Storage of ‘Palmer’ mangoes in low-oxygen atmospheres

Gustavo Henrique de Almeida Teixeira1*, José Fernando Durigan2

1 Fac. Ciênc. Farm. Ribeirão Preto, Univ. São Paulo, Dep. Anál. Clín., Toxicol. Bromatol., Av. do Café, s/n, Campus Univ. USP, Ribeirão Preto, SP-Brasil, CEP: 14.040-903, gustavo@fcfrp.usp.br

Storage of ‘Palmer’ mangoes in low-oxygen atmospheres.

Abstract — Introduction. Mango conservation under traditional refrigeration systems is not totally efficient due to the susceptibility of this fruit to chilling injury, but controlled atmosphere (CA) in association with low temperature can improve its storability and maintain fruit quality during storage. Thus, the aim of our study was to determine the effect of CA with varied concentrations of oxygen during cold storage (12.8 °C) of ‘Palmer’ mango fruit.

Materials and methods. Mature green mango fruit were stored in atmospheres with (1%, 5%, 10%, 15% and 21%) oxygen at (12.8 ± 0.6) °C and RH ~95%) for up to 28 days. A group of fruits without CA was stored in a cold room and served as tray-stored control. Fruit ripening was carried out in ambient conditions [(25.2 ± 0.6) °C, (92.8 ± 2.4)% RH] at intervals of 14 d.

Results and discussion. Fruits stored in low-oxygen concentrations [(1%, 5% and 10%) O2] had significantly lower rates of CO2 production after 14 d of cold storage. Fruit from all treatments were considered immature after 28 d of cold storage, and the mangoes kept at (1%, 5% and 10%) O2 maintained their initial firmness (119.9–125.6 N) when compared with those stored in higher oxygen atmospheres, which underwent a substantial loss of firmness (96.8–109.1 N). At low-oxygen levels, fruit also had lower contents of soluble pectin and total soluble sugars, whereas colour parameters were not affected by the atmospheres. After transfer from CA containers to ambient conditions, even from the lowest oxygen concentrations [(1% and 5%) O2], fruit ripened normally in just 8 d without presenting any low oxygen-related injury.

Brazil / Mangifera indica / fruits / controlled atmosphere storage / respiration / chemicophysical properties / quality / keeping quality
1. Introduction

Brazil is the world’s seventh largest mango producer [1] and in general its commercial production is based on the cultivation of American varieties, such as ‘Tommy Atkins’, ‘Haden’, ‘Kent’, ‘Keitt’ and ‘Palmer’. More recently, late season varieties have increased in importance and ‘Palmer’ has been the most preferable choice due to its quality characteristics and consumer preference.

At ambient temperature mango ripening occurs rapidly and fruit quality can be maintained for just 8 d [2]. Low temperatures can extend mango shelf-life to around 16 d; however, mango conservation under traditional refrigeration systems is not totally efficient due to its susceptibility to chilling injury at temperatures lower than 13 °C [3]. However, controlled atmosphere (CA) in association with low temperature can improve mango storability and maintain fruit quality during long-term storage [2, 4, 5]. Storage under CA has been used to extend the shelf-life of a wide range of mango varieties, e.g., ‘Kensington Pride’ [6], ‘Kent’ [7, 8], ‘Tommy Atkins’ [8, 9] and ‘Haden’ [10].

Kader recommended atmospheres containing 5% oxygen (O₂) and 5% carbon dioxide (CO₂) in order to extend mango shelf-life [11], although it is known that some varieties can tolerate higher CO₂ levels and levels of O₂ lower than 2% [8]. The recommendation of 5% O₂ and 5% CO₂ is commonly used during storage of mangoes under CA conditions. Lizada et al. reported that fruit of the ‘Tommy Atkins’ variety extended its shelf-life up to 31 d in CA containing 5% CO₂ and 5% O₂ [12]. However, these levels did not increase the shelf-life of ‘Kent’ mango, even when the CO₂ level was raised to 10% [7]. On the other hand, Lizada and Ochagavia reported that CA with 10% CO₂ and 5% O₂ increased the shelf-life of fruit of this variety up to 29 d, eight days more than the control [9].

Although promising, the responses of mango fruit to controlled atmosphere have been contradictory, as some varieties show just a tiny increase in shelf-life [13], whereas the shelf-life of other varieties can be extended more than one month [7]. Although it is possible to find studies regarding the use of CA during storage of mangoes, there is no recommendation for the ‘Palmer’ variety. The objective of our study was to determine the best oxygen concentration for the storage of ‘Palmer’ mango during cold storage and its effect on ripening.

2. Materials and methods

2.1. Plant material

Mature green mango fruits (Mangifera indica L. cv. ‘Palmer’) were obtained from a commercial orchard located in Taquaritinga, São Paulo State, Brazil. The maturity index was based on pulp colour (creamy-white = immature and yellow = mature) with fruit presenting titratable acidity (TA) = 0.76 g·100g⁻¹, pH = 3.55, soluble solids content (SSC) = 7.2%, [SSC / TA] ratio = 9.90, firmness = 127.50 N, and purple-green peel (L* = 38.16, Chroma = 9.31 and Hue = 346.90). After harvest, these fruits were transported to the laboratory within 1 h.

2.2. Controlled atmosphere treatments

The ‘Palmer’ mango fruits were placed into hermetic plastic containers (20 L) under a continuous humidified gas flow system of 100 mL·min⁻¹. The flow rate and gas mixtures were established using a mixing board with glass capillary tubes as flow regulators [14]. Compressed air was used as an oxygen (O₂) source and nitrogen (N₂) was obtained from cylinders (White Martins Gases Industriais Ltda., Sertãozinho, Brazil). Both gases were mixed in order to obtain the following oxygen levels: 1%, 5%, 10%, 15% and 21% (container-stored control). Supply and exhaust gas composition were monitored every day using a Dansensor Checkmate 9001 atmosphere analyser (PBI Dansensor, Denmark). A single CA container containing 36 fruit represented an experimental unit and was replicated three times for each CA composition. Similarly, a plastic tray containing 36 fruit served as tray-stored control and was replicated three times (tray-stored control). The storage temperature was (12.8 ± 0.6) °C (RH ~95%) and fruit ripening
was carried out in ambient conditions [(25.2 ± 0.6) °C, (92.8 ± 2.4)% RH] at intervals of 14 d [(0, 14 and 28) d], in order to verify the effect of CA on ripening. The evaluations were performed immediately after the fruits were withdrawn from CA storage [(0, 14 and 28) d] and when the fruit was considered ripe [(0 + 12), (14 + 8) and (28 + 8) d).

2.3. Respiration rate

Every two days during CA storage, thirty-six fruits of each oxygen concentration and the tray-stored control, three replicates, were sealed in 20-L containers at (12.8 ± 0.6) °C for one hour. Similarly, every two days after transfer to ambient conditions, two fruits of each oxygen concentration, three replicates, were sealed in 3.2-L containers at (25.2 ± 0.6) °C for one hour. Initially (0 hour) and after one hour, 0.3-mL gas samples from the container headspaces were analysed for CO₂ using a Finningan 9001 gas chromatograph (Finningan Corporation, San Jose, USA) equipped with a Porapack-N column and a thermal conductivity detector. Nitrogen, at a rate of 30 mL·min⁻¹, was used as the carrier gas. Data were integrated using Borwin 1.20 software (JMBS Developpements, Le Fontanil, France).

2.4. Weight loss

The fruits of all treatments were initially weighed on a Marte AS 2000 balance with 0.01 g precision (Marte, São Paulo, Brazil) and the difference in weight was calculated and expressed as a percentage (%).

2.5. Visual appearance

An untrained panel (10 members) evaluated the fruits based on a 1–5 scale, describing the overall fruit quality: 5, excellent; 4, very good; 3, regular; 2, bad; and 1, very bad.

2.6. Firmness

Pulp firmness was measured using an Effegi fruit tester (Bishop FT 327 Penetrometer, Alfonsine, Italy) with an 8.0-mm diameter tip. The peel from opposite sides was removed and two measurements per fruit were determined in the equatorial region [15].

2.7. Colour

Colour measurements ($L^*, a^*, b^*$) were taken on three fruits of each oxygen concentration and tray-stored control. These values were transformed into chromaticity ($C = \sqrt{a^* + b^*})$

and hue angle ($\text{Hue} = \tan^{-1} \left( \frac{b^*}{a^*} \right)$) according to McGuire [16]. A Minolta CR-400 (Minolta Corp., Osaka, Japan) was used to measure external colour. The mean of two readings per fruit (equator reading) was determined.

2.8. Chemical analysis

Mangoes (4 fruits × 3 replicates per treatment) without peels were homogenised in a blender and the homogenate was used to determine soluble solids content ( SSC), titratable acidity ( TA) and pH [17] (procedures 920.151, 932-12 and 945-27, respectively). The [SSC / TA] ratio was also calculated. The homogenates of all samples were frozen at –20 °C and subsequently used for determining total soluble sugars [18], reducing sugars [19], and total and soluble pectin contents [20, 21].

2.9. Statistical analysis

The experiment was laid out in a completely randomised factorial design with two factors: tray-stored control and the oxygen levels as the first factor and the storage duration [(0, 14 and 28) d] as the second factor. The data of the fruits of all treatments immediately removed from the CA conditions were subjected to analysis of variance, and the treatment means were compared using Tukey’s test at a significance level of $P < 0.05$. The data were analysed using the PROC MIXED procedure of the Statistical Analysis System [22]. Fruits transferred to ambient conditions were also statistically analysed according to a completely randomised design with five treatments (oxygen concentration) and three replicates of six fruits.
The lowest respiration rates presented by the fruits stored in the atmospheres with low oxygen levels ([1%, 5% and 10%] O₂) can be related to its effect on key regulatory steps of glycolysis and, consequently, in respiration, by inhibiting the activity of phosphofructokinase and pyruvate kinase enzymes in the conversion of fructose-6-phosphate into fructose-1,6-bisphosphate and phosphoenol pyruvate into pyruvate, respectively [23].

The adequate use of CA can also contribute to reduced fruit sensitivity to ethylene, mainly at levels of oxygen below 8% [24]. Atmospheres with low O₂ concentrations can inhibit the biosynthesis of ethylene by blocking the ethylene linkage to the receptor responsible for triggering the autocatalytic biosynthesis of this hormone [25]. However, Bender and Brecht reported that the oxygen level had little influence on the ethylene biosynthesis in ‘Kent’ and ‘Tommy Atkins’ mangoes [8], possibly because the oxygen level (3% O₂) was still adequate in relation to the apparent \( K_m \) of ACC oxidase (0.4% O₂) observed in mangoes by Kuai and Dilley [26].

During CA storage, ‘Palmer’ mango fruit, independently of the oxygen concentration, lost less weight than the fruits stored only in the cold room tray-stored control (figure 1). Controlled-atmosphere-stored fruits lost just 0.74% of their original weight, mainly due to the moisture control provided by the humidified gas flow, as the relative humidity in the CA containers was around ~95% compared with the cold room [(68.9 ± 5.9)%] tray-stored control. The difference in relative humidity, water loss, might have caused water stress and a consequent increase in the respiration rate and ethylene production of fruits from tray-stored control (figure 1), similar to results reported by Nakano et al. for ‘Tonewase’ persimmon [27].

Although tray-stored control fruits presented high weight loss values, according to the panellists the fruits were considered very good (4.0 ± 0.2) after 28 d of CA storage (figure 2). On the other hand, the fruits stored in CA received better scores as they were considered excellent throughout cold storage (figure 2), possibly due to the
increase in relative humidity of the gas flow. Shriveling affected the appearance of the tray-stored control treatment just after transfer to ambient conditions (figure 2).

The CA storage with different levels of oxygen did not affect the colour parameters (table I). Fruit were initially considered immature with dark ($L^* = 38.1 \pm 1.5$), not very saturated (chromaticity = $9.3 \pm 3.4$) and purple peel (Hue angle = $346.9 \pm 7.8$). The initial immaturity plus the effects of cold storage and its association with CA could have contributed to the absence of marked modifications in colour parameters (table I). 'Kensington Pride' mango fruit stored at $13 \degree C$ did not reach the maximum characteristic colour of this variety [28], similarly to mangoes of other varieties which presented reduced carotenoid development under cold storage [29, 30]. Therefore, atmosphere control with low $O_2$ concentration did not affect colour modification of 'Palmer' mango fruit during cold storage; in this regard, the temperature should have played the major role.

The fruit stored under CA containing (1%, 5% and 10%) $O_2$ remained firmer (> 120 N) than those of the other oxygen levels and tray-stored control, mainly after 14 d of cold storage (figure 2). Tray-stored control fruits and those stored in CA with (15% and 21%) $O_2$ (container-stored control) showed a sharp reduction in firmness, even though the pulp was considered very hard (~100 N), according to Mitcham and McDonald [31] (figure 2). The absence of great modification in firmness of the fruits stored in CA with (1%, 5% and 10%) oxygen can be related to the inhibition of the expression of many enzymes involved in the breakdown of starch molecules and pectic compounds [23, 32]. In 'Palmer' mangoes that remained firmer, stored in low oxygen, less starch mobilisation occurred, which was related to total soluble sugar (TSS) content, and reduced pectin solubilisation, which is responsible for cell wall integrity. Thus, the lowest TSS content presented in these fruits can be due to the reduced starch breakdown which led the fruits to remain firmer, as starch is responsible for mango pulp firmness [31]. Low levels of sugars (TSS) can also be related to lower respiratory activity of these groups of fruits (figure 1), as a great part of the energy required by the fruit is supplied through aerobic respiration, which involves the breakdown of several organic substances, starch among them [32]. Therefore, at low $O_2$ concentrations the breakdown of starch molecules into glucose, through the action of amylases and/or glucose-1-phosphate, by the enzyme phosphorylase, could have been reduced and have led the fruits of the atmospheres containing (1%, 5% and 10%) $O_2$ to present the lowest TSS contents (figure 3). On the other hand, tray-stored control fruit presented increases in both TSS and reducing sugar contents (figure 3) following what normally happens during mango ripening [33] and in water-stressed persimmon [27].
‘Palmer’ mango fruit stored in (1%, 5% and 10%) O₂ also presented lower soluble pectin contents than the fruits of tray-stored control and the atmosphere containing (15% and 21%) O₂ (container-stored control) (figure 4). Thus, the lower pectin solubilisation of these fruits indicates that the breakdown of cell wall compounds was inhibited, possibly due to a reduction of the activity of pectinolytic enzymes, which could have led these fruits to remain firmer even after 28 d of cold storage (figure 4).

Generally, during mango ripening soluble solids content (SSC) increases due to starch breakdown [34]; however, as previously reported, fruits stored under CA with (1%, 5% and 10%) O₂ presented the lowest sugar contents (TSS), which probably reflected on the SSC (figure 4). During storage, tray-stored control fruits and those stored in the atmosphere containing higher oxygen concentrations (15% O₂ and 21% O₂-container-stored control) showed a sharp increase in SSC, mainly the tray-stored

![Figure 3. Total soluble sugar and reducing sugar contents of ‘Palmer’ mango fruit stored at 12.8 °C under controlled atmosphere with different levels of oxygen for up to 28 d.](image)

### Table I.

Effect of atmospheres on colour, pH, titratable acidity (TA), soluble solids contents (SSC) and [SSC / TA] ratio of ‘Palmer’ mango fruit after 28 d of storage at 12.8 °C.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Colour</th>
<th>pH</th>
<th>Titratable acidity&lt;sup&gt;a&lt;/sup&gt; (g·100 g&lt;sup&gt;−1&lt;/sup&gt;)</th>
<th>[SSC / TA] ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Atmospheres (A)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td>36.75</td>
<td>11.67</td>
<td>249.82</td>
<td>3.58</td>
</tr>
<tr>
<td>21% O₂</td>
<td>37.83</td>
<td>11.12</td>
<td>277.70</td>
<td>3.56</td>
</tr>
<tr>
<td>15% O₂</td>
<td>37.80</td>
<td>10.75</td>
<td>275.03</td>
<td>3.57</td>
</tr>
<tr>
<td>10% O₂</td>
<td>37.95</td>
<td>11.04</td>
<td>299.82</td>
<td>3.57</td>
</tr>
<tr>
<td>5% O₂</td>
<td>36.72</td>
<td>10.61</td>
<td>275.03</td>
<td>3.61</td>
</tr>
<tr>
<td>1% O₂</td>
<td>38.56</td>
<td>10.23</td>
<td>313.17</td>
<td>3.58</td>
</tr>
<tr>
<td><strong>Storage (B)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 d</td>
<td>38.19</td>
<td>9.31 c</td>
<td>240.90 b</td>
<td>3.55 b</td>
</tr>
<tr>
<td>14 d</td>
<td>36.87</td>
<td>12.42 a</td>
<td>299.45 a</td>
<td>3.60 a</td>
</tr>
<tr>
<td>28 d</td>
<td>37.75</td>
<td>10.99 b</td>
<td>305.34 a</td>
<td>3.58 ab</td>
</tr>
<tr>
<td><strong>Interaction</strong></td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>(A) × (B)</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td><strong>Standard deviation</strong></td>
<td>1.84</td>
<td>1.73</td>
<td>58.56</td>
<td>0.07</td>
</tr>
</tbody>
</table>

<sup>a</sup> Citric acid.

Means within a main effect followed by the same letter in the column are not significant by Tukey’s test (p < 0.05).

NS, interaction not significant.
Storage of ‘Palmer’ mangoes

3.2. Ambient ripening

During ripening in ambient conditions (25.2 °C, 92.8% RH), the respiration rates of ‘Palmer’ mango fruits previously stored at different oxygen concentrations for 14 d increased even for those fruits stored at low O₂ concentrations [1%, 5% and 10% O₂], but without presenting any significant difference among treatments (figure 5). In ambient conditions, fruit showed a steady increase in CO₂ production rates without a climacteric peak (figure 5). On the other hand, after 28 d under CA storage, the oxygen concentrations did affect the respiratory activity of the fruit transferred to ambient conditions (figure 5). We observed that the fruit previously stored in atmospheres with low O₂ concentrations (1%, 5% and 10%) initially showed the lowest respiratory activities, which just levelled with the 21% O₂ (container-stored control) rates after 5 d in ambient conditions (figure 5). These differences can be related to the regulation of the metabolic pathways, which had led the mangoes previously stored at (1%, 5% and 10%) O₂ to show reduced respiratory activities and, consequently, delaying ripening. Singh and Pal reported the same trend during guava ripening in ambient conditions after CA storage, and they related the suppression and delay of the climacteric rise to the residual effect of CA on respiration [35]. Despite these differences, the increase in CO₂ production by ‘Palmer’ mango fruit after a short period in ambient
conditions can indicate the non-existence of any irreversible damage on the mitochondrial structure as a consequence of the prolonged exposition to low O₂ concentrations, differently from what happened in extreme conditions (50% CO₂ and 75% CO₂) of atmospheric control [8]. Thus, ‘Palmer’ mango fruits can be stored in atmospheres containing between (1% and 10%) O₂ without affecting their respiratory metabolism.

The fruits previously stored in atmospheres containing low O₂ concentrations (1%, 5% and 10%) for 28 d ripened normally after 8 d in ambient conditions (table II). Soluble solids content increased during the 8-d post-storage holding period especially in these treatments, as observed for those kept in CA with (15% and 21%) O₂ (tray-stored control fruit was destroyed during transfer to ambient conditions – missing data). The titratable acidity declined dramatically during ripening of mango fruit (table II). The data are consistent with what is normally observed during ‘Palmer’ mango ripening [36]. Therefore, ‘Palmer’ mango fruit can be stored in atmospheres containing between (1% and 10%) O₂ without affecting its ripening process. It should be noted that these levels are quite similar to the 5% O₂ recommended for other American mango varieties.

4. Conclusions

The mango cultivar ‘Palmer’ may be stored at low temperature (12.8 °C, ~95% RH) supplemented with atmospheres containing (1% to 10%) O₂ for up to 28 d. The most significant effects of controlled atmosphere in mango included reduction of the respiration rate, delayed ripening and maintenance of the fruit quality. Controlled atmosphere storage seems to be promising for shipping of ‘Palmer’ mango fruit by marine transport to distant markets, which may take three or four weeks. However, the CA requirements need to be evaluated for storing and transporting mangoes in other atmospheres and at temperatures other than 12.8 °C because changing storage conditions may alter fruit quality.

Acknowledgments

The authors would like to thank FAPESP for sponsoring this research (Proc. 05/56159-1) and providing the postdoctoral scholarship (Proc. 05/56160-0) of the first author. We would also like to thank José Maria Sigrist for his special help during the setting up of the flowboard system and Ogata Citrus for providing the fruit for this experiment.
Storage of ‘Palmer’ mangoes

Table II.
Effect of atmospheres on weight loss, initial and final colour, firmness, visual appearance, titratable acidity (TA), soluble solids content (SSC), [SSC / TA] ratio, and pH of ‘Palmer’ mango fruit after 28 d of storage at 12.8 °C plus 8 d in ambient conditions (25.2 °C).

<table>
<thead>
<tr>
<th>Atmosphere</th>
<th>weight loss (%)</th>
<th>Initial colour&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Final colour</th>
<th>Firmness (N)</th>
<th>Appearance&lt;sup&gt;b&lt;/sup&gt;</th>
<th>TA (g 100 g&lt;sup&gt;-1&lt;/sup&gt;)</th>
<th>SSC (%)</th>
<th>[SSC / TA] ratio</th>
<th>pH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control&lt;sup&gt;c&lt;/sup&gt;</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>21% O&lt;sub&gt;2&lt;/sub&gt;</td>
<td>7.31</td>
<td>37.73</td>
<td>10.12</td>
<td>240.75</td>
<td>39.29</td>
<td>13.44</td>
<td>66.71</td>
<td>4.63</td>
<td>4.0</td>
</tr>
<tr>
<td>15% O&lt;sub&gt;2&lt;/sub&gt;</td>
<td>7.35</td>
<td>37.80</td>
<td>9.83</td>
<td>200.27</td>
<td>38.55</td>
<td>13.36</td>
<td>65.41</td>
<td>8.53</td>
<td>4.0</td>
</tr>
<tr>
<td>10% O&lt;sub&gt;2&lt;/sub&gt;</td>
<td>8.51</td>
<td>37.39</td>
<td>10.22</td>
<td>236.00</td>
<td>38.03</td>
<td>14.00</td>
<td>97.79</td>
<td>5.67</td>
<td>4.0</td>
</tr>
<tr>
<td>5% O&lt;sub&gt;2&lt;/sub&gt;</td>
<td>7.77</td>
<td>38.22</td>
<td>11.49</td>
<td>311.04</td>
<td>37.40</td>
<td>14.64</td>
<td>129.31</td>
<td>6.55</td>
<td>4.0</td>
</tr>
<tr>
<td>1% O&lt;sub&gt;2&lt;/sub&gt;</td>
<td>7.03</td>
<td>39.12</td>
<td>10.19</td>
<td>209.84</td>
<td>38.11</td>
<td>14.56</td>
<td>100.28</td>
<td>6.30</td>
<td>4.0</td>
</tr>
<tr>
<td>F value</td>
<td>0.3582</td>
<td>0.1933</td>
<td>0.5872</td>
<td>0.1436</td>
<td>0.3953</td>
<td>0.9065</td>
<td>0.8447</td>
<td>0.7077</td>
<td>0.5121</td>
</tr>
<tr>
<td>LSD</td>
<td>2.41</td>
<td>2.27</td>
<td>3.50</td>
<td>136.67</td>
<td>3.06</td>
<td>5.63</td>
<td>21.57</td>
<td>9.02</td>
<td>0.46</td>
</tr>
</tbody>
</table>

<sup>a</sup> Colour (L*, luminosity; Chroma, chromaticity; Hue, hue angle).
<sup>b</sup> Visual appearance (5, excellent and 1, worst - very bad).
<sup>c</sup> Air: fruit was destroyed during transferral to ambient conditions.

There was no significant different among treatments according to the F value determined by Tukey’s test (p < 0.05).

References


[7] Trinidad M., Bósquez E., Escalona H., Dias de León F., Pérez Flores L., Kerbel C., Ponce de León L., Muñoz C., Pérez L., Controlled atmosphere (5% CO<sub>2</sub> – 5% O<sub>2</sub> and 10% CO<sub>2</sub> – 5% O<sub>2</sub>) do not significantly increase the shelf life of refrigerated Kent Mangoes, Acta Hortic. 455 (1997) 643–653.


Almacenamiento de los mangos ‘Palmer’ en atmósferas pobres en oxígeno.

Resumen — Introducción. La conservación de los mangos en refrigeración tradicional no es del todo eficaz, dado a la sensibilidad al frío de dicho fruto. Sin embargo, un almacenamiento en atmósfera controlada (AC) y a baja temperatura puede mejorar su aptitud de conservación y mantener la calidad de los frutos durante este periodo. Por ello, nuestro estudio quiso determinar el efecto que tienen ciertas atmósferas controladas de diversas concentraciones de oxígeno en los mangos ‘Palmer’ almacenados en frío (12,8 °C). Material y métodos. Se almacenaron frutos maduros de mangos verdes en atmósferas controladas a (1 %, 5 %, 10 %, 15 % y 21 %) de oxígeno, (12,8 ± 0,6) °C y HR ~ 95 % durante 28 días. Un lote de frutos se almacenó sin AC, en cámara fría, como lote testigo. La maduración de los frutos se realizó a temperatura ambiente [(25,2 ± 0,6) °C, y (92,8 ± 2,4) % HR] durante periodos de 14 días. Resultados y discusión. Los frutos conservados en atmósferas de flojas concentraciones de oxígeno [(1 %, 5 % y 10 %) O₂] presentaron las tasas de producción de CO₂ considerablemente más bajas, tras 14 días de almacenamiento en frío. Los frutos de todos los tratamientos se consideraron inmaduros, tras 28 días de almacenamiento frigorífico. Los mangos mantenidos a (1 %, 5 % y 10 %) O₂ conservaron su firmeza inicial [(119,9 a 125,6) N], en relación con los que se almacenaron en atmósfera de concentraciones de oxígeno más elevadas, los cuales, a su vez, padecieron una pérdida sustancial de firmeza [(96,8 a 109,1) N]. En atmósferas de floja concentración de oxígeno, los frutos presentaron también los contenidos más bajos de pectinas solubles y de azúcares solubles totales, mientras que los parámetros de color no se vieron afectados por la atmósfera. Después de transferir a temperatura ambiente los contenedores almacenados en AC, incluso en las concentraciones más flojas de oxígeno [(1 % y 5 %) O₂], los frutos maduraron normalmente en sólo 8 días, sin presentar ningún daño relacionado a los flojos contenidos de oxígeno.

Brasil / Mangifera indica / frutas / almacenamiento atmósfera controlada / respiración / propiedades fisicoquímicas / calidad / aptitud para la conservación