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Effect of Initial Low Oxygen Concentration on Respiration and Quality of Fresh-cut Cabbages

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The effects of perforated film package (PFP) on the gas concentrations and respiration rates of fresh-cut cabbage with initial O2 5, 8, 10 and 21% were studied at 5 and 20°C. The microbial growth, appearance, flavor and ascorbic acid content were also determined. The respiration rates of fresh-cut cabbages were suppressed under initial low O2 during storage of 4 days, especially in 5 and 8% O2 concentrations. The anaerobic respiration did not occur under low O2 levels. It was also found that fresh-cut cabbages had better color retention and quality, reduced respiration rate and microbial population in PFP. Although there was the difference of quality attributes between PFP and MAP, but no significant difference was found.

The effect of initial low O2 5% was the most important among different levels of initial low O2 according to the oxidation of ascorbic acid (AA) and browning of fresh-cut cabbage. Microbial analysis also showed that total count on the surface of fresh-cut cabbage was lowest among initial low O2 treatments. Total ascorbic acid (TAA) decreased by AA oxidation after cutting. Loss in TAA was much lower in PFP than that in MAP. Moreover, the results of sensory evaluation showed that there were no differences in all treatments at 5°C throughout storage. At 20°C, flavor reached 2-grade in all treatments and become 3-grade by 4 days in 21%MAP. The appearances were reached 2-grades after 3 days in 10%PFP and 21%MAP. These suggest that better quality of fresh-cut cabbage could be obtained from the combination of PFP with initial O2 5-8% level and 5°C.

INTRODUCTION

Fresh-cut horticultural products are prepared and handled to maintain their fresh nature while providing convenience to the user. The most food processing techniques stabilize the products and lengthen their storage and shelf life. While fresh-cut vegetables which increase in cut-damaged surface and availability of cell nutrients provide conditions that increase the number and type of microbes that develop and lead to greater opportunity for contamination by pathogenic organisms (Magnuson, et al., 1990; Marchetti, et al., 1992).

Microbial deterioration of fresh-cut vegetables can be controlled by several methods.
Low temperature and packaging technology are vitally important ways to inhibit the proliferation of microorganisms. In particular, the use of permeable polymeric films could retard microbial growth by affecting the concentration of gases, but also maintain higher relative humidity to avoid water loss within the package (Watada et al., 1996). Modified atmosphere packaging (MAP) has been considered to be effective method to extend the shelf life of fresh-cut fruits and vegetables (Kader et al., 1986). The low O₂ and high CO₂ levels that are modified through the product's respiration and permeable film slow down respiration and have an effect of inhibited microbial growth in MAP (Zagory and Kader, 1989; Kader et al., 1989; Lopez-Briones et al., 1993). However, over tolerance limit to low O₂ and high CO₂, it resulted in shift from aerobic to anaerobic respiration and off-flavor and tissue breakdown. The improperly modified atmospheric conditions may adversely affect product quality. Therefore, the selection of suitable packaging materials is critical to obtain the desirable environment for extending the shelf life of fresh-cut vegetables. The optimum storage temperature and gas concentration need to be maintained during whole chain from harvest processing to consumption.

Fresh-cut processing increases the rate of metabolic processes that cause deterioration of fresh products. The physical damage or wounding caused by preparation increase respiration and ethylene production in short time, and associated increases in the rate of other biochemical reactions responsible for changes in color, flavor, texture and nutritional quality (Watada et al., 1999). The reduction of wound-reduced metabolic activity is critical to control these changes. The respiration rate of fresh-cut product is an excellent indicator of metabolic activity of the fresh-cuts to guide the potential storage life. In previous paper, respiration rates of fresh-cut cabbage were investigated in MAP with OPP film, but anaerobic respiration and off-flavor occurred during storage due to beyond the low oxygen limit in MAP with lower permeability of OPP film (Hu et al., 2003). The objectives of this study were to investigate the gas concentrations and the respiration rate of fresh-cut cabbage at initial low O₂ concentrations in PFP and MAP. The effect of initial low O₂ on microbial growth and quality was also determined.

**MATERIALS AND METHODS**

**Plant material**

Cabbages (cv: Akitoku) were obtained from the local supermarket in Fukuoka city, Japan. Samples were transported to the laboratory without refrigeration and immediately processed. Cabbages were selected for uniform size and appearance. Outer and damaged leaves were removed.

**Cabbage slice preparation**

The cabbages were washed with tap water before cutting. Knife and cutting board were washed and treated with 75% ethanol prior to use. The cabbage was cut into 1.5 mm width and 5 mm length by hand with a sharp stainless steel knife. The slices were washed for 6 min with distilled water and then washed with 2°C distilled water for 4 min. The slices were centrifuged for 3 min at 1,720 (rpm) to remove the water on the cabbage surface by washing machine. And then slices were selected at random and samples of
100 ± 1 g were placed in the package (10 × 20 cm) and heat sealed. A silicone cap was applied on each package to permit analysis of package headspace with a syringe. The initial low O₂ concentrations were regulated with 99.99% N₂ gas and the void volume in the package was regulated into 300 ml by 200 ml syringe. All processing operations were conducted at 5–8 °C. The film used for cabbage slices package was oriented polypropylene (OPP) film with thickness 30 μm (Sumitomo Bakelite Co., Ltd, Tokyo Japan). Micro-perforation (0.1 mm in diameter, one hole on each package) was made with needle with 0.1 mm tip. The perforated film packages (PFP) and MAP were stored at 5 and 20 °C for 4 days, respectively. Samples were removed on days 0, 1, 2, 3 and 4 for analyses of gas composition, quality and microbial population.

Gas analysis

Gas sample was withdrawn by gas-tight 1.0 ml syringe from the inside the package. The gas concentrations were measured by injecting 1.0 ml gas sample into gas chromatography (GL Sciences GC–390, Tokyo Japan) equipped with thermal conductivity detector (TCD) and D2000 integrator (Hitachi, Ltd. Tokyo Japan). Helium was used as carrier gas and the flow rate was 30 ml · min⁻¹. The injector and column temperatures were 80 and 50 °C, respectively. The column was WG100 with molecular sieve 5 A and Porapak Q 80/100 mesh.

Mathematical model for respiration

The equations of basic volume balance can be used to determine the respiration rate of fresh-cut cabbage in the packages (Akimoto et al., 1997). The changes in the free volume of gas within package at the short time were calculated (Hu et al., 2003).

Ascorbic acid

Ascorbic acid content was measured by high performance liquid chromatography (HPLC) equipped with Shim-pack SCR–101N column (7.9mm Φ × 30 cm) (Shimadzu Corp., Tokyo Japan). The column is packed with a cation exchanger resin which is sulfonated polystyrene–divinylbenzene copolymer. The mobile phase was mixed solution of 10 mM oxalic acid dihydrate, 15 mM N₂OH and 1 mM EDTA at a flow rate of 1.0 ml min⁻¹. Temperature is 40 °C and detector is RI. Cabbage sample was extracted from excised slices of 5 g cabbage with 5 times the volume of 5% metaphosphoric acid in a mortar with pestle. The homogenate was filtered through filter paper and centrifuged at 3,000 (rpm) for 15 min. The supernatant was taken as sample to measure the ascorbic acid content in cabbage.

Microbial analysis

The Microbial populations were determined by incubating extracts of fresh-cut cabbage on culture medium of potato dextrose agar (PDA) spread with 0.1 ml extraction during 48 h at 30 °C during storage (Babic and Watada, 1996). Colonies were counted and expressed as CFU g⁻¹.

Sensory evaluations

Appearance and flavor were evaluated by five people during storage. Evaluation was
carried out in two items of appearance and flavor. They were both scored on 4-point scale: excellent 1; very good 2; fair 3; poor 4.

**Browning measurement**

A 40 g of sample was weighed, cut and homogenized with 80% ethanol by homogenizer to remove chlorophyll. And then sample was filtered through filter paper. The residue and filter paper was dried at 60°C in the constant temperature drier for 12 h. The residue was placed in a mortar with pestle to make micro-powder. The powder was put on the slide glass 2 mm in thickness and covered with the cover glass. The color of the powder was determined by using a chromameter (model CR 200; Minolta Corp., Japan). A tristimulus colorimeter with an 11 mm aperture and diffuse illumination (C light source) was calibrated with a white standard calibration plate (Y=92.9, x=0.3136, y=0.3200). Expression of color was characterized as Hunter color indexes L*, a* and b*; where L* indicates the lightness, a* means the color axis from green to red, b* the blue-yellow color axis. The changes in the browning of fresh-cut cabbage were shown in following equation (Yano, et al., 1986).

$$\Delta E(\text{Lab}) = \sqrt{(L - L_0)^2 + (a - a_0)^2 + (b - b_0)^2}$$

Where $\Delta E(\text{Lab})$: browning degree; $L_0$, $a_0$, $b_0$: value after cutting cabbage; $L$, $a$, $b$: changing value during storage.

**RESULTS AND DISCUSSION**

**Changes in gas concentration in packages**

Changes in the O$_2$ and CO$_2$ contents in packages at 5°C was shown in Fig. 1. The O$_2$ level in 5% PFP gradually increased to 7.2%, while CO$_2$ level increased to 4.9% by 72 h (Fig. 1A). In 8% PFP, the O$_2$ level was stable in about 8% and CO$_2$ levels was reached 5% (Fig. 1B). For 10% PFP (Fig. 1C), gas composition was increased to 12.3% for O$_2$ and 4.9% for CO$_2$. But O$_2$ was decreased from 21% to 12.5% and CO$_2$ increased from 0.03 to 5.5% in 21%MAP. Among all the samples, highest CO$_2$ content was 5.2% and lowest O$_2$ content was 7.1%, which were within the safe levels from toxicity or anoxia. It was indicated that the perforation on the film played an important role in regulating gas levels based on the relative stable changes in gas concentrations in PFP. For the control, O$_2$ level decreased from 21% to 12.5%, while CO$_2$ increased from 0.03% to 5.5% by 72 h (Fig. 1D). The similar conditions of CO$_2$ and O$_2$ atmospheric were obtained at 5°C by 72 h.

Changes in the O$_2$ and CO$_2$ contents in packages at 20°C was shown in Fig. 2. The remarkable changes in gas compositions were occurred throughout storage. O$_2$ level in 5% PFP decreased to 2.2% within 8 h, then increased more rapidly to 4.2% by 72 h (Fig. 2A). While the CO$_2$ level increased to 12% quickly, and then fluctuated about 13% by hour 72. In 8% PFP, the changes in O$_2$ and CO$_2$ levels were similar to that in 5% PFP (Fig. 2B). In 10% PFP, O$_2$ decreased to 2.2% by 44 h, and then fluctuated in about this level. CO$_2$ increased to 15.2% by 44 h, and then did not change much by 72 h. The O$_2$ decreased from 21% to 1.4% much quickly by 22 h, and then increased slightly. CO$_2$ increased to 14% by 22 h, and then fluctuated about 14.5% (Fig. 2C). Although O$_2$ level decreased to lower than 2.2%, and then increased gradually in all the initial low O$_2$ treatments. It was considered that respiration was suppressed by lower O$_2$ level at initial period, and O$_2$ level
Fig. 1. Changes in gas concentrations in perforated film packages (PFP) with different initial oxygen concentrations and MAP at 5°C.

Fig. 2. Changes in gas concentrations in perforated film packages (PFP) with different initial oxygen concentrations and MAP at 20°C.
was regulated to increase by the hole on the film. For the control, O₂ level decreased at highest speed at initial storage period among all treatments, and then increased slightly due to without getting enough supply through the film (Fig. 2D). CO₂ levels showed similar changing trend in all the samples.

Respiration rate

It has been reported that perforated film package are suitable for high respiration commodities such as strawberry (Chambroy, et al., 1993), mushroom (Lopez-Briones, et al., 1993) and fresh-cut products (Emond, et al., 1991) because of its higher permeability for O₂ and CO₂. In the previous paper, anaerobic respiration was occurred in MAP with OPP film without perforation due to low permeability of film and higher respiration rate of fresh-cut cabbage.

The changes in the respiration rate of fresh-cut cabbage were shown in Fig. 3. The respiration rate decreased slightly at the conditions of increasing O₂ and decreasing CO₂ concentrations at 5 °C by 72 h (Fig. 3A). The respiration rates were decreased at initial 4 h, and then fluctuated in 5–8 CO₂ ml·kg⁻¹·h⁻¹ by the end of experiment. The respiration

![Graph A](image1)

![Graph B](image2)

**Fig. 3.** Respiration rate of fresh-cut cabbage in perforated film packages and MAP at different temperatures (A: 5 °C, B: 20 °C).
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rates remained stable for the duration of the experiment. There was no significant difference among the three levels of initial low O₂. But for control, respiration rate was relative higher than that at initial low O₂ levels.

At 20°C, respiration rate was increased at initial 3 h, and then decreased quickly from 64.6 to 10.3 CO₂ ml⁻¹ kg⁻¹ h⁻¹ by 22 h and slightly increased to 15 CO₂ ml⁻¹ kg⁻¹ h⁻¹ due to anaerobic respiration confirmed by higher respiratory quotient values (4.1) in 21% MAP (Fig. 3B). While in PEP, respiration rates were suppressed markedly with quickly increasing CO₂ and decreasing O₂ levels at initial 10 h, and then fluctuated 5–10 CO₂ ml⁻¹ kg⁻¹ h⁻¹. It was indicated that gas composition in PEP was modified through the low product's respiration and the hole on the package (Fig. 2). It has been reported that the modified atmospheres that best maintain the quality and storage life of fresh-cut products have an O₂ range of 2–8% and CO₂ 5–15%. In this experiment, active modified atmospheric conditions of low O₂ level were established quickly by gas flushing to increase the shelf life and quality of fresh-cut cabbage. The permeability of film was improved by perforated on the package to avoid occurrence of much lower O₂ atmosphere. Therefore, atmospheric conditions were modified through the product's respiration and permeability of perforated film to maintain aerobic respiration in PEP during the storage (Renault, et al., 1994).

The storage life of horticultural products has well been related to the respiration rate of products, the suppression of respiratory metabolism will extend the shelf life of stored fresh product (Lopez-Briones, et al., 1992; Kader, 1986). But O₂ and CO₂ levels beyond its tolerance limits can induce anaerobic respiration and CO₂ damage to result in browning and development of off-flavors. It was suggested that better keeping quality for fresh-cut cabbages could be obtained from low O₂ storage atmosphere above 2.2%. For the effect of initial low O₂, there were no significant differences in changes in gas concentration and respiration rate between 5% PEP and 8% PEP at 5 and 20°C during storage.

MAP can be beneficial in maintaining quality of the fresh-cut product (Gorny, 1997). The suitable gas mixture for MAP has been based on that recommended for the whole commodity (Saltveit, 1997). Fresh-cut products probably can tolerate more extreme levels of O₂ and CO₂ because they do not have as much cuticle or skin to restrict gas diffusion, and the distance of gas diffusion from center to outside of fresh-cut product is much less than that for the whole commodity (King, et al., 1989). It was also reported that an additional benefit of MAP may be attained by actively flushing the package with the desired gas rather than allowing the MAP to develop naturally since the marketing period of fresh-cut product is relatively short (Bai, et al., 2001). In the previous study, fresh-cut cabbage has much higher respiratory activity to consume O₂ and produce CO₂ quickly. Anaerobic respirations were occurred after 3 days of storage at 5°C and after 1 day of storage at 20°C, and lead to browning and development of off-flavors of fresh-cut cabbage (Hu, et al., 2003). Although benefits may occur with modified atmosphere, the extinction point of the fresh-cut products must be recognized to avoid anaerobic respiration (Ko, et al., 1996). In this experiment, relative optimum atmospheric conditions were maintained by regulating initial low O₂ and perforation on the package.

Fresh-cuts generally are much more perishable than intact products because they have been subjected to severe physical stress, such as peeling, cutting, slicing, shredding, trimming and removal of protective cells. Consequently, fresh-cuts probably should be
held at a lower temperature than that recommended for intact commodities. Temperature of 0°C is in most case preferable. However, this is in most cases economically not achievable. Temperature between 5 and 10°C is more commonly found in practice. In this experiment, respiration rate were maintained relative lower levels at 5°C than that at 20°C. The temperature effects on the tissue metabolism and biochemical reactions significantly, it is also of major important to maintain low temperature to prevent microbial growth. It suggests that it is important for handling and storing fresh-cut cabbage at lower temperature or near 0°C (Gorny, 1997), if the product is not sensitive to chilling injury.

**Ascorbic acid**

Fig. 4 shows loss in total ascorbic acid (TAA) contents at different initial O₂ concentrations for fresh-cut cabbage at the end of storage. In 5% and 8% PFP, losses in TAA were similar, that is 9.2 and 9.3%, respectively. It was indicated that initial O₂ 5% and 8% have same effects on suppressing ascorbic acid (AA) oxidation at 5°C. While in 100% PFP and 210% MAP, the rates of loss in TAA were 11.2 and 22.1%, respectively. It demonstrated that initial low O₂ not only suppressed the respiration, but also significantly inhibited activity of ascorbate oxidase to avoid the AA oxidation.

![Graph showing loss in total ascorbic acid in perforated film packages and MAP at different temperatures.](image)

Fig. 4. Loss in total ascorbic acid in perforated film packages and MAP at different temperatures.

It could be also assumed that the role of AA as a reducing power in respect to *de novo* synthesis of polyphenols is possible. Ascorbic acid plays an important role in the defence mechanism against free radicals that induce peroxidation (Barth, *et al.*, 1991). AA can be easily oxidised to dehydroascorbic acid, the latter in turn can either be reduced.

**Browning**

The changes in browning ($\Delta E(\text{Lab})$) of fresh-cut cabbage were shown in Fig. 5. The
browning degree increased to 5.2 slightly with relative higher initial value in 21% MAP at 5°C (Fig. 5A). While in PFP, the browning increased slowly throughout storage. Among the initial low O₂ treatments, 10% PFP had highest value (4.2), but they were lower (2.5–4.2) than that in 21% MAP (5.2). At 20°C (Fig. 5B), the browning degree decreased slightly by 3 days and then increased to 22.5 quickly in 21% MAP. While in PFP, the browning increased slowly throughout storage. Among the initial low O₂ treatments, 10% PFP had highest value (13.2), but they were lower (7.1–13.2) than that in 21% MAP (13.2). It indicated that browning degree was suppressed markedly in initial low O₂ levels. Moreover, the browning was much lower at 5°C than that at 20°C. Low temperature is important essential for storing and maintaining quality of fresh-cut vegetables.

**Sensory evaluation**

Sensory evaluation of fresh-cut cabbage was shown in Table 1. Visual appearances and flavor were not changed much during storage of 4 days in all treatments at 5°C. At 20°C, off-flavor was occurred after 1 day and then not change much in all treatments. But for the control, flavor was reached 3-grade by 4 days. The appearances were reached
Table 1. Sensory evaluation of fresh-cut cabbage at different storage conditions

<table>
<thead>
<tr>
<th>Items</th>
<th>Temperature (°C)</th>
<th>Initial O₂ concentration (%)</th>
<th>Storage time (day)</th>
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<td>1</td>
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<tr>
<td>Flavor</td>
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<td>1</td>
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<td>8</td>
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<td>21</td>
<td>1</td>
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<td>Flavor</td>
<td>20°C</td>
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<td>21</td>
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<td>Appearance</td>
<td>5°C</td>
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<td>Appearance</td>
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<td>21</td>
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2 grades after 3 days. It was explained that the changes in appearance and flavor were due to relative higher activity of physiological metabolism of fresh-cut cabbage at 20°C.

**Microbial analysis**

The effects of initial low O₂ levels on the microbial counts on the surface of fresh-cut cabbage were shown in Fig. 6. At 5°C, bacterial counts increased from $4.6 \times 10^5$ to $1.1 \times 10^6$ CFU g⁻¹ in 21% MAP by 2 days, and reached $2.4 \times 10^6$ CFU g⁻¹ by 4 days. While bacterial counts increased from $4.6 \times 10^5$ to $8.4 \times 10^5$ CFU g⁻¹ in 5% PFP, $4.6 \times 10^5$ to $9.0 \times 10^5$ CFU g⁻¹ in 8% PFP by day 2, respectively (Fig. 6A). After the end of 4-day experiment, bacterial counts were about $1.8 \times 10^6$ CFU g⁻¹ in both PFP. It was indicated that the effect of initial low O₂ on the proliferation and growth of microorganisms was not obvious at 5°C due to remarkable suppression of the microbial proliferation by low temperature. At 20°C, bacterial counts increased from $4.6 \times 10^5$ to $1.4 \times 10^7$ CFU g⁻¹ by 2 days and reached $2.5 \times 10^7$ CFU g⁻¹ by 4 days in 21% MAP. While bacterial counts increased from $4.6 \times 10^5$ to $1.1 \times 10^6$ CFU g⁻¹ in 5% PFP, and then was $3.1 \times 10^6$ CFU g⁻¹ by 4 days. In 8% PFP, from initial value $4.6 \times 10^5$ to $5.5 \times 10^6$ CFU g⁻¹ by 2 days, and then increased to $5.8 \times 10^6$ CFU g⁻¹ by 4 days (Fig. 6B). Microbial analysis showed that the average total plate count on the surface of fresh-cut cabbage was increased during storage, especially at 20°C. The storage temperature has significant effect on the increase in the total plate count. There was lower increasing rate of the total plate count at 5°C than that at 20°C, and the total plate count was lower in PFP than that in MAP.

The microbial populations in PFP were significantly lower than that in MAP. Microbial populations increased most rapidly at the higher storage temperature (Fig. 6B). It was indicated that low levels of O₂ in packages inhibited the growth of aerobic microor-
ganisms. Although the degrees to which CO$_2$ and O$_2$ were modified in PFP and MAP in this study were not enough to inhibit microbial growth completely, the microbial populations were much lower in initial low O$_2$ modifications than that in MAP. One possible explanation for our finding is that the initial low O$_2$ modifications of PFP had an additive retarding effect on microbial growth, and showed the significant effect at higher temperature.

McGill et al. (1966) found that spinach leaves packages in 9.5% CO$_2$ and 3.3% O$_2$ had lower microbial counts than samples packaged in air. Similarly, Babic and Watada (1996) reported that low O$_2$ combined with high CO$_2$ reduced the number of microorganisms on fresh-cut spinach by 10–fold or even 100–fold compared with air at 5°C due to probably decreased oxygen availability. In this experiment, initial low O$_2$ (active MAP) could be used to control microbial development on fresh-cut cabbage at 5°C. Bai et al. (2001) reported that the desired atmosphere was obtained by the active MAP system with low O$_2$ level for retaining shelf life of fresh-cut cantaloupe. Qi et al. (1999) reported that lower microbial population and longer shelf life of honeydew cubes stored at 5°C rather than at 10°C in both air and controlled atmosphere. It indicated that low storage temperature, especially constant low temperature was important for the extending shelf life of
CONCLUSIONS

The effects of perforated film package on the gas concentrations and respiration rates of fresh-cut cabbage with initial O₂ 5, 8, 10 and 21% were studied at 5 and 20°C. The microbial growth and quality were also evaluated. The respiration rates of fresh-cut cabbages were suppressed under an initial low O₂ during storage of 4 days, especially in 5 and 8% O₂ concentrations. The anaerobic respiration did not occur under low O₂ levels. It was also found that fresh-cut cabbages had better color retention and quality, reduced respiration rate and microbial population in PFP. Although there was the difference of quality attributes between PFP and MAP, no significant difference was found.

The effects of initial low O₂ 5% was the most remarkable among different levels of initial low O₂ according to the oxidation of ascorbic acid (AA) and browning of fresh-cut cabbage. Microbial analysis also showed that total count on the surface of fresh-cut cabbage was lowest among initial low O₂ treatments. Total ascorbic acid (TAA) decreased by AA oxidation after cutting. Loss in TAA was much lower in PFP than that in MAP. Moreover, The results of sensory evaluation showed that there were no differences at 5°C throughout storage. At 20°C, flavor reached 2-grade in all treatments and become 3-grade by 4 days in 21%MAP. The appearances were reached 2-grades after 3 days in 10%PFP and 21%MAP. These suggest that better quality of fresh-cut cabbage could be obtained from the combination of PFP with initial O₂ 5–8% level and 5°C.

REFERENCES

Kader, A. A. 1986 Biochemical and physiological basis for effects of controlled and modified
atmospheres on fruits and vegetables. *Food Technol.*, 5: 99–104


