Ripening-induced Chemical and Antioxidant Changes in Bell Peppers as Affected by Harvest Maturity and Postharvest Ethylene Exposure

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Abstract. Greenhouse-grown bell pepper (Capsicum annuum L. ‘Robusta’) were harvested at five stages of maturation (10% red to full red) in early winter 2002 (Expt. 1) and at two stages (10% red and full red) in early Spring 2002 (Expt. 2). The fruit were subsequently stored at 20 °C in a continuous-flow chamber consisting of either 100 µL·L⁻¹ ethylene (balance air) or air-only (control) at 90% relative humidity (RH). Individual fruit were removed from the chambers upon reaching full red color, and stored at ~30 °C until physiochemical analyses were conducted. Harvest maturity, and ethylene exposure had no appreciable effect on pulp soluble solids content, total titratable acidity or pH. Exposure to ethylene hastened ripening time compared to the air control but was independent of fruit maturity at harvest. Fruit exposed to ethylene reached full-red color 6.4 days (Expt. 1) and 4 days (Expt. 2) earlier than air-only fruit, respectively. There were no significant phytochemical and antioxidant differences noted for total carotenoids, total ascorbic acid, and soluble phenolics at various maturity stages due to ethylene exposure. Appreciable differences were observed between the two experiments for phytochemicals and antioxidant, as bell peppers from the latter experiment contained at least twice the concentrations of phytochemicals and antioxidant capacity as those from the first experiment. Differences in these parameters between experiments were attributed to environmental factors such as average temperature, day length, and light intensity. Ethylene was demonstrated to be an effective postharvest treatment for accelerating color change in this bell pepper cultivar, permitting earlier harvest without altering phytochemical synthesis rates.

Bell pepper (Capsicum annuum L.) is a nonpungent fruit that is valued for color, flavor, and nutritional attributes including ascorbic acid, polyphenolics, and various carotenoids. In 2002, U.S. production totaled about $500 million, with Florida being the predominant supplier of fresh-market, bell peppers during the fall, winter and spring seasons, and Florida being the major summer supplier of fresh-market, bell peppers during the fall, winter and spring seasons, and California being the major summer supplier (NASS, 2004). Practically all green peppers are California being the major summer supplier ing the fall, winter and spring seasons, and Traditionally, commercial growers have tare. Bell pepper is also a good source of provitamin A carotenoids and xanthophylls, and numerous studies have focused on improving retention of these compounds during processing and storage. Peppers are also more susceptible to physical and chemical injury during transport and have reduced postharvest shelf life than those harvested at the green stage. Although nonclimacteric with regard to postharvest respiratory pattern, mature-harvested bell pepper will progress through normal ripening processes to degrade chlorophyll while simultaneously synthesizing a variety of red and yellow carotenoids (Saltveit, 1977). Bell pepper is a good source of provitamin A carotenoids and xanthophylls, and numerous studies have focused on improving retention of these compounds during processing and storage. Peppers are also more susceptible to physical and chemical injury during transport and have reduced postharvest shelf life than those harvested at the green stage. Although nonclimacteric with regard to postharvest respiratory pattern, mature-harvested bell pepper will progress through normal ripening processes to degrade chlorophyll while simultaneously synthesizing a variety of red and yellow carotenoids (Saltveit, 1977). Bell pepper is also a good source of provitamin A carotenoids and xanthophylls, and numerous studies have focused on improving retention of these compounds during processing and storage (Howard and Hernandez-Brenes, 1998; Markus et al., 1999; Miguez-Mosquera and Hornero-Mendez, 1994; Simonne et al., 1997). Red bell pepper derives its brilliant color from capsanthin, capsorubin, and capsanthin 5, 6-epoxide (Miguez-Mosquera and Hornero-Mendez, 1994) with concentrations increasing dramatically as the fruit ripens. Bell peppers also contain high concentrations of ascorbic acid (0.15 to 2.0 mg·g⁻¹ fresh weight) compared to other fruit and vegetables (Salunkhe, 1997; Simonne et al., 1997; Howard et al., 2000; Yahia et al., 2001). Ascorbic acid development in pepper and other fruit is related to its characteristic warm winters, high light intensity. Ethylene was demonstrated to be an effective postharvest treatment for accelerating color change in this bell pepper cultivar, permitting earlier harvest without altering phytochemical synthesis rates.

Pepper growers could potentially reduce field production costs by hastening the fruit ripening rate on the plant or by harvesting the fruit before attaining full color and completing the ripening process during storage without appreciable loss in quality or phytochemical attributes. Spray application of ethephon [2-chloroethyloxy phosphonic acid] prior harvest accelerated field ripening of red pepper types and resulted in higher yields (Cantliffe and Goodwin, 1975). However, ethephon application also induced defoliation, limiting this procedure to once-over harvest for the processing market. This is not compatible for fresh-market bell peppers which require multiple harvests. Ethephon is it not registered for postharvest application to peppers (U.S. EPA, 1988). Harvested bell peppers are considered mildly responsive to postharvest exposure to ethylene (Ryall and Lipton, 1979). In a preliminary study, we harvested greenhouse-grown bell peppers at various maturity stages due to ethylene exposure. Appreciable differences were observed between the two experiments for phytochemicals and antioxidants, as bell peppers from the latter experiment contained at least twice the concentrations of phytochemicals and antioxidant capacity as those from the first experiment. Differences in these parameters between experiments were attributed to environmental factors such as average temperature, day length, and light intensity. Ethylene was demonstrated to be an effective postharvest treatment for accelerating color change in this bell pepper cultivar, permitting earlier harvest without altering phytochemical synthesis rates.

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732

HORTSCIENCE VOL. 40(3) JUNE 2005
process used for commercial degreening of fruit during carotenoid synthesis, similar to the effect of ethylene; when ethylene was removed, the fruit turned unmarketable (unpublished data). Also we noticed that accelerated ripening only occurred when the peppers were continuously exposed to ethylene; when ethylene was removed, the ripening rate slowed. Thus, it appeared that the ethylene hastened chlorophyll breakdown during carotenoid synthesis, similar to the process used for commercial degreening of fresh-market citrus, another nonclimacteric fruit (Grierson et al., 1986).

These promising results led to a subsequent, more comprehensive study of two bell pepper cultivars. Red (‘Triple 4’) and yellow (‘Kelvin’) bell peppers were harvested at 10% to 30% color and stored at 20 °C/90% RH. These fruits also ripened to full color, although exposure to ethylene did not accelerate ripening in either of these varieties (Molinari et al., 2000). Peppers ripened faster after the plant than tagged fruit left to ripen on the plant. Together, these studies suggested that beneficial aspects of ethylene treatment were linked to certain cultivars. Little information is available concerning potential beneficial aspects of ethylene exposure during postharvest ripening of bell pepper, such as color enhancement and subsequent changes to antioxidant phytochemicals.

The objective of this study was to determine the feasibility of using continuous ethylene exposure to accelerate the ripening of bell pepper harvested before full color development. Following development of full-red color, fruit quality was evaluated via selected physiochemical and antioxidant properties. This information could be useful to determine optimal harvest times and postharvest handling practices to develop maximum phytochemical content for bell peppers destined for fresh and processed markets.

Materials and Methods

Production methods, harvest and ripening treatments. The greenhouse variety of ‘Robusta’ red bell pepper (De Ruiter Seeds, Inc.) was selected for this study since it responded to ethylene in our preliminary study. Plants were grown hydroponically in a greenhouse at the University of Florida’s North Florida Research and Education Center–Suwannee Valley (Live Oak, Fla.; 30°18’N, 82°54’W) during the Fall 2001 and Spring 2002 growing season. The plants were grown using lay-flat, perforate bag culture and were trained to the two-stem system (Hochmuth and Hochmuth, 2003). Nutrient solution concentrations were graduated throughout the growing season, with averages during fruit growth period of N = 125 ppm, P = 50 ppm and K = 200 ppm. The greenhouse was heated as necessary to maintain ambient temperatures >10 °C.

Fruit for Expt. 1 were harvested (clipped at stem abscission zone) on 28 Jan. at ripeness stages <10% red, 10% to 30% red, 30% to 60% red and full red. Fruit for Expt. 2 were harvested on 4 Apr. at 10% to 30% red and full red. In both experiments peppers were sorted on the day of harvest for uniform quality. Mean fruit size was 88 mm (diameter), 67 mm (length), and 179 g. Peppers were then placed in air or 100 ppm ethylene (continuous-flow chambers) at 20 °C and 90% relative humidity (RH) and stored until each fruit reached full-red stage. As each fruit reached full-red color, it was removed from the treatment chamber for analysis. Peppers harvested at full-red stage were analyzed the day of harvest. External color (CIE color values L*, a*, b*) was then measured using a chromameter (CR 200 series; Minolta Co., Ltd., Osaka, Japan) on opposite sides of the equator (Expt. 1 only). Illuminant angle was D65 with an 8-mm aperture. Hue angle and chroma values were calculated according to Shewfelt et al. (1988).

Sample preparation and analyses. In both experiments, each fruit was sliced in half axially, and seeds, stem, and calyx were removed. The excised pericarp halves were then packaged in reclosable bags of 0.1 mm thickness plastic, and frozen to –30 °C for up to 6 months until further analyses were conducted. Extracts for soluble solids, titratable acidity and pH were obtained by homogenizing the fruit halves with a tissuemizer (PT-10; Brinkman Instruments, Westbury, N.Y.) followed by centrifugation at 15,000 g, for 20 min at 20 °C. The supernatant was filtered through cheesecloth and soluble solids were determined at 20 °C using a digital Abbe Mark II refractometer (model 10480; Delpew, N.Y.). Titratable acidity was determined by potentiometric titration against 0.1 NaOH to pH 8.2 using an automatic titrator (Fisher Tititrimeter II, Pittsburgh, Pa.) and expressed in malic acid equivalents, the predominant acid in ripe bell pepper (Luning et al., 1994). Pulp pH was determined using a digital pH meter (model 140; Corning).

Total ascorbic acid, the sum of L-ascorbic and dehydroascorbic acid, was quantified by reverse-phase HPLC using modified chromatographic conditions described by Gökmen et al. (2000). Each fruit replicate was homogenized using a tissuemizer (PT-10; Brinkman Instruments, Westbury, N.Y.) for 25 s at maximum speed in 50-mL screw cap centrifuge tubes containing 30 mL of 3% malic acid and the resulting isolate filtered for analysis in triplicate (15 g). Dithiothreitol (8 mM) was added to an aliquot of the filtrate as a precolumn reductant and samples kept in the dark for 120 min to convert dehydroascorbic acid to L-ascorbic acid. Samples were then filtered through a 0.45-μm PTFE filter (Whatman, Clifton, N.J.) and analyzed for total ascorbic acid by HPLC. Separation was performed on a Waters 2695 HPLC system using a Supelcosil 4.6 × 250 mm LC-18 column (Supelco, Bellefonte, Pa.), 0.2 M KH₂PO₄ (pH 2.4) in the mobile phase at a flow rate of 0.5 mL·min⁻¹, and UV detection at 254 nm using a Waters 996 PDA detector.

For remaining chemical analyses, macerated fruit (5 g/30 mL) were homogenized in a solution of ethanol and acetone (1:1 v/v) to solubilize major phytochemicals including carotenoids, polyphenolics, and ascorbic acid. The resulting extract was filtered through Whatman #4 filter paper and held in the dark at –20 °C until analysis. This extract was used for the determination of total carotenoids, soluble phenolics, and antioxidant capacity. Carotenoids were determined by recording absorbance values from the two most prevalent spectral bands (452 and 472 nm) that corresponded to the red carotenoid capsanthin (ε = 2,009) and yellow carotenoids such as β-carotene and lutein (ε = 2,450) in a similar manner described by Hornero-Mendez and Mínguez-Mosquera (2001). Total soluble phenolics, including contributions from ascorbic acid, were measured using the Folin-Ciocalteau assay (Talcott et al., 2000) with data expressed as gallic acid equivalents. Antioxidant capacity was determined using the oxygen radical absorbance capacity assay using fluorescein as the fluorescent probe and evaluated against a standard of Trolox as described by Talcott et al. (2003). Isolates were diluted 10-fold in pH 7.0 phosphate buffer (0.75 M K₂HPO₄ and 0.75 M NaH₂PO₄, 61.6:38.9 v/v) before pipetting triplicate samples into a 96-well microplate with corrections made for background interference due to the phosphate buffer and extraction solvents.

The initial ethanol-acetone extract from each bell pepper isolate was then partitioned into two distinct isolates by isolating lipophilic and hydrophilic compounds by the addition of a known volume of petroleum ether (1.3 v/v). Complete partitioning of xanthophylls into petroleum ether was facilitated by the addition of NaCl (1 g) before phase separation. An aliquot of the upper petroleum ether phase, containing lipophilic antioxidant, was evaporated under a gentle stream of nitrogen and redissolved in a known volume of acetone and ethanol (1:1 v/v) to maintain solvent consistency. An aliquot of the remaining lower phase, containing hydrophilic antioxidants, was also recovered for subsequent analysis. All three isolates (initial, lipophilic, and hydrophilic) were then evaluated for antioxidant capacity as described. All data except total soluble solids and acids were expressed on a dry weight basis.

Statistical analysis. Each experiment was set up using a randomized complete block design, blocking for ripeness stage at harvest and ripening method. Sample sizes were n = 3 for all ripeness stages that were stored, and n = 4 (Expt. 1) and n = 7 (Expt. 2) for full-red harvested peppers. Analysis of variance was conducted using GLM procedure (JMP software Version 5; SAS, 2002), with mean separation performed by the LSD test (P < 0.05).

Results and Discussion

Pepper ripening and quality attributes. Continuous exposure to 100 μL·L⁻¹ ethylene significantly accelerated the breakdown of chlorophyll such that these peppers reached full-red stage 3 to 6 d faster than those stored in air (Table 1). Bell peppers do not ripen uniformly, thus ripening rates were highly variable. Peppers
that were not completely red after 6 d storage (air treatment) had good quality; the fruit were very firm, had fresh appearance, stems were green, and were free from shrivel and decay (data not shown). After 10 d storage, the remaining fruit were moderately firm, had acceptable appearance, stems were yellow-green with slight desiccation, and there was no decay.

Peppers developed typical red color during storage, irrespective of harvest maturity or ripening method, in which values were: $L^* = 33.9$ to $35.5$; hue angle $= 24.3$ to $29.0$; chroma $= 24.2$ to $29.9$ (Table 2). These were comparable to color values for bell pepper (cv. Triple 4) harvested at full-red stage ($L^* = 34.2$, hue angle $= 25.7$, chroma value $= 30.1$) reported previously by Molinari et al. (1999).

Soluble solids content (SSC), total titratable acidity (TTA) and pulp pH were not affected by harvest maturity or ripening method. SSC ranged from 6.1 to 7.9 °Brix, TTA from 0.22 to 0.34 meq malic acid/g and pulp pH from 4.9 to 5.1 (Table 3). The higher SSC values for peppers in Expt. 2 can be attributed to more carbohydate accumulation due to longer daylength, higher ambient temperature and stronger light intensity than for peppers in Expt. 1. Peppers from Expt. 2 underwent primary growth and development during March and April when the daylength ranged from 11 h 38 min to 12 h 34 min, the mean outdoor temperature was 18 °C and mean maximum/minimum outdoor temperatures were 27/10 °C. In contrast, peppers for Expt. 1 underwent growth and development during December and January when the daylength ranged from 10 h 13 min to 10 h 40 min, the mean outdoor temperature was 13 °C and maximum/minimum outdoor temperatures were 20/6 °C (U.S. Naval Observatory, 2003). Minor significant differences in pulp pH would not cause perceptible flavor differences to consumers.

**Physicochemical attributes**. Total carotenoids were quantified spectrophotometrically in each fruit at 452 and 472 nm, subjectively corresponding to red (capsanthin) and yellow (β-carotene) carotenoids, respectively (Minguez-Mosquera and Hormero-Mendez, 1994). Red bell peppers were reported to contain six different carotenoids in concentrations ranging from 1.15 to 4.08 μg g⁻¹ dry weight, with lutein and capsanthin the predominant compounds in ripe fruit (Russo and Howard, 2002), while an earlier study found that in addition to capsanthin, β-carotene and violaxanthin were predominant carotenoids (Curl, 1962). In this study, relative changes in concentrations of total carotenoids measured at each wavelength were similar ($r = 0.99$), and indicated no differences due to fruit maturity at harvest or postharvest treatment (Table 4). Carotenoid development occurs as the chloroplast is transformed into chromoplasts during ripening, resulting in novel synthesis of various carotenoids that are not present in green fruit (Hormero-Mendez and Minguez-Mosquera, 2000). Ethylene exposure, despite decreasing the time to reach full red stage, was ineffective at increasing total carotenoids. Data indicate a high degree of variability between bell pepper harvested at 10% to 30% color but due to their nonclimacteric nature, the lack of additional carotenoid synthesis with ethylene exposure was a further indication that chlorophyll destruction, and not alteration of carotenoid biosynthetic pathways, was potentially responsible for fruit color development.

Carotenoid development was higher in peppers from Expt. 2, which had more than twice the carotenoid concentration than peppers from Expt. 1. Fruit harvested at 10% to 30% color in Expt. 2 had 79.6 and 64.2 mg·kg⁻¹ capsanthin for air and ethylene-ripened fruit, respectively, while values from Expt. 1 were 41.9 and 28.3 32 mg·kg⁻¹ for the same respective treatments (Table 4). Russo and Howard (2002) reported that bell peppers grown under greenhouse conditions had higher carotenoid concentrations compared to field-grown fruit, attributable to greater light intensity and lack of stress on the greenhouse-grown plants. Likewise, this study indicated that light intensity might have impacted phytochemical development since daylength was longer, and light intensity and temperature were higher for peppers from Expt. 2. Bell peppers from Expt. 2 that were ripened on the plant also had higher concentrations of carotenoids than those ripened in air or in ethylene.

### Table 1. Effect of harvest maturity and postharvest storage treatment on time for ‘Robusta’ bell peppers to reach full-red color.

| Maturity | Expt. 1 | Expt. 2 |
|----------|---------|---------|
| at harvest | Air | C_H4 | Air | C_H4 |
| <10% Red | 11 a | 6.0 a | --- | --- |
| 10% to 30% Red | 11 a | 4.7 a | 10.0 a | 6.0 a |
| 30% to 60% Red | 11 a | 3.0 a | --- | --- |

*Fruit were stored in air or 100 ppm ethylene at 20 °C and 90% RH until full red.

**Values with different letters within columns of the same postharvest treatment are significantly different (LSD test, $P < 0.05$), and indicate the effect of harvest maturity.**

**For each experiment values within rows for each harvest maturity stage were significantly different (LSD test, $P < 0.05$) and indicate the effect of postharvest treatment.**

### Table 2. Effect of harvest maturity and postharvest storage treatment on external CIE color attributes of ‘Robusta’ bell peppers at full-red stage (Expt. 1).

| Maturity | L* | Hue angle (°) | Chroma value |
|----------|----|--------------|--------------|
| at harvest | Air | C_H4 | Air | C_H4 | Air | C_H4 |
| <10% Red | 33.9 b | 35.5 a | 29.0 a | 25.4 a | 24.2 a | 26.0 a |
| 10% to 30% Red | 34.5 a | 36.8 a | 27.3 a | 24.3 a | 25.0 a | 27.8 a |
| 30% to 60% Red | 34.4 a | 33.9 a | 27.9 a | 26.6 a | 24.4 a | 29.9 b |

*Values with different letters within columns of the same postharvest treatment are significantly different (LSD test, $P < 0.05$), and indicate the effect of harvest maturity.

### Table 3. Effect of harvest maturity and postharvest storage treatment on the soluble solids content (SSC), total titratable acidity (TTA), and pH of ‘Robusta’ bell peppers at full-red stage.

| Maturity | SSC (°Brix) | TTA* | pH |
|----------|-------------|------|-----|
| at harvest | Air | C_H4 | Air | C_H4 | Air | C_H4 |
| <10% Red | 6.3 a | 5.4 a | 0.29 a | 0.34 a | 4.9 b | 5.0 a |
| 10% to 30% Red | 6.1 a | 6.0 a | 0.29 a | 0.28 a | 5.0 a | 5.0 a |
| 30% to 60% Red | 7.4 a | 6.0 a | 0.28 a | 0.27 a | 5.0 a | 5.0 a |
| Red-ripe | 6.5 a | --- | 0.29 a | --- | 5.0 a | --- |

*For each experiment, values with different letters within columns of the same postharvest treatment are significantly different (LSD test, $P < 0.05$), and indicate the effect of harvest maturity.

**Values within rows for each harvest maturity stage were not significantly different (LSD test, $P < 0.05$), indicating there was no effect of postharvest treatment.**

Changes in ascorbic acid and soluble phenolics were similar to trends observed for total carotenoids, with essentially no differences due to ripeness stage at harvest or ripening method (Table 4). As with carotenoids, concentrations of both total ascorbic acid and total soluble phenolics for peppers from Expt. 2 were more than twice those from Expt. 1, indicating that increased synthesis rates were likely a function of temperature and light exposure. Ripening on the plant did not significantly affect total ascorbic acid or total soluble phenolic content, yet concentrations varied appreciably and indicated extensive diversity among individual fruit during ripening (Table 4).

Predominant antioxidant compounds reported in bell peppers include carotenoids, ascorbic acid, flavonoids, phenolics acids, and tocopherols (Howard et al., 2000; Markus et al., 1999; Osuna-Garcia et al., 1998). Despite differences in solubility of these antioxidant compounds, appreciable contributions to total antioxidant capacity were found for both hydrophilic and lipophilic bell pepper extracts. Overall, the antioxidant capacity was not altered by harvest stage or ethylene exposure in either experiment (Table 4), although ripe-harvested fruit from Expt. 2 had a higher antioxidant capacity.“

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734 HORT SCIENCE VOL. 40(3) JUNE 2005
capacity (8.6 μmol Trolox equivalents/g) than those from Expt. 1, and were well correlated to both ascorbic acid and soluble phenolics content (r = 0.84 and 0.89, respectively). When fractionated to subsequently determine relative contributions to antioxidant capacity based on polarity, the hydrophilic fraction (polyphenolics and ascorbic acid) contained 88% ± 2.7% of the total antioxidant activity of the initial isolate (whole fruit) compared to 34% ± 1.0% for the lipophilic fraction (carotenoids and xanthophylls). Capsanthin, the predominant red carotenoid in red bell pepper, was specifically shown to be an effective free radical scavenger even when esterified with fatty acids (Matsufuji et al., 1998), as were the carotenoids in the isolated isolates.

It was previously reported that flavonoids in the presence of polyphenolics or ascorbic acid might adversely impact antioxidant activity or that prooxidant interactions between ascorbic acid and metal ions may exist (Howard et al., 2000). Since the total antioxidant capacity from each fraction was appreciably higher than the initial stock isolate, there is indication that physical and/or chemical interactions among constituents in these fractions unfavorably impacted radical-scapenging properties. Additionally, since the hydrophilic fraction contained the highest contribution to total antioxidant capacity, the difference in values may reflect the high polarity of the radical system, selectivity against peroxyl radicals, or indicate competition or interactions between antioxidant compounds present.

This study demonstrated that ethylene has potential for use as an effective postharvest treatment to facilitate destruction of chlorophyll and accelerate color change without altering the flavor, content of phytochemicals or antioxidant capacity in red bell pepper, cv. Robusta. Additionally, rates of phytochemical synthesis were not affected by fruit maturity at harvest, although fruit harvested with full color had appreciably higher amounts of phytochemicals and antioxidant capacity. These results indicate that ripening of bell pepper for this cultivar could be accelerated by harvest at partially ripe stage and exposure to ethylene at 20 °C, thereby increasing productivity for growers. Use of vapor barriers during ripening and postripening storage at 10°C would likely further extend marketable life.

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### Table 4. Effect of harvest maturity and postharvest storage treatment on individual and total carotenoid contents, ascorbic acid content, total phenolic content, and antioxidant activity of ‘Robusta’ bell peppers at full-red stage.

| Maturity at harvest | Carotenoids at 452 nm (mg·kg⁻¹ as capsanthin, dry wt) | Carotenoids at 472 nm (mg·kg⁻¹ as β-carotene, dry wt) | Ascorbic acid (mg·kg⁻¹, dry wt) | Total soluble phenolics (mg·kg⁻¹ as gallic acid, dry wt) | Antioxidant capacity (µmol Trolox equiv/g, dry wt) |
|---------------------|---------------------------------|---------------------------------|-------------------------------|---------------------------------|---------------------------------|
|                     | Air C H₂ | Air C H₂ | Air C H₂ | Air C H₂ | Air C H₂ |
| Expt. 1             |                     |                     |                     |                     |                     |
| 10% to 20% Red      | 32.7 b        | 30.4 a        | 26.2 ab      | 25.2 a        | 1,150 a      | 749 a    | 36.9 ab      | 30.4 a        | 17.5 b        | 20.2 a        |
| 30% to 60% Red      | 41.9 a        | 28.3 a        | 33.9 b        | 23.8 a        | 1,000 a      | 722 a    | 39.9 ab      | 30.5 a        | 19.3 ab       | 19.0 a        |
| Full-red            | 22.6 b        | 37.3 a        | 19.1 b        | 31.0 a        | 924 a        | 880 a    | 32.6 b        | 45.3 a        | 17.8 b        | 26.1 a        |
| Expt. 2             |                     |                     |                     |                     |                     |
| 10% to 20% Red      | 35.9 a        | ---           | 29.5 ab      | ---           | 1,110 a      | 46.8 a    | ---           | ---           | 25.3 a        | ---           |
| Full-red            | 79.6 b        | 64.2 a        | 99.5 b        | 76.5 a        | 2,560 a      | 2,000 b   | 53.8 a        | 44.5 b        | 29.2 a        | 26.4 a        |
|                     | 121.9 a       | ---           | 150.5 a       | ---           | 2,540 a      | ---      | 60.0 a        | ---           | 35.8 a        | ---           |

*Values with a dagger within columns for each harvest maturity stage and experiment are significantly different (LSD test, P ≤ 0.05), and indicate the effect of fruit maturity at harvest.

*Values with different letters within columns of the same postharvest treatment are significantly different (LSD test, P ≤ 0.05).
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