THE INFLUENCE OF ADDITIVES ON FROZEN SNAKEHEAD FISH SURIMI AND THE APPLICATION OF TRANSGLUTAMINASE TO FISH CAKES

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ABSTRACT

Background. The co-product of cultured snakehead fish protein extraction is an abundant source of myofibrillar protein, with the potential for application in the processing of frozen snakehead fish surimi. The objective of this study was to determine the influence of additives and incubation time on the quality of surimi and surimi-based products.

Materials and methods. Cryoprotectant (a mixture of sucrose and sorbitol at a ratio of 1:1 changed from 2% to 4%), in combination with sodium tripolyphosphate (0.1, 0.15, 0.2 and 0.25%), was added to surimi during its preparation. In addition, the study also investigated the ratio of transglutaminase supplementation (0.5, 0.7 and 0.9%) and incubation time (2, 4 and 6 h) in the processing of high-quality fried fish cakes from frozen snakehead fish surimi.

Results. The results showed that, a combination of 3% cryoprotectant and 0.2% sodium tripolyphosphate helped maintain the quality of snakehead fish surimi after frozen storage. In the processing of fried fish cakes from frozen snakehead fish surimi, the addition of 0.7% transglutaminase (0.28 U/g surimi) with 4 h incubation significantly improved the gel properties of the product.

Conclusions. It is necessary to have appropriate additives and incubation time in the processing of surimi and surimi-based products from the co-product of cultured snakehead fish protein extraction.

Keywords: cryoprotectant, fish cake, sodium tripolyphosphate, surimi, transglutaminase

INTRODUCTION

Snakehead fish (Channa striata) have long been known to be a potential source of the proteins required by humans, especially those with poor health. This is due to its adequate supply of fish protein, and its balance of essential and non-essential amino acids (Mustafa et al., 2013). Surimi is the wet concentrate of the myofibrillar proteins of fish muscle which is light in color, bland in odor and low in fat (Ramadhan et al., 2014). This is an important intermediate product containing stabilized myofibrillar proteins obtained from

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minced fish that is washed to remove blood, pigments, lipids, enzymes and sarcoplasmic protein, blended with cryoprotectants and sodium tripolyphosphate (STPP), and then used in the production of fabricated fish-based products (Balange and Benjakul, 2009; Wu, 2016). Snakehead fish is a popular fish in Vietnam, especially in Vinh Long, but its price is very low. The coproduct of cultured snakehead fish protein extraction is an abundant source of myofibrillar protein (Vo et al., 2017). It possesses some important functional properties, such as gel-forming abilities and water holding capacities, which provide the potential for application in the processing of surimi. Luo et al. (2001) reported that the surimi from freshwater fish species is inferior marine species in terms of gel properties. However, Nousad et al. (1999) observed that surimi from tropical freshwater fish species, for example snakehead fish, wild mullet and Nile tilapia, showed good gel setting abilities. Hultin and Kelleher (2000) also used an alkaline solution to obtain surimi of a better quality, particularly a high gel strength, white color and the right flavor.

Transglutaminase is an enzyme isolated from a variant of *Streptomyces mobaraensis* that forms covalent cross-links between protein molecules (Ramadhan et al., 2014). The application of transglutaminase has created new technological opportunities for producing good fish cakes, even from poor quality fish muscle. Therefore, the aim of the study was to determine the effectiveness of the addition of cryoprotectant and STPP to maintain the gel quality of snakehead fish surimi after freezing and frozen storage. In addition, the study also investigated the ratio of transglutaminase supplementation and the incubation time in the processing of high-quality fried fish cakes from frozen snakehead fish surimi.

**MATERIALS AND METHODS**

The research was conducted at the Laboratory of Food Technology, Faculty of Agriculture and Applied Biology, Can Tho University. Major equipment and chemicals used in the study include Planimeter (Germany), Colorimeter NH300 (China), Rheotex SD305 (Japan).

Activa TG-SR-MH with transglutaminase activity of 40 U/g was used for this research (Determined according to Oteng-Pabi and Keillor, 2013).

Cultured snakehead fish (with average weight from 400€ – 700 g) was purchased directly from the farming area in Vinh Long province, Vietnam. After collection, the fish was transported live (in a bucket of water) to the laboratory, which took about 1 h. At the laboratory, the live fish was kept stable in the water tank for at least 1 h before further processing.

**Sample preparation**

Snakehead fish were weighed prior to preliminary processing. The fish was stunned, cut and its blood was discharged into the water tank. After the blood had been drained, the fins, skin, viscera and head were removed, and the fish was washed in 0.5% NaCl solution. After preliminary processing, the fillet was taken and washed with water at a low temperature (5−10°C). The fillets were then cut into 2 × 2 cm pieces and packed into PE bags (1 kg/bag) for freezing at –18 ±2°C. The fish meat had to be stored frozen at least 24 h before studying to help stabilize and regulate the material source and make it easy to cut afterwards.

**Research procedures**

The frozen fish meat was cut evenly (<5 mm) before washing to recover myofibrillar protein by heated magnetic stirring. One washing cycle was performed with NaCl 0.15 M solution, with a 1:2 ratio of minced fish and solvent at 35°C for 14 min to remove soluble protein and other soluble components, according to Vo et al. (2017). The wash mixture was compressed through a filter cloth to separate the fluid and recover the fish paste with a moisture content of less than 78%. The recovered fish paste was mixed with sugar, sorbitol and tripolyphosphate (experiment 1) to form surimi. The moisture content of the surimi was adjusted to 76% with ice. The final surimi was put into frozen storage for at least 24 h before conducting the next study. The frozen surimi was roughly cut to break down the structure while simultaneously mixing transglutaminase according to the study in experiment 2. The fixed additive and spices of NaCl, modified starch, pepper, garlic powder, monosodium glutamate and ice were 1.32%, 3%, 0.5%, 0.5%, 0.3% and 6.17%, respectively (Nguyen and Dang, 2003; Vo et al., 2018). The surimi were finely cut to create a paste before forming a disk shape (70 × 15 mm). The sample was kept in a refrigerator at 4 ±2°C with the
incubation time in experiment 2 before being pre-steamed at 80–85°C until the central temperature of the product rose to 55°C. The pre-steamed fish cake would be cooled to 2–4°C in a refrigerator for at least 24 h before deep frying at 180°C for 3 min. An analysis of fried fish cake parameters was carried out after keeping the sample in a refrigerator for at least 24 h. The result of experiment 1 was chosen as a fixed factor for experiment 2.

Experiment 1. Investigated the effects of cryoprotectant (mixture of sucrose and sorbitol at ratio of 1:1 changing from 2% to 4%) in combination with STPP (0.1, 0.15, 0.2 and 0.25%) on the quality of frozen snakehead fish surimi.

Experiment 2. Investigated the ratio of transglutaminase (TGase) supplementation (0.5, 0.7, 0.9%; corresponding to TGase activity changed from 0.20, 0.28 and 0.36 U/g surimi) and incubation time (2, 4, 6 h) in the processing of high-quality fried fish cakes from frozen snakehead fish surimi. Other additives were fixed as a result of experiment 1.

Proximate analysis
The proximate analysis was determined according to AOAC methods (AOAC, 1995), the moisture content was determined by AOAC 925.04, the protein content by the Kjeldahl method (AOAC 981.10) and the lipid content by the Soxhlet method (AOAC 920.39). The Water holding capacity (WHC) of the fish samples was determined by using the filter paper press method (FPPM; Muhlisin et al., 2012) with approximately 0.3 g of fish sample being placed between two filter papers and an amount of weight (1 kg) put on it to compress the fish for 10 min. The amount of water expressed to the filter paper due to the compression was indicated by the areas of the purge on the filter paper, which can be measured using a planimeter. Gel strength (GS) was measured using a Rheometer on a cylindrical fish cake (10 × 12 mm) (Nowsad et al., 2000). Whiteness (WI = L* – 3b*) was measured using a Colorimeter NH300 (Luo et al., 2004).

Data analysis
Using Statgraphics Centurion 16.1 program, data were analyzed for the degree of variation and significance of difference based on the analysis of variance (ANOVA) to determine if significant differences ($p \leq 0.05$) existed between treatments using the least significant difference (LSD) or Duncan test. All experiments were carried out in triplicate.

RESULTS AND DISCUSSIONS
Physical and chemical properties of snakehead fish muscle
Snakehead fish from Vinh Long province was used as a raw material for all experiments. The snakehead fish muscle (fillet, paste) was analyzed after each treatment to determine its basic chemical and physical properties (Table 1).

Table 1. Chemical and physical properties of cultured snakehead fish muscle

<table>
<thead>
<tr>
<th>Proximate parameters</th>
<th>Fillet</th>
<th>Paste</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture, %</td>
<td>77.44 ±0.42</td>
<td>78.47 ±0.31</td>
</tr>
<tr>
<td>Protein, %</td>
<td>18.52 ±0.29</td>
<td>16.75 ±0.49</td>
</tr>
<tr>
<td>Lipid, %</td>
<td>2.63 ±0.11</td>
<td>2.02 ±0.13</td>
</tr>
<tr>
<td>WHC, %</td>
<td>67.95 ±0.28</td>
<td>69.97 ±0.36</td>
</tr>
<tr>
<td>pH</td>
<td>6.77 ±0.06</td>
<td>6.83 ±0.06</td>
</tr>
<tr>
<td>Muscle recovery efficiency, %</td>
<td>46.05 ±1.63</td>
<td>70.16 ±0.64*</td>
</tr>
</tbody>
</table>

Different letters in a row indicate significant differences in the test treatments at 95% confidence intervals.
*Calculated according to the amount of fish fillet.

The results also showed that washing increased the water holding capacity of the fish muscle compared to raw fish fillet, as it reached 69.97%. This may have been due to the washing process that removed soluble proteins, lipids (2.02%) and other components from the fish muscle, increasing the muscle myofibrilar protein concentration in the paste (Chaijan et al., 2004). However, the loss of soluble proteins and the fact that the moisture content of the fish paste was maintained at a higher level than that of the fish fillet resulted in the protein content of the paste decreasing to 16.75%. The removal of the sarcoplasmic protein, lipids and pigments, as well as the pH of the fish paste
being maintained at a high level (pH 6.83) by washing with 0.15 M NaCl solution (pH 10), improved the gel formation of the fish paste (Balange and Benjakul, 2009). In addition, the low fluctuation of the raw material index (less than 2%) showed that the uniformity of the material sources had been guaranteed. All of this showed that the washed snakehead fish paste was suitable for processing protein-based products (surimi or fish cakes) but also that it required appropriate solutions to maintain and enhance the gel properties of the product.

The effects of cryoprotectant in combination with STPP on the quality of frozen fish surimi

In the technology of frozen surimi processing, STPP and cryoprotectant have played a key role in maintaining product quality by protecting the structure and the functions of proteins. In the study, an evaluation of the effects of these two additives on the quality of frozen surimi was done using fish cakes processed from surimi after one week of frozen storage. Table 2 shows that the addition of cryoprotectant (2% to 4%) and STPP (0.1% to 0.25%) had a significant effect on the quality of frozen surimi. This influence is reflected by changes in the gel properties of fried fish cakes.

The results showed that low supplementation rates of additives (2% of cryoprotectant, 0.1 or 0.15% STPP) resulted in the lowest gel quality. Research results have also confirmed that it is necessary to increase the rate of addition of additives to improve the quality of frozen surimi. According to Arakawa and Timasheff (1982), the supplementation of cryoprotectant to increase water surface tension and binding energy, prevents the separation of water from proteins, thereby stabilizing the protein of the product. However, the results of Table 2 and Figure 1 showed that the addition rate of the additives was too high (4% cryoprotectant and 0.25% STPP) to disrupt the gel system of the product (water holding capacity and gel strength decrease). This can be explained by the presence of large proportions of additives that reduce the protein concentration in the gel system (Luo et al., 2001). Therefore, it can be concluded that the addition of 3% cryoprotectant combined with 0.25% STPP is suitable

<table>
<thead>
<tr>
<th>Cryoprotectant, %</th>
<th>STPP, %</th>
<th>Whiteness</th>
<th>Gel strength, kgf × mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>0.10</td>
<td>40.11a±0.13</td>
<td>3.61d±0.13</td>
</tr>
<tr>
<td></td>
<td>0.15</td>
<td>40.05a±0.26</td>
<td>3.87d±0.19</td>
</tr>
<tr>
<td></td>
<td>0.20</td>
<td>40.73a±0.83</td>
<td>3.94bc±0.20</td>
</tr>
<tr>
<td></td>
<td>0.25</td>
<td>39.56a±0.68</td>
<td>4.02c±0.24</td>
</tr>
<tr>
<td>3</td>
<td>0.10</td>
<td>43.15c±0.32</td>
<td>3.86d±0.26</td>
</tr>
<tr>
<td></td>
<td>0.15</td>
<td>42.57c±0.32</td>
<td>3.69c±0.16</td>
</tr>
<tr>
<td></td>
<td>0.20</td>
<td>43.50c±0.58</td>
<td>4.40c±0.11</td>
</tr>
<tr>
<td></td>
<td>0.25</td>
<td>44.13c±0.93</td>
<td>4.28c±0.29</td>
</tr>
<tr>
<td>4</td>
<td>0.10</td>
<td>42.72a±0.74</td>
<td>3.40c±0.12</td>
</tr>
<tr>
<td></td>
<td>0.15</td>
<td>42.54c±0.31</td>
<td>3.43c±0.19</td>
</tr>
<tr>
<td></td>
<td>0.20</td>
<td>40.93b±0.51</td>
<td>2.88a±0.22</td>
</tr>
<tr>
<td></td>
<td>0.25</td>
<td>40.09a±0.31</td>
<td>3.12a±0.18</td>
</tr>
</tbody>
</table>

Different letters in a row indicate significant differences in the test treatments at 95% confidence intervals.
The role of transglutaminase supplementation and incubation time in the processing of high-quality fried fish cakes from frozen snakehead fish surimi

The gel-forming process involves cross-linking between myosin chains that are catalyzed by the intracellular transglutaminase. Reactions catalysed by transglutaminase result in significant changes in the physicochemical properties of gel, such as changes in viscosity, thermal stability and elasticity. Yongsawatdigul et al. (2002) investigated the residual transglutaminase activity in threadfin bream surimi wash water, confirming that 44% of the original transglutaminase activity remained in the final surimi after the washing process (Seighalani et al., 2017). All this has shown the need for transglutaminase supplementation during protein gel formation from fish surimi. In addition, the incubation time also plays an important role in promoting the effectiveness of this enzyme during processing. Therefore, the study was conducted on the rate of transglutaminase addition and incubation time to improve the gel quality of the surimi-based product, as shown in Table 3 and Figure 2.

Table 3 showed that different ratios of transglutaminase have a significant effect on the whiteness of the product, which increased with an increase in the ratio of transglutaminase. However, the difference was not statistically significant when the transglutaminase supplementation rate was 0.9% (or 0.36 U/g surimi).

In conclusion, the supplementation of cryoprotectant and STPP in the processing of frozen snakehead fish surimi showed that the addition rate of 3% of cryoprotectant and 0.2% of STPP was appropriate. At this rate of supplementation, the product processed from frozen surimi had an improved water holding capacity, gel strength and whiteness. The result was similar to that of Yoo (2014) in the selection of 3% sorbitol and 0.2% STPP for the processing of frozen surimi from Pacific Sand Lance.

Fig. 1. The effects of cryoprotectant in combination with STPP on the water holding capacity of fried fish cakes processed from frozen snakehead fish surimi.
The results of the study also showed that the whiteness of the product was high for a short incubation time and a high ratio of transglutaminase (highest whiteness with 0.9% transglutaminase and 2 h incubation). This result was similar to that of Karayannakidis et al. (2008), who observed that transglutaminase supplementation improved the whiteness of the gel product from sardine proteins.

The study by Seighalani et al. (2017), and Duangmal and Taluengphol (2010) also showed similar results with transglutaminase supplementation at 0.3 U/g surimi and 3 g/kg surimi, respectively. As transglutaminase catalyzes the cross-linking reaction of myosin and leads to the formation of protein intra- and intermolecular covalent bonds, it was concluded that differences in the whiteness value could correspond to the increased turbidity of gels because of transglutaminase activity. Moreover, an increase in whiteness could be caused by other compounds (lactose, maltodextrin) in commercial transglutaminase.

Table 3. The effects of incubation time and transglutaminase on whiteness, WHC of snakehead fish cakes

<table>
<thead>
<tr>
<th>Incubation time, h</th>
<th>Transglutaminase</th>
<th>Whiteness</th>
<th>WHC, %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>% surimi</td>
<td>U/g surimi</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>0.5</td>
<td>0.20</td>
<td>42.63^bc ±0.37</td>
</tr>
<tr>
<td></td>
<td>0.7</td>
<td>0.28</td>
<td>43.33^abc ±0.29</td>
</tr>
<tr>
<td></td>
<td>0.9</td>
<td>0.36</td>
<td>43.71^c ±0.42</td>
</tr>
<tr>
<td>4</td>
<td>0.5</td>
<td>0.20</td>
<td>41.94^b ±0.51</td>
</tr>
<tr>
<td></td>
<td>0.7</td>
<td>0.28</td>
<td>40.06^c ±0.54</td>
</tr>
<tr>
<td></td>
<td>0.9</td>
<td>0.36</td>
<td>43.22^abc ±0.37</td>
</tr>
<tr>
<td>6</td>
<td>0.5</td>
<td>0.20</td>
<td>40.99^bc ±0.52</td>
</tr>
<tr>
<td></td>
<td>0.7</td>
<td>0.28</td>
<td>42.47^abc ±0.48</td>
</tr>
<tr>
<td></td>
<td>0.9</td>
<td>0.36</td>
<td>43.01^abc ±0.28</td>
</tr>
</tbody>
</table>

Different letters in a row indicate significant differences in the test treatments at 95% confidence intervals.

Fig. 2. The effects of incubation time and transglutaminase on gel strength of fried fish cakes

powder, which cause a light scattering effect, as shown by Seighalani et al. (2017). Besides whiteness, the water holding capacity and gel strength are also important indicators in evaluating fried fish cake quality. Table 3 and Figure 2 show that the water holding capacity and gel strength of the product is significantly affected by the incubation time and the supplementation rate of transglutaminase.

At the appropriate level of incubation time and transglutaminase, the product had the best water holding capacity and gel strength. An increase or decrease of these levels has no positive impact on product quality. This is explained by the effect of transglutaminase, which helps the protein gel to be firmer. The stronger protein gel network was possibly linked to its capacity to hold water. However, transglutaminase can also increase protein to protein interactions, leading to a decrease in protein to water interactions. Therefore, the water holding capacity of gels can decrease when high concentrations of transglutaminase are used (Kaewudom et al., 2013). Figure 2 show that the water holding capacity of fried fish cakes was highest at 4 h of incubation and 0.7% transglutaminase (average activity was 0.28 U/g surimi). At this incubation time and supplementation rate of transglutaminase, the product is statistically different from the other combinations. The results are consistent with the publication of Chaijan and Panpipat (2010) on the effect of transglutaminase on the gelatin formation of mackerel protein (Rastrelliger branchysoma). This study indicated that the gel strength of the product increased with the addition of transglutaminase to 0.25 U/g surimi. The decrease in gel strength of the product with a high transglutaminase supplementation may be explained by the excessive formation of cross-linkages, which lower the gel strength through impeding intermolecular aggregation that reduces formation of formation the gel network (Jongiareonrak et al., 2006).

Figure 2 showed that the incubation time also had a significant effect on the gel strength of the fried fish cakes processed from frozen snakehead fish surimi. Increasing the incubation time from 2 h to 4 h significantly increases the gel strength of the product. However, if the incubation time is increased up to 6 h, there is a rapid decline in the gel strength of the product. At the same time, the results have also shown that increasing the rate of excess transglutaminase results in a breakdown of the gel structure of the product. These results are in accord with those obtained by Jiang et al. (2000) who reported that the gel structure of surimi gels from threadfin bream (Nemipterus virgatus) and pollock (Rastrelliger kanagurta) increased when transglutaminase increased up to a certain level, and a further increase in transglutaminase decreased the gel structure of the product (Seighalani et al., 2017).

From the above results, the combination of 4 h incubation time and 0.7% commercial transglutaminase (or 0.28 U/g of surimi) was the most appropriate condition for fish cake processing from frozen surimi. With the combination of these two factors, fried fish cakes made from frozen snakehead fish surimi had the best whiteness, water holding capacity and gel strength.

The results of the SEM also showed the effectiveness of transglutaminase in improving the structure and water holding capacity of the product (Fig. 3). The images showed that fried fish cakes processed from fish fillet had a bad structure and poor water holding capacity because the water was not distributed evenly.
inside the product. In this product, the water had gathered and formed large holes which were found to be smaller for the surimi origin (Fig. 3A, 3B). Figure 3C shows that, with the participation of 0.7% transglutaminase in combination with a 4 h incubation time, the surface structure of the product was uniform, and the water dispersed and was retained evenly over all parts of the product. As a result, the structure and water retention capacity of the product was significantly improved.

CONCLUSIONS

The results of the study showed the feasibility of using the paste of snakehead fish protein (co-product of soluble protein extraction) for processing frozen surimi. The combination of 3% cryoprotectant (1.5% sorbitol and 1.5% sucrose) and 0.2% STPP helped to maintain the quality of frozen snakehead fish surimi. In addition, the combination of 0.7% transglutaminase (or 0.28 U/g surimi) with 4 h of incubation in the processing of fried fish cakes significantly improved the gel properties of the product from frozen snakehead fish surimi.

REFERENCES


