Characterization of Nuclear and Chloroplast Microsatellite Markers for *Falcaria vulgaris* (Apiaceae)

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ABSTRACT

*Falcaria vulgaris* (sickleweed) is native to Eurasia and a potential invasive plant of the United States. No molecular markers have been developed so far for sickleweed. Characterization of molecular markers for this plant would allow investigation into its population structure and biogeography thereby yielding insights into risk analysis and effective management practices of the plant. In order to characterize the molecular markers, DNA samples were collected from eight populations in Iowa, Nebraska, and South Dakota. Nuclear microsatellite markers developed for other Apiaceae taxa were screened and tested for inter-generic transferability to sickleweed. The chloroplast *trnL* intron and *trnL-F* intergenic spacer regions were sequenced and the sequences were used to design primers to amplify the microsatellites present within each region. We characterized eight polymorphic microsatellite markers for sickleweed that included six nuclear and two chloroplast markers. Our result showed inter-generic transferability of six nuclear microsatellite markers from *Daucus carota* to *F. vulgaris*. The markers we characterized are useful for population genetic study of *F. vulgaris*.

Keywords: *Falcaria vulgaris*; Invasive Species; Microsatellite; *trnL* Intron; *trnL-trnF* Intergenic Spacer; Sickleweed

1. Introduction

Sickleweed (*Falcaria vulgaris* Bernh.; Apiaceae) is native to Europe and Asia [1] and was introduced to the United States in early 1920s [2]. It has been reported from 35 counties across 16 states in the United States and exhibits disjunct distribution in the Midwestern and Eastern USA [3]. Sickleweed exhibits some characteristics of invasive plant species including the production of a large number of seeds, effective seed dispersal mechanism whereby the seeds attached to the stem are carried away by the wind as the stems break at the nodes after the plant senescence, and the ability of the plant to reproduce asexually through root sprouting. These characteristics are perhaps facilitating its emergence as an aggressive weed in the Midwest [4]. Continuous increase in area coverage of sickleweed in Fort Pierre National Grassland (FPNG) and Buffalo Gap national Grassland (BGNG) of South Dakota has attracted attention of ecologists in the Midwest. The plant has also been listed as potential invasive plant by Nebraska Invasive Species Council [5].

Both nuclear and chloroplast DNA markers are commonly used for the genetic analysis of invasive plant populations particularly to predict the invasiveness of the introduced species, identify the source populations and help to design effective control programs for invasive species [6]. Microsatellite markers are one of the most preferred molecular markers because they are co-dominant, hyper-variable and are highly reproducible [7,8]. However, development of novel microsatellite markers is expensive and quite laborious task [9]. Cross-species transferability of microsatellite markers and identification of chloroplast microsatellites using universal chloroplast markers for sequencing chloroplast region are options that can avoid high cost and long time needed for marker development [10,11]. The markers are useful for investigating population structure and phylogeography of the introduced species, and are also useful for the study of comparative studies of different species [10], the process of population divergence and speciation process [12]. No molecular markers have previously been developed for sickleweed that would allow us study the genetic structure of this plant. Here we report on inter-generic transferability of six nuclear microsatellite markers from *Daucus carota* to *F. vulgaris*, and two polymorphic chloroplast microsatellite markers.

2. Methods

2.1. Screening of Microsatellite Markers

We reconstructed a phylogeny of the family Apiaceae based on nuclear ribosomal Internal Transcribed Spacer
(ITS) DNA sequences available in GenBank and searched for microsatellite markers developed for taxa closely related to *Falcaria vulgaris*. Based on our phylogenetic analysis, we decided to screen microsatellite markers developed for *Daucus carota* and *Heracleum mantegazzianum*. We selected 85 microsatellite markers with di-, tri- and tetranucleotide repeats developed for *Daucus* [13,14] and six markers developed for *Heracleum* [15] for the genetic analysis of sickleweed. Fresh leaf tissues were collected in silica gel from eight populations (Table 1) from Iowa (one population), Nebraska (three populations) and South Dakota (four populations). The voucher specimens except for the populations from Boyd County, Nebraska were deposited at South Dakota State University Herbarium (SDC). The silica gel dried leaf samples were ground to a fine powder and total DNA was extracted using DNeasy Plant Minikit (Qiagen Corp., Valencia, CA).

For screening the microsatellite markers, PCR was carried out in a reaction mixture of 15 μl containing 50 ng genomic DNA, 3 μl of 5 X buffer (Promega), 1.2 μl of 10 mM dNTPs (Promega), 2 μl of 25 mM MgCl₂ (Promega), 1 μl each of 10 pM forward and reverse primers and 2 units of Jump Start Taq polymerase (Sigma-Aldrich). The PCR conditions were an initial denaturation of 5 minutes at 72°C, followed by 40 cycles of 1 minute denaturation at 94°C, 20 seconds of annealing temperature and 1 minute extension at 72°C, and final extension of 5 minutes at 72°C. Electrophoresis was carried out in 1.2% agarose gel to evaluate the quality of PCR products and the presence of repeat motif in the amplicons was verified by re-sequencing the PCR products.

For chloroplast microsatellite, we sequenced the trnL intron and trnL-F intergenic spacer (using the primer pair 5'- C G A A T C G T A G A C G T A C G - 3' and 5'- ATTTGAACTGGTGACACGAG-3') of chloroplast gion and found two mononucleotide repeats with more than ten mononucleotide repeats and designed the primers using primer3 software (http://primer3.wi.mit.edu/) to amplify these mono-nucleotide repeats.

### 2.2. Genotyping, Test for Potential Artifacts and Data Analyses
For genotyping, PCR was carried out in a reaction mixture of 15 μl containing 50 ng genomic DNA, 3 μl of 5 X buffer, 1.2 μl of dNTPs, 2 μl of 25 mM MgCl₂, 0.5 μl of 10 pM forward primer, 0.5 μl of 10 pM forward primer tagged with M13 tail, 1 μl of 10 pM reverse primers each and 2 units of Taq polymerase. The PCR conditions were similar to that of primer screening PCR conditions. The PCR products were genotyped using 3730 x1 DNA analyzer (Applied Biosystems) at Iowa State University DNA Facility.

Genemarker V2.4.0 (Softgenetics) was used to visualize the genotyping data and create allele reports. The possible genotyping artifacts such as stuttering, large allele drop-out and presence of null allele were tested using Micro-Checker [16]. The analysis of microsatellite polymorphisms including number of alleles, observed and expected heterozygosity were performed using Arlequin V3.1 [17], the polymorphism information content (PIC) value was computed using the excel microsatellite toolkit [18] and haploid diversity for chloroplast microsatellite was computed using Genalex V. 6.41 [19].

### 3. Results and Discussions
The probability of microsatellite marker transferability reduces with increased phylogenetic relationship [20]. There are several evidences of microsatellite marker transferability within genus but few evidences at higher taxonomic level than genus. For example, microsatellite transferability success rate within the genera in eudicots are approximately 60% compared to 10% across genera [10]. Therefore, it is always beneficial to identify phylogenetically close relatives before screening the microsatellite markers for cross species transferability when microsatellites markers are not available for sister species within the genus. In our phylogenetic analysis, we found *Apium graveolens*, *D. carota*, *Eryngium alpinum* and *H. mantegazzianum* were the only taxa with microsatellite markers developed within the clade and they were distantly related to *F. vulgaris*. *Falcaria* is a monotypic genus. Previous studies have shown transferability of *Daucus* and *Heracleum* microsatellite markers to other species [13,15]; therefore, we chose microsatellite markers developed for these two species to test their transferability to sickleweed.

None of the six *H. mantegazzianum* microsatellites...
markers amplified the *F. vulgaris* DNA; however, all six markers did amplify the DNA of *H. maximum* which was used as a positive control in this test. The transferability of these microsatellite markers (see [15] for primer sequences) from *H. mantegazzianum* to *H. maximum* has not been previously reported. With Daucus microsatellite primers, 26% of the primers amplified *Falcaria* DNA. Based on the quality of electrophoretic bands, Daucus primers tested for the amplification of *Falcaria* were classified: six primers producing a clean band with nearly expected amplicon size (7%), 16 primers producing multiple bands (19%) and 63 primers did not produce any band (74%). The six markers that produced clean bands are presented in Table 2. The identified six nuclear microsatellites and two chloroplast microsatellite primers were used for genotyping sickleweed samples collected from eight populations.

The signals of the genotyping results were clean and did not show stuttering bands. Micro-Checker showed no evidence of scoring error due to stuttering and large allele drop-out. The program did not identify any genotyeing error due to null alleles either. Our test of genotyping artifacts suggest that any deviation from Hardy-Weinberg equilibrium in our analysis is the result of change in allele frequency but not due to genotyping artifacts.

All six nuclear microsatellite loci were polymorphic with mean number of allele per locus of 8.8 (range 3 to 19) and most of these loci except ESSR80 have high polymorphism content value (Figure 1). Three out of six nuclear microsatellite loci were monomorphic for population from Iowa. This could be because of small sample size of the population. Two, three and five loci showed significant deviation from Hardy-Weinberg equilibrium for the populations from Iowa, Nebraska and South Dakota, respectively (Table 3).

Among six nuclear microsatellite loci, three loci (ESSR9, GSSR24 and GSSR25) produced three to four alleles for some samples (Figure 2). These multiple alleles per sample indicates that sickleweed in the novel range might have undergone gene duplications perhaps through polyploidyization [21].

Chloroplast DNA markers are often used for the study at higher taxonomic level as they have very slow evolutionary rate and do not reveal much variation within species [22]. However, chloroplast microsatellites have been effectively used for the study of intra-specific variation of the plant [11]. The two chloroplast microsatellites markers used in our study were also polymorphic and detected two (CSSR1) and three alleles (CSSR2). The average mean haploid diversity for these two loci was 0.23 and these two loci detected three chloroplast haplotypes. The number of alleles and the haploid diversity of these two loci in different populations are presented in Table 4.

These polymorphic microsatellite markers can be used for the study of population structure and gene flow in the introduced as well as native range and will ultimately be useful in the identification of source population(s). Use of non-recombinant chloroplast microsatellite markers and nuclear microsatellite markers will exactly determine if the gene flow is the result of seed or pollen flow [11]. The nuclear and chloroplast microsatellite markers together can provide important insights about the genetic

### Table 2. Characteristics of microsatellite markers for *F. vulgaris*.

<table>
<thead>
<tr>
<th>Locusa</th>
<th>Primer sequences (5'-3')</th>
<th>Repeat motif</th>
<th>Tm (°C)</th>
<th>Size range (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Nuclear microsatellites</strong></td>
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</tbody>
</table>
| ESSR9 (FAM) | F: ATCTGGGGAACTTGCTGTTG  
R: AGCATCAGCAGCAGCTACAA | (TGC)6 | 58 | 290 - 311 |
| ESSR80 (FAM) | F: ACAGCCAGTGAGCAGGACT  
R: GAGATTGTGCAATGTGGGAT | (CA)9 | 53 | 235 - 239 |
| GSSR24 (FAM) | F: GCCAACCATCAAATACACTTCT  
R: GAATACGCGCAATGCAATGCC | (TC)12 | 51 | 281 - 321 |
| GSSR25 (FAM) | F: CCAGAAACTGATTTTATTTTACGG  
R: CTGTTCATCAATAAACCTCTAACTC6 | (CAT)21 | 53 | 166 - 222 |
| GSSR154 (FAM) | F: CCTTATGTGATGCGCTCGAAA  
R: GACTGACCGCTCTCAATCTC | (TC)11 | 53 | 302 - 316 |
| BSSR53 (FAM) | F: GCTTATGAACTCTCTCTATCTCGTCA  
R: CTCACTGAGCTACTCTCCTACTCCTA | (AT)8 | 53 | 193 - 211 |
| **Chloroplast microsatellites** |
| CSSR1 (FAM) | F: GTTTAATGGAAGGAGCTTCTG  
R: TAT CCC CAA AAA GCC CAT T | (A)11 | 53 | 376 - 377 |
| CSSR2 (FAM) | F: CGG AAG TTT CAA TGG AAG GA  
R: TAA TTC CGG GGT TTC TCT GA | (T)11 | 53 | 177 - 179 |

aThe fluorescent dye used to label forward primer is given in parentheses.
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Table 3. Number of samples (n), Number of alleles (A), observed heterozygosity (H_o) and expected heterozygosity (H_e) of different nuclear microsatellite markers for the samples from Iowa, Nebraska and South Dakota.

<table>
<thead>
<tr>
<th>Locus</th>
<th>Iowa</th>
<th>Nebraska</th>
<th>South Dakota</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>A</td>
<td>H_o</td>
</tr>
<tr>
<td>ESSR9</td>
<td>12</td>
<td>3</td>
<td>na</td>
</tr>
<tr>
<td>ESSR80</td>
<td>12</td>
<td>1</td>
<td>na</td>
</tr>
<tr>
<td>GSSR24</td>
<td>12</td>
<td>3</td>
<td>1.00*</td>
</tr>
<tr>
<td>GSSR25</td>
<td>12</td>
<td>4</td>
<td>0.33</td>
</tr>
<tr>
<td>GSSR154</td>
<td>12</td>
<td>3</td>
<td>1.00*</td>
</tr>
<tr>
<td>BSSR53</td>
<td>12</td>
<td>1</td>
<td>na</td>
</tr>
</tbody>
</table>

*Departure from Hardy-Weinberg equilibrium at p < 0.05; na—Locus is monomorphic and test for H-W equilibrium was not done.

Table 4. Number of alleles and haploid diversity of two chloroplast microsatellite markers.

<table>
<thead>
<tr>
<th>Locus</th>
<th>Iowa</th>
<th>Nebraska</th>
<th>South Dakota</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>h</td>
<td>n</td>
</tr>
<tr>
<td>CSSR1</td>
<td>2</td>
<td>0.32</td>
<td>1</td>
</tr>
<tr>
<td>CSSR2</td>
<td>2</td>
<td>0.32</td>
<td>2</td>
</tr>
</tbody>
</table>

Figure 1. Polymorphism information content values displayed by the microsatellite loci.

Figure 2. Electropherograms of a sample at GSSR 25 locus. Similar multiple peaks were present at GSSR24 and ESSR9 loci. Multiple peaks displayed by several loci suggest that sickleweed in the United States might have undergone polyploidization.

from native and introduced range using these markers provide insights into sickleweed evolution [7] and invasion pathways [23] and may contribute in the risk assessment and effective management of this species in the United States.

Screening of more microsatellite markers particularly markers based on Expressed Sequence Tags (ESTs) from other species of Apiaceae family may reveal other microsatellites markers that may be useful for sickleweed because studies have shown that ESTs based microsatellite markers show greater transferability than anonymous microsatellites as the genes are highly conserved across different genera [24,25]. Similarly, there are several other universal chloroplast primers (see [11]) that can be sequenced to identify chloroplast microsatellite markers. These microsatellite markers are useful not only for the population genetic study of sickleweed but may be useful for the comparative study of species across Apiaceae family.

4. Conclusion

Our results demonstrated successful inter-generic transferability of microsatellite markers from *Daucus carota* to *Falcaria vulgaris*. Since these two species belong to two distantly related genera, transferability of microsatellite markers between these species indicates that these microsatellite markers may work for other genera within the Apiaceae family. We are also reporting two chloroplast microsatellite markers for sickleweed. Sequencing other chloroplast region using universal chloroplast markers may reveal more chloroplast microsatellite markers. These nuclear and chloroplast microsatellite markers are polymorphic and are useful for population genetics and phylogeographic study of sickleweed.

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