Genetic differentiation and antioxidant activities of Bouea macrophylla Griffith in Nakhon Nayok province

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Abstract Genetic differentiation and antioxidant activities in ethanolic extracts from leaves of Bouea macrophylla Griffith were determined. The result revealed genetic differentiation among sour ma-praang, ma-yong and sweet ma-praang of B. macrophylla Griffith (\(\Phi_{PT}=0.772, \ p\text{-value}=0.000\)). In addition, high genetic diversities were found in sour ma-praang and sweet ma-praang populations \((P: 51.4 \text{ and } 57.1 \% ; \ He: 0.1900.035 \text{ and } 0.2240.036, \text{ respectively})\), but low genetic diversity was found in ma-yong population \((P: 8.6 \% ; \ He: 0.0350.021)\). Total phenolic contents of sour ma-praang, ma-yong and sweet ma-praang were estimated as 680.51±89.81, 701.03±59.89 and 530.85±41.23 mg gallic acid/g extract, respectively. Free radical scavenging activities of sour ma-praang, ma-yong and sweet ma-praang were found \((1/EC50= 4.17, 1.43 \text{ and } 1.37, \text{ respectively})\) corresponding with metal-chelating activities \((1/EC50=0.83, 0.65 \text{ and } 0.17, \text{ respectively})\). Therefore, the obtained data may be applied to cultivation and utilization of their leaves as source of natural phenolics and antioxidants.

Keywords Antioxidant activity · Bouea macrophylla Griffith · Genetic differentiation · Random amplified polymorphic DNA

Introduction

Marian plum (Bouea macrophylla Griffith) is a fruit tree that belongs to the family Anacardiaceae, and it is grown and widespread in Thailand and Sumatra (Blench 2008). In Thailand, this plant is a popular and important economic fruit tree, well known as ma-praang and ma-yong. Generally, it is classified according to its fruit tastes into 3 groups: sour ma-praang, sweet ma-praang, and ma-yong (Subhadrabandhu 2001). Furthermore, the plant characteristics have height at approximately 27 m and leaf shape like lanceolate or elliptic in length up to 45 cm and width up to 13 cm (Subhadrabandhu 2001). The unripe and ripe fruits of this plant have high nutritional values containing high potassium, essential amino acids, fiber and lipid (Rajan and Bhat 2016). In addition, it shows the existence of antioxidant compounds and antioxidant activities in unripe and ripe fruits (Rajan and Bhat 2016). Antioxidants can help to prevent several diseases that occur from oxidant compounds, such as cancer and alzheimer’s disease (Halliwell et al. 2005; Valko et al. 2006; Gibson et al. 2008). Thus, antioxidants from many plants are good choices to use in food and health promoting of human. Due to most people worry about toxicities of artificial antioxidants used commonly in now.

Nevertheless, there are no scientific reports involving medicinal properties in leaves extracts and genetic structure of B. macrophylla Griffith. This work may help to promote cultivation and heath for further commercial aspect. Therefore, random amplified polymorphic DNA (RAPD) were applied in this study. RAPD is one of popular molecular techniques using PCR method with a single arbitrary primer. This tool provides low cost, convenient, rapid and simplicity (Weder 2002; Suvanchakasem et al. 2012). It has been used to distinguish and identify genotypes in several plant species (Arif et al. 2010; Thummajitasakul et al. 2014).

Nowadays, B. macrophylla Griffith is one of economical fruits but its leaves is not provided importance, in comparison with other plants. However, one of our major purposes is to find natural sources that may contain antioxidants which are useful in health promotion. In addition, some farmers and consumers lack

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knowledge in classification of *B. macrophylla* Griffith. Therefore, this study of genetic markers may assist in identification, breeding and conservation of *B. macrophylla* Griffith in Nakhon Nayok.

**Materials and Methods**

**Sample collection**
Leaves of *B. macrophylla* Griffith (Sour ma-praang, n=5; Sweet ma-praang, n=8; Ma-yong, n=56) were collected from 16 regions of Mueang Nakhon Nayok and Ban Na districts of Nakhon Nayok province in where the plant was popular to cultivate (Fig. 1). Furthermore, global positioning system was recorded by using UTM system. Width, length and tail of each leaf was measured, then cleaned by pure water and kept in a plastic bag at 20°C until used.

**DNA extractions and RAPD method**
Total DNA from fresh leaf of each sample was extracted using plant DNA extraction kit (RBC Bioscience, New Taipei City, Taiwan). Briefly, each sample tissue was grinded by a sterile pestle in 400 mL of GP1 buffer, then reacted with 5 mL of 10 mg/mL RNase A at 65°C for 10 min. Next, GP2 buffer for 100 mL was mixed and left for 3 min at 4°C. The supernatant was used further by adding GP3 buffer for 1.5 time. The purified DNA was obtained by applying to a GD column and centrifuged at 13,000 rpm for 3 min, then washed with 600 mL of W1 buffer and wash buffer, respectively, finally extracted with 100 mL of elution buffer. After that, RAPD reactions were carried out by a LifeECO thermal cycler (Bioer Technology, Hangzhou, China) according to Thummajitasakul et al. (2014a). Thirteen primers (Ashraf et al. 2014; Thummajitasakul et al. 2014a) were screened for RAPD analysis, of these, five primers (OPA03, OPA04, OPA09, OPA15 and C4) providing genetic polymorphisms were selected. Each RAPD condition was performed in a total volume of 20 L consisting of DNA template for 25 ng, 10 L of OnePCR Plus reaction mixture which contained fluorescence dye, Taq DNA polymerase, PCR buffer, dNTP and loading dye (GeneDireX, Las Vegas City, NV, USA), and 1 µM of each arbitrary primer. The reaction was run as one cycle of initial denaturation at 95°C for 5 min, followed by 35 cycles of DNA denaturation at 95°C for 30 s, annealing at 45°C for 1 min and extension at 72°C for 1 min. Finally, the last extension was at 72°C for 5 min, then cooled down at 4°C. The PCR products were checked on a 1.5% electrophoresis gel in 0.5XTAE buffer (Pacific Science Co., Ltd., Bangkok, Thailand) at 100 volts for 20 min, and visualized under UV light (Omega Fluor, Aplegen). A 100 bp DNA ladder containing fluorescence and loading dye was used as marker (GeneDireX).

**Sample extracts and total phenolic contents**
The fresh leaves from sour ma-praang, sweet ma-praang, and ma-yong of *B. macrophylla* Griffith were washed by distilled water and incubated at 50°C until dry. After that, each sample was grinded to a powder by a homogenizer, and kept in a plastic bag at room temperature until analyzed. To obtain extracts, 15 g of dried powder was extracted with 150 mL of 95% ethanol solvent at 50°C for 16 h in an incubator with shaking. Then, the solvent was evaporated in the incubator at 50°C until dry and adjusted the initial concentration into 100 mg/mL. Each extract was collected and kept at 20°C until used. Each extraction was carried out in duplicate.

Total phenolic contents were estimated by the Folin-Ciocalteu colorimetric method (Deetae et al. 2012). Each extract (300 L) was reacted with 1.5 mL of Folin-Ciocalteu reagent for 5 min, and added 1.2 mL of sodium carbonate (7.5% w/v) with incubation at room temperature for 30 min. Each reaction was performed in duplicate. Next, an absorbance was determined at wavelength 765 nm by spectrophotometer (Model T60UV). Water and gallic acid (0-100 g/mL) were used as blank and a standard phenolic compound, respectively. Total phenolic contents were compared in unit of mg gallic acid equivalent per gram extract.

**Antioxidant activities**
The antioxidant activity was performed via an 2,2'-azinobis-(3-ethylbenzothiazoline-6-sulfonic acid (ABTS) free radical method (Deetae et al. 2012). First, the ABTS free radical cation was produced from a reaction of 7 mM ABTS solution (10 mL) with 140 mM potassium persulfate (179 µL) in dark condition at room temperature for 12–16 h. To assay antioxidant activity, the ABTS radical solution was diluted with aqueous solvent to obtain an absorbance of 0.700±0.050 at 734 nm. Next, 3.9 mL of the diluted solution was added in 20 µL of each plant extract and mixed.
gentle, then allowed to stand for 6 min at room temperature in dark condition. After that, the absorbance was determined immediately at 734 nm. Each reaction was tested in duplicate. The percentage of antioxidant capacity was calculated following the method of Thummajitasakul et al. (2014b).

Moreover, a ferrous ion-chelating (FIC) assay of each sample was detected with the protocol of Deetae et al. (2012). Each extract (1 mL) was added with 0.1 mM FeSO$_4$ (1 mL) and 0.25 mM ferrozine (1 mL), and incubated for 10 min in dark condition at room temperature. Then, an absorbance was determined at 562 nm with EDTA used as the positive control. Each reaction was tested in duplicate. The percentage of metal chelating ability was estimated according to the method of Thummajitasakul et al. (2014b). The effective concentration (EC) 50 value, which was an effective concentration of antioxidants at 50% of the free radical scavenging or metal chelating capacities, was calculated from the graph. Then, the EC50 value was modified to 1/EC50 because this value varies directly according to its antioxidant activity.

**Data analysis**

A binary data was scored with the presence (1) or absence (0) of a RAPD band. After that, the values of genetic diversity were determined by haplotype patterns, mean expected heterozygosity (He) and percentages of polymorphic bands (P). The molecular variances among populations (sour ma-praang, sweet ma-praang, and ma-yong) and between two groups (group 1: sour ma-praang and sweet ma-praang, and group 2: ma-yong) of *B. macrophylla* Griffith were analyzed by AMOVA using GenAlEx 6.51 program (Peakall and Smouse 2006; Peakall and Smouse 2012). The principal component analysis (PCA) was performed using excel add-in Multibase version 2015 (Numerical Dynamics 2009).

Analysis of variance (ANOVA) method was used to estimate the difference of antioxidant activities and leaf size (width, length and tail) among three populations at significant level of $p$-value < 0.05 (Soper 2016).

**Results and Discussion**

The RAPD is one of effective techniques having usefulness for many studies in genetic diversities and characterization of several fruit trees, i.e. *Dimocarpus longan* Lour. (Mei et al. 2014), *Litchi chinensis* Sonn. (Cheng et al. 2015). The result showed the RAPD patterns from each selected primer and total 35 loci, and 18 haplotype patterns were obtained (Fig. 2). To elucidate whether genetic differentiation was found in this plant, total samples were divided into 3 populations (sour ma-praang, sweet ma-praang, and ma-yong). High genetic diversities were detected within sour ma-praang and sweet ma-praang providing $p$ values for 51.4 and 57.1 %, and He values for 0.1900.035 and 0.2240.036, respectively. Moreover, low genetic diversity was found in ma-yong population ($p$ values = 8.6 % and He = 0.0350.021) shown in Table 1. The result indicated relatively low level in genetic variation of ma-yong population in comparison to other populations. It may result from cumulative inbreeding within population and habitat restriction (Ellstrand and Elam 1993; Glover and Abbott 1995). In addition to the result above, mean difference of width, length and leaf tail among populations was analyzed by one-way ANOVA. Although, it showed that sour ma-praang and sweet ma-praang were insignificantly detected in size difference for leaf width, length and tail length ($p$-value = 0.874, 0.504, and 0.524, respectively), their leaf sizes were larger than ma-yong, significantly ($p$-value

![Fig. 2 Examples of RAPD patterns amplified by primer OPA04 (A) and primer OPA03 (B). Lanes 1-3, 12-14 were sweet ma-praangs, lanes 4-5 and 15-16 were sour ma-praangs, and lanes 6-11 and 17-22 were ma-yongs of *B. macrophylla* Griffith. The 100 bp DNA ladder was used as marker](image-url)
The mean difference is significant at (p-value=0.000), Table 1. For AMOVA analysis, high genetic differentiation among three populations was significantly found (φPT=0.772, p-value=0.000). While the populations were divided into 2 groups (group 1: sour ma-praang and sweet ma-praang, and group 2: ma-yong), higher quantities than that of sweet ma-praang, significantly (p-value =0.023 and 0.003, respectively). Several researches found.

| Samples                  | He          | P (%) | No. of haplotypes | No. of Different Bands | No. Private Bands* | Leaf size [Mean ± SD (cm)] |
|--------------------------|-------------|-------|-------------------|------------------------|-------------------|---------------------------|
| Sour ma-praang (n=5)     | 0.190±0.035 | 51.4  | 5                 | 27                     | 0                 | 4.50±0.50, 16.7±2.16, 1.34±0.50 |
| Sweet ma-praang (n=8)    | 0.224±0.036 | 57.1  | 8                 | 27                     | 1                 | 4.44±0.72, 15.6±3.24, 1.56±0.63 |
| Ma-yong (n=56)           | 0.035±0.021 | 8.6   | 5                 | 22                     | 3                 | 3.02±0.88, 10.15±2.50, 0.79±0.35 |
| Total (n=69)             | 0.204±0.035 | 68.6  | 18                | 31                     | 31                |                           |

*No. Private Bands=No. of Bands Unique to a Single Population

The mean difference is significant at p-value <0.05.

Table 2 Genetic differentiation among sour ma-praang, sweet ma-praang and ma-yong populations, and between 2 groups (group 1: sour ma-praang and sweet ma-praang, and group 2: ma-yong), which were analyzed by AMOVA.

| Source                  | df | SS         | MS         | Est. Var. |
|-------------------------|----|------------|------------|-----------|
| 3 populations          |    |            |            |           |
| Among Pops              | 2  | 88.417     | 44.208     | 3.869     |
| Within Pops             | 66 | 75.496     | 1.144      | 1.144     |
| 3 populations, 2 groups|    |            |            |           |
| Among Regions           | 1  | 84.197     | 84.197     | 3.554     |
|                         | 1  | 4.219      | 4.219      | 0.500     |
| Among Pops              | 1  | 75.496     | 1.144      | 1.144     |
| Within Pops             | 66 | 75.496     | 1.144      | 1.144     |

*The mean difference is significant at p-value <0.05.

Many phenolics in fruits and plants more than 8,000 compounds with known structures, such as phenols, phenolic acids, flavonoids and tannins (Harbone and Williams 2000; Santos-Buelga and Sacalbert 2000; Naczk and Shahidi 2006). In this work, total phenolic contents were found in 95 % ethanol extracts from leaves of sour ma-praang, sweet ma-praang and ma-yong. There were many reports in other plants with sour taste, such as Mango (Mangifera indica L.) had some phenolic compounds, e.g. high concentration of mangiferin in youg leaves (Barreto et al. 2008). Besides these, leaves extracts of citrus plants revealed high level of phenolic compounds (Al-Anbari and Hasan 2015). Moreover, EC50 values of free radical scavenging activity in each extract between sweet ma-praang and ma-yong showed insignificantly difference (p-value=0.539), and showed higher value than that of sour ma-praang, significantly (p-value=0.000). For metal-chelating activity, it found that EC50 values between sour ma-praang and ma-yong showed insignificantly difference (p-value=0.164), and showed lower values than that of sweet ma-praang, significantly (p-value=0.003 and 0.005, respectively) (Table 3). Many researches have reported antioxidant activities from plant extracts involving phenolic compounds. Of these, polyphenolics can play a major role as antioxidant activity (Lima et al. 2014), such as flavonoids, led to useful for human health by free radical scavenging and metal chelating properties (Harbone and Williams 2000; Mira et al. 2002). In this study, free radical scavenging activity was determined by ABTS method, and metal-chelating activity was estimated by FIC method. The principle of the ABTS method is the production of the ABTS free radical cation for using in reduction reaction with antioxidants, and it can be applied to determine high-pigmented, hydrophilic and lipophilic...
antioxidants (Floegel et al. 2011). For FIC method, metal-chelating activity is based on the prevention of the association between metal and ferrozine (Robu et al. 2012). The chelating activity has a role in treatment of metal overload condition in human body, such as Thalassemia (Ebrahimzadeh et al. 2008). In this study, the ethanol leaves extracts of *B. macrophylla* Griffith, showed free radical scavenging and metal-chelating activities.

Sour ma-praang showed the lowest EC50 value indicating the highest of free radical scavenging activity. Moreover, metal-chelating activities were mostly found in leaves extracts of sour ma-praang and ma-yong. However, plant extracts may content some non-phenolic antioxidants that play importance to contribute antioxidant activities, such as the report of el-Sayed et al. (2008) found four non-phenolic antioxidants (i.e. 1-**O**-beta-**D**-glucopyranosyl-

| Samples          | Total phenolic contents (mg gallic acid/ g extract) | ABTS | FIC |
|------------------|-----------------------------------------------------|------|-----|
|                   |                                                     | EC50 | 1/EC50 | EC50 | 1/EC50 |
| Sour ma-praang   | 680.51±89.81                                       | 0.24±0.08 | 4.17 | 1.21±0.1 | 0.83 |
| Sweet ma-praang  | 530.85±41.23                                       | 0.73±0.07 | 1.37 | 5.75±0.35 | 0.17 |
| Ma-yong          | 701.03±59.89                                       | 0.7±0.06 | 1.43 | 1.55±0.2 | 0.65 |

Fig. 3 The PCA plots. A was the plot of variables (loading) for total phenolic contents, free radical scavenging activity, metal-chelating activity, and the genetic similarity values among sour ma-praang, sweet ma-praang and ma-yong. B was the plot of samples (score) for sour ma-praang, sweet ma-praang and ma-yong of *Bouea macrophylla* Griffith
2-methoxy-3-(2-hydroxy-triaconta-3,12-dienoate)-glycerol) in leaves extracts of Buddleja asiatica Lour. In addition, several antioxidant agents (phenolics, flavonoids, flavonols, tannins, anthocyanins and ascorbic acid) of unripe and ripe fruit extracts of B. macrophylla Griffith have been detected in different solvent extracts (methanol, ethanol, and distilled water) according to the literature (Rajan and Bhat 2016). Of these solvents, the highest antioxidant activity was found in methanol extracts of unripe fruits (Rajan and Bhat 2016). However, there were several factors to increase phenolic contents in plants, such as stressful environments (e.g. insect attack and wound) and organic planting (Woese et al. 1997; Reyes and Cisneros-Zevallos 2003; Mitchell et al. 2007).

The results were confirmed with distribution pattern among three populations of B. macrophylla Griffith of PCA shown in the loading plot using the values of genetic similarities obtained from RAPD analysis. Moreover, interrelation among total phenolic contents, free radical scavenging activity, metal-chelating activity were demonstrated with 1/EC50 values in the loading PCA plot. The PC1 and PC2 revealed 61.8 and 38.2 % of the total variance, respectively. PC1 loadings had total phenolic contents, and three populations, while PC2 loading consisted of free radical scavenging and metal-chelating activities. When the loading and score plots were overlaid, it revealed that sour ma-praang and sweet ma-praang had clustered together, but ma-yong was separated in the plot. Moreover, the ethanolic extract of sour ma-praang showed the highest of total phenolic contents, free radical scavenging and metal-chelating activities (Fig. 3). The PCA result clearly indicated that genetic differentiation existing among sour ma-praang, sweet ma-praang and ma-yong, and it suggested that sour ma-praang showed genetic similarity with sweet ma-praang more than ma-yong. Furthermore, it also confirmed that total phenolic contents, free radical scavenging and metal-chelating activities were mostly found in leaves extracts of sour ma-praang.

In conclusion, based on our research, genetic differentiation was detected among sour ma-praang, sweet ma-praang, and ma-yong of B. macrophylla Griffith. Moreover, high genetic diversities were found within population of sour ma-praang and sweet ma-praang, but low genetic diversity was found in ma-yong population. RAPD technique was certificated to be useful for genetic analysis. It also concluded that high amounts of total phenolic contents, free radical scavenging and metal-chelating activities were detected in leaves extracted with 95 % ethanol solvent. Therefore, more studies are need to identify sequence related amplified polymorphism, bioactive compounds and to extend scope in its cultivation to rise phenolic contents before economical applications, and pharmaceutical useful as natural sources of antioxidants.

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