Gel-Forming Ability of Rohu as Affected by Egg White Powder Addition

Phatthira Sutloet¹, Warangkana Sompongse¹*, Katsuji Morioka²

¹Department of Food Science and Technology, Faculty of Science and Technology, Thammasat University, Pathum Thani, Thailand
²Aquatic Product Utilization Laboratory, Faculty of Agriculture, Kochi University, Nankoku, Japan

Abstract

The gel-forming ability of rohu (Labeo rohita) mince with and without egg white powder (EW) was investigated. Gel from washed mince (washed gel) was prepared under two setting conditions: kamaboko (40°C) and modori (60°C). The gel-forming ability of kamaboko and modori gels was improved by the addition of EW at 2%. The autolytic inhibition of kamaboko gel was obtained in gel added with 2% EW, and 1% EW of modori gel. No marked change was observed in the TCA-soluble peptide content of either gel with the addition of EW above 1%. No effect on the whiteness of both gels was shown after the addition of EW. The addition of EW exhibited smaller cavities and a more compact fibrous network in microstructure.

Keywords

Rohu (Labeo rohita), Gel-Forming Ability, Kamaboko Gel, Egg White Powder, Washed Gel

1. Introduction

The decline in marine fishery resources has affected mince/surimi production, and there have been attempts to utilize freshwater fish as an alternative, including Nile tilapia (Oreochromis niloticus) [1], red tilapia (Oreochromis niloticus × Oreochromis placidus) [2], common carp (Cyprinus carpio), and small-scale mud carp (Cirrhina microlepsis) [3]. Rohu (Labeo rohita) is a freshwater fish species that are widely aquacultured in Thailand. Based on the data for the year 2010, the amount of rohu produced in Thailand was 1167 tons, with a market value of 1.3 million US Dollars [4]. However, rohu contain small pin bones in their flesh, limiting their utilization. Processing the flesh to mince or for use as surimi would expand the range of utilization and market value.
Gel-forming ability is one of the key determinants defining gel quality [2]. Fish muscle cleavage by endogenous protease activity is typically exhibited by the degradation of myofibrillar protein [5]. This degradation, and especially that of myosin, has an adverse effect on gel-forming ability, inhibiting the development of strong three-dimensional networks [6] [7]. Generally, gel weakening is induced at 50°C - 70°C by endogenous thermal stable protease [8] [9]. This phenomenon is called modori. However, the activity of endogenous protease varies from species to species [10].

Egg white powder (EW) is one of the food-grade inhibitors used to overcome the modori softening gel [11] [12]. EW contains ovomucoid, which inhibits trypsin, and ovoinhibitor, which inhibits trypsin and chymotrypsin [13]. It is used as a food ingredient because of its unique functional properties [14]. EW has also been used to improve gel quality [10] [11] [12], inhibiting the softening of the gel by acting as an enzyme inhibitor [10] [15]. Research has reported that the addition of EW improved the whiteness of surimi [16]. However, no study has reported on the addition of EW to the gel from rohu. Consequently, the objective of the present study was to investigate the effect of EW on the gel-forming ability of rohu (Labeo rohita).

2. Materials and Methods

2.1. Materials

Rohu (Labeo rohita), weighing 1000 ± 100 g each, were purchased from Ying Charoen Market (Bangkok, Thailand). Fish were packed in an ice box and transported to the laboratory within 90 min.

Egg white powder (EW) was purchased from Thai Food and Chemical Co., Ltd. (Thailand). Trichloroacetic acid (TCA) was purchased from QRëC (New Zealand). Folin-Ciocalteu’s phenol was purchased from Merck KGaA (Darmstadt, Germany). Bovine serum albumin (BSA) was purchased from Sigma Chemical Co. (St. Louis, MO, USA). Reagents used for gel electrophoresis were purchased from Bio-Rad (Hercules, CA, USA).

2.2. Methods

2.2.1. Preparation of Washed Minces

The minced was prepared according to the method of [17]. The gutted and washed fillet was minced using a meat grinder (2 mm hole diameter; Kenwood, AT950A, England). Then, iced water was added at a ratio of 4:1 (w/w), the mixture was stirred for 5 min and left to stand without stirring for a further 5 min. The temperature of the mixture was maintained below 10°C throughout the process. After that, the mixture was filtered through cheesecloth. The washing process was repeated three times, twice with iced water and the third time with cold 0.3% (w/v) sodium chloride solution.

2.2.2. Preparation of Washed Gels

The procedure for preparation the gels from washed mince (washed gel) fol-
followed the method of [17]. Salt at 2.5% was added, and then EW at concentrations of 1%, 2%, and 3% (w/w) during grinding. Control is a gel without EW. The moisture content of gels was adjusted to 85% by the addition of iced water during grinding. The batter was stuffed into 2.5 cm diameter cellulose casing. The temperature of the batter was maintained below 12°C throughout the process. The kamaboko gel was heated at 40°C for 30 min followed by 90°C for 20 min. The modori gel was heated at 60°C for 30 min then 90°C for 20 min. All gels were immediately cooled in iced water until the core temperature of the samples fell below 10°C after heating. The samples were stored overnight at 4°C ± 2°C prior to further analysis.

2.2.3. Folding Tests
Slices with a thickness of 5 mm were cut from the 2.5 cm diameter cylinder shape, and folded into halves and quarters. The samples were evaluated followed the standard 5-point grade system [17].

2.2.4. Determination of Textural Properties
Gel samples were allowed to reach room temperature (approximately 30°C). Five cylinder-shaped samples 2.5 cm in length were prepared from each gel. The textural properties of the gel samples were determined following the method of [17].

2.2.5. Determination of Expressible Water Content
Gel samples were cut into pieces of 0.5 × 1 × 0.5 cm, and the expressible water content was measured following the method of [18].

\[
\text{Expressible water} \%(\%) = \left( \frac{(X - Y)}{X} \right) \times 100
\]

where, \(X\) is the original weight and \(Y\) is the weight after pressing.

2.2.6. Determination of Whiteness
Gel samples 2.5 cm in thickness and diameter were prepared, and their whiteness was determined using a colorimeter (ColorFlex CX2687, HunterLab, USA). D65 illuminant was used as the light source. CIE \(L^*, a^*, b^*\) values were measured. Whiteness was calculated using the following equation [19]:

\[
\text{Whiteness} = 100 - \left[ \left( 100 - L^* \right) + a^* + b^* \right] / 2
\]

2.2.7. Determination of TCA-Soluble Peptide Content
The TCA-soluble peptide content was measured by the method of [6] with slight modifications. Three grams of gel samples were homogenized with 27 ml of cold 5% (w/v) TCA using ACE homogenizer (AM-8, Nissei, Japan). The homogenate was kept in ice for 1 h and then centrifuged at 8000×g (MX-305, Tomy, Japan) for 10 min at 4°C. Soluble peptides in the supernatant were determined following to the method of [20] and expressed as µmol tyrosine/g sample.

2.2.8. SDS-Polyacrylamide Gel Electrophoresis (SDS-PAGE)
SDS-PAGE was used to analyze the protein pattern of the gel samples using the
method of [21]. Preparation of solubilized sample, stacking and separating gels and analyzing of protein pattern was carried out following the method of [17].

2.2.9. Scanning Electron Microscopy (SEM)
The microstructure of the samples was determined using a scanning electron microscope (JSM-IT300, JEOL, Japan). The specimens were fixed in 2.5% glutaraldehyde in 0.1 M phosphate buffer (pH 7.2) for 2 h. The specimens were rinsed twice with phosphate buffer and once with distilled water for 10 min. After that the specimens were dehydrated with a graded series of ethanol (30%, 50%, 70%, 95%, and 100% v/v) for 10 min. The samples were dried using a critical point dryer (Leica, EM CPD300, Austria), and were mounted and coated with gold (sputter coater, SCD 040, Bulzers, Germany).

2.3. Statistical Analysis
The experiment used a randomized complete block design. Data were subjected to analysis of variance. A Duncan’s new multiple range test was used to determine the differences between sample means at $P \leq 0.05$. All experiments were done in triplicate.

3. Results and Discussion
3.1. Effect of EW on Textural Properties
Figure 1 shows the result of the folding test and gel strength for washed gels containing EW at 0% - 3%. The addition of EW significantly affected the folding test of modori gel ($P \leq 0.05$), but not that of kamaboko gel. However, it significantly affected the gel strength of both modori and kamaboko gels ($P \leq 0.05$). As shown in Figure 1, the lowest folding test and gel strength were found in the modori control gel ($P \leq 0.05$), while the highest gel strength was observed in kamaboko gel with 2% EW ($P > 0.05$). It appeared that setting at 60°C (modori condition) resulted in gel weakening, whereas setting at 40°C (kamaboko condition) resulting in gel strengthening. As gel set at 40°C, endogenous transglutaminase (TGase) was activated, inducing cross-linking of $\varepsilon$-(γ-glutamyl) lysine in myosin heavy chain (MHC) via nondisulfide bonds [22]. In contrast, the weakening of modori gels was induced by endogenous protease. In a previous study on the degradation of washed gels, the lowest gel strength and maximum proteolytic activity in gel set at 60°C [17]. This supported the findings of [8] [9], who reported weakening of gel by endogenous protease under heating at 50°C - 70°C.

In the current study, gel strength increased as the concentration of EW was increased. In the case of kamaboko gel, the addition of 1%, 2%, and 3% EW increased the gel strength by 40.3%, 48.3%, and 45.7% over than that of the control kamaboko gel. In the case of modori gel, the increase was 110.9%, 135.8%, and 159.8%, compared with that of control modori gel. The results were in accordance with those of [23], who reported that the gel strength of arrowtooth
Figure 1. Effect of EW on textural properties of washed rohu gels. (a) folding test, and (b) gel strength. Bars represent the mean and standard deviation. Different letters on each bar indicate significant differences ($P \leq 0.05$).

flounder (Atheresthes stomias) and Alaska pollock (Theragra chalcogramma) surimi increased as the concentration of EW was increased to 2%. [12] found that the addition of 3% EW increased the breaking force and distance of common carp (Cyprinus carpio) surimi gel by 41% and 31%, respectively, compared with a control. Overall these results suggested that the addition of EW improves the textural properties of gels. It might be due to the surface SH groups of EW protein. This enhance gel network formation by forming S-S bonds with other protein molecules [24] [25]. Moreover, [16], who study the interaction between egg white protein and fish protein from Alaska Pollock, found that the decrease in total SH groups corresponding with the increase in force and deformation
value. Therefore, the improvement of gel strength in rohu gels was attributed to
the covalent bonding of sulfhydryl groups. In addition, some proteinase inhibi-
tors in EW have been reported to inhibit proteinases in lizardfish sarcoplasmic
fluid. These include cystatin, ovoinhibitor, and ovomacroglobulin, which inhibit
cysteine proteinase, serine proteinase, and aspartic proteinase, respectively [10]
[15]. This demonstrated that improvement in the strength of the two gels in the
present study was due to covalent bonding of sulfhydryl groups and proteinase
inhibitor. In this study, the gel strength of 2% EW modori gel was increased as
equal to the kamaboko control gel. The result suggested that the addition of 2%
EW improved the textural properties of both kamaboko and modori gels.

3.2. Effect of EW on Expressible Water Content

Figure 2 shows the expressible water content of washed gels at different levels of
EW. The lowest content was observed in kamaboko gel with 3% EW, though this
was not significantly different from the 2% gel (P > 0.05). The gel matrix with
the lowest water-holding capacity, as shown by the highest expressible water,
was that of the modori control gel. This result was in agreement with those for
textural properties, indicating that the strong three-dimensional network of ka-
mboko gel can retain water within its structure. The expressible water content
of both gels decreased as the concentration of EW was increased. For both gels,
the lowest expressible water content was observed with 3% EW. In the case of
kamaboko gel, the content was 24.2% less than control gel (P ≤ 0.05). In modori
gel, it was 37.6% less than control gel.

The modori gel had higher expressible water content than the kamaboko gel.
The expressible water content of the modori control gel had 28.2% higher than
that of the kamaboko control gel. As the concentration of EW increased, the
expressible water content decreased. This has been attributed to modori gel having a poor gel matrix with low water-holding capacity [26], which is improved by the addition of EW. In this study, the expressible water content of 1% and 2% EW modori gel was decreased as equal to the kamaboko control gel. It was suggested that the addition of 2% EW improved the water holding capacity of kamaboko gel and that of 1% EW in modori gel.

3.3. Effect of EW on Whiteness

Figure 3 shows the whiteness of the washed gels containing EW at different levels. The whiteness of neither gel was significantly affected by the addition of EW (P > 0.05). As the concentration of EW was increased, the whiteness of gel did not increase. This might be because the washed gel was naturally white in color, and was not significantly lightened by the light cream of the EW. In addition, it appeared that the effect of modori gel was not different from that of the kamaboko gel, as compared in the same levels of EW (P>0.05). The result was in agreement with [10], which reported that the addition of EW had no effect on whiteness of lizardfish surimi gel prepared from different heating condition.

3.4. Effect of Egg White Powder on TCA-Soluble Peptide Content

Figure 4 shows the TCA-soluble peptide content of both washed gels at different levels of EW. The TCA-soluble peptide content of the kamaboko gel decreased by 38.2%, 39.4%, and 52.5 % with the addition of 1%, 2%, and 3% EW, respectively, compared with that of the control gel. In the case of modori gel, the TCA-soluble peptide content decreased by 23.3%, 22.6%, and 23.3%, respectively. This supported the results from our previous studies, which suggested that the degradation of washed rohu gel was caused by serine protease [27]. Thus, the ovoinhibitor, one of the proteinase inhibitors in EW, could inhibit serine proteinase.

Figure 3. Effect of EW on whiteness of washed rohu gels. Bars represent the mean and standard deviation. Different letters on each bar indicate significant differences (P ≤ 0.05).
Figure 4. Effect of EW on TCA-soluble peptide content of washed rohu gels. Bars represent the mean and standard deviation. Different letters on each bar indicate significant differences (P ≤ 0.05).

3.5. Effect of EW on SDS-PAGE Profile

Figure 5 shows the protein patterns of kamaboko and modori gels containing EW at different levels. The MHC intensity of the kamaboko control gel was slightly lower than that of the unheated control mince (Lane 2, Figure 5(a)), while the MHC band was slightly higher in gel with 2% and 3% EW. In all kamaboko gels, the addition of EW had no effect on the intensity of degraded protein with molecular weights of 97 - 116 kDa, compared with that of kamaboko control gel (Lane 3, Figure 5(a)). In modori gel, the MHC band of the control gel was more severely degraded than that of the unheated control mince (Lane 2, Figure 5(b)). The addition of EW increased the intensity of the MHC band at all levels. These results coincided with the textural properties, and suggested that the degradation of both gels was prevented by the addition of EW. No change of actin band was observed for any gel.

3.6. Effect of EW on Microstructure

Figure 6 shows the microstructure of washed gels containing EW. Kamaboko gel without EW (Figure 6(a)) comprised a fibrous network with aggregation of packed protein, arranged as large clusters. As the concentration of EW was...
K is kamaboko gel and M is modori gel. MHC: myosin heavy chain; AC: actin; S: standard protein; R: raw sample (unheated control mince); CK: kamaboko control gel (without EW); CM: modori control gel (without EW); 1% - 3% K: kamaboko gel with 1%, 2%, and 3% EW; 1% - 3% M: modori gel with 1%, 2%, and 3% EW.

**Figure 5.** Effect of EW on SDS-PAGE pattern of washed rohu gels. (a) SDS-PAGE pattern of kamaboko gel and (b) SDS-PAGE pattern of modori gel.

![SDS-PAGE patterns](image)

**Figure 6.** Effect of EW on microstructure of washed rohu gels. ((a)-(c)) kamaboko gel and ((d)-(f)) modori gel. (a) and (d): kamaboko and modori control gel (without EW), (b) and (e): gel with 1% EW, (c) and (f): gel with 3% EW. Magnification: 10,000×.

increased, the cavities decreased in size and the fibrous network became more compacted (**Figure 6(b), Figure 6(c)**). The gel with 3% EW (**Figure 6(c)**) exhibited a clearly more compacted structure, compared with the control gel (**Figure 6(a)**). A fibrous network was also observed in the modori control gel (**Figure 6(d)**), but not as clearly. Whereas large cavities were found in the structure of the modori control gel, the gel with EW had a more compact structure with smaller cavities (**Figure 6(e), Figure 6(f)**). The modori gel with 3% EW had a
more clearly compacted structure than the control gel. The surface network of the kamaboko gel was smoother and more orderly than that of the modori gel. The higher gel strength of the kamaboko gel was attributed to this structure.

4. Conclusion

The addition of EW was demonstrated to improve the textural properties of washed gels, and to partially inhibit the proteolysis of modori gel. The gel-forming ability of kamaboko and modori gel was improved by the addition of 2% EW. For autolytic inhibition, the kamaboko gel was improved by the addition of EW at 2% and by the addition of 1% EW in the modori. The degradation of MHC in washed gels was prevented by the addition of EW. The microstructure of both gels exhibited smaller protein clusters, and cavities, when EW was added.

Acknowledgements

This research was supported by the National Research Council of Thailand for the 2017 fiscal year. Financial support from Thammasat University through the Ph.D. scholarship is acknowledged.

Conflicts of Interest

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

References


