Polymorphic DNA Markers in Black Cherry (Prunus serotina) Are Identified Using Sequences from Sweet Cherry, Peach, and Sour Cherry

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ABSTRACT. Black cherry (Prunus serotina Ehrh.) is a common secondary forest species with a wide endemic distribution ranging from Nova Scotia south into Mexico, Ecuador, and Peru. Although planted in the United States for its valued lumber, black cherry is essentially a wild species with small fruit (≤6 to 10 mm in diameter). In contrast, in Mexico and Ecuador, domesticates of this species called Capulin, have much larger (2 to 2.5 cm in diameter) edible fruit. To date, no studies of the genetic diversity within North American black cherry or the ancestral origin of the Capulin types have been conducted. Simple sequence repeats (SSRs, also termed microsatellites) would be the marker of choice for such genetic diversity studies due to their hypervariability; however, generation of these sequence-based markers is expensive. Therefore, our objective was to determine if markers already identified in other Prunus L. species would be informative in black cherry. The black cherry germplasm screened consisted of selections originating from Michigan, Mexico, and Ecuador. A chloroplast DNA marker, originally generated from sour cherry (P. cerasus L.), amplified three different sized products in black cherry. Four of the eight nuclear SSR markers tested from peach [P. persica L. Batsch (Peach Group)], sour cherry, and sweet cherry (P. avium L.) also amplified and identified polymorphic markers. Together these four primer pairs resolved 54 putative alleles for the 66 black cherry accessions assayed. Success of the sweet cherry, peach, and sour cherry primers in identification of polymorphic markers in black cherry indicates it should be possible to use these markers for comprehensive molecular genetic studies in black cherry.

There are ≈400 species within the genus Prunus; however, only 25 Prunus species are native to North America (Maynard et al., 1991). Among these 25 species endemic to the New World, only black cherry (P. serotina) is grown in the United States for its high valued hardwood. Black cherry has a wide endemic distribution ranging from Nova Scotia, along the eastern United States coast, with populations extending into west Texas, southern Arizona, and New Mexico. This species is also endemic to central Mexico and into Ecuador and Peru (McVaugh, 1951).

In North America, black cherry is found commonly as a wild species in secondary forests (Maynard et al., 1991), thriving in recently disturbed areas such as hedgerows and logging sites. The fruits of these North American types are small (6 to 10 mm in diameter), nonfleshy, astringent, and have no commercial value (McVaugh, 1951). In Mexico and into the Andean Highlands, however, domesticates of black cherry called Capulin have much larger edible fruit (average 2 to 3.5 cm in diameter) with large pits (Popoeoe, 1924; Popoeoe and Pachano, 1922). While the Capulin cherries are not cultivated in large numbers, they occur commonly in home gardens and along roadsides and fences (Downey, 1999). The fruit is often harvested and sold as fresh fruit in markets in Mexico and the Andean region. It is hypothesized that the large fruited Capulin resulted from domestication and selection by native peoples in Central America (Popoeoe and Pachano, 1922).

Black cherry is a tetraploid (2n = 32) but it is not known if the species is an allotetraploid or autotetraploid and no progenitor species have been proposed. Geographic variation in black cherry has been studied using morphological characteristics (Carter et al., 1983); however, to date, no studies on the genetic diversity within North American black cherry or the ancestral origin of the Capulin types have been reported using molecular markers. The molecular markers of choice for genetic diversity studies are frequently simple sequence repeats (SSR, also termed microsatellites) (Litt and Ludy, 1989) due to their hypervariability, abundance, and relatively simple diagnostic polymerase chain reaction (PCR) procedure (Powell et al., 1996). SSR markers are a group of sequence tagged sites (STSs) where primers of specific sequence are designed to flank hypervariable regions of di-, tri- or tetra-nucleotide repeats (Litt and Ludy, 1989).

Widespread use of SSR markers is often hindered by the cost of identifying these markers since it involves library generation and sequencing. However, if primers designed to amplify loci in one species amplified loci in another species (termed heterologous amplification), the time and expense involved in isolating SSR markers for every species would be reduced. In Prunus, a limited number of SSR primer pairs are available from sweet cherry (P. avium) (C.D. Ryder, K. Edwards, and G.J. King, personal communication), sour cherry (P. cerasus) (A. Iezzoni, unpublished), and peach (P. persica (Peach Group)) (Gannavarapu, 1998).

The success of heterologous amplification using SSR primers depends generally upon the evolutionary distance between the original species and the tested species with decreasing success as genetic distance increases. In many mammalian species (Moore et al., 1991), bird species (Primmer et al. 1996), boids and cervids (Engel et al., 1996), heterologous amplification has been quite successful. This is also the case in plant species with SSRs conserved frequently across related species and sometimes across genera. For example, cross-species amplification was prevalent among three mustard (Brassica L.) species (Szewc-McFadden et al., 1996) and among 10 Actinidia Lindl. species (Weising et al., 1996). Plant taxa where SSR primers amplified loci across genera include the Poaceae (Rodler et al., 1995) and

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Citrinae subtribal group C (Kijas et al., 1995). In contrast, in the Asteraceae there was a lack of SSR conservation across the species tested (Whitton et al., 1997).

Genetic diversity and phylogenetic studies in plants frequently use chloroplast and nuclear markers due to their different modes of inheritance and rates of mutation. Chloroplast DNA (cpDNA) is inherited maternally in cherry (T. Brettin, personal communication) and is especially useful for phylogenetic studies due to its high degree of base sequence conservation (Curtis and Clegg, 1984; Palmer, 1987). Although cpDNA is conserved highly within a species, noncoding regions within the chloroplast genome have higher rates of mutation. Taberlet et al. (1991) identified PCR-based chloroplast polymorphisms by designing primer sets for conserved regions flanking noncoding variable regions of the chloroplast genome. When one of the primer sets, AB, designed by Taberlet et al. (1991), was used in sour cherry, seven length polymorphisms were identified (T. Brettin, personal communication). To better define the insertion/deletion events resulting in these seven polymorphisms, =300 bp of sequence was obtained from each of the seven polymorphic ‘AB’ fragments (T. Brettin, personal communication). The first 300 bp of sequence revealed two polymorphisms which differed for two insertion/deletions. A primer pair spanning these two insertions/deletions was designed and termed the nested AB primers, [AB], to facilitate rapid screening of sour cherry germplasm for the length polymorphism. This nested cpDNA primer pair, [AB], was chosen as a chloroplast marker for use in this study.

The objective of this study was to examine the degree of conservation of a chloroplast marker derived from sour cherry and SSR loci developed from peach, sour cherry, and sweet cherry with a geographically diverse set of SSR loci developed from peach, sour cherry, and sweet cherry. The sweet cherry and peach primers were obtained from G. King or peach were used. The sour cherry primers were obtained from a ‘Erdi Botermo’ genetic library (Iezzoni, unpublished).

Materials and Methods

Black cherry seedlings were collected from Mexico, Ecuador, and Michigan (Table 1). Accessions from central Mexico were collected as open-pollinated half-sib seed from under Capulin trees in August 1996 and therefore represented five open-pollinated seed families (RG, PH, P2, P3, and T1). Two of these families (RG and T1) were collected by collaborators of S. Perez (Queretaro, Mexico). Ten Ecuadorian accessions were obtained from the Ecuadorian Germplasm Institute as open-pollinated seed presumably collected from five trees and were therefore termed five open-pollinated families [Ecu A 1-3, D 1-2, E 1, F 1-2, and H 1-2 (Raul Castillo, Departamento Nacional de Recursos Fitogenetics y Biotecholigia, Quito, Ecuador)]. Ten black cherry plants were obtained as young seedlings from the Michigan State University (MSU) Department of Forestry, East Lansing. These seedlings originated from bulk seed of the black cherry collection at the Kellogg Biological Station, Hickory Corners, Mich. One black cherry accession currently growing on the MSU Campus (Beal) was also included.

One to 10 seedlings per family were scored for the chloroplast fragment and SSR loci (Table 1). DNA was extracted from young leaves following procedures of Stockinger et al. (1996). Amplification of the cpDNA fragment was done using two PCR reactions. The first PCR reaction used the ‘AB’ primers and amplification conditions described by Taberlet et al. (1991). The PCR product from this reaction was diluted 1:100 and used as template in a second PCR reaction with the [AB] nested primers (Table 2). The amplification products were separated on a 6% polyacrylamide gel run at 80 V for 2 h and stained using Silver Sequence staining system (Promega, Madison, Wis.). Amplified fragments were sized using a 10-bp ladder (Gibco BRL, Rockville, Md.).

Eight SSR primer pairs isolated from sour cherry, sweet cherry, or peach were used. The sour cherry primers were obtained from a P. cerasus ‘Erdi Botermo’ genetic library (Iezzoni, unpublished). The sweet cherry and peach primers were obtained from G. King (Wellsbourne, United Kingdom) and A. Abbott (Clemson, S.C.), respectively. Optimization of PCR conditions for each primer pair was done using a temperature gradient PCR thermocycler (Robocycler) (Stratagene, La Jolla, Calif.). The products were separated and stained as described previously. Sequences and annealing temperatures for the successful primers are listed in Table 2.

Pairwise similarity values were calculated using the Nei-Li similarity index (Nei and Li, 1979) and the equation: F = 2 Nxy/(Nx

| Country of origin | Family designation | No. of seedlings tested/family | cpDNA marker | SSR loci | cpDNA length polymorphism (bp) |
|-------------------|--------------------|-------------------------------|--------------|---------|-------------------------------|
| Mexico            | RG                 | 3                             | 10           |         | 250 274 280                   |
|                   | PH                 | 3                             | 10           |         | 3                             |
|                   | P2                 | 3                             | 10           |         | 3                             |
|                   | P3                 | 3                             | 10           |         | 3                             |
|                   | T1                 | 3                             | 10           |         | 3                             |
| Ecuador           | Ecu A              | 1                             | 3            |         | 1                             |
|                   | Ecu D              | 1                             | 2            |         | 1                             |
|                   | Ecu E              | 1                             | 1            |         | 1                             |
|                   | Ecu F              | 1                             | 2            |         | 1                             |
|                   | Ecu H              | 1                             | 2            |         | 1                             |
| Michigan          | MI                 | 3                             | 5            |         | 3                             |
|                   | Beal               | 1                             | 1            |         | 1                             |

Taberlet et al. (1991) identified PCR-based chloroplast polymorphisms for the successful primers are listed in Table 2.
alleles with 19 in total, ranging from 170 to 230 bp in length (Table 2). All putative alleles were shared among all three geographical groups. Only one putative allele was exclusively in Michigan selections. Only four putative alleles were specific to any one particular geographical group. One putative allele was specific to the Ecuadorian group, two were specific to the Mexican group, and one was specific to the Michigan group. Additionally, with all four primer pairs, no putative alleles were found to be shared exclusively between the Michigan and Ecuadorian selections (Fig. 1). Any putative alleles identified in both the Michigan and Ecuadorian selections were also always present within the Mexican selections. Additionally, with sour cherry, sweet cherry, and peach selections (PceGA34, PS12A02, and pchgs2, respectively), the selections from Ecuador were more similar to the Mexican selections than those from Michigan, and the selections from Ecuador and Michigan were the least similar (Table 3). The exception to this pattern is the peach derived primer pair pchgs2, where all 19 putative alleles were identified in the Mexican selections and no novel putative alleles were identified in the selections from Michigan.

A maximum of four putative alleles were amplified for the selections assayed, representing the four alleles in the tetraploid black cherry. None of the putative alleles were specific to the Michigan or Ecuadorian groups. However, five putative alleles were only identified in the Mexican group (176, 196, 214, 226, and 228 bp).

When all four heterologous primer pairs were considered separately, the data obtained from the black cherry selections were generally similar. For example, with all four primer pairs, no putative alleles were identified between the Michigan and Ecuadorian selections (Fig. 1). Any putative alleles identified in both the Michigan and Ecuadorian selections were also always present within the Mexican selections. Additionally, with sour cherry, sweet cherry, and peach selections (PceGA34, PS12A02, and pchgs2, respectively), the selections from Ecuador were more similar to the Mexican selections than those from Michigan, and the selections from Ecuador and Michigan were the least similar (Table 3). The exception to this pattern is the peach derived primer pair pchgs2, where all 19 putative alleles were identified in the Mexican selections and no novel putative alleles were identified in the selections from Mexico and Michigan (Fig. 1).

**Discussion**

One chloroplast primer pair and four out of eight nuclear SSR primer pairs from sweet cherry, peach, and sour cherry exhibited heterologous amplification in black cherry and identified polymorphisms in the black cherry germplasm collection. These results suggest that DNA markers for use in genetic diversity and evolutionary studies in black cherry can be identified using some of the primers developed from peach, sweet cherry, and sour cherry.

The nested chloroplast primers [AB], designed from sour cherry sequence identified three different length fragments within the black cherry germplasm evaluated. Insertion and deletion events that give rise to length variants have been reported to account for a substantial fraction of intraspecific cpDNA variation (Zurawski and Clegg, 1987). Interestingly, two of the three fragments amplified in black cherry (274 and 250 bp) are similar in length to those identified in a French sweet cherry selection (276 bp) and in sweet, ground (P. 

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**Table 2. Summary of primer pairs derived from peach and sweet and sour cherry that identified size polymorphisms when tested on black cherry selections from Michigan, Mexico, and Ecuador.**

| Species source | Target DNA | Primer | Primer sequence (5'–3') | Annealing temp (°C) | Expected product size (bp) | Size range of products (bp) | No. of size polymorphisms/primer pair |
|---------------|------------|--------|-------------------------|---------------------|---------------------------|-----------------------------|--------------------------------------|
| Sour cherry Chloroplast [AB] | GCTGGGAACCGTGAATTCA GGGCCTATCTAAAGTAAAGA CATTTGTGGTCTGCTGTT TCCACTAGGAGGTCAGAAATG | 56 | 249 | 250–280 | 3 |
| Sour cherry Nuclear PceGA34 | GCCACCAATAGTCTTCTCC AGGCCAGATGCACTGTGA AGCCTATGCGTCACTCAG | 60 | 155 | 140–174 | 14 |
| Sweet cherry Nuclear PS12A02 | GCCACCAATAGTCTTCTCC AGGCCAGATGCACTGTGA AGCCTATGCGTCACTCAG | 56 | 200 | 150–178 | 12 |
| Peach Nuclear pchgs2 | ACGCTATGCGTCACTCAG | 60 | 179 | 170–230 | 19 |
| Peach Nuclear pchgs2 | CAACCTGTGATTGCTCCTATTAAAC GTCAAATGCGTTGGTCCGTACACTC AATCATAACATCATTCAGCCACTGC | 60 | 163 | 130–152 | 9 |

+ Nxy, where Nx and Ny are the total number of fragments identified in geographic groups x and y, respectively, and Nxy is the number of fragments shared by the two geographic groups.
fruticosa Pall.), and sour cherry selections (249 bp) (T. Brettin, personal communication). Amplification of the most prevalent 274 bp fragment in families from Michigan, Mexico, and Ecuador suggest that germplasm from these three regions share a common chloroplast type. However, since this primer pair also identified three chloroplast polymorphisms, it may be a useful marker for future analyses of the ancestral polyploid origin of black cherry.

Since cpDNA is inherited maternally in Prunus, it was expected that all the open-pollinated seedlings from each family would have the same chloroplast fragment. However, seedlings from the Mexican family RG exhibited different fragments (250 and 280 bp). For the RG collection family, open-pollinated seed collected by our collaborator was assumed to be derived from only one maternal parent tree. However, the cpDNA data suggest that multiple maternal parents were sampled for this family.

The 280-bp fragment exhibited by the Ecuadorian accession, Ecu D, may have resulted because Ecuadorian accessions were probably derived from multiple maternal parents as well. Morphologically, the RG and Ecu D families were indistinguishable from other collected families and subsequent assays with SSRs showed no further deviation of these accessions from the rest of the Capulin accessions. Therefore, cpDNA divergence is most likely due to collection of open-pollinated seed from multiple maternal trees and not due to sampling of other Prunus species.

It is also theoretically possible that a somatic chloroplast length mutation could have arisen in the maternal plant and that this mutation could have resulted in open-pollinated progeny with different chloroplast types (Hagemann, 1992). In the course of cell divisions, plastids segregate at random. This can result in somatic segregation and sorting out of genetically different plastids. Plant sectors that differ in their chloroplast type would then pass on different chloroplast types to their progeny. However, it is more likely that the chloroplast difference between progeny believed to be from the same maternal parent is due to sampling from different maternal parents since it is a much simpler explanation.

The success in using heterologous PCR primers to amplify SSR loci in black cherry decreases the need to develop new sets of primers and therefore provides opportunities for population and evolutionary genetic studies in black cherry. Even within the Michigan black cherry germplasm, the four SSR primers derived from sweet and sour cherry and peach amplified 29 putative alleles.
suggesting that these SSRs could be used to conduct future population genetic studies within North American black cherry germplasm. Although only eight nuclear primer pairs were tested, the data suggest that within Prunus, the usefulness of a primer pair may depend more upon the primers themselves than the species origin of the primers. For example, the most useful primer pair was from peach, pchgm3. This primer pair amplified the maximum number of putative alleles compared to the other primer pairs tested and even identified tetra-allelic individuals. Identification of four putative alleles per progeny individual with one primer pair indicates that the primer sequences are conserved for all four alleles. In contrast, three other peach primer pairs tested either failed to amplify any bands or only amplified one monomorphic band. Of the two sweet cherry primer pairs tested, one amplified just one monomorphic band and the second amplified twelve putative alleles and the tetra-allelic individuals. It is possible that genetic distance and the usefulness of SSR primers may become more important when amplification is attempted across genera.

Results obtained from each of the four polymorphic primer pairs were similar. For example, the selections from Mexico exhibited the most putative alleles. With the exception of primer pair pchgm2, the Michigan selections had more putative alleles than the Ecuadorian selections. With the exception of peach primer pair pchgm3, selections from Mexico and Ecuador were more similar than selections from Mexico and Michigan, and the Ecuador and Michigan selections were the least similar. Also, with all four primer pairs no putative alleles were found to be shared by the Michigan and Ecuadorian selections that were not also present in the Mexican selections. This last observation suggests that the Ecuadorian and Michigan germplasm may have diverged from a central Mexican population presumably as a result of ecological adaptation and possibly domestication. These trends in the data identified by the four polymorphic primer pairs will be useful for testing hypotheses concerning the evolution of black cherry and the domestication of the Capulin types.

**Table 3. Similarity values (Nei and Li, 1979) for pairwise comparisons of the black cherry selections from North America, Mexico, and Ecuador using four informative primer pairs.**

| Origin of selections used in the pairwise comparison | Primer pairs |
|----------------------------------------------------|--------------|
|                                                     | PceGA34      |
|                                                     | PS12A02      |
|                                                     | pchgm3       |
|                                                     | pchgm2       |
| Mexico and Ecuador                                  | 0.74         |
| Mexico and Michigan                                 | 0.38         |
| Ecuador and Michigan                                | 0.15         |
| Mexico and Ecuador                                  | 0.66         |
| Mexico and Michigan                                 | 0.59         |
| Ecuador and Michigan                                | 0.15         |
| Mexico and Ecuador                                  | 0.64         |
| Mexico and Michigan                                 | 0.77         |
| Ecuador and Michigan                                | 0.66         |
| Mexico and Ecuador                                  | 0.66         |
| Mexico and Michigan                                 | 0.40         |
| Ecuador and Michigan                                | 0.00         |

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