ANTIOXIDANT AND ANTIMICROBIAL ACTIVITIES OF HYDROPHILIC EXTRACTS FROM SHALLOT AND GARLIC BULBS, AND THEIR EFFECTS ON ROUND SCAD DURING ICED STORAGE

HUYNH NGUYEN DUY BAO*, PHAM THI HIEN, VU LE QUYEN

Faculty of Food Technology, Nha Trang University, 02 Nguyen Dinh Chieu street, Nha Trang city 650000, Vietnam.
*Corresponding author: hndbao@ntu.edu.vn, hndbao@yahoo.com

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Abstract
The present study aimed to investigate the in vitro antioxidant and antimicrobial activities of hydrophilic extracts prepared from shallot and garlic bulbs, and their effects on round scad during iced storage. Both the shallot and garlic extracts exhibited antioxidant activity concerning DPPH radical scavenging, total reducing power, and lipid peroxidation inhibition. The extracts also had antimicrobial activity against both Gram-positive (Bacillus cereus, Staphylococcus aureus) and Gram-negative (Escherichia coli, Salmonella typhimurium) pathogenic strains. Effects of treatment with the extracts on quality of round scad were evaluated by monitoring the total lipid hydroperoxides (HPO), thiobarbituric acid reactive substances (TBARS), total volatile basic nitrogen (TVB-N), total viable counts (TVC), and sensory characteristics. The spoilage indicators (HPO, TBARS, TVB-N, TVC) of the round scad treated with either shallot or garlic extracts were significantly suppressed during the ice storage period (P < 0.05). Round scad treated with either shallot or garlic extracts kept their natural sensory characteristics accepted for food-grade longer than 4 days of ice storage, compared with the control (without treatment). These results clearly show that the shallot and garlic extracts are potential natural preservatives, which can extend shelf-life of iced round scad.

Keywords: antimicrobial activity, lipid oxidation, natural antioxidant, fish spoilage; spices

Introduction
In recent years, consumers tend to eat fresh fish rather than processed or frozen ones, while the shelf-life of fresh fish is normally short. The spoilage of fresh fish is mainly due to enzymatic autolysis, microbial growth, and lipid oxidation, and thus limiting the shelf-life. It drastically reduces the product-market value, leading
to considerable financial loss. To extend the shelf-life of fresh fish, the use of preservatives has been believed traditionally to be virtually the only efficacious way. However, the improper use of preservatives can lead to harmful effects on human health (Amit et al., 2017). For this reason, the use of natural preservatives should be a promising designation to extend the shelf-life of fresh fish, thereby minimizing economic loss.

Scad (*Decapterus* ssp.) is a coastal pelagic fish found in the Pacific, Atlantic, and Indian Oceans. The species are substantial sources of food and thus play an important role in the economy of many countries. Scad lipids contain a significant amount of polyunsaturated omega-3 fatty acids (PUFAs) which have been reported to have positive effects on various cardiovascular diseases, including blood clotting and high blood pressure (Metillo and Aspiras-Eya, 2014; Shahidi and Ambigaipalan, 2018). However, because of the presence of PUFAs, the lipids are highly vulnerable towards oxidation. Lipid oxidation is one of the major causes of spoilage of fish and has a detrimental effect on colour, flavour, texture and nutritional value (Bao and Ohshima, 2014). On the other hand, decreased nutritional value, changed texture, discoloration, and off-odour development in fish may also result from bacterial growth (Kuley et al., 2017; Odeyemi et al., 2018). Hence, bacterial growth and lipid oxidation are processes that restrict the shelf-life of fish.

Natural preservatives generally originate from microorganisms, plants, and animals. Mei et al. (2019) suggested that plant extracts might have broad application prospects in the preservation of fish. Numerous *in vitro* studies have been conducted to evaluate the antioxidant and antimicrobial activities of plant extracts. However, only a few studies have been carried out to assess the antioxidant and antimicrobial activities of plant extracts in fish preservation (Pezeshk et al., 2015; Olatunde and Benjakul, 2018; Gokoglu, 2019). Plant extracts for food application are customarily prepared from fruits, vegetables, and other edible natural materials (Sultanbawa, 2014; Weng and Yen, 2015; Gyawali et al., 2015; Pabón-Baquero et al., 2018; Chibane et al., 2019). Among them, the extracts prepared from shallot and garlic bulbs show potential for antioxidant and antimicrobial activities *in vitro* and applicable competency *in vivo* (Pezeshk et al., 2011; Pezeshk et al., 2013; Viswanathan et al., 2014; Mozin et al., 2015; Raeisi et al., 2016; Octaviani et al., 2019). Both shallot and garlic are known to have dietary and therapeutic benefits (Santhosha et al., 2013; Bisen and Emerald, 2016; Zeng et al., 2017). However, many studies were conducted to extract natural bioactive compounds using organic solvents (Viswanathan et al., 2014; Charoenchai et al., 2017; Kang et al., 2018; Octaviani et al., 2019; Thuy et al., 2020). Researchers and experts have considered the environmental impact of using organic solvents to extract natural bioactive compounds to be severe. They are also concerned with residual organic solvent in extracts to contaminate food with added the extracts. The latest development in extraction techniques mainly focuses on finding solutions to minimize the use of organic solvents (Chemat et al., 2012; Tiwari, 2015; McDonnell and Tiwari, 2017). Our previous studies have successfully prepared hydrophilic extracts of shallot and
garlic bulbs using ultrasound-assisted extraction technology (Bao et al., 2017; Phuong and Bao, 2020). The present study was therefore conducted to evaluate the antioxidant and antimicrobial activities of these extracts in vitro and their effects on round scad during iced storage.

Materials and methods

Chemicals

Thiobarbituric acid was obtained from Wako (Wako Pure Chemical Industries, Ltd., Osaka, Japan). All other chemicals were of analytical grade and were obtained from Sigma-Aldrich (Sigma Chemical Co., St. Louis, MO, USA).

Preparation of shallot and garlic extracts

Shallot (Allium ascalonicum) and garlic (Allium sativum) bulbs were purchased from BigC supermarket in Nha Trang, Vietnam. Shallot and garlic extracts were prepared by adapting our previously developed procedure (Bao et al., 2017; Phuong and Bao, 2020). The bulbs of shallot or garlic were individually ground with a Panasonic blender model MX-GM1011 (Panasonic Malaysia), and 100 g of the ground material was separately extracted with 1L of double-distilled water at 30 ± 2 °C for 15 min by ultrasound-assisted extraction technology (frequency 20 kHz). The extract was collected by centrifuging the mixture at 3.000 × g for 15 min at 4 °C and was then filtered through a Whatman grade 1 filter paper.

Treatment of fish with the extracts

Round scad (Decapterus maruadsi, 90 - 110 g body weight and 17 - 20 cm fork length) were purchased from Vinh Luong Fishing Port (Nha Trang, Vietnam). The fish was kept on ice in a Styrofoam box and transported to the laboratory at Nha Trang University. The fish was washed quickly with cold tap water (< 4 °C) within 30 sec to remove any sand and other foreign matter, and was then drained on the screen for 3 min at 4°C. Subsequently, the fish was immersed in either shallot or garlic extracts using a fish/extract ratio of 5/1.5 (w/v) at 2 - 4°C for 10 min. The control fish immersed in double-distilled water at a ratio of 5/1.5 (w/v) for 10 min at 2 - 4°C. All samples were stored in a styrofoam box containing ice using a fish/ice ratio of 1:2 (w/w). To maintain the fish/ice ratio, the molten ice was removed and replaced with an equal amount of ice.

Measurement of 2,2-diphenyl-1-picrylhydrazyl radical scavenging activity

2,2-Diphenyl-1-picrylhydrazyl (DPPH) radical scavenging activity was measured using the method of Fu et al. (2002) with a slight modification. Briefly, a 0.5 mL portion of 0.1 mM DPPH methanol solution was mixed with different volumes of extracts ranging from 0.2 to 0.5 mL, each of which was increased to a final volume of 3 mL by the addition of methanol. The mixtures were held in the dark at 25°C for 30 min. Butylated hydroxytoluene (BHT) was used as a positive standard. The absorbance of the mixtures was measured at 517 nm against a control without DPPH using a Biochrom Libra S50 UV/VIS spectrophotometer (Cambridge, UK). A calibration curve was acquired by measuring various concentrations of the authentic DPPH standard. The DPPH radical scavenging activity was expressed as millimoles of DPPH scavenged.
Measurement of total reducing power ability
Total reducing power ability was measured using the method of Oyaizu (1986) with a slight modification. Briefly, a 0.5 mL portion of 1% potassium ferricyanide was mixed with different volumes of extracts ranging from 0.2 to 0.5 mL, each of which was increased to a final volume of 1.5 mL by the addition of 0.2 M sodium phosphate buffer (pH 6.6). The mixtures were kept at 50°C for 20 min, and then 0.5 mL of 10% trichloracetic acid was added. Subsequently, 2 mL of distilled water was added, followed by the addition of 400 µL of 0.1% ferric chloride. BHT was used as a positive standard. The absorbance of the mixtures was measured at 700 nm against a control, without the extracts, using the spectrophotometer. A calibration curve was acquired by measuring various concentrations of the authentic L-ascorbic acid standard. All data were expressed as milligrams of L-ascorbic acid per millilitre of the extracts.

Measurement of lipid peroxidation inhibitory activity
Lipid peroxidation inhibitory activity was measured following the method of Bao et al. (2014). Briefly, the reaction mixture (0.5 mL) containing 0.1 mL of 50 µM metmyoglobin in phosphate buffer (50 mM, pH 7.4), 0.1 mL of 100 µM H₂O₂ in phosphate buffer (50 mM, pH 7.4), 0.2 mL of 50µM eicosapentaenoic acid in Tween 20, 0.1 mL of the extract was incubated at 37°C for 30 min. BHT was used as a positive standard. Thiobarbituric acid reactive substances (TBARS) formations in the incubation mixture were determined spectrophotometrically according to the procedure of Uchiyama and Mihara (1978).

Evaluation of antibacterial activity
Antibacterial activity was evaluated using the well diffusion method on Trypticase soy agar (TSA). Four species of foodborne pathogenic bacteria were used as references for the antibacterial assay of the extracts, including Bacillus cereus, Escherichia coli, Salmonella typhimurium, and Staphylococcus aureus. All tested microorganisms were supplied by the Microbiology Laboratory, Center for Experiments and Practices, Nha Trang University. Briefly, TSA agar plates were inoculated with bacterial strain under aseptic conditions, and wells (diameter = 5mm) were filled with 30 µl of the extracts or 30 µl of double-distilled water for control. The plates were incubated at 37°C for 24 hours under aerobic conditions. After the incubation period, the diameter of the growth inhibition zone around the wells was measured. The inhibition zones were reported in millimetres (mm).

Measurement of total phenolic compounds
The total amount of phenolic compounds was quantitatively measured following the description of Fu et al. (2002). The reaction mixture containing 4 mL of 10% aqueous Folin–Ciocalteu reagent solution, 0.2 mL of the extracts, and 0.8 mL distilled water was mixed thoroughly in a test tube. Subsequently, 5 mL of 7.5% sodium carbonate solution in distilled water was added, followed by the incubation at 25°C for 30 min. The absorbance of the mixture was measured with the spectrophotometer at 765 nm against a control without the extracts. A calibration curve was acquired by measuring various concentrations of the authentic gallic
acid standard. All data were expressed as milligrams of gallic acid per millilitre of the extracts.

**Measurement of total lipid hydroperoxides**

Total lipid hydroperoxides were measured according to the procedure of Shantha and Decker (1994) with a slight modification. Briefly, lipids were extracted from 5 g of minced fish muscle following the method of Bligh and Dyer (1959). The filtrate obtained was raised to a final volume of 10 mL by addition of chloroform/methanol (2/1). Subsequently, 50 µL of 30% ammonium thiocyanate solution was added, following by the addition of 50 µL of 2% ferrous chloride solution. The mixture was vortexed for 2-4 s and then incubated at room temperature for 5 min. The absorbance of the mixture was measured at 500 nm against a blank without the sample using the spectrophotometer. A calibration curve was acquired by measuring various concentrations of the authentic cumene hydroperoxide standard. All data were expressed as nmol of cumene hydroperoxide per gram of the fish muscle.

**Measurement of thiobarbituric acid reactive substances**

Thiobarbituric acid reactive substances (TBARS) were measured following the method of Uchiyama and Mihara (1978). Briefly, 0.5 g of the minced fish meat was homogenized with 4.5 mL of 1.15 % KCl solution. The reaction mixture containing 0.5 mL of the homogenate, 0.3 mL of 1 % phosphoric acid, and 1.0 mL of 0.6 % 2-thiobarbituric acid solution was mixed thoroughly in a screw-capped test tube. The mixture was subsequently incubated at 95 °C for 45 min using a water bath. After cooling to ambient temperature, 4.0 mL n-butanol was added to the test tube. The tube was vortexed and then centrifuged at 3000 rpm for 10 min. The absorbances of the mixture were measured at 535 nm and 520 nm against a blank without the sample using the spectrophotometer. The TBARS value was calculated using the difference between absorbances at 535 nm and 520 nm. A calibration curve was acquired by measuring various concentrations of the authentic 1,1,3,3'-tetraethoxypropane standard. The TBARS values are expressed in terms of malondialdehyde (MDA) equivalents.

**Measurement of total volatile basic nitrogen**

Total volatile basic nitrogen (TVB-N) was measured according to described in the Commission Regulation (EC) No 2074/2005 of 5 December 2005, Chapter III, “Determination of the concentration of TVB-N in fish and fishery products” (2005).

**Determination of total viable count**

The fillets with skin were separated from whole round scad and minced under aseptic conditions. A portion of 25 g the minced fish was mixed with 225 mL of sterilized saline peptone solution (0.1% w/v peptone, 0.9% w/v NaCl) in a stomacher for one minute. Successive 10-fold dilutions were made as required. Total viable count (TVC) was performed on Plate Count Agar (Merck, Germany) incubated at 30°C for 72 h according to ISO 4833-2 (2013). The results are
expressed as decimal logarithmic average values of colony-forming units per gram (log cfu/g).

**Sensory evaluation**

Sensory quality of round scad was evaluated according to the sensory scheme (Table 1) for bluefish described in the Council Regulation (EC) No 2406/96 of 26 November 1996 (1996) using a five-member panel. Panelists were trained to assess according to a scale from 0 to 3, in which a score of 2.7 to 3.0 = extra quality (E); a score of 2.0 to 2.6 = good quality (A); a score of 1.0 to 1.9 = fair quality (B); a score of 0.0 to 0.9 = poor quality (not admitted).

**Table 1.** EU scheme for the sensory evaluation of bluefish freshness.

<table>
<thead>
<tr>
<th>Parts of fish inspected</th>
<th>Criteria</th>
<th>Freshness category</th>
<th>Not admitted</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skin</td>
<td>Bright pigmentation, bright, shining iridescent colours; clear distinction between dorsal and central surfaces</td>
<td>Loss of lustre and shine; duller colours; less difference between dorsal and ventral surfaces</td>
<td>Dull, lustreless, insipid colours; skin creased when fish curved</td>
</tr>
<tr>
<td>Skin mucus</td>
<td>Aqueous, transparent</td>
<td>Slightly cloudy</td>
<td>Milky</td>
</tr>
<tr>
<td>Consistency of flesh</td>
<td>Very firm, rigid</td>
<td>Fairly rigid, firm</td>
<td>Slightly soft</td>
</tr>
<tr>
<td>Gill covers</td>
<td>Silvery</td>
<td>Silvery, slightly red or brown</td>
<td>Brownish and extensive seepage of blood from vessels</td>
</tr>
<tr>
<td>Eye</td>
<td>Convex, bulging; blueblack bright pupil, transparent ‘eyelid’</td>
<td>Convex and slightly sunken; dark pupil; slightly opalescent cornea</td>
<td>Flat; blurred pupil; blood seepage around the eye</td>
</tr>
<tr>
<td>Gills</td>
<td>Uniformly dark red to purple. No mucus</td>
<td>Less bright colour, paler at edges. Transparent mucus</td>
<td>Becoming thick discoloured opaque mucus</td>
</tr>
<tr>
<td>Smell of gills</td>
<td>Fresh seaweed; pungent; iodine</td>
<td>No smell or seaweed. Neutral smell</td>
<td>Slightly Sulphureous fatty smell, rancid bacon cuttings or rotten fruit</td>
</tr>
</tbody>
</table>
Statistical analyses
All the experiments were performed in triplicates. Means and standard deviations for all multiple measurements were calculated using Microsoft Excel 2013. Statistical difference between the samples at p<0.05 was determined by a one-way ANOVA and Tukey’s Multiple Comparisons of Means using R software version 3.5.2 (http://cran.R-project.org).

Results and discussion
DPPH radical scavenging activity, total reducing power ability, and total phenolic compounds
DPPH radical scavenging activity (RSA) and total reducing power ability (RPA) of BHT and the extracts prepared from the shallot and garlic bulbs with different amounts are presented in Figure 1.

Figure 1. DPPH radical scavenging activity (a) and total reducing power ability (b) of butylated hydroxytoluene (BHT) and extracts prepared from the shallot and garlic bulbs. Data are presented as mean ± SD (n = 3). Values with different superscript letters represent a significant difference (p < 0.05).
The amounts of the extracts in the reaction mixtures ranged from 0.2 to 0.5 mL. It is evident that, in general, both the extracts displayed increased RSA and RPA by their amounts in the reaction mixture. There was no significant difference ($p > 0.05$) between these two extracts with respect to RSA (Figure 1a). However, the RPA of shallot extract was higher ($p < 0.05$) than that of garlic extract at the same concentration (Figure 1b). Both RSA and RPA of the extracts were significantly lower ($p < 0.05$) than that of BHT (1 mg/mL) at the same volume.

Antioxidant activity of extracts from shallot and garlic has been investigating by many researchers (Benkeblia, 2005; Leelarungrayub et al., 2006; Othman et al., 2008; Chaithradhyuthi et al., 2009; Rahman et al., 2012; Bisen and Emerald, 2016). The antioxidant activity was largely contributed by the phenolic compounds (Bozin et al., 2008), especially quercetin presented in the shallot extract (Pudzianowska et al., 2012; Thuy et al., 2020) and phenolic acids presented in the garlic extract (Beato et al., 2011; Drozd et al., 2011). The results obtained in the present study showed that the concentrations of total phenolic compounds in shallot and garlic extracts were 8.82 ± 0.06 µg/mL and 14.20 ± 0.07 µg/mL, respectively.

**Lipid peroxidation inhibitory activity**

Percentage inhibition of lipid peroxidation by BHT and the extracts prepared from the shallot and garlic bulbs are presented in Figure 2. Both the extracts inhibited more than 80% of lipid peroxidation in the reaction mixtures.

![Figure 2](image)

**Figure 2.** Lipid peroxidation inhibitory activity of butylated hydroxytoluene (BHT) and extracts prepared from the shallot and garlic bulbs. Data are presented as mean ± SD ($n = 3$). Values with different superscript letters represent a significant difference ($p < 0.05$).

Nuutila et al. (2003) reported that extracts from onion and garlic inhibited lipid peroxidation, and the inhibitory activity correlated positively with the radical scavenging activity of these extracts. The present study found that there was no
significant difference ($p>0.05$) between the shallot and garlic extracts concerning the lipid peroxidation inhibitory activity. The lipid peroxidation inhibitory activity of the extracts was lower ($p<0.05$) than that of BHT (1 mg/mL) at the same volume. These results confirm a positive correlation between lipid peroxidation inhibitory activity and radical scavenging activity of the extracts.

**Antimicrobial activity**

The antimicrobial activity of extracts prepared from the shallot and garlic bulbs against pathogenic bacteria is presented in Figure 3. The results showed that the extracts have antimicrobial activity against both Gram-positive (*Bacillus cereus, Staphylococcus aureus*) and Gram-negative (*Escherichia coli, Salmonella typhimurium*) pathogenic strains. Some researchers investigated the antimicrobial activity of shallot and garlic extracts (Chaithradhyuthi *et al.*, 2009; Amin *et al.*, 2009; Viswanathan *et al.*, 2014; Saenthaweesuk *et al.*, 2015). The major active compounds were essential oils, organosulfur and phenolic compounds (Benkeblia, 2004; Benkeblia *et al.*, 2005; Block, 2010), especially allicin presented in the garlic extract (Ankri and Mirelman, 1999; Rahman, 2007). The present study has shown that the antimicrobial activity of garlic extract was significantly higher ($p<0.05$) than those of shallot extract. This may probably be due to the garlic extract containing a certain amount of allicin which actively inhibits a wide range of Gram-negative and Gram-positive bacteria (Ankri and Mirelman, 1999).

![Antimicrobial activity of extracts prepared from the shallot and garlic bulbs. Data are presented as mean ± SD (n = 3). Values with different superscript letters represent a significant difference ($p < 0.05$).](image)

**Chemical, microbiological and sensory changes of round scad stored in ice**

Changes in total lipid hydroperoxides (HPO) and thiobarbituric acid reactive substances (TBARS) of round scad with treated shallot extract, garlic extract, and control during ice storage are presented in Figure 4.
Figure 4. Changes in total lipid hydroperoxides (HPO, a) and thiobarbituric acid reactive substances (TBARS, b) of round scad during iced storage. (■), The round scad was immersed in shallot extract; (▲), The round scad was immersed in garlic extract; (●), The round scad was immersed in distilled water. Data are presented as mean ± SD (n = 3). Values with different superscript letters represent a significant difference (p < 0.05).

The induction period of lipid oxidation in the muscle of round scad with treated shallot extract or garlic extract was prolonged significantly as shown in Figure 4a. Meanwhile, the accumulated amount of HPO in the muscle of fish with treated shallot extract or garlic extract was significantly (p<0.05) suppressed after 3 days of ice storage, while the HPO in the muscle of control fish increased significantly (p<0.05) after one day under similar storage conditions. Rancid odor development has a significant correlation with the HPO content of fish muscle during post-
mortem storage and HPO at a higher level of 1500 nmol/g was noted rancid odor development (Sohn et al., 2005). The present study showed that the accumulated amount of HPO in the muscle of round scad with treated either shallot or garlic extracts was virtually controlled below the level of 1200 nmol/g of fish muscle after 5 days of ice storage.

The accumulated amount of TBARS in the muscle of all round scad increased significantly after 1 day of ice storage (Figure 4b). However, the TBARS value of fish in the control group was significantly (p<0.05) higher than that in the shallot extract group and the garlic extract group during 5 days of ice storage. TBARS were measured as an indicator of secondary lipid oxidation products. The scad contains highly polyunsaturated fatty acid (Hale, 1984; Metillo and Aspiras-Eya, 2014) and the improper handling after harvesting can induce lipid oxidation. The onset of lipid oxidation in fish leads to a loss of quality. The loss of quality is usually evident in the later stages of lipid oxidation and is associated with the characteristics of flavour, colour, texture, and nutritional value (Bao and Ohshima, 2013). Therefore, the shelf-life of fresh fish is limited. Previous studies reported that TBARS values have a correlation with sensory assessments (Hoyland and Taylor, 1991; Raharjo et al., 1993), fish with TBARS above the level of 200 nmol MDA/g will probably have rancid odours (Ke et al., 1976). The present study found that the immersing round scad in either shallot or garlic extracts kept the TBARS content of the treated fish below the level of 200 nmol MDA/g after 5 days of ice storage.

Changes in total volatile basic nitrogen (TVB-N) of round scad with treated shallot extract, garlic extract, and control during ice storage are presented in Figure 5.

![Figure 5](image-url) Changes in total volatile basic nitrogen (TVB-N) of round scad during iced storage. (■), The round scad was immersed in shallot extract; (▲), The round scad was immersed in garlic extract; (●), The round scad was immersed in distilled water. Data are presented as mean ± SD (n = 3). Values with different superscript letters represent a significant difference (p < 0.05).
TVB-N is one of the most widely used measurements of the fresh fish quality because the increased TVB-N value is related to the activity of endogenous enzymes and the growth of spoilage bacteria (Howgate, 2010a; Howgate, 2010b). The increase in TVB-N in the muscle of round scad with treated either shallot or garlic extracts remarkably retarded after 5 days of ice storage, while the TVB-N in the muscle of control fish increased significantly (p<0.05) after one day under similar storage conditions (Figure 5). The limit of TVB-N for fresh fish is 25 mg N/100 g of fish muscle (Connell, 1990). In the present study, the TVB-N in the muscle of fish with treated either shallot or garlic extracts was retained below the level of 25 mg N/100 g of fish muscle after 5 days of ice storage. The lower TVB-N value of fish with treated either shallot or garlic extracts could be due to the antimicrobial effect of the extracts.

Changes in total viable count (TVC) of round scad with treated shallot extract, garlic extract, and control during ice storage are presented in Figure 6.

Figure 6. Changes in total viable count (TVC) of round scad during iced storage. (■), The round scad was immersed in shallot extract; (▲), The round scad was immersed in garlic extract; (●), The round scad was immersed in distilled water. Data are presented as mean ± SD (n = 3). Values with different superscript letters represent a significant difference (p < 0.05).

Figure 6 shows that the TVC in all of the round scad increased with storage time. The immersing round scad in either shallot or garlic extracts inhibited bacterial growth in the fish during 5 days of ice storage. The TVC of the fish treated with garlic extract were significantly (p<0.05) lower than those treated with shallot extract. Fish spoilage is more often characterized by off-odours caused by the metabolism of bacteria. When the number of microorganisms grows to more than $10^7$–$10^8$ cfu/g, significant amounts of volatile sulfur-containing compounds are produced and the spoilage becomes evident (Gram and Huss, 1996; Gram and Dalgaard, 2002). In the present study, the TVC was lower than $10^6$ cfu/g in all of
samples after 5 days of ice storage. This result may be due to the scad that was washed with cold tap water prior to treatment in the present case. Inácio et al. (2003) also reported that there was a positive effect of washing with tap water on the quality of whole scad (Trachurus trachurus) and the TVC of whole scad washed with tap water was lower than $10^6$ cfu/g after 5 days of ice storage.

Changes in sensory scores of round scad with treated shallot extract, garlic extract, and control during ice storage are presented in Figure 7.

![Figure 7](image-url)

**Figure 7.** Changes in sensory scores of round scad during iced storage. (■), The round scad was immersed in shallot extract; (▲), The round scad was immersed in garlic extract; (●), The round scad was immersed in distilled water. Data are presented as mean ± SD (n = 5).

Figure 7 shows that the sensory scores of round scad in all samples decreased with storage time. The round scad immersed in either shallot or garlic extracts retained a good quality (A) longer than 1.5 days of iced storage and was accepted for food-grade after 5 days of iced storage. Contrary to this, round scad of the control group was not accepted for food-grade after only one day under similar storage conditions. In the first-three days of iced storage, the sensory scores of round scad treated with garlic extract were slightly higher (p<0.05) than that of round scad treated with shallot extract. However, there was no significant difference between the sensory scores of round scad treated with shallot extract and round scad treated with garlic extract at the later stage of storage. The appearance of round scad in the first-three days of ice storage as shown in Figure 8. Shallot and garlic flavours of the treated fish were not recognized in this particular experiment. Volatile organosulfur compounds are major responsible for the characteristic flavours of *Allium* species, including shallot and garlic. However, they are relatively unstable and undergo rapid decomposition to produce lipophilic organosulfides such as diallyl sulfide, diallyl disulfide, and allyl methyl sulfide. The lipophilic organosulfides can be readily oxidized by hydrogen peroxide to form allicin and odourless substances (allyl methyl sulfoxide, allyl methyl sulfone, diallyl sulfone)
In this particular case, HPO formed in the fish with treated shallot extract or garlic extract might react with organosulfides of the extracts to form odourless substances. Thus, shallot and garlic flavour intensity of the treated fish decreased with storage time.

![Figure 8. Apparent changes occurred in round scad during ice storage. Shallot extract, the round scad was immersed in shallot extract; Garlic extract, the round scad was immersed in garlic extract; Control, the round scad was immersed in distilled water.](image-url)
Conclusions

The hydrophilic extracts from shallot and garlic exhibited antioxidant and antimicrobial activities in vitro and in post-mortem round scad stored on ice. The immersion of round scad in the shallot or garlic extracts effectively extended their shelf-life. Shallot and garlic bulbs are often used as elementary spices with no known toxic effects. This study clearly showed that the hydrophilic extracts prepared from shallot and garlic bulbs are promising sources of natural preservatives for fish and fish products.

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