Controlled-Atmosphere Effects on Postharvest Quality and Antioxidant Activity of Cranberry Fruits

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The effects of controlled-atmosphere (CA) storage on the firmness, respiration rate, quality, weight loss, total phenolics and flavonoids contents, and total antioxidant activities of the Pilgrim and Stevens cultivars of cranberries (Vaccinium macrocarpon Aiton) have been studied during storage in atmospheres of 2, 21, and 70% O2 with 0, 15, and 30% CO2 (balance N2); and 100% N2 at 3 °C. Elevated CO2 concentrations decreased bruising, physiological breakdown, and decay of berries, thereby reducing fruit losses. Respiration and weight loss of fruits decreased, but fruit softening increased, at higher CO2 concentrations. Accumulations of acetaldehyde, ethanol, and ethyl acetate varied by cultivar and storage atmosphere but were generally highest in the 2 and 70% O2 and 100% N2 atmospheres and increased in response to elevated CO2 concentrations. Overall, the 30% CO2 plus 21% O2 atmosphere appeared optimal for the storage of cranberries. Sensory analysis is required, however, to confirm that accumulations of fermentation products at this atmosphere are acceptable for consumers. Stevens fruits had a higher phenolics content and total antioxidant activity than Pilgrim fruits. The storage atmosphere did not affect the content of total phenolics or flavonoids. However, the total antioxidant activity of the fruits increased overall by about 45% in fruits stored in air. This increase was prevented by storage in 30% CO2 plus 21% O2.

KEYWORDS: Cranberry; Vaccinium macrocarpon Aiton; controlled atmosphere; antioxidants; phenolics; flavonoids

INTRODUCTION

Cranberry fruits are grown mainly for processing purposes, but in the U.S., strong markets for fresh fruit exist for the Thanksgiving and Christmas holiday seasons. In addition to the traditional aspects of cranberry consumption, health benefits of the fruit have become increasingly important. Fruits of Vaccinium sp. including cranberries are particularly rich in flavonoids and other phenolic compounds (1, 2). These compounds have health-promoting benefits as antioxidants and anticarcinogens (1, 3, 4). Wang and Stretch (2) found that the concentrations of antioxidants, anthocyanins, and total phenolics in 10 cranberry cultivars ranged from 8.6 to 14.1 µmol of Trolox equivalents per gram, from 0.20 to 0.66 mg of cyanidin-3-galactoside per gram of fresh fruits, and from 1.20 to 1.76 mg of gallic acid equivalents per gram of fresh fruits, respectively.

Cranberries are harvested by wet or dry methods. The most common method, wet-harvesting, involves scooping or raking of the fruit in flooded bogs with a motorized water reel. Dry-harvesting involves hand picking, hand scooping, and machine scooping (5). Recently, a dry-harvester that works in a fashion similar to a vacuum cleaner has been developed in upstate New York (6). Harvest methods influence storage life through the different degrees of bruising and stress imposed on the fruits (7). Cranberries can be stored for 2–3 months with adequate cooling and ventilation, but quality losses can be high because of shrinkage due to moisture loss, physiological breakdown, end rot, and fungal diseases (8). Cultivar and fruit maturity can also affect storage quality (8). Optimum storage temperatures were reported to be 1.7–4.4 °C (9).

Controlled-atmosphere (CA) storage can extend the storage life of fruits and vegetables by decreasing metabolism and suppressing postharvest decay, but reports on the CA storage of cranberries are limited. Cranberries responded relatively poorly to CA conditions of 0, 5, 10% CO2 with 3, 10, and 21% O2 (10), perhaps because of limited gas diffusion of the berry and naturally high internal CO2 concentrations (11). Initial treatments with N2 gas have been shown to improve cranberry storage life by reducing fungal decay (12). These studies were carried out before the advent of rapid CA storage technology in which CA regimes are imposed within a few days of harvest, and in addition, further research is needed to identify potential benefits and limits of tolerance of cranberries to CO2 (13). Also, although Wszelaki and Mitcham (14) found that superatmo-
spheric O₂ reduced the decay of strawberries, no studies on the
effect of this technology are available for cranberries.

Although information about the effects of air storage on
antioxidant capacity and associated antioxidant compounds of
strawberries, raspberries, and blueberries is becoming available
(15), little is known about the effects of postharvest treatments,
especially CA storage, on the antioxidant and phenolics contents of
fruit. Wang and Stretch (2) found that the antioxidant
activities and anthocyanin and phenolic contents of cranberries
increased during storage, with maximum increases occurring
at 15 °C compared with 0, 5, 10, and 20 °C.

The objective of this research was to determine the effect of
a wider range of CO₂ and O₂ concentrations, including super-
atmospheric O₂, than those previously studied on the quality,
storage life, total phenolic and flavonoid contents, and total
antioxidant activities of cranberries.

MATERIALS AND METHODS

Reagents. Folin–Ciocalteu’s phenol reagent, (+)-catechin, sodium
nitrite (NaNO₂), and acetaldehyde were obtained from Sigma (St. Louis,
MO). Gallic acid, sodium carbonate, aluminum chloride, ethyl acetate,
and sodium hydroxide were purchased from Fisher Scientific (Pitts-
burgh, PA). Acetone and ethanol were obtained from Mallinkrodt
(Paris, Kentucky) and Pharmacomproducts Inc. (Brookfield, CT), respec-
tively.

Fruit Material. Cranberries were purchased from the Oswego
Cranberry Company LLC (Pulaski, NY). The berries from each cultivar
were dry-harvested by vacuum on two occasions: Pilgrim cultivar
cranberries were harvested in 2000 on October 3 (harvest 1) and October
16 (harvest 2), and Stevens cultivar cranberries were harvested on
October 25 (harvest 1) and November 7 (harvest 2). The berries were
sorted and packaged in clamshell containers (each approximately 450 g)
and transported to our laboratory on the day of harvest in cardboard
boxes.

CA Storage. Berries were weighed (500 g) in half-gallon mason jars
and placed in a flow-through system kept at 3 °C. The gases (O₂,
CO₂, and N₂) were mixed in a main flowboard to obtain the desired
atmospheres of 0% O₂, 10% O₂, and 2, 21, and 70% O₂ with 0,
15, and 30% CO₂. Each gas mixture was distributed to the jars (75
mL/min) through smaller flowboards, each with 12 outlets. Atmospheres
were verified daily by gas chromatography (model 1200 gas partitioner,
Fisher Scientific, Springfield, NJ). Three replicates of fruits (three jars)
from each treatment were evaluated at each sampling time.

Determination of Weight Loss and Fruit Losses. The fruit weight
was recorded, and the percentage weight loss from harvest was
ca
culated. The fruits were inspected to sort out the sound berries from
decayed/spoiled berries. Unsound berries were those with mechanical
damage, poppers (turgid berries with internal tissue breakdown), and
fungal and physiological breakdown. Both the sound and the unsound
berries were weighed, and the percentage of unsound berries was
calculated. Only sound fruits were used for subsequent measurements.

Respiration Rate and Ethylene Production Rate. Respiration rates of
fruits from harvest 1 were measured by a closed system. Fruits (100 g),
removed from CA storage, were placed in a 500-mL jar and kept
overnight in air at 3 °C to equilibrate, and then the jar was sealed with
a cap containing a rubber septum for headspace gas sampling. The initial
and final (after 3–4 h storage at 3 °C) headspace gas compositions
were analyzed using a gas chromatograph with a thermal conductivity
detector (Fisher Gas Partitioner, model 1200, Fisher Scientific,
Springfield, NJ). The gases were separated in a dual-column config-
uration consisting of a 1.8 m × 0.32 cm stainless steel column packed
with 80/100 mesh poropak Q and a 3.3 m × 0.48 cm stainless steel
column packed with 60/80 mesh molecular sieve 13X (Supelco,
Bellefonte, PA). The column and injector temperatures were 90 and
135 °C, respectively. Ethylene concentrations were determined by
headspace sampling using a gas chromatograph equipped with a flame
ionization detector and a 1 m × 3 mm stainless steel column packed
with 80/100 Alumina F-1 (Hewlett-Packard 5890, Wilmington, DE).

The column, detector, and injector temperatures were 190, 240, and
220 °C, respectively.

Firmness Measurement. Firmness of the berries was measured by
puncture test using an Instron Universal Testing Machine (model 1122,
Instron Corp., Canton, MA.). The puncture test was carried out by a
flat-point probe (4-mm diameter) installed on the Instron with a cross-
head speed of 50 mm/min. The maximum force to punch a hole through
a fruit was recorded in newtons. Ten fruits randomly selected from
each treatment replicate were measured.

Volatilie Analysis. Selected berries (50 g) were frozen in liquid N₂
and kept at −80 °C prior to analysis. The acetaldehyde, ethanol, and
ethyl acetate concentrations in the tissues were measured using a
modification of the procedure described by Gunes et al. (16). Five grams
of frozen powdered fruits was placed in each 20-mL glass vial (Fisher
Scientific, Pittsburgh, PA), which was then sealed with Tellon-lined
septum caps. The vials were heated in a water bath (Precision Scientific
Co., Chicago, IL) at 70 °C for 25 min. A 0.1-mL-headspace gas sample
was injected into a gas chromatograph with a FID (HP 5890, Hewlett-
Packard, Wilmington, DE). The volatiles were separated in a 15 m ×
0.53 mm Stabilwax wide-bore capillary column with 1.0 µm coating
(Restek Corp., Bellefonte, PA). The column temperature was set to 40
°C for the initial 4 min and then increased to 220 °C at a 20 °C/min
rate and held at that temperature for 10 min. The areas of the peaks
corresponding to acetaldehyde, ethyl acetate and ethanol were quantified
with an integrator (HP 3394, Hewlett-Packard, Wilmington, DE).

Standard curves were obtained using authentic compounds.

Extraction of Phenolics and Flavonoids. Frozen berries (only the
first harvest from both cultivars) were crushed into coarse pieces, and
30 g of fruit was homogenized with 60 mL of 80% acetone using a
Waring blender for 3 min. The samples were further homogenized using
a homogenizer (Ultra-Turrax T25, IKA Works, Inc., Wilmington,
NC) for an additional 2 min and filtered through 82 Whatman paper.
The filtrate was recovered and the acetone was evaporated off by a
rotary evaporator (model 78820-00, Labconco, Kansas City, MO) at
45 °C for 30 min. The samples were then brought to the desired volume
(50 mL) with deionized water and kept frozen at −80 °C prior to
analysis.

Determination of Total Phenolics. The total phenolic contents of
the berries were measured using a modified Folin–Ciocalteu colori-
metric method (17). A 0.2-mL sample of fruit extract was added to a
25-mL flask containing 5 mL of deionized water. One milliliter of
Folin–Ciocalteu reagent was added to the mixture, and the flask was
shaken and allowed to stand at room temperature for 5 min. Ten
milliliters of 7% Na₂CO₃ was added and the solution was diluted to
do the desired volume (25 mL) with deionized water. Absorbance
was measured at 750 nm versus a blank after 2 h at room temperature.
The results are expressed as gallic acid (GA) equivalents using the
standard curve (absorbance versus concentration) prepared from authentic
gallic acid.

Determination of Total Flavonoids. The total flavonoid content
of the berries was determined by a colorimetric method developed by
Jia et al. (18). A 0.3-mL fruit extract was added to a 10-mL flask
containing 5 mL of deionized water. A 0.3-mL portion of 5% NaNO₂
was added to this mixture, which was allowed to stand for 5 min at
room temperature. A 0.3-mL portion of 10% AlCl₃·6H₂O was added,
and the mixture was allowed to stand for 6 min at room temperature.
Two milliliters of 1 M NaOH was added, and the solution was diluted
to the desired volume (10 mL) with deionized water. The absorbance
of the solution versus a blank at 510 nm was measured immediately.
The results are expressed as catechin equivalents using a standard curve
(absorbance versus concentration) prepared from authentic catechin.

Quantification of Total Antioxidant Activity. The total antioxidant
activity of the cranberry extracts (at harvest and after storage for 2
months under air and 21% O₂ plus 30% CO₂) was measured by the
total oxyradical scavenging capacity (TOSC) assay (19). Antioxidant
activity was assessed at four different time points (15, 30, 45, and 60
min) and six concentrations to determine the TOSC value. The TOSC
value was quantified from the integration of the area under the kinetics
Table 1. Firmness (N) of Pilgrim and Stevens Cranberries Stored under Different Combinations of O₂ and CO₂ at 3 °C for 2 Months

<table>
<thead>
<tr>
<th>O₂ (%)</th>
<th>CO₂ (%)</th>
<th>Pilgrim</th>
<th>Stevens</th>
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<tr>
<td>0</td>
<td>13.3</td>
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<td>–</td>
</tr>
<tr>
<td>2</td>
<td>14.9</td>
<td>12.7</td>
<td>12.9</td>
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<td>14.1</td>
<td>12.7</td>
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</tr>
<tr>
<td>70</td>
<td>14.6</td>
<td>12.9</td>
<td>12.1</td>
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</table>

<table>
<thead>
<tr>
<th>O₂ (%)</th>
<th>CO₂ (%)</th>
<th>Pilgrim</th>
<th>Stevens</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
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<td>–</td>
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<tr>
<td>70</td>
<td>13.6</td>
<td>11.9</td>
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* Data for both harvests have been combined. Pooled standard deviations are 2.69 and 2.34 for Pilgrim and Stevens, respectively. Firmness at harvest was 13.3 and 12.6 N for Pilgrim and Stevens, respectively.

curve according to the equation

\[
TOSC = 100 - \left(\frac{\text{SA}}{\text{CA}}\right) \times 100
\]

where \(\text{SA}\) and \(\text{CA}\) are the integrated areas from the sample and the control reaction, respectively. Total antioxidative activity is expressed as the median effective dose (EC₅₀ in micrograms of tissue per milliliter) for the samples.

**Statistical Analysis.** An analysis of the variance was performed using the general linear model procedure to determine the main effects and interactions (Minitab Release 13.1, State College, PA). Multiple comparisons of individual treatments were also performed by Fisher’s least significant difference (LSD) procedure in one-way analysis of variance.

**RESULTS**

**Fruit Firmness.** The firmness of the cranberries was found to be greater after storage in 0% CO₂ than at harvest, but overall, firmness was not affected by cultivar or O₂ in the storage atmosphere (Table 1). However, fruits stored in 15 and 30% CO₂ were softer in both cultivars compared with fruits from CO₂-free atmospheres. The overall firmness readings of fruits stored under 0, 15, and 30% CO₂ were 14.2, 12.8, and 12.6 N for Pilgrim and 13.9, 12.3, and 12.1 N for Stevens, respectively. Thus, 15 and 30% CO₂ levels resulted in up to 10 and 14% reductions in firmness compared to 0% CO₂. Differences in firmness between fruits kept at 15 and 30% CO₂ were not significant.

**Respiration and Ethylene Production.** The respiration of the cranberries, in terms of both CO₂ production rates (RCO₂) and O₂ consumption rates (RO₂), was measured at only one time point after removals of the berries from CA storage, and therefore, dynamic changes in fruit after CA treatments over time were not taken into account. However, respiration was affected by cultivar and CO₂ and O₂ concentrations in the storage atmosphere (Table 2). Both RCO₂ and RO₂ were lower for the 30% CO₂ atmosphere than for the 0 and 15% CO₂ atmospheres (\(P \leq 0.05\)), but there was no difference between the 0 and 15% CO₂ treatments. The overall mean RCO₂ values of fruits stored at 0, 15, and 30% CO₂ were 5.0, 4.7, and 4.8 mL of CO₂/kg/h, respectively, for Pilgrim and 2.8, 2.8, and 2.12 mL of CO₂/kg/h, respectively, for Stevens. The overall mean RO₂ values of fruits stored at 0, 15, and 30% CO₂, respectively, were 3.4, 2.2 and 1.9 mL of O₂/kg/h for Pilgrim and 2.1, 1.3, and 1.5 mL of O₂/kg/h for Stevens. Storage of fruits at 2% O₂ resulted in a lower respiration rate than the ambient and superatmospheric O₂ storage (\(P \leq 0.05\)). However, the respiration rates were similar at 21 and 70% O₂. Overall, the mean RCO₂ values for 2% O₂ were 4.2 and 2.2 mL of CO₂/kg/h for Pilgrim and Stevens, respectively, whereas for 21 and 70% O₂, they were 4.9 and 5.4 mL of CO₂/kg/h, respectively, for Pilgrim and Stevens. The overall mean RO₂ values were 5.0, 4.7, and 4.8 mL of O₂/kg/h for Pilgrim and 2.8, 2.8, and 2.12 mL of O₂/kg/h for Stevens.

**Fruit Bruising and Physiological and Fungal Breakdown.** In general, losses of fruits due to damage, whether from the presence of bruising or physiological or fungal breakdown, decreased as the CO₂ concentration in the 2 and 21% O₂ atmospheres increased (Figure 1A–D). However, the reduced losses associated with the CO₂ atmospheres were lower in fruits from the second harvest. Superatmospheric O₂ increased the

![Figure 1](image-url)
loss of fruits compared with the 2 and 21% O2 atmospheres. The fruits from both cultivars were damaged extensively by the end of the fourth month of CA storage (>97% unsound fruit; data not shown).

Among the atmospheres tested, 21% O2 plus 30% CO2 resulted in the least loss of quality. The incidence of unsound fruits in this atmosphere was about 15% in contrast to 75% in air for Stevens (Figure 1C). An atmosphere of 2% O2 plus 30% CO2 also resulted in better-quality fruits than did air storage, but it was not as beneficial as 21% O2 plus 30% CO2.

Weight Loss. The weight loss of the fruits was less than 1% for all atmospheres (data not shown), probably because of the high relative humidity (>95%) in the storage compartments. However, fruits stored in 30% CO2 atmospheres had up to 30% less weight loss compared to those stored in CO2-free atmospheres (P ≤ 0.05), and overall, Pilgrim had a greater weight loss (0.85%) than Stevens (0.70%).

Fermentation Products. Accumulations of acetaldehyde, ethanol, and ethyl acetate were affected by atmosphere, as well as interactions between cultivar and atmosphere (Figures 2–4A,B; Table 3). These fermentation products accumulated to similar extents in both cultivars. Because no harvest time effect was detected, data for both harvests were combined within each cultivar. Accumulation of acetaldehyde increased under elevated CO2 conditions in 70% O2 (Figure 2A,B). However, there was no additional effect of CO2 at 2% and 21% O2. Fruits kept in 0% O2 also accumulated large concentrations of acetaldehyde.

Accumulation of ethanol showed patterns similar to those found for acetaldehyde, with a significant atmosphere effect (Figure 3A,B). Greater concentrations of ethanol were found in fruits stored in 15 and 30% CO2 atmospheres compared with 0% CO2 for Pilgrim, but no significant effect of CO2 was detected for Stevens. The ethanol concentrations for the 15 and 30% CO2 levels were similar. Superatmospheric O2 resulted in higher ethanol accumulations than did 2 and 21% O2 for both cultivars (P ≤ 0.05). The accumulation of ethanol was greatest under 0% O2 for both Pilgrim and Stevens (Figure 3A,B).

Ethyl acetate showed little accumulation at 0, 2 or 21% O2, irrespective of the CO2 concentration, but it did accumulate in 70% O2 (Figure 4A,B). For Pilgrim, but not Stevens, increasing CO2 concentrations in 70% O2 resulted in higher ethyl acetate accumulations.

Table 3. Analysis of Variancea for Acetaldehyde, Ethanol, and Ethyl Acetate in Pilgrim and Stevens Cranberries Stored under Different Combinations of O2 and CO2 at 3 °C for 2 Months

<table>
<thead>
<tr>
<th>source</th>
<th>DFa</th>
<th>acetaldehydeb</th>
<th>ethanold</th>
<th>ethyl acetateb,c</th>
</tr>
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<tr>
<td>O2</td>
<td>2</td>
<td>***</td>
<td>***</td>
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</tr>
<tr>
<td>CO2</td>
<td>2</td>
<td>***</td>
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</tr>
<tr>
<td>cultivar (C)</td>
<td>1</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>O2*CO2</td>
<td>4</td>
<td>*</td>
<td>*</td>
<td>***</td>
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<td>O2*C</td>
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<td>*</td>
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<td>NS</td>
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<tr>
<td>CO2*C</td>
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<td>NS</td>
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<td>NS</td>
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<tr>
<td>O2<em>CO2</em>C</td>
<td>4</td>
<td>*</td>
<td>*</td>
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* DF, degrees of freedom. ** NS, *, **, and *** indicate nonsignificant and significant at P ≤ 0.05, 0.01, and 0.001, respectively. a Natural logarithm of the transformed data was used to satisfy the assumptions of the model.

Total Phenolic and Flavonoid Contents. Stevens cranberries had a slightly higher total phenolic content than Pilgrim fruits (Table 4), the average values over all storage conditions being 3.45 and 3.27 mg/g of tissue, respectively. The total phenolic contents of the fruits were not affected by CA storage conditions.

The storage atmosphere affected the content of flavonoids in the cranberries. Flavonoid contents were lower in 15% CO2 than in 0 and 30% CO2 (P ≤ 0.05), but the magnitude of the overall difference was very small (0.09 mg/g). Neither the O2 level nor its interactions with the CO2 level affected the flavonoid content. The total flavonoids content of Pilgrim did not change over 2 months of CA storage, but that of Stevens increased slightly in some atmospheres, such as 2% O2 with 0 and 30% CO2, air, and 70% O2 with 0 and 15% CO2. Overall, the flavonoid contents of Pilgrim and Stevens were similar.

Total Antioxidant Activity. We selected the optimum atmosphere of 21% O2 plus 30% CO2 indicated from the above
are 0.06 and 0.11 for Pilgrim and Stevens, respectively. A increase. 50% from the initial values in Stevens and Pilgrim, respectively, 5. The antioxidant activities of the fruits increased by 36 and 30% CO2 in 21% O2, and to a lesser extent 30% CO2 with 2% CO2 in 21% O2, and to a lesser extent 30% CO2 with 2%.

DISCUSSION

Our results indicate that CA atmospheres can beneficially affect cranberry fruit quality. Among the atmospheres tested, 30% CO2 in 21% O2, and to a lesser extent 30% CO2 with 2%
method, which employs vacuum technology to collect the fruits (6). The extent of bruising associated with this method compared with alternative methods has not been studied. In addition, the berries might have been bruised during subsequent sorting operations employed in the elevator, bounce board, and conveyor belt used for grading. It has been reported that cranberries stored in chaff undergo less spoilage than those that are cleaned or sorted before storage (10). An additional factor in the high losses found in our experiments might have been the high relative humidity under CA conditions. Anderson et al. (10) found that CA storage was only as good as air storage for maintaining fruit quality when the humidity and the O₂ and CO₂ concentrations were regulated.

The effects of fermentation product accumulation at 21% O₂ and 30% CO₂ on fruit acceptability was not tested by formal sensory analysis in this study. Informal assessment indicated that flavor was not detrimentally affected; the acetaldehyde, ethyl acetate, and ethanol concentrations were less than 6, 0.30, and 161 mg/kg, respectively, and close to the olfactory threshold concentrations of 5, 0.015, and 100 μg/L for acetaldehyde, ethyl acetate, and ethanol, respectively, in apple essence for human perception (23). However, although the concentrations of these compounds at which the fruits become unacceptable can be expected to be much higher than the threshold concentrations, the effects of the fermentation product accumulation on fruit acceptability remain to be confirmed.

The total phenolic contents of the Pilgrim and Stevens cranberries at harvest were 3.33 and 3.45 mg/g, respectively, and therefore much higher than the values of 1.20 and 1.26 mg/g of fruits, respectively, reported by Wang and Stretch (2). The higher values might be due to inclusion of seeds in our study or to environmental influences.

The total phenolic contents of either cranberry cultivar did not increase during storage. Wang and Stretch (2) also found little increase in phenolic concentrations at 0 °C, but at 5 °C and above, increases were substantial. Our storage temperature of 3 °C was intermediate between these temperatures and indicates that storage temperature might be an important factor in phenolic metabolism.

Total antioxidant activity in Pilgrim and Stevens increased by 50 and 36%, respectively, during air storage for 2 months. Changes in total antioxidant activity were also affected by temperature (2). They were lowest at lower storage temperatures but increased by about 20% in both cultivars at 0 °C during three months of air storage. The correlations between antioxidant activity and concentrations of phenolics and anthocyanins were lowest at the lower storage temperatures (2).

The mechanism by which CA storage prevented the increase in total antioxidant activity that was found in air-stored fruits is not clear, but CA conditions might affect the release of bound phytochemicals that contribute to antioxidant activity. Heat-processed tomatoes have higher bioaccessible lycopene, apparently because thermal processing disrupts cell membranes and cell walls to release lycopene from the cell matrix and insoluble fibers (24). Although elevated CO₂ concentrations in the storage atmosphere did not maintain berry firmness relative to low CO₂ atmospheres, maintenance of cellular integrity by CA storage conditions might prevent the release of bound phenolics and flavonoids from the cell matrix of cranberry fruits and maintain lower antioxidant activities.

Overall, our data indicate that CA storage might have benefits for storage of cranberry fruits that are greater than previously realized. The storage life of cranberries can be extended using a CA of 21% O₂ plus 30% CO₂ through reductions in decay and respiration rate. This increase in storage life could provide extended availability of fresh cranberries with premium quality in the market, provided that the storability of unsorted fruits is greater than that shown for sorted and packed fruits and that accumulations of fermentation products do not reach unacceptable levels. The concentrations of phytochemicals and total antioxidant activity in cranberries were maintained at harvest levels during storage in this atmosphere.

**ABBREVIATIONS USED**

- CA, controlled atmosphere; CO₂, carbon dioxide; O₂, oxygen; N₂, nitrogen; RCO₂, CO₂ production rate; RO₂, O₂ consumption rate; RQ, respiratory quotient; P, probability; EC₅₀, median effective dose; NaNO₂, sodium nitrite, AlCl₃·6H₂O, aluminum chloride; NaOH, sodium hydroxide; KOH, potassium hydroxide; L, liter; mL, milliliter; μL, microliter; min, minute; h, hour; kg, kilogram; g, gram; °C, degrees Celsius; nm, nanometer; M, molar; N, Newton.

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**LITERATURE CITED**


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