He) and polymorphism information content (PIC) values based on 40 SSR loci developed from S. lazica.

| Accession number | Motif       | Na  | Ne  | Ho  | He  | PIC  | Number of private alleles |
|------------------|-------------|-----|-----|-----|-----|------|---------------------------|
| 1 K234457        | (AG)₉       | 7   | 4.07 | 0.71 | 0.75 | 0.72 | 91.3                      |
| 2 K234458        | (CT)₄       | 5   | 3.43 | 0.50 | 0.71 | 0.66 | 69.6                      |
| 3 K234459        | (AGA)₂      | 2   | 1.74 | 0.10 | 0.43 | 0.34 | 71.2                      |
| 4 K234468        | (AG)₁₂      | 12  | 7.04 | 0.64 | 0.86 | 0.84 | 84.5                      |
| 5 K234470        | (AG)₁₄      | 3   | 1.35 | 0.25 | 0.26 | 0.24 | 78.3                      |
| 6 K234485        | (CT)₁₅      | 6   | 3.99 | 0.40 | 0.75 | 0.71 | 84.6                      |
| 7 K234491        | (AG)₂₁      | 12  | 6.26 | 0.76 | 0.84 | 0.82 | 82.4                      |
| 8 K234493        | (GA)₁₆      | 6   | 2.93 | 0.38 | 0.66 | 0.6  | 66.7                      |
| 9 K234495        | (AG)₁₄      | 3   | 2.47 | 0.43 | 0.60 | 0.53 | 53.1                      |
| 10 K234496       | (TC)₁₄      | 5   | 3.13 | 0.36 | 0.68 | 0.62 | 62.1                      |
| 11 K234497       | (GT)₁₅      | 6   | 2.56 | 0.46 | 0.61 | 0.57 | 57.1                      |
| 12 K234499       | (AC)₁₄      | 8   | 3.83 | 0.51 | 0.74 | 0.7  | 70.0                      |

Novel SSR markers and genetic diversity in S. lazica

Due to the codominant nature and high polymorphism, SSR markers are reliable tools for genetic diversity and population structure analysis as well as genetic mapping, fingerprinting and parental identification in plants. However, sequence information is needed for primer development for each species; therefore, the development process is expensive and time-consuming. Next-generation sequencing technology provides an opportunity to generate thousands of SSR loci in a very short time at a lower cost than Sanger sequencing. However, testing the loci for polymorphism in the lab is still expensive and time-consuming. However, it is a very powerful technique, once the SSR loci have been developed.

The Apiaceae family includes several agronomic and medicinal vegetable species, one of which is S. lazica in the genus Scaligeria. There are a limited number of SSR markers in the literature for Apiaceae family [21,39,40].

Table 3. Mean of population genetic parameters in this study.

| Populations | Sample size | Na  | Ne  | Ho  | He  | PIC  | Polymorphism (%) | Number of private alleles |
|-------------|-------------|-----|-----|-----|-----|------|------------------|---------------------------|
| RCH         | 10          | 3.10 | 2.12 | 0.43 | 0.42 | 0.37 | 92.4              | 65                        |
| RCE         | 10          | 4.10 | 2.76 | 0.57 | 0.57 | 0.52 | 97.3              | 44                        |
| TSM         | 10          | 2.53 | 1.74 | 0.42 | 0.35 | 0.30 | 91.3              | 4                         |
| TAD         | 10          | 2.93 | 2.11 | 0.44 | 0.46 | 0.40 | 94.7              | 16                        |
Till now, to the best of our knowledge, there is no study on development of SSR markers in *Scaligeria* in the literature. Thus, this is the first report that presents polymorphic SSR loci for *Scaligeria*.

In this study, from 15.17G clean Illumina data, 1982 SSR primer pairs were successfully designed and the most frequent repeats were identified to be dinucleotides (70.63%), in agreement with previous reports. In Apiaceae family, Rijal et al. [21] and Cavagnaro et al. [17] reported polymorphic microsatellite markers in *Heracleum persicum* Desf. ex Fisch and *Daucus carota* L., respectively, and they found that the most frequent repeats are dinucleotides, as in our study. The most abundant dinucleotide motif in *Scaligeria* was TC/GA. The GA motif is generally one of the most abundant dinucleotide repeats in plants [41–43]. There are several reports on SSR development in Apiaceae: Michalcyzk et al. [39] generated 12 SSR loci from *Cnidium dubium* (Schkuhr) Thell. and obtained an average of 8.3 alleles per locus by characterizing 40 individuals. The authors calculated the Ho and He values as 0.62 and 0.63, respectively. Tew et al. [40] reported eight polymorphic SSR loci from *Lilaeopsis schaffneriana* subsp. recurva (A.W.Hill) Affolter by fingerprinting 48 individuals from two natural populations and obtained 4.4 alleles per locus. Maksylewicz and Baranski [44] developed 27 polymorphic SSR loci from carrot (*D. carota* L. ssp. *sativus*) and tested in two F1 hybrids, ten open pollinated cultivars and six landraces, obtaining an average of 9.4 alleles. The PIC, Ho and He values were reported as 0.56, 0.39 and 0.61, respectively. Rijal et al. [21] generated 25 polymorphic microsatellite markers in *H. persicum* by characterizing eight geographically distant samples of *H. persicum* and obtained Na of 2.3 alleles per locus. The average values of Ho and He for the 25 polymorphic SSR loci were 0.57 and 0.42, respectively. In this study, 84 polymorphic SSR loci were developed and 40 were used in genetic diversity analysis of *S. lazica*. The selected 40 SSR loci

Figure 4. An UPGMA dendrogram of 40 *S. lazica* accessions from four locations obtained by analyzing 40 SSR loci.
produced 264 polymorphic alleles. The number of generated and characterized polymorphic SSR loci was higher than those in the above-mentioned studies for Apiaceae family [21,40–44]. The mean PIC value obtained in this study was higher (PIC = 0.60) than those in the previous reports discussed above. The average $Ho$ value for the 40 polymorphic loci in this study was 0.47, which was higher than that for $L. schaffneriana$ subsp. recurva and

![Figure 5](image-url)

**Figure 5.** Population structure of 40 accessions estimated from 40 SSRs using STRUCTURE program. $K = 2$ to $K = 5$ represent the sub-population of the 40 accessions originating from different places of Turkey.

![Figure 6](image-url)

**Figure 6.** Values of $\Delta K$, with its modal value detecting a most probable $K$ of two subgroups ($K = 2$), including a small peak at four subgroups ($K = 4$).
**Cluster analysis and genetic structure of *S. lazica***

To identify the true optimal number of subsets (K) in STRUCTURE, LnP(D) and ΔK were chosen, as described in Ren et al. [45]. The K value that provides the maximum likelihood, called LnP(D) in STRUCTURE, is generally considered as the optimal number of subdivisions [37]. Our results from two main genetic sites can be summarized with the maximum of ad hoc measure ΔK at K = 2 (Figure 6). All the accessions separated and consistently grouped well in both cluster and structure analysis. The dendrogram and structure analysis at K = 2 divided *Scaligeria* accessions into two main clusters: the first cluster included all RCH accessions, whereas the second cluster contained the accessions from the other collection sites. The second cluster divided into two subclusters: the first subcluster contained RCE with 10 accessions, whereas the second subcluster contained TSM and TAD which included 10 accessions in each group.

According to both cluster and structure analyses, the accessions from RCH and TAD locations were the most distant group, whereas the closest groups were TSM and TAD. These two locations were also geographically the closest (Table 1 and Figure 1). The high number of private alleles separated the RCH location from the others, which was also supported by cluster and structure analyses at K = 2 (Figures 4 and 5). In the structure analysis at K = 4, all accessions were classified into four different clusters according to their collection sites. The structure analysis demonstrated that the accessions in RCE location were the most diverse ones, which was supported by structure analysis at K = 5. The results presented in this study, therefore, provide information about the genetic diversity levels of *S. lazica* from different locations in Turkey. Similar diversity can also be observed in the production of trans-anethol by the accessions, which is the most important product of *Scaligeria* species.

**Conclusions**

Here we present novel SSR markers for the first time in *S. lazica* by next-generation sequencing. Dinucleotide repeats were most abundant (70.63%), followed by trinucleotide (11.5%) and hexanucleotide repeats (11.3%) in *S. lazica*. SSRs are a co-dominant marker system and can be applied not only as a useful molecular tool for the analysis of genetic variation, population structure and phylogenetic relationships of *S. lazica* accessions collected from different locations, but also may contribute to development of DNA markers linked to the agronomic and medicinal characters in *S. lazica*. Furthermore, novel SSR markers developed from this study can be also analyzed to test their transferability to other *Scaligeria* species.

**Disclosure statement**

No potential conflict of interest was reported by the authors.

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