Enhancement of the antioxidant capacity and reduction of chilling injury in ‘Douradão’ peaches refrigerated under pre-storage and modified atmosphere

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ABSTRACT. The present study aimed to investigate the effects of pre-storage for 24h and 48h and different modified atmosphere packaging-MAP treatments (using low-density polyethylene – LDPE) on the antioxidant compounds and the postharvest quality of ‘Douradão’ peaches under cold storage. The peaches were submitted to the following treatments: a control (no packaging or pre-storage) and pre-storage for 24h or 48h at 25°C combined with passive MAP (LDPE 60 µm or 80 µm). After the application of the treatments, the fruits were stored at 0°C for 30 days plus 3 days at 25°C to simulate marketing conditions. The results showed that on the simulated marketing period, all the treatments (except for the control) reduced woolliness and internal browning. Furthermore, both pre-storage and LDPE MAP enhanced the polygalacturonase (PG) activity, the levels of phenolic compounds and the antioxidant capacity. Thus, these results suggest that the pre-storage associated with LDPE packaging increases the shelf life of peaches by up to 30 days under cold storage.

Keywords: Prunus persica (L.) Batsch; stone fruit; woolliness; internal browning; postharvest.

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Introduction

The peach [Prunus persica (L.) Batsch] is the most important species of the genus Prunus and one of the most significant fruits in terms of production (Cantin et al., 2010). It is a highly perishable climacteric fruit that deteriorates rapidly at room temperature (Lurie & Crisosto, 2005). Storage at temperatures of approximately 0°C has been used to extend the shelf life of peaches by 3 - 4 weeks depending on the cultivar. However, long storage periods at low temperatures may cause chilling injuries such as woolliness and internal browning. These injuries are only perceived when the fruit is exposed to room temperature during marketing (Lurie & Crisosto, 2005). According to Zhou, Ben-Arie, and Lurie (2000), the woolliness in stone fruits is a chilling injury caused by an imbalance in the activity of pectinolytic enzymes during the maturation stage. This imbalance is caused by increased pectinmethylsterase (PME) activity and decreased polygalacturonase (PG) activity. It results in the accumulation of pectic substances with a low degree of esterification that can bind to the water present in the cells, forming a gel and leading to a lack of juiciness (Zhou, Dong, Ben-Arie, & Lurie, 2001). These partially degraded substances can also trespass into intercellular space and mesocarp cells and capture the water that is present (Bron, Jacomino, & Appezzato-da-Glória, 2002).

To prolong the cold storage time and maintain fruit quality, it is necessary to use additional technologies such as pre-storage and a modified atmosphere. Pre-storage consists of exposing the harvested fruits to temperatures between 20°C and 30°C for 1 or 2 days before cold storage (Infante, Meneses, & Crisosto, 2009). Crisosto, Garner, Andris, and Day (2004) observed that the pre-storage of ‘Summer Lady’ and ‘Ryan Sun’ peaches for 48 hours at 20°C reduced the woolliness and increased the storage period. The pre-storage treatment also enhanced PG activity in peaches (Von Mollendorff & De Villiers, 1988) and nectarines (Zhou et al., 2000), which reduced the woolliness in the fruits when compared to those submitted to cold storage immediately after harvest. Peaches and nectarines that were submitted to pre-storage for 1 or 2 days presented a reduction in woolliness, which improved the sensory quality of these fruits and increased the
consumer acceptance (Sasaki, Cerqueira, Sestari, Del Aguila, & Kluge, 2010; Neves, Campos, Prill, & Roberto, 2013; Infante et al., 2009). Nevertheless, one of the problems reported by these authors is the reduction in firmness and the increase in weight loss during the storage period.

Modified atmosphere is a widely used technique for limiting water loss, delaying ripening and suppressing postharvest diseases in fruits, and it contributes to prolonging shelf life during cold storage (Malakou & Nanos, 2005). It reduces the levels of O₂, while CO₂ levels increase inside the packaging due to the combination of the product’s mass and the respiration rate, the gas transmission rate through the packaging and the permeation area (Kartal, Aday, & Caner, 2012). However, changes in the levels of O₂ and CO₂ should not exceed the critical limit to prevent the occurrence of physiological disorders such as fermentation (Beaudry, 1999). Studies indicate that the use of passive modified atmosphere in peaches under cold storage reduces the loss of weight and firmness and may decrease chilling injuries in some cultivars (Santos, Castro, Picoli, & Rolim, 2008; Santana, Benedetti, Sigrist, & Sarantopóulos, 2010). After evaluating the use of polypropylene and polyethylene, Akbudak and Eris (2004) reported that these packages were efficient in maintaining the quality of ‘Flavorcrest’ peaches and ‘Fantasia’ nectarines for 30 and 45 days of storage, respectively.

Although there are several studies regarding chilling injuries, they are still the major problem associated with the conservation of stone fruits around the world (Fruk, Cmelik, Jemric, Hribar, & Vidrih, 2014; Brummell, Dal Cin, Lurie, Crisosto, & Labavitch, 2004). Through the use of pre-storage and controlled atmosphere, Murray, Lucangeli, Polenta, and Budde (2007) observed that combined techniques for the conservation of peaches and nectarines under refrigeration are more significant in the reduction of chilling injuries than the same treatments when separately applied. The same was observed by Jin, Zheng, Tang, Rui, and Wang (2009) with the use of pre-storage and methyl jasmonate, Malakou and Nanos (2005) with pre-storage and modified atmosphere, and Cao, Hu, Zheng, and Lu (2010) with pre-storage and salicylic acid. ‘Douradão’ peaches are widely cultivated in Brazil with great consumer acceptance. As different varieties of peach present diverse responses to postharvest treatments (Ju, Duan, Ju, & Guo, 2001), the study of new postharvest technologies that prolong the storage of ‘Douradão’ peaches is extremely relevant. This work aimed to evaluate the effects of the combination of pre-storage and passive modified atmosphere in reducing the wooliness and prolonging the shelf life of ‘Douradão’ peaches under cold storage.

**Material and methods**

‘Douradão’ peaches were harvested in the region of Itupeva, São Paulo State, at physiological maturity (breaking of the green background color) and were transported to the Laboratory of Postharvest Physiology and Biochemistry at the Biological Sciences Department of the “Luiz de Queiroz” College of Agriculture in Piracicaba, São Paulo State, Brazil. The fruits were sorted by size and selected according to the absence of mechanical injuries or visible signs of pathogenic agents.

The following treatments were applied: control (no packaging or pre-storage used before cold storage); pre-storage for 24h at 25°C + LDPE 60 µm (CO₂ transmission of 9,577 mL m⁻² day⁻¹ and O₂ transmission of 2,872 mL m⁻² day⁻¹ with a moisture transmission that was 5.9 g water m⁻² day⁻¹); pre-storage for 48h at 25°C + LDPE 60 µm; pre-storage for 24h at 25°C + LDPE 80 µm (CO₂ transmission of 7,425 mL m⁻² day⁻¹, O₂ transmission of 1,705 mL m⁻² day⁻¹, and the moisture transmission was 4.1 g water m⁻² day⁻¹); and pre-storage for 48h at 25°C + LDPE 80 µm. The fruits submitted to pre-storage were placed in hermetic boxes with a 186 L capacity and stored at 25°C. The control fruits were placed in open plastic boxes. After pre-storage, the fruits were placed in LDPE packages to promote passive modified atmosphere during cold storage. All the treatments were stored at 0°C and 90 ± 5% RH for 30 days. After this period, the packages were opened, and the fruits were kept at 25°C and 60 ± 5% RH for 5 days to simulate marketing conditions.

The following variables were determined:

**Package Internal Gas Composition:** A gas analyzer (Illinois Instruments, model 6600 Headspace Oxygen/Carbon dioxide analyzer) was used to determine the gas composition within the package during cold storage. Samples of the inner atmosphere were collected using silicon septa previously fixed to the packages. The results were expressed in percentages of O₂ and CO₂.

**Wooliness Index:** Wooliness was determined according to Nava and Brackmann (2002), with adaptations. The evaluation was subjective, considering the appearance and quality of the manually extracted juice. The fruits were divided into categories: 1 = fruits with high juice content (without wooliness), 2 = fruits with moderate juice content (moderate wooliness) and 3 = fruits without juice (totally wooly).
Internal browning index: This index was evaluated according to an adapted methodology of Ben-Arie and Sonego (1980). The index of internal browning was calculated through a visual scale as follows: fruits without browning (0% of the fruit surface), slightly brown flesh (< 25% of the fruit surface), moderately brown flesh (25-50%), and severely brown flesh (> 50% of the fruit). The index of internal browning (IB) was calculated as the following quotient: IB (0 × % of fruits without browning) + (1 × % of slightly brown fruits) + (2 × % of moderately brown fruits) + (3 × % severely brown fruits)/4. The higher the value that was obtained, the more intense the internal browning.

PME activity (EC 3.1.1.11): The activity was determined by spectrophotometry and according to the adapted methodology of Ratner, Goren, and Monselise (1969). For the extraction, 5 g of frozen fruit flesh was triturated with 20 mL of NaCl (0.2 N) at 4°C (enzymatic extract). The activity was quantified using 30 mL of a pectin solution (1% - diluted in NaCl 0.2 N) with the pH adjusted to 7.0 with NaOH (0.1 N). A total of 5 mL of enzymatic extract was added, and the pH was again adjusted to 7.0 with NaOH (0.1 N). Then, the sample was titrated with NaOH (0.01 N) for 10 minutes to maintain the pH at 7.0. The PME activity was considered the content of the enzyme able to catalyze pectin demethylation, which corresponds to 1 nmol of NaOH per gram of fresh weight per minute (nmol g⁻¹ min⁻¹).

PG activity (EC 3.2.1.15): The enzymatic extract was obtained according to Denès, Baron, and Drilleau (2000) with small modifications. A total of 0.5 g of lyophilized flesh was weighed and homogenized with 15 mL of cold acetone using a high-speed blender (Ultraturrax, UT) for 1 minute. Subsequently, the homogenate was kept in an expanded polystyrene box containing ice for 30 minutes. Then, the supernatant was discarded. The pellet was resuspended in 10 mL of cold acetone and kept in an expanded polystyrene box containing ice for 15 minutes. Next, the supernatant was discarded, and the pellet was used for enzyme determination. The pellet was resuspended in 10 mL of 0.2 M Tris (hydroxymethyl)-aminomethane buffer (Tris buffer), pH 7, containing 500 mg L⁻¹ of sodium metabisulphite, 1% polyvinylpolypyrrolidone (PVPP) and 1 M of NaCl. The extraction was conducted for 2 hours using an orbital shaker at 200 × g in the dark and inside an expanded polystyrene box containing ice. The extract was transferred to Eppendorf tubes and centrifuged at 20,000 × g for 15 minutes at 4°C. The supernatant was used as an extract for the enzymatic assay. The PG activity was determined according to the method proposed by Gross (1982), which measures the reduction of the formed groups using polygalacturonic acid as the substrate. The substrate contained 0.6 mL of a solution prepared with 0.4% (w/v) polygalacturonic acid dissolved in 0.05 M of sodium acetate buffer diluted in NaCl 0.2 N) with the pH adjusted to 4.5. The reaction occurred by the addition of 0.15 mL of enzymatic extract followed by incubation at 37°C for 10 minutes under agitation at 30 × g. The reaction was stopped by adding 2 mL of 10 mM borate buffer at pH 9 and 0.4 mL of 1% (w/v) cyanoacetamide. The mixture was placed in a water bath at 100°C for 10 minutes followed by a cooling period at room temperature. A total of 200 µL of the sample was pipetted to an ultraviolet-flat-bottom microplate and read in a multiple microplate reader at 22°C. The absorbance was read at 276 nm. The calibration curve was conducted with D-galacturonic acid, and the PG activity was expressed as mmol of galacturonic acid per gram of dry weight per minute (mmol g⁻¹ min⁻¹).

Flesh firmness: This parameter was determined using a manual penetrometer with an 8-mm diameter flat tip. After removing the peel, two measures were taken from each fruit from opposite sides of the equatorial section. The results were expressed in Newtons (N).

Total phenolic compounds: Samples of 0.2 g of lyophilized flesh were weighed and homogenized with 6 mL of methanol using a high-speed blender (Ultraturrax, UT) for 1 minute. The extraction was conducted for 1 hour in an orbital shaker at 200 × g in the dark and inside an expanded polystyrene box containing ice. Then, 2 mL of extract was transferred to Eppendorf tubes and centrifuged at 15,000 × g for 10 minutes at 4°C. The content of the phenolic compounds solubilized in the supernatant was determined according to Swain and Hillis (1959) with modifications. A total of 19.2 µL of extract was placed in a flat-bottom microplate, and subsequently, 29 µL of Folin-Ciocalteu reagent (1N) was added. After 3 minutes of incubation in the dark and at room temperature, 192 µL of a solution containing Na₂CO₃ (0.4%) and NaOH (2%) was incorporated and then the samples were kept in the dark for 1 hour at room temperature. The absorbance was measured at 750 nm in a multiple microplate reader (Tecan Infinite M200, Männedorf, Switzerland) at 22°C. The results are expressed in ng of gallic acid in 100 g of dry weight.

Antioxidant capacity (FRAP): Samples of 0.2 g of lyophilized flesh were weighed and homogenized with 6 mL of methanol using a high-speed blender (Ultraturrax, UT) for 1 minute. The extraction was conducted for 1 hour in an orbital shaker at 200 × g in the dark and inside an expanded polystyrene box.
containing ice. After that, 2 mL of extract was transferred to Eppendorf tubes and centrifuged at 15,000 \( \times \) g for 10 minutes at 4°C. The total antioxidant capacity was determined based on the methodology of Benzie and Strain (1996). A buffer solution composed of 500 mmol L\(^{-1}\) of acetate, pH 5.6 (3.1 g C\(_2\)H\(_3\)NaO\(_2\)•3H\(_2\)O) and 16 mL of C\(_6\)H\(_6\)O\(_2\) per liter of buffer solution was prepared. Two other solutions were made: one using 10 mmol L\(^{-1}\) of 2, 4, 6-tripyridyl-s-triazina (TPTZ) in 40 mmol L\(^{-1}\) of HCl and the other using 20 mmol L\(^{-1}\) of FeCl\(_3\)•6H\(_2\)O. The FRAP reagent was prepared by the addition of 10 mL of acetate buffer, 1 mL of TPTZ and 1 mL FeCl\(_3\)•6H\(_2\)O followed by incubation for 2 hours at 37°C. A 6 \( \mu \)L aliquot of the extract was placed in a flat-bottom microplate, and subsequently, a total of 198 \( \mu \)L of FRAP solution was added. The microplate was then kept in the dark for 50 minutes at room temperature, and the absorbance was measured at 593 nm in a multiple microplate reader (Tecan Infinite M200, Männedorf, Switzerland). The results were expressed in mg of trolox in 100 g of dry weight.

For each treatment, all these analyses were conducted on day 0, after 30 days of cold storage and after 30 days of cold storage plus 3 days at room temperature. Each treatment was comprised of 4 repetitions of eight fruits except for the gas composition analysis, which occurred on days 0, 5, 10, 15, 20, 25, and 30 of cold storage. In this case, each treatment comprised 5 repetitions of 4 fruits.

To determine the effect of each factor on each individual parameter, a one-way analysis of variance (ANOVA, \( p \leq 0.05 \)) was conducted (statistics software SAS, version 9.2, SAS Institute, Cary, NC, USA). The mean values were compared by Tukey’s test when significant differences among treatments and interactions between factors were found.

**Results**

**Gas composition**

\( \text{CO}_2 \) production increased, while \( \text{O}_2 \) was reduced for all the pre-storage treatments in association with LDPE packaging when compared to the control. The LCPE with 60 \( \mu \)m packaging showed contents of \( \text{CO}_2 \) and \( \text{O}_2 \) approximately 6% and 5%, respectively. The LDPE with 80 \( \mu \)m packaging showed contents approximately 16% for \( \text{CO}_2 \) and 1% for \( \text{O}_2 \) (Figure 1A and B).

**Figure 1.** The \( \text{O}_2 \) (A) and \( \text{CO}_2 \) (B) composition inside the modified atmosphere packages (LDPE 60 or 80 \( \mu \)m) of ‘Douradão’ peaches after pre-storage (24 or 48h at 25°C) and during storage for 30 days at 0°C and 90 ± 5% RH. The bars represent a standard deviation of the mean \((n = 5)\).

**Woolliness index, Internal browning index, PME, and PG activity**

No significant difference was found between the treatments for the index of woolliness when the fruits were removed from the cold storage. However, all the combined treatments of pre-storage with modified atmosphere using LDPE 60 and 80 \( \mu \)m showed a lower index of woolliness, 2 and 1.20, respectively, when compared to the control (2.8) in the simulated marketing period (Figure 2A). The MAP using LDPE of 80 \( \mu \)m provided the lowest index of woolliness.
Figure 2. The index of woolliness (A), Internal browning index (B), PME activity (C) and PG activity (D) in 'Douradão' peaches submitted to pre-storage (24 or 48h at 25°C), packaged under modified atmosphere (LDPE 60 or 80 µm) and stored for 30 days at 0°C and ± 5% RH followed by a simulated marketing period of 3 days at 25°C and 60 ± 5% RH. Means followed by the same letter do not differ between treatments by the Tukey’s test (p ≥ 0.05). The index of woolliness: 1 = fruits with a high content of juice (without woolliness); 2 = fruits with a moderate content of juice (moderate woolliness) and 3 = fruits without juice (totally woolly).

The internal browning did not affect the fruits when they were removed from the cold storage. However, in the simulated marketing period, the control treatment presented the highest index of internal browning (Figure 2B). PME showed low activity in the control treatment when the fruits were removed from cold storage. Nonetheless, in the simulated marketing period, the control treatment presented the highest activity of this enzyme when compared to the other treatments except for the pre-storage for 24h + LDPE 80 µm (Figure 2C). PG presented lower activity for the control treatment when the fruits were removed from the cold storage and during the simulated marketing period when compared to all the pre-storage treatments (Figure 2D).

**Flesh firmness, total phenolic compounds, and antioxidant capacity**

The firmness decreased with the time of storage in all treatments. In addition, in the simulated marketing period, lower values of firmness were found in all the treatments with combined pre-storage and modified atmosphere when compared to the control (Figure 3A).

The treatments of pre-storage for 48h in combination with LDPE packaging of 60 or 80 µm presented the highest contents of total phenolic compounds on the day the fruits were removed from cold storage. After the simulated marketing period, the contents of the total phenolic compounds of all the treatments submitted to pre-storage and LDPE packaging were higher than the control treatment. In addition, the treatments submitted to pre-storage of 48h presented higher contents of total phenolic compounds in relation to the treatments with only 24h, regardless of the thickness of the film used in the package (Figure 3B).

The antioxidant capacity measured by FRAP was higher for the treatment of pre-storage for 48h + LDPE 80 µm when the fruits were removed from cold storage. In the simulated marketing period, all the treatments of pre-storage in association with LDPE packaging showed a higher antioxidant capacity compared to the control treatment (Figure 3C).
Figure 3. Firmness (A), total phenolics compounds (B) and antioxidant capacity (FRAP) (C) in ‘Douradão’ peaches submitted to pre-storage (24 or 48h at 25°C), packaged under modified atmosphere (LDPE 60 or 80 µm) and stored for 30 days at 0°C and ± 5% RH, followed by a simulated marketing period of 3 days at 25°C and 60 ± 5% RH. The means followed by the same letter do not differ between treatments by the Tukey’s test (p ≥ 0.05).

Discussion

Gas composition

Reductions in the concentrations of CO₂ and O₂ were observed for all the treatments of pre-storage and packaging. The pre-storage for 24 and 48h associated with LDPE packaging of 80 µm presented the largest differences in the concentrations of CO₂ and O₂ when compared to the control. As expected, a thicker LDPE packaging presented a lower permeability to gasses: 80 µm versus 60 µm. According to Kartal, Aday, and Caner (2012), the use of passive MAP promotes a reduction in O₂ levels and an increase in CO₂ levels inside the packaging. This occurs due to the mass and respiration rate of the fruits, the surface area and the permeability rate of the packaging material. In this study, the use of MAP did not cause fermentation of the fruits. The levels of O₂ and CO₂ achieved inside the package were under the limits of tolerance to prevent anaerobic fermentation in peaches, which is less than 2% for O₂ and up to 10 to 25% for CO₂ depending on each peach cultivar (Zagory & Kader, 1988; Fernandez-Trujillo, Salmeron, & Artés, 1997). Retamales, Cooper, Streif, and Kania (1992) achieved great results in maintaining the quality of refrigerated nectarine using 20% CO₂ without the presence of any deleterious effects in the fruit. Zoffoli, Rodriguez, Aldunce, and Crisosto (1997), after studying different packages for cold stored peaches, observed that the levels of CO₂ varied from 10 to 25% and the levels of O₂ varied from 1.5 to 10%. They also noted that greater reductions in chilling injuries occurred in packages with high levels of CO₂. Lee (2014) also observed the quality maintenance and reduction of chilling injuries in ‘Mibaekdo’ peaches submitted to a pretreatment with 30% CO₂.
Woolliness index, Internal browning index, PME, and PG activity

The effect of pre-storage combined with LDPE packaging on the cold storage of ‘Douradão’ peaches was shown to be effective in reducing woolliness when the fruits were cold stored for 30 days at 0°C followed by 3 days at room temperature. The pre-storage also reduced the woolliness in different varieties of peaches and nectarines when the fruits were exposed to room temperature before the cold storage (Zhou et al., 2000; Sasaki et al., 2010; Neves, Campos, Prill, & Roberto, 2013).

Choi and Lee (1997) observed a significant effect on the reduction of woolliness using a passive modified atmosphere in ‘Yumyeong’ peaches when the fruits were compared with unpackaged ones. Santana, Benedetti, Sigrist, and Sarantopóulos (2010) and Santana, Benedetti, Sigrist, and Sato (2011) observed a significant reduction in woolliness with the use of active modified atmosphere and LDPE packaging (with values of 1% O₂ and 3% CO₂).

The internal browning presented the highest values for the control treatment in the simulated marketing period. This result is probably due to the deterioration and senescence of tissues after a long period under refrigeration, which leads to changes in cell membrane permeability. These changes in permeability modify both the cellular metabolism and the normal activity of membrane enzymes, especially the oxidative enzymes. The modification of flesh color is due to the damages caused in the cells by toxic intermediary products that accumulate during cold storage and to the oxidation of phenolic compounds by the polyphenoloxidase enzyme (Macheix, Fleuriet, & Billot 1990; Espin, Morales, Váron, Tudela, & Garcia, 1997).

The exposition of stone fruits to pre-storage before cold storage promotes the stability of cellular tissues and reduces the activity of oxidative enzymes such as polyphenoloxidase (Neves, Campos, Prill, & Roberto, 2013). This enables the peach to resist a longer period of exposition to low storage temperatures without presenting chilling injuries such as internal browning (Crisosto, Garner, Andris, & Day 2004). The results found in this study corroborate the findings of Jin, Zheng, Tang, Rui, and Wang (2009), who observed a reduction in internal browning in ‘Baifeng’ peaches after heat treatment at 38°C for 12 hours. Neves, Tosin, Silva, Vasconcelos, and Roberto (2012) reported the efficiency of pre-storage for 24 or 48 hours in preventing internal browning in three peach cultivars that had been stored for 28 days under cold storage. Retamales, Cooper, Streif, and Kania (1992) found that the application of pre-storage for 48 hours associated with a modified atmosphere with a high CO₂ concentration (20%) significantly reduced internal browning.

The increase in PME activity and the decrease in PG activity observed in this study are related to the high index of woolliness in the control fruits in the simulated marketing period. The index of woolliness in stone fruits is considered to occur due to an imbalance between the activities of the pectinolytic enzymes PME and PG during cold storage (Ben-Arie & Sonego, 1980). The low activity of PG combined with the de-esterification of pectin molecules from the action of PME results in the accumulation of pectins with high molecular weight and low degrees of esterification that bind calcium ions, which is what generates the pectates. These compounds remove the free water from the cells and form a pectic gel, thus promoting the loss of juiciness in fruit and a farinaceous aspect (Obenland, Crisosto, & Rose, 2003). Brummett et al. (2004) observed that PG synthesis is regulated by ethylene. The low ethylene production during cooling reduces the synthesis and activity of the enzyme and causes chilling injuries such as woolliness in stone fruits. Other authors found similar results to this study and reported an increase in PME activity at the same time as the increase in woolliness (Artés, Cano, & Fernández-Trujillo, 1996). Choi and Lee (1997) and Santana et al. (2011) also noticed low PG activity and a high index of woolliness in peaches without packaging when compared to packaged fruits stored for 4 weeks at 0°C. Zhou et al. (2000) and Brummett et al. (2004) also found a lower PG activity in woolly peaches and nectarines after cold storage.

Flesh firmness, total phenolic compounds, and antioxidant capacity.

The decrease in firmness found during the simulated marketing period in treatments that combined pre-storage and LDPE were expected since the exposure to higher temperatures before the cold storage accelerates the metabolism of stone fruits and affects the firmness and all the other aspects related to ripening (Vitti, Kluge, Jacomino, & Lima, 2007). Camargo, Lima, Scalon, and Siqueira (2000) reported that the loss of flesh firmness during ripening is due to the action of enzymes from the wall such as PG. The reduction in the firmness of the fruits submitted to the pre-storage and MAP may be related to the higher
PG activity (Figure 2D). According to Evangelista, Chitarra, and Chitarra (2000), PG catalyzes the hydrolysis of the β-1,4 bonds between the galacturonic acid residues in the pectin chain and consequently promotes the reduction of firmness in the fruits. The lower firmness found in the treated fruit during the simulated marketing period may also be related to the higher juiciness and lack of woolliness in these fruits (Figure 2A). Other authors observed similar results in peaches and nectarines submitted to pre-storage and concluded that the decline in firmness is related to an increase in juiciness and to a decrease in woolliness (Infante et al., 2009; Sasaki et al., 2010). Murray et al. (2007) reported that the combination of pre-storage and controlled atmosphere reduced the fruit firmness to values of approximately 6 N, as well as the woolliness, thus maintaining the fruit quality. Cano-Salazar, López, Crisosto, and Echeverría (2013) also reported a reduction in firmness in 'Early Rich' peaches to 4.5 N after exposure to room temperature (20°C) for 36 hours, whereas the untreated fruits presented values of 14.9 N for firmness.

The control treatment presented a lower content of phenolic compounds. Genetic and environmental factors are determining factors in the content of phenolic compounds (Robards, Prenzler, Tucker, Swatsitang, & Glover, 1999). However, oxidative reactions that occur during processing or storage can also modify these compounds because the increase in their content is related to responses to stress conditions. In general, oxidative regulation is a process that promotes dynamic balance in the system and produces and scavenges reactive oxygen species (ROS). According to Toivonen (2004), the tissue initially responds to the stress, leading to the production of ROS and consequently activating the system responsible for the production of antioxidant compounds. Nevertheless, if there is a high level of stress, the tissue may not be able to maintain the necessary balance and causes damage to the cells. Phenolic compounds are one of the main mechanisms for the protection of cellular tissue (Mittler 2002) and act directly as antioxidant agents or substrates for superoxide (Robards et al., 1999). The internal browning is generally related to the increase in the activities of polyphenoloxidase and peroxidase that can lead to the oxidation of phenolic compounds to quinones or to quinine compounds, resulting in brown polymerized pigments (Lil, O'Donagheue, & King, 1989). Jin et al. (2009) also related the high index of internal browning with the low content of phenolic compounds in peaches when they were exposed to room temperature after refrigeration.

According to Malakou and Nanos (2005), the exposure of fruits to high temperatures before cold storage leads to an increase in the content of phenolic compounds and is a way to protect the vegetal tissue from the stress caused by the temperature. The same authors found an increase in the content of phenolic compounds in 'Royal Glory' peaches during the simulated marketing period with the application of heat treatment (immersion in hot water) before cold storage. However, Neves et al. (2013) did not find any difference between the treatments both with and without the application of pre-storage in peaches kept under refrigeration for 28 days.

The use of pre-storage associated with LDPE packaging promoted a higher antioxidant capacity in 'Douradão' peaches during the simulated marketing period. This finding can be related to the lower internal browning index and the higher content of total phenolic compounds observed in these treatments over the same period. Studies report that the antioxidant capacity of phenolic compounds is due to the reducing power of the aromatic hydroxyl groups that reduce reactive free radicals, such as “singlet” oxygen, or decomposed peroxides, which produce the radical phenoxy that is less reactive (Campos, Martino, Sabarense, & Pinheiro-Sant’Ana, 2008; Silva, Costa, Santana, & Koblitz, 2010). These results are in accordance with those found by Gil, Tomas-Barberan, Hess-Pierce, and Kader (2002) and Mokrani et al. (2016), who observed a high correlation between the antioxidant capacity and the content of total phenolic compounds. According to these authors, phenolic compounds are mainly responsible for the antioxidant capacity in peaches.

**Conclusion**

The combination of pre-storage for 24 and 48 hours and LDPE packaging of 60 and 80 µm is a strategy to prevent or reduce chilling injuries in 'Douradão' peaches while maintaining the quality of fruits during cold storage for 30 days. The LDPE packaging of 80 µm associated with pre-storage for both 24 and 48 hours showed the best results in the reduction in woolliness. All the treatments that used a combination of pre-storage and LDPE were efficient in inhibiting internal browning, highest content of phenolic compounds and antioxidant capacity during the simulated marketing period the fruit.
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