Comparative Study of the Physico- and Biochemical Properties of Two Types of Salted Japanese Apricot (Prunus mume) Pickles

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Two types of commercial salted Japanese apricot (ume) pickle products with different textures were studied and their physicochemical and biochemical properties were compared. Considering the effects of fruit raw material ripeness and the pickle processing method, a pickled unripe-hard texture fruit (9% salinity) called “Karikari-ume” and a pickled ripe-soft texture ume fruit (10% salinity) called “Umeboshi” were used as sample materials. The results showed that the pH and moisture content of Karikari-ume (3.18 and 81.99%, respectively) were higher than that of umeboshi (2.84 and 74.08%, respectively). Meanwhile, the TSS and TA of citric acid and the TA of lactic acid value of the Karikari-ume (4.45, 0.92, and 1.30%, respectively) were lower than the Umeboshi (7.17, 1.79, and 2.52%, respectively). Karikari-ume also showed higher bioactive compounds and antioxidant activities assessed by DPPH•, ABTS•+, FRAP, and MIC assays (17.48–130.58 unit per gram of sample dry weight). These results suggested that the ripeness of the fruit material used in pickle processing could influence the physicochemical and biochemical properties of salted Japanese apricot pickles.

Keywords: Japanese apricot, salted pickle, firmness, acidity, bioactive compounds, antioxidant activity

INTRODUCTION

The Japanese apricot (Prunus mume) known as “ume” in Japan is rich in organic acids, edible fiber, minerals, and phenolic compounds. Owing to the high citric acid content of the mature fruit (up to 6–7% of fresh weight) in edible parts, the fresh fruit is extremely sour (Mitani et al., 2018). Ume fruit also contains very high contents of organic acids and phenolic compounds, mainly hydroxycinnamic acid derivatives (Mitani et al., 2013). Meanwhile, the immature green fruit contains cyanogenetic glycosides such as amygdalin (Go et al., 2018). Thus, ume fruit is not commonly consumed as fresh produce due to its sourness and toxicity; most of the fruit requires processing.

Traditionally, ume fruit is simply pickled with salt and processed by drying, because it can be used for medicinal purposes (Chung et al., 2013), e.g., as a hangover remedy, for liver rejuvenation, for detoxification, to treat nausea, and as an appetite stimulant. Besides consumption as traditional ume pickles, processed ume fruits are widely consumed in Asian and Southeast Asian countries.
as a sauce, juice, food garnish or decoration, and liquor (Chen et al., 2017). Owing to their high content of catalytic enzymes, probiotics, and bioactive compounds such as flavanols, ume could also provide antioxidative and antibacterial effects, as well as anti-inflammatory, hepatoprotective, and anticancer properties. Regarding the health aspect, the consumption of foods rich in flavanols was proven to decrease the risk of certain chronic diseases (Bailly, 2020; Gossard and Lipski, 2020). Salted ume pickles can also prevent the growth of some microbes (Mouritsen, 2018; Tylewicz et al., 2020).

There are two types of ume pickles: (1) pickled unripe ume fruit, commercially called “Karikari-ume,” and (2) pickled ripe ume fruit, normally called “Umeboshi.” These products show a different texture related to ripeness of the initial raw fruit material; Karikari-ume has a hard and crispy mouthfeel from pickling at the unripe green stage, while Umeboshi has a soft and slight mushy texture. Both salted pickles are often marinated with aromatic red perilla leaves (Shiso) as a natural coloring (red-purple color). As a pickle processing method for longer preservation, a salt-based osmotic agent is used to reduce the free water in raw materials. This type of osmotic agent normally absorbs free water and becomes a hypertonic solution, which generates a water transfer process by which free water in water-containing tissue is transferred to a more concentrated hypertonic solution. Meanwhile, the osmotic agent also diffuses into the material. Thus, a salt osmotic treatment influences the salt and moisture content of the fruit and can also affect its water activity (Osman and Faruk, 2016). For more acidic fruits like ume, osmotic salt treatment can affect the pH of processed material that includes an exuded salty solution, which could change the physicochemical and biochemical properties of processed pickle products. The rate of osmosis could also influence the acidity of the processed material (Ramya and Jain, 2017).

This study aimed to investigate and assess the physicochemical and biochemical properties of two different types of commercialized salted Japanese apricot pickles with different textures. Firmness, acidity, bioactive compound contents, and antioxidant activities were also investigated and compared between fruit types.

**MATERIALS AND METHODS**

**Chemicals and Reagents**

All chemicals and reagents used were of analytical grade or above. Chemicals used in the determination of antioxidative activity, such as 1,1-diphenyl-2-picrylhydrazyl (DPPH®), 2,4,6-tri(2-pyridyl)-s-triazine (TPTZ®), 3-(2-pyridyl)-5,6-diphenyl-1,2,4-triazine-p, p’-disulfonic acid monosodium salt hydrate (FerroZine®), FeCl₃, gallic acid monohydrate, and (+)-catechin hydrate, were obtained from Sigma-Aldrich Ltd. (St. Louis, MO, USA). Folin-Ciocalteu phenol reagent, Trolox® (6-hydroxy-2,5,7,8-tetramethylchroman-2-carboxylic acid), 2,2’-Azino-bis (3-ethylbenzothiazoline-6-sulfonic acid) diammonium salt (ABTS⁺), FeCl₂, and FeSO₄ were purchased from Wako Pure Chemical Co. (Tokyo, Japan), and ethylenediaminetetraacetic acid disodium salt (EDTA) was acquired from Dojin Chemical Laboratory Co., Ltd. (Tokyo, Japan).

**Pickle Samples and Physicochemical Property Determination**

A commercial Karikari-ume (hard texture) and Umeboshi (soft texture) were purchased from a supermarket in Matsudo, Chiba, Japan. Figure 1 displays the appearance of two types of ume pickles. The moisture content (MC) of the pickle samples was measured using a hot air oven (WFD-400; Eyela, Tokyo, Japan) at 100 ± 5°C for 16 h according to AOAC method number 930.15 (Horwitz and Latimer, 2005). pH, total soluble solids (TSS), and titratable acidity (TA) were determined using a pH meter (AS800; As One, Osaka, Japan), a digital handheld pocket refractometer (PAL-1; Atago, Tokyo, Japan), and the titration method according to Islam et al. (2013). The salinity (%) was measured using a salt concentration refractometer (RAS-28, As One, Tokyo, Japan). The TA was also determined in the same juice by titration: 15 g of juice was added to a beaker, and the pH values were measured from the start until the endpoint (pH 9.0) of titration. The volume of NaOH solution required for titration was recorded and the percent titratable acidity (expressed as % acid equivalent) was calculated from the molecular weight of citric acid and lactic acid of 192.1 and 90.08 g/mol, respectively, with the following formula:

\[
\text{Weight of acid} = \frac{\text{Conc. of NaOH} \times \text{Vol. of used NaOH} \times \text{molecular weight of acid}}{\text{Exchanged electrons}}
\]

\[
\% \text{ Total acidity} = \frac{\text{weight of acid}}{\text{weight of sample}} \times 100
\]

All above determinations were performed in triplicate. Pickle firmness was determined using a Creep Meter (RE2-3305S,
Yamaden, Tokyo, Japan), and the conical probe was equipped with a 20 N load cell and a 2.00-mm-diameter probe for the karikari-ume (hard texture) and a 3.00-mm-diameter probe for the umeboshi (soft texture). Pickle seeds were removed before testing, and 10 replications were performed for each sample. The forces and cross-sectional area of loaded probes were recorded for the computation and converted to stress (N m⁻²) (Moore and Booth, 2015).

Biochemical Property Determinations

The prepared pickles were extracted and quantified by the procedure described by Ketnawa et al. (2020). The determinations of all spectrometric methods were measured by a Thermo Scientific Multiskan® FC microplate photometer (Thermo Fisher Scientific, MA, USA), and the assays were completed in 96-well microplates with slight modifications as described by Reginio et al. (2020). The extracted samples were stored at −20°C until further determinations.

Total Phenolic Content (TPC)

Total phenolic content (TPC) was determined using the Folin-Ciocalteu reagent test method based on the assay of Singleton et al. (1999) with slight modifications according to Reginio et al. (2020). The extracted sample (25 µL) or gallic acid standard (0–120 mg/L) was mixed with 125 µL of 10% (v/v) Folin-Ciocalteu solution and 100 µL of 7.5% (w/v) sodium carbonate solution. The mixture was incubated for 1 h at room temperature in the dark, and the absorbance was then measured at 740 nm; distilled water was used as a blank. TPC was reported as milligrams (mg) of gallic acid equivalents (GAE) per gram of sample dry weight (DW).

Total Flavonoid Content (TFC)

The total flavonoid content (TFC) was determined using an aluminum chloride colorimetric assay according to Zhishen et al. (1999) with slight modifications for 96-well microplate assays. A 33 µL of extracted sample or catechin standard solution (0–100 mg/L) was added to 113 µL distilled water, and 10 µL of 5% NaNO₂ was mixed in. After 5 min, 10 µL of 10% AlCl₃ was added. At 6 min, 67 µL of 1M NaOH was added to the mixture. Immediately, 80 µL of distilled water was added to the reaction mixture and mixed well. The absorbance of the mixture was determined at 520 nm. The TFC was expressed as milligrams (mg) of catechin equivalents (GAE) per gram of sample DW.

DPPH* Scavenging Activity (DPPH)

The DPPH* radical scavenging activity of the sample was investigated by a later modification of the assay of Brand-Williams et al. (1995) with slight modifications by Ketnawa et al. (2020). Briefly, 5 µL of extracted sample was added to 195 µL of 60 µM DPPH*. The mixture was vigorously shaken and allowed to stand in the dark for 30 min at 23°C. The absorbance of the resulting solution was measured at 517 nm. The DPPH radical scavenging activity was calculated by comparison with a Trolox standard curve (0–1,000 µmol/L) and expressed as micromoles (µmol) of Trolox equivalents (TE) per gram of sample DW.

ABTS** Scavenging Activity (ABTS)

ABTS** scavenging activity was evaluated according to Ketnawa and Ogawa (2019), with slight modifications. ABTS solution was prepared by dissolving 7 mM ABTS** in 2.45 mM potassium persulfate, allowing the mixture to react in the dark for 12 h at room temperature before analysis. The ABTS solution was diluted with distilled water to obtain an absorbance of 0.7 ± 0.03 at 740 nm. At the beginning of the reaction, 10 µL of the extracted sample was mixed with 320 µL of diluted ABTS solution. The absorbance was measured at 740 nm after incubation for 10 min at 30°C in the dark. An ascorbic acid standard (0–100 µmol/L) was prepared in the same manner as a reference standard, and ABTS** radical scavenging activity was determined in millimoles (mmol) of ascorbic acid equivalents (AsA) per gram of sample DW.

Ferric-Reducing Antioxidant Power (FRAP)

The ferric-reducing antioxidant power (FRAP) was measured by the FRAP assay of Benzie and Szeto (1999) with slight modifications by Reginio et al. (2020). FRAP reagent (300 mM acetate buffer [pH 3.6], 10 mM TPTZ® solution in 40 mM-HCl, and 20 mM FeCl₃·6H₂O solution in a ratio of 10 : 1 : 1) was prepared fresh and 130 µL mixed with 20 µL of the extracted sample. The mixture was incubated at the same temperature in the dark for 30 min. The absorbance at 595 nm was determined after incubation with distilled water instead of the sample as a blank. FRAP was calculated from the standard curve of FeSO₄ (0–500 µmol/L) and expressed as millimoles (mmol) of FeSO₄ equivalents per gram of sample DW.

Metal Ion Chelating (MIC) Activity

The chelating activity on Fe²⁺ was identified regarding Ketnawa and Ogawa (2019) with some modifications. The diluted sample (300 µL) was mixed with 2 mM FeSO₄ (5 µL) and 5 mM Ferrozine® (5 µL). The mixture was vigorously mixed and allowed to stand for 10 min at room temperature. The absorbance was then measured at 560 nm with distilled water as a blank. EDTA standard (0–10 mmol/L) was prepared and used as a standard curve. Ferrous chelating activity was expressed as millimoles (mmol) of EDTA equivalents per gram of sample DW.

Statistical Analysis

Triplicate determinations were analyzed throughout the experimental period using IBM SPSS Statistics for Windows, Version 26.0 (Armonk, NY: IBM Corp). Data and mean comparisons were performed using a t-test. Differences were considered statistically significant at p < 0.05.

RESULTS AND DISCUSSION

Physicochemical Properties

The physicochemical properties of the two pickles used are shown in Table 1. The pH of Umeboshi (10.0% salinity) was lower than that of Karikari-ume (9.0% salinity), which could be related to the lower MC of Umeboshi. According to a previous study by Odake et al. (1999), MC of fresh fruit (~90%) decreased to ~77% due to the salted processing. Thus, it was considered
that Karikari-ume would be a less salted condition than the umeboshi. The TSS and TA of citric acid and the TA of lactic acid for Karikari-ume were 4.45, 0.92 ± 0.01, and 1.30 ± 0.02%, respectively, whilst Umeboshi showed higher percentages of 7.17, 1.79 ± 0.03, and 2.52 ± 0.04%, respectively. These results relate to the lower in pH value of Umeboshi relative to Karikari-ume. A previous report by Wu et al. (2005) showed that malic acid, citric acid, quinic acid, and shikimic acid are major acids in Prunus fruits. Besides, Yu et al. (2015) also reported that the dominant organic acid in ume fruits was citric acid, which reaches 39.3 g/kg. Odake et al. (1999) reported that 9.87% NaCl of Umeboshi pickle had TA and moisture contents of ~3.8 and 77%, respectively. Karikari-ume produced significantly higher pickle firmness than Umeboshi. According to the processing method, Umeboshi has longer pickling time using the ripe ume fruit, meanwhile, Karikari-ume made from unripe small green ume fruit was pickled by a different processing method depending on the fruit conditions. The differences in physicochemical parameters between them could depend not only on the original fresh product properties but also the processing conditions.

### Bioactive Compounds
Bioactive compounds for both types of salted pickles were determined and reported as total phenolic content (TPC) and total flavonoid content (TFC) (Figure 2). TPC values for Karikari-ume and Umeboshi were 1.99 and 2.10 mg GAE/g DW, respectively, with no significant differences (p > 0.05), whereas TFC was considerably higher in Umeboshi than in Karikari-ume. This may be due to the difference in the initial level of TFC in the fresh raw material and could be attributed to the different biochemical, physiological, and structural reactions and modifications that occur during the fruit maturation process. Previous research found that a fresh Japanese apricot fruit contained high values of phenolic acids such as 3-O-caffeoylquinic acid, 5-O-caffeoylquinic acid, chlorogenic acid, and tetra-O-acylated sucrose-related compounds (Mitani et al., 2013). The range of contents was from 150 to 320 µmol per gram trolox equivalents, which depended on the stage of fruit ripeness (Mitani et al., 2013). The observation of the decrement of phenolic contents and antioxidant activities as ripening
progressed was reported in previous studies (Macheix et al., 1990; Reginio et al., 2020), thus affecting the contents of polyphenols and their antioxidant activities of ume pickles in the study.

It was reported that a Japanese traditional pseudo-medicinal product named Bainiku Ekisu, a paste extracted and concentrated from Japanese apricot fruit, could inhibit bacterial growth (E. faecalis, S. aureus, and E. coli) in concentrations between 1.0 and 10.0 mg/mL (Yang et al., 2014). Additionally, Mitani et al. (2018) reported that Umezu, a salty extracted juice by-product of umeboshi processing, contained high amounts phenolic compounds, further clarified by microbial growth inhibitory activity research. Umezu was reported to have antimicrobial activity against enterobacteria due to the action of phenolic compounds (Mitani et al., 2018). Phenolic compounds modify the permeability of bacterial cell membranes and cause changes in various intracellular functions by hydrogen bonding to enzymes or modification of cell wall rigidity with integrity losses due to different interactions with the cell membrane. This may induce irreversible damage of the cytoplasmic membrane and coagulation of the cell contents that can even lead to the inhibition of intracellular enzymes of microorganisms (Cushnie and Lamb, 2011). Although it could not be simply compared with the current results, Karikari-ume might have more functional potential than Umeboshi, which is indicated by the significant differences in flavonoid content. The antimicrobial properties of pickles prepared under standard laboratory conditions in comparison with commercial ones, therefore, need to be investigated in the future.

Antioxidant Activities
Antioxidant activities were evaluated by means of DPPH, ABTS, FRAP, and MIC assays, and the results are depicted in Figure 3. Even though the difference in TPC between Umeboshi and Karikari-ume was non-significant (Figure 2), the antioxidant activity did not follow the same trend. All the antioxidant activity measurements on Karikari-ume were significantly greater than those of Umeboshi ($p < 0.05$, Figure 3). Karikari-ume had the highest ABTS and free-radical-scavenging ability: DPPH and ABTS ranged from 17.48 to 130.58 units per gram of sample DW. In addition, the FRAP and MIC values of Karikari-ume were 37.29 mmol FeSO$_4$ and 23.24 mmol EDTA equivalents per gram of sample DW, respectively, while the FRAP and MIC of Umeboshi were 6- to 8-fold lower. Debnath et al. (2012) reported that mume fructus extract inhibited scavenging activity against free radicals including DPPH*, ABTS*+, hydroxyl, and superoxide with half maximal inhibitory concentrations ($IC_{50}$) of 0.40, 0.36, 1.75, and 1.60 mg/mL, respectively. We assumed that the types of phenolics, which depended on the raw materials, maturity, and pickle processing method, could influence the antioxidant activity. For instance, the number of hydroxyl groups in phenolic molecules have an effect on antioxidant activities (Shahidi and Ambigaipalan, 2015). The lactic acid fermentation process for plant-based foods improved antioxidative activity due to microbial hydrolytic breakdown of plant cell walls and developed phenolic compounds (Hur et al., 2014). However, the hard cell wall of unripe ume fruit material may help to preserve bioactive compounds in their active form, whereas soft-type fruits may be prone to loss of activity to extracted juice during pickle manufacturing. Thus, further investigations of optimization of the ume pickle processing, characterization of phenolic acids, and flavonoid profiles are needed in the future.

CONCLUSION
In this study, differences were observed in the physicochemical properties between two types of ume pickles. The bioactive compounds present in ume pickles depended on the ripeness of the raw material and acidification processing, thus affecting the antioxidant properties. Karikari-ume- presented higher concentrations of total flavonoids and antioxidant activities. Considering that the raw material conditions and manufacturing process contributed to differential physicochemical properties, especially the lower moisture content and pH, these factors may affect water activity and antimicrobial agents in ume fruit. This was also related to acidification processes, and further research is needed on pickle manufacturing process optimization, the relationship between phenolic profiles and antioxidant activities, as well as other bioactivities. These data could benefit further research and the industrial sector by optimizing the pickling process, providing the highest amount of bioactive compounds with potential bioactivities. Moreover, they could help researchers to discover and develop novel healthy products and further study the release of bioactive compounds through digestion and antimicrobial properties.

DATA AVAILABILITY STATEMENT
The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

AUTHOR CONTRIBUTIONS
JS designed the experiment, performed the experiments, analyzed the data and prepared the figures, and wrote the manuscript. SK performed, verified the analytical assay, and reviewed the manuscript. YO supervised and critically reviewed the manuscript. All authors contributed to the article and approved the submitted version.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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