Phenotypic and Genetic Diversity Evaluation of Sengon (Falcataria moluccana (Miq.) Barneby & JW Grimes) from Solomon Provenance on Progeny Trial in Cirangsad Experimental Forest, West Java

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Abstract

Initial evaluation of phenotypic variability and genetic diversity was conducted on the progeny test of 2 years-old sengon from Solomon provenance, which derived from nine families of mother tree and were then planted in 4 blocks in the Cirangsad Experimental Forest. Phenotypic assessment on eight traits was conducted on 36 trees using a scoring system, while genetic diversity of 13 selected individuals which represented high and low-score phenotypic traits was analyzed using 5 selected RAPD primers. The result on phenotypic assessment showed that family 3 (37.25 points) has the highest average score of phenotypic quality and family 4 has the lowest average score (7.50 points). Furthermore, genetic analysis showed that the low-scoring sengon population had a greater mean genetic diversity (He = 0.2535) than the high-score population (He = 0.2345). The analysis of molecular variance (AMOVA) revealed a significant genetic differences (p-value < 0.001) among high and low-score populations and the dendrogram of genetic distance revealed clustering of individuals having similar superior phenotypic against those having non-superior, indicating the selection based on phenotypes in this study had succeeded in pooling the good quantitative alleles in the selected population. This evaluation results can be used as a reference in determining the best families to produce superior sengon (from Solomon provenance) offspring in the future that have desired adaptability, productivity, and diversity.

Keywords: Falcataria moluccana, genetic diversity, phenotypic variability, RAPD, sengon, Solomon provenance

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Introduction

Sengon (Falcataria moluccana (Miq.) Barneby & J. W. Grimes) is a fast-growing tree species belonging to the Fabaceae family which is native to Moluccas, New Guinea, New Britain, and the Bismarck Archipelago and Solomon Islands (Wagner et al., 1999; Setiadi et al., 2014; Baskorowati et al., 2017). The tree can grow up to 40 m tall with a clear bole up to 20 m, a diameter of up to 100 cm, and a horizontal canopy (Krisnawati et al., 2011). This species does not require specific growing requirements and is easy to adapt, thus it is widely planted and become one of the highest priority species in industrial forest plantations and community forest in Indonesia especially in Java Island as raw material for pulp, paper, and timber that most used in people's sawmills (Priadi & Hartati, 2015; Baskorowati et al., 2017; Hidayat et al., 2017a; 2017b; Mahbub et al., 2020). The provinces of Central Java and West Java are reported to have contributed more than 60% of the total number of sengon trees planted by community forest (Krisnawati et al. 2011). In 2019, the number of sengon stands harvested has produced 62,270 m³ of wood making this species one of the 10 highest-produced timber in Indonesia after Acacia, Eucalyptus, teak, Shorea, pine, and mahogany (BPS, 2021).

The cultivated sengon is very diverse in terms of growth performance and its productivity is still low (Setiadi et al., 2014; Rostini et al., 2018), therefore a sengon stands that has relatively homogeneous growth and high productivity is much sought for. One of the recommended stands is the population from Solomon provenance, which is reported to have higher productivity than local sengon (Ikhfán & Wijayanto, 2019), which the productivity of Solomon sengon can reach 3 times compared to local sengon which are currently found in Indonesia, especially Java (Setiadi et al., 2014). However, sengon from Solomon provenance is very rarely cultivated by the community in Indonesia due to lack of availability of the seeds that must be imported from Solomon Islands (Setiadi et al., 2014). Therefore, the Rumpin Seed Source and Nursery Centre (RSSNC) in collaboration with State-Owned Forestry Company (Perum Perhutani) has established the progeny trial plot in the Cirangsad Experimental Forest which is intended to build seed sources, conserve genetic material and test the suitability of land on the species collected, one of which is sengon from Solomon provenance. Information on genetic diversity is necessary to formulate conservation strategies and marker-based assessment can be used such as DNA fragment analysis (Loera-Sánchez et al., 2019). The use of RAPD markers rather than other molecular markers for the analysis of genetic diversity in this study was based on the inexpensive, simplicity, and ease of use of these markers. In addition, this marker is dominant and does not require specific knowledge of DNA sequences (Williams et al.,...
In order to select the best offspring of *sengon* from Solomon provenance planted in the progeny trial plot in Cirangsad Experimental Forest, an initial evaluation of offspring based on phenotypic variability and genetic diversity are needed. Therefore, the objective of this study was to evaluate the phenotypic variability and the genetic diversity of the Solomon *sengon* population grown in the Cirangsad Experimental Forest through Random Amplification of Polymorphic DNA (RAPD) genetic markers. The results of this evaluation study are expected to be able to recommend the best families that can be used to supply the needs of good quality Solomon *sengon* seeds in the future.

**Material and Methods**

**Study site and plant material** This study was divided into 2 activities, namely (i) fieldwork consisting of phenotypic assessment and leaf samples collection for further DNA analysis, which were carried out in the Experimental Forest of Cirangsad, and (ii) laboratory work consisting of genetic diversity analysis using Random Amplification of Polymorphic DNA (RAPD) marker, which were conducted at the Forest Genetics Laboratory, Department of Silviculture, Faculty of Forestry, IPB University, Bogor. Geographically, the Experimental Forest of Cirangsad located in E 106°33'19.07771”–E 106°34'02.3985” and S67°30'36.6411”–S67°29'33.1894” with an altitude of 600–700 m above sea level (Figure 1). The forest is divided into several areas based on research objectives, namely progeny test, species trial, demonstration plot and untested seed orchard. Cirangsad experimental forest area has a sloping and hilly topography. The texture of the soil is clay with a CEC of 25.02 (meq 100 g of soil). The soil is yellowish brown in color and has a pH of 4.84. The soil nitrogen content was 0.14%, while the C-organic content was 1.30%. Then the organic matter contained in the soil is 2.23%. The climate in Cirangsad is classified as type A (Schmidt & Ferguson, 1951) with an average rainfall of 2,000–2,500 mm year⁻¹.

Assessment on phenotypic variability was carried out on a 2-years old *sengon* from Solomon provenance tree stand planted in a single tree plot with a randomized complete block design at the progeny trial area of the Cirangsad Experimental Forest. The trees are located in four blocks which each block consists of nine individual trees (Figure 2). These nine individual trees are offspring from nine different mother trees (hereinafter referred to as a family), which originated from Solomon Island and were then planted in Pompok Landak Permanent Nursery located in Haurwangi Village-Bojongpicung Sub-District-Cianjur Regency-West Java Province. In total, 36 individual trees were used in this study (Table 1).

![Figure 1 Map of study site in Cirangsad Experimental Forest.](image-url)
Phenotypic assessment: The phenotypic quality assessment of 2-years old sengon from Solomon provenance tree was divided into measurement of quantitative parameters and qualitative parameters, from which those parameters were selected based on desirable traits of economic interest of sengon tree such as stem straightness, clear bole height, disease resistance, etc. Quantitative parameters include survival percentage, tree height, clear bole height, stem diameter, branch diameter and branch angle. Meanwhile, qualitative parameters include stem shape, bark color, and resistance to pests & diseases. Tree height and branch free height were measured by 250 cm long-tree height measuring stick with a marker for every 10 cm. While, stem diameter (diameter at breast height) and branch diameter were measured by tree caliper.

The measurement data that has been obtained was then scored based on the scoring system method with a comparison tree for several important trait according to Zobel and Talbert (1984) with some modification such as excluding crown and natural pruning parameters. In this study, 18 individuals of 2-years old common sengon tree and growing 4 m from the tested sengon tree were used as comparison trees. The comparison tree used were located near block 1 and block 3 of the tested sengon. Scoring on the sengon from Solomon provenance tree was intended to obtain a total score of all parameters measured for each individual. Each measured parameter has a specific score and the maximum total score for all parameter is 90 points (Table 2). The method of scoring analysis is by giving a rank to the average scoring rate for each family. Rank 1 shows the highest average score in the family, while ranking 9 shows the lowest average score in that family.

Genetic assessment: Leaves from each individual sengon tree selected based on the phenotypic score were collected and dried in the field with silica gel for further genetic analysis. Silica gel-dried leaves were ground to a fine powder in liquid nitrogen with a mortar and pestle and subsequently used for DNA extraction. Total genomic DNA was extracted from each sampled leaf using the modified CTAB method (Weising et al., 2005).

The genomic DNA of each individual trees was amplified by Random amplified polymorphic DNA (RAPD) using five selected primers, from Operon Technologies Inc. with average size of 10 bases: OPA 05 (5' AGGGGTCTTG '3), OPA 14 (5' TCTGTGCTGG '3), OPO 10 (5' TCAGAGCGCC '3), OPU 05 (5' TTGGCGGCCT '3), and OPY 16 (5' GGCCCAATGT '3). These five primers were selected from the screening results of 23 primers kit A (OPA), B (OPB), O (OPO), U (OPU) and Y (OPY) which were successfully amplified, had clear bands and were polymorphic.

Polymerase Chain Reaction (PCR) amplification was performed in a volume of 13.5 μl, containing about 40 ng of genomic DNA, 2.5 μM of primer, and Taq DNA Polymerase (GoTaq® Green Master Mix, Promega). The reactions were

![Figure 2 A sketched map of 2-years old sengon (from Solomon provenance) progeny trial blocks in Cirangsad Experimental Forest.](image)

Table 1  
| Family | Block |
|--------|-------|
|        | 1     | 2     | 3     | 4     |
| 1      | S1B1  | S1B2  | S1B3  | S1B4  |
| 2      | S2B1  | S2B2  | S2B3  | S2B4  |
| 3      | S3B1  | S3B2  | S3B3  | S3B4  |
| 4      | S4B1  | S4B2  | S4B3  | S4B4  |
| 5      | S5B1  | S5B2  | S5B3  | S5B4  |
| 6      | S6B1  | S6B2  | S6B3  | S6B4  |
| 7      | S7B1  | S7B2  | S7B3  | S7B4  |
| 8      | S8B1  | S8B2  | S8B3  | S8B4  |
| 9      | S9B1  | S9B2  | S9B3  | S9B4  |

Note: S = sengon tree; B = block; example: S1B1 = sengon tree from family 1 in block 1.
Table 2  Method of phenotypic quality assessment for sengon (from Solomon provenance) trees

| No | Trait              | Full score (point) | Evaluation system | Score (point) |
|----|--------------------|--------------------|-------------------|--------------|
|    |                    |                    | Value Scale       |              |
| 1  | Stem height        | 10                 | Percentage of values above the average of control trees <5% | 0            |
|    |                    |                    | Percentage of values above the average of control trees 5<10% | 2            |
|    |                    |                    | Percentage of values above the average of control trees 10<15% | 4            |
|    |                    |                    | Percentage of values above the average of control trees 15<20% | 6            |
|    |                    |                    | Percentage of values above the average of control trees 20<25% | 8            |
|    |                    |                    | Percentage of values above the average of control trees ≥25% | 10           |
| 2  | Stem diameter      | 20                 | Percentage of values above the average of control trees <5% | 0            |
|    |                    |                    | Percentage of values above the average of control trees 5<10% | 5            |
|    |                    |                    | Percentage of values above the average of control trees 10<15% | 10           |
|    |                    |                    | Percentage of values above the average of control trees 15<20% | 15           |
|    |                    |                    | Percentage of values above the average of control trees ≥20% | 20           |
| 3  | Clear bole height  | 10                 | Percentage between clear bole height and total height <30% | 0            |
|    |                    |                    | Percentage between clear bole height and total height <30% | 2            |
|    |                    |                    | Percentage between clear bole height and total height <30% | 4            |
|    |                    |                    | Percentage between clear bole height and total height <30% | 6            |
|    |                    |                    | Percentage between clear bole height and total height <30% | 8            |
|    |                    |                    | Percentage between clear bole height and total height <30% | 10           |
| 4  | Stem shape         | 30                 | Straight stem     | Subtract 3–20 points |
|    |                    |                    | S-shaped stem     | Subtract 3–20 points |
|    |                    |                    | Bow-shaped stem   | Subtract 3–20 points |
|    |                    |                    | J-shaped stem     | Subtract 2–8 points  |
|    |                    |                    | Spiral steam      | Subtract 2–8 points  |
|    |                    |                    | Forked stem       | Subtract 5–20 points  |
| 5  | Branch diameter    | 5                  | Percentage between the diameter of the branch and the diameter of the stem where the branch is located >50% | 0            |
|    |                    |                    | Percentage between the diameter of the branch and the diameter of the stem where the branch is located 30–50% | 2            |
|    |                    |                    | Percentage between the diameter of the branch and the diameter of the stem where the branch is located <30% | 5            |
| 6  | Branch angle       | 5                  | Branch angle <50° | 0            |
|    |                    |                    | Branch angle 50–70° | 2            |
|    |                    |                    | Branch angle >70° | 5            |
| 7  | Survival rate      | 5                  | Dead tree         | 0            |
|    |                    |                    | Living tree       | 5            |
| 8  | Pest and disease resistance | 5 | Unhealthy (area of signs of pest attack >20%) | 0 |
|    |                    |                    | Healthy (area of signs of pest attack <20%) | 5 |

Phenotypic variability In this study, phenotypic variability of the studied sengon was estimated using POPGENE 32 (Yeh et al., 1997) and the presence of molecular variance within and between the sengon population structure was assessed through Analysis of molecular variance (AMOVA) by using GenAIEx 6.5 software (Peakall & Smouse, 2006). In addition, the un-weighted pair group method with arithmetic averages (UPGMA) based dendrogram was constructed using the NTSYS version 2.0 (Rohlf, 1998).

Results and Discussion

Phenotypic variability In this study, phenotypic variability of the studied sengon was estimated by scoring 8 phenotypic parameters on 36 individual 2-years old trees derived from 9 families. The results showed that 9 out of 36 trees died, therefore the overall full score of the parameters for these nine individuals was 0 (Table 3). The families that had a high survival rate (100%) were families 1, 2, 3, and 8. While family 4 had a lower survival rate (25%) in each block than other families indicated that individual tree from family 4 had a lack of ability to survive in low or hilly areas such as the area in the Cirangsad Experimental Forest.
The scoring results of all parameters at the family level (Table 3) showed that family 3 has the highest average score (57.25 points) and family 4 has the lowest average score (7.50 points) indicated that family 3 has superior traits in all parameters and family 4 has less superior traits in all growth parameters, especially the survival rate parameter where only 1 individual tree was still alive. Meanwhile, the score for all parameters at the individual level showed that individual tree from family 1 planted in block 2 (S1B2) received the highest score (83 points) followed by family 3 in block 3 (S3B3) and family 6 in block 3 (S6B3) which have 76 points, respectively. Additionally, individual tree from family 4 planted in block 2, 3, and 4 (S4B2, S4B3, and S4B4) received the lowest score (0 points). The results at the individual level are almost in line with the results at the family level where family 3 has a high average score and family 4 has a low average score among the other 9 families. The results of this evaluation can be used as a reference in determining the best families as seed sources as seen from the performance of the offspring.

The number of individuals of studied sengon that had a scoring value of ≥60 points was 7 individuals and the number of individuals that had a scoring value of <60 points was 28 individuals, of which there were only 8 individuals who had a score of <45 points. Based on these scoring results, 15 of the 27 living individuals in 4 blocks were selected with the highest (7 individuals) and lowest (8 individuals) scores for further genetic analysis with the RAPD marker. The selected individuals are presented in Table 4, which shows that the trees classified into the high scoring population are derive from the families 1, 2, 4, 7, 8, and 9, while the trees classified into the low scoring populations are derive from the families 1, 2, 4, 7, 8, and 9. This pattern also indicates that the mother trees from families 1, 7, 8, and 9 could produce offspring with different phenotypic performances, which may be due to outcrossing and heterozygosity.

**Genetic diversity** Genetic analysis of sengon from Solomon provenance with RAPD markers was carried out on individual trees that had been selected through phenotypic quality scoring system. The five RAPD primers used in this study showed good amplification pattern and presented a total of 63 loci (Figure 3).

The calculated genetic diversity parameters of the 2-years old studied sengon population were the number of alleles expected (Na), the number of alleles observed (Ne), gene diversity (He) and percent polymorphic loci (PLP). The overall results of RAPD analysis of studied sengon population in this study showed that the expected average number of alleles (Na) was 1.9683, the average number of alleles observed (Ne) was 1.4523, the mean value of genetic diversity (He) was 0.2813, and the percent polymorphic locus (PLP) was 97% (Table 5). The mean value of genetic diversity in these 2-years old studied sengon population provides an implicit overview of the genetic diversity of the nine parent tree families that grow in Pompok Landak Permanent Nursery. In the present study, the mean genetic diversity (He) of sengon from Solomon provenance was slightly higher when compared to the genetic diversity of the common sengon population (He = 0.2757) in Cianjur community forest and the genetic diversity of the sengon population (He = 0.2349) in nine community forests in Java Island reported by Siregar and Olivia (2012) who also used RAPD markers. In addition, this study also had a slightly

### Table 3  The results of the phenotypic quality score for 2-years old sengon (from Solomon provenance)

| Block | 1   | 2   | 3   | 4   | 5   | 6   | 7   | 8   | 9   | Average | Rank | Survival rate (%) |
|-------|-----|-----|-----|-----|-----|-----|-----|-----|-----|---------|------|-------------------|
| 1     | 41  | 35  | 41  | 30  | 0   | 59  | 71  | 57  | 36  | 55.50   | 2    | 100               |
| 2     | 83  | 40  | 55  | 0   | 53  | 47  | 0   | 36  | 0   | 47.50   | 3    | 100               |
| 3     | 59  | 55  | 76  | 0   | 68  | 76  | 50  | 40  | 73  | 57.25   | 4    | 100               |
| 4     | 39  | 60  | 57  | 0   | 49  | 0   | 0   | 60  | 0   | 57.25   | 8    | 100               |

### Table 4  List of sengon (from Solomon provenance) samples used for further RAPD analysis

| No. | Family | Block | Tree code | No. | Family | Block | Tree code |
|-----|--------|-------|-----------|-----|--------|-------|-----------|
| 1   | 1      | 2     | S2B1      | 1   | 7      | 1     | S7B1      |
| 2   | 4      | 1     | S4B1      | 2   | 1      | 2     | S1B2      |
| 3   | 9      | 1     | S9B1      | 3   | 3      | 3     | S3B3      |
| 4   | 2      | 2     | S2B2      | 4   | 5      | 3     | S5B3      |
| 5   | 8      | 2     | S8B2      | 5   | 6      | 3     | S6B3      |
| 6   | 7      | 3     | S7B3      | 6   | 9      | 3     | S9B3      |
| 7   | 8      | 3     | S8B3      | 7   | 8      | 4     | S8B4      |

Note: S = sengon tree; B = block; example: S1B1 = sengon tree from family 1 in block 1
Table 5 Summary of genetic diversity in 2-years old *sengon* (from Solomon provenance) populations

| Population | N  | Na (SD)         | Ne (SD)         | PLP (%) | He (SD) |
|------------|----|-----------------|-----------------|---------|---------|
| High score | 7  | 1.7143 (0.4554) | 1.3762 (0.3219) | 71      | 0.2345 (0.1725) |
| Low score  | 8  | 1.7460 (0.4388) | 1.4181 (0.3418) | 75      | 0.2535 (0.1780) |
| Overall    | 15 | 1.9683 (0.1767) | 1.4523 (0.3101) | 97      | 0.2813 (0.1455) |

Note: N = sample size; Na = number of alleles observed; Ne = effective allele number; He = Nei's gene diversity (Nei, 1973); and PLP = percentage of polymorphic loci. Values in parentheses are standard deviations.

Figure 3 The example of *sengon* from Solomon provenance DNA profile using RAPD primers on 7 high-score trees and 8 low-score trees (the sample order is based on Table 4). (a) The amplified DNA of *sengon* using primer OPA-05, (b) The amplified DNA of *sengon* using primer OPA-14, and (c) The amplified DNA of *sengon* using primer OPY-16.
higher mean of genetic diversity than the sengon half-sib progeny population (He = 0.244) analyzed with the RAPD marker by Rostini et al. (2018) and genetic diversity of healthy-growing sengon populations (He = 0.262) in various locations on the Java Island which was analyzed by Lelana et al. (2018) using the RAPD marker. This pattern indicated that sufficient of the diversity source in sengon from Solomon provenance that grows in Pompos Landak Permanent Nursery-Haurwangi Village at this time.

In this study, genetic diversity analysis was partitioned between populations that had high phenotypic parameter scores and the other populations that had low phenotypic parameter scores (Table 5). The results of the analysis showed that the studied sengon population with low scores had a slightly greater mean value of genetic diversity parameters (\( Na = 1.7460, Ne = 1.4181, \) PLP 75% and He = 0.2535) than the studied sengon population with high scores (\( Na = 1.7143, Ne = 1.3762, \) PLP 71% and He = 0.2345), indicating the genetic diversity of the sengon from Solomon provenance having low-score population is greater when compared to the high-score population genetic diversity. The small genetic diversity in the high-score sengon population is expected because a selected population that has superior traits would have smaller diversity. Smaller genetics diversity means the population is getting more uniform, and if these trees will be used as plus tree in the seed orchard, the resulting offspring will have more uniform phenotypic traits as their parents.

**Genetic distance**

The differences in genetic diversity between populations indicates the genetic distance between the two Solomon sengon populations. The result of the genetic distance analysis between a high-score and low-score populations (Table 6) showed that the high-score tree from family 1 in block 2 (S1B2) and low-score from family 4 in block 1 (S4B1) and also the low-score tree from family 1 in block 4 (S1B4) and low-score from family 4 in block 1 (S4B1) have the highest genetic distance value (\( d = 0.6774, \) respectively) indicated that family 1 have a distant relationship with the low-score tree from family 4. Meanwhile, the low-score trees from family 8 in block 3 (S8B3) and family 7 in block 3 (S7B3) have the lowest genetic distance value (\( d = 0.1358 \)) indicated close relationship between the two low-score families. The results of present study also revealed that the offspring derived from the same population or family do not always have close genetic distances or are in the same group (Widyastuti, 2007). For example, in this study (Table 6), it was found that the genetic distance between the studied sengon trees that belonged to the same family (family 9), but differed in the phenotypic trait scores, namely S9B3 (high score) and S9B1 (low score), had a large genetic distance (\( d = 0.4055 \)). This indicates that family 9 comes from the mother tree that might be heterozygous and self crossed or might be underwent outcrossed.

In addition, the comparison of genetic distance between high- and low-score individuals can also be described from the percentage of individuals in each genetic distance range (Table 7), where the results showed that 36.3% of the high-score-low score individuals have large genetic distance (range = 0.5861–0.7361) and this was the highest percentage when compared to the percentage between individuals with the highest score (0%) or between individuals with the lowest score (3.7%) indicating high and low-score individual have more genetic distance. Moreover, 38.10% of individuals with the highest scores have a close genetic distance (range = 0.158–0.2858) and this is the highest percentage compared to individuals with low scores or the high-low score individuals, indicating a close relationship between high score individuals. Furthermore, significant genetic differences (\( p \text{-value} <0.001 \)) were detected in the analysis of molecular variance (AMOVA) among high and low-score

| Population | S7B1 | S1B2 | S3B3 | S5B3 | S6B3 | S9B3 | S8B4 | S2B1 | S4B1 | S9B1 | S2B2 | S8B2 | S7B3 | S8B3 | S1B4 |
|------------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|
| **High score** | **** | 0.3365 | 0.2311 | 0.3589 | **** | 0.2719 | 0.4055 | 0.2513 | 0.2930 | **** | 0.2311 | 0.4543 | 0.3365 | 0.4796 | 0.4296 | 0.1919 | 0.4055 | 0.2513 | 0.4296 | 0.3365 | 0.1728 | **** |
| **Low score** | 0.2113 | 0.2113 | 0.3145 | 0.4296 | 0.2930 | 0.2113 | 0.4543 | 0.4296 | 0.5055 | 0.4055 | 0.2113 | 0.3589 | 0.3365 | 0.4796 | 0.5055 | 0.3365 | 0.4296 | 0.4296 | 0.4926 | 0.3365 | 0.1728 | **** |

Note: S = sengon tree; B = block; example: S1B1 = sengon tree from family 1 in block 1
population (Table 8). AMOVA analysis comparing high and low-score population also revealed that 11% of the molecular variance existed between the population, whereas 89% was within population (Table 8).

The calculated genetic distance obtained in the present study were then used to construct a dendrogram based on UPGMA (Figure 4), which clustered the individuals and families studied. The dendrogram showed that high-score

### Table 7 Summary of sengon individual percentage based on mean genetic distance range

| Phenotypic score category | Mean genetic distance range | % Individu |
|--------------------------|----------------------------|------------|
| High score–High score    | 0.1358–0.2858              | 38.10      |
|                         | 0.2859–0.4359              | 52.38      |
|                         | 0.4360–0.5860              | 9.52       |
|                         | 0.5861–0.7361              | 0.00       |
| High score–Low score     | 0.1358–0.2858              | 21.43      |
|                         | 0.2859–0.4359              | 46.43      |
|                         | 0.4360–0.5860              | 26.79      |
|                         | 0.5861–0.7361              | 5.36       |
| Low score–Low score      | 0.1358–0.2858              | 32.14      |
|                         | 0.2859–0.4359              | 42.86      |
|                         | 0.4360–0.5860              | 21.43      |
|                         | 0.5861–0.7361              | 3.57       |

Note: df = degree of freedom, φPT = recorded population genetic differentiation, and an asterisk indicates significance (p-value < 0.001).

### Table 8 AMOVA table of molecular variation among and within high and low-score sengon populations from GenAlEx

| Source            | df | Sum of squares | Mean square | Estimated varian | % | φPT       |
|-------------------|----|----------------|-------------|------------------|---|-----------|
| Among population  | 1  | 17.415         | 17.415      | 1.142            | 11% | 0.114*** |
| Within population | 13 | 115.518        | 8.886       | 8.886            | 89% |           |
| Total             | 14 | 132.933        | 10.028      |                  | 100%|           |

Note: df = degree of freedom, φPT = recorded population genetic differentiation, and an asterisk indicates significance (p-value < 0.001).

Figure 4 UPGMA dendrogram of genetic distance of sengon (from Solomon provenance) families based on RAPD profiles.
tree populations tended to cluster together and do not merge with low-score tree populations. Likewise, the low-score tree population tended to form a separate cluster with the high-score tree population. This pattern suggested that even though there is no evidence that certain RAPD marker was linked or associated with the phenotypic characters observed, the RAPD marker used in this study has clustered individuals with high-scored phenotypes separately with lower scored ones, thus the selection based on phenotypes had succeeded in pooling the good quantitative alleles in the selected population.

Conclusion
The present study revealed that the family 3 has the highest average score of phenotypic quality (57.25 points) and family 4 has the lowest average score (7.50 points). This evaluation results can be used as a reference in determining the best families to produce superior sengon (from Solomon provenance) offspring in the future that have desired adaptability, productivity and diversity, as seen from the performance of the offspring. Furthermore, the medium level of genetic diversity (He = 0.2813) for the total sengon (from Solomon provenance) population in this study suggested the sufficient diversity source in sengon from Solomon provenance that grows in Pompok Landak Permanent Nursery. Besides that, the genetic diversity of the sengon from Solomon provenance having low-score population was slightly greater (He = 0.2535) when compared to the high-score population genetic diversity (He = 0.2345) indicated that the sengon mother tree experienced outcrossing. In addition, the comparison of genetic distance between high- and low-score individuals showed that high and low-score individual have more larger genetic distance then the high-score individuals or the low-score individuals. This pattern supported by AMOVA results that showed significant genetic differences (p-value <0.001) among high and low-score population and also UPGMA dendrogram of genetic distance which illustrated high-score tree populations tended to cluster together and do not merge with low-score tree populations, thus the selection based on phenotypes had succeeded in pooling the good quantitative alleles in the selected population. Clustering together of individuals having similar superior phenotypic traits against those having non-superior traits indicated that those individuals have distinctive different genotype.

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