

## PERMANENT GENETIC RESOURCES

Isolation and characterization of microsatellite markers for Carolina hemlock (*Tsuga caroliniana*)

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## Abstract

We describe the isolation and characterization of 31 polymorphic di- and trinucleotide microsatellite marker loci for Carolina hemlock (*Tsuga caroliniana* Englem.). In addition, primer pairs for 16 loci amplified scoreable alleles in six other *Tsuga* species. In eastern North America, both Carolina hemlock and eastern hemlock (*Tsuga canadensis* [L.] Carr.) populations are declining due to infestation by hemlock woolly adelgid, *Adelges tsugae*. The markers described here should enhance population genetic studies of hemlocks, providing valuable information for conserving and restoring these important forest tree species.

**Keywords:** conservation genetics, hemlock, SSR markers

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Carolina hemlock is endemic to the southern Appalachian Mountains of eastern North America. It is a wind-pollinated species with a restricted range confined to isolated rocky slopes and ridges (Farjon 1990). In recent years, this rare and vulnerable species, which has a G3 global conservation priority rank ([www.natureserve.org/explorer/](http://www.natureserve.org/explorer/)), has begun to experience mortality due to infestation by the hemlock woolly adelgid (HWA), *Adelges tsugae* Annand, an exotic insect that has caused a rapid decline of eastern hemlock across much of its range (McClure *et al.* 2003).

Microsatellite markers were developed from fresh leaf tissue DNA collected from a single *Tsuga caroliniana* tree (tree #247, Linville Falls, North Carolina). Enrichment, cloning and sequencing were performed (Genetic Identification Services) as previously described (Josserand *et al.* 2006), with the exception that the clone libraries were enriched for AC, GA and ATG simple sequence repeats (SSR). From each library, 53 recombinant colonies were selected at random and their cloned fragments were sequenced. Relatively long SSRs that were centrally located in the sequence were identified for 135 of these inserts: 48, 47 and 40 from the AC, GA and ATG libraries, respectively. Polymerase chain reaction (PCR) primer pairs for 99 of these inserts were selected using DesignerPCR version 1.03 (Research Genetics) and synthesized (Integrated DNA Technologies) for further evaluation. All forward primers were tailed on their 5' end with the

M13forward(-29) sequence, CACGACGTTGTAACGAC, allowing fluorescent dye (6-FAM, VIC, NED or PET, Applied Biosystems) incorporation in PCR when using a dye-labelled M13forward(-29) primer. All reverse primers were tailed on their 5' end with a PIG tail, GTTTCCTT (Brownstein *et al.* 1996), forcing nontemplated dA additions to the amplified fragments.

Synthesized primer pairs were screened for polymorphism against 23 *T. caroliniana* samples collected from seven typical hemlock sites distributed across the mountainous area of North Carolina and adjacent South Carolina. In addition, a subset of 63 markers that cleanly amplified in *T. caroliniana* was evaluated for use in six additional *Tsuga* species: eastern hemlock (*T. canadensis*),  $n = 31$ ; Chinese hemlock (*T. chinensis*),  $n = 5$ ; mountain hemlock (*T. mertensiana*),  $n = 3$ ; northern Japanese hemlock (*T. diversifolia*),  $n = 3$ ; southern Japanese hemlock (*T. sieboldii*),  $n = 2$ ; and western hemlock (*T. heterophylla*),  $n = 4$ . All DNA samples were isolated from fresh leaf samples using DNeasy 96 Plant Kit (QIAGEN). PCR and allele separation were performed as previously described (Josserand *et al.* 2006), with the exception that a hot-start monoclonal antibody (GeneCraft, GmbH) was used in the PCR. An internal LIZ 600 size standard was run with each sample and alleles were sized using the local southern algorithm and binned with GeneMapper 3.7 software (Applied Biosystems). Analyses of genotype frequencies, genotypic disequilibrium and exact tests for  $P$  values were conducted with GenePop 3.4 (<http://genepop.curtin.edu.au/>).

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Of the 99 primer pairs tested with *T. caroliniana* samples, 34 amplified consistent allelic profiles and of these, 31 were polymorphic (Table 1). Among the polymorphic loci, the number of alleles per locus ranged from two to seven

(average = 3.4), with observed and expected heterozygosities ranging from 0.043 to 1.0 (average = 0.208) and from 0.043 to 0.768 (average = 0.475), respectively. Tests of Hardy–Weinberg equilibrium found that 23 loci were not in equi-

**Table 1** Results for 31 microsatellite loci isolated from *Tsuga caroliniana* and tested on a sample of 23 *T. caroliniana* trees

Locus name	Source library	GenBank Accession	Observed repeat motif	Primer sequences (5'–3')	Number of alleles: range in size (bp)	$H_O$	$H_E$	HWE (P value)	Significant LD with TcSI locus
TcSI_002	AC	BV726470	(AC) <sub>23</sub>	F: GCATTAGGATAATCATTCAGAG R: AGAGGGAAAAATAACCACTGAG	2: 233–235	0.043 <sup>xs</sup>	0.122	0.002	
TcSI_004	AC	BV726471	(AC) <sub>27</sub>	F: GCCAGATTTTGTCTAATCC R: TTGGTGTCTTATCTAATCAGTG	5: 270–283	0.174 <sup>xs</sup>	0.414	0.068	080
TcSI_006	AC	BV726538	(GA) <sub>11</sub>	F: ATCCCCCAATTAGCCATG R: AGGGCGGATCAATGTTTG	3: 113–161	1.000	0.619	0.000*	
TcSI_007	AC	BV726472	(CA) <sub>16</sub> (AT) <sub>8</sub>	F: CAAGGCAACAACATATCTTTAGC R: CAACTTTCTTCCAACCTCTGT	2: 261–265	1.000	0.500	0.000*	
TcSI_008	AC	BV726473	(AC) <sub>16</sub> (AT) <sub>9</sub>	F: ACGGCTTTTCATTTGATG R: CATTCCGTAACATTTGATGC	3: 119–155	1.000	0.619	0.000*	
TcSI_009	GA	BV726474	(TC) <sub>30</sub>	F: TGGTTACAAGGTTCTGACAGTC R: CTACCCAATGCTAAGGTGATT	6: 250–272	0.087 <sup>xs</sup>	0.438	0.000*	
TcSI_012	GA	BV726476	(TC) <sub>12</sub>	F: CCATTTGCTAGACATGCAATAC R: TTTCAGATGGACAGACGTAATG	2: 249–251	0.087	0.159	0.030	
TcSI_014	GA	BV726478	(TC) <sub>18</sub> (CA) <sub>8</sub>	F: CCAGATGGAAATGGTAGGTG R: AATGGGTGAGAGATAATGTGAG	2: 167–169	0.130 <sup>xs</sup>	0.364	0.002	040, 076
TcSI_022	AC	BV726483	(AC) <sub>19</sub>	F: CCAAATTCAGAATGTCTATCC R: TTGAAGGAATAGTGGTTGATG	3: 221–225	0.136 <sup>xs</sup>	0.501	0.000*	
TcSI_027	AC	BV726487	(TG) <sub>15</sub>	F: AAATCCGTCAGTGTGCTAC R: CTCCAGTCCACATAAACCTTC	3: 180–184	0.043 <sup>xs</sup>	0.417	0.000*	078
TcSI_030	AC	BV726489	(AC) <sub>13</sub>	F: GCTGCATACTGTTTACCCTAAA R: TAGGCTTGTGTTCCCTCTGC	2: 201–203	0.043 <sup>xs</sup>	0.405	0.000*	
TcSI_032	AC	BV726490	(AT) <sub>10</sub> (TG) <sub>6</sub> (TG) <sub>9</sub>	F: TACGGACCATCGGCTATG R: ATGATTATGCCCCGTCAAT	2: 185–187	0.217 <sup>xs</sup>	0.405	0.026	
TcSI_040	GA	BV726493	(AG) <sub>27</sub>	F: CAATCCCAAAGAACAAGTC R: GGAGGAAGGACACGAGAAG	5: 113–123	0.087 <sup>xs</sup>	0.740	0.000*	
TcSI_044	GA	BV726495	(TC) <sub>7</sub> (CT) <sub>7</sub>	F: TCTTCTCTTCTACTGCCTCTAA R: AGGATAAGTGTGTTTCTGAG	5: 150–158	0.087 <sup>xs</sup>	0.400	0.000*	
TcSI_055	AC	BV726502	(CA) <sub>20</sub> (AT) <sub>6</sub>	F: TTCCTCAACCCATTATACAATC R: TCAAGGATGAACCATTAGAATC	2: 268–270	0.087 <sup>xs</sup>	0.386	0.000*	
TcSI_057	AC	BV726504	(AC) <sub>13</sub>	F: TAACCCATCCCTTTACAAGAG R: TAGCCGTGTGGTCATAGTTAG	2: 135–137	0.130 <sup>xs</sup>	0.405	0.001*	066, 075, 079
TcSI_060	AC	BV726507	(AC) <sub>20</sub>	F: CAACAATCATGTCACTCCATA R: GTCGTGCGAATTTGTCTC	3: 99–103	0.087 <sup>xs</sup>	0.322	0.001*	
TcSI_061	AC	BV726508	(AC) <sub>24</sub>	F: TGTGCTTGGGAATTATCTTAAC R: CTTCAAGACCCTGCCTAAAG	2: 136–138	0.043	0.043	0.915	
TcSI_062	AC	BV726509	(AT) <sub>5</sub> (TG) <sub>32</sub>	F: ACGCCAACAACTTCCCTTT R: GAGGGTATGGAGATGFC	6: 268–299	0.136 <sup>xs</sup>	0.689	0.000*	066, 080
TcSI_066	AC	BV726513	(TG) <sub>5</sub> (GT) <sub>27</sub>	F: TGCACCATGCAATGAAAC R: GCCCATCAAATGAAATG	2: 217–218	0.043 <sup>xs</sup>	0.485	0.000*	075, 080
TcSI_067	AC	BV726514	(AC) <sub>16</sub>	F: AATATGGGATGTGGTAACTGTG R: CTCGTGTGTTTGTGTTATGGT	2: 242–244	0.043 <sup>xs</sup>	0.315	0.000*	
TcSI_074	GA	BV726519	(GA) <sub>17</sub>	F: CAAAGATTGGAGACGAACTTG R: CCTTACAGAAATACCCATCAC	5: 169–179	0.217 <sup>xs</sup>	0.584	0.002	076
TcSI_075	GA	BV726520	(GA) <sub>20</sub>	F: CTCGTGAGCATGATGTGTAAC R: ATAGTTCCACTGCAACCTCATA	3: 174–188	0.130 <sup>xs</sup>	0.559	0.000*	080, 089
TcSI_076	GA	BV726521	(GA) <sub>30</sub>	F: AGAGCACACGAGAAATAGATTG R: TATTCCAACCATAGGTCTCAAC	7: 158–176	0.217 <sup>xs</sup>	0.768	0.000*	
TcSI_078	GA	BV726522	(CT) <sub>20</sub>	F: AACCCATACAAAGGATGCC R: GAGAGGTAGGAAGGATGTTG	5: 235–243	0.217 <sup>xs</sup>	0.719	0.000*	
TcSI_079	GA	BV726523	(AG) <sub>16</sub>	F: CCCAACCAACCTTTTGTGTA R: AGCACACTCTCACACTTCACTC	3: 290–294	0.130 <sup>xs</sup>	0.523	0.000*	089

Table 1 Continued

Locus name	Source library	GenBank Accession	Observed repeat motif	Primer sequences (5'–3')	Number of alleles: range in size (bp)	$H_O$	$H_E$	HWE ( $P$ value)	Significant LD with TcSI locus
TcSI_080	GA	BV726524	(CT) <sub>25</sub>	F: TCCTCCTCATTTCACACTCAC R: CCAAATCCAACATTCCTAGTT	4: 224–230	0.087 <sup>xs</sup>	0.717	0.000*	087
TcSI_083	GA	BV726527	(AG) <sub>25</sub>	F: GTGCTCTGAGGTGTGTGTC R: TTGGATGTTTACGATGATGG	3: 234–239	1.000	0.572	0.000*	
TcSI_085	GA	BV726528	(TC) <sub>20</sub>	F: TCITCCCTTCTCCTCATAGGTC R: GATTTTCTGTGCATGTGTGTTTC	3: 280–284	0.174 <sup>xs</sup>	0.436	0.014	
TcSI_087	GA	BV726530	(GA) <sub>23</sub>	F: TTTGTTCTTGGAACTTGACATC R: CTTTTATGCAGCAAATTCACCTC	4: 279–285	0.217 <sup>xs</sup>	0.606	0.001*	
TcSI_089	GA	BV726532	(TC) <sub>19</sub>	F: GGGAAATGTGCCTTAGTTTAA R: GGAAAGGAGAAAACACACAG	3: 115–134	0.130 <sup>xs</sup>	0.517	0.000*	

Repeat structure and primer sequences and observed number of alleles, allele size range (base pairs, bp), heterozygosity values (observed,  $H_O$  and expected,  $H_E$ ), and  $P$  values for exact tests of Hardy–Weinberg equilibrium (HWE), and TcSI locus number exhibiting significant ( $P < 0.05$ ) genotypic disequilibrium in paired tests (LD). Loci with  $H_O$  values marked with <sup>xs</sup> had an excess of homozygotes, as determined by Micro-Checker analysis (van Oosterhout *et al.* 2004).  $P$  values marked with \* are significant following sequential Bonferroni correction for  $P < 0.05$  (Holm 1979).

Table 2 Marker transferability results for 16 *Tsuga caroliniana*-derived markers tested with six *Tsuga* species of various sample sizes ( $n$ ), observed number of alleles and allele size range in base pairs

Locus	<i>T. canadensis</i> $n = 31$	<i>T. chinensis</i> $n = 5$	<i>T. mertensiana</i> $n = 3$	<i>T. diversifolia</i> $n = 3$	<i>T. sieboldii</i> $n = 2$	<i>T. heterophylla</i> $n = 4$
TcSI_004	—	6: 235–291	1: 304	2: 230–338	2: 254–316	3: 226–232
TcSI_008	—	—	3: 103–115	1: 146	—	6: 106–144
TcSI_012	11: 243–268	6: 230–259	2: 239–241	2: 209–239	1: 241	3: 239–247
TcSI_014	—	5: 164–195	—	—	—	—
TcSI_029	3: 119–125	5: 105–123	2: 109–121	2: 103–105	1: 108	3: 107–113
TcSI_030	—	—	1: 201	—	—	—
TcSI_040	10: 91–121	2: 80–85	—	2: 129–131	1: 117	6: 95–115
TcSI_052	3: 207–230	1: 219	1: 225	1: 218	1: 222	—
TcSI_060	—	4: 113–129	1: 109	—	2: 111–124	1: 109
TcSI_061	—	—	—	—	—	2: 104–108
TcSI_067	—	—	—	—	—	3: 245–252
TcSI_071	—	—	2: 112–114	—	—	—
TcSI_076	—	6: 126–192	—	—	2: 149–182	—
TcSI_080	6: 216–232	5: 209–234	—	—	—	6: 221–239
TcSI_083	8: 247–265	—	—	—	—	—
TcSI_089	—	6: 117–142	—	2: 134–140	1: 128	—

—, no amplification or difficult to interpret amplification products. The following loci (GenBank Accession) were monomorphic in *T. caroliniana*: TcSI\_029 (BV726488), TcSI\_052 (BV726500) and TcSI\_071 (BV726517).

librium ( $P < 0.05$ , Table 1). Tests for two-locus genotypic disequilibrium found 16 significant locus pairs out of 435 tests after sequential Bonferroni correction for  $P < 0.05$  (Holm 1979). Tests for null alleles, using the Dunn-Sidak correction for Monte Carlo simulations in the program Micro-Checker (van Oosterhout *et al.* 2004), found 25 loci that have an excess of homozygotes (Table 1). Their mean estimated proportion of null alleles across all null estimators was 0.35 with a 95% confidence interval of  $\pm 0.03$ , indicating little variation in the proportion of excess homozygosity among the loci. The

results of these three tests together are consistent with inbreeding within geographically isolated stands (Farjon 1990) and do not support the presence of widespread linkage disequilibrium or high null allele frequencies.

Microsatellite primers developed from some conifer species can be amplified in related taxa (Echt *et al.* 1999; Chagné *et al.* 2004; Josserand *et al.* 2006). Of the 34 primer pairs that cleanly amplified in *T. caroliniana*, 16 (47%) amplified in the other six species (Table 2), with the greatest proportion being transferable to *T. chinensis* (29%, 10 markers) and

the least to *T. canadensis* (18%, 6 markers). Of the three markers that were monomorphic for *T. caroliniana*, TcSI\_029, TcSI\_052 and TcSI\_071, each successfully amplified polymorphic loci in at least one other *Tsuga* species. TcSI\_029 appeared polymorphic in five other species and TcSI\_052 and TcSI\_071 appeared polymorphic in one other species each (Table 2).

Clearly the microsatellite markers described in this paper will enhance population genetic studies of Carolina hemlock and other hemlock species, as well as assist inter- and intraspecies breeding programmes (Bentz *et al.* 2002; Pooler *et al.* 2002) working to develop HWA resistant populations for species conservation and restoration.

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