Genetic diversity and population structure of *Eurycoma apiculata* in Eastern Sumatra, Indonesia

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**Abstract.** Zulfahmi, Parjanto, Purwanto E, Yunus A. 2021. Genetic diversity and population structure of *Eurycoma apiculata* in Eastern Sumatra, Indonesia. Biodiversitas 22: 4431-4439. Information on genetic variation within and among populations of *Eurycoma apiculata* plants is important to develop strategies for their conservation, sustainable use, and genetic improvement. To date, no information on genetic variation within and among populations of the *E. apiculata* has been reported. This study aims to assess genetic diversity within and among populations of *E. apiculata* based on RAPD markers, and to determine populations to collect *E. apiculata* genetic material for conservation and breeding programs. Young leaves of *E. apiculata* were collected from six natural populations. Fifteen RAPD primers were used to assess the genetic diversity of each population. The data obtained were analyzed with POPGEN and Arlequin software. The amplification results of 15 selected primers produced 3-16 loci with all primers 100% polymorphic. At the species level, the mean allele per locus (Na), number of effective alleles (Ne), percentage of polymorphic loci (PPL), Nei’s gene diversity index (He) and Shannon information index (I) were 2,000, 1,244, 100%, 0.167, and 0.286, respectively. At the population level, the mean values for Na, Ne, PPL, He and I were 1.393, 1.312, 39.27%, 0.119, and 0.186, respectively. The highest value of gene diversity within population (He) was found in the Lingga-1 population and the lowest value was found in the Rumbio population. The value of genetic differentiation among populations (Gst) of *E. apiculata* is 0.284, consistent with the results of the AMOVA analysis which found that genetic variation among populations was 23.14%, indicates that the genetic variation of *E. apiculata* was more stored within populations than among populations. The gene flow (Nm) value of *E. apiculata* was 1.259 migrants per generation among populations. The Nm value of this species was high category, and could inhibit genetic differentiation among populations. The clustering of *E. apiculata* population based on the UPGMA dendrogram and PCA was inconsistent with its geographic distribution, reflecting the possibility that genes migration occurred between islands in the past. The main finding of this study was the genetic variation of the *E. apiculata* mostly stored within the population. Therefore, the population with the highest genetic diversity is a priority for in-situ conservation, and collection of *E. apiculata* genetic material for ex-situ conservation and breeding programs should be carried out minimum from Lingga-1 and Pokomo populations.

**Keywords:** Pasak bumi, genetic variation, gene flow, conservation and breeding strategy

**INTRODUCTION**

*Eurycoma apiculata*, A.W. Benn is a member of the Simaroubaceae family. *E. apiculata* is only grown in Sumatra island and Malaysian Peninsular in the primary and secondary forest, as well as sandy and acid soils of tropical forest (Nooteboom 1962; Padua et al. 1999; Zulfahmi et al. 2019a). Specifically, existing of *E. apiculata* in Sumatra island has been reported by Zulfahmi et al. (2019a) and Zulfahmi et al. (2020) in the Riau Province and Riau Islands Province. *E. apiculata* is one of the pivotal medicinal plants to develop in the future as the source of herbal medicine. Traditionally, the root extract of the *E. apiculata* is used as a drink to tonic, diarrhea, febrifuge, as well as to decline the boneaches whereas its leaves decoction is used to decline the skin itchiness (Nooteboom 1962; Padua et al. 1999; Zulfahmi et al. 2019a).

Knowledge of genetic variation within and among plant populations is pivotal to develop strategies for optimal management of genetic resources for conservation, sustainable use, and genetic improvement (Medhi et al. 2014; Saini et al. 2018). Besides, genetic variation is needed for plants to adapt to environmental conditions, mainly facing climate change running. Information on genetic diversity within and among populations of *E. apiculata* has not been reported yet. Meanwhile, the natural population of the *E. apiculata* continues to experience degradation due to forest fires and various exploitative human activities, consequently, this species has been established by the Indonesian government as protected species based on the regulation number of P.20/MENLHK/SETJEN/KUM.1/6/2018. Therefore, an assessment of the genetic diversity of *E. apiculata* is urgent to be implemented and this scientific report is expected to become a consideration in the compilation of strategy conservation of this species. The eastern Sumatra region is the epicenter of the spreading of *E. apiculata* reported in Sumatra (Zulfahmi et al. 2018; Zulfahmi et al. 2019a; Zulfahmi et al. 2019b; Zulfahmi et al. 2020) so this area can be considered as a target for *E. apiculata* conservation areas in the future.
Information on the diversity of *E. apiculata* based on morphological markers has been carried out by Zulfahmi et al. (2019b) and Zulfahmi et al. (2020), but the diversity information obtained is not accurate enough to be considered in developing management and breeding strategies for *E. apiculata* plants because morphological markers are strongly influenced by environmental factors and plant growth (López-Caamal and Tovar-Sánchez 2014; Nadeem et al. 2018; Uslan & Pharmawati 2020). To overcome the weaknesses of this morphological marker, analysis of diversity using DNA molecular markers is necessary.

DNA-based molecular markers are often used to determine plant genetic diversity. One of the DNA-based markers was Random Amplified Polymorphic DNA (RAPD) (Williams et al. 1990). This marker allows us to obtain large amounts of data on genetic variation within and between populations without prior detailed knowledge of DNA sequences, the number of primers practical unlimited that can be used to provide information about variation across the genome, as well as relatively cheap, fast, and easy compared to other DNA markers (Williams et al. 1990; Weising 2005; Kumari 2014; Dhutmal et al. 2018). RAPD markers have weaknesses, namely low reproductive ability (low reproducibility) and dominant properties (Weising 2005), but it can be overcome through improved laboratory techniques, scoring procedures, and the use of analysis of molecular variance (AMOVA) (Nybom 2004; Weising 2005; Excoffier et al. 2007). Although RAPD has several drawbacks, the use of RAPD markers in population genetic diversity studies has been popular in various of plants species such as *Plumbago zeylanica* (Panda et al. 2015), *Pinus merkusii* (Tuong et al. 2016), *Cassia tora* (Tilwari et al. 2016), *Panax ginseng* (Wang et al. 2016), Aloe species (Adienge et al. 2019), *Silybum marianum* (L.) Gaertn (Hamouda 2019). This study aims to assess genetic diversity within and among populations of pasak bumi (*E. apiculata*) based on the Random Amplified Polymorphic DNA (RAPD) marker, and to determine populations to collect *E. apiculata* genetic material for conservation and breeding programs.

**MATERIALS AND METHODS**

**Sample collection**

Young leaves of *E. apiculata* were taken from six natural populations in Riau and Riau Islands (Eastern Sumatra), Indonesia as shown in figure 1. The longitude and latitude positions, as well as status of each population was displayed in Table 1. The number of samples per population was five individuals. Field distance between individuals collected was at least 20 m. The collected leaves are put into a plastic bag that has given silica gel with a ratio of leaves and silica gel was 1:5 (w/w). Silica gel served to reduce the water contents of the leaves and prevent the samples from being attacked by fungi. The samples were sent to the laboratory and stored in a freezer at -20 °C until DNA extraction was carried out.

![Figure 1. Location of sample collection Eurycoma apiculata in Riau, Indonesia](image-url)
DNA extraction

Genomic DNA of *E. apiculata* was isolated from leaf tissue using CTAB (cetyltrimethyl ammonium bromide). The isolation procedure followed the method of Doyle and Doyle (1990) with slight modifications (using 0.20% PVP and 1.0% mercaptoethanol). The quality DNA was determined by electrophoresis on agarose gel with ethidium bromide. Electrophoresis was carried out using 1x TAE solution (Tris Acetate EDTA) for 45 minutes at a voltage of 120 volts. The DNA banding patterns were observed under ultraviolet (UV) light and the gel documentation was performed using GelDoc (BioRad). The extracted DNA was stored in a freezer at -20 °C until PCR amplification was carried out.

Primer selection and DNA amplification

Thirty-six (36) random primers were tested and selected for DNA amplification of *E. apiculata*. Two DNA samples were mixed (bulk), and then used as samples for primer selection. Fifteen (15) primers that gave high polymorphism will be selected and used for DNA amplification of all samples. The PCR machine was set up as follows: initial denaturation for 5 minutes at 95 °C, then followed by 39 cycles with denaturation for 1 minute at 94 °C, annealing for 1 minute at 37 °C, extension for 1 minute at 72 °C, and final extension for 10 minutes at 72 °C. The total volume of the PCR reaction was 10 μl, consisting of 1.30 μl of template DNA (5-10 ng), 1.00 μl of primer (5 pmol/μl), 2.70 μl of free RNase water, and 5.00 μl of HotStar Taq Master Mix (Qiagen).

The PCR amplification results were separated by electrophoresis on agarose gel with agarose concentrations of 1.50% (w/v) at a voltage of 120 volts for 45 minutes. The 100 bp DNA ladder (Vivantis) was also included in electrophoresis as a measurement standard or reference. The DNA banding patterns obtained in agarose were observed under ultraviolet (UV) light and documented using the GelDoc system (BioRad). Band patterns analysis was performed using Image Lab software (BioRad) version 2.0.1.

Data analysis

The band patterns obtained from PCR amplification of each sample were scored with value of 1 for band present and 0 for band absent. The scoring results are compiled as binary data to be analyzed with software. The calculated genetic parameters included average number of alleles per locus (\(N_a\)), Number of effective alleles per locus (\(N_e\)), percentage of polymorphic loci (\(PPL\)), Nei’s gene diversity (\(H_s\)), Shannon’s information index (\(I\)), total gene diversity (\(H_T\)), gene diversity within population (\(H_S\)), coefficient of genetic differentiation among populations (\(G_{ST} = [H_s - H_s]/H_T\)), and gene flow among populations (\(N_m = [1 - G_{ST}]/4G_{ST}\)). All of these parameters were calculated using the POPGEN Software Version 32 (Yeh et al. 1999). Molecular analysis of variance (AMOVA) was also performed to estimate component variation among populations and within populations using ARLEQUIN software version 3.01 (Excoffier et al. 2007). Dendrogram of UPGMA (Unweighted Pair-Group Method Arithmetic Mean) was constructed using NTSYSpc Version 2.00 Software (Rohlf 1998) and principal component analysis (PCA) was performed using SAS software version 9.01 (SAS Institute 2002).

RESULTS AND DISCUSSION

Primer polymorphism

Of the thirty-six (36) primers tested, 15 primers were selected due to high polymorphism and clear DNA banding as shown in Table 2. The number of bands generated ranging from 3-16 bands with the size of the DNA bands ranging from 200-2000 bp, depending on the type of primer used, and the plant genotype tested. Seven primers (X-01, OPY-16, P-08, OPY-15, OPD-03, OPY-19, OPJ-20, and D-11) produced a higher number of bands than the other primers.

The percentage of polymorphic bands of each primer is 100% which indicates that the polymorphism of the genomic DNA of this species is high. Fifteen selected primers resulted in a total number of DNA bands i.e. 132 bands, with an average of 8.8 bands per primer. This result was similar to those reported by Rosmaina and Zulfahmi (2013) in *E. longifolia* Jack (8.8 bands/primer), higher than those reported by Rosmaina et al. (2015) in *E. longifolia* Jack (2.5 bands/primer), but it was lower than that reported by Razi et al. (2013) on *E. longifolia* Jack plants in Malaysia (12.8 bands/primer). These differences were caused by the different types of primers used, the plant genotypes and the origin of the population studied.

| Table 1. Characteristic of *Eurycoma apiculata* research sites in Eastern Sumatra, Indonesia |
|-----------------------------|-----------------------------|-----------------------------|
| Population                  | Status of research sites    | Longitude                  | Latitude        |
| Pokomo, Kampar District, Riau Province | Protected forest | 100°57′9″ E | 0°15′7″ N |
| TAHURA, Siak District, Riau Province | Forest Park | 101°25′46″ E | 0°40′21″ N |
| Rumbio, Kampar District, Riau Province | Protected forest | 101°8′20″ E | 0°19′40″ N |
| Lingga-1, Lingga District, Riau Islands Province | Natural forest | 104°40′26″ E | 0°10′41″ S |
| Lingga-2, Lingga District, Riau Islands Province | Protected forest | 104°34′52″ E | 0°12′42″ S |
| Sentajo, Kuantan Singingi District, Riau Province | Protected forest | 101°34′2″ E | 0°31′37″ S |
Table 2. The selected primer and their sequences, the size and the number of amplified bands of *Eurycoma apiculata*

| Primer name | Sequences                  | Fragment size (bp) | Number of band | Number of fragment polymorphic | % fragment polymorphic |
|-------------|---------------------------|--------------------|----------------|-------------------------------|------------------------|
| X-01        | 5’CTCACCGTGC3’             | 300-2000           | 15             | 15                            | 100                    |
| OPY-16      | 5’GGGCCAATG3’              | 300-1200           | 9              | 9                             | 100                    |
| P-08        | 5’ACATCGCCCA3’             | 250-1500           | 12             | 12                            | 100                    |
| OPY-15      | 5’AGTCGCCCTT3’             | 300-1500           | 11             | 11                            | 100                    |
| Z-13        | 5’GACTAAGCC3’              | 400-1500           | 6              | 6                             | 100                    |
| OPD-03      | 5’GTGCGCATCA3’             | 350-1100           | 10             | 10                            | 100                    |
| OPY-19      | 5’TGAGGTCGCC3’             | 250-650            | 4              | 4                             | 100                    |
| OPY-08      | 5’AGCAGAGCA3’              | 250-900            | 8              | 8                             | 100                    |
| OPD-08      | 5’AGGCCCTACAG3’            | 400-750            | 5              | 5                             | 100                    |
| OPJ-20      | 5’AAGCGGCCCT3’             | 290-1200           | 16             | 16                            | 100                    |
| D-11        | 5’AGCGCCATTG3’             | 200-1500           | 16             | 16                            | 100                    |
| D-08        | 5’GTGTCGCCCA3’             | 350-800            | 8              | 8                             | 100                    |
| OPT-07      | 5’GCAGGCGTGT3’             | 250-950            | 6              | 6                             | 100                    |
| K-02        | 5’GTCTCCGCAA3’             | 1000-1300          | 3              | 3                             | 100                    |
| Mean        |                           |                    | 8.8            | 100                           | 100%                   |

Table 3. The genetic variation index of six populations of *Eurycoma apiculata*

| Population | Na  | Ne  | PPL (%) | He  | I   |
|------------|-----|-----|---------|-----|-----|
| Rumbio     | 1.371 | 1.168 | 37.12   | 0.107 | 0.169 |
| Pokomo     | 1.439 | 1.222 | 43.94   | 0.135 | 0.208 |
| Sentajo    | 1.386 | 1.900 | 38.64   | 0.181 | 0.184 |
| Tahura     | 1.303 | 1.195 | 30.30   | 0.112 | 0.166 |
| Lingga-1   | 1.508 | 1.210 | 50.76   | 0.139 | 0.222 |
| Lingga-2   | 1.349 | 1.174 | 34.85   | 0.108 | 0.167 |
| Mean population | 1.393 | 1.312 | 39.27   | 0.119 | 0.186 |
| Species level | 2.000 | 1.244 | 100.00  | 0.167 | 0.286 |

The PCR amplification results of *E. apiculata* with primers OPT-07, OPJ-20, and D-11 are shown in Figure 2. There are differences in the number and size of bands produced between individual samples as a result of several events, namely: i) insertion or small deletion of DNA strands that cause changes in the size of the amplification fragment, (ii) deletion occurred at the primer annealing site resulting in loss of fragments or increased fragment size, (iii) insertion of large DNA fragments between the primer annealing sites which exceeds the PCR capability so that no fragments are detected, (iv) nucleotide substitution at one or two primary target sites that affects the annealing process, which results in the presence or absence of polymorphisms or changes the size of the fragments (Weising et al. 2005).

Genetic diversity within populations

The number of alleles per locus (Na), number of effective alleles per locus (Ne), percentage of polymorphic loci (PPL), Nei’s gene diversity (He), and Shannon’s information index (I) of *E. apiculata* are shown in Table 3. At the species level, the Na, Ne, and PPL values were 2.00, 1.244, and 100%, respectively. The PPL value of *E. apiculata* in this study (100%) was higher than the PPL value of *E. longifolia* (18.50%) reported by Loc et al. (2016), and other medicinal plants such as *Nepeta kotschyi* Boiss (PPL = 30.80%) (Hadi et al. 2020), *Panax ginseng* (PPL = 78.90%) (Wang et al. 2016), and *Retama raetam* (PPL = 56.09%) (Abdellaoui et al. 2014).

The value of the Nei’s gene diversity (He) and Shannon’s information index (I) of *E. apiculata* species is 0.167 and 0.286, respectively. The genetic diversity values of *E. apiculata* in this study were lower than the genetic diversity values of the *E. longifolia* reported by Rosmaina and Zulfahmi (2013) (He = 0.29), Rosmaina et al. (2015) (He = 0.181), and cross-pollinated species (He = 0.27) (Nyborg and Bartish 2000; Nyborg 2004). This is closely related to differences in plant genotypes and species distribution. Species with a wide distribution have higher genetic diversity values than species with a narrow distribution (Levy et al. 2016; Chung et al. 2018; Li et al. 2020). *E. longifolia* has a wider distribution (covering Sumatra and Kalimantan) than *E. apiculata* which has a narrow distribution (only in Sumatra) (Nootenboom 1962; Zulfahmi et al. 2019). Moreover, these results are in agreement with our assumption that restricted distribution of *E. apiculata* and protected species will have lower genetic diversity compared to widely dispersed species. This is due to the influence of directional selection that encourages adaptation to the local environment, inbreeding, and genetic drift that occur in small populations (Gibson et al. 2008).
Figure 2. The results of PCR amplification of *Eurycoma apiculata* used primer OPT-07 [A], OPJ-20 [B], and D-11 [C]. M: DNA ladder, Rumbio population [1-5], Sentajo population [6-10], Pokomo population [11-14], Lingga-1 population [15-19], Tahura population [20-24], and Lingga-2 population [25-28].
At the population level, the number of alleles per locus (Na) and the percentage of polymorphic loci (PPL) of *E. apiculata* ranged from 1.303-1.508 and 30.30% – 50.76%, respectively, which the highest value was observed in the population of Lingga-1 and the lowest value was observed in the Tahura population. The number of effective alleles per locus (Ne) ranged from 1.168 (Rumbio population) to 1.222 (Pokomo population). The Nei’s gene diversity (He) values of the *E. apiculata* population ranged from 0.107-0.139 (Table 3). The highest value of gene diversity was observed in the Lingga-1 population and the lowest value was observed in the Rumbio population. Populations with high genetic diversity are valuable due to provide diverse gene pools for genetic conservation and plant breeding programs.

The low value of genetic diversity in the Rumbio population compared to other populations is due to a large number of conversions of Rumbio forest areas into rubber plantations by the surrounding community, as the consequences this population is fragmented. The results of this study are in line with those reported by Panda et al. (2015) on *Plumbago zeylanica* L. plants in which fragmented populations have lower genetic diversity values than other populations that are not fragmented. According to Azman et al. (2020) that populations that have low genetic diversity values indicate that these populations are in a condition of threatening, fragmented, and damaged by human activities. In contrast, The high genetic diversity in the Pokomo population could be caused by several factors, namely i) the genetic diversity has been high from the beginning of the population established, ii) the population has not been a lot disturbed by human activities, so its condition is more maintained, and iii) the occurrence of random mating among individuals resulting in genetic recombination and increasing genetic diversity within population.

If populations of *E. apiculata* are grouped according to regional distribution (Sumatra and Riau Islands), the average value of genetic diversity of the *E. apiculata* population from the Riau Islands region (Lingga-1 and Lingga-2) is 0.124, higher than the population genetic diversity of *E. apiculata* from Sumatra region (0.118). This result contradicts the general hypothesis that the plants genetic diversity of the mainland populations is higher than that of the island populations. But this result is in line with those reported by Garcia-Verdugo et al. (2015) on the *Periploca laevigata* plant who found that genetic diversity in the island populations was higher than in the mainland populations.

Genetic diversity is important for plants to adapt to changes in environmental conditions so that plants can survive for a long time. Lack of amount of genetic diversity will limit the ability of plants to cope with environmental changes and their role in the ecological and evolutionary development of the biosphere (Runo et al. 2004) so that the maintenance of genetic diversity is considered important as a carrier of diversity for ecological adaptation and microevolution.

### Genetic differentiation and population structure

The value of genetic differentiation among populations (*G_{st}*)) of *E. apiculata* was 0.284 (*H_{T} = 0.167, H_{S} = 0.120*), indicates that 28.40% of the total genetic diversity is stored among populations and 71.60% of the total genetic diversity existed within the population. The results of the analysis of molecular variance (AMOVA) also confirmed that the genetic diversity of *E. apiculata* is more stored within population than among populations, in which the percentage of genetic diversity among populations and within the population of *E. apiculata* is 23.14% and 76.86%, respectively (Table 4). AMOVA results showed a highly significant difference (*P < 0.001*) in genetic differentiation among populations. The high genetic diversity stored within the population is due to the fact that *E. apiculata* is a cross-pollinated species, which pollination is assisted by insects and bees (Zulfihami et al. 2020) so that random mating can occur among individuals and results in high genetic variability within the population. The high genetic diversity within population will be a consideration in developing a conservation strategy and selecting genetic

### Table 4. Results of molecular analysis of variants (AMOVA) of *Eurycoma apiculata*

| Source of variation | Degree of freedom | Sum of square | Variance component | Variation (%) | P value |
|---------------------|-------------------|--------------|--------------------|--------------|---------|
| Among populations   | 5                 | 152.833      | 3.673              | 23.14        | < 0.001 |
| Within population   | 24                | 292.800      | 12.200             | 76.86        |         |
| Total               | 29                | 445.633      |                    |              |         |

### Table 5. The value of the coefficient of genetic distance (below the diagonal) and genetic similarity (above the diagonal) of genetic similarity of *Eurycoma apiculata* is based on Nei 1978

| Population | Rumbio | Pokomo | Sentajo | Tahura | Lingga-1 | Lingga-2 |
|------------|--------|--------|---------|--------|----------|----------|
| Rumbio     | ****   | 0.9743 | 0.9511  | 0.9468 | 0.9794   | 0.9564   |
| Pokomo     | 0.0260 | ****   | 0.9668  | 0.9441 | 0.9486   | 0.9413   |
| Sentajo    | 0.0502 | 0.0337 | ****    | 0.9413 | 0.9242   | 0.9030   |
| Tahura     | 0.0547 | 0.0575 | 0.0604  | ****   | 0.9614   | 0.9338   |
| Lingga-1   | 0.0208 | 0.0328 | 0.0788  | 0.0394 | ****     | 0.9729   |
| Lingga-2   | 0.0446 | 0.0605 | 0.1020  | 0.0685 | 0.0275   | ****     |


material to build a breeding population. The $G_{ST}$ value of *E. apiculata* in this study was lower than the average $G_{ST}$ of the *E. longifolia* (0.31) reported by Rosmaina & Zulfahmi (2013), *Panax ginseng* plant ($G_{ST} = 0.430$) (Wang et al. 2016) and higher than cross-pollinated plants ($G_{ST} \leq 23\%$) (Nybom 2004) and other species such as *Pinus merkusii* ($G_{ST} = 0.186$) (Tuong et al. 2016), and *Retama raetam* ($G_{ST} = 0.260$) (Abdellaaoui et al. 2014).

According to Nei (1978), the $G_{ST}$ value can be grouped into three categories, low if the $G_{ST}$ value is <0.05, moderate if the $G_{ST}$ value is 0.05-0.15, and high if the $G_{ST}$ value is > 0.15. Based on this category, the $G_{ST}$ value of the *E. apiculata* in this study is the high category. The high $G_{ST}$ value indicates that gene flow among populations of *E. apiculata* through seeds and pollen is limited. The seed size of *E. apiculata* is relatively large and heavy so that the spread of seeds is only limited to the forest floor near the mother tree, reflected most of seedlings found around the mother tree. The gene flow of the *E. apiculata* through the pollen depends on the pollinator (bees and beetles). Rader et al. (2011) reported the ability of beetles carried pollen of *Brassica rapa* L as far as 400 from a pollen source while Tani et al. (2009) reported that pollen dispersal distance of *Shorea parvifolia* Dyer and *Shorea leprosula* Miq is about 250-450 m and more than 700 m, respectively, from the mother tree in the tropical forests of Peninsular Malaysia.

Gene flow is one of the important parameters that determine plant genetic diversity. Gene flow is the transfer of genes within and among populations. This gene flow is determined by pollinators, seed dispersers, stand density, flowering phenology, plant sex distribution, outcrossing rate and inbreeding depression (Dick et al. 2008). The gene flow value ($Nm$) of *E. apiculata* in this study was 1.259 migrants per generation among populations. According to Govindaraju (1989) that the value of gene flow can be categorized into three levels, namely low if $Nm <0.25$, moderate if $0.25 < Nm < 0.99$, and high if $Nm > 1$. Based on these categories, the value of gene flow *E. apiculata* is the high category. The gene flow value of this study was higher than that reported by Rosmaina and Zulfahmi (2013) on *E. longifolia* ($Nm = 1.11$) and other medicinal plants such as *Zanthoxylum* spp ($Nm = 1.31$) (Medhi et al. 2014), *Retama raetam* ($Nm = 1.42$) (Abdellaaoui et al. 2014).

Theoretically, a high $Nm$ value is considered sufficient to inhibit genetic drift and prevent genetic differentiation between populations, whereas $Nm$ value < 1 is not sufficient to counteract the effects of genetic drift, which is the dominant factor causing genetic differentiation between populations (Li et al. 2018). Based on the value of gene flow in this study, genetic drift has not been the dominant factor influencing the genetic structure of the populations of the *E. apiculata*. However, the population of *E. apiculata* is currently in a threatening condition due to habitat fragmentation which will slowly affect their genetic structure. High values of $Nm$ between populations of *E. apiculata* also indicate that geographic barriers do not significantly affect gene flow, but geographic distances may influence genetic relationships.

Values of genetic similarity and genetic distances between populations of *E. apiculata* are shown in Table 5. The value of genetic similarity between populations of *E. apiculata* ranged from 0.9030 - 0.9794, with the highest value of genetic similarity observed between the Lingga-1 population and the Rumbio population, while the lowest genetic similarity value was found between the Sentajo and Lingga-2 populations (Table 5). The Lingga-1 and Lingga-2 populations have closer genetic similarities to the Rumbio population than other populations from Sumatra.

The results of the UPGMA dendrogram based on genetic similarity Nei (1978) grouped the *E. apiculata* populations into two main groups at the 0.945 or 94.50% genetic similarity, namely the first group consisted of Pokomo and Sentajo populations, while the second group consisted of Tahura, Lingga-2, Rumbio, and Lingga-1 populations. At the genetic similarity of 0.96 (96.00%), the second group was divided into two sub-groups, namely the first sub-group consisted of the Tahura population, the second sub-group consisted of the Lingga-2, Rumbio, and Lingga-1 populations (Figure 3).

![Figure 3](attachment:image3.png) **Figure 3.** UPGMA dendrogram for the population of *E. apiculata* based on the genetic similarity value of Nei 1978

![Figure 4](attachment:image4.png) **Figure 4.** Scatter plot *Eurycoma apiculata* population based on PCA
PCA analysis was also performed to better understand the genetic structure of the population of species. The PCA results of *E. apiculata* showed that the first and second main components (PC-1 and PC-2) explained the cumulative variation percentage of 63.64% of the total variation, in which the first main component (PC-1) explained 35.71% of the total variation with an eigenvalue of 2.67 while the second main component (PC-2) explained 27.94% of the total variation with an eigenvalue of 2.09. According to Banda & Kumarasamy (2020), the acceptable threshold value for the percentage of the cumulative variation of PCA is greater than 60%. The percentage of cumulative variation of PC1- and PC2 of *E. apiculata* (63.64%) in this study was above this threshold and was higher than the percentage of cumulative variation of PC-1 and PC-2 with RAPD markers reported by Dillipan et al. (2017) on seaweed (58.55%). Scattered plot PC-1 and PC-2 of PCA grouped the population of *E. apiculata* into four groups, namely the first group was the population of Pokomo and Sentajo, the second group was the Tahuara population, the third group was the Rumbio population, and the fourth group was the Lingga-1 population and the population Linga-2 (Figure 4).

The results of the UPGMA dendrogram and PCA scatter plot of *E. apiculata* exhibited that population clustering does not reflect the geographic distribution of the species. Population groupings not reflecting geographic distribution reported by Zulfahmi et al. (2015) on Meranti species (*Shorea* spp.), Zulfahmi et al. (2020) on *E. apiculata*, and Zulfahmi et al. (2021) on *E. longifolia*. A mixture of population from the Riau Islands and the population from the island of Sumatra in one group indicated the genetic flow of the *E. apiculata* between the Sumatra island (Rumbio) and the Riau Islands (Lingga-1 and Lingga-2). This genetic flow probably occurred during the glacial period, in which the islands of Sumatra and Riau Islands were still joined which were connected by a stretch of savanna forest (Slik et al. 2011; Wurster et al. 2019).

In conclusion, the fundamental genetic information regarding the natural population of *E. apiculata* was obtained using RAPD marker in this study, in which the mean value of genetic diversity within the population was 0.120, the value of genetic differentiation among populations (*G_{st}*). 0.284, as well as geographic patterns of genetic variation among populations, were also detected. These genetic findings have important implications for conservation and breeding programs. The Lingga-1 and Pokomo populations can be selected as a target for the collection of *E. apiculata* genetic material for ex-situ conservation and breeding programs. Finally, this study suggests using molecular markers with high variability in future studies to obtain detailed genetic information that more facilitates the conservation and management of genetic resources of *E. apiculata*.

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