Effect of Ginger’s (Zingiber officinale) Aqueous Extract on Characteristics of Transglutaminase Mediated Sausages from Thai Pangas (Pangasianodon hypophthalmus) Surimi During Refrigerated Storage

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Abstract

The effect of ginger’s aqueous extract (GAE) during refrigerated storage of the restructured products from Pangasius (Pangasianodon hypophthalmus) was evaluated. Protein and lipid oxidation, protein pattern, TPC as well as WHC, gelling properties, texture profiles and whiteness of the surimi gel was evaluated periodically during a refrigerated storage period of 20 days. Increase of water holding capacity in GAE added gels indicated stronger protein network formation, whereas, decrease of protein solubility suggested formation of protein aggregates during gelation. Lipid oxidation decreased in treated samples but the rate of increase varied, depending upon the concentration of GAE. Protein carbonyl increased during storage, but slowly in treated samples. Gel strength in treated samples increased and accompanied by thickening of myofibrillar head chain. Hardness, adhesiveness and gumminess parameters affected most due to addition of GAE. Sensory analysis revealed that the sausage with 1% GAE preferred most and control was acceptable up to 16 days.

Keywords: Pangasius; Sausage; Ginger; Protein oxidation; Lipid oxidation

Practical Application

In order to avoid any chemical gel enhancer or preservative for checking lipid oxidation in sausage type of products, use of microbial transglutaminase and natural antioxidant rich spice like ginger is a good option. From consumers’ acceptance point of view this type of products would fetch a level of consumers in city and big towns who are health conscious.

Introduction

Lipid oxidation is considered as a major cause of deterioration of muscle food which ultimately affect colour, flavour, texture and nutritional value of food. Lipid peroxidation and resultant lipid degraded products pose a great concern in respect of safety of fish and fish products. Oxidation thus limits the storage time and thereby also affects the marketing of both fish and meat products. Lipid oxidized products, particularly aldehydes, can react with specific amino acids to form carbonyls [1]. These can also interact with protein aggregates [2], causing additional nutritional losses. Antioxidants are usually incorporated to food to prevent development of off-flavour compounds in order to avert oxidative destability of food components [3]. Until recently, synthetic antioxidants such as butylated hydroxyanisole (BHA), butylated hydroxytoluene (BHT) and tertiary butylhydroquinone (TBHQ) were used to control lipid oxidation in foods. But, present day consumers do not prefer addition of any chemicals in food because of their health risks and toxicity. The fear of adverse effects of incorporating synthetic antioxidants in food products prompted a demand for novel natural antioxidants in food products [4]. A number of studies have been conducted to discover and develop novel antioxidative preservatives with different physicochemical properties needed for diversified food systems [5,6]. A wide variety of herbs, spices and fruits are used more and more as additives with antioxidative capacity [7,8].
For surimi products, the technique mostly used for obtaining a good gel depends on solubilizing and extracting myofibrillar proteins with 2 to 3g/100g salt and the solubilized expanding form a continuous matrix and then undergo thermal aggregation, cross-linking and develop into fine three dimensional solid-like networks resulting in elastic gel. The cross-linking of myosin promoted by a calcium-dependent endogenous transglutaminase (TGase) contained in fish muscle, which catalyses an acyl transfer reaction between γ-carboxymamide groups of glutaminyl residues in proteins. Fish with high content of lipid and myoglobin results difficulties in making high quality surimi [9]. To enhance the gel strength of surimi or fish mince, various food-grade ingredients and cross-linking enzymes such as microbial transglutaminase have been used [10]. Due to adverse effects of some ingredients on the surimi gel, particularly on its flavour or colour, the need of natural additives with an ability of protein cross-linking has been paid increasing attention for the surimi industry.

Polyphenolics are one of the compounds that are found in both edible and inedible plants and herbs/spices and it could be the source of a good antioxidant agent. These can act as reducing agents, free radical scavengers and Fe$^{2+}$ chelators or quenchers in the formation of singlet oxygen [11,12]. Thus phenolics are of increasing interest in the food industry because they retard the oxidative degradation of lipids and thereby improve the quality and nutritional value of food [13]. The high antioxidant capacity of these plant parts is particularly due to their content of different phenols, anthocyanins and ascorbic acid, which can act as radical scavengers [14]. The interactions between phenolic compounds and proteins play a very important role in the processing of certain food products. In an alkaline solution phenols may be oxidised easily to their corresponding quinones which can readily undergo attack by nucleophiles such as lysine, methionine, cysteine and tryptophan residues in a protein chain [15]. The formation of rigid molecular structures by reactions of ortho-quinones with proteins has been demonstrated by Strauss & Gibson [16]. Interactions of different phenolic acids and flavonoids with soy proteins were reported by Rawel et al. [17]. Significant increase in the gel strength of bigeye snapper surimi was found when oxidised phenolic compounds were added [18]. Antioxidant and antimicrobial characteristics of ginger extract have been reported by different workers [19,20].

The in vitro antioxidant activity of gingerol and other constituents of ginger [21] had been reported. Gingerol, the pungent factor in ginger oleoresin, inhibited phospholipids peroxidation induced by the FeCl$_3$ asorbate system [22]. Inhibition of xanthine oxidase activity responsible for the generation of reactive oxygen species, such as superoxide anion was documented with gingerol [23]. Sekiwa et al. [24] reported that glucosides related to gingerdiol from ginger has antioxidative activity.

The surimi making ability of many freshwater species is relatively low than that of its marine counterpart, but, could be upgraded by manipulating processing techniques. Thai pangas (Pangasianodon hypophthalmus) is extensively cultured in India and Bangladesh. The fish has a great aquaculture potential due to its very high growth rate compared to other popular major carps. The abundant catch of Thai pangas might be utilized as an alternative source of surimi raw material for development of restructured products. This study aims to determine the effects of ginger’s (Zingiber officinale) aqueous extract on lipid oxidation and microbial propagation on sausage from Thai pangas surimi during refrigerated storage by evaluating the characteristics such as textural, chemical and microbial besides acceptability.

Materials and Methods

Preparation of aqueous extract of ginger

To prepare aqueous extract of locally available ginger (Zingiber officinale), the ginger was peeled, cut into pieces and dried in an hot air oven at temperature 40±2 °C. It was then ground using an electric blender. Twenty grams of the ground material was soaked in 100ml of hot sterile water and allowed to stand for 48h. The crude extracts were obtained by filtration. The process was repeated twice and all the filtrates were collected and subjected to evaporation at 50°C in rotary vacuum evaporator (OSAKA J.P. Selecta, Spain). The powdered aqueous extract of ginger was kept in aluminium pouch and stored at -20 °C for future use.

Preparation of surimi

Fresh pangasius for the study was collected from the local fish farm at Lembucherra, Tripura. Length and weight range of fish were 37-49cm and 636-809g respectively. Fishes were washed with chilled water, gutted, dressed, filleted by hand and minced by employing a mechanical meat mincer with a 3mm-hole plate. Washing of the minced meat was performed in wash tanks maintaining a water temperature of 10 °C using a fish mince to water ratio of 1:4 (w/v) for three times with ten min duration of each wash (twice with potable water and last one with 0.1% NaCl solution to facilitate dewatering). The slurry was stirred for 3min and allowed to settle for 2min before water was decanted. Final dewatering was carried out using a screw press (Deb Enterprise, India). Sorbitol (4g), sucrose (4g) and polyphosphate (0.3g) were added to 100g of dewatered mince as cryoprotective agents and then mixed for 5min in a silent cutter (Sunlabz, India) at a temperature below 10 °C. The washed mince (surimi) was packed in low density polyethylene (LDPE) pouches (500g per pouch) and quickly frozen at -35 °C for 2h in air blast freezer (Sanyo, Japan) and stored at -20 °C in a deep freezer (Vest Frost, Denmark) for development of restructured products within a week.

Preparation of surimi gel

Frozen surimi was tempered for about 2h at 20±2 °C until it reached 5±1 °C, followed by chopping for 1 min at high speed in a silent cutter (Sunlabz Equipments, Chennai, India). Moisture of surimi was adjusted to 80% by using ice water. Salt (NaCl) was...
added@2.5% and mixed in a silent cutter for five min followed by microbial transglutaminase (MTGase) @1.0% and mixed for another five min. Aqueous extract of ginger (GAE) was added to each 500 g part@0.5, 1.0 and 1.5% and designated as PSgr-0.5, PSgr-1.0 and PSgr-1.5 respectively. Throughout the mixing operation temperature of surimi sol was kept below 10 °C. The control (CON) was made without addition of ginger extract. The surimi paste was then stuffed into vinylidene chloride casing (10 cm length, 2.0 cm diameter). Thermal setting was done according to the two-step heating method suggested by Luo et al. [25]. The casings were immersed in water at 40 °C for 30 min followed by immersion in water at 85 °C for 30 min. After cooking, the casings were immediately removed, placed in iced water, and cooled at 4-5 °C for 30 min. The gels were then stored overnight at 4 °C in a refrigerator. For storage study, the gels were stored at 4 °C in a refrigerator for 20 days and storage changes were analysed at 4 days interval.

**Moisture, ash, protein, fat content and pH**

Moisture, ash, protein and fat content of pangasius surimi were determined according to AOAC [26]. For determination of the pH, 10 g of sample were homogenized with 50 mL distilled water and pH value was measured by a digital pH-meter (Sartorius, PB-20).

**Determination of protein solubility, WHC**

Gel (0.5 g) were homogenised in 10 mL of 0.6M KCl in 50mM pH 7.4 tris-HCl buffer for 1 min in a tissue homogenizer (IKA, Germany). The homogenate was centrifuged at 10,000 rpm for 10 min at 4 °C (Remi, India). The supernatant was diluted ten-fold with 0.6M KCl and protein determination was performed using the Biuret method [27]. Analyses were performed in triplicate and the solubility was expressed in mg of soluble protein/100mg of gel.

WHC was evaluated by the technique outlined by Barrera et al. [28]. A portion of 5g of each gel was weighed and placed on 8 layers of filter paper (Whatman No. 1). Samples were placed in 50mL centrifuge tubes and centrifuged at 5000g at 4 °C for 15min (make REMI, India). Immediately after centrifugation, the gels were removed and re-weighed. WHC was expressed as the weight of the centrifuged gels relative to the original weight of samples.

\[
\text{WHC} (%) = \frac{(W2 / W1) \times 100}{100}
\]

where W1 represents the weight of the gel before centrifugation and W2 represents the weight of the gel after centrifugation.

**Determination thiobarbituric acid reactive substances (TBARS)**

The 2-thiobarbituric acid (TBA) assay was carried out according to the procedure of Schmedes & Holmer [29]. Sausage sample (10g) was mixed with 25 mL of trichloroacetic acid solution (200g/l of TCA in 135mL/1 phosphoric acid solution) and homogenized in a blender for 30 s. After filtration, 2 mL of the filtrate were added to 2 mL TBA solution (3g/l) in a test tube. The test tubes were incubated at room temperature in the dark for 20 h; then the absorbance was measured at 532 nm by using UV-VIS spectrophotometer (model UV-1200, Shimadzu, Japan). TBA value was expressed as mg malonaldehyde per kg of sausage. The values of three independent experiments were recorded as mean±SD.

**Determination of protein carbonyls**

Protein carbonyls were estimated according to the method suggested by Levine et al. [30]. Surimi gel (0.5 g) was homogenised in 10 mL 50 mmM tris-buffer (pH: 7.4) containing 1mM EDTA and 0.01% BHT. 200 μL of the homogenate was precipitated with 50 μL of TCA (100%) followed by centrifugation (10,000 g) for 5 min and the pellet was incubated in the dark for 1 h with 500 μL of 10 mmM dinitrophenylhydrazine (DNPH) in 2M HCl. A blank without added DNPH was made in 2M HCl and incubated in similar way. The samples were precipitated with 50 μL TCA (100%) and centrifuged (10,000 g) for 5 min followed by washing of the pellets three times with 1 mL ethanol/ethyl acetate 1:1 (v/v). The pellet was re-dissolved in 1 mL of 6M guanidine chloride in 20 mmM KH2PO4. Carbonyl content was calculated using the molar absorption coefficient of 22,000 M⁻¹ cm⁻¹.

**Determination of whiteness**

Colour of gel was determined in triplicate using spectrophotometer (Colourflex EZ, Hunter Associates Laboratory, Inc, Reston, VA) with illuminant of D 65/10°. This instrument was calibrated with black and white reference tiles before analysis. A horizontal section of gel measuring approx. 5mm was placed above the light sources and post processing L* (lightness), a* (redness/greenness) and b* (yellowness/blueness) values were recorded. The CIELAB (L*, a*, b*) colour scale was used for the study. Whiteness was calculated as described by Lanier et al. [1991] as follows:

\[
\text{Whiteness} = 100 - (100 - L^*) + a^*2 + b^*2 ½
\]

**Texture profile analysis (TPA)**

Texture profiles of gel were determined using a TA-XT2 Stable Micro Systems Texture meter (Surrey, England, UK). Restructured fish products (surimi gel) were removed from the casings and equilibrated to room temperature for 30 min in a plastic bag to avoid dehydration before the mechanical properties were measured. Textural profile analysis (TPA) was performed using an aluminium cylindrical probe (P/50) with 50mm diameter. Samples were compressed to 60% of the initial height using a compression speed of 60 mm min⁻¹. Hardness, springiness, cohesiveness and gumminess were reported for each treatment. Six samples were analysed for each treatment at room temperature (25-27 °C).
Changes in biochemical properties during refrigerated storage

Changes in pH, WHC and protein solubility: Changes in pH, water holding capacity (WHC) and protein solubility (PS) during refrigerated storage are given in Figure 1. The initial pH value (Figure 1a) ranged from 7.65 (in control sample) to 7.27-7.42 (in treated samples). Significant (p<0.05) reduction of pH in GAE treated samples may be attributed to the interaction of organosulphur/phenolic compounds of ginger with fish muscle proteins. In all GAE added formulations, storage had a significant (p<0.05) effect on the pH values, which tended to increase with storage time.

Indices such as water holding capacity (Figure 1b), is often used to assess the textural quality of the restructured fish products and it also indicates the deterioration of protein quality during low temperature storage. Water holding capacity increased (p<0.05) proportionately with the addition of GAE. The increases in WHC in day-1 in treated sausages may be explained as the formation of stronger network induced by GAE might imbied more water. However, the variation in WHC between the treatments may be due to differences in the concentration of phenolics in different treatments. From day-1 onwards, the differences between the treatments were significant (p<0.05). This may be explained as the result of protein denaturation induced by refrigerated storage leading to low affinity for water and it was accompanied by gradual loss of protein solubility. Moreover, modification of protein-phenolics interaction with the gradual denaturation and/or degradation of protein during storage may also be responsible for changes in WHC of protein as observed in this study. The texture of gel is also dependent on WHC which affect or influence sensory acceptability. So WHC is important to maintain at higher level during the storage period for better sensory quality.

The decrease in solubility (Figure 1c) suggests the formation of protein aggregates during gelation process. During heating, proteins underwent denaturation and aggregation to form a three dimensional structure. The alteration of protein extractability is a useful factor which may be used to determine the textural quality of fish muscle, as protein aggregation is accompanied by a significant decrease in their solubility [40]. In the present study, protein solubility not changed in Day-1, but as the storage progressed the PS values were found to be decreased significantly (P<0.05) in all the groups indicating the formation of protein aggregates during refrigerated storage.

Sensory evaluation

Sensory evaluation was performed by a panel of 6 judges. The panel evaluated each treatment within each replication in triplicate, and the evaluation was performed with the samples at room temperature. The panel judges were trained on the attributes of the sausage type products such as colour, flavour, taste and texture. Based on those attributes they were instructed to evaluate acceptability using ten-point Hedonic Scale (like extremely-9, like very much-8, like moderately-7, like slightly-6, neither like nor dislike-5, dislike slightly-4, dislike very much-3, dislike moderately-2, dislike slightly-1). A score below 6 was considered as rejected.

Statistical analysis

The data obtained were analyzed using analysis of variance (ANOVA), and when significant differences were found, comparisons among means were carried out by using Duncan’s Multiple Comparison Test (p <0.05) by Statistical Package for Social Sciences (SPSS, version 11.0 for windows).

Results and Discussion

Proximate analyses of fish muscle and surimi

The principal biochemical constituents of raw material fish Pangasius (Pangasianodon hypophthalmus) were moisture (74.04±2.5g kg⁻¹), protein (163.9±3.4g kg⁻¹), fat (75.7±1.4g kg⁻¹) and ash (10.9±0.2g kg⁻¹). The proximate analysis showed that the fish had low moisture and high protein and moderate fat content. Lower moisture and higher lipid content in pangas muscle was also reported by Hossain etal. [32], Silverstein etal. [33] reported the proximate composition of channel catfish as moisture (74-76%), protein (17-19%) and fat (3-6.9%). Whereas, in another study, proximate composition of catfish (Clarias gariepinus) was reported moisture, protein and fat content as 71.85%, 19.51% and 14.28% respectively [34]. There are influences of various factors such as nutrition, living area, and fish size, catching season, seasonal and sexual variation as well as other environmental condition on the proximate composition of fish species [35]. The surimi had moisture (795.7±1.8g kg⁻¹), protein (146.8±2.7 g kg⁻¹), fat (13.3±0.4 g kg⁻¹) and ash (33.6±1.4 g kg⁻¹). Washing reduced total protein content which may be explained as the removal of sarcoplasmic protein during washing which makes up to 20% to 25% of total protein of fish muscles [36,37]. Majumdar et al. [38] also reported a significant decrease of protein content after washing of silver carp mince. Removal of sarcoplasmic proteins may result increased concentration of myofibrillar proteins followed by improved water holding capacity of meat, because of increased hydration of protein due to removal of blood, pigments, proteins etc. during washing [39].
aggregates. The formation of disulphide bond which results in the aggregation of proteins [41] might have contributed to low solubility of proteins. Hydrogen bonds might involve in the interactions between hydroxyl groups of phenolic compounds and the nitrogen or oxygen of amino acids. From the result, the decreased protein solubility over the storage period indicated the aggregation as well as denaturation of proteins caused by low temperature.

Changes in total volatile basic nitrogen (TVB-N) and thiobarbituric acid reactive substances (TBARS)

Changes in TVB-N in GAE treatment as well as during storage period of surimi gel are presented in Figure 1d. Although, increase of TVB-N values in GAE treated samples compared to control in Day-1 was not significant (p>0.05), but changes during storage
Changes in the concentration of such compounds is a meaningful indicator of the oxidative status of muscle proteins [48]. Changes in the protein carbonyls of different treatments during refrigerated storage are shown in Fig. 1f. There was no significant difference between treatment and control in Day-1. As storage progressed protein oxidation was evident by a significant (P<0.05) increase in the protein carbonyl content. But the treated gel showed significant difference (P<0.05) with control in all sampling days. At the end of the storage period, control showed the highest and the surimi gel with 1.5% GAE showed lowest levels of carbonyl content. The order of protection offered against the formation of carbonyl compounds was PSgl-1.5> PSgl-1.0> PSgl-0.5> control. These results are consistent with lipid oxidation products emphasizing a possible relationship between the protein carbonyl formation and lipid peroxidation (Figure 1f).

The lowest carbonyl contents in GAE treated restructured products suggest a protective role of this extract against oxidation of proteins. Certain phenolic acids have been reported to inhibit formation of protein carbonyls in meat products as measured by DNP method [49]. Likewise, in our study also the phenolic rich extracts of ginger prevented the formation of protein carbonyls and thereby protein oxidation. Phenolic compounds have been suggested to inhibit the oxidation of proteins either by retarding the lipid oxidative reactions or by binding to the proteins or by forming complex with them [50].

Changes in whiteness

Whiteness of the gels was found to be reduced (P<0.05) upon addition of GAE (Figure 1g). That may be due to moderately grayish colour of GAE. This may be due to interaction of ginger’s organosulphur compounds with the muscle pigments leading to increase of the whiteness of the gel. But there were no significant changes (P>0.05) in whiteness during refrigerated storage of all the treatments as well as control except treatment with higher percent of GAE (PSgr1.5). The differences in colour alteration between treatments were possibly caused by the differences in pigment content in muscle. The result also indicates that the surimi treated with GAE prevented the oxidation of heme proteins present in the gel which are red in their reduced form and brown in their oxidized ferric form leading to maintain the whiteness during storage.

Changes in texture profiles

Amongst the textural attributes; hardness, springiness, cohesiveness and gumminess were measured periodically and presented in Figure 2. Hardness, springiness and gumminess of sausage were affected most due to addition of GAE. Hardness, peak force required for the first compression, increased (P<0.05) with increase of GAE up to 1.0%, but decreased thereafter in sausage having 1.5% GAE. Maximum hardness was found in sausage containing 1.0% GAE. This may be explained that more cross links following more GAE might reduce the flexibility of protein aggregates. The decreased gel strength at higher concentrations (1.5%) of GAE in the present study might be

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associated with self-aggregation of phenolic compounds, leading to the loss in capability of protein cross-linking. Cao et al. [51] reported the polymerisation of protein molecules as a possible subsequent reaction of different proteins with phenolic substances. This study also supports the observation by Ngapo et al. 1996 i.e., more interactions or cross links restrict the flexibility of the protein aggregates, and the gels become less springy and more rigid. The lower solubility of large phenolic compounds at high concentration causes the difficulty to interact with proteins [52]. It is also possible that the size of the phenolic compound can decrease its conformational flexibility in protein–phenolic compound interactions.

As the storage period increased, the hardness decreased with varying rate. On Day-20, the hardness of the sausages was found to be in the order of PSgl-1.0 > PSgl-0.5 > PSgl-1.5 > CON. Such behaviour of gels may be because of protein denaturation during low temperature storage and this was also found to be associated with decrease of WHC and protein solubility. Gumminess and springiness decreased (P<0.05) during the refrigerated storage of surimi gels, whereas no significant (P>0.05) change was observed in case of cohesiveness. Such changes of texture profile was presumed to be due to the increased denaturation of proteins, induced by extended refrigerated storage accompanied by loss of WHC and protein solubility.

Changes in TPC

Total plate count increased in all the groups during the period of refrigerated storage, but difference between control and GAE treated RPs was not significant upto day-4 (Figure1d). There were no significant differences (P > 0.05) with respect to TPC between the samples and ranged from 1.73 to 1.85 log cfu/g. Whereas, on 20th day, the TPC of control increased to 5.09 log cfu/g which significantly (P< 0.05) differed from the treated groups, wherein the count ranged from 4.49 to 4.97 log cfu/g with lowest recorded from sample treated with 2% GAE. Although the antimicrobial activity of ginger and ginger derived
organismsulfur compounds were reported in culture media, few reports are available on its effect in meat products. Whereas, the antimicrobial activity of organosulfur compounds was widely investigated against both food spoilage bacteria and food-borne pathogens [53].

**Acceptability**

The acceptability scores of the samples were assigned based on the attributes such as appearance, flavour, taste and texture. The acceptability of the product over the storage time as affected by different concentration of ginger’s aqueous extract is presented in Figure 2. The trend of overall acceptability observed in the day-1 was maintained even on day-20. All the treated groups were acceptable up to the end of the storage, however, they were scored in the order of PSgl-1.0 > PSgl-1.5 > PSgl-0.5. The control was found acceptable only up to Day-16 and sample PSgl-0.5 and PSgl-1.5 scored between slightly liked and moderate at the end of storage period. The consumers’ acceptability is usually based on the cumulative effect of all the sensory qualities, viz., appearance, texture, flavour, taste etc. Textural properties of sausage type of products are regarded as an important criterion for the consumer’s acceptability is concerned. In this study, the hardness of treatment PSgl-1.0 estimated to be highest amongst the samples. Next to texture, the sensory attribute which attracts the consumer more is flavour. Ginger has got some pungency in its flavour and too much ginger flavour is not accepted by the consumers. The result indicated that minimum ginger flavour in treatment with 1.0% GAE was highly accepted compared to others.

**Effect of GAE on protein patterns**

![Protein patterns of GAE treated surimi gel in day-1*](image)

Protein patterns of gel in control and GAE treated groups in day-1 is depicted in Figure 3. The thickness of the myofibrillar head chain (MHC) was found to be more in treated samples and also tended to increase with increasing concentration of GAE. Act in was found to be the dominant protein in the gel, suggesting that act in was more resistant to proteolysis or could not be polymerised during gelation. The result was in agreement with Benjakul et al. [54] who reported that act in Pacific whiting muscle was more resistant to proteolysis than MHC. The result suggests that GAE might be able to inhibit the degradation of MHC to some extent, as evidenced by the more retained MHC in treated gels. Protein cross-links might be more resistant to proteolysis caused by indigenous proteases. Phenolics present in GAE might have exerted some protective activity against certain proteases. Kroll et al. [55] reported that the interactions between phenolic compounds and proteins may result in inhibiting certain proteases [56,57].

**Conclusion**

The study revealed that addition of ginger’s aqueous extract in surimi sausage reduced lipid and protein oxidation during refrigerated storage. Moreover, the textural quality of the surimi sausage also improved with incorporation of ginger’s aqueous extract. Therefore, for development of sausages from freshwater fish like pangasius with inherent low gelling capacity, ginger in the form of aqueous extract may be used for safety of the product as well as for making the product as health food due to its enrichment with antioxidants from ginger.

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