Chemical and physical characterization of mume fruit collected from different locations and at different maturity stages in São Paulo State

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Abstract

*Prunus mume* is widely studied due to its health benefits regarding increase of blood fluidity and consequent improvement of the cardiovascular system and the prevention or even the fight against different types of cancer. However, in Brazil this culture is found only among oriental descendants. The present study aimed to characterize mume fruit collected from three different locations in the State of São Paulo regarding general aspects such as pH, total titratable acidity (TTA), total soluble solids (TTS), pectin content and yield of pulp and chemical characteristics: total phenolic compounds (TPC) and antioxidant capacity. Mume fruit were collected unripe and analyzed until maturation about 88 days after flowering. Fruit collected in Botucatu came from a commercial mume fruit producer and had average weight of 16.9 g, while in fruit from other locations weight varied from 5.7-6.9 g. TSS ranged from 9.5 to 10.0 Brix, total solids was 10.2-12.2% and pH showed values between 2.5 and 2.7 for all locations. TTA expressed in citric acid decreased from 4.0-5.7 g (100g⁻¹) at unripe stage to 2.0-3.8 g (100g⁻¹) in mature-stage fruit. Pectin content decreased from 11.2 to 10.8% during fruit maturation, TPC content was 147-226 mg catechin (g⁻¹) on a dry matter basis and the antioxidant capacity was 96-169 µMol Trolox (g⁻¹) on a dry matter basis or 21-34 µMol Trolox (g⁻¹) on a wet matter basis.

Keywords: mume; fruit characterization; health product.

1 Introduction

The family Rosaceae is one of the largest Angiosperm family. Regarding the economic aspect, it is one of the most important families, including apple (*Malus sylvestris*), pear (*Pyrus communis*), peach (*Prunus persica*), plum (*Prunus salicina*), strawberry (*Fragaria vesca*), yellow plum (*Eriobotrya japonica*), cranberry (*Rubus idaeus*) and mume (*Prunus mume*). Plants from this family are cultivated in South and Southeast regions of Brazil because of the milder weather conditions, favorable to the development of temperate fruit (SOUZA; LORENZI, 2005). Mume tree exhibits an arboreal growth and 4-6 meters high, with white and androgynous flowers that bloom from June to August. Fruit are drupaceous, with firm pulp with bitter and sour taste and maturity from October to December (LORENZI et al., 2006).

Asia, in particular China, is the birthplace of mume culture, where production, consumption and acceptance are high. Further, in Asia there are different cultivars that include early and late fruit producing plants, so the harvest interval increases to three or four months. For the same plant, the interval for harvesting does not exceed three weeks.

In the State of São Paulo, mume trees were introduced in 1960’s by Asian descendants and even though in Brazil there are no studies to improve yield and quality of fruit, there are over 200 cultivars of *Prunus mume* Siebold et Zucc. worldwide, whose fruit are consumed mainly in Korea, China and Japan (TSUBAKI; OZAKI; AZUMA, 2010). In Japan fruit price is about 1.68 US dollar (kg⁻¹), and present an average yield of 6.7 ton (ha⁻¹) (JUN; CHUNG, 2008; TOPP; NOLLER; RUSSEL, 2007).

*P. mume* has been studied in Brazil as rootstocks for peach (*Prunus persica*) and plums (*Prunus salicina*), showing promising characteristics, such as rusticity, high resistance to pests and diseases (MAYER; PEREIRA; MÔRO, 2008). These characteristics suggest that mume fruit can be produced in large scale without or with less use of pesticides.

Mume fruit can be collected directly from the tree, shaken to fall in large clothes, nets or trimmed grass at unripe-stage or dropped by their own while still green but physiologically mature. At this stage, maturation occurs at temperature above 20 °C ventilated (LUO, 2006) because mume fruit display a typical climacteric pattern of respiration and ethylene biosynthesis. The removal of over-ripe and injured fruit reduces significantly the presence of insects and flies during harvest and maturation. Texture, color and aroma of these fruits change significantly during maturation (MIYAZAWA et al., 2009).

Several mume processed products have been consumed as healthy foods and for the treatment of several diseases because they are rich in bioactive compounds, such as anti-cancer and antioxidant substances (TSUBAKI; OZAKI; AZUMA, 2010; LIU et al., 2009; SHI et al., 2009; ADACHI et al., 2007; SHI; MOY, 2005). Given the similarity of unripe apricot and mume...
Mume fruit characterization at different maturity

fruit, Jun and Chung (2008) developed sequence-characterized amplified region (SCAR) markers to differentiate mume fruit due to its higher value and demand. This identification can be used both for germplasm classification and to detect apricot unripe fruit commercialized as mume fruit. Jo et al. (2006) used mume fruit extract as a natural source of antioxidant compounds, which can inhibit lipid oxidation and formation of “warmed over flavor” volatiles during storage of cooked chicken meat. Mume extract inhibits free radical formation during initial stages of oxidation. At high concentrations, these highly reactive free radicals break triglyceride chains to produce free carboxylic acid that provide product alteration.

The present study aimed to characterize mume fruit collected from three different locations in the State of São Paulo regarding general aspects such as pH, total titratable acidity (TTA), total soluble solids (TTS), pectin content and yield of pulp and chemical characteristics: total phenolic compounds (TPC) and antioxidant capacity. These results can help to decide the best option: breeding or adaptation of new cultivars, for the introduction of mume to locations where currently peach and plums are grown in Brazil. Also, fruit characterization is needed to permit its use as ingredient for food production.

2 Materials and methods

2.1 Plant material

Fruit used were green-to-yellow color soon after natural fruit dropping occurred, approximately 88 DAF (days after flowering) and were stored for up to 8 days at 26 °C ventilated until complete ripening. In previous studies, fruit were collected from the trees weekly from 60 DAF and fruit did not mature properly.

Mume fruit were collected in São Paulo State, Brazil, from different locations:

- S – Angatuba, SP- Latitude 23° 30’ 43” S; Longitude 48° 16’ 38” W; Elevation 737 m;
- CB – Capão Bonito, SP- Lat. 24° 02’ 52” S; Long. 48° 21’ 19” W; Elev. 705 m;
- SUN – Botucatu, SP- Lat. 22° 57’ 56” S; Long. 48° 27’ 25” W; Elev. 843 m.

According to Koppen climate classification, all locations are Cwa, characterized as highland tropical climate, with rainfall in summer and dry winter, with average temperature of the warmest month above 22 °C. In Angatuba, mume trees were planted as windbreak for peach and plum culture and in Capão Bonito, trees were planted for ornamental purpose. Botucatu was the only commercial mume production found in the State of São Paulo, with plants brought from China in the 1960’s. SUN was the only location where the pruning of the tree was done. Pesticides were not applied to any fruit and plants used in the present work.

Physical analysis were performed in 40 fruit from each location using a caliper (Marberg) accurate to 0.05 mm and digital scale (Mettler Toledo, AB204) accurate to 0.1 mg. Pulp yield was determined for each fruit manually by separating into three different fractions: skin, flesh and kernel with use of latex gloves and knife. Each fraction was weighed separately soon after separation.

2.2 Chemical analysis

Full-mature fruit were used for total soluble solids (TSS) and total titratable acidity (TTA) content analysis. Analyses were performed in duplicate for each sample using 4 different samples for each location. For each sample, 8 fruit were manually pulped. TSS results were obtained using a digital refractometer Reichert, AR200, expressed in Brix. TTA was expressed in percentage of citric acid, analyzed in duplicate for each sample, using 4 different samples for each treatment or location. Total solids content were analyzed with four repetitions of 10 g of fruit from each location and dried in circulation heater at 105 °C for 24 hours (INSTITUTO..., 2008).

Pectin content was examined by mixing 4 g of lyophilized mume with 1:50 nitric acid 50 mM at 80 °C for 25 minutes. After filtration and cooling to 4 °C, the acid extract was mixed with 1:2 ethanol 96°GL at 4 °C and allowed to stand still for 30 minutes. After this period, was filtered and kept inside permeable bags overnight with ethanol 70% under agitation. Then, washed again with ethanol 95%, and dried at 40 °C. Pectin analysis was performed in triplicate.

2.3 Sample preparation for phenol and antioxidant analysis

For total phenol content and antioxidant power analysis, an extraction solution was previously prepared by mixing ethanol 70%: distilled water: formic acid 3% (80:20:1) as described by McGhie, Hunt and Barnett (2005) with modifications. Samples containing 8 fruit were frozen at -86 °C for at least 2 days before being lyophilized (Terroni, LD1500A). Next, 2.5 grams of dried pulp with peel were weighed in a centrifuge tube and mixed with 25 mL of the extraction solution. The product was placed in contact with the extraction solution at 15 °C for 24 hours and then immediately centrifuged (Celm, Combate) for 10 minutes at 2,232 g (3,500 rpm). Previous studies were performed to adjust the composition of the extraction solution, ratio of the dried product to the amount of extraction solution and extraction time to ensure the extraction of most of the phenolic compounds and reproducibility of the results. Samples were identified and maintained in closed flasks inside a freezer for up to 4 months for analysis of total phenol content and antioxidant activity.

For each location, fruit samples were prepared from 1 day after harvest (DAH) until complete maturation, up to 8 DAH.

2.4 Ferric-reducing antioxidant power (FRAP) assay

The reducing ability was determined by using the FRAP assay (BENZIE; STRAIN, 1996). The FRAP reagent was freshly prepared from 300 mmol (L⁻¹) acetate buffer (pH 3.63), 10 mmol (L⁻¹) tripyridyltriazine (TPTZ) made up in 40 mmol (L⁻¹) HCl and 20 mmol (L⁻¹) FeCl₃. All three solutions were mixed together in the ratio of 10:1:1 (v/v/v). An aliquot of 0.1 mL of the tested
sample solution was mixed with 3.0 mL of FRAP reagent. The absorption of the reaction mixture was measured at 593 nm after 15 minutes incubation at 26 °C. The reducing capacity was expressed as trolox equivalent (TE) concentration.

FRAP analyses were performed in duplicate.

2.5 Determination of total phenolic content

Total phenolic content of ethanol extracts was determined with the Folin-Ciocalteu colorimetric method (SINGLETON; ROSSI JUNIOR, 1965) with some modifications. Briefly, 0.1 ml extract was diluted with 8.4 ml distilled water and mixed with 0.5 ml Folin-Ciocalteu reagent by manual shaking for 15 20 seconds. After 3 minutes, 1.0 ml of 20% sodium carbonate solution was added. The reaction mixture was incubated at room temperature for 1 hour and absorbance was measured at 720 nm using a dual beam UV-Vis spectrophotometer (Shimadzu, Mod. UV-Mini 1240). Catechin (SIGMA, C-1251) was used as an analytical standard for total phenolic quantification and it was expressed in milligrams of catechin equivalents (CE) per gram of dried fruit.

TPC analyses were performed in triplicate.

2.6 Statistical analysis

Analysis were performed in random sequence of experiments and significant differences in treatment means were checked using STATISTICA software (version 5.5, StatSoft Inc., Tulsa, OK, USA) using Tukey’s test (p<0.05).

3 Results and discussion

3.1 Plant material

Fruit from CB and S were not significantly different (p<0.05) in size and weight; CB fruit presented diameter, height and mass of 2.239 ± 0.125 cm, 2.310 ± 0.125 cm and 6.892 ± 1.058 g, and S fruit, 2.130 ± 0.174 cm, 2.304 ± 0.199 cm and 5.701 ± 1.298, respectively. Fruit from SUN location were significantly larger in size and heavier in weight, with diameter 3.153 ± 0.264 cm, height 3.164 ± 0.215 cm and mass 16.909 ± 3.667 g.

The location SUN is the only commercial mume fruit producer and although fruit were larger and heavier compared to other locations, SUN fruit were still at the lower limit of weight for 1 hour and absorbance was measured at 720 nm using a dual beam UV-Vis spectrophotometer (Shimadzu, Mod. UV-Mini 1240). Catechin (SIGMA, C-1251) was used as an analytical standard for total phenolic quantification and it was expressed in milligrams of catechin equivalents (CE) per gram of dried fruit.

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The location SUN is the only commercial mume fruit producer and although fruit were larger and heavier compared to other locations, SUN fruit were still at the lower limit of weight as reported by Jun and Chung (2008) probably due to genetic improvement in Asian countries. Mume plants studied presented ratio values below 5, making product unacceptable by most consumers of fresh fruit. Thus, the consumption of mume fruit is recommended as processed-food products, where the ratio can be increased by sugar incorporation or acid dilution. These products show a healthy appeal, due to the high antioxidant capacity of the fruit (QUAST et al., 2011).

Pectin content decreased slightly during maturation, from 11.2 ± 0.1% to 10.8 ± 0.1% expressed on a dry matter basis. This provides approximately 1.32% on a wet matter basis that represents about half of pectin content in citrus fruit and is similar to apple pomace (BAKER, 1997).

3.2 Ferric-reducing antioxidant power (FRAP) assay

Fruit collected from the CB location showed in general higher FRAP values during maturation. FRAP values did not vary significantly during maturation of fruit in the CB location. Another tendency was observed in fruit from the S and especially from the SUN location, where antioxidant capacity tend to lower during maturation of mume fruit. Antioxidant capacity from the CB location ranged from 145 to 170 µMol TE per gram of dry fruit. In S and SUN locations, these results ranged from 96 to 160 µMol TE g⁻¹ DF. Previous studies performed in samples “hot-filled” and kept inside a glass container sealed for one year at room temperature resulted in an antioxidant capacity of about 70-80 µMol TE g⁻¹ DF. This indicates that the compounds responsible for the antioxidant capacity of mume products can be kept sealed for one year without need of refrigeration and maintain about 50% of its original antioxidant capacity.

The antioxidant capacity ranged from 21-35 µMol TE g⁻¹ of fresh fruit for all locations. These values are shown in Table 1 and are comparable to that of guavas and blueberries, 18-32 and 13-46 µMol TE g⁻¹ of fresh fruit, respectively (THAIPONG et al., 2006).

3.3 Determination of total phenolic content (TPC)

According to Table 2, CB fruit presented a trend of higher TPC values compared to other locations. This difference is probably a result of differences among cultivars, since climate is similar for all locations. It can also be observed that TPC did not vary during maturation of fruit, except for SUN location, which showed similar TPC values as fruits collected...
Table 1. FRAP assay of mume fruit collected from different locations and in different days after harvest – DAH.

<table>
<thead>
<tr>
<th>Samples</th>
<th>Antioxidant capacity (µMol TE g⁻¹ DF)*</th>
<th>Antioxidant capacity (µMol TE g⁻¹ FF)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>CB - 1 DAH</td>
<td>145.65 ± 18.62ab</td>
<td>27.24 ± 3.44ab</td>
</tr>
<tr>
<td>CB - 2 DAH</td>
<td>154.21 ± 24.18ab</td>
<td>30.51 ± 4.07ab</td>
</tr>
<tr>
<td>CB - 3 DAH</td>
<td>166.63 ± 11.09b</td>
<td>32.94 ± 2.73ab</td>
</tr>
<tr>
<td>CB - 4 DAH</td>
<td>153.01 ± 9.55b</td>
<td>31.17 ± 2.84ab</td>
</tr>
<tr>
<td>CB - 5 DAH</td>
<td>169.26 ± 5.84a</td>
<td>34.45 ± 4.85a</td>
</tr>
<tr>
<td>CB - 8 DAH (ripe fruit)</td>
<td>162.31 ± 3.88a</td>
<td>33.27 ± 1.51ab</td>
</tr>
<tr>
<td>S - 3 DAH</td>
<td>96.48 ± 22.59b</td>
<td>21.35 ± 5.47b</td>
</tr>
<tr>
<td>S - 4 DAH</td>
<td>125.48 ± 9.51ab</td>
<td>28.74 ± 2.25ab</td>
</tr>
<tr>
<td>S - 5 DAH</td>
<td>131.55 ± 10.14ab</td>
<td>30.81 ± 2.58ab</td>
</tr>
<tr>
<td>S - 6 DAH</td>
<td>116.52 ± 0.86a</td>
<td>26.91 ± 0.39ab</td>
</tr>
<tr>
<td>S - 7 DAH (ripe fruit)</td>
<td>116.34 ± 14.81ab</td>
<td>26.96 ± 4.65ab</td>
</tr>
<tr>
<td>SUN - 3 DAH</td>
<td>159.34 ± 6.95a</td>
<td>31.42 ± 1.39ab</td>
</tr>
<tr>
<td>SUN - 4 DAH</td>
<td>136.59 ± 12.33ab</td>
<td>28.18 ± 2.36ab</td>
</tr>
<tr>
<td>SUN - 5 DAH</td>
<td>119.21 ± 21.10b</td>
<td>23.57 ± 3.95b</td>
</tr>
<tr>
<td>SUN - 6 DAH</td>
<td>116.99 ± 32.44b</td>
<td>22.35 ± 6.43ab</td>
</tr>
<tr>
<td>SUN - 7 DAH (ripe fruit)</td>
<td>117.25 ± 10.63b</td>
<td>23.78 ± 1.88ab</td>
</tr>
</tbody>
</table>

*Least significant difference was 25.50 and 5.88 µMol trolox-equivalent (g⁻¹) of dry fruit and fresh fruit, respectively. **Means with different letters in the same column differ significantly (p<0.05) in Tukey's test.

Table 2. Total phenolic content of mume pulp+peel collected from different locations and in different days after harvest – DAH.

<table>
<thead>
<tr>
<th>Samples</th>
<th>Phenolic content (mg CE g⁻¹ DF)*</th>
<th>Phenolic content: Antioxidant capacity (mg CE:µMol TE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CB - 1 DAH</td>
<td>199 ± 21abc</td>
<td>1.366</td>
</tr>
<tr>
<td>CB - 2 DAH</td>
<td>207 ± 27abc</td>
<td>1.342</td>
</tr>
<tr>
<td>CB - 3 DAH</td>
<td>228 ± 16a</td>
<td>1.368</td>
</tr>
<tr>
<td>CB - 4 DAH</td>
<td>203 ± 31abc</td>
<td>1.327</td>
</tr>
<tr>
<td>CB - 5 DAH</td>
<td>226 ± 2ab</td>
<td>1.335</td>
</tr>
<tr>
<td>CB - 8 DAH (ripe fruit)</td>
<td>215 ± 15ab</td>
<td>1.325</td>
</tr>
<tr>
<td>S - 3 DAH</td>
<td>147 ± 27a</td>
<td>1.524</td>
</tr>
<tr>
<td>S - 4 DAH</td>
<td>161 ± 8a</td>
<td>1.283</td>
</tr>
<tr>
<td>S - 5 DAH</td>
<td>162 ± 3a</td>
<td>1.231</td>
</tr>
<tr>
<td>S - 6 DAH</td>
<td>144 ± 11a</td>
<td>1.236</td>
</tr>
<tr>
<td>S - 7 DAH (ripe fruit)</td>
<td>160 ± 31a</td>
<td>1.375</td>
</tr>
<tr>
<td>SUN - 3 DAH</td>
<td>226 ± 7a</td>
<td>1.418</td>
</tr>
<tr>
<td>SUN - 4 DAH</td>
<td>178 ± 20a</td>
<td>1.303</td>
</tr>
<tr>
<td>SUN - 5 DAH</td>
<td>164 ± 15a</td>
<td>1.376</td>
</tr>
<tr>
<td>SUN - 6 DAH</td>
<td>148 ± 11a</td>
<td>1.265</td>
</tr>
<tr>
<td>SUN - 7 DAH (ripe fruit)</td>
<td>167 ± 30a</td>
<td>1.441</td>
</tr>
</tbody>
</table>

*Expressed in mg of catechin-equivalent per gram of dry fruit. Different letters indicate significant difference (p<0.05). Least significant difference (LSD) was 22. **Means with different letters in the same column differ significantly (p<0.05) in Tukey's test.

from CB location. During ripening TPC values for SUN location where similar to those of fruits collected at S location. This information is highly relevant because most studies about Prunus mume products are related to health benefits of unripe fruit consumption. Therefore, it is possible to develop different products with ripe mume fruit, keeping health appeal.

Mume fruit collected from the CB location presented significantly higher TPC values but the antioxidant capacity was similar to collected from other locations. This suggests that CB fruit probably present different phenolic composition with lower antioxidant power that can be result of differences in cultivar or soil. In fruit collected from all locations, the increase of antioxidant power was directly proportional to the TPC increase and the ratio TPC:FRAP (mg CE:µMol TE) ranged from 1.23 to 1.52 for mume fruit for all locations, similar to studies in different European plum genotypes (RUPASINGHE; JAYASANKAR; LAY, 2006).

4 Conclusions

The present work showed that mume trees grown in São Paulo State give fruits significantly different with average diameter of 2.1-2.3 cm in CB and S location and 3.1-3.2 cm in SUN location. Mass of fruit ranged from 5.7 to 6.9 g in CB and S, 16.9 g in SUN. Yield of fruit flesh and TSS was not significantly different between locations and represented about 71.3% of total weight and 9.5-10.0°Brix, respectively. TTA decreased during maturation of fruit ranging from 4.0-5.7 to 2.0-3.8 g (100g⁻¹) citric acid. Pectin content ranged from 10.8 to 11.2% on a dry matter basis. The antioxidant capacity evaluated by the FRAP assay ranged from 96 to 170 µMol TE(g⁻¹) dry fruit and it was observed a direct proportionality to the TPC, that ranged from 144 to 228 mg CE (g⁻¹) DF.

References


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