Antioxidant Activities of Purple Rice Bran Extract and Its Effect on the Quality of Low-NaCl, Phosphate-Free Patties Made from Channel Catfish (*Ictalurus punctatus*) Belly Flap Meat

B. Min, M.-H. Chen, and B.W. Green

ABSTRACT: Purple rice bran contains high amounts of natural antioxidants that consist of water- and lipid-soluble compounds. Hexane-insoluble and hexane-soluble fractions were separated from 100% methanolic extract from purple rice bran (RBE-HI and RBE-HS, respectively). Total anthocyanin, tannin, flavonoid, and phenolics contents were determined in those fractions, and their antioxidant capacities were evaluated by measuring 1,1-diphenyl-2-picrylhydrazyl (DPPH) radical scavenging capability, oxygen radical absorbance capacity (ORAC), and iron chelating capability (ICC). RBE-HI and RBE-HS were also added to restructured patties made from minced channel catfish (*Ictalurus punctatus*) belly flap meat. Lipid oxidation, color, and/or textural properties were determined for raw and cooked patties during a 12-d storage at 4 °C. All antioxidant indices, except for ICC, of RBE-HI were significantly higher than those of RBE-HS due probably to its higher anthocyanin content (*P* < 0.05). RBE-HS showed higher ICC (*P* < 0.05). However, both fractions showed similar antioxidant activity in raw and cooked patties during storage, resulting from the complexity of antioxidant action in food systems. Textural properties (hardness, cohesiveness, chewiness, and springiness) in cooked patties with RBE-HS and RBE-HI were well maintained during storage, but changed significantly in the control (*P* < 0.05). Only RBE-HS limited microbial growth in raw patties during storage (*P* < 0.05), but its inhibitory effect was marginal because of low-dose and physical interactions with the matrix. \(L^*\) (lightness) and \(a^*\) (redness) of raw and cooked patties decreased significantly by both fractions, whereas \(b^*\) (yellowness) was significantly decreased by RBE-HI and increased by RBE-HS (*P* < 0.05). In conclusion, we suggest that purple rice bran extract is applicable to meat products as a natural preservative, but color change in the products may limit its application.

Keywords: antimicrobial activity, antioxidant activity, channel catfish belly flap meat, purple rice bran extract, textural properties
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and others 2001; Ahn and others 2007; Bozin and others 2007). Rosemary extract, which is available commercially in the United States and Europe, is extensively investigated for use at 0.02% to 1% in various meat products to prevent lipid oxidation and microbial growth (Nissen and others 2004; Fernández-López and others 2005; Sebranek and others 2005; Ahn and others 2007).

Rice bran, a byproduct of rice milling, is a constituent (approximately 10%) of the whole rice grain and consists of the bran layers (pericarp, seed coat, nucellus, and aleurone) and the germ (Rohrer and Siebenmorgen 2004). The typical color of rice bran is light brown. Rice bran is the most nutritious part of rice grain and a rich source of bioactive phytochemicals. Rice bran shows strong antioxidant activities in various food systems (Nanua and others 2000; Kim and Godber 2001) and health beneficial effects, such as reduced total plasma cholesterol, increased HDL cholesterol level, and inhibited platelet aggregation (Gerhardt and Gallo 1998; Bramley and others 2000; Cicero and Gaddi 2001; Iarivalla 2001). The high antioxidant capacity of light brown rice bran is mainly attributed to its lipophilic antioxidants, which include γ-oryzanol, tocopherols, and tocotrienols (Quereshi and others 1997; Cicero and Gaddi 2001; Wu and others 2004). These lipid-soluble antioxidants consist of a phenolic compound with hydroxyl groups, which are responsible for antioxidant activity and a hydrocarbon side chain or phytosterol, which provides these compounds with hydrophobic characteristics. Wu and others (2004) indicate that the antioxidant capacity measured by oxygen radical absorbance capacity (ORAC) value in lipophilic fraction of the rice bran is significantly greater than that in its hydrophilic fraction.

Unlike the light brown rice bran, red and purple-pigmented rice bran contains high amounts of hydrophilic phenolic compounds such as anthocyanins, in addition to the lipophilic antioxidants (Hu and others 2003). Ling and others (2001) report that the development of ROS in liver, lipid oxidation in aorta, and the area of atherosclerotic plaque is lower in rabbits fed with red and purple rice bran, compared to those fed with light colored bran. Goffman and Bergman (2004) report that the concentration of total phenolic contents in red and purple rice bran is approximately 10 times higher than in light brown rice bran, indicating that the antioxidant activity of rice bran is closely related to rice bran color. Anthocyanins, a member of classes of flavonoids, are responsible for the color of purple rice bran. Major components of anthocyanins in black rice with purple bran are cyaniding-3-glucoside and peonidin-3-glucoside and are mainly located in the aleurone layer of the rice bran (Hu and others 2003). Anthocyanins are reported to possess remarkable antioxidant activities (Satue-Gracia and others 1997; Nam and others 2006; Phlipot and others 2006) as well as anti-inflammatory properties (Hu and others 2003). Therefore, these findings suggest that compared to light-brown rice bran, purple rice bran may be a well-balanced source of both hydrophilic and lipophilic antioxidants for foods like meat products, which are complex mixtures of hydrophilic and lipophilic components. In addition, inclusion of natural extracts can provide the health-beneficial phytonutrients to the meat products.

We developed low NaCl (1%), phosphate-free restructured patties made from minced channel catfish (I. punctatus) belly flap meat using microbial transglutaminase and isolated soy protein (Min and Green 2008). A product of catfish fillet processing, belly flap meat is a low-valued meat with high fat content. Its high fat content is responsible for its limited shelf life and undesirable flavor and texture qualities in this further processed product. Thus, the inclusion of antioxidants would retard lipid oxidation in processed products, extend the shelf life, and maintain sensory quality properties of the patties during storage. The objectives of this study were to determine the phenolic compounds content, including anthocyanins, tannins, flavonoids, and total phenolics, in purple rice bran extract, to evaluate the antioxidant capacities of the purple rice bran extract, including free radical scavenging capacity and metal chelating capability, and to determine the effects of the purple rice bran extract on the development of lipid oxidation, changes in textural properties, and microbial growth in catfish patties during cold storage.

Materials and Methods

Chemicals and reagents

Trolox (6-hydroxy-2,5,7,8-tetramethylchromane-2-carboxylic acid), sodium nitrite, aluminum chloride, 2-thiobarbituric acid (TBA), ferrozine (3-[2-pyridyl]-5,6-bis [4-phenyl sulfonyl acid]-1,2,4-triazine), ammonium ferrous sulfate, 1,1-diphenyl-2-picrylhydrazyl (DPPH), fluorescein disodium (FL), dimethyl sulfoxide (DMSO), α-tocopherol, and butylated hydroxytoluene (BHT) were purchased from Sigma-Aldrich (St. Louis, Mo., U.S.A.). Kuromannin, (+)-catechin, and Folin-Ciocalteau reagent were obtained from Fluka (Milwaukee, Wis., U.S.A.). 2,2′-Azobis(2-amidinopropane) dihydrochloride (AAPH) was purchased from Wako chemicals (Richmond, Va., U.S.A.). Randomly methylated β-cyclodextrin (RMCD) was purchased from Cyclodextrin Technologies Development Inc. (Highsprings, Fla., U.S.A.). Methanol and acetone were high-performance liquid chromatography (HPLC) grade from Fisher Scientific (Fair Lawn, N.J., U.S.A.). Hexane was UltimaAR grade from Mallinckrodt Baker Inc. (Phillipsburg, N.J., U.S.A.). Rosemary extract (Fortium R10WS; water-soluble type) was obtained from Kemin Food Ingredients Inc. (Des Moines, Iowa, U.S.A.). Microbial transglutaminase (MTGase; Activia® TI) was obtained from Ajinomoto Food Ingredients (Chicago, Ill., U.S.A.) and isolated soy protein (ISP; Supro® 500E) from Solae LLC (St. Louis, Mo., U.S.A.). All other chemicals and reagents used were of reagent grade.

Preparation of rice bran extract

Rice bran extract (RBE) was prepared from a purple bran cultivar IAC600, developed by Dr. Bastos (Instituto Agronômico, Campinas, SP Brazil). The IAC600 was grown in 2005 at Beaumont (Tex., U.S.A.) using the standard cultural management practices for this region. After removing husks from the rough rice, the resulting brown rice (with bran layers and germ still intact) was stored at ~20 °C until use. The brown rice was milled using Satake Motor one-pass mill (Tokyo, Japan) and the bran fraction, including bran and germ, was sieved through an 840-micron sieve (nr 20, E.H. Sargent & Co., Chicago, Ill., U.S.A.). The rice bran was extracted with 100% methanol overnight with shaking at a bran-to-soil ratio of 1:10 under nitrogen gas at room temperature. The next day, the mixture was centrifuged at 4000 × g for 10 min at 20 °C, and then the supernatant was collected. The pellet was extracted with 100% methanol one more time for 2 h. Both supernatants were pooled and filtered through 0.45-μm polyvinylidene fluoride (PVDF) membrane (Waters Corp., Milford, Mass., U.S.A.), and then methanol was evaporated under reduced pressure (400 mTorr at ~130 °C) using cold trap (TitanTM Vapor Trap, FTS systems Inc., Stone Ridge, N.Y., U.S.A.). The 100% methanol extract was further separated into two fractions: hexane-soluble (RBE-HS) and hexane-insoluble fractions (RBE-HI). The extract was mixed with an equal volume of hexane by vortexing for 5 min and sonicating for 2 min, followed by centrifugation at 4000 × g for 10 min at 20 °C. The hexane extraction was repeated one more time. Hexane layers from both extractions were pooled and then hexane was evaporated...
under the reduced pressure to collect RBE-HS. The lower layer was lyophilized for RBE-HI. Both fractions were flushed with nitrogen gas and stored in freezer until use.

Various indexes for antioxidant activities of rice bran extracts

Rosemary extract (RE), RBE-HI, and RBE-HS were reconstituted and diluted with acidified ethanol (85% ethanol and 15% 1.5 N HCl, v/v) to determine total anthocyanin content according to the method of Abdel-Aal and Huel (1999). Total anthocyanin content was expressed as microgram kuromanin equivalents per milligram sample. Total tannin content was quantified by the quinoline assay of Sun and others (1998), using the reconstituted solution of RBE-HI, RBE-HS, and RE with 100% methanol and expressed as microgram catechin equivalents per milligram sample.

For total flavonoid content, total phenolic content, DPPH scavenging capability, ORAC values, and iron chelating capability, RBE-HI, RBE-HS, and RE were reconstituted with DMSO. Total flavonoid content was determined by the method of Zhishen and others (1999) and expressed as microgram catechin equivalents per milligram sample. Total phenolics content was analyzed by the method of Goffman and Bergman (2004) with modification. Briefly, 0.6 mL of sample or standard was diluted with DMSO appropriately and mixed with 0.25 mL of 20% Folin–Ciocalteau reagent. After 5 min, 0.5 mL of 0.5M ethanolamine was added to the mixture. After 90 min of color development at room temperature, the absorbance was measured at 760 nm against the reagent blank. Total phenolic content was expressed as microgram gallic acid equivalents per milligram sample. Methodology of Goffman and Bergman (2004) was used to assay DPPH scavenging capability by monitoring the scavenging activity of DPPH radicals by samples and was calculated by a linear regression equation between a series of concentrations of gallic acid as a standard and their absorbance and was expressed as microgram gallic acid equivalents per milligram sample.

The procedure of Huang and others (2002a, 2002b) was modified by using hydrophilic for RBE-HI and lipophilic ORAC values for RBE-HS and RE. An Infinite 200 microplate reader equipped with Magellan reader software (version 6.4; Tecan, Grödig, Austria) was used with a fluorescence filter for an excitation wavelength of 485 nm and an emission wavelength of 520 nm. RBE-HI, Trolox as a standard, FL, and AAPH were diluted with or prepared in 75 mM phosphate buffer, pH 7, whereas RBE-HS, RE, and Trolox were diluted with 7% RMCDC in 50% acetone solution (w/v). Aliquots of the diluted samples or standards (50 μL) were added to a 96-well black polystyrene flat bottom plate with a lid (Corning, Corning, N.Y., U.S.A.). FL solution (100 μL; final conc. 0.048 μM) was added to samples or standards in the 96-well plate and preincubated at 37 °C for 15 min in the microplate reader. After AAPH (50 μL; final conc. 63.4 mM) was rapidly added to each well, the plate was immediately transferred to the plate reader and agitated for 10 s prior to the first reading and each subsequent reading. Fluorescence was measured at every 2.5-min interval for 60 min for hydrophilic ORAC and 120 min for lipophilic ORAC at 37 °C. The ORAC values were calculated using a linear regression equation (AUC = \( \sum f_i/\alpha + \sum f_i/\beta \)) between a series of Trolox concentrations (\( \gamma = \alpha \times \beta \)) and the net area under the fluorescence decay curve (net AUC) (\( x \)). The area under the curve (AUC) was calculated as:

\[
AUC = (0.5 + f_1/f_1 + f_3/f_1 + f_5/f_1 + \cdots + f_i/f_1 + \cdots + f_n/f_1) \times CT
\]

where \( f_i = \) the initial fluorescence reading, \( f_i = \) the fluorescence reading at cycle \( i \), and \( CT = \) cycle time in minutes.

The net AUC was calculated by subtracting the AUC of blank from the AUC of samples or standards. ORAC value was expressed as microgram Trolox equivalents per milligram sample.

Iron chelating capability was assayed using the ferrozine method (Carter 1971). In brief, 100 μL of sample or standard were mixed with 100 μL of 1 mM ammonium ferrous sulfate solution and 2.7 mL 10% ammonium acetate buffer solution. After 5 min, 50 μL of 6.1 mM ferrozine color reagent were added to the mixture and mixed thoroughly. After 10 min, the absorbance was determined at 562 nm. Iron chelating capability was calculated by using a linear regression equation between a series of concentrations of gallic acid as a standard and their absorbance and expressed as microgram gallic acid equivalents per milligram sample.

Production of patties

Fresh channel catfish belly flap meat was purchased from an Arkansas processor. Upon receipt, the meat was washed using a pressure washer (2200 psi, Troy-Bilt Model 020295, Briggs & Stratton Power Products Group, LLC, Jefferson, Wis., U.S.A.) for 5 min to remove excessive fat and de-pigment the adhered peritoneal membrane. The washed meat was drained for around 2 h in the refrigerator before storing in vacuum-packaging bags (Cryovac® HP2700 vacuum bag, Sealed Air, Duncan, S.C., U.S.A.). The vacuum-packaged meat was stored at ~20 °C until use.

The formulation of ingredients and patty preparation conditions in this study were based on our previous study (Min and Green 2008). Belly flap meat was thawed overnight in a refrigerator and ground separately twice using a grinder (Model 722, Biro Manufacturing Co. Ltd., Marblehead, Ohio, U.S.A.) fitted with a plate with six 18-mm diameter openings. Six treatment groups were prepared with: (1) control (no antioxidant), (2) 0.02% BHT (w/w), (3) 0.1% \( \alpha \)-tocopherol (w/w), (4) 0.1% rosemary extract (RE) (w/w), (5) 0.1% RBE-HI (w/w), and (6) 0.1% RBE-HS (w/w). BHT and \( \alpha \)-tocopherol were used as references for synthetic and natural antioxidants, respectively, and RE was used as a reference for natural extract. The RBE-HI, RBE-HS, and RE were dissolved in 70% ethanol (71 mg/mL) and others were dissolved in 100% ethanol. The ground meat was mixed with sodium chloride (1%), water (10%), and antioxidants for 2 min in a mixer (Model KSM150PSOB, KitchenAid, St. Joseph, Mich., U.S.A.). Ethanol was added to the control and other treatments, if needed, to equalize the ethanol level in every treatment. MTGase–water (1:3, w/w) and ISP–water (1:4, w/w) mixtures were subsequently added to the mince and mixed for 2 min. The final concentration of MTGase and ISP were 0.1% and 1.7%, respectively. Total mixing time was standardized to 4 min and the final temperature of the meat mixture in all treatments was kept below 12 °C throughout the mixing process. Four replicate batches (500 g per batch) were prepared for each treatment. Patties (100 g; 100-mm diameter) were formed using a hamburger press (Model 1404, Univex, Salem, N.H., U.S.A.) and stored overnight at 4 °C in the refrigerator for the reaction of MTGase. For the cooked patty study, the same experimental set of patties as previously mentioned was prepared on a different day. Patties were cooked in a superheated steam oven (Model AX700R, Sharp Electronics, Mahwah, N.J., U.S.A.) at 227 °C to an internal temperature of 80 °C. Raw and cooked patties were individually packaged in oxygen-permeable zipper bags (polyethylene, 4 × 6, 2 mil.; Associate Bag Co. Milwaukee, Wis., U.S.A.) and stored at 4 °C in the refrigerator until used. Raw and cooked patties were analyzed for 2-thiobarbituric acid-reactive substances (TBARS), color values, and textural profile analysis (for only cooked patties) at 0, 3, 6, 9, and 12 d of storage. Aerobic plate count (APC), coliform plate count (CPC), and yeast
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plate (YPC) were determined for only raw patties at 0, 2, 4, 6, and 8 d of storage.

2-Thiobarbituric acid reactive substances

Raw and cooked patties were analyzed for TBARS value, as a lipid oxidation index, according to the method of Min and Ahn (2008). Briefly, 5 g of ground meat were homogenized with 15 mL DDW and 100 μL BHT solution (6% in 100% ethanol, w/v) using a homogenizer (Fisher Scientific, Pittsburgh, Pa., U.S.A.) for 15 s at speed 7. The meat homogenate (1 mL) was mixed with 2 mL TBA/TCA solution (20 mM TBA/15% trichloroacetic acid [TCA; w/v]). The mixture was incubated for 15 min in a boiling water bath. After cooling, the mixture was centrifuged at 3000 × g for 15 min. The absorbance of the supernatant was determined at 531 nm against a reagent blank. The amount of TBARS was expressed as milligrams malondialdehyde (MDA) per kilogram meat.

Color measurement

Commission Internationale de l’Eclairage (CIE) L∗ (lightness), a∗ (redness), and b∗ (yellowness) for raw and cooked patties during storage were determined using a HunterLab Miniscan XE plus colorimeter (Model 4500L, Hunter Associate Laboratory Inc., Reston, Va., U.S.A.) with 25-mm aperture, CIE D65 illuminant, and 10° standard observer. The colorimeter was calibrated against black and white reference tiles. Color values were measured on the surface of 2 patties per each batch and 6 measurements per each patty (3 on each side of the patty). Whiteness (Park 2000) was calculated by the following equations:

\[ \text{Whiteness} = 100 - [(100 - L^*)^2 + a^*^2 + b^*^2]^{1/2} \]

Texture profile analysis

An Instron universal testing machine (Model 3342, Instron Co., Norwood, Mass., U.S.A.) equipped with a 500-N load cell was used to analyze texture properties of the cooked patties during storage. Texture profile analysis (TPA) was performed using a 90-mm diameter aluminum plunger according to the methods of Bourne (2002). Two cylindrical samples (20 mm in diameter and 18 mm in thickness) were extracted from each patty and compressed through two repeated cycles with 80% compression of the initial height at a crosshead speed of 60 mm/min. There was a 5-s rest period between the 2 cycles to allow the sample to recover its height. Hardness, cohesiveness, chewiness, and springiness were calculated from the TPA curve. Hardness (N) was defined as the peak force during the first compression cycle, cohesiveness as the ratio of the positive force area under the second (A2) compression to the area under the first (A1) compression, chewiness (N·mm) as the energy required to compress the sample and the product of hardness × cohesiveness × springiness, and springiness (%) as the ratio of the height recovered during the rest period between the 1st and 2nd cycles to the original height to the sample (Bourne 2002).

Microbiological analysis

APC, CPC, and YPC were determined for raw patties using Petrifilm™ aerobic count plate, coliform count plate, and yeast/mold count plate (3M Co., St. Paul, Minn., U.S.A.), respectively. At each storage interval, a 25-g sample of the patties was homogenized with 225 mL sterile Butterfield’s phosphate buffer (0.3 mM potassium phosphate buffer, pH 7.2; 3M Co.) for 2 min at room temperature. Appropriate serial dilutions were made with sterile Butterfield’s phosphate buffer, and then 1 mL of each dilution was spread duplicate on appropriate plates for APC, CPC, and YPC. The plates for APC were incubated at 35 °C for 48 h, for CPC at 35 °C for 24 h, and for YPC at 20 °C for 5 d (AOAC 2005).

Data were expressed as log_{10} colony forming units (CFU) per gram sample.

Statistical analysis

All data were analyzed by one-way or two-way analysis of variance using the SAS program (version 9.1 for windows, SAS Inst. Inc., Cary, N.C., U.S.A.) and reported as means and standard error of the mean (SEM) in tables and means and standard deviation (SD) in figures. The analyses for all antioxidant indexes for RBE-HI, RBE-HS, and RE were performed in triplicate (n = 3). In the study using catfish patties, 4 batches for each treatment were produced for TBARS, surface color, and TPA measurements (n = 4). For microbiological analysis, 3 batches from each treatment were randomly selected and analyzed (n = 3). Student–Newman–Keuls’ multiple range test was used to compare the mean values of treatments (P < 0.05) (Kuehl 2000).

Results and Discussion

Phenolic compounds and antioxidant capacity of purple rice bran extract

Methanolic extract from purple rice bran was further extracted by hexane, resulting in hexane-insoluble and hexane-soluble fractions (RBE-HI and RBE-HS, respectively) to compare the antioxidant capacities of nonlipophilic and lipophilic components in purple rice bran extract. The lipophilic components such as tocopherols, tocotrienols, and γ-oryzanol have been well recognized as major antioxidants in typical light brown rice bran extract (Quereshi and others 1997; Cicero and Gaddi 2001). Rosemary extract (RE) was chosen as a reference of natural extract in this study because its antioxidant activity in meat products has been well demonstrated and commercially available in the United States. Carnosol and carnosic acid, phenolic diterpenes, were reported to account for around 90% of antioxidant capacity in commercially available rosemary extract (Wijeratne and Cuppett 2007).

Total anthocyanin content, total flavonoid content, and total phenolics content in RBE-HI was significantly higher than those in RBE-HS and RE (P < 0.05) (Table 1). Anthocyanins appeared to be primarily responsible for total flavonoid and total phenolics contents in RBE-HI. Anthocyanins are one of the most abundant natural colorants in nature and belong to the class of flavonoids. Hu and others (2003) indicated that the major anthocyanins found in black rice, cyanide-3-glucose, and peonidin-3-glucose, have strong antioxidant activities in an in vitro model system. Compared to those of RBE-HS and RE, the ORAC value and DPPH scavenging capability of RBE-HI was significantly higher and appeared to be closely related to its higher total anthocyanins, flavonoids, and phenolics contents (P < 0.05) (Table 1). Total tannin content was much lower than total flavonoids and total phenolics content in RBE-HI and RBE-HS, but not detected in RE. Tannins were reported to have free-radical scavenging activity (Oki and others 2002). Therefore, tannins in RBE-HI and RBE-HS may be contributing to the DPPH scavenging ability and ORAC values. However, profiles of phenolic compounds present in purple rice bran have not been fully investigated yet because most studies on the antioxidants in rice bran have focused on its lipophilic compounds. Further studies on the identification of phenolic compounds in purple rice bran are needed.

The iron chelating capability in RBE-HS was significantly higher than that in RBE-HI, but was not detected in RE (P < 0.05) (Table 1). The status of free iron ion is very important for the development of lipid oxidation in meat products because the reactivity of ferric ion is negligible (Ahn and Kim 1990). Free-iron ion catalyzes
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due to the production of hydroxyl radical (·OH) from H₂O₂ as well as the degradation of lipid hydroperoxides to produce peroxyl and alkoxyl radicals, which can initiate lipid oxidation and/or be self-degraded to the secondary products of lipid oxidation (Min and Ahn 2005). Min and others (2006) reported that the ferric ion reducing catalytic activity for catalyzing lipid oxidation, present in cooked meat. Therefore, free ionic iron is regarded as a primary catalyst for lipid oxidation in cooked meat products because other catalysts such as hemoglobin and myoglobin are denatured and degraded to release free ionic iron by heating (Min and Ahn 2005). The chelation of free ionic iron is a promising approach to retard the development of lipid oxidation in meat products because many further-processed meat products are cooked before distribution or consumption. Phenolic compounds have been considered as good chelators (Hider and others 2001) because the negatively charged phenoxide groups of deprotonated phenolic compounds chelate free ionic iron and stabilize the form of the free ionic iron. However, it is uncertain which compounds contributed to higher iron chelating capacity in RBE-HS and further study is needed.

Effect on lipid oxidation in raw and cooked patties during storage

The effect of purple rice bran extract on the development of lipid oxidation, expressed as TBARS values, in raw and cooked catfish patties during 12 d of storage at 4 °C is shown in Figure 1A and 1B.

### Table 1 — Various indexes for antioxidant activities of hexane-insoluble and hexane-soluble fractions from 100% methanol extract from purple rice bran (RBE-HI and RBE-HS, respectively) and rosemary extract (RE).a

<table>
<thead>
<tr>
<th>Sample</th>
<th>Total anthocyanin content (μg kuromain eqv./mg)</th>
<th>Total tannin content (μg catechin eqv./mg)</th>
<th>Total flavonoid content (μg catechin eqv./mg)</th>
<th>Total phenolics content (μg gallic acid eqv./mg)</th>
<th>DPPH scavenging capability (μg Trolox eqv./mg)</th>
<th>ORAC value (μg Trolox eqv./mg)</th>
<th>Iron chelating capability (μg gallic acid eqv./mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RBE-HI</td>
<td>90.05a</td>
<td>5.89a</td>
<td>50.96a</td>
<td>113.31a</td>
<td>169.95a</td>
<td>582.68a</td>
<td>28.89a</td>
</tr>
<tr>
<td>RBE-HS</td>
<td>16.77b</td>
<td>0.57b</td>
<td>14.37b</td>
<td>9.49b</td>
<td>15.04c</td>
<td>322.14b</td>
<td>40.39b</td>
</tr>
<tr>
<td>RE</td>
<td>n.d.</td>
<td>n.d.</td>
<td>2.55b</td>
<td>14.22b</td>
<td>21.85b</td>
<td>158.88c</td>
<td>n.d.</td>
</tr>
<tr>
<td>SEM</td>
<td>1.03</td>
<td>0.14</td>
<td>0.42</td>
<td>0.19</td>
<td>0.27</td>
<td>8.42</td>
<td>0.32</td>
</tr>
</tbody>
</table>

a Means with different letters (a to c) within the same column are significantly different (P < 0.05). SEM = standard error of the mean. n = 3.

b Hydrophilic ORAC values were measured for RBE-HI, and lipophilic ORAC values were measured for RBE-HS and RE.

c n.d. = not detected, indicating that the values were below minimum in the standard curve.

DPPH = 1,1-diphenyl-2-picrylhydrazyl; ORAC = oxygen radical absorbance capacity.
The initial (day 0) TBARS values for raw patties with RBE-HI and RBE-HS was significantly lower than those for control and patties with other treatments (P < 0.05) (Figure 1A). TBARS value of control sample increased significantly during storage, and at a much higher rate than in the antioxidant-treated patties (P < 0.05). The standard synthetic antioxidant, BHT, showed the strongest antioxidant activity in the raw patties during storage, inhibiting increases in the TBARS value. Increases in TBARS values of patties treated with natural antioxidants were observed throughout the storage period, but their rates of increase were significantly lower than those of the control (P < 0.05). The raw RE-treated patties showed the lowest TBARS values among natural antioxidant-treated patties after 12 d of storage. Antioxidant activity of RE was similar to BHT until day 6, but gradually lost its antioxidant capacity thereafter. In raw patties, antioxidant activity of BHI was similar to that of BHT-treated patties until day 6 and to RE-treated patties until day 9, after which its antioxidant activity decreased significantly (P < 0.05). At day 12, the TBARS value of raw RBE-HI-treated patties was the same as that of raw Toco- and BHI-HS-treated patties. In general, an antioxidant becomes an antioxidant radical after...
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off-flavor development (Min and Ahn 2005). In all cooked patties, TBARS values increased during storage, but the addition of antioxidants slowed the rate of increase. After 12 d of storage, BHT-treated patties had the lowest TBARS value, whereas RE-treated patties had the highest among antioxidant-treated patties. On day 12, TBARS values of Toco-, RBE-HI-, and RBE-HS-treated patties did not differ significantly ($P > 0.05$). The pattern of increases in TBARS values during storage differed among cooked patties treated with natural antioxidants. The TBARS value of RE-treated patties increased significantly after day 3, whereas those of RBE-HS-treated patties increased significantly only until day 6 ($P < 0.05$). During storage, TBARS values of RBE-HI- and Toco-treated patties increased gradually. In addition, the change in TBARS value during storage within treatment differed between raw and cooked patties. We attribute these differences to different modes of antioxidant action by the natural extracts given their different profiles of phenolic compounds. Heating also can change these profiles in natural extracts by inactivating heat-labile phenolic compounds. However, the antioxidant mechanisms of the components in natural extracts in real food systems have not been studied.

Antioxidant activity of RBE-HI in raw and cooked patties after 12 d of storage did not differ from those of RBE-HS and were even lower than those of RE in raw patties (Figure 1), whereas its antioxidant indexes, such as DPPH scavenging capability and ORAC values (Table 1), were much greater than those of RBE-HS and RE. Frankel and Meyer (2000) suggested limitation to the use of one-dimensional methods to predict the effectiveness of natural extracts as antioxidants in complex food systems. They indicated that test methods such as the DPPH scavenging capability and the ORAC method, which measures the free radical trapping ability of antioxidants, do not consider the complexity of antioxidant action. In real food systems, like meat products, the antioxidant action becomes more complicated and the various mechanisms, including free radical chain breaking, oxygen scavenging, metal chelation, and inhibition of oxidative enzymes, are effective. In addition, Porter (1993) suggested the paradoxical behavior of antioxidants in

![Figure 2](image-url)
food systems on the basis of phenomenological observation. He indicated that hydrophilic antioxidants are more active in bulk oil systems but lipophilic antioxidants are more active in lipid suspended in aqueous systems like an oil-in-water emulsion. Meat products, including the catfish patties in this study, are similar to lipid suspended in aqueous systems. Therefore, the higher iron chelating capability (Table 1) and hydrophobicity of RBE-HS may account for its higher effectiveness as an antioxidant in the patties, compared to RBE-HI.

Effect on changes of color and textural properties in the catfish patties

The addition of BHT, Toco, and RE did not change any color attributes ($L^*$, $a^*$, $b^*$, and whiteness) in the raw patties, compared to the control (Table 2). However, RBE-HI- and RBE-HS-treated patties had significantly lower $L^*$, $a^*$, and whiteness values compared to the control and Toco- and RE-treated patties ($P < 0.05$), and the $b^*$ value of RBE-HI-treated patties was significantly lower ($P < 0.05$). These changes were attributed to the presence of anthocyanins, natural colorants, responsible for the dark purple color of the pigmented rice bran. The different concentration of total anthocyanins of RBE-HI- and RBE-HS-treated patties (Table 1) likely caused the difference in patty color attributes between these 2 treatments. The presence of carotenoids, natural lipid-soluble colorants, identified in rice bran oil (Stoggl and others 2005) likely resulted in the increased yellowness ($b^*$ value) and decreased redness ($a^*$ value) in the RBE-HS raw patties. In addition, all color attributes in RBE-HI-treated patties changed significantly during 12 d of storage ($P < 0.05$) due probably to a loss of monomeric anthocyanins and an increase of polymeric color, resulting from the polymerization of anthocyanins during storage (Hager and others 2008). Meanwhile, cooking increased $b^*$ values, but decreased whiteness of all patties because of relative increases in pigment concentration by loss of water during cooking (Table 3). In addition, heating may increase anthocyanin polymerization (Hager and others 2008), resulting in changes of color attributes in cooked RBE-HI-treated patties.

Figure 2 shows the effect of antioxidants on changes in textural properties (hardness, cohesiveness, chewiness, and springiness) of cooked catfish patties during 12 d of storage. Initial (day 0) values of all textural parameters in all treatments did not differ. However, all textural parameters of the control patties changed during storage. Hardness of control patties increased significantly ($P < 0.05$), whereas cohesiveness, chewiness, and springiness decreased significantly after day 4 ($P < 0.05$). On the other hand, textural parameters of antioxidant-treated patties remained unchanged during storage and no differences were observed among treatments during storage. Free radicals generated in the patties can cause not only the development of lipid oxidation but also protein oxidation, resulting in denaturation of protein molecules (Min and Ahn 2005). In addition, the lipid oxidation secondary products can also cause protein denaturation, which deteriorates texture properties of meat products (Kanner 1994). Myofibrillar proteins play crucial roles in functional properties, such as water holding properties and textural properties, in meat products. Oxidative damage of proteins may cause loss of protein solubility, aggregation, and complex
formation, and consequently loss of protein network (Karel and others 1975, Estévez and others 2006). These changes can result in reduced spaces between the protein molecules in the patties, leading to greater loss of moisture during storage and consequently a harder texture in meat products. In addition, broken protein networks can cause loss of textural properties such as cohesiveness, chewiness, and springiness. Therefore, we assume that inclusion of antioxidants may prevent the development of protein oxidation in and maintain the textural properties of the patties during storage.

Effect on microbial growth in raw catfish patties

The initial (day 0) aerobic plate counts (APC), coliform plate counts (CPC), and yeast plate counts (YPC) did not differ between the control and natural extract-treated patties and the APC, CPC, and YPC increased significantly in all the patties during 8 d of storage (P < 0.05) (Figure 3). Microbial growth in the patties during storage was not inhibited by RE and RBE-HI. Several studies have shown the antimicrobial effect of RE on foodborne pathogens such as E. coli (Del Campo and others 2000; Fernández-López and others 2005; Bozin and others 2007). Del Campo and others (2000) indicated that compounds responsible for antimicrobial activities in RE are probably lipid-soluble phenolic diterpenes which are the major antioxidants of RE. This suggestion explains the absence of antimicrobial activities in RBE-HI. However, these researchers used in vitro microbiological test such as agar diffusion method and used undiluted natural extracts in their studies. The doses of natural extracts in their studies were much higher than those used in this study. In addition, it was suggested that the physical interaction among antimicrobial agents and “real” food matrix may interfere their antimicrobial activity in food systems (Fernández-López and others 2005). Microbial growth was inhibited in the RBE-HS-treated patties after 12 d of storage (P < 0.05; Figure 3A, 3B, and 3C). Lipid-soluble phenolic compounds, such as γ-oryzanol, in the RBE-HS may account for its antimicrobial activities. However, low dose and physical interaction with the matrix of the patties may restrict the antimicrobial activities of RBE-HS in the patties during storage.

Conclusions

RBE-HI showed higher total phenolics content, DPPH scavenging capability, and ORAC value, but lower iron chelating capability, compared to RBE-HS. On the other hand, both fractions showed similar antioxidant activity in raw and cooked catfish patties and maintained better textural properties of cooked patties during 12 d of storage, compared to the control. Only RBE-HS presented antimicrobial activities in raw patties during storage, but its effect was marginal due to low dose and interaction with the matrix of the patties. However, inclusion of RBE-HI and RBE-HS significantly changed color attributes in raw and cooked catfish patties. We suggest that purple rice bran extract is a good candidate as a natural preservative for extending shelf life and preventing deterioration of meat product quality. Furthermore, purple rice bran extract can add health benefits to meat products. Color change of the patty, however, should be considered and the mechanism of this color change requires further research.

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References


Rice bran extract in catfish patty...


