INTRODUCTION
Banana blossom (Musa acuminata), also commonly called banana flower, is considered as a leftover product after cultivation, which is widely consumed as a vegetable in the Southeast Asian countries, including Indonesia, Malaysia, Thailand, and the Philippines (Liyanage et al., 2016; Wahab, Ismail and Abidin, 2020). Banana blossom contains high nutritional quality, especially dietary fibre. According to Sharma, Shukla and Golani (2019), the banana blossom (100g) contained dietary fibre (5.74g), protein (1.6g), fat (0.6g), carbohydrate (5.7g), calcium (73.3mg), phosphorous (56.4mg) and vitamin E (1.07mg). Dietary fibre plays an essential role in lowering serum cholesterol levels, preventing obesity, normalizing blood glucose and insulin levels (Bhaskar et al., 2012; Elaveniya and Jayamuthunagai, 2014). In addition, banana blossom possesses some bioactive components, such as vitamin C, saponin, tannin, flavonoid, alpha-tocopherol, and myoinositol phosphates, which could promote health benefits (Somsub et al., 2008; Sheng et al., 2010). Thus, due to being rich in a range of nutrients and biologically active components, the banana blossom has gained considerable attraction as an alternative source of functional food ingredients.

Abon, also known as shredded fish or fish floss, is one of the traditional dried meat products popular among Indonesia and the Asian community. It is recognized by different local names such as serunding in Malaysia, mahu in the Philippines, moo yong in Thailand, ihtieosteitc in Vietnam, and rousing in China (Hang, 2015). Popular raw materials for producing floss are beef and chicken; however, some fish species are also suitable materials for shredded meat processing. The processing of floss product generally begins with steaming of washed meat until it is tender. The steamed meat is then shredded finely and mixed with some formulated spices, followed by adding coconut milk. After that, the mixture is fried and stirred continuously under heat until the mixture is dry. Finally, excessive oil in the dried meat product is then separated and removed automatically (Huda et al., 2012). Shredded...
meat is consumed as part of a daily dish or served as filler of lemper (traditional food made of glutinous rice and wrapped in a banana leaf) (Suryani, Abdurrahim and Alindah, 2016). As a dried meat product, it contains a range of nutrients, including protein/amino acids, fat/fatty acids, and ash/mineral. Protein and fat found in meat floss products are generally high around 23.99 – 32.93% and 18.31 – 32.30%, respectively (Huda et al., 2012; Wazir et al., 2019; Fahmi and Purnamayati, 2020). A high level of fat is mainly caused by cooking oil absorbed during frying. As reported by Wazir et al. (2019), the meat floss contained approximately 87.73 – 91.65% of saturated fatty acids. For ash value, it contains around 3.17 – 6.67%. Among raw materials, a meat floss made of fish is much preferred by many people following the study from Huda et al. (2012), which revealed that fish floss has the highest score in overall sensory parameters compared to other meat floss samples.

Little tuna (Euthynnus affinis) belongs to the Scombridae family that is categorized as a medium-sized tuna (Ahmed et al., 2015). It is one of the most commercially important marine fish species in Indonesia. According to the Ministry of Marine and Fisheries Affairs (2021), the total production of little tuna significantly increased around 61.27%, from 366,900 tonnes in 2010 to 592,056 tonnes in 2019. In terms of nutritional value, little tuna is rich in protein and contains the amount of polyunsaturated fatty acids (PUFA). Kannaiyan et al. (2019) reported the protein content found in the white and dark muscle of little tuna was around 23.15% and 23.12%, respectively, while PUFA was around 51.86% and 55.87% in the white and dark muscle, respectively. Due to nutrients rich, little tuna is extensively employed as an essential raw material for sashimi and canned products in the seafood processing industry. Also, other diversifications from little tuna have been developed, including shredded meat, nugget, dumplings, fish ball, and fish cake (Suprayitno, Adi and Sulistiyati, 2016; Hizbullah et al., 2020). Fish floss derived from little tuna or other fish species mainly uses muscle part with adding some condiments. As a result, it has high protein content, but low dietary fibre. Moreover, histamine content would be high in the end-product because of the raw materials prepared particularly from the dark muscle of tuna and it could give rise to food-borne poisoning (Lee et al., 2016; Colombo et al., 2018). Therefore, there has been increasing interest in formulating food products with functional ingredients to deal with the unbalanced nutrition and undesirable component. Previous studies reported different quality aspects of banana blossom added nugget, noodle, biscuit, and meat floss ( Wahab, Ismail and Abidin, 2020; Elaveniya and Jayamuthunagai, 2014; Komal and Kaur, 2019; Novidiyanto et al., 2020; Puspita, Kartikaningsih and Dayuti, 2019). Among them, the added banana blossom could elevate the amount of dietary fibre, antioxidant, and other functional properties.

Scientific hypothesis
There is a lack of information regarding the use of banana blossom incorporated with tuna floss in increasing fibre content and decreasing histamine levels. Therefore, this study hypothesized that incorporating little tuna floss with banana blossom may give favorable nutritional features, particularly enhancing fibre content and lowering histamine levels. Banana blossom was added up to 500 g.kg⁻¹ in the formulation of little tuna floss. Nutritional aspects such as chemical composition, amino acids, fatty acids, and dietary fibre contents of the formulated little tuna floss were analyzed. Heavy metals content as safely required in the food developments was evaluated. The histamine content and pathogenic bacteria test were also determined. In addition, the sensory properties of the little tuna floss were studied to provide a basis for consumer acceptance and commercial applications. This study may stimulate further study in developing healthy fish floss products for nutritional and functional applications.

MATERIAL AND METHODOLOGY

Samples
Little tuna (E. affinis) used in this study was obtained from Sedang Biru fish market (Malang, East Java, Indonesia). The size of little tuna samples was approximately 202.8 ± 3.3 g and 27.6 ± 1.9 cm for weight and length, respectively. Banana blossom (M. acuminata) was purchased from a local market (Malang, East Java, Indonesia), and its weight was around 2.2 ±0.2 kg.

Chemicals
All chemicals and reagents used were of analytical grade. H:SO₄, petroleum ether and nitric acid (Merck, Germany) were supplied from a local supplier. Kjeldahl catalyst selenium tablet (Fisher Chemical, USA).

Bacteria and biological material
Bacteria used, including Escherichia coli, Salmonella sp. and Staphylococcus aureus were obtained from the Indonesian Culture Collection (InaCC), Research Center for Biology, Indonesian Institute of Sciences (LIPI). Microbial media used in a recent study were Oxoid (Basingstoke, UK)-based Nutrient agar (NA), MacConkey agar (MCA), Rappaport-Vassiliadis broth (RV), Xylose Lysine Deoxycholate (XLD) agar and Baird Parker agar (BPA).

Instruments
The instrument used consists of the oven (M720, Germany), Soxtect 2050 (FOSS Analytical, Denmark), ultra-pressure liquid chromatography (UPLC) (Waters, US), gas chromatography (GC) (Agilent Technologies, California, US), High-Performance Liquid Chromatography (HPLC) (Agilent Technologies, California, US) and Atomic Absorption Spectrophotometer (AAS) (GBC Scientific Equipment, USA).

Description of the experiment
Sample preparation
Around 15 kg of little tuna samples were purchased from Sendang Biru fish market (Malang, East Java, Indonesia). Fish samples were kept under cold conditions in an insulated box with ice during transportation (around 2-3 h). Upon arrival, the little tuna samples were washed in running water and were then filleted manually using a sharp knife. The filleted samples, including white and dark muscles, were stored in a freezer until use. For banana blossom, about 5 pieces of samples used were was obtained from a local market (Malang, East Java, Indonesia). Due to fresh banana blossom samples used, the...
detailed preparation was described in the following procedure for addition into little tuna floss.  

**Preparation of little tuna floss incorporated with banana blossom**

Table 1 presents the distinct formulations for all little tuna floss samples. The banana blossoms added with banana blossom in preparing little tuna floss products were 12.5% to 50% based on the weight of the whole little tuna sample (Figure 1). The little tuna meat used consists of white and dark muscles with distinct proportions i.e., 80% and 20%, respectively. These treatments were chosen in compliance with the formulated proportion for fish floss recommended by the previous study from Puspita, Kartikaningsih and Dayuti (2019). The control little tuna floss (0%) was used as a control and compared to the treated little tuna floss samples (12.5 – 50%) to evaluate the distinct characteristics of the formulation process and its corresponding attributes on the treated little tuna floss produced. During preparation, the thawed little tuna muscles were steamed at 100°C for 15 min (Halco, Indonesia). Separately, the banana flowers were cut into small sizes and then steamed at the same condition of fish treatment. After the steaming process, the steamed tuna was shredded using a fork and all ingredients used were thoroughly mixed into the pan. Afterward, the mixture samples were fried and stirred continuously under heat until the mixtures were dried. The excessive oil in the dried fish floss products was then separated and removed automatically using a spinner machine (Spiner Abon, Indonesia). All experimentations were carried out in triplicate.

**Nutritional attribute**

**Proximate analysis**

The proximate analysis of little tuna floss samples was determined according to the method of the Association of Official Analytical Chemists (AOAC, 2000). The Kjeldahl method was used to obtain the crude protein from all dried fish floss samples, while fat content was analyzed using the Soxhlet extractor method. The ash and moisture contents of the samples were examined by the gravimetric method. Carbohydrate content was measured by subtracting total crude protein, fat, ash, and moisture contents from 100%. For energy value (kcal.100g⁻¹), the calculation used the following formula provided by AACC (2000):

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\text{Energy value} = (4 \times \text{carbohydrate content}) + (4 \times \text{protein content}) + (9 \times \text{lipid content}) \quad (1)
\]

**Dietary fibre analysis**

The dietary fibre analysis used in this study referred to the method of AOAC 985.29 (AOAC, 2000). The little tuna floss samples were dried at 100 °C to constant weight using air-oven (M720, Binder, Germany). Around 0.5 g of all samples were added α-amylase enzyme and kept in accordance with its optimum condition (viz. pH 6; at 100 °C, for 30 min). Afterward, the treated samples were added protease with incubating at 60 °C for 30 min at optimum pH (7.5). The protease-treated samples were then added amyloglucosidase and incubated at pH 6 for 30 min at optimum temperature. Total dietary fibre of all treated samples was measured by precipitating using ethanol, followed by washing and drying. The obtained residues were then weighed manually.

**Amino acids analysis**

The amino acids composition was analyzed using ultra-pressure liquid chromatography (UPLC) (Waters, US) according to the method of Nollet and Fidel (2015). About 5 mL of hydrochloric acid 6 N was mixed to 0.1 g of samples. The prepared samples were then hydrolyzed at 110 °C for 24 h (M720, Binder, Germany). The hydrolysed samples were transferred to prepared distilled water. After that, the mixtures were filtered with a 0.45 μm filter paper. The 500 μL filtrates were mixed with 40 μL of 2-Amino-4-boronobutanoic acid (ABBA) and 460 μL of double distilled water. Then, 10 μL of the solution was mixed with 70 μL of AccQ Fluorine Borate and 20 μL of fluorine. The homogenized solutions were incubated at 55 °C for 10 min. Finally, the solutions were injected into the UPLC system to calculate the amino acid composition.

**Fatty acids analysis**

The fatty acid composition of little tuna floss was analysed by PT. Saraswanti Indonegetech, Bogor, Indonesia. Fatty acid methyl esters (FAMEs) from extracted fat of tuna floss samples were prepared by basic transesterification following the official method (AOAC, 2000), using hexane and hydroxide potassium 2N.

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**Table 1** Formulation for preparation of little tuna floss products.

| Ingredients                  | Control little tuna floss | Banana blossom addition |
|------------------------------|---------------------------|-------------------------|
|                              | 12.5%                     | 25%                     | 37.5%                    | 50%                     |
| Little tuna meat (g)         | 1000                      | 870.5                   | 750                      | 620.5                   | 500                     |
| Banana blossom (g)          | 0                         | 120.5                   | 250                      | 370.5                   | 500                     |
| Coconut milk (L)            | 1                         | 1                       | 1                        | 1                       | 1                       |
| Onion (g)                   | 40                        | 40                      | 40                       | 40                      | 40                      |
| Garlic (g)                  | 10                        | 10                      | 10                       | 10                      | 10                      |
| Ginger (g)                  | 10                        | 10                      | 10                       | 10                      | 10                      |
| Sugar (g)                   | 10                        | 10                      | 10                       | 10                      | 10                      |
| NaCl (g)                    | 80                        | 80                      | 80                       | 80                      | 80                      |
| Candlenut (g)               | 5                         | 5                       | 5                        | 5                       | 5                       |
| Brown sugar (g)             | 100                       | 100                     | 100                      | 100                     | 100                     |
| Lemongrass leaves (10 g)    | 20                        | 20                      | 20                       | 20                      | 20                      |
| Lime leaves (8 pieces)      | 8                         | 8                       | 8                        | 8                       | 8                       |
Figure 1 Preparation of tuna floss incorporated with banana floss.

Other spices and materials were proportionally prepared, including garlic, onion, ginger, candlenut, lemongrass leaves, lime leaves, coconut milk, salt, brown sugar and refined sugar.
FAMEs were analyzed by gas-chromatography (GC) (Agilent Technologies, California, US) described by Aquilani et al., (2018). The GC with a flame ionization detector (FID) was used to analyze the amount of fatty acids in the little tuna floss samples. The GC analysis was conducted using capillary column polyethylene glycol, equivalent variants CP-Wax 52 CB (30m × 0.25mm × 0.25µm film thickness). The carrier gas was nitrogen at a flow rate of 1 mL.min⁻¹. The initial oven temperature was set at 120 °C with an increase of 5 °C per minute until reached 240 °C. The injector and detector temperatures were 260 °C. Individual peaks were measured by comparison of their retention times (RT) with those of standards. Peak areas were calculated, and FAMEs were expressed as the area percentage of total area FAMEs (%).

**Heavy metal analysis**

Heavy metals analyzed in this study were composed of cadmium (Cd), lead (Pb), arsenic (As), mercury (Hg), and tin (Sn). These microelements were conducted by the method of AOAC (2000) using the Atomic Absorption Spectrophotometer (AAS) (GBC Scientific Equipment, USA).

**Histamine determination**

The histamine content was estimated by High-Performance Liquid Chromatography (HPLC) (Agilent Technologies, California, US) according to the method from Muscarella et al., (2005). Around 10 g of homogenized little tuna floss was extracted by 20 mL of 5% trichloroacetic acid (TCA). After filtration, 1 mL of the extracted sample was purified with 2 mL of chloroform solution. The mixture was then centrifuged and 20 µL of the supernatant was analyzed by HPLC. The HPLC column used was a Supelcosil LC ABZ 4.6 × 150 mm, 5 µm film thickness (Supelco, Bellefonte, PA, USA). The deansulphonate phosphate buffer pH 6.9/acetonitrile mixture (85/15) was used as a mobile phase. Isocratic elution was used; the flow rate was 1.2 mL.min⁻¹. The UVDAD was regulated at the absorption wavelength of 210 nm; the injection loop was 20 µL.

**Bacterial count**

Bacterial evaluations in the tuna floss consist of total bacteria count, total coliform, *Escherichia coli*, *Salmonella* sp., and *Staphylococcus aureus* with the methods developed by Anang et al. (2018) with some modifications. For total bacteria count, serial dilutions of 10 – 1 to 10 – 5 were prepared by diluting around 1 g of the little tuna floss samples into 9 mL of distilled water. Approximately 0.1 mL of aliquots from 10 – 3 to 10 – 5 dilutions were inoculated into Petri dishes containing the Nutrient Agar media. The plates were then incubated at 37 °C for 24 h. After incubation, bacteria colonies formed were counted using the colony counter and recorded as total viable counts. Total coliform was analyzed by dropping 0.1 mL of diluted samples into MacConkey broth with Durham tubes and incubated at 37 °C for 24 h. After incubation, the inoculated tubes were identified as positive total coliform by changing color from purple to yellow, and gas was collected in the Durham tubes. The positive tubes were then transferred into a 5 mL test tube of tryptone solution and incubated at 44 °C for 24 h to test *E. coli*. Afterward, a drop of Kovacs’ reagent was added into the treated tubes. The suspected *E. coli* in each tube was shown by a red ring color development, indicating the presence of indole. *Salmonella* sp., the count was evaluated using pre-enrichment of bovine peptone water, followed by enrichment of Rappaport-Vassiliadis (RV) broth. After enrichment, around 0.1 mL of aliquots were inoculated into Xylose Lysine Deoxycholate (XLD) agar. The suspected *Salmonella* colonies appear colorless with a black center because of H₂S production. For S. aureus count, about 1 g of fish floss samples were added into test tubes containing 9 mL of distilled water and the mixtures were serially diluted from 10 – 1 to 10 – 5. Around 0.1 mL of aliquots (10 – 3 to 10 – 5) were inoculated into plates prepared with Baird Parkeragar media. Afterward, the inoculated samples were incubated at 36 °C for 48 h. After the incubation period, colonies formed were identified with the suspected colonies were round, convex with a diameter of 2 – 3 mm, greyish black color with a clear circle (halo).

**Sensory evaluation**

Sensory parameters, including appearance, aroma, color, taste, and texture were evaluated using 30 untrained sensory panelists randomly selected among students and this method was adopted from Sęczyk, Świeca and Gawlik-Dziki (2016). The panelists were initially presented with the little tuna floss samples in identical but labeled containers with a three-digit code for their evaluation. Before the sensory session, the tuna floss samples were prepared in triplicate in a randomized permutation. This study used a hedonic test with a 9-point scale to obtain the acceptability score of the little tuna floss products. The likeness scale was arranged in accordance with the above sensory parameters was as follows: 1 = dislike extremely; 2 = dislike very much; 3 = dislike moderately; 4 = dislike slightly; 5 = neither like nor dislike; 6 = like slightly; 7 = like moderately; 8 = like very much; 9 = like extremely.

**Statistical analysis**

The experimental design applied in this study was a completely randomized design (CRD). All measurements were performed in triplicate. Data were expressed as the mean values ± standard deviation (SD). The differences were determined using a one-way analysis of variance (ANOVA), followed by Duncan’s test. The significant difference was established at p <0.05 using SPSS, Version 27, statistical software program (SPSS Inc., Chicago, Ill., USA).

**RESULTS AND DISCUSSION**

**Nutritional attributes**

*Proximate composition* The proximate composition of little tuna floss (as a control) and floss samples incorporated with banana blossom with different levels (12.5 – 50%) is presented in Table 2. Results showed a significant (p <0.05) decreased in protein and fat contents of treated tuna floss samples with the increase in the percentage of banana blossom.
Interestingly, the histamine contents of treated adults. The high energy level and protein content in the FAO/WHO/UNU (2007) samples (285.28 ± 0.14 kcal) were higher than the 200 mg kg⁻¹ allowable limit recommended by the FAO/WHO (2012), which would not cause an adverse effect. This histamine limit is also under the European Union Regulation (EC) No 2073/2005 (EC, 2005) for fishery products. It can be inferred that all treated tuna floss samples are safe as a food product.

### Amino acids composition

Table 3. presents the profile of amino acids found in the control sample and the tuna floss incorporated with different concentrations of banana blossom. Results exhibited that glutamic acid (Glu) was the most dominant amino acid in all little tuna flosses. However, 15-amino acid analyzed in the control sample was a higher amount of total amino acids both essential and non-essential amino acids than found in the treated tuna floss products. This is due to the lower amino acids content observed in banana blossom incorporated with tuna floss samples (Table 3). In terms of essential amino acids, the control and treated samples showed higher values compared to the non-essential amino acids. Furthermore, the content of essential amino acids, including arginine, histidine, isoleucine, leucine, lysine, phenylalanine, threonine, and valine, found in the treated tuna floss samples did not meet the standard recommended by Food and Agriculture Organisation/World Health Organisation/United Nations.

### Table 2 Proximate composition (g 100g⁻¹), energy value (Kcal) and histamine content (mg kg⁻¹) of the control sample and added little tuna floss with banana blossom.

| Parameter          | Control little tuna floss | Banana blossom addition | SNI* |
|--------------------|---------------------------|--------------------------|------|
|                    | 12.5%                     | 25%                      | 37.5%| 50%  |
| Moisture           | 16.03±0.01a               | 16.45±0.38a              | 16.74±0.27a | 17.23±0.16b | 17.39±0.06b | <7   |
| Crude protein      | 43.08±1.14c               | 30.27±1.41c              | 29.51±1.24c | 28.68±0.30c | 28.13±0.16c | >15  |
| Crude fat          | 18.70±0.31c               | 18.02±0.07e              | 16.62±0.06b | 15.20±0.18b | 14.79±0.08b | <30  |
| Ash                | 4.12±0.03b                | 4.45±0.11a               | 4.52±0.06a  | 4.78±0.22a  | 5.68±0.13b  | >7   |
| Crude fibre        | 1.8±0.42a                 | 2.6±0.21a                | 2.8±0.50b   | 3.1±0.43b   | 3.5±0.59c   | <1   |
| Carbohydrate       | 15.07±0.58c               | 27.81±1.04b              | 29.61±0.43c | 31.11±0.27d | 31.01±0.20d | -    |
| Energy             | 271.66±1.01a              | 303.69±1.84d             | 297.53±0.52c | 289.92±0.25b | 285.28±0.14b | -    |
| Histamine          | 319.48±1.12c              | 144.16±1.18d             | 110.56±1.68c | 85.09±1.68b | 52.25±1.35a | <200** |

Note: Values are given as mean ± standard deviation from triplicate determinations (n = 3). Different superscript letters in the same row indicate significant differences (p < 0.05). * SNI 01-3707-1995 is used for fish floss specification. ** Histamine limit referred to FAO/WHO (2012).
University (FAO/WHO/UNU) for children and adult humans (FAO/WHO/UNU, 2007).

Nevertheless, fish floss is a non-staple food product like bread, noodle, and rice, which is routinely consumed in large quantities to provide adequate energy, but it is generally combined with other staple foods to enhance nutritional values.

**Fatty acids composition**

The effect of the formulation of tuna floss with banana blossom on the fatty acid composition is presented in Table 4. The content of fatty acids in the tuna floss samples, both control and treated samples with different levels of banana blossom, varied considerably (p < 0.05). The most abundant fatty acids in all treated floss samples were oleic acid (C18:1) (34.03 – 40.63%), followed by palmitic acid (C16:0) (27.19 – 31.84%) and linoleic acid (C18:2) (11.20 – 13.30%), but these fatty acids were slightly lower compared to the control sample’s fatty acids. It might be due to a low-fatty acids content observed in the banana blossom (0.39 – 1.28%). In addition, all tuna floss incorporated with banana blossom had lower content of saturated (33.84 – 41.44%), monounsaturated (34.37 – 41.00%), and polyunsaturated (13.72 – 16.29%) fatty acids than those contained in the control. However, among the identified fatty acids, omega-3 (n-3) and omega-6 (n-6) fatty acids seem to be the most important, due to their multiple biological roles, such as reducing oxidative stress, influencing the inflammatory cascade, presenting neuroprotection, and cardiovascular protection. The total n-6 and n-3 in the treated samples ranged at around 11.28 – 13.40% and 2.44 – 3.23% respectively, with an n-6/n-3 ratio of about 3.89 – 4.62%. These values within the range (1 – 5) of omega-6 and omega-3 ratio per day recommended by some food experts that should be consumed to prevent undesirable diseases related to the lack of essential fatty acids intake (EFSA, 2010). Moreover, amongst n-3 fatty acids, eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA) are required essentially by the human body and approximately 0.2-2.0 g/day recommended by most health organizations (Desai et al., 2018), and all formulated tuna floss samples met the requirements.

**Heavy metal content**

Heavy metals analyzed in this study are cadmium (Cd), lead (Pb), arsenic (As), mercury (Hg), and tin (Sn). These are classified as toxic metals and endanger human health if the total content of the metals exceeds the recommended exposure limits (Sajib et al., 2014; Lukáčová et al., 2014; Timoracká, Vollmannová and Ismael, 2017). As presented in Table 5, the formulation of tuna floss with different levels (12.5 – 50%) of banana blossom had a wide range of metal element compositions such as Cd (<16×10⁻⁴ – 4 mg kg⁻¹), Pb (<20×10⁻⁴ – 4 mg kg⁻¹), As (8×10⁻⁴ – 4 mg kg⁻¹), Hg (1×10⁻² – 4 mg kg⁻¹) and Sn (36.15 – 37.67 mg kg⁻¹).

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**Table 3** Amino acids composition (mg g⁻¹ protein) of control sample and tuna floss incorporated with banana blossom.

| Amino acids | Banana blossom | Control tuna floss | Banana blossom addition | RDA* |
|-------------|----------------|-------------------|-------------------------|------|
|             | 12.5%          | 25%               | 37.5%                   | 50%  |
| Ala         | 0.88±0.00      | 12.60±0.04        | 10.43±0.03              | 6.92±0.03 |
| Arg         | 1.36±0.01      | 32.81±0.05        | 26.73±0.07              | 16.05±0.04 |
| Asp         | 1.58±0.00      | 34.90±0.13        | 28.94±0.07              | 20.54±0.10 |
| Glu         | 2.08±0.02      | 55.88±0.17        | 46.82±0.12              | 32.25±0.08 |
| Gly         | 1.05±0.01      | 24.43±0.05        | 20.34±0.03              | 12.76±0.01 |
| His         | 0.62±0.01      | 25.31±0.10        | 19.71±0.09              | 13.22±0.06 |
| Ile         | 0.94±0.01      | 24.29±0.08        | 19.81±0.05              | 11.93±0.04 |
| Leu         | 1.11±0.01      | 40.84±0.15        | 33.57±0.08              | 20.25±0.08 |
| Lys         | 0.78±0.01      | 32.12±0.10        | 26.41±0.08              | 18.57±0.06 |
| Phe         | 0.76±0.01      | 26.69±0.15        | 21.02±0.03              | 13.63±0.00 |
| Pro         | 0.90±0.01      | 15.14±0.05        | 12.76±0.04              | 8.11±0.03  |
| Ser         | 1.40±0.01      | 24.42±0.08        | 19.47±0.06              | 11.53±0.03 |
| Thr         | 0.86±0.01      | 28.99±0.09        | 23.60±0.09              | 14.09±0.06 |
| Tyr         | 0.67±0.00      | 16.27±0.07        | 13.67±0.03              | 8.25±0.04  |
| Val         | 0.94±0.01      | 27.32±0.09        | 22.24±0.09              | 13.98±0.02 |
| TEAA*       | 8.05           | 254.56            | 206.76                  | 129.97    |
| TNEAA       | 7.89           | 167.37            | 138.76                  | 92.12     |

Note: Values are given as mean ± standard deviation from triplicate determinations (n = 3). Different letters in the same row indicate significant differences (p < 0.05). * RDA: Recommended dietary allowance for children and adult humans by FAO/WHO/UNU (2007). ** TEAA: total essential amino acids (arginine, histidine, isoleucine. leucine, lysine, phenylalanine, threonine and valine). *** TNEAA: total non-essential amino acids (aspartic acid, glutamic acid, serine, tyrosine, glycine and alanine).
### Table 4 Fatty acids profile (%) of control tuna floss and treated tuna floss with addition of banana blossom.

| Fatty acids | Banana blossom | Control tuna floss | Banana blossom addition |
|-------------|-----------------|--------------------|------------------------|
| C6:0        | n.d.            | 0.04 ±0.000b       | 0.03 ±0.001a           |
|             |                 | 0.04 ±0.000b       | n.d.                   |
| C8:0        | n.d.            | 0.38 ±0.007c       | 0.37 ±0.006c           |
|             |                 | 0.44 ±0.009d       | 0.23 ±0.001b           |
| C10:0       | n.d.            | 0.25 ±0.002c       | 0.26 ±0.003c           |
|             |                 | 0.33 ±0.006d       | 0.16 ±0.002b           |
| C12:0       | n.d.            | 2.15 ±0.051c       | n.d.                   |
|             |                 | 2.68 ±0.063d       | 1.35 ±0.005b           |
| C14:0       | 0.02 ±0.000     | 1.71 ±0.014c       | 1.76 ±0.041a           |
|             |                 | 1.85 ±0.012b       | 1.20 ±0.012b           |
| C15:0       | n.d.            | 0.05 ±0.000b       | 0.04 ±0.000a           |
|             |                 | 0.04 ±0.004b       | 0.04 ±0.001c           |
| C16:0       | 0.92 ±0.010     | 32.52 ±0.056c      | 31.84 ±0.43c           |
|             |                 | 28.83 ±0.115b      | 27.47 ±0.224a          |
| C16:1       | 0.02 ±0.000     | 0.19 ±0.000        | 0.17 ±0.004            |
|             |                 | 0.16 ±0.001        | 0.16 ±0.001            |
| C17:0       | n.d.            | 0.12 ±0.000        | 0.10 ±0.001            |
|             |                 | 0.10 ±0.000        | 0.10 ±0.001            |
| C17:1       | n.d.            | 0.04 ±0.001        | 0.03 ±0.000            |
|             |                 | 0.03 ±0.001        | 0.02 ±0.000            |
| C18:0       | 0.12 ±0.000     | 4.53 ±0.004c       | 4.31 ±0.061d           |
|             |                 | 3.93 ±0.004c       | 3.68 ±0.024b           |
| C18:1n9     | 0.39 ±0.001     | 41.99 ±0.020c      | 40.63 ±0.323d          |
|             |                 | 35.86 ±0.001c      | 34.97 ±0.139b          |
| C18:2n6     | 1.28 ±0.000     | 14.02 ±0.008d      | 13.30 ±0.106c          |
|             |                 | 12.42 ±0.001b      | 11.25 ±0.042a          |
| C18:3n3     | 0.36 ±0.000     | 2.45 ±0.000c       | 2.17 ±0.023c           |
|             |                 | 2.28 ±0.002d       | 1.68 ±0.008b           |
| C20:0       | 0.24 ±0.003     | 0.38 ±0.006d       | 0.35 ±0.007c           |
|             |                 | 0.30 ±0.002b       | 0.29 ±0.007b           |
| C20:1       | 0.14 ±0.000     | n.d.               | 0.17 ±0.004            |
|             |                 | 0.16 ±0.000        | 0.16 ±0.002            |
| C20:4n6     | n.d.            | 0.11 ±0.001c       | 0.10 ±0.002b           |
|             |                 | 0.12 ±0.002d       | 0.12 ±0.003d           |
| C20:5n3     | n.d.            | 0.20 ±0.003c       | 0.18 ±0.004a           |
|             |                 | 0.19 ±0.002b       | 0.21 ±0.003c           |
| C21:1       | n.d.            | 0.18 ±0.001        | n.d.                   |
|             |                 | n.d.               | n.d.                   |
| C22:6n3     | n.d.            | 0.64 ±0.012c       | 0.55 ±0.008b           |
|             |                 | 0.76 ±0.006d       | 0.67 ±0.016c           |
| C24:0       | 0.02 ±0.000     | 0.13 ±0.002        | 0.10 ±0.001            |
|             |                 | 0.10 ±0.002        | 0.12 ±0.000            |
| Σ SFA       | 1.38 ±0.005     | 42.25 ±0.031c      | 41.44 ±0.443d          |
|             |                 | 38.63 ±0.030c      | 34.65 ±0.197b          |
| Σ MUFA      | 0.62 ±0.001     | 42.40 ±0.019c      | 41.00 ±0.323d          |
|             |                 | 36.21 ±0.001c      | 35.31 ±0.139b          |
| Σ PUFA      | 1.64 ±0.000     | 17.42 ±0.009d      | 16.29 ±0.143c          |
|             |                 | 15.77 ±0.008b      | 13.93 ±0.073a          |
| Σ n6        | 1.28 ±0.000     | 14.13 ±0.004d      | 13.40 ±0.004c          |
|             |                 | 12.54 ±0.001b      | 11.37 ±0.004a          |
| Σ n3        | 0.36 ±0.000     | 3.29 ±0.007c       | 2.90 ±0.011c           |
|             |                 | 3.23 ±0.005d       | 2.56 ±0.009b           |
| PUFA/SFA    | 6.54 ±0.025     | 2.27 ±0.001b       | 2.16 ±0.042a           |
|             |                 | 2.25 ±0.003b       | 2.21 ±0.024ab          |
| EPA+DHA     | n.d.            | 0.84 ±0.015c       | 0.73 ±0.012b           |
|             |                 | 0.95 ±0.009d       | 0.88 ±0.019c           |
| n6:n3       | 3.56            | 4.29               | 4.62                   |

Note: Values are given as mean ± standard deviation from triplicate determinations (n = 3). Different superscript letters in the same row indicate significant differences (p <0.05).

### Table 5 Heavy metals content (mg.kg⁻¹) of control and added tuna floss samples with banana blossom.

| Elements | Control little tuna floss | Banana blossom addition | SNI* |
|----------|----------------------------|-------------------------|------|
|          |                            | 12.5%                   | 25%  | 37.5% | 50%  |      |
| Cd       | <0.0016                    | <0.0016                 | <0.0016 | <0.0016 | <0.05 |
| Pb       | <0.0020                    | <0.0020                 | <0.0020 | <0.0020 | <2.00 |
| As       | <0.0008                    | <0.0008                 | <0.0008 | <0.0008 | <1.00 |
| Hg       | <0.0001                    | <0.0001                 | <0.0001 | <0.0001 | <0.05 |
| Sn       | 27.53±0.41                 | 36.15±0.120b            | 34.21±0.10b | 37.67±0.20b | 37.3±0.20b | <40.00 |

Note: Values are given as mean ± standard deviation from triplicate determinations (n = 3). Different superscript letters in the same row indicate significant differences (p <0.05). *SNI 01- 3707-1995 is used for fish floss specification.
Compared to the control treatment, all identified elements were relatively the same in the contents except amount of Sn, which the content of Sn increased significantly (p <0.05). However, all heavy metals contained in the treated samples agreed with the Indonesian National Standard for fish floss specification (SNI 01-3707, 1995). It indicates that the tuna floss incorporated with a banana blossom is potentially safe for human beings.

**Bacterial test**

Table 6. shows the bacterial counts for all tuna floss samples evaluated with different tests, including total count, *Salmonella* sp., *E. coli*, *S. aureus*, and total coliform. For total bacterial count, a non-selective media was used, and the results presented that no significant difference (p >0.05) was exhibited in the total bacterial count and slightly higher compared to scad fish (*Decapterus* sp.) floss (Kasmiat et al., 2020). However, all treated samples did not exceed the maximum permissible limits of microbial contamination in food products regulated by the Indonesian National Standard (SNI 7988, 2009). The identification of *Salmonella* sp., *E. coli*, *S. aureus* in the tuna floss samples using specific media showed no growth (negative) after incubation and these results agreed with the requirement for microbial contamination in the dried fish product (SNI 7988, 2009). Furthermore, all treated samples, both augmented and un-augmented with banana blossom, showed a similar count (<3 MPN.g⁻¹) to the total coliform. These results revealed that the total coliform in all tested samples were within the acceptable microbial quality of the Indonesia National Standard. From those microbial evaluations, the tuna floss product used in the present study agrees with the SNI 7988: 2009.

**Sensory attribute**

Sensory evaluation is a suitable tool for developing food products with assessing consumer’s acceptance (Anang et al., 2018; Fiorentini, Kinchla and Nolden, 2020; Witczak, Jaworska and Witczak, 2020).

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**Table 6** Bacterial count of control tuna floss and tuna flosses added with banana blossom.

| Bacterial test                  | Control tuna floss | Banana blossom addition | SNI* |
|--------------------------------|--------------------|-------------------------|------|
|                                | 12.5%              | 25%                     | 37.5%| 50%   |
| Total count (cfu.g⁻¹)           | 3.4 × 10⁴          | 1.6 × 10⁴               | 3.3 × 10⁴ | 5.2 × 10⁴ | 1.1 × 10⁴ | <1× 10⁵ |
| *Salmonella* sp. (cfu.g⁻¹)      | negative           | negative                | negative | negative | negative | negative |
| *E. coli* (cfu.g⁻¹)             | negative           | negative                | negative | negative | negative | <3      |
| *S. aureus* (cfu.g⁻¹)           | negative           | negative                | negative | negative | negative | <1×10² |
| Total coliform (MPN.g⁻¹)        | <3                 | <3                      | <3     | <3     | <3      | <10     |

Note: *SNI 7988: 2009: maximum permissible limits of microbial contamination in food products.

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**Figure 2** Physical appearance and sensory attributes of control tuna floss and tuna floss incorporated with different levels (12.5 – 50%) of banana blossom. The same letters denote the lack of statistically significant differences between the results at p <0.05 (n = 30).
Assessment of sensory attributes (appearance, aroma, color, taste, texture, and overall quality) of tested tuna floss by the panelist is depicted in Figure 2. Generally, the addition of banana blossom at the level from 12.5% to 50% to tuna flosses had a significant influence (p < 0.05) on its sensory characteristics and consequently on consumer acceptance. Amongst these formulations, the 37.5% added floss showed the highest scores in almost assessed attributes compared to other treatments and control samples.

In addition, the mean score of overall acceptability in the tested floss samples was approximately 7.11, which implies that the treatment of tuna flosses is preferred moderately among the consumers. This acceptability value agreed with the requirement recommended by the Indonesian National Standard (5) (SNI 01-3707, 1995), and even much higher than the score accepted by the SNI. Also, these results are in accordance with the studies from Candra and Tunoq (2018), Puspita, Kartikaningsih and Dayuti (2019) and Novidiyanto et al., (2020) used the addition of banana flowers into snakehead (Channa striata), Indian scad (Decapterus russelli) and chicken floss products, respectively.

**CONCLUSION**

Taken together, the addition of tuna floss with different levels (12.5% – 50%) of banana blossom reduced histamine level and increased the dietary fibre content in the little tuna floss samples. In addition, microbial and toxic elements were permissible limits regulated by the Indonesian National Standard (SNI) in all the treated tuna flosses. Our results suggested that 37.5% of banana blossom incorporated into the tuna flosses was selected as an appropriate formula due to its attributes, such as being more acceptable for overall sensory evaluation, high contents of EPA+DHA, and dietary fibre, as well as a low histamine content. However, our further research concerns the nutritional value of the selected tuna floss sample, especially on the enrichment of essential amino acids composition through fortification strategy. Also, functional properties like antioxidant and antimicrobial activities will be further studied.

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Contact Address:
Hartati Kartikaningsih, Department of Fishery Product Technology and Bioseafood Research Group, Faculty of Fisheries and Marine Science, Universitas Brawijaya, Jl. Veteran 65145, Malang, Indonesia, Tel.: +62341551611, E-mail: hartatikartikan@ub.ac.id
ORCID: https://orcid.org/0000-0002-5124-1512

Yahya, Department of Fishery Product Technology and Bioseafood Research Group, Faculty of Fisheries and Marine Science, Universitas Brawijaya, Jl. Veteran 65145, Malang, Indonesia, Tel.: +62341551611, E-mail: yahya.mp@ub.ac.id
ORCID: -

Yuniar Tri Hartita, Department of Fishery Product Technology, Faculty of Fisheries and Marine Science, Universitas Brawijaya, Jl. Veteran 65145, Malang, Indonesia, Tel.: +62341551611, E-mail: trihartitaunyiar@gmail.com
ORCID: https://orcid.org/0000-0002-6349-6774

Abdul Aziz Jaziri, Department of Fishery Product Technology and Bioseafood Research Group, Faculty of Fisheries and Marine Science, Universitas Brawijaya, Jl. Veteran 65145, Malang, Indonesia, Tel.: +62341551611, E-mail: azizizajiri@ub.ac.id
ORCID: https://orcid.org/0000-0001-7121-4055

Wahidu Zaman, Department of Food Engineering and Tea Technology, Shahjalal University of Science and Technology, Sylhet-3114, Bangladesh, E-mail: wahidanfl@yahoo.com / wahid-ttc@sust.edu
ORCID: https://orcid.org/0000-0003-1513-7301

Rovina Kobun, Faculty of Food and Nutrition, Universiti Malaysia Sabah, Jalan UMS, 88400 Kota Kinabalu, Sabah, Malaysia, Tel.: +6088320000, E-mail: rovinarub@ums.edu.my
ORCID: https://orcid.org/0000-0002-4985-9145

*Nurul Huda, Faculty of Food and Nutrition, Universiti Malaysia Sabah, Jalan UMS, 88400 Kota Kinabalu, Sabah, Malaysia, Tel.: +6088320000; Department of Food Science and Technology, Faculty of Agriculture, Universitas Sebelas Maret, Jl. Ir. Sutami, Surakarta, 57126, Central Java, Indonesia,
E-mail: drmurullah@ums.edu.my
ORCID: https://orcid.org/0000-0001-9867-6401

Corresponding author: *