QUALITY AND ANTIOXIDANT PROPERTIES OF WHOLE AND FRESH CUT ‘CHERRY’ PEPPERS DURING STORAGE AT 10°C

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Abstract: Changes on quality and antioxidant properties of whole and fresh cut (without the core) peppers (Capsicum annuum, L. cv. ‘Cherry’), stored for 10 days at 10°C in PET trays covered with PVC film were studied. During the storage their general appearance, fungal decay, color, respiration rate, sugar content and firmness were evaluated. Changes on carotenoids, total phenols, ascorbic acid and antioxidant capacity were also studied. Immediately after cutting and throughout the storage, an increase in respiration rate was found. No differences in color evolution were observed during storage between whole and fresh cut fruits. There were no differences in firmness, total sugars and carotenoids between whole and cut fruits either. Total phenols, ascorbic acid and antioxidant capacity did not change largely in both intact and without core peppers during the experiences. However at the end of the storage period (10 days), whole pepper fruit displayed good general appearance and overall higher quality than de-cored fruit. Results suggest that Cherry peppers could be marketed as fresh-cut peppers in the type of without core fruits, although they would be only stored for 6-7 days at 10°C.

Palabras claves: Pimientos Cherry; capacidad antioxidante; fenoles totales; ácido ascorbico

Key words: Cherry peppers, antioxidant capacity, total phenols, ascorbic acid

INTRODUCCIÓN

Cherry peppers (Capsicum annuum, L. cv. ‘Cherry’) are small (diameter of 26 to 32 mm), fruits with bright red color and sweet taste. While their size and appearance might be appealing, they present a large number of seeds making tedious home prepara-

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tion operations. Fresh-cut products are lightly processed fruits and vegetables with increased functionality maintaining most properties of fresh products (Salunkhe et al., 1991). However, in general fresh cut products deteriorate rapidly and show reduced shelf-life compared to whole fruits and vegetables (Conesa et al., 2007b). During their preparation there are different steps as selection, washing, cutting, seed separation, draining and packaging, which have a great influence in the organoleptic, nutritional and microbiological characteristics of the final product. These vegetables can show symptoms of deterioration as dehydration, color and texture changes, loss juice and/or microbial growth during the storage. In order to minimize these effects and to achieve a high quality product, it is crucial to select properly the variety and maturity of the raw material, process, packaging and storage conditions (Watada et al., 1999).

Peppers (*Capsicum annuum*, L.) are a good source of bioactive compounds (Howard et al., 2000). Ascorbic acid, phenolic and carotenoid compounds contribute to the nutritional value and to the health-promoting properties of sweet peppers (Raffo et al., 2007).

There are no studies about the postharvest behavior of Cherry peppers and less of the evolution of physical and compositional parameters of lightly processed-Cherry peppers during refrigerated storage. The purpose of this paper is to evaluate the changes on quality parameters and antioxidant properties of whole and cut ‘Cherry’ peppers during storage at 10°C.

**MATERIALS AND METHODS**

**Plant material.** Cherry peppers (*C. annuum*, L. cv. ‘Cherry’) produced in Corrientes city (Argentina), having 95-100% red color, were used for this study. Fruit free of damage and uniform size (weight: 10 ± 1.1 g) were selected.

**Processing and storage.** Peppers were carried to the lab within 24 hours of being harvested. Fruits were washed with chlorinated water (100 mg/L, 20 seconds) and then divided into three lots. Cores of fruits from one of the lots were removed, a small area of tissue was extracted around of their peduncle with a 13 mm diameter core-borer and their seeds and placental tissue were eliminated (without core). Peppers from the second group were cut into halves (from which seeds were also removed), and those from the third lot were not cut. Fruits from the three lots were packaged in crystalline PET trays and covered with PVC film. Twelve trays for each treatment were prepared, containing each one approximately 70 g (7 fruits). The trays were stored at 10°C for 10 days and were sampled at day 0 and during storage (3 trays separated by type of cut and storage time). Samples were immediately processed or otherwise frozen at -20°C until analysis. The complete experiment was repeated three times. Since results from the different experiments showed a similar trend data from the first one is shown. Measurements determinations were done in triplicate on vegetal material corresponding to different combinations type of cut/storage time.

**Sensory quality.** A subjective evaluation of the whole or cut- fruits were realized, in a storage time determined, examining visually their general appearance (taking into
account color, brightness, firmness, dehydration). The percentage of fruits which presented macroscopic fungal growth was visually evaluated.

In order to evaluate the general appearance, the following scale was used: 1 = very good or fresh appearance, 2 = good, 3 = fair, 4 = poor. The index of general appearance (I) was calculated for a determined time, through this formula:

\[ I = \frac{1n + 2n + 3n + 4n}{N} \]

where:
- n: number of fruits or pieces in that category.
- N: total number of examined fruits or pieces.

It was considered that fruits lost market quality when I >2.

**Color.** Fruit color was evaluated with a colorimeter (Minolta, Model CR-300), by measuring the \( L^* \), \( a^* \) and \( b^* \) parameters in three zones of each fruit. Ten fruits from each type of cut and storage time were analyzed.

**Respiration rate.** Ten fruits were placed in a hermetic jar, the CO\(_2\) produced by fruits was determined, during an hour, realizing the test every 5 minutes, using a sensor IR (Alnor Compu Flow® Model 8650). From the obtained straight line the CO\(_2\) production rate was calculated. Results were expressed as µL CO\(_2\)/g h.

**Total sugars.** Five grams of tissue was ground with 30 mL of ethanol. The mixture was homogenized for 15 min in a homogenizer stirrer (GLAS-COL, Model 099C K4424, Terre Huate, IN, USA) and then centrifuged at 5000 x \( g \) for 10 min. The supernatant was utilized to determine total sugar content by the anthrone method (Southgate, 1976). The absorbance was measured at 620 nm. A calibration curve was prepared by using glucose as a standard. Results were expressed as grams of glucose per 100 grams of fresh weight (f.w.).

**Firmness.** The maximum force of penetration was recorded. Firmness was measured by using a Texture Analyzer (TA-XT2i) by compression tests. Each fruit piece was penetrated 3 mm with a 3 mm diameter flat probe at a rate of 0.5 mm/s on the equatorial zone. Fifteen fruits of similar size were used. Eight determinations were done in each fruit.

**Total carotenoids.** Carotenoids determination was performed by visible spectrophotometry. Five grams of tissue was ground in 35 mL of acetone and homogenized for 15 min. The mixture was vacuum filtered, washing the residue with acetone until a complete disappearance of color in the tissue. The volume of the acetonic extract was measured. Afterwards, 25 mL of it were transferred to a separating funnel and then the carotenoids were extracted with 25 mL of petroleum ether 35-60 bp. The volume of the ethereal extract was measured and its absorbance was determined at 450 nm. The extinction coefficient was 2.5x10\(^5\) mL/g cm (Davies et al., 1970). Results were expressed as micrograms of \( \beta \)-carotene per gram of fresh weight.

**Total phenols.** Five grams of tissue was ground in 30 mL of ethanol. The mixture was homogenized for 15 min in a homogenizer stirrer (GLAS-COL, Model 099C K4424, Terre Huate, IN, USA) and then centrifuged at 5000 x \( g \) for 10 min. Total phenolic content in the supernatant was determined using Folin-Ciocalteau reagent (Singleton et al.,
The absorbance was measured at 760 nm and the total phenols content was calculated using 1.13x10^{-3} mol/L clorogenic acid as standard. Results were expressed in milligrams of clorogenic acid per gram of fresh weight.

Ascorbic acid content. The content in ascorbic acid of peppers was assessed by HPLC using a Shimadzu LC-10AT chromatograph, with a UV-visible detector Shimadzu SPD-10A. Ten grams of sample were homogenized in 0.05 mol/L H_3PO_4 (Nisperos-Carriedo et al., 1992). The mixture was filtered and centrifuged at 10000 x g for 20 min at 4 ºC. The resultant supernatant was filtered prior to the injection into the chromatograph, through nylon membranes of 0.45 µm. The mobile phase was methanol:H_2O (30:70), pH: 2.8 (with 0.05 mol/L H_3PO_4), at a flow rate of 1mL/min and the chromatographic column used was Supelcosil LC 18.5 µm x 250 mm. Absorbance at 260 nm was determined. For identification and quantification a 1.53 x 10^{-3} mol/L standard ascorbic acid solution was used. Results were expressed in milligrams of ascorbic acid per 100 grams of fresh weight.

Antioxidant capacity. In order to determine the antioxidant activity of fruits the radical chromogen 2,2-diphenyl-1-picrylhydrazyl (DPPH●) in methanol solution was used (Brand-Williams et al., 1995). Five grams of tissue were ground in 30 mL of ethanol by using a homogenizer stirrer (GLAS-COL, Model 099C K 4424, Terre Huate, IN, USA). The extract was centrifuged at 5000 x g for 10 min. Different aliquots of supernatant (0-600 µL) were added to test tubes containing 3.4 mL of 7.61x10^{-5} mol/L DPPH● in methanol prepared daily (final volume of reaction: 4 mL). A bsorbance at 517 nm was measured with a spectrophotometer Metrolab 1700 at different times until the reaction reached a plateau. The % of remaining DPPH● against the volume of extract was then plotted to obtain the amount of extract necessary to decrease the initial DPPH● concentration by 50%, which was defined as EC_{50}. Results were expressed in milligram of fresh weight.

Statistical analysis. Experiments were performed according to a factorial design. The factors were the type of cut and the storage time. Data were statistically analyzed by the analysis of variance (ANOVA), and means were compared by the test LSD at a significance level of 0.05 (InfoStat, 2002).

RESULTS AND DISCUSSION

Fig. 1 shows the index of general appearance (I) during storage at 10°C. The general aspect of whole peppers relatively unchanged during storage, reaching a good general appearance (I = 2) after 10 days at 10°C. In contrast the cut fruits showed a continuous and more rapid decline in quality, reaching an index between good and fair (I = 2.5-2.7) after 10 days. However, the cut peppers were still marketable at 6 days of storage.
Fig. 1: Index of general appearance (I) in whole (□), de-cored (■) and halved ( □ ) Cherry peppers stored during 10 days at 10°C. (LSD_{0.05} = 1.2).

Fig. 2: Changes in (a) lightness (L*) and (b) a* value of whole (□) and de-cored (■) fruit stored at 10°C. *Value significantly different from that corresponding to whole peppers (LSD_{L*} = 0.46; LSD_{a*} = 1.14).
In the case of decay, higher incidence was observed in cut halves fruits after 10 days at 10°C (approximately 50% of fruits) (data not shown). Since the halved peppers were the most deteriorated, the following experiences were performed with only a type of cut (without core fruits).

Lightness parameter (L\(^*\)) presented an initial value of 35.6 ± 1.0 (Fig. 2a), which was close to the reported for peppers cv. Zafiro (90% red) (Vicente et al., 2005). No differences in fruit lightness (L\(^*\)) were found between intact and without core fruits until the eight day of storage at 10°C when without core peppers showed a value of L\(^*\) 2.5% lower than that of control fruits. The parameter a\(^*\) showed a similar evolution (Fig. 2b), for both treatments. Concerning parameter b\(^*\), there were no differences between control and cut fruits during storage (data not shown). No variation in superficial color parameters during storage could be due to the fact that the fruits were already at advanced ripening stage.

Fig. 3: Changes in (a) respiration rate, (b) sugar content and (c) firmness of whole (□) and decored (■) fruit stored at 10°C. (LSD\(_{\text{respiration rate}}\) = 28.99; LSD\(_{\text{sugar content}}\) = 0.1; LSD\(_{\text{firmness}}\) = 1.16).
Peppers are fruits which generally show a low respiration rate, (Barth et al., 2004). Initial levels of CO₂ production in whole Cherry peppers were 32.26 ± 2.07 µL/g h, and did not change significantly during storage at 10ºC (Fig. 3a). These results were similar those reported by other authors for other pepper varieties (González-Aguilar et al., 1999).

One of the changes promoted as response to the wound during the preparation of fresh-cut vegetables is the respiration increase (Barth et al., 2004; Beaulieu et al., 2003). Respiration rate increased to value of 44.84 ± 2.54 µL/g h immediately after the cut, presenting a continuous increment throughout the storage (Fig. 3a). At the end of the storage period, the CO₂ production of cut peppers increased about 4 fold, which may be attributed to the deterioration observed. A similar behavior was informed for green chilly peppers stored for 6 days at 10 ºC (Kang et al., 1997), and for diced green and red bell peppers (El-Bassuoni et al., 1994).

The main sugars influencing peppers fruit taste are glucose, fructose and to a lower extent sucrose. Initial values of total sugars were 1.87 ± 0.08 g glucose/100 g fresh weight. During storage sugar content remained practically invariable. Pepper processing did not cause variations in total sugar contents (Fig. 3b). Similarly, Conesa et al. (2007a) did not find changes in total sugars in red peppers cut in cubes after 10 days at 5ºC. While, Raffo et al. (2007) informed that glucose and fructose of red peppers were constant during 21 days at 7.5ºC. Contrarily, Raffo et al. (2008) observed a significant increase for sliced red peppers stored for 9 days at 8 ºC, due to the effect of concentration related to the water loss. In our experiences there was only a slight dehydration.

Tissue softening is a very serious problem with fresh-cut fruit products that can limit shelf-life. Vicente et al. (2005) reported a reduction in Zafiro peppers upon storage for 12 days at 10º C. González Aguilar et al. (2004) found a continuous decrease of firmness in green cut peppers, stored in MAP during 14 days at 10ºC. The authors related this decrease to fungal growth. Toivonen et al. (2004) found a decrease in the firmness of green cut peppers stored in modified atmosphere at 10ºC for 10 days. However in the present study the firmness virtually did not change in whole fruits during storage at 10ºC. In contrast, firmness decreased slightly in de-cored peppers, reaching values 13% lower than the whole fruits ones at the end of storage (Fig. 3c) which could be related to the slight dehydration and damage observed.

The Cherry peppers presented initially total carotenoid content expressed as β–carotene of 276 ± 9 µg/g fresh weight. At the end of the storage at 10ºC, a slight increase of 8% relative to the initial carotenoids content was detected in whole fruits (Fig. 4). Even so, this increase was not reflected in superficial color evolution. Vicente et al. (2005) found an increase of carotenoids in cv. Zafiro 90% red whole peppers at day 12 of storage at 10º C.
No significant differences in pigments content were observed between control and without core fruits (Fig. 4). In other assays, fruits with initial carotenoids content of 204 ± 4 µg β-carotene/g fresh weight showed an increase of 40% during the storage of whole peppers similar to the evolution of without core fruits.

Total phenols initial level was 3.90 ± 0.20 mg chlorogenic acid/g fresh weight. Total phenols values did not change significantly during storage at 10ºC for both assays conditions (whole and without core fruits) (Fig. 5a). Similar results were reported in cut red peppers cv. ‘Festos’ (Sgroppo et al., 2005), whereas a decrease in total phenols level in sliced green peppers stored at 7ºC for 10 days of storage were observed by Toivonen et al. (2004).

The initial ascorbic acid content was 62.7 ± 1.7 mg/100g fresh weight and it remained relatively constant in whole fruits during the storage period (Fig. 5b). Hussein et al. (2000) did not find changes in the content of vitamin C in green peppers stored at 4ºC. Otherwise, Toor et al. (2006) found that a relative stability of ascorbic acid content in stored tomatoes at 7ºC could be related to the acidity and the presence of phenols and flavonoids in tissue. On the other hand, Raffo et al. (2007) informed a slight increase of ascorbic acid content in whole red peppers stored for 21 days at 7.5ºC.
Fig. 5: Changes in (a) total phenols, (b) ascorbic acid content and (c) antioxidant capacity of whole (□) and de-cored (■) fruit during storage at 10°C. Bars represent standard error of the mean.

The content of ascorbic acid remained almost constant in without core Cherry peppers during storage at 10°C. In addition, no differences were observed between control and de-cored fruits (Fig. 5b). González-Aguilar et al. (2004) found that ascorbic acid content of cut green peppers (cv. ‘Wonder’) did not change during the storage at 10°C, while Raffo et al. (2008) reported an increment in ascorbic acid content in sliced red peppers stored at 8°C, relating those changes to the concentration effect associated to water loss. On the other hand, Hussein et al. (2000) informed a vitamin C content decreased in sliced green peppers stored at 4°C.

Antioxidant capacity expressed as $EC_{50}$ was determined using the radical chromogen DPPH$^\bullet$. The initial $EC_{50}$ was $41.2 \pm 3.6$ mg fresh weight. Higher values of $EC_{50}$ represent low values of antioxidant activity. Whole fruits showed an $EC_{50}$ increase of 35% at the third day of storage at 10°C, and then it decreased until the end of storage reaching a no different significant value compared to the initial one at day 10 (Fig. 5c.).
Vicente et al. (2005) in another variety of whole sweet peppers (cv. 'Zafiro'), did not observe modifications in the antioxidant capacity at day 12 of storage at 10°C. EC50 in without core Cherry peppers relatively unchanged during the experience. (Fig. 5c). A antioxidant activity decrease more than 20% of the initial level was detected in diced red peppers cv. 'Margarita' at day 7 of storage at 11°C (Sgroppo et al., 2004).

According to the obtained results, it is evident that no changes relative to the initial antioxidant activity were observed in both whole and without core peppers at the end of storage. Lana et al. (2006) did not find differences in the total antioxidant activity, measured using a lipid peroxidation inhibition assay, between whole and sliced tomatoes during the refrigerated storage. Although, Odriozola-Serrano et al. (2008) informed an antioxidant capacity decrease in fresh cut tomatoes during the storage.

CONCLUSIONS

Whole Cherry peppers maintained their quality during storage for 10 days at 10°C, presenting slight variation of chemical-physical parameters analyzed. However, de-cored fruits conserved their quality very well for 6-7 days of storage at 10°C without showing noticeable changes in the carotenoid content, total phenols, ascorbic acid and antioxidant capacity. Due to the scarce variation in the antioxidant characteristics, light processing did not decrease the nutritional quality of the product. Results suggest that Cherry peppers could be marketed as fresh-cut Cherry peppers in the type of without core fruits. Further complementary treatment to refrigeration might be evaluated in fresh cut peppers intended for storage periods longer than 6-7 days.

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