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# Development of 32 EST-SSR markers for Abies firma (Pinaceae) and their transferability to related species ${ }^{1}$ 

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- Premise of the study: We developed simple sequence repeat (SSR) markers from expressed sequence tags (ESTs) for Abies firma, a conifer endemic in Japan, to facilitate evaluation of the population genetic structure in this species.
- Methods and Results: We designed primers for 153 EST-SSRs identified from 486322 ESTs from A. sachalinensis ESTs, and tested 96 of them for PCR amplification. Thirty-two primers provided clear amplification, and 14 of those 32 displayed clear polymorphic patterns in multiple populations of A. firma and in two closely related species. The number of alleles per locus and mean expected heterozygosity ranged from one to six and 0 to 0.476 , respectively.
- Conclusions: The EST-SSR markers developed in this study may be useful for phylogeography and population genetic studies of A. firma. Successful amplifications were obtained for two other Abies species, suggesting that these markers may also be useful for similar applications in other fir species.

Key words: Abies; cross-amplification; expressed sequence tag; microsatellite; Pinaceae; pyrosequencing.

In the family Pinaceae, Abies is the genus with the second highest number of species. Approximately 40 species are widely distributed in the northern hemisphere in regions ranging from temperate to subarctic zones. Four of the five species that grow in the Japanese archipelago are endemic to Japan. Abies firma Siebold \& Zucc. is a major tree species occurring only in warmtemperate forests in Japan. This species is frequently found in mixed forest along with species such as Tsuga sieboldii Carrière and Fagus crenata Blume, but it sporadically forms pure stands at the late succession stage (Farjon, 1990). In recent years, the area covered by A. firma forest has been significantly reduced by logging and exploitation. Moreover, since the early 1960s, forest decline and tree dieback in A. firma forests in many areas of Japan have been observed as a consequence of environmental stress factors such as air pollution (Suzuki, 1992). For effective genetic conservation of these forests, it is necessary to understand the phylogeographic pattern and the genetic diversity within and among A. firma populations. Population genetic studies to date have relied on allozyme markers (Saito et al., 2005) and mitochondrial DNA markers (Tsumura and Suyama, 1998), and have not made use of microsatellites.

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Microsatellite markers are recognized as versatile molecular tools for inferring genetic structure and gene flow. In recent years, expressed sequence tag (EST)-based markers have been increasingly used in studies of genetic variation because large numbers of polymorphic markers can be developed with relative ease using EST data and markers of this type are less susceptible to null alleles than are anonymous simple sequence repeats (SSRs). Moreover, because ESTs correspond to coding DNA, the flanking sequences of EST-SSRs are located in wellconserved regions across phylogenetically related species, making them markers of choice for comparative mapping and relevant functional and positional candidate genes to study their colocation with quantitative trait loci. In the work described here, we developed EST-SSR markers for A. firma from published expressed sequence data, and evaluated the extent of the polymorphism that they exhibit and their potential for transfer to two other closely related Japanese Abies species (A. homolepis Siebold \& Zucc. and A. veitchii Lindl.).

## METHODS AND RESULTS

A total of 486322 A. sachalinensis F. Schmidt (a species related to A. firma) ESTs were downloaded from the National Center for Biotechnology Information (NCBI) database and used for PCR primer design. First, polyA and adapter sequences were removed from the cDNA sequences using the program Cross_ match (http://bozeman.mbt.washington.edu/phrap.docs/phrap.html) and the TIGR SeqClean sequence trimming pipeline (http://compbio.dfci.harvard.edu/ tgi/software/). EST sequences were then assembled de novo using MIRA (Chevreux et al., 2004), resulting in a total of 38953 contigs (hereafter referred to as unigenes). Using the resultant unigene library, PCR amplicon primers were designed using MISA (Thiel et al., 2003) and Primer3 (Rozen and Skaletsky,
Table 1. Characteristics of the 32 EST-SSR primers used for Abies firma.

| Locus | Primer sequences ( $5^{\prime}-3^{\prime}$ ) | Repeat motif | Size range (bp) | Polymorphism | GenBank accession no. | BLAST top hit description [organism] | BLAST top hit accession no. | $E$-value |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| As_c10422 | F: TCTGAGTGCTAACCTGTGGACTGC | (CTG) ${ }_{5}$ | 184 | no | FX334335 | no hit | - | - |
|  | R: CGGGAGTATGAGGAGTTGTTGACTC |  |  |  |  |  |  |  |
| As_c14033 | F: GACCACACAATTCAAATGATTGCC | $(\mathrm{AG})_{6}$ | 151-156 | yes | FX334336 | no hit | - | - |
|  | R: GTAATGAGCTGGAAGCTGGTCTCC |  |  |  |  |  |  |  |
| As_c14394 | F: GTATGTTGCCTCTGTtttGATGGC | (TGC) ${ }_{5}$ | 103-111 | yes | FX334337 | no hit | - | - |
|  | R: AGCCTGCCACATCTCTCAATATCC |  |  |  |  |  |  |  |
| As_c14606 | F: TGTTATTTCGGGTGGAGTTTTTTGG | (TAA) ${ }_{5}$ | 294-296 | yes | FX334334 | unknown [Picea sitchensis] | ABK2196.1 | 5.25E-21 |
|  | R: ССTCAGACCAACCAAAAGAGAGGA |  |  |  |  |  |  |  |
| As_c23058 | F: AACGTTTTGGATCGACTCCATGTT | $(\mathrm{TGC})_{5}$ | 230 | no | FX334338 | no hit | - | - |
|  | R: GTAACAGCTGAACTACCAGCCACG |  |  |  |  |  |  |  |
| As_c28104 | F: CGAGGAAGAAGCCAAGTTATCAGG | (ATA) ${ }_{5}$ | 153-181 | yes | FX334339 | no hit | - | - |
|  | R: CACAGTTAAAAAGGCGGCCTACAG |  |  |  |  |  |  |  |
| As_c28696 | F: TAAGCAAGGACAGCTTGCATACCC | $(\mathrm{TA})_{8}$ | 234 | no | FX334340 | no hit | - | - |
|  | R: TCTTGTACGCACAACCCTGTCAAT |  |  |  |  |  |  |  |
| As_c32410 | F: CTGAGCACGTGAGGAAGCAAAAT | $(\mathrm{AT})_{6}$ | 117-123 | yes | FX334341 | no hit | - | - |
|  | R: TGGGAGATAGCCTCATTAGGTTGC |  |  |  |  |  |  |  |
| As_c35493 | F: AAGGACCTGGTCAAAAAGCATTCA | (AAG) ${ }_{6}$ | 288 | no | FX334332 | heat shock protein [Picea mariana] | AAC32131.1 | $8.52 \mathrm{E}-15$ |
|  | R: CCGGTGTtACATAACCAGGACCAT |  |  |  |  |  |  |  |
| As_rep_c49 | F: GACGAAGATCAGTACAAGGCACGA | (AGGAGA) ${ }_{7}$ | 257-284 | yes | FX334333 | no hit | - | - |
|  | R: GCGATCCTTCAATTTGTCCTTCTC |  |  |  |  |  |  |  |
| As_rep_c66 | F: GTTGGGGTCGTGAAGAGGACACT | (GTG) ${ }_{6}$ | 251-284 | yes | FX334318 | unknown [Picea sitchensis] | ABK22207.1 | $1.18 \mathrm{E}-29$ |
|  | R: GGCATCGTAGCCATAACTGTAGCC |  |  |  |  |  |  |  |
| As_rep_c4656 | F: TCCTCGTCGTGTTCTACTCССТСТ | $(\mathrm{CTC})_{5}$ | 228-251 | yes | FX334319 | putative syntaxin 1 A <br> [Tanystylum orbiculare] | ABV81823.1 | $4.35 \mathrm{E}-21$ |
|  | R: ACAAATCCAACAATGTCGACAGGA |  |  |  |  |  |  |  |
| As_rep_c5215 | F: GATTCTGATCATGATAGGGGCAGG | (AG) ${ }_{6}$ | 247 | no | FX334320 | RNA-binding protein, putative [Ricinus communis] | XP_002532972.1 | $4.19 \mathrm{E}-08$ |
|  | R: TСТСССтTGTGGСТTTСТTСTTTG |  |  |  |  |  |  |  |
| As_rep_c5432 | F: TGGGTGAAGAGAGAACCAGAAAGG | $(\mathrm{ATG})_{5}$ | 225 | no | FX334321 | unknown [Zea mays] | ACL54598.1 | 3.92E-73 |
|  | R: TCCAATGCGACATAATGATTCCAC |  |  |  |  |  |  |  |
| As_rep_c5928 | F: GGTCTCGAGTTCGAGGACAAAGAA | $(\mathrm{AGG})_{5}$ | 164 | no | FX334322 | 60S ribosomal protein L44 [Elaeis guineensis] | ACF06522.1 | 3.32E-41 |
|  | R: TGCAAAGTGTGCTTTCTACAAGCC |  |  |  |  |  |  |  |
| As_rep_c7912 | F: TAGAGGAAATGCTTGCTCGTCTCG | $(\mathrm{GAA})_{6}$ | 294-299 | yes | FX334323 | PREDICTED: uncharacterized protein LOC100267326 [Vitis vinifera] | XP_002285773.2 | $5.34 \mathrm{E}-13$ |
|  | R: AGGACTTCCTCTGCAAATCCACAC |  |  |  |  |  |  |  |
| As_rep_c10703 | F: GCAGCTGCATCAGTCGCTAAGG | $(\mathrm{GCA})_{5}$ | 152 | no | FX334342 | no hit | - | - |
|  | R: GCCTTCAAGCAATCCAACTTCACT |  |  |  |  |  |  |  |
| As_rep_c10904 | F: TCCATGTCATTTATGGAGCACCTG | $(\mathrm{CAAT})_{5}$ | 125 | no | FX334324 | dormancy/auxin associated-like protein, partial [Picea sitchensis] | ADP94920.1 | $8.93 \mathrm{E}-15$ |
|  | R: CCAATCCAACAGAACATAAATGCAG |  |  |  |  |  |  |  |
| As_rep_c11017 | F: GTtTCAttcgctgttacgatgitga | $(\mathrm{AT})_{6}$ | 234-246 | yes | FX334343 | no hit | - | - |
|  | R: GGAACTTGTCTAAGATTCCGCCAT |  |  |  |  |  |  |  |
| As_rep_c11401 | F: CGGCAACACAGACAGAAGAAAGAA | $(\mathrm{GAA})_{5}$ | 151 | no | FX334344 | no hit | - |  |
|  | R: GGGGATACCTCACATCCACTCAAC |  |  |  |  |  |  |  |

Table 1. Continued.

| Locus | Primer sequences ( $5^{\prime}-3^{\prime}$ ) | Repeat motif | Size range (bp) | Polymorphism | GenBank accession no. | BLAST top hit description [organism] | BLAST top hit accession no. | E-value |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| As_rep_c12415 | F: ACTCCTCCTCCTGGCCTTAAATTG | $(\mathrm{TA})_{10}$ | 285 | no | FX334345 | no hit | - | - |
|  | R: GTGGATTCTTCTCTTCCTGGATCG |  |  |  |  |  |  |  |
| As_rep_c12939 | F: TCCCAATAGAATTTGGGGGATAGC | $(\mathrm{TTC})_{5}$ | 233 | no | FX334346 | no hit | - | - |
|  | R: CTTAGAAGAAGCAGCAGCTCAGCC |  |  |  |  |  |  |  |
| As_rep_c13048 | F: ATGCACAAGGGCCAGAAGTTAGAG | (TGA) ${ }_{5}$ | 267 | no | FX334325 | unknown [Picea sitchensis] | ABK24403.1 | $8.16 \mathrm{E}-60$ |
|  | R: TCATGTTTGCTTCCTCTGCATCTC |  |  |  |  |  |  |  |
| As_rep_c13359 | F: CGGCTTCCTGCTATTACTGTTGCT | $(\mathrm{GCAACG})_{5}$ | 210-235 | yes | FX334326 | unknown [Picea sitchensis] | ADE76551.1 | $2.41 \mathrm{E}-39$ |
|  | R: CATCATGTGATCGTGGTCCTCAC |  |  |  |  |  |  |  |
| As_rep_c14053 | F: TAATATGAGACAGCCTTCGGGCTT | $(\mathrm{AT})_{10}$ | 85 | no | FX334347 | no hit | - | - |
|  | R: CTCCAGGTTACCATCCTTTGGTTG |  |  |  |  |  |  |  |
| As_rep_c14410 | F: ACTGAACTGAGGCACCGGAATTAG | $(\mathrm{CT})_{7}$ | 152 | no | FX334348 | no hit | - | - |
|  | R: AGAGGAGTAGAGAGTGTGGGGACG |  |  |  |  |  |  |  |
| As_rep_c16096 | F: CATCCTTTCGGTGCCTATTATTCG | $(\mathrm{AGA})_{5}$ | 200-203 | yes | FX334327 | unknown [Picea sitchensis] | ABK25258.1 | 4.11E-06 |
|  | R: AACTCTGGTAGAAGAAGCGCAGGA |  |  |  |  |  |  |  |
| As_rep_c17556 | F: GTGAGACAGTTGCCCCTTTCAGTT | $(\mathrm{CAG})_{6}$ | 242-256 | yes | FX334328 | predicted protein [Populus trichocarpa] | XP_002332355.1 | $3.79 \mathrm{E}-35$ |
|  | R: TAAGCTTTCGGAGGCGTTGTATGT |  |  |  |  |  |  |  |
| As_rep_c18764 | F: TGTATTCTTAGAGCCTGTGCAGCAA | $(\mathrm{ATAAG})_{5}$ | 257 | no | FX334349 | no hit | - | - |
|  | R: TAAAGGAGGAAATGGCACGTGAAC |  |  |  |  |  |  |  |
| As_rep_c27580 | F: TCCAAAGGTGGAAGAGAAGCAATC | $(\mathrm{CTT})_{5}$ | 230 | no | FX334329 | unknown [Picea sitchensis] | ABK25146.1 | $1.81 \mathrm{E}-13$ |
|  | R: CTTTGGAGAAAGCCTCATGGAGAA |  |  |  |  |  |  |  |
| As_rep_c32446 | F: CAATTGAAGATGTGCGAAAGTTGC | $(\mathrm{CTG})_{5}$ | 258-265 | yes | FX334330 | unknown [Picea sitchensis] | ADE75915.1 | $9.35 \mathrm{E}-20$ |
|  | R : CTGCTTGCCCCTACATTCACATTT |  |  |  |  |  |  |  |
| As_rep_c33168 | F: TCAACAACGTCGTCAGTGTATAGTCG | $(\mathrm{ATC})_{7}$ | 86 | no | FX334331 | unknown [Picea sitchensis] | ADE75720.1 | 4.16E-22 |
|  | R: CGGATGATGCCATACTTCGGTTAT |  |  |  |  |  |  |  |

Table 2. Characteristics of the 14 polymorphic EST-SSR markers used for three Abies species.

| Locus | A. firma |  |  |  |  | A. homolepis |  |  |  |  | A. veitchii |  |  |  |  | Size range (bp) | Total A |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | $N$ | A | $H_{\text {o }}$ | $H_{\text {e }}$ | $F_{\text {IS }}$ | $N$ | A | $H_{\text {o }}$ | $H_{\text {e }}$ | $F_{\text {IS }}$ | $N$ | A | $H_{\text {o }}$ | $H_{\text {e }}$ | $F_{\text {IS }}$ |  |  |
| As_c14033 | 18 | 2 | 0.333 | 0.284 | -0.172 | 22 | 1* | 0.000 | 0.000 | - | 24 | 2 | 0.375 | 0.361 | -0.040 | 151-156 | 3 |
| As_c14394 | 17 | 1* | 0.000 | 0.000 | - | 22 | 2 | 0.273 | 0.240 | -0.135 | 24 | 2 | 0.042 | 0.042 | 0.000 | 103-111 | 3 |
| As_c14606 | 17 | 1 | 0.000 | 0.000 | - | 22 | 1* | 0.000 | 0.000 | - | 22 | 2 | 0.227 | 0.431 | 0.472 | 294-296 | 2 |
| As_c28104 | 20 | 3 | 0.300 | 0.267 | -0.123 | 24 | 2 | 0.042 | 0.042 | 0.000 | 22 | 1* | 0.000 | 0.000 | - | 153-181 | 3 |
| As_c32410 | 20 | 3 | 0.150 | 0.145 | -0.036 | 23 | 2 | 0.087 | 0.085 | -0.023 | 22 | 1* | 0.000 | 0.000 | - | 117-123 | 3 |
| As_rep_c49 | 20 | 2 | 0.100 | 0.097 | -0.027 | 24 | 3 | 0.458 | 0.368 | -0.246 | 24 | 4 | 0.417 | 0.476 | 0.124 | 257-284 | 6 |
| As_rep_c66 | 20 | 2 | 0.150 | 0.142 | -0.056 | 22 | 1 | 0.000 | 0.000 | - | 22 | 2 | 0.136 | 0.130 | -0.050 | 251-284 | 3 |
| As_rep_c4656 | 20 | 1* | 0.000 | 0.000 | - | 22 | 1* | 0.000 | 0.000 | - | 22 | 2 | 0.364 | 0.476 | 0.236 | 228-251 | 2 |
| As_rep_c7912 | 20 | 1 | 0.000 | 0.000 | - | 24 | 1 | 0.000 | 0.000 | - | 24 | 1* | 0.000 | 0.000 | - | 294-299 | 2 |
| As_rep_c11017 | 18 | 1* | 0.000 | 0.000 | - | 24 | 1* | 0.000 | 0.000 | - | 22 | 1* | 0.000 | 0.000 | - | 234-246 | 1 |
| As_rep_c13359 | 20 | 2 | 0.050 | 0.050 | 0.000 | 24 | 2 | 0.083 | 0.082 | -0.022 | 24 | 1* | 0.000 | 0.000 | - | 210-235 | 3 |
| As_rep_c16096 | 19 | 2 | 0.105 | 0.102 | -0.029 | 22 | 2 | 0.045 | 0.045 | 0.000 | 22 | 1* | 0.000 | 0.000 | - | 200-203 | 2 |
| As_rep_c17556 | 19 | 1* | 0.000 | 0.000 | - | 24 | 1 | 0.000 | 0.000 | - | 22 | 1* | 0.000 | 0.000 | - | 242-256 | 1 |
| As_rep_c32446 | 19 | 1 | 0.000 | 0.000 | - | 22 | 1* | 0.000 | 0.000 | - | 22 | 1* | 0.000 | 0.000 | - | 258-265 | 1 |

Note: $A=$ number of alleles per locus; $F_{\text {IS }}=$ fixation index; $H_{\mathrm{e}}=$ expected heterozygosity; $H_{\mathrm{o}}=$ observed heterozygosity; $N=$ number of individuals genotyped.
*Monomorphic in this population but polymorphic in other populations.
2000), after trimming low quality regions using the qualityTrimmer command in the Euler-SR package (Chaisson and Pevzner, 2008). The criteria applied to identify microsatellite loci were at least six dinucleotide repeat units, or five tri- to hexanucleotide repeat units. To eliminate redundancy (i.e., multiple sets of primers for the same locus), all assembled sequences containing microsatellites were subjected to a BLAST search against the NCBI nonredundant (nr) protein database using the BLASTX algorithm with an $E$-value cutoff of 1.0E-3. A total of 153 EST-SSR primer pairs bordering sequence regions with more than four di- to hexanucleotide repeats were designed. Ninety-six of the 153 primers, for nonredundant loci with large numbers of repeats, were selected for further evaluation. For each primer pair, genomic DNA from one individual of A. firma was used to check PCR amplification. The PCR reaction was carried out following the standard protocol supplied with the QIAGEN Multiplex PCR Kit (QIAGEN, Hilden, Germany), in a final volume of $10 \mu \mathrm{~L}$, which contained approximately 5 ng of DNA, $5 \mu \mathrm{~L}$ of $2 \times$ Multiplex PCR Master Mix, and 0.2 $\mu \mathrm{M}$ of each primer. The PCR thermal profile involved denaturation at $95^{\circ} \mathrm{C}$ for 3 min , followed by 35 cycles of $95^{\circ} \mathrm{C}$ for $30 \mathrm{~s}, 55^{\circ} \mathrm{C}$ for $1 \mathrm{~min}, 72^{\circ} \mathrm{C}$ for 1 min , and a final 7 -min extension step at $72^{\circ} \mathrm{C}$. PCR products were labeled with ChromaTide Alexa Fluor 488-5-dUTP (Invitrogen, Carlsbad, California, USA) according to Kondo et al. (2000), and loaded onto an automated sequencer (ABI Prism 3100 Genetic Analyzer; Applied Biosystems, Carlsbad, California, USA) to determine fragment lengths, which were analyzed using GENOTYPER software (Applied Biosystems). Thirty-two loci exhibited clear PCR amplification with fragment sizes ranging from 50 to 500 bp (Table 1). The polymorphism of these fragments was evaluated using eight individuals of each of three Abies species (A. firma, A. homolepis, and A. veitchii) sampled across the species' geographical range. Fourteen of the 32 loci were polymorphic and provided clear fragment patterns. The genetic variation at these 14 loci was evaluated using 20 individuals from the A. firma population. Information about the populations sampled is provided in Appendix 1, and specimen vouchers were deposited in the Forestry and Forest Products Research Institute herbarium. To characterize each EST-SSR marker, the following four genetic diversity statistics were calculated using FSTAT 2.9.3 (Goudet, 2001): number of alleles per locus $(A)$, observed heterozygosity $\left(H_{0}\right)$, expected heterozygosity $\left(H_{\mathrm{e}}\right)$, and fixation index $\left(F_{\text {IS }}\right)$. In addition, the significance of Hardy-Weinberg equilibrium and genotypic equilibrium were tested by 1000 randomizations with adjustment of the resulting $P$ values by sequential Bonferroni correction, using FSTAT 2.9.3. Cross-amplification was conducted on one population each for two Abies species (Table 2, Appendix 1) following the protocol described above. Of the 14 polymorphic loci, As_rep_c4656, As_rep_c32446, As_c14394, As_rep_c 11017, and As_rep_c 17556 were not polymorphic in this population, but they were polymorphic in other populations (data not shown). As_c14606 was also monomorphic in A. firma but polymorphic in A. veitchii. As_rep_ c7912 was monomorphic in all three species but polymorphic in other populations of A. veitchii.
$A$ ranged from one to three and $H_{\mathrm{e}}$ ranged from 0 to 0.284 . The results of cross-species amplification showed that all 14 loci were amplified successfully
in A. homolepis and A. veitchii. The total number of alleles ranged from one to six. Analysis of the 14 polymorphic loci indicated no significant deviation in $F_{\text {IS }}$ or genotype disequilibrium among locus pairs for any of the three species.

## CONCLUSIONS

The EST-SSR markers described here will be useful for future genetic studies of A. firma. Interspecific amplification of these markers also shows their potential for use in closely related species. These markers may therefore provide a tool for understanding population demography, population structure, gene flow, and mating systems in Abies species.

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Appendix 1. Information about the populations of three Abies species sampled in this study.

| Species | Locality | Geographic coordinates | Accession no. |
| :--- | :---: | :---: | :---: |
| A. firma | Onzui, Shiso City, Hyogo Prefecture, Japan | $35.249^{\circ} \mathrm{N}, 134.523^{\circ} \mathrm{E}$ | $\mathrm{TF}-\mathrm{K} 11-0098$ |
| A. homolepis | Yamanaka, Yamanaka-ko Village, Minami Tsuru County, | $35.438^{\circ} \mathrm{N}, 138.885^{\circ} \mathrm{E}$ |  |
| A. veitchii | Yamanashi Prefecture, Japan | $35.442^{\circ} \mathrm{N}, 138.902^{\circ} \mathrm{E}$ |  |
|  | Yamanaka, Yamanaka-ko Village, Minami Tsuru County, | TWTw 20773 |  |
|  | Yamanashi Prefecture, Japan |  |  |


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