Development of microsatellite markers for the Japanese endemic conifer Thuja standishii and transfer to other East Asian species

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Abstract

Objective
Design polymorphic microsatellite loci that will be useful for studies of the genetic diversity, structure and reproduction in the Japanese endemic conifer Thuja standishii and test the transferability of these loci to the two other East Asian species, T. sutchuenensis and T. koraiensis.

Results
Fifteen loci were developed which displayed 3 to 21 alleles per locus (average = 9.2) among 97 samples from three populations of T. standishii. Observed heterozygosity for all samples varied between 0.33-0.75 (average 0.54) while expected heterozygosity values were higher with an average over the 15 loci of 0.62 (0.37-0.91). Low multi-locus probability of identity values (< 0.00001) indicate that these markers will be effective for identifying individuals derived from clonal reproduction. All 15 loci amplified in T. sutchuenensis, the sister species of T. standishii, with 1 to 11 alleles per locus (average = 4.33) while 13 loci amplified in the more distantly related T. koraiensis with 1 to 5 alleles per locus (average = 2.15).

Introduction
Thuja is a small Cupressaceae genus consisting of five extant species with three species in East Asia and two in North America (LePage 2003). Thuja standishii (Gordon) Carriere (kurobe or nezuko in Japanese and Japanese arbor-vitae in English) is endemic to Japan where it has a scattered distribution from 40.67° N to 33.49° N on the islands of Honshu, Shikoku and Dogo (Shimane Prefecture). The
species is most commonly a component of subalpine forests but can occur in a variety of habitats including warm temperate evergreen forest (rarely), cool temperate forest, moorland and near the alpine zone. The species can be a large tree to 35 m in height to a multi-stemmed shrub under 1 m tall at the limits of its elevational range (Worth et al. 2019). *Thuja standishii* can purportedly reach a great age with trunks up to 3.5 m in diameter (Giant Tree Database, Biodiversity Center of Japan (http://kyoju.biodic.go.jp)) and individuals over 1000 years old.

Unlike most other Cupressaceae conifers of Japan, *T. standishii* has received little research attention with basic information on its conservation (including the impact of past logging), reproductive biology, and genetic diversity lacking. The rarity of forest dominated by *T. standishii* and its current insignificant role in forestry probably underlies this lack of research. However, the species is undoubtedly an important part of Japan’s biodiversity and cultural heritage being one of the five precious timber trees (Kiso go-boku) that from the early 18th century were strictly protected from cutting in the Kiso region of central Honshu (Mertz 2011) and, in some parts of Japan, forests containing *T. standishii* are considered to be some of the most untouched forests remaining (e.g. Wada 1994). Small population size and geographic isolation, its vulnerability to ring barking by deer and the impacts of past logging (Worth et al. 2019) has resulted in some populations being of conservation concern, especially in western Japan where the species is very rare (Worth et al. 2019). One key aspect of the species biology that is poorly understood is the role of asexual reproduction in its regeneration. However, similar to two other Japanese Cupressaceae conifers, *Thujopsis dolabrata* and *C. pisifera* that have been proven to regenerate clonally (Hayakawa et al. 2004; Hasegawa et al. 2015), *T. standishii* also forms dense understory banks of juveniles (Worth personal...
observation) which may be clonally derived.

This study describes the development of Expressed Sequence Tagged (EST) nuclear microsatellite markers for *T. standishii* using next generation sequencing that will be useful for investigating the species range-wide genetic diversity, gene flow and reproductive biology. In addition, we tested the transferability of the developed markers in the Chinese restricted endemic *Thuja sutchuenensis*, the sister species of *T. standishii* (Li and Xiang 2005; Peng and Wang 2008), and the only other East Asian species, *T. koraiensis*, for which no microsatellite markers have yet been developed.

Materials and methods

**Materials and Methods**

Total RNA was extracted from an individual of *T. standishii* collected from the Forestry and Forest Products Research Institute Arboretum using a plant RNA isolation mini kit (Agilent Technologies, USA). An RNA-seq data set was constructed by the Beijing Genomics Institute on an Illumina HiSeq 4000 platform. The *T. standishii* RNA-seq data consisted of 38,076,160 paired-end reads of 100 bp length. **De novo** assembly was undertaken in CLC Genomics Workbench 8.5.1 and the 53,614 resultant contigs (*N*50 = 1,503 bp) were mined for microsatellite regions. Primers were developed bordering these regions with default settings using PrimerPro (http://webdocs.cs.ualberta.ca/~yifeng/primerpro/). Microsatellites were selected if the number of tandem repeat units was greater than eight and if the microsatellite was located less than 25 bp from the beginning or end of the contig. These criteria resulted in 64 microsatellite primer pairs which were trialled for amplification in four samples. A total of 36 primer pairs successfully amplified and
were subsequently tested for size heterogeneity in eight samples representative of the species range. For all loci, the forward primer was synthesized with one of three different M13 sequences (5’ GCCTCCCTCGCGCA 3’, 5’ GCCTTGCCAGCCGC 3’, and 5’ CAGGACCAGGGCTACGTG 3’), and the reverse was tagged with pig-tail (5’ GTTTCTT 3’; (Brownstein et al., 1996)). The PCR reactions were performed following the standard protocol of the Qiagen Multiplex PCR Kit (Qiagen, Hilden, Germany), and consisted of a 10 uL reaction volume, containing approximately 5 ng of DNA, 5 uL of 2x Multiplex PCR Master Mix, and 0.06 uM of forward primer, 0.1 uM of reverse primer, and 0.08 uM of fluorescently labelled M13 primer. The PCR thermocycle consisted of an initial denaturation at 95°C for 3 min; followed by 35 cycles of 95°C for 30 sec, 60°C for 3 min, 68°C for 1 min; and a 20 min extension at 68°C. The PCR products were separated by capillary electrophoresis on an ABI3130 Genetic Analyzer (Life Technologies, Waltham, MA, USA) with the GeneScan 600 LIZ Size Standard (Life Technologies) and genotyping was done in GeneMarker (SoftGenetics, LLC, PA, USA). Overall, 15 loci were found to amplify reliably, display polymorphism and were readily scorable. The genetic variability of these 15 markers were tested in three populations from Atebi Daira Small Bird Forest, Mt Chausu Nature Park, in Nagano Prefecture (35.2286° N, 137.6673° E), Mt Torigata in Kouchi Prefecture (33.4936° N, 133.0638° E) and Mt Yamizo in Fukushima Prefecture (36.9343° N, 140.2679° E). The 15 primer pairs were also tested in 13 samples of T. sutchuenensis and four of T. koraeensis (Table S1). Genetic analyses were undertaken in GenAlEx 6.5 (Peakall and Smouse, 2006) and Genepop 4.2 (Raymond and Rousset 1995). In addition, a similarity search of the contigs containing the 15 loci was conducted by the BLASTX algorithm (Altschul et al. 1990) against the National Center for Biotechnology Information (NCBI) non-redundant protein
sequences (nr) database.

The multi locus probability of identity (PID) for the 15 markers, that is, the probability that two individuals drawn at random from a population will have the same genotype (Waits, et al. 2001), was calculated in Gimlet version 1.3.3 (Valiere 2002). Three PID estimates outlined by Waits et al. (2001) were estimated: biased PID which assumes individuals mate randomly; unbiased PID which corrects for sampling a small number of individuals and, sibs PID which assumes the population is composed of siblings.

Results

In *T. standishii*, the 15 loci (Table 1) displayed 3 to 21 alleles over the three populations with an average of 9.2 alleles per locus (Table 2). Overall observed heterozygosity varied between 0.33–0.75 (average = 0.54) while expected heterozygosity values were generally higher (0.37–0.91 with an average of 0.62). At the population level, the number of alleles observed per locus varied from 2–15 (average = 5.43) with six loci showing more than four alleles in each of the three populations. No significant deviations from Hardy–Weinberg equilibrium expectations were detected for any loci except for Kurobe_4219 in the Atebi Daira and Mt Yamizo populations (P = < 0.0004). Additionally, allele frequencies appeared independent among loci with no significant linkage disequilibrium detected after Bonferroni correction. Multi-locus probability of identity values across all 15 loci were below the threshold value of 0.01 considered by Waits et al. (2001) to be required to reliably distinguish between individual genotypes, even under the sibs PID (Table S2). This indicates that our markers will be effective for both identifying individuals derived from clonal
reproduction and sexually derived individuals in populations even where inbreeding is prevalent.

All 15 loci amplified in *T. sutchuenensis*, two loci being monomorphic, with 1 to 11 alleles per locus (average = 4.33) while observed and expected heterozygosity were 0.43 and 0.48, respectively (Table 3). On the other hand, only 13 loci amplified in *T. koraiensis* with three loci being monomorphic. In this species, 1 to 5 alleles per locus (average = 2.15) were observed with observed and expected heterozygosity of 0.44 and 0.31, respectively (Table 3).

**Discussion**

The development of EST microsatellites for *Thuja standishii* will enable new genetic research into this important Japanese endemic conifer including studies of range-wide level genetic diversity and gene flow and also stand-level processes including inbreeding and clonality. The development of molecular markers may help to foster research into this species, which because of its wide ecological range, from warm temperate forests to near the alpine zone, is an ideal species to investigate ecological and genetic processes under strongly contrasting climates.

The transferability of the 15 loci was consistent with the phylogenetic relationships of the East Asian *Thuja* (Li and Xiang 2005; Peng and Wang 2008). Thus, all 15 loci successfully amplified in the sister species of *T. standishii, T. sutchuenensis* and displayed considerable allelic diversity with up to 11 alleles per locus. These loci, therefore, may be particularly applicable for use genetic studies of this geographically restricted endangered species (Yang et al. 2013). In contrast, two of the fifteen loci did not amplify in the more distantly related *T. koraiensis* and the number of alleles per locus (ranging from 1 to 5 alleles) was low although this low
allelic diversity could be due to the low number of samples tested.

Limitations

The number of published microsatellite markers may be too low for optimal performance of some genetic analyses. These microsatellite loci have not been tested in the two North American species, T. plicata and T. occidentalis. We did not afford much time optimizing loci, therefore some polymorphic loci that may have worked with further effort may have been excluded.

Abbreviations

bp: base pair; EST: expressed sequence tag; RNA: ribonucleic acid.

Declarations

Authors’ contributions

JRPW, KC, YH and AQ devised the study and did the sample collection. JRPW designed the primers, undertook fragment analysis and genotyping. JRPW did the data analysis and wrote the manuscript. All authors participated in the draft and read and approved the final manuscript.

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Competing interests

The authors declare that they have no competing interests.

Availability of data and materials

The 38,076,160 paired-end reads are deposited in the NCBI BioProject Database, Accession number: PRJNA554973. The genotype data for every individual are available on demand to the corresponding author.
Consent for publication

Not applicable

Ethics approval and consent to participate

Not applicable

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### Table 1 Characteristics of the 15 microsatellite markers developed for *Thuja standishii*

| Locus         | Primer (5′-3′) | Repeat motif | Allele size range (bp) | Bl                      |
|---------------|----------------|--------------|------------------------|-------------------------|
| Kurobe_18480  | F: TCAGGACCCCAACAAATAAA | (AT)9        | 135-189                | unknown                 |
|               | R: GATGCTGGAGATGGTATTTCG  |              |                        |                         |
| Kurobe_2969   | F: ACACAATCTTGCCCAGAAGC | (TA)8        | 269-291                | WRI2 [L]                |
|               | R: ACACAATCTTGCCCAGAAGC  |              |                        |                         |
| Kurobe_23700  | F: CCGACGCCTCTACATTCATT | (TCT)10      | 250-268                | hypothetical protein    |
|               | R: ATGGAAAAGAGGCAGAGCAA |              |                        | [Marchantia]             |
| Kurobe_44557  | F: GATCCCTTTTGATGGGTTCA | (TC)11       | 313-349                |                         |
|               | R: TTATCCCCAACCTCAGCGAG  |              |                        |                         |
| Kurobe_15129  | F: TTAACATCCACAAATGGCAA | (TC)12       | 205-233                | unknown                 |
|               | R: CAAATGCATGGAATGTGGTC  |              |                        |                         |
| Kurobe_16758  | F: TGCAACGTCAAAACACGAT   | (CT)8        | 272-283                | CYP866B′                 |
|               | R: CACCCCTCTAGAGCGTGGTC  |              |                        |                         |
| Kurobe_6943   | F: CTACGTCTTAGCCTCCACGC  | (CT)11       | 159-169                | early no tric           |
|               | R: ATGCCATACTGGGTTCCTGAG |              |                        |                         |
| Kurobe_23263  | F: CCAGGCCCTTAAATTTCCACTC | (AG)10      | 290-314                |                         |
|               | R: CCATGGCACAATGGGATACAGA |              |                        |                         |
| Kurobe_51603  | F: CAAAGCAACAGAGCCAACAA  | (TG)13       | 162-201                |                         |
|               | R: AAATAGGGTGCTGAGGTTG   |              |                        |                         |
| Kurobe_31302  | F: TCCCAAGTCATGGGCCAGT   | (GT)8        | 264-274                |                         |
|               | R: GTCACTCGGAGCAATTTT     |              |                        |                         |
| Kurobe_38308  | F: TTTGCTGAAACCCAGGAATC  | (TG)11       | 272-297                |                         |
| Microsatellite | Primer Combination | Repeat Unit | Size (bp) | Description |
|---------------|--------------------|-------------|-----------|-------------|
| Kurobe_41636 | F: TTGAACCTTTGCAGTGGGAAA | (CA)8 | 157-165 |  |
|               | R: GAACCAGACAGGAATCGGAA |       |         |  |
| Kurobe_42400 | F: TTCACACAAGGGCCTACTCC | (TC)9 | 254-272 |  |
|               | R: GCCGACAGGTCTCTTAGCAC |       |         |  |
| Kurobe_40825 | F: TTATCCCATGAGCGTGCTTT | (CT)8 | 232-276 |  |
|               | R: TAGGTGCGGTGACTATGCAG |       |         |  |
| Kurobe_4219 | F: ACGCATCGGACAGTCGTATT | (TC)8 | 276-306 | MACP At4 |
|               | R: CCGTGCGAGAACAAAGAAAT |       |         |  |

- denotes that no BlastX hits were found

Table 2  Genetic diversity of the 15 polymorphic nuclear microsatellites assessed across three populations of *Thuja standishii*.
| Locus          | Atebi Daira (32) | Mt Torigata (31) | Mt Yamizo (34) | All (97) |
|----------------|------------------|------------------|----------------|----------|
|                | Na   | Ho   | He  | Na   | Ho   | He  | Na   | Ho   | He  | Na   | Ho   | He  | Na   | Ho   | He  |
| Kurobe_18480  | 13   | 0.75 | 0.77 | 11   | 0.81 | 0.82 | 8    | 0.38 | 0.43 | 21   | 0.64 |
| Kurobe_2969   | 6    | 0.53 | 0.45 | 5    | 0.52 | 0.60 | 7    | 0.85 | 0.77 | 9    | 0.64 |
| Kurobe_23700  | 5    | 0.78 | 0.77 | 5    | 0.42 | 0.38 | 6    | 0.59 | 0.57 | 7    | 0.60 |
| Kurobe_44557  | 14   | 0.68 | 0.89 | 9    | 0.68 | 0.73 | 15   | 0.88 | 0.88 | 21   | 0.75 |
| Kurobe_15129  | 7    | 0.66 | 0.71 | 2    | 0.03 | 0.03 | 5    | 0.68 | 0.69 | 8    | 0.46 |
| Kurobe_16758  | 4    | 0.50 | 0.51 | 3    | 0.29 | 0.26 | 4    | 0.85 | 0.73 | 7    | 0.56 |
| Kurobe_6943   | 3    | 0.28 | 0.30 | 3    | 0.65 | 0.51 | 4    | 0.35 | 0.33 | 7    | 0.42 |
| Kurobe_23263  | 2    | 0.41 | 0.46 | 3    | 0.48 | 0.44 | 3    | 0.38 | 0.51 | 4    | 0.42 |
| Kurobe_51603  | 7    | 0.75 | 0.75 | 3    | 0.55 | 0.54 | 8    | 0.82 | 0.70 | 9    | 0.71 |
| Kurobe_31302  | 3    | 0.63 | 0.55 | 3    | 0.52 | 0.51 | 2    | 0.12 | 0.11 | 3    | 0.41 |
| Kurobe_38308  | 6    | 0.59 | 0.66 | 3    | 0.55 | 0.45 | 5    | 0.50 | 0.58 | 9    | 0.55 |
| Kurobe_41636  | 2    | 0.41 | 0.36 | 2    | 0.39 | 0.46 | 3    | 0.21 | 0.21 | 4    | 0.33 |
| Kurobe_42400  | 7    | 0.50 | 0.48 | 5    | 0.58 | 0.53 | 6    | 0.38 | 0.36 | 9    | 0.48 |
| Kurobe_40825  | 7    | 0.50 | 0.63 | 6    | 0.71 | 0.71 | 5    | 0.68 | 0.58 | 11   | 0.63 |
| Kurobe_4219   | 4    | 0.28 | 0.51 | 6    | 0.61 | 0.57 | 6    | 0.62 | 0.70 | 9    | 0.51 |
| Average       | 6.00 | 0.55 | 0.59 | 4.60 | 0.52 | 0.50 | 5.80 | 0.55 | 0.54 | 9.20 | 0.54 |

*Note: Na = the number of alleles, Ho = observed heterozygosity and He = expected heterozygosity.*
Table 3. Genetic diversity of the 15 microsatellite loci in *T. sutchuenensis* and *T. koraiensis*.

| Locus          | *T. sutchuenensis* (13) | *T. koraiensis* (4) |
|----------------|-------------------------|---------------------|
|                | Na | Ho | He | Na | Ho | He  |
| Kurobe_18480   | 7  | 0.46 | 0.80 | 1  | 0.00 | 0.00 |
| Kurobe_2969    | 7  | 0.61 | 0.79 | 2  | 1.00 | 0.50 |
| Kurobe_23700   | 3  | 0.15 | 0.14 | 2  | 0.25 | 0.22 |
| Kurobe_44557   | 11 | 0.61 | 0.89 | 3  | 0.50 | 0.41 |
| Kurobe_15129   | 6  | 0.69 | 0.76 | 2  | 0.25 | 0.22 |
| Kurobe_16758   | 4  | 0.69 | 0.56 | -  | -    | -    |
| Kurobe_6943    | 1  | 0.00 | 0.00 | -  | -    | -    |
| Kurobe_23263   | 3  | 0.38 | 0.33 | 2  | 0.50 | 0.50 |
| Kurobe_51603   | 4  | 0.38 | 0.38 | 5  | 1.00 | 0.75 |
| Kurobe_31302   | 1  | 0.00 | 0.00 | 2  | 0.75 | 0.47 |
| Kurobe_38308   | 6  | 0.61 | 0.69 | 2  | 0.25 | 0.22 |
| Kurobe_41636   | 2  | 0.46 | 0.43 | 2  | 0.25 | 0.22 |
| Kurobe_42400   | 2  | 0.31 | 0.26 | 1  | 0.00 | 0.00 |
| Kurobe_40825   | 6  | 0.69 | 0.80 | 1  | 0.00 | 0.00 |
| Kurobe_4219    | 2  | 0.38 | 0.39 | 3  | 1.00 | 0.59 |
| Average        | 4.33 | 0.43 | 0.48 | 2.15 | 0.44 | 0.31 |

Denotes that the loci did not amplify

**Supplementary Files**

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