Factors Influencing Male Reproductive Success in a Cryptomeria japonica Seed Orchard Revealed by Microsatellite Marker Analysis

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Abstract
We investigated the influence of male flower production, floral synchrony and inter-tree distances on male reproductive success in a miniature seed orchard of Cryptomeria japonica. We used six microsatellite markers to determine the paternity of each seed. In the seed orchard, the average pollen contamination and clonal self-fertilization rates were 38.7% and 1.7%, respectively. The level of male reproductive success of constituent clones varied from 0.0 to 15.7%. Five clones showing the highest male reproductive success contributed ca. 30% of all analyzed seeds as a pollen donor after excluding contamination by external sources of pollen. The statistical analyses showed that male reproductive success was strongly influenced by male flower production of each clone and, possibly, by their distance to the mother trees. The linear regression which included male flower production and floral synchrony as independent variables, however, accounted for only 14.7% of variation of reproductive success. Since we found no significant correlation between male reproductive and female reproductive success, we may be able to equalize male and female reproductive success independently.

Key words: conifer, floral phenology, paternal contribution, SSR.

Introduction
Seed orchards, which consist of superior tree clones, are important sources of materials for afforestation. Ideally, such materials should have high genetic value and diversity. In order to maintain high genetic value of their seeds, rates of contamination by external pollen and self-fertilization within the orchards should be minimal, while the male reproductive success of the constituent clones should be roughly equal to ensure the seeds have high diversity.

Recently, many studies have assessed the quality of seed yields using molecular markers. Pollen contamination rates exceeding 30% have been found in most seed orchards of dominant forestry species (WHEELER and JECH, 1992 for review; ADAMS et al., 1997; STOEHR et al., 1998; PAKKANEN et al., 2000; MORIGUCHI et al., 2004, 2005), and self-fertilization rates appear to be around 5% in most seed orchards examined to date (RITLAND and EL-KASSABY, 1985; RUDIN et al., 1986; GOTO et al., 2002; MORIGUCHI et al., 2004, 2005). Furthermore, male reproductive success has deviated from equality in all studies in which it has been examined (STOEHR et al., 1998; GOTO et al., 2002; STOEHR and NEWTON, 2002; MORIGUCHI et al., 2004, 2005). These findings suggest that the quality and diversity of seeds in seed orchards may be affected by multiple factors, and several such factors have been identified. First, various authors have proposed that the pollen contamination rate among ramets within seed orchards may be related to their spatial distance from neighboring artificial forests (FRIEDMAN and ADAMS, 1985; YAZDANI and LINDOREN, 1991; MORIGUCHI et al., 2004). Second, the average contamination rate of seed orchards is influenced by the area of surrounding artificial forests (MORIGUCHI et al., 2005). Third, differences in self-fertilization rates of ramets within a seed orchard are related to the temporal separation of seed cone receptivity and pollen shed within clones (STOEHR et al., 1998), the amounts of pollen produced (SCHOEN et al., 1986), and variations in both spacing and crown size (RITLAND and EL-KASSABY, 1985; PAKKANEN et al., 2000). Fourth, the average self-fertilization rate within seed orchards is influenced by the number of ramets per constituent clone (MORIGUCHI et al., 2005). However, there have been few studies on factors that influence male reproductive success, because previously used genetic markers such as allozymes provide insufficient resolution to determine the pollen donors of the progeny, due to the small number of allozyme loci available for analysis and their low polymorphism.

Therefore, in the present study, we used microsatellite markers to investigate pollen-mediated gene flow in a Cryptomeria japonica seed orchard. Microsatellite markers are powerful molecular tools for determining parentage due to the high levels of variation in the repeat units. Our objective was to assess the relationships between male reproductive success of all constituent clones and biological traits such as male flower production, floral synchrony and inter-tree distances. The results of such studies could be used to further improve seed orchards.

Materials and Methods
Investigated seed orchard
This study was carried out in a miniature seed orchard in Niigata Prefecture (37°34′N, 138°46′E). The
height of the trees of each clone was about 2 m, they were spaced 1.5 x 1.5 m apart, and the orchard was 15 years old. The orchard covers 2.8 ha and was divided into 26 blocks. Gibberellin was applied to a study block (0.10 ha) containing 1144 trees (11,555/ha) representing 54 clones to induce the flowering of seed trees at the end of July 1999. Data on biological parameters such as male flower production and flowering phenology in this miniature seed orchard were collected in the flowering season of the year 2000.

**Airborne pollen survey**

Airborne *C. japonica* pollen was collected 1.3 m above the ground by Durham samplers (DURHAM, 1946) at two observation points, one 300 m north of the orchard (outside the seed orchard) and one in the center of the gibberellin-treated study block (Figure 1). The pollen was collected from 7 March to 11 April. A glass slide smeared with petrolatum was collected every day at 9:00 AM, and the pollen grains within a 1 cm x 1 cm area on the slide were counted under a microscope at x 200 magnification. Daily airborne pollen counts were expressed as counts/cm².

![Figure 1. – Location of the nine mother trees in studied block and pollen samplers (black circles).](image)

**Male flower production**

First, the amount of male flowers on each ramet was visually graded in the following five classes: (0) none, (1) very few, (2) light, (3) medium, and (4) high. Second, the quantity of male flowers on a single ramet was estimated for each specific class by random sampling measurements. That is, we randomly sampled three ramets from each class, and measured the total weight of male flowers on the single ramet after drying at 105°C for three days. Then, male flower production of a single ramet was estimated for each class as the average weight of these three ramets. Finally, male flower production of each clone was calculated as the sum of male flower production of all ramets belonging to the clone.

**Floral synchrony**

Floral synchrony between male and female flowers of different clones was calculated as follows. First, we assessed the temporal pollen dispersal capacity of each clone by examining five ramets per clone every three days from March 6 to April 11. We tapped the male flowers on the five ramets and graded the pollen dispersal capacity of each clone in four classes: (0) pollen not yet ready to shed, (1) small amounts of pollen ready to shed, (2) medium amounts of pollen shed, and (3) considerable amounts of pollen shed. The temporal pollen dispersal capacity on the observation day for each clone was estimated from the average of the pollen dispersal scores of its five ramets. The temporal pollen dispersal capacity was standardized so that the total score over the flowering period be unity.

Next we assessed the temporal reproductive ability of female flowers for each clone. A twig with female flowers (10 to 20 per twig) was covered by a paper bag once every three days from March 12 to April 4. We assumed that there were no phenological differences among ramets belonging to the same clone, since the variation in floral phenology among ramets within clones is usually minor relative to clonal differences in many conifers (JONSSON et al., 1976; GRIFFIN, 1982; EL-KASSABY et al., 1984). All the covering bags were removed in the middle of May when the male flowers finished shedding pollen. Bagged cones were collected at the end of October, and non-bagged (i.e., open-pollinated) cones were also collected randomly as controls. We then carried out a germination test of seeds extracted from the cones in an incubator at 23°C for 34 days using 100 seeds per clone with three replicates. The pollination rate of each observation day was defined as the ratio of the germination rate of bagged seeds covered on the observation day to the germination rate of open-pollinated seeds. The reproductive ability of female flower for each clone during the period between two successive observation days can be calculated as the difference in the pollination rate between the two days. This temporal reproductive ability was also standardized so that the total score over the flowering period be unity.

Floral synchrony between two clones can be calculated as degree of overlap between the temporal pollen dispersal capacity of one clone and the temporal reproductive ability of another. That is, the degree of floral synchrony between male flowers of one clone and female flowers of another clone can be calculated as the sum of products between the temporal pollen dispersal capacity and the temporal reproductive ability over all observation days.

**Total seed production**

Total seed production of each clone was calculated as follows. First, the amount of cones produced by each ramet was graded in four classes: (0) none, and: (1) light, (2) moderate, and (3) high. Second, all cones were harvested from three randomly selected ramets in each class, and dried. Total weight of seeds extracted from the cones was measured for each ramet. Then, seed production of a single ramet of each specific class was estimated as the average seed weight over the three ramets. Finally, total seed production of each clone was calculated as the sum of seed production of all ramets belonging to the clone.
DNA extraction

We collected needles from all clones in the scion garden that provided scions to establish the miniature seed orchard. The total DNA of all constituent clones in the orchard was then extracted from needle tissues using the CTAB method (MURRAY and THOMPSON, 1980) or modified CTAB method (TSUMURA et al., 1995). The extracted crude DNAs were purified using a Fast DNA kit (BIO 101).

We collected seeds from nine ramets clones (represented by eight clones) to investigate the pollen contamination rate and male reproductive success of each clone in this miniature seed orchard (Figure 1). Seeds were sown on sterilized paper on plastic plates, and DNA was extracted from germinated embryos using the CTAB method (MURRAY and THOMPSON, 1980).

Microsatellite genotyping

We used six microsatellite markers that showed high stability and polymorphism (Table 1; MORIGUCHI et al., 2003 and TANI et al., 2004) to determine the pollen donors and pollen contamination rates of seeds from nine mother trees. These markers have shown co-dominant segregation in a three-generation pedigree of *C. japonica*. PCR amplifications were carried out using Model 9600 and 9700 GeneAmp PCR Systems (Applied Biosystems) in mixtures, with a total volume of 10 µL, containing 20 mM Tris-8.0, 50 mM KCl, 1.5 mM of MgCl₂, 0.2 µM of each primer, 5 ng of template DNA, and 0.625 units of Taq polymerase (PRIMEGEN), with the following program: 5 min denaturation at 94°C; followed by 30–35 cycles of 30 sec denaturation at 94°C, 30 sec annealing at 55–60°C, and 30 sec extension at 72°C; with a final extension step of 72°C for 5 min. Genotypes of paternal candidate *C. japonica* trees and 100 offspring from each of nine plus trees based on the microsatellite markers were determined using ABI prism 310 and 3100 genetic analyzers (Applied Biosystems).

Data analysis

From the genotype data for the paternal candidates of the constituent clones, we calculated the number of alleles per locus (NA), polymorphism information content (PIC; BOTSTEIN et al., 1980), the paternity exclusion probability and multi paternity exclusion probability (WEIR, 1996) for each locus using the G-DIVERSE software developed by Hiroyoshi Iwata (NARC, Tsukuba, Japan).

We determined the pollen donor clones of the seeds that had not originated from pollen contamination or self-fertilization, using the method of MORIGUCHI et al. (2004). Deviations from equal male reproductive success were evaluated using the χ² goodness of fit test, as follows. First, we examined the null hypothesis of equal male reproductive success for each ramet to exclude the effect of the number of ramets per clone using the formula ECV₁ (see below). Second, we examined the null hypothesis of equal male reproductive success for each ramet to exclude the effect of the difference of total male flower production among clone using the formula ECV₂:

ECV₁ = (total number of seeds fertilized by plus-trees) x [(number of ramets in i-th clone)/(total number of ramets)].

ECV₂ = (total number of seeds fertilized by plus-trees) x [(total male flower production of i-th clone)/(total male flower production of all clones)].

The observed male reproductive success values were based on the number of seeds contributed by each clone.

To assess the influence of biological traits such as total male flower production and floral synchrony on the male reproductive success of each clone, we performed a multi linear regression analysis, including total male flower production and floral synchrony as independent variables and male reproductive success as a dependent variable, using SPSS ver. 11.5 (SAS Institute Inc., Cary, N.C.).

Table 1. - Microsatellite markers used in this study and their polymorphism.

<table>
<thead>
<tr>
<th>primer</th>
<th>motif</th>
<th>primer sequence</th>
<th>NA a</th>
<th>PIC b</th>
<th>Q c</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cggsr 78</td>
<td>(GA)₃₀</td>
<td>AAGGAAGGAGTCCAGAGGAAGTGA ACTGCGAGTAACTGAATGCT</td>
<td>18</td>
<td>0.916</td>
<td>0.841</td>
</tr>
<tr>
<td>Cs 1525</td>
<td>(CA)₃₀</td>
<td>ATGAAGTGCCCCTGTTGTG ATCAGCTCTCTTTATCC</td>
<td>32</td>
<td>0.929</td>
<td>0.866</td>
</tr>
<tr>
<td>Cggsr 181</td>
<td>(GA)₃₀</td>
<td>AGAGGGAGGGAGGAAATACAT GGCAGAGTGTTATGTTAC</td>
<td>42</td>
<td>0.950</td>
<td>0.905</td>
</tr>
<tr>
<td>Jsc20</td>
<td>(TG)₃₀</td>
<td>TCCCTTTTTGTTATTTTACAT ACTCAAATCGGATATCT</td>
<td>13</td>
<td>0.773</td>
<td>0.615</td>
</tr>
<tr>
<td>Cj 333</td>
<td>(GA)₃₀</td>
<td>AGGAGATTAGGATTGGTGGG GGTGTTACCTCTATGAT</td>
<td>33</td>
<td>0.935</td>
<td>0.876</td>
</tr>
<tr>
<td>Cjggsr 77</td>
<td>(CT)₃₀</td>
<td>CCTTGATACATTTAATTTGACCT AGGGAGGAGGAGATA</td>
<td>29</td>
<td>0.855</td>
<td>0.748</td>
</tr>
</tbody>
</table>

a Number of alleles; total NA = 167
b Polymorphism information content; average PIC = 0.893
c Multi-paternity exclusion probability = 0.999
To assess the effect of spatial distance to the mother tree on male reproductive success, we carried out a randomization test as follows. First, we calculated the mean number of seeds of fertilized by the $i$-th of $k$ surrounding clones (where $k = 8$, in general). Second, we randomly selected constituent clones of the same number, $k$, and calculated the mean number of seeds fertilized by the $i$-th investigated clone. This randomization process was carried out 10,000 times, generating an empirical distribution of the mean number of seeds under the null hypothesis that male reproductive success does not depend on the distance between maternal and paternal trees. Finally, using the obtained empirical distribution, we calculated the probability of the mean number of seeds fertilized by the $i$-th clone being greater or equal to the number observed, and if the significance level, $P$, of the difference between the randomized and actual distributions was $\leq 0.05$, the alternative hypothesis was accepted.

To assess the relationship between male reproductive success and female reproductive success, we calculated Spearman’s $\rho$ rank correlation coefficients between these variables (using SPSS ver. 11.5; SAS Institute Inc., Cary, N.C.) for all constituent clones.

**Results and Discussion**

**Microsatellite markers**

For paternity analysis markers are required that can provide high multi-paternity exclusion probabilities (Weir, 1996). The six microsatellite markers we used here had sufficient polymorphism for this purpose, yielding a multi-paternity exclusion probability of 0.999 (Table 1), and thus sufficient resolution to determine pollen donors in the seed orchard.

**Differences in male reproductive success**

Airborne dispersal of *C. japonica* pollen had begun by 7 March at each observation point and increased rapidly from 11 March. The highest peaks were recorded on 19 and 28 March (Figure 2). Little airborne pollen was recorded after the end of April. The pollen captured at the two monitoring points (one inside and one outside the orchard) showed similar dispersal patterns.

We were able to determine a paternal clone for each seed after excluding seeds originating from contamination (i.e., fertilization by pollen from outside the orchard) (38.7%) and self-fertilization (1.7%). The $\chi^2$ test for the goodness of fit to the null hypothesis of equal male reproductive success of each clone showed that the level of male reproductive success, in terms of their contributions to seed production by the constituent clones, differed significantly ($p < 0.001$ in both cases). Although in both cases the departure from the expectation on equal male reproductive success were highly significant, the $\chi^2$ value was smaller in the case considering the total male flower production of each clone than that in the case considering the total number of ramets of each clone. This indicates that total male flower production of each clone may be more important than total number of ramets of each clone when we will equalize the contribution of each clone as a pollen donor. Eight clones (nos. 5, 6, 10, 26, 31, 52, 54, and 57) showed no male reproductive success at all. The clone with the highest male reproductive success was clone 53, which fertilized 9.8% of the examined seeds (Figure 3). The five clones with the highest male reproductive successes (nos. 53, 3, 58, 71, and 11) collectively accounted for about 30% of the paternity of all assessed seeds that did not originate from fertilization by contaminating pollen. Similar results have been found in seed orchards of other conifer species, such as *Pinus contorta* Doug., *Pinus thunbergii* Parl. and *Pseudotsuga menzeisii* Franco (Stoehr et al., 1998; Goto et al., 2002; Stoehr and Newton, 2002). However, the male reproductive success of each constituent clone should ideally be roughly equal to ensure high genetic diversity amongst the resulting seeds and to help protect the progeny against disease and environmental changes (Zhu et al., 2000; Burdon, 2001).

**Relationships between male reproductive success and biological traits**

The total male flower production of the clones was strongly associated with their male reproductive suc-
cesses (Figure 4; Table 2), but there was no clear relationship between male reproductive success and the floral synchrony (Figure 5). The multilinear regression model we derived accounted for ca. 15% of the variance in the male reproductive success of the clones. The coefficient of regression was significant at the 0.5% probability level for total male flower production, but non-significant for floral synchrony (Table 2). A single linear regression of male reproductive success on total male flower production showed a significant positive slope ($P < 0.005$; Figure 4). This finding is consistent with previous reports that the total number of male cones correlates with male reproductive success in *Picea glauca* Voss (SCHOEN and STEWART, 1986), *Pseudotsuga menziesii* Franco (BURCZYK and PRAT, 1997) and *Pinus thunbergii* Parl. (GOTO et al., 2005). In our study, there was a ca. 20-fold difference in pollen production between the clones that produced the most and least pollen. Similarly, KUSABA (1985) reported that the amount of male flower production differed 100-fold between clones in another *C. japonica* seed orchard. Thus, the influence of the differences in total male flower production among clones on male reproductive success might be unexpectedly large in conifers.

ERICKSON and ADAMS (1989) suggested that both relative male flower production and floral overlap could strongly influence inter-mating patterns amongst seed orchard clones, and GOTO et al. (2002) have shown that a lack of flowering phenology overlap with other clones can inhibit male reproductive success. However, in our study, there was no apparent relationship between male reproductive success and floral synchrony (Figure 5; Table 2). In *C. japonica*, pollen grains are pulled into the micropyle rapidly after capture by the pollination droplet (the fluid in the canal). We investigated the flowering period of female flowers using paper bags, because it is impractical to observe pollination droplets directly. The data we gathered in this manner show that the male reproductive success of each clone was not significantly influenced by the floral synchrony per se, but it was significantly affected by the density of their pollen-grains in the air at the time that the pollination droplets were extruded. Therefore, it may be difficult to detect

**Figure 3.** – Observed (solid bars) and expected male reproductive success (open bars) of each clone. The ECV values in text were used as expected male reproductive success.

**Figure 4.** – The relationship between male reproductive success and total male flower production. The male reproductive success of each clone is represented by the proportion of seeds originating from fertilization by its pollen (relative to all seeds except those originating from fertilization by contaminating pollen).

**Table 2.** – Results of the multilinear regression, including male reproductive success as a dependent variable, and total male flower production and floral synchrony as independent variables.
effects of floral synchrony due to the stronger effect of total male flower production.

The low $R^2$ values obtained in the regression analysis imply that factors other than male flower production and floral synchrony influenced male reproductive success (Table 2). Male reproductive success depends not only on flowering traits, such as male flower production and floral synchrony, but also on inter-tree distances and traits of the pollen grains, such as germination vigor and time, pollen growth rate and selective fertilization (Pfaehler, 1975). In this study, the substantial male reproductive success of clone 53 was mainly due to its high contribution to seed production in clone 6 (accounting for about 64% of its total fertilizations). The floral synchrony period between these clones is not very long, so other factors such as pollen competition may also influence male reproductive success (Stoehr et al., 1999; Nikkanen et al., 2000; Aronen et al., 2002). Further study is required to assess this possibility.

Table 3. – Significance of the distance to each of the mother trees considered in the randomization test.

<table>
<thead>
<tr>
<th>Mother tree</th>
<th>$P$</th>
</tr>
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<tbody>
<tr>
<td>BG</td>
<td>0.329</td>
</tr>
<tr>
<td>AP</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>AU</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>AM</td>
<td>0.696</td>
</tr>
<tr>
<td>AH</td>
<td>0.112</td>
</tr>
<tr>
<td>AO</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>AA</td>
<td>0.159</td>
</tr>
<tr>
<td>AI</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>AX</td>
<td>&lt; 0.01</td>
</tr>
</tbody>
</table>

Effect of spatial distance to mother trees on male reproductive success

In the randomization test, we detected significant distance effects on male reproductive success in five of the nine investigated trees (Table 3), implying that male reproductive success was related to the distance between the mother trees and pollen donors. This distance may slightly influence male reproductive success, although the observed tendency was not strong. Shen et al. (1981) found that as much as 31% of the fertilizing pollen could be received from neighboring trees when the wind direction and flowering times were favorable. Erickson and Adams (1989) also suggested that the distance between mother trees and pollen donors is an important factor for mating success when the combined effects of floral overlap and relative male flower production of potential male parents is high. Our results are consistent with these reports.

Seed orchard improvement

The results of the present study indicate that total male flower production strongly affects male reproductive success and the inter-tree distance also has some effect. The distance effects may be balanced out when all the seeds from all ramets representing the same clone are considered, because seed orchards are generally designed to promote random mating. To reduce inequalities in male reproductive success in seed orchards, we should pay more attention to the total male flower production of each clone. However, total reproductive success includes not only male reproductive success, but also female reproductive success. We investigated the relationship between male reproductive and female reproductive success, but found no correlation between them (Figure 6, Spearman $\rho = -0.79$, $P = 0.568$), sug-
gesting that we should concentrate on equalizing male flower production and/or synchronizing floral phenology between clones in seed orchards. To reduce the variations in reproductive success we could harvest equal numbers of seeds from each of the clones, or adjust the numbers of the various clones to equalize male flower production, if the labor costs are not too high. The information from our study should facilitate the future improvement of *C. japonica* seed orchards.

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References


Genetic Relationships among *Schizolobium parahybum* (Vell.) Blake (Leguminosae) Ecotypes from Ecuador and other Countries

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Abstract

Fifteen ecotypes of *Schizolobium parahybum* (Vell.) Blake collected in Ecuador (9), Brazil (3), Bolivia (1) Costa Rica (1), and Peru (1) were analyzed using Random Amplified Polymorphic DNA (RAPDs), Amplified Fragment Length Polymorphisms (AFLPs) and micro-satellites (SSRs) in order to determine their genetic relationships and diversity patterns among ecotypes and to identify the origin of cultivated germplasm in Ecuador. Although AFLP markers were the most informative technique based on amplified products, SSRs clearly differentiated the ecotypes of Ecuador based on their geographical origin or genetic status into two groups: commercial ecotypes growing at western Ecuador very similar to the ecotype from Costa Rica, and native germplasm from eastern Ecuador and ecotypes from Brazil, Peru and Bolivia.

Key words: AFLP, genetic relationships, DNA-fingerprinting, RAPD, *Schizolobium parahybum* (Vell.) Blake, SSR.

Introduction

Forestry industry for wood production of Ecuador extensively exploits *Schizolobium parahybum* (Vell.) Blake (“pachacho”, “guanacaste” or “palo de picho”). This species is largely distributed through America due it shows good adaptation to variable climate conditions and has great possibilities to continue its exploitation for wood production and other uses. Farmers from

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**Introduction**

Forestry industry for wood production of Ecuador extensively exploits *Schizolobium parahybum* (Vell.) Blake (“pachacho”, “guanacaste” or “palo de picho”). This species is largely distributed through America due it shows good adaptation to variable climate conditions and has great possibilities to continue its exploitation for wood production and other uses. Farmers from Ecuador are interested to establish orchards which show fast growth and the best wood quality for production of cellulose (Montenegro, 1987). In 1950, *S. parahybum* was introduced to Quevedo, Ecuador by the Experimental Laboratory “Pichilingue” of the Instituto Nacional de Investigaciones Agrícolas y Pecuarias (INIAP). Although no data are available about the origin of the introduced germplasm, it has been assumed that it originated from Costa Rica. Therefore, no evidence is known about the origin of the most of commercial plantations in Ecuador, from Costa Rica or native germplasm. In Ecuador, *S. parahybum* is commonly propagated by seeds but no plant or seed pre-selection is done for the establishment of orchards and this fact has increased the genetic variability on commercial and native plantations (Tipan, 1982).

The knowledge of genetic diversity and the relationships among genotypes will be important for the development of appropriate strategies for the *in situ* conservation of natural woods and the regeneration of partially logged forests of *S. parahybum* in Ecuador. This information can also serve as a baseline for determining whether genetic diversity has been lost through the sampling or conservation involving *ex situ* propagation, since those facts have been done in germplasm from Ecuador for plant improvement. Genetic diversity characteristics will be useful for planning a breeding strategy for commercial purposes. Genetic diversity of *S. parahybum* could be characterized based on morphological traits or using molecular markers which detect variation at the DNA sequence level. In particular, DNA-based polymorphisms are a powerful tool for assess genetic similarity between natural and breed genotypes.

The PCR-based techniques provide a representative sample of the genome and a virtually unlimited number