Combined Treatment of Carbon Monoxide and Chitosan Reduced Peach Fruit Browning and Softening During Cold Storage

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Abstract: The effects of postharvest application of chitosan and carbon monoxide (CO) on fruit browning and softening during cold storage (8 °C) were evaluated. CO (10 µmol/L) significantly delayed the internal browning of peach fruit, and the effect was enhanced in combination with chitosan (1%, w/w). Further studies showed that treatment with CO and chitosan reduced the increase of PPO and POD activities, maintained PAL activity and total phenolics content at a higher level. Moreover, it also reduced fruit tissue softening by retarded the increase of PE, PG activities and water soluble pectin content, inhibited the decline of flesh firmness as well as sustained the balance of PG and PE activities, improved the ability of chilling injury tolerance. Therefore, peaches treated with chitosan and CO obviously delayed the fruit browning and softening during cold storage, and it indicates that combined treatment with chitosan and CO can be effective in reducing browning and softening of peach fruit and inhibited chilling injury during cold storage.

Keywords: Peach, CO, Chitosan, Browning, Softening, Cold Storage

1. Introduction

Peach is one of the most popular fruit in the world because of its high nutrient level and pleasant flavor. Peaches ripen in summer, and its respiratory metabolism is strong after harvest, which lead to soften, browning and decay quickly at ambient temperature [1, 2]. Cold storage remains the main method to slow the product deterioration in terms of consumer perception and nutritional value. However, low temperature results in chilling injury symptoms, manifested as mealy (soft and dry, with lack of juice) or leathered flesh (hard textured, with no juice), with/without browning and bleeding in the flesh) in some peach cultivars during or after cold storage [3] and these influence its commercial value and economic benefit. Chemical treatments have been used to delay fruit senescence and prolong postharvest shelf life. However, the use of chemicals has been minimized for food safety and environmental reasons. Many physical methods including modified atmosphere packaging, heat and UV-C pretreatments, are being extensively studied as substitutes for the current chemical methods in commercial applications [4-6].

Chitosan and its derivatives like oligochitosan, have been reported to control postharvest diseases effectively. Chitosan is safe, nontoxic, biocompatible, and biodegradable natural alkaline polysaccharide derived from the deacetylation of chitin [7]. Chitosan has been successfully used in many post harvested fruits and vegetables, such as grape, strawberries, berry, jujube and fresh cut lotus root through single coating or comprehensive treatments [8, 9]. Chitosan can form a film on fruit and vegetable surfaces and reduces respiration rate by adjusting the permeability of carbon dioxide and oxygen. The NH$_2$-group of chitosan may also restrain the propagation of harmful germs, thus effectively controlling fruit decay [10].

Carbon monoxide is a stable diatomic gas molecule. As another gas signal transducer molecule of plant apart from NO, it involved in various physiological and metabolic regulation [11]. Recent research suggested that CO could enhance the anti-senesence ability of plant leaf, improve the SOD, POD and CAT activities of plant tissue, and reduce the MDA content of fresh cut Chinese rose flower [12, 13]. In addition, CO fumigation could prevent the browning of fresh cut lotus root [14]. At present, the researches of the mechanism and the
physiological effects of CO on postharvest fruits and vegetables are still not very clear.

In the present work, we treated peaches with CO gas, chitosan and CO plus chitosan, and determined the relevant parameters of flesh browning and softening during cold storage. Our objectives were to disclose the possible mechanism of CO alone and plus chitosan treatments on fruit browning and softening of postharvest peach during cold storage.

2. Materials and Methods

2.1. Materials

Peach fruit (Prunus persica cv. Hongburuan) were harvested at commercial maturity from local orchard situated in Yaodu district, Linfen city, Shanxi province, China and immediately transported to our laboratory within 2 h after harvest. Peaches with uniform shape, size, and color, as well as no insect pest and mechanical damage, were selected as materials.

Chitosan (water-soluble, molecular weight of about 200 kDa and 85% deacetyl degree) was purchased from AK Biotech Co., Ltd. (Shandong, China). CO gas with a purity of 99.99% was purchased from Beijing Huaneng Special Gases Co., Ltd. (Beijing, China). Carbazole and pyrocatechol (analytical grade) were purchased from Sinopharm Chemical Reagent Co., Ltd. (Shanghai, China). Pectin, sodium carboxymethyl cellulose, and galacturonic acid were purchased from Sigma (USA). Other reagents of analytical grade were purchased from Alfa Aesar Company (Tianjin, China).

2.2. Fruit Treatment

The peaches were randomly divided into 4 groups, each with 60Kg fruits. Based on our previous experiments, the first group was immersed in 1% chitosan solution for 3 min, and then dried under natural wind; the second group was fumigated with 10 µmol/L CO gas for 2 hours; the third group was firstly fumigated with 10 µmol/L CO gas for 2 hours, and then dipped into 1% chitosan solution for 3 min, afterward the peaches were dried under natural wind; the fourth group was untreated jujubes served as control samples. After treated, all samples were stored at 8 ℃ and the relevant parameters for the analyses of peaches were measured periodically.

2.3. Firmness and Water Pectin

Firmness was determined by a penetrometer (mode GY-4, Mudanjiang, China) with a 8-mm-long probe with a diameter of 10 mm. 6 fruits were used and two points per fruit were selected to determine the firmness of each replicate sample.

Pulp (5 g) was homogenized in 40 ml of 85 ℃ ethanol (95%), and the homogenate was incubated for 10 min at 85 ℃, then centrifuged (10 000 ×g). The residue was re-extracted with 25 ml of 63% ethanol at 85 ℃ for a further 10 min. The residue was again centrifuged, the water-soluble pectin extracted from the ethanol washed residue. The extract is used to determined water-soluble pectin [15].

2.4. IB Index

The degree of IB was visually assessed on the mesocarp surface following a double cut parallel to the axial diameter [16]. The extent of flesh browning was divided into five classes: 0, no browning; 1, browning covering <25% of the cut surface; 2, browning covering ≥25% but <50% of cut surface; 3, browning covering ≥50% but <75% of cut surface; 4, browning covering ≥75% of cut surface. The IB index was calculated using the following formula:

\[
IB\ index = \sum[(IB\ level) \times (number\ of\ fruit\ at\ the\ IB\ level)]/(4 \times \text{total number of fruit in the treatment})
\]

2.5. PG and PE Activities

PG and PE activities were determined according to the modified method of Zhang et al. (2015) [17]. One unit of PG activity was expressed as nmol of galacturonic acid per min per mg protein, and the PE enzyme activity was expressed as U·mg⁻¹ protein, one unit activity was calculated as 1 mmol L⁻¹ NaOH consumed per mg protein.

Protein content in the enzyme extracts was estimated using the Bradford (1976) method, with bovine serum albumin as a standard. Specific activity of the enzymes was expressed as units per milligram protein.

2.6. Total Phenolic Content

The total phenolic content was measured according to the FolinicCiocalteu procedure [18], with little modifications. 5 g mortar of flesh tissues were extracted three times with 50 mL of methanol, and centrifuged at 10,000 g for 10 min at 4 ℃ with centrifuge (Eppendorf 5417R, Hamburg, Germany). The supernatants (1 mL) were mixed with 1.5 mL of the FolinicCiocalteu reagent. After mixing thoroughly for 6 min, 6 mL of 1.0 mol/L Na₂CO₃ solution were added into the mixture. After 90 min at 25 ℃, the absorbance was measured at 765 nm in a spectrophotometer. The total phenolic content was determined by comparison with the absorbance of gallic acid, which was used at different concentrations as the standard.

2.7. PAL, PPO and POD Activities

PAL activity was analyzed by using a modified method of Zhang et al. (2013) [14], 2 g flesh of peaches was homogenized in a cold 10 mL solution of 0.05 mol/L sodium borate buffer (pH 8.0) containing 0.5 g polyvinylpyrrolidone, 5 mmol/L 2-mercaptoethanol and 2 mmol/L EDTA at 4 ℃. The homogenate was centrifuged at 12,000 g for 20 min at 4 ℃. The supernatant was used as the crude enzyme extract and stored at 4 ℃ for the determination. 0.5 mL of enzyme extract was incubated with assay medium containing 3.5 mL of 100 mmol/L sodium borate buffers (pH 8.7) and 1 mL of 10 mmol/L L-phenylalanine as substrate at 37 ℃ for 1 h. The reaction was terminated by adding 0.2 mL of 6 mol/L HCl. PAL activity was measured by change in absorbance at 290 nm. One unit of enzyme activity was defined as the change in absorbance of 0.01 per hour.

POD and PPO activities were measured in pulp tissues
following the method as described by Zheng et al. (2007) [19]. POD activity was based on the determination of guaiacol oxidation at 470 nm by H2O2. The change in absorbance at 470 nm was followed every 30 s by a spectrophotometer (UNIC UV-2100, UNIC (Shanghai) Equipment Co. Ltd., China). One unit of POD defined as the amount of enzyme caused 0.01 absorbance increase per min under the conditions of assay. PPO activity was measured by incubating 0.5 ml of enzyme extract to 2.5 ml of buffered substrate (100 mM sodium phosphate, pH 6.4 and 50 mM Catechol), and then monitoring the change of absorbance at 398 nm. One unit of activity of PPO was defined as the amount of enzyme causing 0.01 absorbance increase per minute under the conditions of assay. The content of protein in crude enzyme extraction was assayed as mentioned above.

2.8. Statistical Analysis

Each experiment was repeated three times and the data was processed by analysis of variance (ANOVA) using DPS7.05 statistical software (Refine Information Tech. Co., Ltd, Hangzhou, China). The treatments were compared at P = 0.05 using Tukey’s test, which indicates the multi-comparison value in each case. The data were expressed as mean ± standard deviation (SD). Differences at P≤0.05 were considered as significant.

3. Results and Analysis

3.1. IB index and Total Phenolic Content

Fig. 1. Effects of different treatments on IB index (A) and total phenolic content (B) of postharvest peach.

The control fruit showed IB on the 10th day and thereafter the index increased rapidly. CO and chitosan treatment applied separately effectively reduced the development and severity of IB by 50% and 28%, respectively, after 30 days of storage at 8 °C. However, it was found that the combined application of the two treatments was much more effective than the application of either treatment alone, and significantly (P≤0.05) reduced the development of IB by 62% after 30 days (Fig. 1A). During 30 days of storage at 8 °C, total phenolics of control peaches was significantly lower than that of the treatments. Obviously, CO and chitosan treatment alone could maintain total phenolics of flesh at a high level, and treated with CO was more effective than that with chitosan. The content was significantly (P≤0.05) higher in the combined treatment than that in the separate treatments and control peaches after 10th day (Fig. 1B).

3.2. PAL, PPO and POD Activities

PAL activity in the control peach showed a little decrease during the first 10 days, then increased, and declined after 20 days storage. PAL activities in the CO-treated, chitosan-treated and combined treated fruit showed a trend of first increased and then decreased, the peak appeared on 15th, 15th and 20th day, respectively; all treatments maintained PAL activity at a higher level during cold storage(Fig. 2A); and the differences between the treatment with CO, chitosan and CO plus chitosan were significant (P≤0.05).

PPO activities increased in the control and all the treatments fruit during the storage period. Compared with the control, the treatments with CO, chitosan and CO plus chitosan obviously inhibited the increase of PPO activities. After 30 days of storage, the activities in the treatments were 75%, 84% and 63%, respectively, of the control (Fig.2B).

POD activities in the control and chitosan-treated peach fruit increased slightly during the first 15 days; for the treatments with CO or plus chitosan, the activities seemed little decreased after 10 days storage; and it were increased thereafter in all treatments and the control peach fruit. All treatment slowed down the changes of POD activity; however, the treatment with CO plus chitosan was most effective in all treatments, and the activity was only 67% of the control at day 30 (Fig. 2C).
3.3. Firmness and Water Soluble Pectin Content

Fruit firmnesses of all treatments fruit were declined during the whole storage period. But for the control firmness showed a little increase after 20 days storage (Fig. 3A). Firmness of the treatments (with CO, chitosan, and CO plus chitosan) fruit were significantly higher compared to the control before 25, 20 and 30, respectively, days storage (P≤0.05).

Water soluble pectin content in the control peaches increased in the first 20 day, and then decreased (Fig. 3B). Water soluble pectin content in all treatments peaches were increased during storage time and it were lower than in the control. Obviously, the combined treatment was more effective than other.

3.4. PE and PG Activities

Generally, PE activities in the control, as well as in the treatments peaches increased with the extension of storage time (Fig. 4A). However, all treatments could distinctly inhibit the increase of PE activity in the flesh of peach. PE activity in combined treatment peach was significantly lower than other treatments and the control after 5 day, was 37% lower than in the control.

PG activity in the control and treatment with chitosan fruit, as shown in figure 4B, firstly ascended and then declined, and the peaks appeared at day 15 and 20, respectively. The activities in the CO-treated and combined treated peaches gradually increased during storage time. The treatment with CO, chitosan and CO plus chitosan could inhibit the change of PG activity, and the combined treatment was more effective than with either treatment alone.

4. Discussion

4.1. Effect of CO and Chitosan on Flesh Browning of Postharvest Peach

Peaches are easy softening, browning and rot after harvested at room temperature. However, Peaches are very sensitive to low temperature and exhibit chilling injury after
long periods of refrigeration. The main symptoms of chilling injury are leathered flesh or internal browning and flesh mealy that lead to a dry, grainy sand-like texture [3]. Carbon monoxide enhances the chilling tolerance of recalcitrant Baccaurea ramiflora seeds via nitric oxide-mediated glutathione homeostasis [20].

Fig. 4. Effects of different treatments on PE (A) and PG (B) activities of postharvest peach.

Our results showed that the symptom of flesh browning emerged at 10th day after cold storage in the untreated ‘Hongburuan’ peach fruit, and IB index was increased with the extension of storage time. The treatments of 10 µmol/L CO and 1% chitosan alone could inhibit the increase of flesh browning, and IB index in the combined treatment was lower than other treatment and control. Flesh browning was one of the main CI symptoms and oxidation of phenolic substrates by polyphenol oxidase (PPO) was thought to be a major cause of the discoloration in many fruits [3]. In general, phenolic compounds with regards to their antioxidant capacity were accumulated in plant tissue under cold stress [21]. Thus especial attention should be paid to the dual action of phenols of the antioxidant activity and oxidative browning. Peroxidase was one of the main antioxidant enzymes and catalyzed the polymerisation of phenolic compounds as well [22], and was usually estimated as main factor to illustrate the changes of phenol metabolism.

Internal browning of fruit is probably related to the increase in PPO and POD activities, which could oxidize phenolic compounds to quinone or quinine-like compounds, finally appearing as polymerized brown pigments [5]. Qi et al. (2011) [23] reported that chitosan-coating treatment Extended the shelf-life of Fresh-cut ‘Fuji’ apples by controlling its cut-surface browning. Zhang et al. (2013) [14] found that CO retarded the browning of fresh-cut lotus by inhibited PPO and POD activities, at the same time CO also promoted PAL activity and all of these were conducive to maintaining the quality and shelf life. Our results showed that CO alone of combined with chitosan could induced PAL activity, increased the total phenolics, which could enhance the resistance to cold damage and strengthen the tolerance for chilling injury. In addition, CO reduced PPO and POD activity, and inhibited enzymatic browning in flesh of peach.

4.2. Effect of CO and Chitosan on Fruit Softening of Postharvest Peach

Peach fruits softening characterized as firmness decline in ripening process was associated with cell wall modification. Polygalacturonase (PG) is one of the pectin-degrading enzymes and plays a central role in ripening process [24]. Pectin methyl esterase (PME), catalyzing hydrolysisation of methyl ester group of pectin, is a key control enzyme for both assembly and disassembly of pectin network [25] by determining tissue integrity during fruit senescence. Zhou et al (2000a) [26] found an increase in PE activity and inhibition of PG in cold stored peach fruit relative to their activities in normally ripening fruit. The occurrence of flesh mealyness and wooliness in peaches or nectarines has been found to be associated with imbalance of PG and PME activities [27]. Firmness of untreated fruit did not decrease after 15 days cold storage, and a little increased at day 30, but for treated fruit firmness reduced in whole storage period, but the decline was not too big, the firmness decreased by 50% (from 7.12 to 3.19 Kg/cm²), obviously, ‘hongburuan’ peach was a variety more like non-melt peach. The water soluble pectin in the control fruit increased before 20th day then decreased. CO, chitosan or combined treatment retarded the increase of water soluble pectin in the fruit but not inhibited the decline of flesh firmness, and peaches softened as usual. The treatments were effective to resist the chilling injury refer to the condition of flesh browning. The further research result showed that the treatment inhibited the increase of PE and PG activities as well maintain the balance between PE and PG activity which is necessary for fruit softening [28].

5. Conclusion

CO (10 µmol/L) significantly delayed the internal browning of peach fruit, and the effect was enhanced in combination with chitosan (1%, w/w). The treatment with CO and chitosan reduced the increase of PPO and POD activities, maintained PAL activity and total phenolics content at a higher level. And, it also retarded the increase of PE, PG activities and water soluble pectin content, inhibited the decline of flesh firmness as well as sustained the balance of PG and PE activities. The combined treatment with CO and chitosan can be effective in reducing browning and softening of peach fruit and inhibited
chilling injury during cold storage.

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