Quality changes and shelf life of salted duck egg white meringues stored in alternative packages at two temperatures

Paramee Noonim, Karthikeyan Venkatachalam*

Faculty of Innovative Agriculture and Fishery Establishment Project, Prince of Songkla University Surat Thani Campus, Makham Tia, Mueang, Surat Thani 84000, Thailand

*Corresponding author: Karthikeyan Venkatachalam, Faculty of Innovative Agriculture and Fishery Establishment Project, Prince of Songkla University Surat Thani Campus, Makham Tia, Mueang, Surat Thani 84000, Thailand. Emails: karthikeyan.v@psu.ac.th; drkarthikeyan.v@outlook.com

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Abstract

Salted duck egg white meringues stored in alternative packages (paper control, metalized low-density polyethylene (M-LDPE), and polyethylene terephthalate (PET)) and at two alternative storage temperatures (30°C and 40°C) were tested for changes in quality over a period of 120 days. Every 15 days, the meringues were tested for physicochemical and microbial qualities. Package type, storage temperature, and storage time all influenced the overall meringue quality. This study revealed that meringues kept at 40°C experienced more loss of quality than those kept at 30°C. During storage, meringue's lightness and yellowness diminished while redness increased, and the sample volume shrunk. The control samples were significantly more affected than the other cases. The sample's weight, moisture, $a_w$, and pH were highest in the control samples when stored at 40°C. Textural profiles such as hardness, chewiness, cohesiveness, fracturability, gumminess, and springiness decreased in all cases with storage time, whereas adhesiveness and resilience increased. The M-LDPE and PET packages maintained the sample texture. The radical scavenging abilities of the samples did not significantly differ by the alternatives tested, but a gradual increase was noted during prolonged storage. The control samples had significant levels of pathogenic bacteria (Escherichia coli and Salmonella) and spoilage microorganisms (yeast and mold), reducing the shelf life to 90 days. In contrast, M-LDPE and PET packages maintained the sample qualities throughout the testing period of 120 days. It was found that M-LDPE and PET packages prolonged the shelf life of meringues when stored at 30°C, more so than at 40°C.

Keywords: meringue; packaging; physicochemical qualities; storage; temperature

Introduction

The most traditional and popular preserved egg product in Thailand is salted duck egg, prepared by brining and/or mud coating. Thailand is well-known for its mud-coated salted duck eggs (Chi and Tseng, 1998; Lai et al., 1999). Properties of these eggs depend on the length of curing with salt. More prolonged curing could improve the taste and structure of the egg yolk, which is the main component, as it has a wide range of applications and higher economic value than the egg white, which is frequently discarded owing to extreme saltiness; this leads to economic loss and an increase in waste and pollution (Venkatachalam, 2018; Wang, 2017; Xu et al., 2017). Therefore, food products made from salted duck egg white, while avoiding expensive desalination processes, have recently become popular (Xiao et al., 2019), resulting in several food products, including meringue, noodles, and jelly products. Among the various products, meringue is one of the most straightforward products to
adopt this new philosophy, as it uses the egg white efficiently as its primary ingredient and along with sugar, salt, acid, and flavors, followed by baking to give it additional attractiveness (Bennion et al., 1997). The egg plays a crucial role, as the quality of the foam is mainly determined by the foaming capacity and volume of the foam and the egg white's stability (Lomakina and Mikova, 2006).

Bakery items are a low to intermediate moisture commodity, often sensitive to any condition that might induce spoilage by physical, chemical, or microbiological decomposition that reduces the shelf life (Amit et al., 2017). In high moisture products, the primary issue is microbiological deterioration caused by bacteria, yeasts, and/or molds (Rawat, 2015). Many industrially produced bakery products have a surface that is essentially sterile, but post-bake handling can quickly lead to fungal and microbial surface contamination by exposure to airborne contaminants as well as by equipment contact (Saranraj and Geetha, 2012). Therefore, storage under ambient temperature is preferable for storing bakery products. The meringues can be kept at room temperature for several weeks, although it may cause color, texture, and flavor changes (Punidades and McKellar, 1999). Food packaging can preserve food and help delay degradation, extend the shelf life, and maintain, improve, and ensure product quality and safety.

Paper packaging is typically not the best solution for protecting food for longer because it lacks barrier properties and heat stability. However, paper packaging is frequently coated and/or impregnated with materials such as wax, resin, or lacquer to improve its usefulness as a barrier (Marsh and Bugusu, 2007). On the other hand, packages made of metalized low-density polyethylene (M-LDPE) or polyethylene terephthalate (PET) are low-cost and lightweight with a variety of physical (including optical) characteristics (Andrady and Neal, 2009). In addition, plastics have varying permeability to light, gases, and vapors, which is a critical drawback in their usage. LDPE and M-LDPE, on the other hand, are two of the most widely used thermoplastic polymers in the food packaging industry. These are frequently utilized in the packaging of food, milk, and agricultural products (Bastarrachea et al., 2011), having diverse mechanical, physical, and chemical characteristics. Typically, the meringue products are stored in paper-laminated PE packages, as they are inexpensive and can have information printed on them. Yuceer and Caner (2021) observed that storing meringue products in high-density polyethylene (HDPE) package could also be able to store the meringues for up to 90 days with fewer loss in the physicochemical characteristics.

Salted duck egg white meringues are a newly developed product and have not yet been studied for shelf-life and stability. Therefore, the present study was aimed to examine the quality changes in meringue stored in alternative packages and at two storage temperatures for a prolonged period.

## Materials and Methods

### Materials and chemical reagents

Salted duck eggs (from 20 days of salt curing) were obtained from farmers in the Chaiya district of Surat Thani province, Thailand. The eggs were adequately cleaned and split open so that the egg white (EW) could be carefully separated for use in meringue preparation. The additional components in the recipe, including organic coconut sugar powder, pure vanilla extract, and stabilizer (cream of tartar), were acquired from a local store in Surat Thani province. Chemical and reagents such as 2, 2'-diphenyl-1-picrylhydrazyl, ferrous ammonium sulfate, tartaric acid, peptone, ethylenediamine tetra acetic acid, dimethyl sulfoxide, L-ascorbic acid, trichloroacetic acid (TCA), Nash reagent, 2,2'-azino-bis (3-ethylbenzthiazoline-6-sulphonic acid), ferrous chloride, ferrozine, ferric ammonium citrate, and ammonium sulfate were purchased from Sigma Aldrich (St. Louis, MO, USA). Ethanol was purchased from J. T. Baker (Phillipsburg, NJ, USA). Peptone, potato dextrose agar, and tryptic soy agar were purchased from HI Media Laboratory (Mumbai, India).

### Meringue preparation and storage

The appropriate ingredient proportions for making meringue were established as follows: egg white (EW, 100%); coconut sugar (75%); stabilizer (0.01%); and food flavors (0.01%). The EW was added to the mixing bowl and beaten using an electric kitchen mixer (RBSFOODMIXERPRO, Cuizimate, Thailand) with a 4.5 L stationary bowl and rotating beaters to make the meringues. The speed was set to number 4 (100 rpm) for 10 min, and during that time, the remaining components, namely, sugar, stabilizer, and food flavoring agent, were gradually added to achieve homogenous whipping and flavor dispersion. After getting homogenous whipping, the mixture was put into a pastry bag with a 3 cm nozzle, and a consistent cone ball shape was slowly formed on a baking pan covered with parchment paper. The meringues were then baked for 25 min at 163°C in a double stack infrared oven. Finally, the cooked meringues were allowed to cool down to room temperature before packing and storage. For packing and storing the meringue products, three alternative packages were used: polyethylene (PE)-laminated paper package (the control treatment), M-LDPE package, and PET package.
(Packaging materials used in this study were purchased from PPM Pack Co., Ltd., Thailand). In each replication, 25 meringues were included per package. The samples were securely packed and placed on a rack for 120 days at two different storage temperatures (30°C and 40°C). The samples were tested for various attributes at 15-day intervals until the end of storage, as indicated below. The storage was stopped when the microbial growth exceeded the acceptable limit.

Quality determinations

Measurement of color characteristics
Color measurement of meringue surface based on CIE L*, a*, b* color system was carried out using a Hunter Colorimeter fitted with an optical sensor (AOAC, 2003).

Measurement of diameter, height, and weight
Meringue diameter and height were measured with a Vernier caliper at two places, and the average values were recorded. The weight of the meringue was measured by using a weight balance (Zoulias et al., 2000).

Measurement of moisture, water activity, texture, and pH
The moisture contents of the cracker samples were measured using an infrared moisture analyzer (MA160, Sartorius, Germany). The water activity was measured at 25°C with a dew point water activity analyzer (Series 4TEV, Aqua Lab, WA, USA). Texture analysis (including hardness) of the baked meringue was assessed using a texture analyzer in a compression mode with a sharp-blade cutting probe. Pretest and posttest speeds were set at 1.5, 2, and 10 mm/s, respectively. For measuring pH, 10 g of sample was homogenized with 100 mL of distilled water and measured using a handheld digital pH meter (Clean, pH30, China).

Texture profile analysis
Meringue samples were tested using a texture analyzer (Texture Analyzer, model TA XT plus, Stable Micro Systems Ltd., UK) (Yuceer and Asik, 2020). The settings of the texture analyzer to measure the meringues were as follows: pretest speed 5 mm/s, test speed 1.0 mm/s, posttest speed 5 mm/s, penetration distance 10 mm, stopping time between directions 5 s, and trigger force 10 g. Stable Micro Systems aluminum probe (P/36R, 36 mm) was used to test the texture profile. The textual profiles that included hardness, adhesiveness, chewiness, cohesiveness, fracturability, gumminess, resilience, and springiness were obtained from the Texture Exponent Software made for texture analysis.

Antioxidant activities
Meringue samples were extracted before analyzing the antioxidant activities. For extraction, a 5 g sample was suspended in 10 mL of 95% ethanol and then mixed vigorously by vortexing for 3 min. After that, it was centrifuged at 10,000 g for 20 min at 4°C. The supernatant was collected and stored in the dark and refrigerated until analysis. For DPPH (2,2-diphenyl-1-picrylhydrazyl) radical scavenging assay (Brand-Williams et al., 1995), a 100 µL aliquot was placed into a test tube containing 3.9 mL of 60 µmol/L DPPH and mixed well. After that, the reaction mixture was incubated for 30 min in the dark at ambient temperature. Then, the reaction mixture was measured at 515 nm using a spectrophotometer. The results are expressed as a percentage of DPPH radical scavenging ability. For hydroxyl radical scavenging assay (Halliwell et al., 1987), a 1 mL aliquot was added in a test tube containing 1 mL of ferrous ammonium sulfate (0.13%- Ethylenediaminetetraacetic acid (EDTA) (0.26%) solution, 0.5 mL of 0.018% EDTA, 1 mL of 0.85% dimethyl sulfoxide, and 0.22% ascorbic acid). After that, the reaction mixture was mixed well and incubated in a water bath at 90°C for 10 min. Then, the reaction was terminated by adding 1 mL of ice-cold TCA. After that, 3 mL of Nash reagent was mixed in the reaction mixture and kept at room temperature for 15 min to develop a yellow color. Then, the reaction mixture was measured at 412 nm using a spectrophotometer. The results are expressed as a percentage of hydroxyl radical scavenging ability. For ABTS (2,2’-azino-di-3-ethylbenzthiazoline sulfonic acid radical cation scavenging) assay, a 100 µL aliquot was placed in a test tube and mixed with 100 µL ABTS reagent (as described in Lee et al., 2015) in a 96-well microplate, and then it was incubated for 6 min at room temperature. After incubation, the sample was measured at 734 nm using a microplate reader. The results are expressed as a percentage of ABTS radical scavenging ability. For ferrous ion chelating activity (Singh and Rajini, 2004), a 200 µL aliquot was mixed with 0.8 mL of 95% ethanol containing 2 mM FeCl₂, and 5 mM ferrozine. After that, the reaction mixture was kept at room temperature for 10 min. Then, the absorbance of the reaction mixture was measured at 562 nm against a blank. A control was prepared using 0.2 mg of EDTA in 95% ethanol. The results are expressed as a percentage of ferrous ion chelating activity.

Microbiological analysis
A 25 g sample was aseptically weighed and dissolved in 225 mL of sterile distilled water, followed by centrifugation at 500 g for 5 min, and then the supernatant was collected, and 5 mL was placed into a test tube. It was diluted at 1:10 using 0.1% peptone water, followed by microbiological analysis using the pour plate method. For total plate count, 1 mL of diluted sample was aseptically inoculated on a Petri dish with 20 mL of sterilized agar (plate count agar) and mixed thoroughly. After solidification of agar, the Petri dish was inverted and incubated for 48 h at 30°C. Colonies were counted and reported as
log CFU/g (American Public Health Association, 1978). For yeast and mold, 1 mL of diluted sample was aseptically inoculated on a Petri dish with 20 mL of sterilized, acidified (using 10% tartaric acid to adjust the pH to 3.5) potato dextrose agar, and mixed thoroughly. After that, the agar was allowed to solidify completely, and then the dish was incubated at 30°C for 48 h. Colonies were counted and reported as log CFU/g (Lekjing and Venkatachalam, 2021). For *Escherichia coli* (*E. coli*), the samples were determined using Petri film *E. coli* Count Plate by following the AOAC (2002) method. Diluted sample (1 mL) was placed on the Petri film plate and left for 1 min to let the gel solidify, and then incubated for 2 days at 37°C. After incubation, the colonies were counted and reported as log CFU/g. For *Salmonella*, diluted sample (1 mL) was aseptically inoculated on a Petri dish containing 20 mL of tryptic soy agar yeast extract along with 0.05% ferric ammonium citrate and 0.03% sodium thiosulfate (Amanda et al., 2014). After that, the plate was allowed to solidify and was incubated at 35°C for 48 h. Colonies were counted and are reported as log CFU/g.

**Statistical analysis**

A completely randomized design was used in this study. Sampling and experiments were done in triplicates. Data shown are mean values with standard deviations. One-way analysis of variance was carried out, and means comparisons were performed using Duncan’s multiple range test. Statistical analysis was performed using SPSS (SPSS Inc., Chicago, IL, USA) for Windows.

**Results and Discussion**

**Color characteristics and physical dimensions**

Figure 1 shows the time profiles of color, height, diameter, and weight of the meringues in long-term storage in the various packages and the tested storage temperatures. Overall, this study discovered that the storage period and storage temperature substantially impacted the quality of the meringue. Typically, protein-rich products are susceptible to changes in color (Chevallier et al., 2000). The prolonged storage affects the color coordinates, including lightness, redness, and yellowness (Figure 1A–C). In a comparison of the treatments, the control samples showed significant changes in meringue color. The lightness and yellowness gradually decreased, while the redness steadily increased with storage, indicating that the meringues were affected by nonenzymatic browning reactions (Charoen et al., 2014). In comparison to the lower temperature storage, the higher temperature gave more browning to the samples. The differences in color changes between the treatments were minimal. However, the M-LDPE samples retained their color better and developed browning more slowly than the PET samples. On-nom et al. (2015) also reported a similar finding, as the prolonged storage decreased the color of samples when stored in PET more than with the M-LDPE package. Moisture and water activity of the samples play crucial roles in browning (Gonzalez, 2017). This study found that the range of water activity in samples was susceptible to the onset of the Maillard reaction. Bassey et al. (2013) also reported that water activity is the key factor in stored samples allowing activation of the browning-related reactions. The increase in the moisture level of bakery products or any dried products in a package depends on the water vapor transmission through the barrier package (Shakerardakani and Karim, 2012). When meringue samples were stored at the lower temperature, they retained their height and diameter (Figure 1D–E), which tended to decrease during storage. Prolonged storage had a noticeable effect on the height and diameter of the meringue. While no significant differences were observed between the various treatments, compared to control, substantial differences were observed. Abasi et al. (2009) reported that changes in the structure of baked food materials are mainly attributed to time and temperature. In addition, the choice of treatment had no discernible effect on the weight of meringues (Figure 1F). However, control samples held at either storage temperature had a gradual weight increase over time, albeit a relatively small increase. The increased weight in control samples could be due to sugar in the meringue recipe, which is hygroscopic and likely absorbs free moisture from the air while the sample firmness should be degraded.

**Moisture, water activity, and pH**

Figure 2 shows the meringue moisture, water activity, firmness, and pH changes during storage in the alternative packages and at two temperatures. The moisture content of the samples steadily increased with time (P<0.05), and the water activity followed a similar pattern. In a comparison of the packages, the results indicate that meringue samples stored in paper packages at both temperatures were highly affected by the atmospheric moisture content (Figure 2A). In addition, the temperature was a significant factor affecting the changes in the meringue moisture content. Turan (2021) revealed that the moisture absorption of the food product through the package is significantly impacted by the rate of water vapor transmission, the thickness of the package, and the environment’s relative humidity (RH). At the higher storage temperature, the elevated relative humidity in the environment increased the moisture transmission rate across the packaging material and the food product (Kurek et al., 2014). At temperatures above ambient, airier food products such as meringue absorb moisture...
faster than other products (Aviara, 2020). Water activity is one of the significant parameters associated with the shelf life via effects on storage stability, microbial growth, and rheological behavior of dehydrated food products (Syamaladevi et al., 2016; Yuceer, 2020). Yuceer (2020) has shown that prolonged storage of meringue elevated its water activity dramatically, mainly attributed to its high hygroscopic nature. The water activity of the control samples increased gradually over time, whereas the samples stored in M-LDPE and PET packages remained stable (Figure 2B). A steady increase in pH was observed in all samples stored in different packages and at different temperatures (Figure 2C). The typical pH of meringue is 7.0 (Kim et al., 2016), but this study found that the addition of salted duck egg white slightly decreased the pH to 6.8. Storage temperature did not affect the pH changes
Quality changes of meringues during storage

Figure 2. Time traces of moisture content, water activity, and pH of the meringue samples stored in alternative packages at two storage temperatures for a prolonged period.

Textural characteristics

Figure 3 shows the time profiles of textural characteristics of meringues stored in the different packages and at two temperatures. The amount of force required to compress food products by a certain amount is referred to as hardness (Singh et al., 2015). The hardness in all cases tended to decrease throughout storage. Control samples had the most severe loss of firmness, followed by M-LDPE and PET samples (Figure 3A). The lower temperature retained firmness better than the higher temperature. The differences in the sample's hardness could be caused by the absorption of moisture. Firmness is directly related to density, viscosity, surface tension, and other physical features of the food product (Day and Golding, 2016; Pascua et al., 2013). The water activity and the specific volume could significantly impact the firmness of the meringue over prolonged storage. The adhesiveness of the meringue increased with storage time, and the control samples had higher adhesiveness than the others (Figure 3B). Storage temperature significantly impacted the adhesiveness of the meringues in the control package, whereas it did not affect samples in M-LDPE and PET packages. On the other hand, the chewiness and cohesiveness decreased with storage time (Figure 3C–D). Samples in the M-LDPE and PET packages retained chewiness and cohesiveness better, whereas the control samples significantly lost those attributes. Storage temperature and storage period significantly affected these values, and the control samples had lower chewiness when stored at 30°C and slightly higher cohesiveness when stored at 40°C. Moisture content could adversely influence the textural properties. Grigelmo-Miguel et al. (1999) reported that textural properties, particularly the chewiness, were adversely affected by the moisture content in muffins. Wada et al. (2017) reported that hardness, cohesiveness, and adhesiveness shared a similar trend in baked food products. The fracturability of meringue indicates the ease of breaking and decreases gradually with the storage time (Figure 3E). Storage temperature and packaging material did not significantly affect the meringue's fracturability. Singh et al. (2015) reported that baked food products enriched with fiber exhibited more fracturability than protein-rich products. On the other hand, the gumminess of the meringue gradually decreased with storage time (Figure 3F). The control samples lost more gumminess than the other cases, and there were no significant differences among the treatments. Storage temperature did not influence the meringue's gumminess much. Resilience indicates the ability of a food product to retain its original shape. The results showed that increased storage time increased the resilience of the meringue (Figure 3G). The control samples had the least resilience, less than M-LDPE and PET packaged samples. Furthermore, the temperature did not affect the meringue's resilience. On the other hand,
Figure 3. Time traces of textural characteristics of the meringue samples stored in alternative packages at two storage temperatures for a prolonged period.
the springiness of meringues gradually decreased with storage duration, and the samples in M-LDPE and PET packages did not differ much during the storage and were not affected by the storage time (Figure 3H). In contrast, the control samples significantly lost springiness, and the samples stored at 30°C were more affected. Yuceer and Asik (2020) reported that prolonged storage significantly affected meringue's rheological and textural properties.

Antioxidant activities

Figure 4 illustrates the antioxidant activity of meringues held for an extended period in alternative packages at two temperatures. Overall, a sharp rise in antioxidant activity was seen in the meringues in all cases. Prolonged storage considerably affected the radical scavenging activity. The storage temperature significantly affected antioxidant properties, particularly DPPH radical scavenging, hydroxyl radical scavenging, and ferrous ion chelating activities. All meringue samples showed a linear trend in increasing DPPH radical scavenging activity (87–96%) (Figure 4A). In comparison to the other antioxidant activities, DPPH radical scavenging activity was the highest. The control samples had significantly less scavenging activity than samples in the actual treatments. The samples held in the M-LDPE package at both temperatures had increasing DPPH scavenging activity, followed by samples in PET packages. Storage temperature had no significant impact on the meringue's DPPH radical scavenging abilities; however, for samples stored at 30°C, the DPPH radical scavenging somewhat increased. Similarly, the hydroxyl radical scavenging activity (81–89%) and ABTS radical scavenging activity (55–67%) in meringue increased gradually with storage time (Figure 4B–C), and prolonged storage significantly increased these activities in the meringues, while the choice of packaging material and storage temperature did not affect them. Toward the end of storage, meringue samples stored in the PET package at either storage temperature had slightly higher hydroxyl and ABTS radical scavenging activities than the other cases. On the other hand, the ferrous

![Figure 4](image_url)

Figure 4. Time traces of various radical scavenging abilities of the meringue samples stored in alternative packages at two storage temperatures for a prolonged period.
meringues had a high level of total phenolics, which is widely regarded as an antioxidant source. According to Devi et al. (2015), coconut sugar contains a high level of ascorbic acid and polyphenolic contents. Coconut sugar was employed as a component in the meringue in this study, which may have contributed to the antioxidant activity of the samples. Purlis (2010) found a continuous development of condensation and polymerization via the Maillard reaction in samples during storage, owing to the sample's high amino acid and sugar contents and their pH, which are optimal for Maillard reactions. This is consistent with the findings of the current investigation.

**Microbial growth**

Figure 5 shows the time profiles of microbiological growth in meringues stored in different packages and at two storage temperatures. At both temperatures, considerable microbial growth was seen in the control samples; however, there was a noticeable trend in the ferrous ion chelating activity of samples, such that the meringue samples stored at 30°C showed a higher ferrous ion chelating activity than the samples stored at 40°C. The PET-packaged samples demonstrated more significant chelating activity than the M-LDPE-packaged samples when stored at 30°C, but this trend was not consistent at 40°C. Typically, eggs and egg-related products contain high-quality proteins such as ovalbumin, ovotransferrin, cystatin, and phosvitin, all of which have demonstrated significant antioxidant activity (Nimalaratne and Wu, 2015). Charoen et al. (2014) revealed that meringues have high antioxidant activity, attributed to melanoidin and D-ketohexoses, and found greater concentrations in the baked meringues. Kim et al. (2016) reported that

![Graphs showing microbial growth](image-url)
although there were no significant differences between the control samples, and those held at 40°C exhibited slightly higher microbial growth. The total plate counts of the meringue samples demonstrated that M-LDPE and PET packages could maintain the samples without microbial growth (Figure 5A), although after 60 days of storage, a slight onset of microbial growth was also found in these samples. However, the growth remained within acceptable bounds. Similarly, yeast and mold proliferation were extremely high in the control samples and lesser in those packaged in M-LDPE and PET (Figure 5B). In comparison to PET, M-LDPE packages gave slightly greater yeast and mold growth. The PET package may limit oxygen transport, thereby limiting the yeast and mold growth in meringues during storage. Among the many microorganisms found in bread products, mold is known to be the primary cause of deterioration. Low moisture content, water activity, and pH of baked food products may inhibit bacterial development (Guynot et al., 2005a, 2005b), while yeast and molds remain unaffected. Saeed et al. (2018) revealed that mold was more prominent in bakery products than yeast during storage. This could be due to the low moisture and water activity requirements for mold, whereas the requirements are higher for the growth of yeast. In addition, Salmonella and E. coli were also detected in meringue samples during the storage (Figure 5C–D). E. coli was observed in the control samples from day 15 onward, whereas the M-LDPE and PET packages successfully eliminated the growth of E.coli for most of the storage period. Samples held at 30°C had lesser E. coli growth than those stored at 40°C. Similarly, Salmonella development was seen in meringues stored for an extended period, particularly in control samples. In contrast, no Salmonella was observed in the M-LDPE and PET packaged samples at either temperature. Podolak et al. (2010) reported that Salmonella and E.coli are potential pathogens seen in food products baked in low moisture. According to Wu (2016), pathogenic bacteria can survive in extreme conditions, and under stressful conditions, they adapt by activating stress-induced responses (narZ, dadA, stiC, and rpoS), allowing them to survive in the long-term storage.

Conclusion

Salted duck egg white meringues showed good resistance against deterioration during prolonged storage. The alternative packages and storage temperatures played significant roles in the changes in meringue quality. The color of the meringue was significantly affected by the duration of storage. The higher 40°C storage temperature increased the browning of the samples compared with the lower 30°C temperature. Meringues also had a gradual weight increase with storage time at higher temperatures. Meringue samples stored in the control package with poor barrier properties were strongly impacted by the absorption of atmospheric moisture at both temperatures. The firmness of all samples tended to decrease throughout storage. Meringues exhibited the highest DPPH radical scavenging activity as compared to the other radical scavenging activities. Choice of packaging material or storage temperature did not significantly impact the meringue’s radical scavenging abilities. Microbial growth significantly impacted the meringue’s shelf life, especially for the control samples. Samples stored at 30°C had slightly less E. coli growth than those at 40°C. Salmonella and E. coli were also observed in the meringues during storage, but only minimally compared with yeast and mold growth. Control samples were severely affected by microbial growth, whereas the M-LDPE and PET packaged cases had only very minimal growth. This study concludes that M-LDPE and PET packages are well suited for packaging meringue for a prolonged storage period, and storage below 40°C would help retain the quality for a longer time.

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Conflict of Interest

The authors have declared no conflict of interest in the submission of this article.

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