Evaluation of *Pulasan (Nephelium ramboutan-ake)* Genetic Diversity in Bogor, West Java, Using Microsatellite Markers

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ABSTRACT

Pulasan (Nephelium ramboutan-ake (Labill.) Leenh) fruit is highly similar to *rambutan rapiah* (*Nephelium lap-paceum*) fruit with ovate shape, sweet and sour fresh taste, but it has a thick rind. The diversity of *pulasan* is little informed including in Bogor. The objective of this study was to analyze the genetic diversity of *pulasan* from Bogor revealed by microsatellite marker. The DNA of 63 individuals from 10 populations of *pulasan* were extracted using CTAB method and amplified using two primer sets, LMLY6 (GA)9(CA)2(GA)4 dan LMLY12 (CT)11. DNA amplification product was visualized and arranged in a matrix of binary data then analyzed the value of the number of different alleles (Na), the number of effective alleles (Ne), Shannon information index (I), heterozygosity (He), and the percentage of polymorphism (PLP). The results of the analysis showed the highest genetic diversity was found in North Bogor (He=0.313). The genetic diversity within a population (61%) was higher than that among populations (39%). A dendrogram was constructed using the Unweighted Pair Group Method with arithmetic Mean (UPGMA). The similarity index ranged from 52 to 100% that means there are close relationships among individuals. Cluster analyses grouped some individuals originated from different locations in the same group. The levels of heterozygosity within a population was determined by the history of each individual in a population.

Keywords: Bogor, genetic diversity, Nephelium ramboutan-ake, microsatellite

INTRODUCTION

In Java, *pulasan* is known as rambutan babat, rambutan ake, and rambutan leci. The plant has leaves and branches that similar to those of rambutan, but it has a smaller size. *pulasan* fruit is like rambutan, ovate shape with a sweet taste and a slightly sour, but it has a thick rind and with no hairs. *Pulasan* has many uses. It has quite a hard stem, so it is often used for making household appliances. The seeds contain vegetable oils that can be used to make candles and soaps [1]. In addition, the seeds can be eaten after baked, and the taste likes beans. The baked seeds can be made into a powder and used it like cocoa powder. *Pulasan* fruit can be eaten directly like rambutan or as an additional ingredient in ice cream, pudding, jam, syrup and mixed in drinks such as cocktails [2].

Pulasan is a native species of Java, Borneo, and the

Microsatellite marker can be used to describe the genetic diversity of *pulasan*. This marker is often used to describe the diversity of organisms at the species level because its existence is abundant, it produces high polymorphisms, and co-dominant marker [4]. Microsatellite markers had been successfully used to describe the genetic diversity at the species level, for example in rice [5], sorghum [6], and apple [7]. This study aimed to analyze the genetic diversity of *pulasan*

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Philippines [3]. Unfortunately, it is now hard to find in Java Island since people's interest to grow *pulasan* tree decreases due to its low production. *Pulasan* given its many benefits, and has huge potential for its development as an alternative fruit in the future, however, its diversity in Java island, especially in Bogor, has not been identified. Therefore, it is important to evaluate its genetic diversity.

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Primer	Forward	Reverse
LMLY6	AAGGAATAAAGCTATCAATAAA	GATCTCTATCTCATCAAACCT
LMLY12	GAAGCTGTCTTAACACTCCAC	ACAAACCTAGAAACCAAAAG

in Bogor, West Java, using microsatellite markers.

MATERIALS AND METHODS

Sampling

pulasan leaves were taken from four districts in Bogor Municipality (West Bogor Barat, North Bogor, South Bogor and Central Bogor) and six districts in Bogor Regency (Cileungsi, Sukaraja, Babakan Madang, Ciomas, Parung, and Cibinong). Total samples of the plant taken are 63 trees. From each tree, as many as four to five mature leaves were taken. The leaves were put into plastic bags, added with silica gel, covered, and stored in a -20°C temperature.

DNA isolation, Polymerase Chain Reaction (PCR) and visualization of PCR results

Isolation of DNA used the CTAB method [8] with some modifications. PCR amplification method was based on the method conducted by Sim et al. [9] using two pairs of primers, namely LMLY 6 and LMLY 12 (Table 1).

Amplification process was carried out in 25 μ L mix solution contained 12.5 μ L of Go Taq Green master mix, 1 μ L each of primer forward and reverse, 0.25 μ L BSA, 2 μ L of DNA, and 8.25 μ L ddH₂O. This process was performed for 35 cycles consisting of several stages: initial denaturation at a temperature of 94°C for 35 seconds, denaturation at 94°C for minute, annealing at 46°C for the LMLY6 primer and 46.3°C for LMLY12 for 35 seconds, and elongation at a temperature of 72°C for 10 minutes. Amplification results were then examined using horizontal electrophoresis and used 2.5% of agarose gel for 100 minutes. Results of electrophoresis were visualized using UV light and then photographed.

Data analysis

The resulted visualization of PCR was observed by looking at the presence or absence of band considered microsatellite alleles on *pulasan* accession. When there was a band, then a score of one was given, and when there was none, a score of zero was given. The resulted scores were arranged in a matrix of binary data and then analyzed using the software GenAlex to generate the value of the number of different alleles (Na), the number of effective alleles (Ne), Shannon information index (I), heterozygosity (He), the percentage of polymorphism (PLP) and the analysis of molecular variation (AMOVA). The sample from the Parung District was not analyzed because only one sample was found. Polymorphism percentage value is calculated with the following formula:

$$\begin{split} N_e &= \frac{1}{p^2 + q^2} \\ I &= -(p \times Ln(p) + q \times Ln(q)) \\ H_e &= 2 \times p \times q \end{split}$$

With the $q = \sqrt{frequency of band absence}$ and p = 1 - q according to Hardy-Weinberg equilibrium [9]. The dendrogram was constructed using the Jaccard coefficient and Unweighted Pair Group Method with Arithmatic Mean (UPGMA), it performed using the software NTSYS-pc ver 2.1 [10].

RESULTS AND DISCUSSION

Pulasan exploration

Based on the resulted exploration in Bogor Regency and Municipality, the study found *pulasan* plants in ten districts (Table 2). *pulasan* trees found on all locations are 63 trees in number. The district with the highest number of trees is Cileungsi with as many as 13. The district with the least number of trees is Parung. The number of locations where *pulasan* trees found varies with each district.

Pulasan DNA amplification

All *pulasan* plants of Bogor have successfully been amplified using LMLY6 and LMLY12. LMLY6 Primer generated PCR products in the form of two DNA bands of 140 bp and 160 bp (Figure 1). The bands with the size of 160 bp were amplified in all plants, while the size of 140 bp only amplified in some plants. Thus, the 140 bp band shows variations in LMLY6 primer. Unlike LMLY6 primer, LMLY12 produced PCR products with 2 - 7 DNA bands per individual *pulasan* (Figure 2). The band sizes generated by LMLY12 were of 150 bp, 200 bp, 225 bp, 250 bp, 400 bp, 450 bp and 500 bp. Among the resulted PCR of LMLY12, two bands tend to be amplified in each plant, namely the sizes of 225 bp and 450 bp. While the other bands produced using LMLY12, show variation among indi-



Figure 1. DNA amplification of *pulasan* genome in several districts in Bogor using LMLY6 primer and an annealing temperature of 46.3°C on agarose gel of 2.5%. (M: DNA Marker 1000 bp; E: North Bogor; C: Ciomas.)



Figure 2. Variations in DNA amplification results of *pulasan* genome in several districts in Bogor using LMLY12 primer and an annealing temperature of 46°C on agarose gel of 2.5%. (M: DNA Marker 1000 bp; A: Cileungsi; D: Sukaraja; E: North Bogor; F: Babakan Madang; G: West Bogor; H: Central Bogor. I: Cibinong)

vidual *pulasan* tree. Thus, the higher level of polymorphism is generated by the LMLY12 primer.

Genetic diversity of pulasan

The analysis of genetic diversity on two microsatellite primers showed that *pulasan* in the District of North Bogor has the highest values of heterozygosity (He = 0313), Shannon information index (I = 0458), and percentage of polymorphism (77.78%) (Table 3). The lowest values of heterozygosity, Shannon information index and the percentage of polymorphism were found in Ciomas District. Cileungsi District is the district with the highest number of *pulasan* trees (13 trees). However, the genetic diversity value in the District is low (He = 0102, I = 0175, and PLP = 55.56). This value is lower than those of heterozygosity and polymorphism percentage in North Bogor District where only six trees are found. *pulasan* in Babakan Madang District is also the same as that of Cileungsi District, with a large population size but it has low genetic diversity. In contrast to those locations, *pulasan* found in Central Bogor and Cibinong districts are less than five trees, but they have high values of heterozygosity and polymorphism. *pulasan* is also found less than five trees in Ciomas, but the values of heterozygosity (He = 0.026) and polymorphism percentage (11 : 11%) of *pulasan* in Ciomas District are lower than those of Central Bogor District and Cibinong.

The average percentage of polymorphism of Bogor pulasan is 44.44%. A locus is categorized as polymorphic if the frequency of an allele is <99% [11]. Thus, all loci generated by Bogor pulasan is polymorph. Evaluation of the genetic diversity using microsatellite markers on annual plant been made. An observation of genetic diversity of apples coming from the Swiss Federal Research Staton, Switzerland and Philip Forsline at The National germplasm repository for Apple and Grape, USA, showed that the average of heterozygosity value was 0.78, which is categorized as very high [7]. In contrast to the observation on apples, the genetic diversity of woody plants in tropical forests was lower, He=0.149 [12]. Comparing to the previous studies, the average value of *pulasan* heterozygosity is quite high, He=0.161.

The higher the value of heterozygosity in a population, the higher the genetic variation in the population [13]. Pulasan population in North Bogor District has the highest genetic variation, whereas that of Ciomas District has the lowest genetic variation. According to Nei [14], one of the factors affecting the level of heterozygosity is the population size. The population in North Bogor District is classified low (N=6). However, Cileungsi District with the largest population size (N=13) also has a quite low value of heterozygosity. In contrast to both districts, a small population of Cibinong District (N=2) showed a high heterozygosity, He=0.230. The low heterozygosity of pulasan in Cileungsi district may occur because all samples were originated from the same location, Taman Buah Mekar Sari. All *pulasan* trees in this place were propagated by grafting from the experimental garden in Cipaku. Thus, all individuals are genetically alike. Whereas, pulasan individuals in the Districts of North Bogor and Cibinong *pulasan* came from different locations. Thus,

Districts	Location	Number of trees
West Bogor	Cibalagung	3
	Gunung Batu	1
	Curug Mekar	1
South Bogor	Cipaku field trial	9
Center Bogor	Panaragan	1
	Bogor Botanical Garden (KRB)	3
North Bogor	Cimahpar	2
	Makam Bodas	4
Babakan Madang	Cijayanti	11
Cibinong	Sukahati	1
	Nanggewer	1
Cileungsi	Taman Buah Mekar Sari	13
Ciomas	Kota Batu	4
Parung	Jabon Mekar	1
Sukaraja	Cikeas	8
Total		63

Table 2. Exploration locations and the number of trees found

Table 3. Genetic Diversity of pulasan in ten districts in Bogor

Populations	Ν	Na	Ne	Ι	He	PLP
West Bogor	5	1.444	1.387	0.319	0.218	55.56%
South Bogor	9	0.889	1.118	0.111	0.073	22.22%
Center Bogor	4	1.222	1.368	0.309	0.209	55.56%
North Bogor	6	1.778	1.541	0.458	0.313	77.78%
Babakan Mandang	11	1.333	1.334	0.260	0.180	44.44%
Cibinong	2	1.556	1.393	0.336	0.230	55.56%
Cileungsi	13	1.333	1.142	0.175	0.102	55.56%
Ciomas	4	0.778	1.034	0.044	0.026	11.11%
Sukaraja	8	0.778	1.179	0.141	0.099	22.22%
Average	6.889	1.235	1.277	0.239	0.161	44.44%

Note: N = number of samples; Na = number of different alleles; Ne = number of effective alleles; I = index of Shannon information; He = heterozygosity; PLP = percentage of polymorphisms

Table 4. Analysis of molecular variation (AMOVA) of *pulasan* populations

	df	SS	MS	Est. Var.	%	
Among populations	8	33.256	4.157	0.503	39%	
Within populations	53	42.486	0.802	0.802	61%	
Total	61	75.742		1.304	100%	

Note: df = degrees of freedom; SS = sum of squares; MS = middle square

in addition to the population size, two other possibilities are affecting the genetic diversity level of *pulasan*, the origin of the individuals making up the population and the multiplication through vegetative reproduction. According to the previous study, other factors may affect the value of heterozygosity within a population, such as mutation, reproductive and recombinant traits, random mating, and natural selection [15].

AMOVA results show that the variation in the population (61%) is higher than the variation between populations (39%) (Table 3). Variations in large populations can occur as a result of interbreeding (Hamrick



Figure 3. Dendrogram of *pulasan* in Bogor constructed by the Jaccard coefficient and UPGMA method based on microsatellite data. A1-13: Cileungsi; B1-9: South Bogor; C1-4: Ciomas; D1-8: Sukaraja; E1-6: North Bogor; F1-11: Babakan Madang; G1-6: West Bogor; H1-4: Center Bogor; I1-2: Cibinong; J1: Parung.

1989). In addition, a broader distribution of plants, a large size, and an adjacent location of populations will produce considerable variation in the population because it will facilitate the occurrence of gene flow [16].

Pulasan relationship

The degree of similarity in the 63 samples of *pulasan* varies from 0.52 to 1.00. In the similarity level of 68%, all of the samples split and form into three main groups. The first group is *pulasan* of the Districts: Cibinong, Babakan Madang, Ciomas, and North Bogor. The second group is *pulasan* of the Districts: Ciomas, South Bogor, North Bogor, Parung, West Bogor, Babakan Madang, Cileungsi, and Central Bogor. The third group is *pulasan* of the Districts: Cileungsi, Babakan Madang, North Bogor, West Bogor, South Bogor, Cibinong, Sukaraja, and Central Bogor.

The results of cluster analysis (Figure 3) shows that individual *pulasan* do not cluster based on the origin of the population but they are scattered randomly, except for the population of Sukaraja District. *pulasan* from this district is identical, with the similarity level of 100%. The same individual trees occur since all *pulasan* of Sukaraja comes from trees grafted from the oldest tree in the area. *pulasan* B3 from the Experimental Garden of Cipaku is grouped together with *pulasan* of Cileungsi District, indicating that it is the progenitor of *pulasan* in Taman Buah Mekar Sari, Cileungsi.

Pulasan Individual from Bogor Botanical Gardens (KRB) and Cibinong District are grouped into populations from different locations. This result can be explained since KRB is a garden of plant collection that gathers all plants from different places. *Pulasan* individuals of Cibinong is also found from various and distant locations.

CONCLUSION

As many as 63 *pulasan* trees were found in ten Districts and Municipality of Bogor. All *pulasan* trees have been successfully amplified using the primers LMLY6 and LMLY12 with the annealing temperature of 46.3° C and 46° C respectively. Primer LMLY6 produces two bands of 140 bp and 160 bp, while, Primer LMLY12 produced seven sizes of DNA bands, 150 bp, 200 bp, 225 bp, 250 bp, 400 bp, 450 bp and 500 bp. The highest genetic variation of *pulasan* population is found in North Bogor (He = 0.313). The percentage of variation within a population (61%) is higher than between populations, 39%. The level of genetic similarity between *pulasan* trees ranged 52 - 100%, which means that the relationship among *pulasan* individuals is quite close. The origins of the individuals that make up the population determine the level of the population heterozygosity.

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