Qualities of Cookie Made With Beeswax-Coconut Oil Organogels as Replacement for Shortening

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Abstract The effects of beeswax-coconut oil organogels (7.5%, 10.0%, 12.5%, 15.0% and 17.5% beeswax, wt) quality-related characteristics of shortening replaced cookies were evaluated. The following quality attributes were measured: organogels melting points, fatty acids contents, proximate composition, color, texture, spread factor, sensory evaluation, antioxidant activity and storage stability. Peroxide values of cookie extracts made with organogels were dramatically lower than that of cookie made with shortening (p<0.05) which had a value of more than 10 meq/kg lipids, after 12 days storage at 60°C and strong oil rancid odor. All cookies made with organogels stored at 60°C for 20 days had peroxide values lower than 1.5 meq/kg lipids and no rancid odor, indicating the cookie was still in low oxidation condition. No significant differences were found in cookie dough texture, proximate composition, spread factor, reducing power. The melting point, texture profile of hardness and adhesiveness increased significantly with beeswax ratio increase in organogel. The total phenolic content of cookie extracts increases slightly (p>0.05), nevertheless, the antioxidant activity of cookie extracts with light or without light exposure for 2 hours decreases slightly. No significant differences in sensory attributes of cookies made with shortening and 12.5% beeswax-coconut oil organogel except crisp were revealed by 72 panelists. For cookie made with coconut oil, the oil separate out problem after baking was solved by adding more than 12.5% beeswax in organogel. Accordingly, cookies made with 12.5% beeswax-coconut oil organogel is considered the most acceptable and appropriate practice in processing cookies for commercial purposes of shortening replacement.

Keywords: Coconut oil, beeswax, organogel, cookie, antioxidation


1. Introduction

Epidemiologic study demonstrated that the consumption of high amounts of saturated fat and cholesterol induces high blood cholesterol [1]. Coconuts have served mankind as important foods for thousands of years. Coconut oil contains mostly (>90%) of saturated fatty acids and it has received bad reputation to be hypercholesterolemic in action [2,3]. Clinical studies have revealed the consumption of coconut was not associated with cardiovascular disease and heart attacks [4]. Virgin coconut oil (VCO), which refers to coconut oil produced from coconut milk by natural or mechanical method (wet method). VCO is increasing in popularity as functional oil [3]. The VCO is rich in medium chain fatty acids which are associated with increase in the serum triacylglycerol but incorporation of structured lipid and other functional substances improving the lipid profile [5]. Nevin and Rajamohan [6] found low density lipoprotein (LDL) cholesterol in VCO fed animals was decrease and high density lipoprotein (HDL) cholesterol was increase. Although, only trace amount α-tocopherol in VCO, which contained high amount of total phenolics [7] and VCO was shown superior in antioxidant action [2].

Shortening prevents the formation of gluten during pastry, cake and cookie dough mixing. It also gives the tender texture and mouthfeel of baked products [8]. Shortening also imparts other functional characteristics such as aeration, stability, positively contributing to the geometry and structure of the baked products [9]. A great deal of effort has been made to reduce the use of shortening due to the possible presence of trans fatty acids and a high level of saturated fatty acids with the recent healthy trend [9]. It is possible to apply virgin coconut oil (VCO) to baked products instead of shortening, however, the use of VCO might produce baked products with more greasy and less crispy characteristic because its low melting point.

Organogels are defined as 3-dimensional networks of an organic phase produced by adding some organogelators into the liquid phase. Organogels have some advantages such as not changing the fatty acid composition; therefore, no trans and saturated fatty acids are produced [10,11,12]. The organogels were stable against oxidation during storage and their melting point was higher [13]. Coconut oil organogels might be used as margarine-like or shortening foot stock. Beeswax is regarded as a food additive approved by the US FDA. Through organolation with beeswax, coconut oil can be entrapped in organogelator and produced a solid-like gel network, organogel.
In this study, the oleogels of virgin coconut oil with beeswax were prepared and they were incorporated into the formulation of cookies as a shortening replacer. The effects of organogels on the physicochemical properties of the cookies were investigated. The effect of the amount of beeswax supplement on the dough texture, baked cookie, cookie storage stability, antioxidant ability and consumer acceptance was evaluated.

2. Materials and Methods

2.1. Raw Materials and Organogel Preparation of Coconut Oil with Beeswax

Virgin coconut oil was obtained from Bakersking International Corp (Taipei, Taiwan). Beeswax was purchased from Bee-World Company (Taoyuan city, Taiwan). Sucrose, brown sucrose, nonfat dry milk, high-fructose corn syrup, and salt were purchased from a local market. Cake flour used in this study was purchased from the Cha Hwa corporation (Taichung, Taiwan). Sodium bicarbonate, all-purpose shortening, food grade sodium bicarbonate and ammonium bicarbonate were obtained from a local store (Fu Shen baking ingredient store, Keelung, Taiwan). Kjeldahl catalyst tablets, sodium hydroxyl, boric acid, sulfuric acid, methyl red, 1,1-diphenyl-2-picrylhydrazyl, ferrous chloride 4-hydrate, and ferrozine (3-(2-pyridyl)-5,6-bis(4-phenylfulfonic acid)-1,2,4-triazine were purchased from Panreac Appli Chem (Gatersleben, Saxony-Anhalt, Germany). All reagents were obtained from analytical grade. Acetone, methanol, ethanol, trichloracetic acid, and acetic acid were purchased from Sigma Aldrich (St. Louis Missouri, USA). Monobasic sodium phosphate dibasic sodium phosphate, methyl red, and sulfuric acid were purchased from Merck (Whitehouse station, NJ, USA). Ethyl ether was purchased from Nihon Shiayaku industries (Taiwan, R.O.C.).

Virgin coconut oil was replaced with beeswax at 7.5%, 10.0%, 12.5%, 15.0%, and 17.5% levels. The mixtures were heated at 80°C in a water bath to dissolve the beeswax and poured separately into 250 ml beaker and cooled down to room temperature (22.5°C) for 24 hours. Hardness and adhesiveness test was performed following the method of Lima and Guraya [17]. A TA-XT2 Texture Analyzer (Stable Micro Systems Co., Ltd., Haslemere, England) with a cylinder probe (P/25, 25 mm in diameter) was used to compress organogel samples at 10.00 mm/s to 50% target value (trigger load: 0.5 g) from the sample surface and withdrawn at the same speed. The maximum force during compression was recorded. The test was performed in triplicate and the average maximum force is reported as hardness in Newton (N). Adhesiveness is the force required to remove the probe from the organogel.

2.2. Thermal Analysis, Fatty Acid Content and Texture Profile Analysis of Organogels

The method of AOCS [14] was adapted for thermal analysis of organogels. All samples were analyzed with a Mettler Toledo DSC822e Differential Scanning Calorimeter (DSC) (Mettler Toledo, Swiss) according to the method of Dodd and Tonge [15]. Around 10-15 mg of each organogel mixture was weighed into an aluminum pan. The following temperature program was used: heating from 10°C to 80°C by 10°C /min. The pan was purged with dry nitrogen to prevent oxidation of organogel sample at flow rate of 80 ml/min. Lipid extraction was determined by homogenizing 15 g cookie samples with 150 ml chloroform-methanol mixture (2:1; v/v) containing 0.2% butylated hydroxytoluene (BHA) as antioxidant. Fatty acids were esterified into methyl esters according to AOAC [16].

VCO with beeswax at different levels were heated at 80°C in a water bath to dissolve the beeswax and poured separately into 250 ml beaker and cooled down to room temperature (22.5°C) for 24 hours. Hardness and adhesiveness test was performed following the method of Lima and Guraya [17]. A TA-XT2 Texture Analyzer (Stable Micro Systems Co., Ltd., Haslemere, England) with a cylinder probe (P/25, 25 mm in diameter) was used to compress organogel samples at 10.00 mm/s to 50% target value (trigger load: 0.5 g) from the sample surface and withdrawn at the same speed. The maximum force during compression was recorded. The test was performed in triplicate and the average maximum force is reported as hardness in Newton (N). Adhesiveness is the force required to remove the probe from the organogel.

2.3. Cookie Formulation and Preparation

A single-bowl mixing procedure was used for making cookies according to AACC [18] method 10-54, with some modification. Appropriate amounts of dry ingredients (sucrose (20.0 g), brown sugar (16.0 g), salt (2.0 g), sodium bicarbonate (1.6 g) and nonfat dry milk (1.6 g) were weighed and mixed to give a uniform mixture. All-purpose shortening (160.0 g) was added and mixed at low speed for 1 min and scraped down after each minute. The flour (35.2 g) was mixed at low speed for 1 min and scraped down every 20 s. High-fructose corn syrup (2.4 g) and ammonium bicarbonate (0.8 g) was dissolved in the water (35.2 g) and added into the mixing bowl. The shortening in the cookie formula was replaced by 100% coconut oil, and 7.5%, 10.0%, 12.5%, 15.0% and 17.5% levels of beeswax organogels. After the mixing was completed, the dough was removed from the mixing bowl and placed on an ungreased baking sheet between two identical cutting boards, and then rolled by rolling pin to desired thickness of 2 mm. The dough was cut using an aluminum cookie cutter (60 mm diameter). The cut dough sheet was baked at 205°C for 7 min. After baking the cookies were cooled to room temperature and stored in Ziploc bags at refrigerator (7°C) until evaluated.

2.4. Proximate Chemical Composition and Color of Cookie, Texture Profile Analysis of Cookie Dough and Cookie

The proximate chemical composition of cookies was determined according to the Association of Official Analytical Chemists method [16]. Spread factor of cookies was measured from the ratio of average value of diameter and average value of thickness of cookies with some modification (The diameter of ten cookies was measured again after rotating each cookie to 90°C and then the average value of cookie diameter was calculated). Ten cookies were stacked on each other and their thickness was measured [19].

Texture profile analysis of cookie doughs were performed following the method of Tarancón et al. [20]. A TA-XT2 Texture Analyzer with an aluminum cylinder P/25 probe was used to double compress cookie doughs,
which were rolled into 10 mm thickness and cut with cookie cutter (60 mm in diameter). Dough samples were compressed at 10.00 mm/s to 30% depth (trigger load: 0.5 g and waiting time between the two cycles: 5 s) from the dough surface and withdrawn at the same speed. The maximum force during compression was recorded. The test was performed in triplicate and the average maximum force is reported as hardness in Newton (N). Adhesiveness is the force required to remove the probe from the dough. Springiness was expressed as a ratio of the height during the second compression divide by the original height during the first compression.

Hardness test of cookie was performed following the method of Tarancón et al. [20]. A TA-XT2 Texture Analyzer with a round probe (P/25A, 4 mm in diameter) was used to compress cookie samples at 1.00 mm/s to 50% target value (trigger load: 20.0 g and supports apart: 50 mm). The maximum force during compression was recorded. The test was performed in triplicate and the average maximum force is reported as hardness in Newton (N).

The color of the cookie samples were examined with a spectrophotometer (TC-1800 MK II, Tokyo, Japan) using L (lightness), a (redness/greenness) and b (yellowness/blueness) color scale according to the method of Cruz-Romero et al. [21]. Both a white tile and a black cup were examined before the test to standardize the spectrophotometer. The color of the cookie samples was recorded after taking three measurements for each sample, and triplicate determinations were recorded for each treatment. The color difference

$$\Delta E = ((\Delta L)^2 + (\Delta a)^2 + (\Delta b)^2)^{1/2}$$

where $$\Delta L = L_{sample} - L_{control}$$;

$$\Delta a = a_{sample} - a_{control}$$; $$\Delta b = b_{sample} - b_{control}$$.

2.5. Total Phenolics Content, Reducing Power and Antioxidant Activity Measured by β-Carotene Bleaching Method

Cookie samples were milled and screened through a sieve (0.5 mm). The ground cookie samples were stored at -20°C. The ground cookie samples (3 g) were extracted with 22.5 ml of a methanol:acetone:water (1:1:1; v/v/v) using a stirring hot plate (Model PC-420D, Corning, NY, USA) at 1000 rpm for 30 min according to the method of Seczyk et al. [22] with slight modification. The extracts were centrifuged (6800×g) at 4°C for 30 min and extraction procedure was repeated two times. All extracted solutions were combined and stored in darkness at -20°C until analysis.

Total phenolics were evaluated using Folin-Ciocalteau reagent described by Singleton and Rossi [23]. The cookie extracted solution (0.1 ml), water (0.1 ml) and Folin-Ciocalteau reagent (0.4 ml) were mixed, and then 2 ml of sodium carbonate (100 g/1000 ml) was added after 3 min and mixed thoroughly. It was allowed to stand for 30 min and measured the absorbance at 700 nm in a microplate Reader (Model AMR-100, Allsheng Instruments Co., Ltd., Hangzhou City, China). Total phenolics content was calculated as gallic acid equivalents in mg/g of dry weight.

A 1 ml of cookie extracted solution was added to 1 ml of sodium phosphate buffer (0.2 mM, pH 6.6) followed by 1 ml of 1% potassium ferricyanide. The reaction mixture was incubated for 20 min in a water bath at 50°C. After incubation, 1 ml of 10% trichloroacetic acid was added, followed by centrifugation at 1700×g for 10 min at 4°C. The upper layer (1 ml) was mixed with 1 ml distilled water and 0.2 ml of 0.1% ferric chloride. Absorbance of the resulting solution was measured at 700 nm. A reaction mixture containing 125 μl of DI water served as the blank and 125 μl ascorbic acid (500 μM) served as the positive control. A high absorbance was indicative of strong reducing power [24].

Oxidative loss of β-carotene in a β-carotene linoleic acid emulsion was used to assess the antioxidant activity of the examined cookie extracts [25]. β-carotene (1 mg) was dissolved in 10 ml of chloroform and 3 ml of β-carotene solution was mixed with 40 mg of purified linoleic acid and 400 mg of Tween 20 in around-bottom flask. Chloroform was removed by purging with nitrogen. Pure water (100 ml) was added into the β-carotene/linoleic acid emulsion and subjected to Vertex shaking for 60 s. Cookie extracts (0.1 ml) and aliquots (3 ml) of the β-carotene/linoleic acid emulsion were placed in capped culture tubes and mixed thoroughly. The tubes were placed with light exposure and without light exposure for 0, 60, and 120 minutes. Oxidation of β-carotene/linoleic acid emulsion was monitored spectrophotometrically by measuring the absorbance at 465 nm after 0, 60, and 120 minutes (AMR-100, All sheng Instruments Co., Ltd., Hangzhou city, China). A control was prepared using 0.1 ml of 95% ethanol instead of the extract.

Degradation rate (DR) = ln(A₀ / A_sample) / t

The antioxidant activity (%AOA)

$$\frac{[\text{DR}_{blank} - \text{DR}_{sample}]}{\text{DR}_{blank}} \times 100\%$$

Where A₀ is the initial absorbance (465 nm) at time zero, A_sample is the absorbance (465 nm) at time 120 min and t is time (min).

2.6. Cookie Lipid Fraction Oxidation Measurements

The lipids of ground cookies (100 g) were extracted for 60 min with the use of petroleum ether in a laboratory shaker at ambient conditions [26]. After filtration and separation of lipid fraction, solvent was removed by evaporation under reduce pressure on rotary evaporator at 50°C (IKA RV-10 basica, IKA, Germany). The lipids obtained were frozen (-18°C) until further use.

The antioxidant activity of the cookie prepared with coconut oil and beeswax organogel was evaluated by measuring the peroxide value (PV) in the solvent extracts of cookies [14]. The PV was measured by putting the cookie sample (5 g) into an Erlenmeyer flask (250 ml) and adding a glacial acetic acid/isooctane solution (3:2, v/v, 30 ml). After nitrogen gas was flushed into the Erlenmeyer flask, a saturated potassium iodide solution (1 ml) was added. The flask was immediately sealed, shaken gently for 1 min, and allowed to stand in the dark for 5 min.
After distilled (30 ml) was added with vigorous stirring, the solution was titrated with 0.01 N sodium thiosulphate solution, using a 1% starch solution (400 μl) as the indicator.

$$\text{PV (mg/kg lipid) = (V2 - V1) x 0.01 x 1000 / W.}$$

$$V_2 - V_1 = \text{ml of sodium thiosulphate (Na}_2\text{S}_2\text{O}_3)$$

(blank corrected).

Stability of cookies was followed periodically after 0, 4, 8, 12, 16, and 20 days of storage at 60°C, by determining peroxide value (PV) and recording odor change during 20 days according to the method of Mildner-Szkudlarz et al. [27].

## 2.7 Sensory Evaluation

Thirty-two male and forty female undergraduate and graduate students from the Department of Food Science, were instructed to evaluate the appearance, odor, crisp, flavor, coconut flavor and overall acceptability using a seven-point hedonic scale ranging from "1=extremely dislike" to "7=extremely like" according to the method of Sudha et al. [28].

## 2.8 Statistical Analysis

Data was examined with an analysis of variance using the SPSS statistic program for Windows Version 12 (SPSS Inc., Chiago, IL, USA). Duncan’s multiple range test was used to identify the difference between treatments at a 5% significance level (p<0.05). Differences between the means were evaluated using Duncan’s Multiple Range Test. [29]

## 3. Results and Discussion

### 3.1 Thermal Analysis of Organogels and Fatty Acid Contents of Organogels

The melting points of 7.5% and 10.0% beeswax organogels (35.3°C and 35.8°C, respectively) were close to that of shortening (34.3°C) (Table 1). The main fatty acids of cookie made with shortening were palmitic acid (43%) and oleic acid (40%) with 0.4% trans fatty acid (Table 2). However, there were no trans fatty acid was found and the main fatty acids of beeswax-coconut oil organogels were lauric acid (46-48%), myristic acid (18%), palmitic acid (10-12%) with less than 1% of lignoceric acid (Table 2).

### Table 1. Peak temperature and melting point of different concentrations of coconut oil and beeswax organogels

<table>
<thead>
<tr>
<th>Sample</th>
<th>Control</th>
<th>CNO</th>
<th>CNO + BW (7.5%)</th>
<th>CNO + BW (10%)</th>
<th>CNO + BW (12.5%)</th>
<th>CNO + BW (15%)</th>
<th>CNO + BW (17.5%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>T&lt;sub&gt;peak&lt;/sub&gt; (°C)</td>
<td>34.29 ± 2.60°</td>
<td>32.50 ± 0.11°</td>
<td>20.02 ± 0.26°</td>
<td>19.67 ± 2.37°</td>
<td>19.57 ± 1.44°</td>
<td>19.97 ± 1.15°</td>
<td>23.18 ± 1.57°</td>
</tr>
<tr>
<td>Melting point (°C)</td>
<td>34.29 ± 2.60°</td>
<td>25.58 ± 0.14°</td>
<td>35.32 ± 0.32°</td>
<td>35.85 ± 1.16°</td>
<td>36.66 ± 0.43°</td>
<td>37.04 ± 0.42°</td>
<td>39.50 ± 0.93°</td>
</tr>
</tbody>
</table>

Control: commercial shortening; CNO: coconut oil; CNO + BW (7.5%): 7.5% beeswax-coconut oil organogels; CNO + BW (10%): 10% beeswax-coconut oil organogels; CNO + BW (12.5%): 12.5% beeswax-coconut oil organogels; CNO + BW (15%): 15% beeswax-coconut oil organogels; CNO + BW (17.5%): 17.5% beeswax-coconut oil organogels.

### Table 2. Fatty acids contents of commercial shortening, beeswax and organogels

<table>
<thead>
<tr>
<th>Fatty acids (%)</th>
<th>Control</th>
<th>CNO</th>
<th>CNO + BW (7.5%)</th>
<th>CNO + BW (10%)</th>
<th>CNO + BW (12.5%)</th>
<th>CNO + BW (15%)</th>
<th>CNO + BW (17.5%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>6:0</td>
<td>ND*</td>
<td>0.63</td>
<td>0.66</td>
<td>0.65</td>
<td>0.65</td>
<td>0.65</td>
<td>0.63</td>
</tr>
<tr>
<td>8:0</td>
<td>ND*</td>
<td>8.19</td>
<td>8.29</td>
<td>8.11</td>
<td>8.09</td>
<td>8.05</td>
<td>7.94</td>
</tr>
<tr>
<td>10:0</td>
<td>ND*</td>
<td>6.33</td>
<td>6.24</td>
<td>6.14</td>
<td>6.11</td>
<td>6.06</td>
<td>6.00</td>
</tr>
<tr>
<td>12:0</td>
<td>0.21</td>
<td>49.02</td>
<td>47.74</td>
<td>47.3</td>
<td>46.89</td>
<td>46.47</td>
<td>45.99</td>
</tr>
<tr>
<td>14:0</td>
<td>1.12</td>
<td>19.05</td>
<td>18.27</td>
<td>18.20</td>
<td>18.00</td>
<td>17.85</td>
<td>17.67</td>
</tr>
<tr>
<td>16:0</td>
<td>42.98</td>
<td>8.40</td>
<td>9.63</td>
<td>10.23</td>
<td>10.69</td>
<td>11.16</td>
<td>11.76</td>
</tr>
<tr>
<td>9c-16:1</td>
<td>0.16</td>
<td>0.09</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
</tr>
<tr>
<td>17:0</td>
<td>0.10</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
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</tr>
<tr>
<td>18:0</td>
<td>4.40</td>
<td>2.54</td>
<td>3.04</td>
<td>3.01</td>
<td>3.00</td>
<td>3.01</td>
<td>2.99</td>
</tr>
<tr>
<td>11c-18:1</td>
<td>0.71</td>
<td>0.13</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
</tr>
<tr>
<td>9c,12c-18:2</td>
<td>9.31</td>
<td>0.78</td>
<td>0.64</td>
<td>0.68</td>
<td>0.68</td>
<td>0.68</td>
<td>0.67</td>
</tr>
<tr>
<td>9c,12c-18:2</td>
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<td>ND*</td>
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<tr>
<td>9c,12c-18:2</td>
<td>0.17</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
</tr>
<tr>
<td>11c-15c-18:3</td>
<td>0.17</td>
<td>0.28</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
</tr>
<tr>
<td>20:0</td>
<td>0.39</td>
<td>0.35</td>
<td>0.09</td>
<td>0.10</td>
<td>0.10</td>
<td>0.10</td>
<td>0.11</td>
</tr>
<tr>
<td>11c-20:1</td>
<td>0.15</td>
<td>0.68</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
</tr>
<tr>
<td>22:0</td>
<td>0.09</td>
<td>1.21</td>
<td>ND*</td>
<td>0.05</td>
<td>0.06</td>
<td>0.08</td>
<td>0.08</td>
</tr>
<tr>
<td>24:0</td>
<td>0.09</td>
<td>14.67</td>
<td>0.36</td>
<td>0.51</td>
<td>0.62</td>
<td>0.76</td>
<td>0.91</td>
</tr>
<tr>
<td>Trans fatty acids</td>
<td>0.35</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
<td>ND*</td>
</tr>
</tbody>
</table>

ND* = not detectable.
3.2. Texture Properties of the Organogel, Cookie Dough and Cookies

Hardness and adhesiveness of the organogels were measured at room temperature (22.5 °C) and presented in Figure 1. Organogels made with 15% beeswax have shown a significant increase in the hardness and adhesiveness values. The hardness and adhesiveness values of all-purpose shortening (control) were 0.99 N and 0.35 N, respectively. Among all organogel samples, the hardness and adhesiveness of 10% beeswax organogel were close to those of shortening. It showed no differences (p>0.05) compared to the hardness and adhesiveness of shortening and organogels containing less than 12.5% beeswax. Hardness is the force required to compress a material under certain defined conditions. The stabilization of organogel structure is via non-covalent interactions like van der Waals interactions hydrogen bonding and others [12]. Beeswax contains straight-chain acids with carbon skeletons of up to 36, including some C18 hydroxyl acids that can be esters, diesters and triesters and straight-chain monohydric alcohol compounds with carbon chains from C24 to C36 [30]. The increase of beeswax concentration leads to higher values of hardness and adhesiveness, which could be associated with strengthened network of gels formed during crystallization process (Figure 1). The highest hardness and adhesiveness values were 13.75 N and 2.83 N for organogel made with 17.5% beeswax. Gels were shown to become stronger and sticker after adding beeswax. Similar trend was reported in cod liver oil organogels formed with beeswax and carnauba wax [31]. It was proposed due to progression of aggregates among the sterol-sterol ester assemblages of the β-sitosterol and γ-orzanol organogelator used by Calligaris et al. [32]. The gel strength increases in the carnauba wax-sunflower oil organogels as organogelator (the Candelilla wax) concentration increases [10]. Those results agree with our findings. Very similar trends of changes were observed at another important texture parameter, adhesiveness for all samples and the above researches. Adhesiveness is the force required to remove a material from its surface. Some moderate levels of both hardness and adhesiveness are necessary for the shortening and margarine products for baking industry.

However, the low viscosity of the coconut oil when dough mixing and it melts causing a difficulty in shaping and handling, when coconut oil melts out the dough and it will enhance the cohesion of gluten strand forming during cookie dough mixing and it form a greasy and firmer cookie texture. There was no significant effect was found on cookie dough hardness, adhesiveness and springiness compared to that of cookie dough made with shortening (data not shown).

![Figure 1](image)

Control: commercial shortening; CNO + BW (7.5%): 7.5% beeswax-coconut oil organogels; CNO + BW (10%): 10% beeswax-coconut oil organogels; CNO + BW (12.5%): 12.5% beeswax-coconut oil organogels; CNO + BW (15%): 15% beeswax-coconut oil organogels; CNO + BW (17.5%): 17.5% beeswax-coconut oil organogels. Expressed as mean ± standard deviation (n=3). Values followed by the different letter within each column are significantly different (p<0.05).

Figure 1. Properties of commercial shortening and organogels: hardness (A) and adhesiveness (B)
3.3. Cookie Chemical Composition and Spread Factor

No significant differences were found between cookie control and tested groups for crude protein (1.68% to 2.0%), fat (22.79% to 26.91%) and carbohydrate (64.54% to 68.15%) (p>0.05) except moisture (3.72% to 8.17%) and ash (1.57% to 1.76%) content of cookies. The cookie of 7.5% beeswax-coconut oil organogels showed highest moisture content (8.17%) and lowest ash content (1.57%).

The spread factors of cookies made with beeswax coconut oil organogel were in the range of 7.9 to 11.9, which were not statistically significant (p>0.05) to the spread factor (10.36) of cookie with the shortening (data not shown). The similar spread factor to the cookies made of shortening shows the amount of beeswax in this study would not affect the spread factor at application of cookie products. The spread factor was not much affected by 7.5% to 17.5% beeswax-coconut oil organogel. Our results agree with finding of that when added 2% to 10% of natural wax to olive oil and flaxseed oil organogel and their cookie spread factors were not affected [19].

Cookies made with organogel showed a decrease in cookie hardness compared to that with coconut oil (Figure 2), they were not statistically significant (p>0.05) comparing to cookies made with shortening. Increasing the amount of beeswax from 7.5% to 17.5% in the organogel did not show a significant decrease in cookie hardness. The hardness of cookies with 12.5%, 15.0% and 17.5% beeswax coconut oil organogel (4.90 N, 4.75 N and 4.95 N, respectively) was similar to the hardness of the cookies with shortening (4.71 N, Figure 2) (p>0.05). Jang et al. [9] proposed the cookies with the candelilla wax-canola oil organogel showed the replacement of shortening with organogel produced cookies with soft eating characteristics.

However, there was oil separate out of cookies (Figure 3) when the cookie left on paper towel after baking specially for the cookies made with coconut oil and 7.5% to 10% beeswax organogel. The phenomenon was less observed at cookies made of shortening and 12.5% to 17.5% beeswax organogel (Figure 3). This result could be also attributed to the high hardness and adhesiveness of the organogel which play a positive role in prevent liquid coconut oil leak out during and after baking. The experiment results showed that the beeswax helps the formation of the three-dimensional network of the solid phase in the cookie matrix prevent liquid oil separating out [10].

The color of cookies play an important role in consumers’ acceptability and perception of the cookie. Higher L value of 7.5% beeswax coconut oil organogel was observed. \( \Delta E \) of cookie made with 15.0% organogel was smallest (1.69) comparing to that of cookie made with shortening, which indicated the consumers might not be able to tell the color difference between these two products (data not shown).

3.4. Determination of Total Phenolics Content, Reducing Power and Antioxidative Activity

Addition of beeswax organogel increased the total phenolics content (Figure 4) of cookie. In comparison to the control, which is made with shortening, the content of phenolics in cookie extracts of 15.0% and 17.5% beeswax-coconut oil organogels were significantly higher (46.67 and 50.66 GAE/100 g sample, respectively) than that of control (23.47 GAE/100g sample) (Figure 4). It is due to the flavonoids present in beeswax [31]. These flavonoids are polyphenolics originate from honey and propolis. Tomas-Barberan et al. [33] examined the phenolic components of beeswax and found that it contained a family of flavonoids, the composition of which was identical with that of honey and propolis. The flavonoids were not characteristic of beeswax but, like propolis, were derived primarily from popular resin and from the honey. Beeswax is a complex mixture of chemical compounds predominantly based in straight-chain monohydric alcohol compounds with carbon chains from C24 to C36 and straight-chain acids with carbon skeltons of up to C36, including some C18 hydroxyl acids that can be esters, diesters and triesters [30]. Beeswax consists of 70 to 71% total esters, 1 to 1.5% free alcohols, 9 to 11% free acids and 12 to 15% hydrocarbons [34,35].
Control: commercial shortening; CNO: coconut oil; CNO + BW (7.5%): 7.5% beeswax-coconut oil organogels; CNO + BW (10%): 10% beeswax-coconut oil organogels; CNO + BW (12.5%): 12.5% beeswax-coconut oil organogels; CNO + BW (15%): 15% beeswax-coconut oil organogels; CNO + BW (17.5%): 17.5% beeswax-coconut oil organogels.

Figure 3. The oil spread pattern of cookie made with different concentrations of coconut oil and beeswax organogels

Figure 4. Comparison on the total phenolic content of cookie extracts

Control: commercial shortening; CNO: coconut oil; CNO + BW (7.5%): 7.5% beeswax-coconut oil organogels; CNO + BW (10%): 10% beeswax-coconut oil organogels; CNO + BW (12.5%): 12.5% beeswax-coconut oil organogels; CNO + BW (15%): 15% beeswax-coconut oil organogels; CNO + BW (17.5%): 17.5% beeswax-coconut oil organogels. Expressed as mean ± standard deviation (n=3). Values followed by the different letter within each column are significantly different (p<0.05).
The reducing powers of cookie extracts are shown in Figure 5. The reducing power of cookies made with 15% and 17.5% beeswax-coconut oil organogel exhibited lower value and they were lower than that of 10 μM ascorbic acid (p<0.05). It seems that beeswax addition decrease the reducing power of cookie extracts. However, the reducing power of cookies made with shortening and different organogels were not remarkable different (p>0.05).

Antioxidative activities of cookie extracts with and without light exposure during 2 hours period at room temperature are demonstrated in the Figure 6A and 6B, respectively. Clearly, antioxidative activities of cookies made with virgin coconut oil were always higher than those of control and 17.5% beeswax organogel. It may be due to beeswax replaced coconut oil in the organogel and it dilutes the antioxidative activity of cookie extract. There was a decrease trend in the antioxidative activity of cookie made with higher amount of beeswax under light exposure (Figure 6A). Antioxidative activity (38.16 and 43.39) of cookie made with 12.5% beeswax organogel was similar to that of control (40.38 and 39.01) with and without light exposure for 1 hour. However, the antioxidative activity of cookie made with 12.5% beeswax organogel was steady higher than that of control after 2 hours (Figure 6).

Control: commercial shortening; CNO: coconut oil; CNO + BW (7.5%): 7.5% beeswax-coconut oil organogels; CNO + BW (10%): 10% beeswax-coconut oil organogels; CNO + BW (12.5%): 12.5% beeswax-coconut oil organogels; CNO + BW (15%): 15% beeswax-coconut oil organogels; CNO + BW (17.5%): 17.5% beeswax-coconut oil organogels. Expressed as mean ± standard deviation (n=3). Values followed by the different letter within each column are significantly different (p<0.05).

**Figure 5.** Comparison on the reducing power of cookie extracts

**Figure 6.** Comparison on the antioxidant activity of cookie extracts with light exposure (A) and without light exposure (B)
Control: commercial shortening; CNO: coconut oil; CNO + BW (7.5%): 7.5% beeswax-coconut oil organogels; CNO + BW (10%): 10% beeswax-coconut oil organogels; CNO + BW (12.5%): 12.5% beeswax-coconut oil organogels; CNO + BW (15%): 15% beeswax-coconut oil organogels; CNO + BW (17.5%): 17.5% beeswax-coconut oil organogels. Expressed as mean ± standard deviation (n=3). Mean with superscripts of different small letters in the same columns are significantly different (p < 0.05). Mean with superscripts of different capital letters in the same storage days of each sample are significantly different (p < 0.05).

3.5. Oxidative Stability of Cookies Made with Organogels during Storage

At the beginning, peroxide value of extracted liquids from cookie made with shortening was 5 meq/kg lipids (Figure 7). Among all organogel samples, the peroxide values were in the range of 0.5 to 0.78 meq/kg lipids (Figure 7) after baking. There was a dramatically increase in peroxide value of control stored at 60°C for 20 days storage. Obviously, peroxide values of the cookies made with organogel were far below the 10 meq/kg liquids limit during 20 days storage at 60°C. It was indicated that coconut oil and its beeswax organogels was effective against the development of first step oxidation.

Cookie made with commercial shortening still had strong butter aroma after 4 days storage at 60°C. Nevertheless cookies made with organogels smelled more coconut aroma when it containing more beeswax. There was rancid aroma was detected in cookie made with shortening after 8 days storage at 60°C. The odor of cookies made with organogels was similar to that of 4 days storage. When cookies made with shortening stored at 60°C for 12 days, the oil rancid aroma and butter flavor were strong. Among all organogel samples, coconut aroma and rancid flavor were barely detected after 12 days storage at 60°C. Studies also showed the ability of beeswax organogels to maintain aroma characteristics through time, as volatile compounds were successfully incorporated [36].

3.6. Consumer Sensory Analysis

The results for appearance, odor, flavor, crisp, coconut flavor and overall acceptability for the differ samples is shown in Table 3. The appearance of the cookie made with shortening was rated higher (4.94) than that of cookies made with 7.5% and 15% beeswax organogels (4.54 and 4.51, respectively) (Table 3) (p<0.05). The odor, flavor, crisp and overall acceptability of cookies made with coconut oil and 15% beeswax organogels were significantly lower than that of control. There was no difference between control and 12.5% beeswax organogel, it was still on acceptable range (>4). The coconut flavor of all organogel samples was not significant higher than that of control. It indicates the difference was small (p>0.05). The mean coconut flavor scores of all the cookies made with organogels were not over 4, indicating they were acceptable to the consumers and the coconut flavor was not strong.

Table 3. The sensory evaluation analysis of cookie with different concentrations of coconut oil and beeswax organogels

<table>
<thead>
<tr>
<th></th>
<th>Appearance</th>
<th>Odor</th>
<th>Flavor</th>
<th>Crisp</th>
<th>Coconut flavor*</th>
<th>Overall acceptability</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>4.94 ± 0.96a</td>
<td>4.67 ± 1.11a</td>
<td>4.61 ± 1.22a</td>
<td>3.88 ± 1.49a</td>
<td>2.99 ± 1.56a</td>
<td>4.47 ± 1.05a</td>
</tr>
<tr>
<td>CNO</td>
<td>4.57 ± 1.17ab</td>
<td>4.03 ± 1.15a</td>
<td>3.88 ± 1.35a</td>
<td>2.96 ± 1.46ab</td>
<td>2.82 ± 1.60a</td>
<td>3.96 ± 1.43ab</td>
</tr>
<tr>
<td>CNO + BW (7.5%)</td>
<td>4.54 ± 1.05ab</td>
<td>4.44 ± 1.06ab</td>
<td>4.4 ± 1.17ab</td>
<td>3.24 ± 1.34bc</td>
<td>3.04 ± 1.44a</td>
<td>4.15 ± 1.04ab</td>
</tr>
<tr>
<td>CNO + BW (10%)</td>
<td>4.63 ± 0.96b</td>
<td>4.47 ± 0.98b</td>
<td>4.49 ± 1.11ab</td>
<td>3.63 ± 1.40ab</td>
<td>3.04 ± 1.55ab</td>
<td>4.31 ± 1.23ab</td>
</tr>
<tr>
<td>CNO + BW (12.5%)</td>
<td>4.72 ± 0.10ab</td>
<td>4.42 ± 1.15ab</td>
<td>4.47 ± 1.02ab</td>
<td>3.33 ± 1.47bc</td>
<td>3.24 ± 1.52ab</td>
<td>4.35 ± 1.14ab</td>
</tr>
<tr>
<td>CNO + BW (15%)</td>
<td>4.51 ± 1.07b</td>
<td>3.96 ± 1.12b</td>
<td>3.72 ± 1.10b</td>
<td>2.63 ± 1.18b</td>
<td>2.81 ± 1.30b</td>
<td>3.58 ± 1.06b</td>
</tr>
<tr>
<td>CNO + BW (17.5%)</td>
<td>4.69 ± 1.08b</td>
<td>4.29 ± 1.20b</td>
<td>4.11 ± 1.31b</td>
<td>3.19 ± 1.39bc</td>
<td>2.99 ± 1.53a</td>
<td>4.19 ± 1.18bc</td>
</tr>
</tbody>
</table>

Control: commercial shortening; CNO: coconut oil; CNO + BW (7.5%): 7.5% beeswax-coconut oil organogels; CNO + BW (10%): 10% beeswax-coconut oil organogels; CNO + BW (12.5%): 12.5% beeswax-coconut oil organogels; CNO + BW (15%): 15% beeswax-coconut oil organogels; CNO + BW (17.5%): 17.5% beeswax-coconut oil organogels. Expressed as mean ± standard deviation (n=72). Values followed by the different letter within each column are significantly different (p<0.05).

1 – 7 scale: 1 = dislike very much, 7 = like very much
1* – 7 scale: 1 = none, 7 = too strong.
feasible for utilization of beeswax-coconut oil organogel of lignoceric acid. It is suggested 12.5% beeswax is lauric acid, myristic acid, palmatic acid with less than 1% cookie made with beeswax-coconut oil organogels were the storage stability, prevents oil oxidation and gives no trans fatty acid was found and the main fatty acids of cookie made with shortening were palmatic acid and shortening. The use of coconut oil produces cookies with no significant difference was observed on cookie dough hardness, adhesiveness and springiness compared to that of cookie made with shortening. There was also no significant difference was observed on cookie hardness compared to that of cookie made with shortening. The use of coconut oil produces cookies with more greasy and less crispy characteristics. The hardness of cookie was significant decrease when beeswax organogel containing higher amounts of total phenolic content but lower reducing power. The antioxidant activity of cookie was  mainly coming from total phenolic content but lower reducing power. The hardess and adhesiveness of organogels were significantly increased by adding up 15.0% beeswax. The significant increase on the organogel melting point was also observed in above 12.5% beeswax supplemented organogels. The melting points of 7.5% and 10.0% beeswax organogels were close to that of shortening (34.3°C). There was no significant effect was found on cookie dough hardness, adhesiveness and springiness compared to that of cookie dough made with shortening. There was also no significant difference was observed on cookie hardness compared to that of cookie made with shortening. The use of coconut oil produces cookies with more greasy and less crispy characteristics. The hardness of cookie was significant decrease when beeswax organogel containing higher amounts of total phenolic content but lower reducing power. The antioxidant activity of cookie was mainly coming from coconut oil and it can prevent lipid oxidation of cookie stored at 60°C for 20 days. It was indicated that coconut oil and its beeswax organogels was effective against the development of first step oxidation. Coconut oil increases the storage stability, prevents oil oxidation and gives coconut flavor of the baked products. The main fatty acids of cookie made with shortening were palmitic acid and oleic acid with 0.4% trans fatty acid. However, there were no trans fatty acid was found and the main fatty acids of cookie made with beeswax-coconut oil organogels were lauric acid, myristic acid, palmitic acid with less than 1% of lignoceric acid. It is suggested 12.5% beeswax is feasible for utilization of beeswax-coconut oil organogel in cookie manufacture as replacement for commercial shortening.

A correlation analysis was evaluated between the physicochemical properties in order to better understand the relationships among different quality attributes (Table 4). Beeswax ratio in organogel was negative correlated with the antioxidant activity of cookie extracts (Table 4).

4. Conclusion

Hardness and adhesiveness of organogels were significantly increased by adding up 15.0% beeswax. The significant increase on the organogel melting point was also observed in above 12.5% beeswax supplemented organogels. The melting points of 7.5% and 10.0% beeswax organogels were close to that of shortening (34.3°C). There was no significant effect was found on cookie dough hardness, adhesiveness and springiness compared to that of cookie dough made with shortening. There was also no significant difference was observed on cookie hardness compared to that of cookie made with shortening. The use of coconut oil produces cookies with more greasy and less crispy characteristics. The hardness of cookie was significant decrease when beeswax organogel containing higher amounts of total phenolic content but lower reducing power. The antioxidant activity of cookie was mainly coming from coconut oil and it can prevent lipid oxidation of cookie stored at 60°C for 20 days. It was indicated that coconut oil and its beeswax organogels was effective against the development of first step oxidation. Coconut oil increases the storage stability, prevents oil oxidation and gives coconut flavor of the baked products. The main fatty acids of cookie made with shortening were palmitic acid and oleic acid with 0.4% trans fatty acid. However, there were no trans fatty acid was found and the main fatty acids of cookie made with beeswax-coconut oil organogels were lauric acid, myristic acid, palmitic acid with less than 1% of lignoceric acid. It is suggested 12.5% beeswax is feasible for utilization of beeswax-coconut oil organogel in cookie manufacture as replacement for commercial shortening.

Table 4. Correlations of beeswax concentrations, total phenolic content, reducing power and antioxidant activity

<table>
<thead>
<tr>
<th>Beeswax concentrations</th>
<th>Total phenolic content</th>
<th>Reducing power</th>
<th>AOA (light 1hr)</th>
<th>AOA (light 2hr)</th>
<th>AOA (dark 1hr)</th>
<th>AOA (dark 2hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beeswax concentrations</td>
<td>1</td>
<td>0.349</td>
<td>-0.177</td>
<td>-0.739**</td>
<td>0.809**</td>
<td>-0.695**</td>
</tr>
<tr>
<td>Total phenolic content</td>
<td></td>
<td></td>
<td></td>
<td>-0.370</td>
<td>-0.391</td>
<td>-0.259</td>
</tr>
<tr>
<td>Reducing power</td>
<td></td>
<td></td>
<td></td>
<td>0.525*</td>
<td>0.263</td>
<td>0.368</td>
</tr>
<tr>
<td>AOA (light 1hr)</td>
<td></td>
<td></td>
<td></td>
<td>1</td>
<td>0.663**</td>
<td>0.498*</td>
</tr>
<tr>
<td>AOA (light 2hr)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1</td>
<td>0.515*</td>
</tr>
<tr>
<td>AOA (dark 1hr)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1</td>
</tr>
<tr>
<td>AOA (dark 2hr)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.099</td>
</tr>
</tbody>
</table>

AOA (light 1hr): antioxidant activity with light exposure in 1 hour; AOA (light 2hr): antioxidant activity with light exposure in 2 hours; AOA (dark 1hr): antioxidant activity without light exposure in 1 hour; AOA (dark 2hr): antioxidant activity without light exposure in 2 hour. * and ** indicate significance at p<0.05 and 0.01 respectively.

References


