Research Article

Effects of Freeze Vacuum Drying Combined with Hot Air Drying on the Sensory Quality, Active Components, Moisture Mobility, Odors, and Microstructure of Kiwifruits

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In this study, freeze vacuum drying (FVD), hot air drying (AD), and FVD combined with AD (FVD-AD) were used to dry kiwifruits. Dried products were analyzed comprehensively on their sensory quality, active components, moisture mobility, odors, and microstructure. Results showed that the FVD-AD saved time by 38.22% compared with FVD while maintaining an acceptable product quality. The antioxidant properties of FVD-AD were lower than those of FVD but significantly higher than those of AD. Moreover, compared with FVD products, FVD-AD products were moderately hard (5252.71 ± 33.53 g) and improved in color, bound water, and microstructure. Additionally, FVD-AD consumed lesser drying time and energy than FD. According to cluster analysis, the odors of FVD-AD products were similar to those of the fresh ones. Principal component analysis of physicochemical and drying cost indicated that FVD-AD was a promising processing technique for functional kiwifruit snacks.

1. Introduction

Originated in Asia, kiwifruits became popular worldwide due to its sensory and nutritional properties, such as a high level of fibers and bioactive compounds with antioxidant activity. However, kiwifruits were prone to a series of biotic and abiotic stresses, which caused physiological and biochemical changes, leading to fruit quality deterioration, nutrient loss, and decay. For spoilage prevention, chemical reactions must be reduced and shelf life extended.

Hot air drying (AD) was the most common drying method in food production because of its affordability. However, long drying periods often result in inferior product quality due to nutrient losses [1]. Meanwhile, freeze vacuum drying (FVD) was another drying method to remove water from a frozen solution by sublimation under reduced pressure [2], giving rise to high-quality dried products [3]. The main difference with respect to hot air drying was that water was not removed by evaporation but by sublimation from a completely frozen product. The pore structure produced by freeze vacuum drying (FVD) products contributed to brittle texture and rapid rehydration [4]. Therefore, developing FVD technique can reduce the production cost without compromising product quality was necessary [5]. The sublimation drying was the key step in the FVD process, and it affected the quality of the products directly [6]. However, the desorption process took up nearly half the time of the whole process although it can only...
remove a little water. In order to guarantee the quality of the products and reduce drying time, many different combinations of drying methods have been recently used to dehydrate fruits and vegetables and avoid the disadvantages of freeze vacuum drying method, such as high energy consumption, low drying efficiency, and high cost. For example, to achieve the same moisture content in products, the production cost of freeze drying can be 200–500% higher than that of hot air drying [5]. Combining hot air and freeze vacuum drying in fruits and vegetables has been reported to improve product quality in the form of better aroma, faster and better rehydration, considerable savings in energy, and much shorter drying times, compared with hot air drying [7]. Donsi et al. [8] dried apples, potatoes, and carrots by freeze vacuum drying (FVD) combined with AD (FVD-AD), which can potentially obtain high-quality dehydrated fruits and vegetables.

2.2. Drying Methods. Kiwifruits (5 kg) were subjected to three different drying methods, respectively, until the moisture contents were below 4% (dry weight).

2.2.1. Freeze Vacuum Drying (FVD). The FVD process was performed using a laboratory-scale freeze dryer (Scientz-18ND, Martin Christ, Germany). Kiwifruits were spread uniformly on the metal trays and then frozen at −40°C followed by freeze drying for 42 h under the condition of absolute pressure of 20–40 Pa, chamber temperature of 20°C, and condenser temperature of −60°C.

2.2.2. Hot Air Drying (AD). Kiwifruits were spread uniformly on the metal trays and dried using a laboratory electric oven dryer (DHG-9123A, Shanghai Jinghong Test Equipment Co., Ltd., Shanghai, China). The AD processing parameters were set as air temperature of 70°C and air velocity of 0.1 m/s for 24 h, and the limit maximum internal temperature of samples was 60°C.

2.2.3. Freeze Vacuum Drying-Hot Air Drying (FVD-AD). Samples were first dried by FVD for 10 h (the moisture content of the samples was 30 ± 5%) and then processed by AD for 2 h (the moisture content was <4%).

2.3. Determination of Specific Energy Requirements, Color, Texture, Moisture Content, and Water Activity (a_w). The energy required for drying each kilogram of kiwifruit was calculated by

\[ E_{kg} = \frac{E_t}{W_0} \times 1000, \]  

where \( E_{kg} \) is the specific energy requirement (kWh), \( E_t \) is the total energy consumed (kWh), and \( W_0 \) is the initial weight of the kiwifruit sample (g).

A color meter (Model CR-400, Konica Minolta Holdings, Inc. China) was used to measure the color of the kiwifruits, with a white reference tile used for calibration. The hunter-Lab units \( L^∗, a^∗, \) and \( b^∗ \) were used to define the colors, which were expressed in terms of hue angle (°). Color changes between the fresh and dried samples (\( \Delta E \)) were measured 5 times.

A texture analyzer (TA.XT 2i/50, Stable Micro System Ltd., Surrey, UK) with a P/2 probe was used for test analysis. The probe was used to measure the maximum force required to penetrate an individual rehydrated piece of kiwifruit, to a depth of 30%, positioned horizontally over a heavy duty platform, with 5, 1, and 5 mm/s as the prespeed, test-speed, and postspeed settings, respectively. For each experiment, the hardness (g) was recorded.

Moisture content was determined by the oven method. The water activity of the samples was measured using a water
activity meter (LabMaster-aw neo, Novasina, Switzerland) at a constant temperature of 20 ± 1°C. A total of three readings were made for each sample.

2.4. Determination of Ascorbic Acid Content (AA). The amount of AA was determined by means of a test, using Folin–Ciocalteu reagent, derived from what was first proposed by Jagota and Dani [16]. The results were reported as milligrams of AA (mgAA) per 100 g of dried matter.

2.5. Determination of the Total Flavonoid Content (TFC) and DPPH Radical-Scavenging Assay. TFC was measured as described by An et al. [17], with minor modifications. Extract of kiwifruit by ethanol (1 mL) and rutin standard solutions were mixed, respectively, with 1 mL of 5% NaNO2, and the total volume was made up to 12.5 mL with 70% ethanol and then allowed to mix for 5 min. After that, 1 mL of 10% Al(NO3)3 was added and allowed to stand for 6 min. The further 5 mL of 1 mol/L NaOH was added, and the total volume was made up to 25 mL with 70% ethanol. The absorbance of the reaction mixture was read at 510 nm.

DPPH radical-scavenging ability of the kiwifruit extracts was determined using previously described studies [18], with modifications. Briefly, 3.0 mL of 0.065 mmol/L DPPH and 0.5 mL of 70% ethanol were mixed. Then, 3.0 mL of 0.065 mmol/L DPPH and 0.5 mL of extract of kiwifruit by ethanol were mixed. Finally, 3.0 mL of 70% ethanol and 0.5 mL of extract of kiwifruit by ethanol were mixed, and after standing for 30 min, the absorbance of the mixture was measured at 517 nm.

2.6. Low-Field Nuclear Magnetic Resonance (LF-NMR) Measurements. The relaxation times using NMR (Niumag Co., Ltd., Shanghai, China) were evaluated to perform LF-NMR measurements. Approximately 4 g of samples were placed in cylindrical glass tubes (2 cm diameter, 4 cm high). The glass tubes with samples were placed into a CTHI-100 B/150 B/250B box at 25°C (STIK Co., Ltd., Shanghai, China). Transverse relaxation times (T2) were measured using the Carr–Purcell–Meiboom–Gill (CPMG) pulse sequence. The parameters of CPMG were as follows: corresponding resonance frequency (SF) for protons, 20 MHz; spectral width (SW), 100 kHz; echo time (TE), 0.250 ms; pulse widths at 90° (P1) and 180° (P2), 8.0 and 15.52 μs, respectively; waiting time (TW), 800 ms; radio frequency delay time (RFD), 0.08 ms; analog gain (RG1), 20 db; and digital gain (DRG1), set 3. Data from 3,000 echoes were acquired as 32 scan repetitions.

2.7. Electronic Nose (E-Nose) Analysis. The E-nose system composed of a measuring chamber with 18 metal oxide sensors (Alpha M.O.S., FOX4000, France). This equipment was used to distinguish the differences on the aroma profile of kiwifruits. Approximately 2 g of samples were placed into a 5 mL glass vessel that was immediately sealed by a metal screw cap. Firstly, the samples were balanced after 300 s of equilibration at 60°C, and the headspace reached a steady state. The injection speed was 150 mL/min to ensure an injection period of 1 s. Filtered and dried air (purity > 99.999%) with a flow rate of 150 mL/min was used as a carrier gas for E-nose detection. The data acquisition lasted for 120 s, and system rebalance needs 300 s. For each sample, the E-nose detection was performed three times under the same condition.

2.8. Microstructural Analysis. Microstructures of the longitudinal section of FVD, AD, and FVD-AD kiwifruit samples were acquired using a field emission scanning electron microscope (FE-SEM) (SU8010, Hitachi, Tokyo, Japan) at an accelerating voltage of 1.0 kV. The SEM micrographs were obtained at 30× and 100× magnifications.

2.9. Experimental Design and Statistical Analysis. The experiment was designed as a randomized complete block, and individual kiwifruits were the experimental unit for quality. Figure 1 provides an overview of the order of preparation and testing. All experiments were run at least three times, and the data were expressed as mean ± standard deviation (SD). Statistical analyses were performed by Microcal Origin 9.0 software (Microcal Software, Inc., Northampton, USA). Analysis of variance (ANOVA) and Duncan’s multiple-range test (p < 0.05) were used to evaluate the differences among the samples. XLSTAT-Excel was used to perform principal component analysis (PCA).

3. Results and Discussion

3.1. Effect of the Drying Method on Color Parameters. The fruit color can be easily changed during the drying process; this change was one of the negative quality attributes that affect customers’ perceptions of dried products [19]. In kiwifruits, significant differences in L*, a*, and b* values between the fresh and dehydrated samples were observed (p < 0.05) (Table 1). Fresh kiwifruits were characterized by high luminosity, with a tendency to green and yellow. Regarding a* (redness) and b* (yellowness), higher L* and lower a* values were found in samples dried by FVD and FVD-AD, as compared with those samples dried by AD. Similarly, Lin et al. [20] observed a higher L* value for carrot slices under FVD than that for those under AD. However, the b* value of dried kiwifruits was higher than that of fresh. The color of vegetables was determined by natural color compounds that can be oxidized during the drying treatment, and the important factors accelerating degradation were high temperature and presence of oxygen [21]. The moisture migration rate can reflect the changes of color in drying samples. During drying of kiwifruits, the moisture of samples' surface was evaporated gradually and the internal moisture transferred from the center of the samples to the surface. The color of the dried fruit generally changed because of browning reaction, which was always related to the Maillard reaction [22]. The total color difference (ΔE) demonstrated that the color closely follows fresh kiwifruits. Overall, drying using FVD-AD resulted in the lowest total color change (ΔE), while the highest total color (ΔE) change
of kiwifruit slices was observed during AD. This was clearly depicted in the images of the fresh and dried samples (Figure 2). As observed from the image, the color of samples dried using FVD-AD looked very similar to that of the fresh sample, whereas the color of samples dried using FVD and AD was sharply different from that of the fresh sample. This may be due to the vacuum environment during FVD and FVD-AD. However, oxidative reactions and enzymatic browning possibly occurred during AD process, making the products turn brown easily [23]. The $h^\circ$ value showed significant differences ($p < 0.05$) among fresh and dried kiwifruits. The fresh and FVD-AD kiwifruits were close to kelly, and FVD and AD were close to orange. Hence, the color of FVD-AD was the closest to fresh kiwifruits.

### 3.2. Effect of the Drying Process on Drying Time, Hardness, Moisture Content, and Water Activity

The moisture content and water activity of kiwifruits manufactured with different drying methods are shown in Table 2. Significant differences were observed among drying time, hardness, moisture content, and water activity of the selected drying methods ($p < 0.05$). The initial moisture content of fresh kiwifruit was at 87.55% ± 1.64%, dry basis. Finally, the moisture content of kiwifruit was about 3.00%. The $a_w$ values of AD, FVD, and FVD-AD kiwifruits were low, ranging among 0.3048 and 0.3335 compared with fresh samples (0.9927). If the $a_w$ value was below 0.60, then the growth of microbes and lipid oxidation in food products were totally restrained during storage [24, 25]. As shown in Table 2, FVD-AD significantly reduced the drying time by 38.22% ($p < 0.05$), compared with the FVD alone; FVD-AD also consumed lesser energy than in FVD. This result could be due to the slower vaporization of moisture in FD than in AD. Mass transfer within the kiwifruits should be rapid during AD because a large vapor pressure was produced by air on the surface of

### Table 1: Color parameters of kiwifruits samples dried by different drying methods.

<table>
<thead>
<tr>
<th>Drying method</th>
<th>$L^*$</th>
<th>$a^*$</th>
<th>$b^*$</th>
<th>$\Delta E$</th>
<th>$h^\circ$</th>
</tr>
</thead>
<tbody>
<tr>
<td>FVD</td>
<td>62.45 ± 1.42&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1.34 ± 0.22&lt;sup&gt;c&lt;/sup&gt;</td>
<td>31.54 ± 1.49&lt;sup&gt;b&lt;/sup&gt;</td>
<td>10.27 ± 0.25&lt;sup&gt;b&lt;/sup&gt;</td>
<td>82.71 ± 1.54&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>AD</td>
<td>55.88 ± 0.35&lt;sup&gt;c&lt;/sup&gt;</td>
<td>3.05 ± 0.18&lt;sup&gt;a&lt;/sup&gt;</td>
<td>38.73 ± 1.36&lt;sup&gt;a&lt;/sup&gt;</td>
<td>12.28 ± 0.52&lt;sup&gt;a&lt;/sup&gt;</td>
<td>76.00 ± 1.08&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td>FVD-AD</td>
<td>60.30 ± 1.17&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.97 ± 0.20&lt;sup&gt;d&lt;/sup&gt;</td>
<td>28.64 ± 0.79&lt;sup&gt;b&lt;/sup&gt;</td>
<td>6.55 ± 0.14&lt;sup&gt;c&lt;/sup&gt;</td>
<td>88.16 ± 1.12&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Fresh</td>
<td>67.62 ± 0.51&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.42 ± 0.32&lt;sup&gt;b&lt;/sup&gt;</td>
<td>24.36 ± 0.21&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0</td>
<td>97.30 ± 2.24&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

Note: Data are shown as the mean ± SD ($n = 3$). Means within a column with the same letter are not significantly different as indicated by Duncan’s multiple-range test ($p < 0.05$). FVD: freeze vacuum drying; AD: air drying; FVD-AD: combined drying consisting of freeze vacuum drying and air drying.
kiwifruits. The hardness of AD products was significantly higher \((p < 0.05)\) than of FVD and FVD-AD, whereas the FVD-AD products were slightly hard at 5252.71 ± 33.53g, which was in between the values for FVD and AD kiwifruits. This finding was consistent with the results of moisture content and water activity. Several earlier reported samples were produced using AD and characterized by greater hardness \([26, 27]\).

### 3.3. Effect of Drying Methods on the Antioxidant Components

Table 3 shows the comparison of the active composition analysis in the samples subjected to different drying methods with that in the fresh samples. Clearly, significant differences were noted among kiwifruits \((p < 0.05)\). The active component contents of AA, TFC, and DPPH decreased significantly \((p < 0.05)\) upon drying, and FVD samples showed the highest values, followed by FVD-AD samples. The reducing
power could be attributed mainly to the bioactive compounds associated with antioxidant activity [28]. The contents of phenolic compounds in the tested samples may be affected by the losses due to the drying process and oxidation and also by the de novo synthesis. These results could be due to the influences of temperature and oxidation [29, 30]. Low-temperature processing caused minor effect on the drop of active component contents. However, high-temperature treatment led to tremendous decrease in the content of active components [31]. This was somehow consistent with the previous reports [32]. The low-oxygen and temperature environment during FVD and FVD-AD could effectively decrease the wastages of AA and flavonoids. However, compared with that in FVD treatment, FVD-AD treatment exposed more oxygen to kiwifruit samples. Moreover, in case of AD, temperature of samples increased rapidly due to the presence of an electric heater as the sole source for providing energy. Therefore, AA and TFC of kiwifruits significantly decreased in AD and FVD-AD [28].

3.4. Free and Bound Water. To evaluate the water content of samples, the $T_2$ relaxation curves of EWP after multi-exponential fitting analysis with the CPMG sequence are presented in Figure 3. Generally, the relaxation time $T_2$ of water was closely related to its molecular dynamics. Water with higher degrees of freedom corresponds to longer $T_2$ and vice versa. Consequently, different transverse relaxation time $T_2$ components represented water in different mobile states in ginger samples rich in water [33]. All samples showed four peaks, and these peak times are identified as $T_{21}$, $T_{22}$, and $T_{23}$, according to research. The signals in the range of 0–10 ms (I, $T_{21}$) indicated the bound water, which was closely integrated with polar groups of molecules; those in the range of 10–100 ms (II, $T_{22}$) reflected the immobilized water; and those within the limits of 100–1000 ms (III, $T_{23}$) represented the free water [34]. This allowed the separate observation on raw kiwifruit tissue of extracellular space, cell wall, cytoplasm, and vacuole, together with their modifications upon technological treatments [35]. As shown in Table 4, in raw material, water was found to be distributed as follows: $86.15 \pm 2.37\%$ in vacuole ($T_{23}$, 1072.26 ± 14.12 ms), 9.98 ± 0.26% ($T_{22}$ 75.64 ± 0.31 ms) in cytoplasm/extracellular space, and 3.88 ± 0.36% ($T_{21}$ 1.52 ± 0.02 ms, 8.11 ± 0.02 ms) was ascribed to the structural water of the cell wall. The significance ($p < 0.05$) was surveyed in $A_{21}$, $A_{22}$, and $A_{23}$ between the fresh and the dried samples. The transverse relaxation time of free water and immobilized water reduced significantly in the dried samples (Figure 2), indicating that their mobility decreased gradually and drying treatment restricted water mobility. The $T_{21}$ fractions representing bound water of AD samples accounted for $84.398\% \pm 1.30\%$ of the total signal, which was higher than the other samples, followed by FVD-AD samples (75.494% ± 0.42%) and last, the FVD samples. This could indicate that tonoplast integrity was compromised increasing the exchange of water between cell vacuole and extracellular/cytoplasm volumes in the drying process [36]. This was in accordance with the difference in the amount of the NMR signal (or in other words, in the amount of water) associated to the peaks.

3.5. E-Nose Analysis. In order to legibly describe the aroma profile of kiwifruits, a PCA plot was performed (Figure 4(a)). Electronic nose combined with PCA can be used to distinguish the flavor characteristics of kiwifruit according to different drying methods. The discrimination index was 92, and the score plots of kiwifruit showed that the first principal component (PC 1) accounted for 99.196%, whereas the second principal component (PC 2) accounted for 0.3573%. The cumulative contribution of the first two PC variances was 99.553%, which was enough to represent variables. According to Figure 4(a), the fresh and FVD-AD samples appeared on the negative electrodes of PC 1 and PC 2, but samples of AD showed the opposite. At the same time, FVD samples appeared on the negative side of PC 1 and the positive side of PC 2. The region of fresh samples was close to the FVD-AD, indicating that possible similarities could be present among these samples.

The raw response data generated by the 18 sensors of the electronic nose were collected and converted into radar graphs as shown in Figure 4(b). Generally speaking, the
Table 4: Effect of drying methods on transverse relaxation time (T<sub>2</sub>) and the percentages for different relaxing components (A<sub>2</sub>).

<table>
<thead>
<tr>
<th>Method</th>
<th>T&lt;sub&gt;21a&lt;/sub&gt; (ms)</th>
<th>T&lt;sub&gt;21b&lt;/sub&gt; (ms)</th>
<th>T&lt;sub&gt;22&lt;/sub&gt; (ms)</th>
<th>T&lt;sub&gt;23&lt;/sub&gt; (ms)</th>
<th>A&lt;sub&gt;21&lt;/sub&gt; (%)</th>
<th>A&lt;sub&gt;22&lt;/sub&gt; (%)</th>
<th>A&lt;sub&gt;23&lt;/sub&gt; (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FVD</td>
<td>0.16 ± 0.01&lt;sup&gt;b&lt;/sup&gt;</td>
<td>9.82 ± 0.01&lt;sup&gt;a&lt;/sup&gt;</td>
<td>65.79 ± 3.98&lt;sup&gt;b&lt;/sup&gt;</td>
<td>265.61 ± 3.56&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2402.33 ± 6.53&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1049.33 ± 5.48&lt;sup&gt;a&lt;/sup&gt;</td>
<td>606.19 ± 5.48&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>AD</td>
<td>0.16 ± 0.01&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2.66 ± 0.28&lt;sup&gt;c&lt;/sup&gt;</td>
<td>16.30 ± 0.46&lt;sup&gt;c&lt;/sup&gt;</td>
<td>100.21 ± 2.46&lt;sup&gt;c&lt;/sup&gt;</td>
<td>3255.34 ± 7.84&lt;sup&gt;a&lt;/sup&gt;</td>
<td>337.14 ± 1.71&lt;sup&gt;c&lt;/sup&gt;</td>
<td>264.66 ± 1.73&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td>FVD-AD</td>
<td>0.14 ± 0.01&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2.66 ± 0.04&lt;sup&gt;c&lt;/sup&gt;</td>
<td>14.16 ± 0.42&lt;sup&gt;c&lt;/sup&gt;</td>
<td>86.98 ± 3.17&lt;sup&gt;d&lt;/sup&gt;</td>
<td>1721.49 ± 10.26&lt;sup&gt;c&lt;/sup&gt;</td>
<td>212.32 ± 8.67&lt;sup&gt;d&lt;/sup&gt;</td>
<td>346.50 ± 2.67&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Fresh</td>
<td>1.52 ± 0.02&lt;sup&gt;a&lt;/sup&gt;</td>
<td>8.11 ± 0.02&lt;sup&gt;b&lt;/sup&gt;</td>
<td>75.64 ± 0.31&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1072.26 ± 14.12&lt;sup&gt;a&lt;/sup&gt;</td>
<td>153.17 ± 2.98&lt;sup&gt;d&lt;/sup&gt;</td>
<td>394.48 ± 4.42&lt;sup&gt;b&lt;/sup&gt;</td>
<td>3405.08 ± 10.42&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

Note: Data are shown as the mean ± SD (n = 3). Means within a column with the same letter are not significantly different as indicated by Duncan’s multiple range test (p < 0.05). FVD: freeze vacuum drying; AD: air drying; FVD-AD: combined drying consisting of freeze vacuum drying and air drying; T<sub>21</sub> refers to the transverse relaxation time of the free water; T<sub>22</sub> and T<sub>23</sub> refer to the transverse relaxation time of the bound water and the immobilized water, respectively. A<sub>21</sub> represents the corresponding water fraction to T<sub>21</sub>, A<sub>22</sub> represents the corresponding water fraction to T<sub>22</sub>, and A<sub>23</sub> represents the corresponding water fraction to T<sub>23</sub>.
similar shapes of these radar graphs implied the similarity of these samples from different drying methods. Results suggested that the fingerprints outlined the fresh samples during the drying period. The relative values of P30/2 (alcohol), P30/1 (solvents), PA/2 (ammonia, amines), and P10/1 (hydrocarbons, methane) of fresh samples were significantly lower than of the dried samples. Moreover, a radar fingerprint chart of fresh samples and FVD-AD samples almost overlapped, indicating that similar volatile ingredients existed in these samples. Pei et al. [37] reported that the odors in button mushroom dehydrated by FVD were significantly lost than those by freeze vacuum combined drying, which was consistent with our result.

3.6. Effect of Drying Methods on Microstructure. Figure 5 shows cross-sectional SEM images of dry kiwifruits that were dried with FVD, AD, and FVD-AD (Figures 5(a)–5(c)). Generally, the microstructural change during kiwifruit dehydration was related to the water migration of cells. This phenomenon caused cell stress and turgor loss, resulting in contraction and structural collapse at different levels. The surface structures of the AD samples were tightly shrunk and flattened, while those of the FVD-AD samples showed less shrinkage, and the bowl-shaped structure of the cell walls was maintained. A similar honeycomb network was observed in the FD sample, which exhibited the least amount of tissue shrinkage or cell collapse. The cell walls of freeze-dried samples appeared to be comparatively smooth and thin, explaining the noncrisp and spongy texture obtained by freeze-dried kiwifruits [38]. These phenomena occurred because the samples of FVD and FVD-AD were in vacuum and could easily generate porous structures. In addition, the FVD samples had the lowest hardnes because the material was frozen during drying: Therefore, it reduced cell damage and had a porous honeycomb structure with weak force resistance [39]. Moreover, the kiwifruits dried by FVD-AD had a more porous and less collapsed structure, compared with those of FVD samples because the FVD-AD sample was heated by hot air and water evaporated more rapidly, resulting in puffing and creating larger pores within the samples [40]. The larger pore size and higher porosity allowed this sample to absorb more water during rehydration.

3.7. PCA. Three drying methods for kiwifruit samples were evaluated by exploratory PCA technology. The physical and chemical properties and oxidation resistance of any cluster were evaluated. The color index was expressed by E, and the drying efficiency index was expressed by drying time. As shown in Figure 6, the cumulative contribution of PC 1 and PC 2 accounts for 93.16% of the total variance (PC1 = 87.34%, PC2 = 8.81%). PC 1 was highly correlated with E (−0.876), A23 (0.897), water activity (0.910), AA (0.968), TFC (0.967), and DPPH (0.962). PC 2 was mainly correlated with TPC (0.541), A23 (−0.438), and aw (−0.412).

The PC 1 and PC 2 scores of fresh samples were considerably the highest among the other drying methods because they had higher contents of AA and TFC, as well as higher antioxidant activity. The factor score of fresh samples was the highest (2.07), followed by FVD-AD samples (−0.81) and FVD samples (−0.90). Meanwhile, the sample of AD showed the lowest factor score (−5.22). Therefore, AD samples were negatively correlated with PC1 and PC2, representing that AD had a negative effect on the active
Figure 5: Scanning electron micrographs of kiwifruit samples dried by FVD (a), AD (b), FVD-AD (c). FVD, freeze vacuum drying; AD, air drying; FVD-AD, combined drying consisting of freeze vacuum drying and air drying.

Figure 6: Principal component analysis plot of data for different sensory quality, antioxidant properties, and drying efficiency of kiwifruit samples. FVD: freeze vacuum drying; AD: air drying; FVD-AD: combined drying consisting of freeze vacuum drying and air drying.
components, antioxidant activity, and color quality. The FVD process was positively correlated with PC1 and PC2, representing that it had a significant effect on the antioxidant activity of kiwifruits. However, the FVD-AD process was negatively correlated with PC2 and positively correlated with PC1 because it had negative effects on water activity and free water and positive effects on TPC, AA, DPPH, and color.

4. Conclusion

The application of FVD-AD in the drying of kiwifruit was evaluated. Compared with the AD method, the FVD-AD method yielded more active component contents of AA, TFC, and antioxidant properties. The color of FVD-AD product was close to that of fresh kiwifruit. According to cluster analysis, it was proved that the volatile compounds of FVD-AD products were more similar with those of fresh ones compared to FVD and AD products. In addition, the application of FVD-AD contributed to form porous microstructure in dried kiwifruits. The total drying time of FVD-AD was 38.22% lower than that of FVD, and the bound water of FVD-AD was more than that of FVD. From the perspectives of color, energy consumption, moisture, odors, and microstructure, FVD-AD was the most promising processing method for functional kiwifruit snacks.

Data Availability

The data used to support the findings of this study are included within the article.

Conflicts of Interest

The authors declare that there are no conflicts of interest regarding the publication of this paper.

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