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PRIMER NOTE

## ISOLATION AND CHARACTERIZATION OF 22 EST-SSR MARKERS FOR THE GENUS *THUJOPSIS* (CUPRESSACEAE)<sup>1</sup>

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- Premise of the study: Expressed sequence tag-simple sequence repeat (EST-SSR) markers were developed from Thujopsis
  dolabrata var. hondae (Cupressaceae) using Illumina sequencing to investigate the genetic diversity and population structure
  of the genus Thujopsis.
- *Methods and Results:* Twenty-two primer pairs were developed from ESTs of *T. dolabrata* var. *hondae*. The primers amplified di- and trinucleotide repeat-containing sequences. Polymorphisms were assessed in 81 individuals from two *T. dolabrata* var. *hondae* populations and one *T. dolabrata* population. The number of alleles ranged from one to 17 per locus. The observed and expected heterozygosities ranged from 0.000 to 1.000 and from 0.000 to 0.926, respectively.
- Conclusions: These new EST-SSR markers will be useful in analyses of the genetic diversity and population structure of the genus *Thujopsis*.

**Key words:** Cupressaceae; expressed sequence tag; microsatellite; next-generation sequencing; *Thujopsis*.

The Cupressaceae clade has the broadest diversity of habitats and morphologies of any conifer family, and the genus Thujopsis, which belongs to the Cupressaceae, is considered to be one of the early diverging genera (Pittermann et al., 2012). Thujopsis is native to Japan and includes one species (T. dolabrata Siebold & Zucc.) and one northern variety (T. dolabrata var. hondae Makino) (Hayashi, 1960). The two varieties of Thujopsis are distinguished by variation in their cone morphology. The varieties also differ in their geographic ranges, although their distributions overlap in the central region of the Japanese archipelago (Hayashi, 1960). *Thujopsis dolabrata* has antimicrobial properties and demonstrates a strong antifeedant effect on termites (Inamori et al., 2006). Moreover, its essential oil is used as an antibacterial agent and as an aromatic substance. Thus, T. dolabrata is an important tree species in Japanese forestry and future forest tree breeding because of its superior wood properties. However, there have been few reports about genetic differences between the two varieties.

Recently, expressed sequence tag (EST)–based markers have been used increasingly in studies of genetic diversity and population structure in tree species (e.g., Fageria and Rajora, 2013). EST-based markers are less susceptible to null alleles than anonymous simple sequence repeats (SSRs). Moreover, because ESTs correspond to coding DNA, the flanking sequences

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of EST-SSRs are located in well-conserved regions across phylogenetically related species (Uchiyama et al., 2013). Nuclear SSR markers have been developed for *T. dolabrata* var. *hondae* (Mishima et al., 2012), and population genetics and phylogeographic analyses of this variety were performed using these markers (Higuchi et al., 2012). However, the nuclear SSR markers have not been tested on *T. dolabrata*. An analysis of the genetic diversity and population structure for *Thujopsis* plants in the Japanese archipelago is urgently required. Therefore, it is necessary to obtain molecular markers with high transferability within the genus *Thujopsis* that also exhibit a low frequency of null alleles. In this paper, we describe the development and characterization of 22 EST-SSR markers for the genus *Thujopsis* from expressed sequence data of *T. dolabrata* var. *hondae*.

### METHODS AND RESULTS

One T. dolabrata var. hondae individual (voucher no. TF-K12-001) was used for the RNA sequencing experiment. Leaves and cambiums were sampled from a population in the Aomori Prefecture (Owani: 40°27′21″N, 140°34′08″E) and were immediately frozen in liquid nitrogen and stored at -30°C. The cetyltrimethylammonium bromide (CTAB) method was used to extract the total RNAs (Chang et al., 1993). A TruSeq RNA Sample Prep Kit (Illumina, San Diego, California, USA) was used to create the RNA sequencing library, according to the manufacturer's protocol. A HiSeq 1000 (Illumina) was used to sequence the library with 2 × 101-bp paired-end reads. More than 237.24 million paired-end raw reads were obtained. After removal of low-quality reads, 233.92 million clean reads remained (accession no.: DRA002435). Using the short reads assembly programs Velvet 1.2.08 (Zerbino et al., 2009) and Oases 0.2.08 (Schulz et al., 2012), the clean reads were assembled de novo into 76,377 contigs and 41,182 unigenes, from 100 to 14,834 bp, with a mean length of 1525 bp. MSATCOMMANDER 1.0.8 (Faircloth, 2008) was used to screen for microsatellite loci containing di- and trinucleotides. Primer3 (Rozen and Skaletsky, 1999) was then used to design PCR primers. The minimum number of repeats was set as nine and 11 for di- and trinucleotide repeats, respectively. All

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TABLE 1. Characteristics of 22 EST-SSR loci for use in the genus Thujopsis.

						DDBJ acc	ession no.
Locus		Primer sequences (5′–3′)	Repeat motif	$T_{\rm a}(^{\circ}{\rm C})$	Allele size range (bp)	1 <sup>a</sup>	2 <sup>b</sup>
Tdest1	F:	GCCTCCCTCGCGCCATCAGGATTTTCTGACAGGCTTTGTTCTC	(CT) <sub>11</sub>	60	137–173	LC010288	LC010310
	R:	GTTTCTTAATTCCCAAGAGTGCTTATGAGTTC					
Tdest3		GCCTCCCTCGCGCCATCAGCGGCCCAGGTTTCTGTACTC	$(AT)_{11}$	60	169–186	LC010289	LC010311
m1 .11		GTTTCTTGCCCATTAAAGTCGGGTATTG	( A TT)	60	120 161	1 6010200	1 0010212
Tdest11		GCCTCCCTCGCGCCATCAGTGGGATACATACTGCATTTGTTAGG	$(AT)_{12}$	60	138–161	LC010290	LC010312
Tdest14		GTTTCTTCTCCCCAAGCAAGTCACCAC	(AG) <sub>12</sub>	60	157–185	LC010291	LC010313
Tuest14		GCCTCCCTCGCGCCATCAGCAGTAGACAATTTCTGCAAATCACC GTTTCTTTCCCTTTTGTTGGCATTATAGG	$(AG)_{12}$	00	137-163	LC010291	LC010313
Tdest17		GCCTCCCTCGCGCCATCAGGCTTTTGATGTCCGCTATATCCTC	$(AG)_{12}$	60	159–168	LC010292	LC010314
raestr,		GTTTCTTGGAGATTCCAATGTTTGTCATGC	(110)12	00	137 100	EC0102)2	Leorosii
Tdest21		GCCTCCCTCGCGCCATCAGGTCCATCCATTCTCACTCCAAAG	$(AG)_{13}$	60	231-274	LC010293	LC010315
	R:	GTTTCTTAGCAGACCCTATTTCACAGCATC	· /13				
Tdest24	F:	GCCTCCCTCGCGCCATCAGATACCATACAGCTTTCAGCCAG	$(AT)_{15}$	60	243-267	LC010294	LC010316
	R:	GTTTCTTGCAGAACAAACGAATCAATGAGAG					
Tdest29 F		GCCTCCCTCGCGCCATCAGAAACGACTCTGCTGGATTTCAC	$(AC)_{16}$	60	219–246	LC010295	LC010317
		GTTTCTTTTCCGCTCTTGATTTTCTCTCC					
Tdest35		GCCTCCCTCGCGCCATCAGAAGCTATTGACCCTTCTCAGGATAC	$(CT)_{15}$	60	194–230	LC010296	LC010318
m1 .07		GTTTCTTCCATGTTGAATTGTTCCCTTTC	(AFFC)	60	164 176	1 6010207	1 0010210
Tdest37		GCCTCCCTCGCGCCATCAGCCAAGCGACAGAAAACCATTC	$(ATC)_9$	60	164–176	LC010297	LC010319
Tdest38		GTTTCTTTCAGTCTCTTCCTCCTCCTCCTC	(ACC)	60	117 124	LC010298	LC010320
Tuestoo		GCCTCCCTCGCGCCATCAGTGACCATTCCTCCTCCTCCTC GTTTCTTCATGTTTGCAGTTGAGAGAGACC	$(ACC)_9$	00	117–134	LC010298	LC010320
Tdest39		GCCTCCCTCGCGCCATCAGGCAGCACAGGAGAAGAAGATG	(GCT) <sub>o</sub>	60	153–165	LC010299	LC010321
Tuesta		GTTTCTTACAACAGCCACAACGTGTCC	(001)9	00	133-103	LC010277	LC010321
Tdest42		GCCTCCCTCGCGCCATCAGCTCCCTATCCCAACACCAACAC	(ACC) <sub>9</sub>	60	226–255	LC010300	LC010322
1 4050.12		GTTTCTTTGCCTACCTATCCTTCTTCTCC	(1100)9		220 200	20010000	20010022
Tdest43		GCCTCCCTCGCGCCATCAGGGTCCAATGCAGGTAATACAAGAAG	(CGG) <sub>9</sub>	60	137-153	LC010301	LC010323
	R:	GTTTCTTTCCCCGCCAAGATACTCAAC					
Tdest44	F:	GCCTCCCTCGCGCCATCAGTTTGGTGGTGGAGGTGGTG	$(GAT)_9$	60	134-137	LC010302	LC010324
	R:	GTTTCTTCGCTTATGCCAAGCAGTCATC					
Tdest45		GCCTCCCTCGCGCCATCAGTGAGGGTGGTGAGACAATTC	$(GGT)_{12}$	60	211–236	LC010303	LC010325
		GTTTCTTCAAGATTTGGAACTCCTGCAAC					
Tdest49		GCCTCCCTCGCGCCATCAGGTGCCCTCAAAGTTACAGCAGTC	$(GAT)_{10}$	60	233–248	LC010304	LC010326
TI 150		GTTTCTTGCAATCACCTCATCCTCACTTC	(CCT)	60	220, 251	1.0010205	I CO10227
Tdest52		GCCTCCCTCGCCCATCAGTTCAGGAAGGCCAAGGAGAG	$(GGT)_{11}$	60	239–251	LC010305	LC010327
Tdest53		GTTTCTTGATCCTCCTGCATCATTTTGTTC GCCTCCCTCGCGCCATCAGCCAAAGCCCTTCCAGTAACATC	(CTT) <sub>13</sub>	60	244–284	LC010306	LC010328
Tuesiss		GTTTCTTGATGGAATGAGTGAATCTCAGGAAC	(C11) <sub>13</sub>	00	244-204	LC010300	LC010326
Tdest54		GCCTCCCTCGCGCCATCAGCCCTGTATTATTCTCAACATCATCG	(CTT) <sub>11</sub>	60	182-203	LC010307	LC010329
1 000to 1		GTTTCTTGGGATTCAGACAAGGGCAAG	(011)11	50	102 200	20010007	2001002)
Tdest56		GCCTCCCTCGCGCCATCAGCATTGCCCTTTGGAATATAGGATC	(AAG) <sub>9</sub>	60	153-165	LC010308	LC010330
		GTTTCTTGTTGCCCATCTGCTCTTCTTC	` //				
Tdest58	F:	GCCTCCCTCGCGCCATCAGCTGAACGGCGCCCTAATCTC	$(AAG)_{13}$	60	151-180	LC010309	LC010331
	R:	GTTTCTTGCCCACTCCTCAAATCCAAC					

*Note*: DDBJ = DNA Data Bank of Japan;  $T_a$  = annealing temperature.

forward primers were fluorescently labeled using FAM (carboxyfluorescein) with the 454A adapter sequence (5'-GCCTCCCTCGCGCCATCAG-3') at the 5'-end (Blacket et al., 2012). Additionally, all reverse primers were attached to a PIG-tail sequence (5'-GTTTCTT-3') at the 5'-end of the sequence (Brownstein et al., 1996).

The CTAB method (Murray and Thompson, 1980) was used to extract genomic DNAs from needles sampled from 32 T. dolabrata plants in the Nagano Prefecture (Kiso: 35°43′38″N, 137°37′15″E) and 49 T. dolabrata var. hondae needles were sampled from two populations in the Aomori Prefecture (Owani: 40°27′21″N, 140°34′08″E; and Okoppe: 41°28′42″N, 140°57′10″E). These populations were located in the typical range of each variety. To confirm PCR amplification, we used eight DNA samples from four individuals per variety. PCR was performed in a final volume of 10  $\mu$ L, containing 5  $\mu$ L of 2× GoTaq Hot Start Colorless Master Mix (Promega Corporation, Madison, Wisconsin, USA), 1  $\mu$ M of each primer, and 60 ng of template DNA. Reactions were performed with initial denaturation at 94°C for 2 min; followed by 40 cycles of 94°C for 30 s, 60°C for 30 s, and 72°C for 30 s; and

then 72°C for 5 min using an ABI 9700 (Applied Biosystems, Foster City, California, USA). The PCR products were separated electrophoretically on 1% agarose gels and stained with GelRed Nucleic Acid (Nacalai Tesque, Kyoto, Japan). To confirm the presence of SSRs, we sequenced the PCR products from two DNA samples using an ABI PRISM 3100 Genetic Analyzer (Applied Biosystems). For fragment analysis, the PCR conditions were modified. The concentration of the primers were 0.15, 0.5, and 0.2  $\mu M$  for the forward primer, reverse primer, and 454A-FAM primer, respectively, and the number of PCR cycles was 30. PCR products for 81 DNA samples from two varieties were electrophoresed on an ABI PRISM 3100 Genetic Analyzer (Applied Biosystems) and Geneious 7.0.4 software (Biomatters Ltd., Auckland, New Zealand; http://www.geneious.com/) was used to assess the fragment sizes. To characterize the microsatellite loci, CERVUS 3.0 software (Kalinowski et al., 2007) was used to calculate the number of alleles (A), the observed heterozygosity  $(H_0)$ , the expected heterozygosity  $(H_e)$ , and the frequency of null alleles (r). GenAlEx 6.501 (Peakall and Smouse, 2012) assessed the probability of identity  $(P_{ID})$ .

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a 1 = T. dolabrata var. hondae.

b2 = T. dolabrata.

Table 2. The results of final primer screening on samples from three populations of the genus *Thujopsis*.

		Thujopsis dolabrata var. hondae										Thujopsis dolabrata					
		Okoppe $(N = 27)$					Owani (N = 22)				Kiso (N = 32)						
Locus	$\overline{A}$	$H_{\rm o}$	$H_{\rm e}$	$P_{ m ID}$	r	$\overline{A}$	$H_{\rm o}$	$H_{\rm e}$	$P_{ m ID}$	r	$\overline{A}$	$H_{\rm o}$	$H_{\rm e}$	$P_{ m ID}$	r	Total A	
Tdest1	12	1.000	0.897	0.026	-0.066	11	0.955	0.883	0.034	-0.052	12	0.938	0.898	0.024	-0.030	16	
Tdest3	6	0.444	0.484	0.305	-0.002	2	0.409	0.333	0.508	-0.113	9	0.813	0.830	0.056	-0.001	10	
Tdest11	12	0.963	0.860	0.040	-0.071	9	0.682	0.809	0.070	0.086	9	0.719	0.810	0.069	0.056	14	
Tdest14	11	0.852	0.885	0.031	0.009	12	0.864	0.884	0.032	0.001	11	0.875	0.880	0.032	-0.005	14	
Tdest17	8	0.778	0.747	0.102	-0.028	8	0.545	0.678	0.152	0.100	7	0.625	0.579	0.229	-0.073	8	
Tdest21	17	0.852	0.903	0.023	0.023	11	0.818	0.770	0.080	-0.060	15	0.969	0.926	0.015	-0.031	21	
Tdest24	8	0.852	0.832	0.058	-0.024	9	0.727	0.846	0.050	0.064	6	0.719	0.740	0.116	0.008	10	
Tdest29	5	0.593	0.622	0.193	0.013	5	0.545	0.540	0.278	-0.008	1	0.000	0.000	1.000	ND	6	
Tdest35	17	0.852	0.922	0.016	0.035	15	0.909	0.900	0.025	-0.024	13	0.813	0.850	0.044	0.010	23	
Tdest37	4	0.333	0.372	0.438	0.062	4	0.409	0.411	0.398	-0.029	3	0.281	0.294	0.539	0.009	5	
Tdest38	3	0.593	0.639	0.215	0.015	5	0.727	0.743	0.125	-0.001	4	0.625	0.615	0.234	-0.016	6	
Tdest39	4	0.815	0.613	0.226	-0.159	3	0.682	0.654	0.206	-0.027	2	0.125	0.119	0.786	-0.023	4	
Tdest42	6	0.519	0.558	0.277	0.011	5	0.636	0.630	0.210	-0.030	7	0.563	0.652	0.173	0.053	10	
Tdest43	6	0.778	0.806	0.077	0.011	8	0.909	0.768	0.097	-0.100	4	0.406	0.455	0.342	0.080	8	
Tdest44	1	0.000	0.000	1.000	ND	1	0.000	0.000	1.000	ND	2	0.031	0.031	0.940	-0.002	2	
Tdest45	7	0.704	0.643	0.206	-0.058	4	0.545	0.579	0.249	0.024	2	0.094	0.091	0.833	-0.015	7	
Tdest49	2	0.148	0.140	0.754	-0.030	3	0.091	0.090	0.834	-0.014	4	0.563	0.495	0.319	-0.082	5	
Tdest52	2	0.926	0.507	0.376	-0.301	2	1.000	0.512	0.375	-0.333	2	0.281	0.246	0.604	-0.072	2	
Tdest53	12	0.926	0.897	0.025	-0.025	10	0.818	0.885	0.033	0.030	9	0.719	0.783	0.080	0.039	14	
Tdest54	3	0.370	0.545	0.325	0.169	3	0.455	0.563	0.274	0.078	4	0.094	0.632	0.212	0.743	5	
Tdest56	3	0.778	0.645	0.212	-0.104	4	0.500	0.506	0.293	-0.014	4	0.531	0.563	0.257	0.018	4	
Tdest58	5	0.519	0.524	0.320	-0.005	5	0.364	0.499	0.307	0.129	3	0.281	0.298	0.528	0.011	9	

Note: A = number of alleles;  $H_e = \text{expected heterozygosity}$ ;  $H_o = \text{observed heterozygosity}$ ; N = sample size; ND = not determined;  $P_{\text{ID}} = \text{probability of identity}$ ; r = null allele frequency.

Primer3 designed 58 primer pairs, of which 33 showed amplification for all eight samples in both varieties. For 29 of the 33 primer pairs, the SSR sequence in the PCR product was identified for both varieties. Finally, 22 of the 29 amplified primers showed clear fragment patterns, thus they were selected as the developed markers. These primer pairs show different fragment sizes and the same annealing temperature (Table 1). All 22 loci were polymorphic in both varieties. The observed number of alleles per population ranged from one to 17,  $H_0$  ranged from 0.000 to 1.000, and  $H_e$  ranged from 0.000 to 0.926. The  $P_{\rm ID}$  ranged from 0.015 to 1.000 (Table 2). One of the loci (Tdest54) showed high r values relative to the other loci. These 22 EST-SSR markers had lower r and higher  $P_{\rm ID}$  values compared with reported SSR markers developed from the genomic DNA of T. dolabrata var. hondae (Mishima et al., 2012)

#### **CONCLUSIONS**

In this study, we developed 22 EST-SSR markers for the two varieties of *Thujopsis*, using expressed sequence data of *T. dolabrata* var. *hondae*. These markers have two advantages: high ability to detect genetic polymorphisms in *Thujopsis* varieties, and a low null allele frequency. Accordingly, these EST-SSR markers will be valuable tools for investigating the genetic diversity and population structure of the genus *Thujopsis*. Moreover, these markers will help to advance breeding programs for species in the genus *Thujopsis*.

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