Genetic variation in *Rhodomyrtus tomentosa* (Kemunting) populations from Malaysia as revealed by inter-simple sequence repeat markers

T.S. Hue¹, T.L. Abdullah¹, N.A.P. Abdullah² and U.R. Sinniah¹

¹Department of Crop Science, Faculty of Agriculture, Universiti Putra Malaysia, Serdang, Selangor, Malaysia
²Department of Crop Science, Faculty of Agriculture and Food Sciences, Universiti Putra Malaysia Bintulu Campus Sarawak, Jalan Nyabau, Bintulu, Sarawak, Malaysia

Corresponding author: T.L. Abdullah
E-mail: thohirah@upm.edu.my

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**ABSTRACT.** Kemunting (*Rhodomyrtus tomentosa*) from the Myrtaceae family, is native to Malaysia. It is widely used in traditional medicine to treat various illnesses and possesses significant antibacterial properties. In addition, it has great potential as ornamental in landscape design. Genetic variability studies are important for the rational management and conservation of genetic material. In the present study, inter-simple sequence repeat markers were used to assess the genetic diversity of 18 *R. tomentosa* populations collected from ten states of Peninsular Malaysia. The 11 primers selected generated 173 bands that ranged in size from 1.6 kb to 130 bp, which corresponded to an average of 15.73 bands per primer. Of these bands, 97.69% (169 in total) were polymorphic. High genetic diversity was documented at the species level (\(H_r = 0.2705\); \(I = 0.3973\); PPB = 97.69%) but there was a low diversity at population level (\(H_s =...\)
0.0073; I = 0.1085; PPB = 20.14%). The high level of genetic differentiation revealed by $G_{st}$ (73%) and analysis of molecular variance (63%), together with the limited gene flow among population ($N_m = 0.1851$), suggests that the populations examined are isolated. Results from an unweighted pair group method with arithmetic mean dendrogram and principal coordinate analysis clearly grouped the populations into two geographic groups. This clear grouping can also be demonstrated by the significant Mantel test ($r = 0.581, P = 0.001$). We recommend that all the *R. tomentosa* populations be preserved in conservation program.

**Key words:** Contracting population; Gene erosion; Molecular marker; Multi-usage; Native plant; Rose myrtle

**INTRODUCTION**

*Rhodomyrtus tomentosa* (Aiton) Hassk. of the Myrtaceae family, is an edible, sun-loving flowering shrub that can grow up to 2-3 m tall. Its common name is kemunting (Malay) and apart from Malaysia, it is distributed in southern China, Philippines, Japan, India, Malaysia and Indonesia (Chen, 1984). It is usually found growing at open, often degraded sandy sites, along the seashores and open banks of river (Latiff, 1992) where it is used as an indicator of acidic soil (Wei et al., 2009). Naturally, *R. tomentosa* is easily propagated through seed dispersal by frugivorous birds and ants (Wei et al., 2004).

*R. tomentosa* is appreciated for its high value in traditional medicine (Ong and Nordiana, 1999; Wei, 2006), as well as its huge potential as an antibacterial agent against a broad range of gram positive and pathogenic bacteria (Limsuwan et al., 2009). Its showy pinkish five-petal flowers are produced all year round and possess a high aesthetic value. Together with the pinkish blooms which turn white one day after blooming and the exquisite maroon traits of its flowers have made *R. tomentosa* a suitable candidate as an ornamental and landscaping plant (Zhao et al., 2006). The high content of carbohydrates, fat and energy of its juicy ripened dark purple berries serve to attract birds into the landscape (Liu and Deng, 1997), as well as a filling for making pies and jam (Latiff, 1992). The locals also consume its fruit raw due to the high mineral and vitamin content (Latiff, 1992).

*R. tomentosa* is still underutilized in Malaysia despite its multiple uses due to lack of exposure and documentation. Its population size in Malaysia has become severely reduced and fragmented in recent years, mainly due to anthropogenic pressures. Where its natural populations are located in areas with heavy human activity, this is causing the population size to decline at a more severe rate compared to if natural populations existed at locations with less human activity. In view of the above matter, immediate effort to collect, conserve and document this valuable plant in Malaysia should be carried out before it goes into the fate of extinction. Before an effective conservation program can be initiated on *R. tomentosa*, the genetic variation within and between its populations should be studied. According to Falk and Holsinger (1991), knowledge of genetic variation within and between the populations of a plant species is a prerequisite to formulate an effective and efficient conservation program, where the maximum genetic diversity of the species in the center of its origin and distribution is preserved. Maintenance of maximum and sufficient amounts of genetic diversity, which is the key to the diversity of a species, is crucial for a plant species to buffer sudden environmental changes imposed by unstable surroundings, and it is
important for the long term survival and evolution of a species (Barrett and Kohn, 1991). The survival and evolutionary capability of a species will be severely restricted with the loss of genetic diversity, occurring mainly through reduction in population size (Barrett and Kohn, 1991). The extent of genetic diversity in natural populations can be accessed through detailed population genetic analyses (Hamrick and Godt, 1996). In addition to assisting in the formulation of an efficient and effective conservation program, the knowledge of genetic diversity within natural populations can also be utilized in future breeding programs where the genetically diverse gene pool of landraces can be employed in hybridization programs to develop improved individuals (Sandhu et al., 2006).

Recently, the potential and applicability of ISSR markers in the study of natural plant populations has been demonstrated and proven (Zhang et al., 2006), owing to its advantages compared with other marker techniques. ISSRs are polymerase chain reactions (PCR) based markers that can be applied without prior knowledge of the plant genome sequence, where it greatly saves time and costs (Wolfe and Liston, 1998). It can be employed using a single primer consisting of a di- or trinucleotide simple sequence repeat, with or without a 3'- or 5'-end anchoring sequence of 1 to 3 nucleotides, targeting microsatellites regions that are abundant and dispersed throughout the genome. This technique exhibits more polymorphic fragments than other dominant markers such as random amplified polymorphic DNA (RAPD) (Qian et al., 2001). Primers are usually 16 to 25 nucleotides long resulting in higher stringency annealing (Wolfe and Liston, 1998) and overcoming the problem of reproducibility encountered by RAPD (Yang et al., 1996).

Previous studies conducted on *R. tomentosa* have been mainly concentrated on its ecology (Ren et al., 1997), antibacterial activities (Limsuwan et al., 2009), phytochemical analysis (Hiranat et al., 2012) and reproductive biology (Wei et al., 2009). To date, there is no study on the genetic diversity and structure of its natural population in Malaysia. The genetic pressure exerted on *R. tomentosa* imposed through its declining population size has warranted a thorough study on its population genetic structure, where the knowledge of the partitioning of genetic diversity within and between populations will be important to gain insight into the extent of the severity of population decline. Although *R. tomentosa* is still not widely exploited economically, however it possesses great future prospects in landscaping as well as in the food and beverage industry, in addition to its potential as natural antibacterial agent and traditional medicine. The once abundant *R. tomentosa* is more difficult to find in the wild nowadays. Thus, a study has been undertaken where inter simple sequence repeat (ISSR) was utilized to study the genetic variation among populations of *R. tomentosa*, and subsequently to suggest possible conservation strategies based on its population genetic structure.

**MATERIAL AND METHODS**

**Plant material sampling**

Young leaves of *Rhodomyrtus tomentosa* were collected from 15 natural populations (Table 1 and Figure 1), which represented the natural distribution of this plant in Malaysia from 2009 to 2012. In each of these populations, random samples of 5 to 10 plants were obtained. The total sample included 95 individuals across 10 states in Malaysia. The number of individuals sampled from each population depending on its size. Overall, the populations occurred in small isolated patches. The leaf samples were wrapped in wet paper and put in polythene bags to ensure freshness. These were then transported back to the laboratory where their DNA was immediately extracted. Longitude, latitude and altitude for each of the populations were also recorded.
DNA extraction and polymerase chain reaction (PCR) amplification

Leaf tissue was ground to powder form in liquid nitrogen. Total genomic DNA was extracted using a GeneAll® DNA extraction kit (GeneAll® Biotechnology, Korea) according to the manufacturer protocol, quantified using Nanodrop 200c spectrophotometer (Thermo Scientific, USA) and quality was determined through electrophoresis on a 0.8% agarose gel. Extracted DNA was stored at -20°C until use. PCR amplifications were performed in a TProfessional Standard Thermocycler (Biometra, Germany) in a total volume of 25 µL, containing 15 ng template DNA, 12.5 µL 2X Type-it Microsatellite PCR Master Mix (Qiagen, Germany), 0.4 µM primer, and 4 µL RNase-free water (Qiagen, 2008). Thermal cycling conditions for amplification were as follows: initial denaturation at 95°C for 5 min followed by 10 touch-down cycles at 95°C for 30 s, 90 s annealing temperature, starting at 5°C above the calculated melting temperature of each of the primers and decreasing 1.0°C per cycle and 30 s at 72°C. The program continued for another 25 cycles using the final annealing temperature of each primer reached in the touchdown phase, and finishing with a final extension step of 10 min at 68°C and kept at 4°C at ∞. A total of 22 primers previously used to study the genetic diversity in *Eucalyptus* sp of the same family, Myrtaceae (Balasaravanan et al., 2005;
Okun et al., 2008) and 19 primers from University of British Colombia (UBC) primer set no. 9 were synthesized and initially screened for amplifications. Each sample was replicated twice to ensure reproducibility. Each PCR run included a negative control. For screening purposes, the 41 primers were initially tested on one individual from each of the population. Amplification products were separated via electrophoresis on 1.5% (w/v) agarose gel with 1X TBE buffer at 70 V for 75 min, stained with GelRed™ Nucleic Acid Stain (Biotium Inc., USA) and visualized under UV light using the Alpha Innotech imaging system (Alpha Innotech, USA). GeneRuler 100bp ladder (Fermentas, Lithuania) was used as DNA molecular weight markers.

Statistical analysis

Since ISSR are dominant markers, it was assumed that each band represented the phenotype at a single biallelic locus. Amplified fragments were scored for presence “1” or absence “0” to construct the binary data matrix. The resulting binary data matrix was analyzed using POPGENE version 1.31 (Yeh et al., 1997) to estimate the following genetic diversity parameters, assuming Hardy-Weinberg equilibrium (HWE): percentage of polymorphic bands (PPB), expected heterozygosity ($H_e$), mean gene diversity within populations ($H_s$), total population gene diversity ($H_t$), Shannon information index (I) (Lewontin, 1972), coefficient of gene differentiation ($G_{st}$) (Nei, 1973), and gene flow ($N_m$). Gene flow was calculated from $G_{st}$ according to the equation $N_m = (1 / G_{st} - 1) / 4$ (Nei, 1973). A dendrogram was generated based on Nei’s (1978) unbiased measure of genetic distance among the populations with the UPGMA algorithm using NTSYS-pc 2.11 (Rohlf, 1998). In order to evaluate the partitioning of genetic variability between and within populations, analysis of molecular variance (AMOVA) was performed. The graphical representation of the relationship of the populations was revealed by principal coordinate analysis (PCoA). A Mantel test (Mantel, 1967) was performed to reveal the correlation between genetic and geographic distances (km) among populations. AMOVA, PCoA, Mantel test results, and geographic distances were calculated using the package GenAlEx 6.1 software (Peakall and Smouse, 2006).

RESULTS

Genetic diversity revealed by ISSR markers

Of the 41 primers tested, 11 of them (Table 2) yielded strong, clear and reproducible bands and were selected for further study of all 95 individuals. A total of 170 bands were amplified by the 11 selected primers, which ranged in size from 1.6 kb to 130 bp, corresponded to an average of 24.29 bands per primer. Of these bands, 95.29% (162 in total) were polymorphic (Table 2). Of the 11 primers, four of them generated 100% of polymorphic bands. Primer ISO 15 exhibited the lowest polymorphisms, which were only 70% (Table 2). The percentage of polymorphic bands (PPB) for a single population ranged from 6.47% (T01) to 41.62% (N01), with an average of 23.88%, and the average effective number of alleles per locus was 1.42. The average gene diversity was estimated at 0.087 within populations ($H_s$), and 0.25 for the total population ($H_t$). Expected heterozygosity ($H_e$) and Shannon Information Index (I) exhibited a similar trend to PPB. As shown by these three indices, among the 15 populations, population N01 exhibited the greatest level of variability ($H_e = 0.1513$; PPB = 42.35%; I = 0.2254) (Table 3).
Population structure

The mean $G_{ST}$ for all populations was estimated at 0.65, which indicated that 65% of the genetic variability was distributed across populations. The number of migrants ($N_m$) was estimated at 0.2652 individuals per generation between populations (Table 3), and strong genetic differentiation in $R.$ tomentosa suggests that the populations examined are isolated and that gene flow between the populations is restricted. The population genetic differentiation calculated by AMOVA was in agreement with the $G_{ST}$ value, where 63% of the total variation was partitioned among populations while the within populations variation was only 37% (Table 4).

Genetic relationships

Based on Nei’s (1978) unbiased measure of genetic distance, the populations Q01 and T01 were the farthest apart with a distance measure of 0.3408; while populations SA01 and SA02 shared the closest relationship, with a distance measure of 0.0363 (Table 5).
The dendrogram from UPGMA analysis based on Nei’s (1978) genetic distance clustered the 15 populations into two main clusters at a genetic distance of 0.13 (Figure 2). Cluster I included populations from C01, C02, D01, D02, K01, J01, J02, J03, M01, N01, T01 and T02. All of these populations originated from Peninsular Malaysia. Cluster II consisted of populations from Sabah and Sarawak from East Malaysia (SA01, SA02 and Q01). Under cluster I, the population from Bera, Pahang (C02), both of the populations from Kelantan (D01 and D02), and the population from Naka, Kedah (K01) were more closely related compared to their respective relationships with the population from Kuantan, Pahang (C01). Populations from Melaka (M01) and Negeri Sembilan (N01), both originating from the southwestern part of Peninsular Malaysia were closely related to each other. Both of the populations from Terengganu (T01 and T02) were the most distantly related to the others within cluster I, and these were followed by the populations J01, J02 and J03, all of which originated from the state of Johor. There was a clear differentiation between populations from Peninsular Malaysia and East Malaysia. Overall, populations from the same state were highly related while populations from the different states were more distantly related (Figure 2). The relationship suggested by the two-dimensional PCoA plot (Figure 3) was consistent with the dendrogram generated through Nei’s (1978) unbiased measure of genetic distance. The first two axes of the PCoA plot accounted for 26.78% and 20.40% of the total genetic variability respectively. The populations were divided into two groups in a PCoA plot, with group I consisting of populations from Peninsular Malaysia and group II consisting of populations from East Malaysia (Sabah and Sarawak). The populations can be grouped together according to their state of origin in group I. Two subgroups can be identified in group II, one consisting of populations from Sabah and the other consisting of populations from Sarawak.

The result of the Mantel test with 999 permutations showed that in *R. tomentosa*, the genetic distance of populations was significantly correlated with geographic distance ($r = 0.581$, $P = 0.001$).
DISCUSSION

The genetic structure and genetic diversity within and among plant populations can be considerably affected by its breeding system (Hamrick and Godt, 1996). Wei et al. (2009) has

Figure 2. UPGMA dendrogram based on Nei’s (1978) genetic distance among populations of *Rhodomyrtus tomentosa*.

Figure 3. PCoA plot on 95 individuals of 15 populations of *Rhodomyrtus tomentosa*.
shown that R. tomentosa possesses a mixed mating system (outcrossing and geitonogamy) with geitonogamy being the predominating system. ISSR assessment of the 15 populations of R. tomentosa from Malaysia revealed that this species contains a relatively high level of genetic diversity among its populations ($H_t = 0.2510; I = 0.3897; PPB = 95.29\%$). However, within each population, a rather low level of genetic diversity was observed ($H_s = 0.087; I = 0.1291; PPB = 23.88\%$). In addition, a high level of genetic differentiation ($G_{ST} = 0.6534$) and a low level of gene flow ($N_m = 0.2652$) was also seen among the R. tomentosa populations. The results from $G_{ST}$ were further supported by the results from AMOVA, where 63\% of the genetic variation was distributed among populations. The above findings concur with Hamrick and Godt (1996) and Nybom and Bartish (2000) who state that selfing species usually possess lower genetic diversity within populations and higher genetic differentiation among populations compared to outcrossing species. This is also comparable to other selfing plant species, such as Sinojackia dolichocarpa C.J. Qi ($H_s = 0.2255; PPB = 72.99\%; I = 0.3453$) (Cao et al., 2006) and Tacca chantieri André ($h = 0.264; PPB = 90.62\%$) (Zhang et al., 2006).

R. tomentosa has a wide range of distribution across the South East Asian region, as well as some of the East Asian region (Chen, 1984). It has been reported that the genetic diversity for widespread species is higher than that of rare and endangered species (Verma et al., 2009). In Malaysia, R. tomentosa was once growing profusely with somewhat contiguous large population sizes (Latiff, 1992), contributing to its large gene pool with abundant genetic diversity. During that time, considerable diversity levels had been accumulated. Recently, it has been subjected to habitat destruction due to anthropogenic disturbances and its populations have become fragmented. Despite this, high variability has been and continues to be well conserved in these severely isolated populations, thus leading to the high level of genetic diversity among the populations. According to Zawko et al. (2006), for plant species that experienced isolation of populations and reduction in the population size which was occurring recently due to anthropogenic pressure, insufficient timeframes were provided for genetic diversity to be reduced. The recent occurrence of habitat fragmentation in populations of R. tomentosa can be inferred from the groupings found in the populations from M01 and N01, both from different states. We can assume that previously, populations from each state occurred in a continuous pattern with moderately frequent gene flow, and due to recent development activities, the once continuous distribution of R. tomentosa has been broken down and become genetically subdivided. This can also be understood from the groupings found in the two populations from Sabah, where the genetic variation within populations is low. The close relationships between populations from the same state may be attributed to the close distance between populations sampled from the same state (5.93-99.13 km) as compared with the wider distance between populations sampled from different states (125.96-1792.40 km). However, the possibility of restricted gene flow cannot be ruled out due to physical distance and the geitonogamous pollination system of the species leads to closely related populations from the same state and distantly related populations from different states.

Slatkin (1985) has proposed that significant population differentiation may be maintained under strong differential selection, which will occur if the migration rate is less than one successful migrant per generation ($N_m < 1$). Compared with the RAPD-derived value for selfing plant species ($G_{ST} = 0.59$) (Nybom and Bartish, 2000), the genetic differentiation value obtained from this study was higher ($G_{ST} = 0.65$), indicating significant genetic divergence among populations. Fluctuating population number and size, shifts in distribution, habitat fragmentation and population isolation may be involved in shaping the genetic structure of plant populations (Hamrick and Godt, 1996).
Furthermore, the genetic structure may also be shaped by local adaptations due to genotype x environment interactions (Cao et al., 2006).

Reduced effective population size is a characteristic in selfing species (Ingvarsson, 2002). In small isolated populations, the genetic differentiation caused by genetic drift and inbreeding is more pronounced (Barett and Kohn, 1991). Recent field observations have shown that *R. tomentosa* occurs in severely fragmented and isolated habitats, with small population sizes. As inferred from the low level of gene flow among populations of *R. tomentosa* in Malaysia, genetic drift may be a contributing factor to the possible high levels of genetic differentiation in its future populations. Due to the predominant geitonogamous mating system of *R. tomentosa*, together with the low gene exchange between populations, severe inbreeding may occur in the future thus contributing to the loss of genetic variation within populations. In the long term, genetic drift and the high genetic differentiation will cause alleles to be fixed in isolated populations, which may lead to new variations or the total extinction of the populations depending on whether advantageous or non-advantageous alleles have been fixed. The low gene flow among *R. tomentosa* populations has also caused lack of hybridization among individuals from other populations leading to a low, within population variability, and restricted gene pool. Together with the homogenous environmental conditions of local habitats, this has caused lack of speciation. In addition, although the highest value of genetic distance recorded among *R. tomentosa* populations was 0.3408, these populations were still considered to be closely related to each other. The observed genetic variation existing among populations is probably caused by the adaptation of the populations to the local environmental pressures.

Seed dispersal mechanisms also contribute to the genetic structuring of a plant population (Hamrick and Godt, 1996). Hamrick and Godt (1996) performed a combined analysis of mating systems and seed dispersal mechanism and revealed that for mixed mating species with ingested seed dispersal mechanisms, the genetic differentiation among populations is higher than in species with wind dispersal mechanisms from the same breeding system. The seeds of *R. tomentosa* are ingested and dispersed by frugivorous birds, mainly bulbul (*Pycnonotus sinensis*), which consume the juicy ripened berries (Wei et al., 2004). Bulbuls are known to live in pairs or small groups where they have defined territories and small home ranges and thus tend to disperse the seeds within 100 m from the mother plant (Spiegel and Nathan, 2007). The seed dispersal distance by bulbuls rarely exceeds 500 m (Spiegel and Nathan, 2007). In addition to ingested seed dispersal by bulbuls, the seeds of *R. tomentosa* can also be dispersed by ants, which move the seeds from ripened berries fall to the ground or from feces deposited by bulbuls to their colonies. The average maximum dispersal distance by ants is only 4-5 m (Wei et al., 2004). The seed dispersal limitations of bulbuls and ants may have contributed to the patchy populations of *R. tomentosa*.

Gene flow may also occur via pollen dispersal. Wei et al. (2009) has shown that *R. tomentosa* is mainly pollinated by bees from the Apidae family with *Amegilla florae* and *Xylocopa nasalis* as the primary pollinators. It has been shown that bees belonging to the Apidae family have relatively short foraging distance, averaging 1526.1 m, and they tend to revisit the same plant on which they have fed previously (Tepedino, 2005). The foraging habits of the pollinators are dependent on the properties of the plant population, where the ability of the plant to attract and support pollinators will be affected by its small population sizes (Tepedino, 2005). This condition is further exaggerated by urbanization where the habitat needed to support pollinators is threatened (Tepedino, 2005). In view of the above, the small and fragmented populations of *R. tomentosa* are a result of urbanization and may not be sustainable for pollinator subsistence in the long run. The limited gene flow as inferred from the short seed dispersal and pollen exchange has led to a higher genetic differentiation among *R. tomentosa* populations, in line with the findings of Hamrick and Godt (1996).
It is well known that the amount of gene flow through pollination and seed dispersal can be limited by geographic isolation (Pfeifer and Jetschke, 2006). In R. tomentosa, the short distance of seed dispersal by bulbuls (100-500 m) and ants (4-5 m), and the short pollen dispersal distance by bees (1.5 km) are well below the minimum geographic distance between J02 and J03 (5.93 km). This phenomenon has further exaggerated the high level of genetic differentiation among R. tomentosa populations. The pattern of genetic distance was corresponded to the geographic distance (Mantel test: r = 0.581, P = 0.001). This suggested that apart from genetic drift, isolation by distance (Wright, 1943) could in part contribute to the present observed genetic diversity. Populations from East Malaysia (SA01, SA02 and Q01) and West Malaysia (the remaining populations) are separated by the South China Sea and the genetic distances between populations from the two regions are high as shown by UPGMA dendrogram and the PCoA plot.

From the results of the population genetic structure of R. tomentosa obtained from this study, insights can be gained on conservation strategies for this species in Malaysia. The high level of genetic differentiation and the limited gene flow among R. tomentosa populations suggests that all the extant populations should be preserved to conserve the maximum genetic variability of the species. It is well known that maintaining the genetic diversity is one of the main objectives for a conservation program (Avise and Hamrick, 1996). Keeping this in view, for ex situ conservation, a small number of plants should be sampled from all the populations. The severe habitat fragmentation and small population sizes of R. tomentosa has made it vulnerable to the loss of genetic diversity due to genetic drift and inbreeding. Thus, efforts have to be made to protect its natural habitat and to increase its population size for in situ conservation. Population from Negeri Sembilan (N01) contains a significant amount of the genetic diversity and should be given priority to be conserved both in situ and ex situ. Nevertheless, other populations should not be neglected in the conservation program as they might contain rare alleles that could lead to desired characteristics.

In conclusion, the high level of genetic variation found among the populations and low levels of genetic variation within populations has indicated that R. tomentosa has undergone habitat fragmentation recently, thus efforts should be taken promptly by authorities to ensure the continuing existence of this plant in their natural habitat in the future. The genetic variation exerted by the populations has enabled them to respond to the environmental change of their surroundings, but still not enough to trigger the speciation process. Care should be taken during ex situ conservation to ensure that the populations are maintained under the same environments.

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