Efficacy of puffer fish (Takifugu rubripes) sauce in reducing hydroxyl radical damage to DNA assessed using the apurinic/apyrimidinic site method

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Abstract. Apurinic/apyrimidinic (AP) sites are frequently observed DNA lesions when cells are exposed to hydroxyl radicals. We developed a new method for measurement of the antioxidative activity of foods using the occurrence frequency of AP sites on DNA. Combined with the electron spin resonance (ESR) method as a standard method, we examined whether fish and soy sauces including puffer fish [Takifugu rubripes (Temminck et Schlegel)] sauce could protect DNA from damage caused by hydroxyl radicals. The results showed that the ratios of DNA protection by puffer fish sauce, salmon fish sauce, sandfish fish sauce (Shottsuru), colorless soy sauce, squid fish sauce (Ishiru), dark color soy sauce and light color soy sauce were 68.9, 67.0, 60.1, 49.7, 34.1, 28.2 and -4.4%, respectively. Puffer, salmon, and sandfish fish sauces showed high ratios of DNA protection against hydroxyl radicals. On the other hand, IC50 values of hydroxyl radical scavenging of the puffer, salmon, sandfish, squid fish sauces and colorless, dark and light color soy sauces were 0.20, 0.09, 4.16, 0.26% and 0.28, 0.14 and 0.18%, respectively. Though the puffer fish sauce exhibited the highest level of DNA protection among the examined samples and a high hydroxyl radical scavenging capability, a correlation between the radical scavenging capability and DNA protection against hydroxyl radicals among the examined fish and soy sauces was not found.

Introduction

Radicals are molecules that contain one or more unpaired electrons. Oxidative damage caused by oxygen-derived species, such as superoxide anion (O2·-) and H2O2, has been implicated in the initiation of cancer (1), and foods are implicated in 30% of the cases. There is considerable interest in the possibility that O2·- and H2O2 exert their toxicity by being converted into the highly reactive hydroxyl radical (•OH) in reactions that require metal ions (2). Therefore, antioxidants in the human diet are of great interest as possible protective agents for reducing oxidative damage, and a wide variety of supplements have been produced as such agents. As well, many researchers have sought radical scavengers from food products (3-6).

On the other hand, fish sauces are traditional Japanese and Asian fermented food seasonings. In Japan, ‘Shottsuru’ is made from sandfish, ‘Ishiru’ is made from squid and ‘Ikano Shoyu’ is made from sand lance fish. Also, ‘Jeotgal’ in Korea, ‘Patis’ in the Philippines, ‘Nam plaa’ in Thailand, ‘Nouc mam’ in Vietnam and ‘Yuiru’ in China are famous fish sauces from East and Southeast Asia. We have examined the high antioxidative activities of seafoods including fish sauces by examining the peroxyl and hydroxyl radical scavenging capability by using the chemiluminescence and
the electron spin resonance (ESR) methods (7-10). However these activities have not yet been examined based on a biological system.

Tamm et al (11) found for the first time that acid treatment of DNA protonated the N-glycosidic bonds of purines in purified DNA, resulting in the hydrolysis of purines from the pentose sugar without breaking the phosphodiester backbone directly. Greer and Zamenhof (12) reported that purine bases are also slowly released from DNA during incubation at a neutral pH and high temperature. Based on the determined rate constant for depurination of *Bacillus subtilis* DNA of 3x10^-11 sec^-1 per nucleotide at 37˚C and pH 7.4, Lindahl and Nyberg (13) considered that depurination of DNA may occur at a physiologically significant rate under *in vivo* conditions. It was reported that the number of depurination sites on DNA in microbial cells of *B. subtilis* (14,15) and the radioreistant bacterium *Deinococcus radiodurans* (16) increased with a marked decrease in pH value and a marked increase in temperature. In this study, we found the apurinic/apyrimidinic (AP) site phenomenon as one of the DNA lesions frequently observed when the extracted DNA is exposed to hydroxyl radicals generated from the Fenton's reaction.

We developed a new method for measurement of antioxidant activity using the occurrence frequency of AP sites on DNA, and we examined whether some seafood, especially fish and soy sauces could protect the DNA from hydroxyl radical damage *in vitro*.

**Materials and methods**

**Samples of fish and soy sauces.** The puffer fish [Takifugu rubripes (Tennminck et Schlegel)] sauce was made in our laboratory using the meat, skin and bones without the internal organs containing the poison, i.e. tetrodotoxin. The puffer fish materials were added to soybean, wheat, soy sauce koji mold, NaCl and water. These were fermented for ~1 year at room temperature. As the puffer fish sauce, we used the supernatant of the fermented original sauce mash after heat sterilization. Salmon fish sauce 3, sandfish fish sauce (Shottisuru) and squid fish sauce 1 (Ishiru) were described in our previous study (8), and extremely dark color soy sauce 2, light color soy sauce 4 and colorless soy sauce 2 were similarly described in our previous study (17).

**Chemicals.** Hydrogen peroxide (H₂O₂) was obtained from Santoku Chemical Industries Co., Ltd. (Tokyo, Japan); iron (II) sulfate heptahydrate (FeSO₄·7H₂O) was from Sigma-Aldrich Japan K.K. (Tokyo, Japan); and 5,5-dimethyl-1-pyrroline N-oxide (DMPO), as a spin trapping reagent, was from Toyko Kasei Kogyo Co., Ltd. (Tokyo, Japan). Tris(hydroxymethyl)aminomethane and deoxyribonucleic acid (DNA) sodium salt from salmon testes was resolved in TE buffer (10 mM Tris-HCl, 1 mM EDTA, 2Na, pH 7.5) at the concentration of 1 mg/ml as the DNA solution. Hydroxyl radical-damaged DNA and anti-oxidized DNA samples were prepared according to the modified method described previously (19). DNA solution (300 μl) in a 1.5-ml microtube was added to 50 μl of a 1.0-mM FeSO₄ solution to initiate the Fenton's reaction, which occurs as in the following chemical equation: Fe³⁺ + H₂O₂ → Fe²⁺ + •OH + OH⁻.

After 1 min of the Fenton's reaction, the hydroxyl radical generation, i.e. spin adduct DMPO-OH was measured using the ESR spectrometer (JES-FR30, JEOL Ltd., Tokyo, Japan). The ESR measurement conditions were as follows: output, 4 mW (9.4 GHz); magnetic field, 342.790±5 mT; modulation amplitude, 0.079 mT; response time, 0.1 sec; sweeping time, 1 min; and amplification ratio, 32-125.

**Calculation of the IC₅₀ value of hydroxyl radical scavenging.** As an indicator of the antioxidative activity, the inhibition of the hydroxyl radical peak in the ESR pattern was measured by the change of the peak height ratio of the sample compared with the inner standard manganese peak height. The lower the peak height ratio, the more inhibition of hydroxyl radical generation occurred. The value IC₅₀ was defined as the concentration of fish or soy sauce reducing the control peak height ratio of ESR to half. First, the antioxidative value was calculated using the following formula: (log Io/I) x 100; Io = peak height ratio of the control; I = peak height ratio of each concentration of the fish or soy sauce sample.

When the value of this formula reached 30.103, the I value corresponded to half-inhibition. Next, from the relationship between the antioxidative value and the concentration of the fish or soy sauce, the IC₅₀ value was calculated (18).

**Preparation of DNA sample.** DNA sodium salt from salmon testes was resolved in TE buffer (10 mM Tris-HCl, 1 mM EDTA, 2Na, pH 7.5) at the concentration of 1 mg/ml as the DNA solution. Hydroxyl radical-damaged DNA and anti-oxidized DNA samples were prepared according to the modified method described previously (19). DNA solution (300 μl) in a 1.5-ml microtube was added to 50 μl of a 1.0-mM FeSO₄ solution, 250 μl of fish or soy sauce as the antioxidant DNA sample, or pure water as a positive control, i.e. hydroxyl radical-damaged DNA sample, and immediately incubated at 37˚C for 5 min in a water bath (ThemoMax Tm-1, As One Corporation, Osaka, Japan). Next, this mixing DNA solution was added to 50 μl of an 8.8-mM H₂O₂ solution, and mixed quickly with a micropipette, initiating the Fenton’s reaction. As a negative control, the mixing DNA solution (Kamimashi-Gun, Kumamoto, Japan) were used as a DNA damage quantification kit. Hydrochloric acid (HCl) and sulfuric acid (H₂SO₄) were from Wako Pure Chemical Industries, Ltd. (Osaka, Japan).

**Electron spin resonance (ESR) experiment procedure.** The electron spin resonance (ESR) method was described previously (18) and explained as follows. Hydroxyl radicals were generated by the Fenton's reaction. First, 50 μl of a 1.0-mM FeSO₄ solution was added to 20 μl of a 90-mM DMPO solution as a spin trapping reagent, and this mixing solution was further added to 250 μl of fish or soy sauce as the sample or to ultra pure water as the control. The ultra pure water was made using the ultrapure water purification system (Mili-Q Jr., Nihon Millipore Kogyo K.K., Yonezawa, Japan). Next, the sample or control solution was added to 50 μl of an 8.8-mM H₂O₂ solution to initiate the Fenton's reaction, which occurs as in the following chemical equation: Fe⁺² + H₂O₂ → Fe³⁺ + •OH + OH⁻.
was added to 50 μl of pure water instead of H2O2 solution. All the solutions were again incubated at 37˚C at 15 min in the water bath. These solutions were cooled in ice to be used as each DNA sample.

Preparation of ARP-labeled DNA. ARP (aldehyde reactive probe; N’-aminoxyxymethylcarbonylhydrazino-D-biotin)-labeled DNA solution was prepared according to a modified method described previously (20,21). A 100-μl/ml concentration solution of each DNA sample was prepared by dilution with TE buffer, and measured at 260 nm absorbance using an UV-VIS spectrophotometer (UV mini 1240, Shimadzu Corporation, Kyoto, Japan) with a quartz cuvet of 10-μl volume (105.210-QS-1, Hellma GmbH and Co. KG, Müllheim, Germany). Ten microliters of prepared DNA solution was mixed with 10 μl of ARP solution in a 1.5-ml microtube. Biotinylation of AP sites on DNA was completed by incubating the mixed solution sample at 37˚C for 1 h in the water bath. The incubated mixture was then added with 380 μl of TE buffer, and transferred to the 0.5-ml filtration tube Microcon YM-30 (molecular weight; 30,000, Bedford, MA, USA). The filtration tube was centrifuged at 2,500 x g for 20 min, and the filtrated solution was discarded. The ARP-labeled DNA on the filter was re-suspended in 400 μl of TE buffer with a micropipette, and then centrifuged at 2,500 x g for 20 min and the filtrate was discarded. The ARP-labeled DNA was finally prepared in 400 μl of TE buffer by serially re-suspending with 200 μl of TE buffer, then transferred to a 1.5-ml microtube, and stored at 0-5˚C.

Determination of the number of AP sites in DNA. Ninety microliters of the ARP-labeled DNA solution was diluted with 310 μl of TE buffer, and then 60 μl of the solution was placed in one well of a 96-well microplate with a U-shaped bottom. The ARP-labeled DNA was adsorbed to the well by mixing with 100 μl of a DNA binding solution from a DNA damage quantification kit (Dojindo Laboratories) and incubated overnight at room temperature. After discarding the solution, the well was washed 5 times with 250 μl of PBST washing buffer using an auto-miniwasher (BioTec Co., Ltd., Tokyo, Japan). The plate was inverted and tapped on a paper towel several times to remove the solution completely. A 150-μl aliquot of horseradish peroxidase (HRP)-streptavidin solution, which was diluted 4,000-fold with PBST washing buffer, was added to each well, and the plate was incubated at 37˚C for 1 h. The reaction solution in the well was discarded and the well was washed with 250 μl of PBST washing buffer 5 times, and tapped on a paper towel. One hundred microliters of the substrate solution was added to each well, mixed with a micropipette, and incubated at 37˚C for 1 h. Fifty microliters of reaction mixture, which developed a blue color at this point, was mixed with 50 μl of 1 M sulfuric acid, and changed to yellow. The absorbance was determined at 450 nm using the multilabel microplate reader Mithras LB940 (Berthold Technologies GmbH and Co. KG). Dual samples were determined for each set of data, and ARP-labeled DNA solution was replaced with TE buffer for the background determination. For the calibration curve, 60 μl each of standard [0, 2.5, 5, 10, 20 and 40 AP sites per 1x10^5 base pairs (bp)] was placed in two wells of the 96-well plate per one concentration sample of the standard solution. We were able to determine 1-40 AP sites per 1x10^5 bp.

Calculation of the ratio of DNA protection. The ratio of DNA protection was defined as the following equation: Ratio of DNA protection (%) = [(A - B) - (C - B)] x 100/(A - B) = (A - C) x 100/(A - B); A, number of AP sites per 100,000 bp after being exposed to hydroxyl radicals in the Fenton’s reaction solution as a control; B, number of AP sites per 100,000 bp without the exposure; and C, number of AP sites per 100,000 bp after the exposure with a fish or soy sauce.

Results

IC50 of hydroxyl radical scavenging capability of fish and soy sauces. The IC50 of the sauce was defined, in this experiment, as the dilution level of the sauce product effective in depressing the strength of electron spin resonance (ESR) of hydroxyl radicals by half. This level was calculated from the linear calibration curve obtained between the levels of the depression and the dilution of the sauce products added as a hydroxyl radical scavenger. The antioxidative value of 30.103 (log2 x 100), which corresponded to half-depression, was calculated to be 0.20% in the case of puffer fish sauce, as shown in Fig. 1. Similar linear relationships were obtained for the other 7 sauces, and their IC50 values were calculated by the same method.

Quantity of AP sites on DNA damaged by hydroxyl radicals. The quantity of AP sites on DNA damaged by hydroxyl radicals was obtained from the calibration curve using the ARP-DNA standard solution (Fig. 2). From the calibration curve, the number of AP sites produced without exposure to hydroxyl radicals, i.e., spontaneous DNA damage, was calculated to be 9.66 per 100,000 base pairs (bp) and that of DNA exposed to hydroxyl radicals produced by the Fenton's reaction was 42.34 (Fig. 3).
Evaluation of the antioxidative level of fish and soy sauces. The IC$_{50}$ values were compared among the various fish and soy sauces. The order of the strength of hydroxyl radical scavenging capability is shown in Fig. 4; the lower the height of the bar in the figure the stronger the radical scavenging capability. Based on the data in the figure, salmon fish sauce, dark color soy sauce, light color soy sauce, puffer fish sauce, squid fish sauce (Ishiru) and colorless soy sauce showed high hydroxyl radical scavenging capabilities. The IC$_{50}$ values of these sauces were 0.09, 0.14, 0.18, 0.20, 0.26 and 0.28%, respectively. On the other hand, sandfish fish sauce (Shottsuru) showed weak antioxidative activity compared with the sauces described above; the IC$_{50}$ value of this fish sauce was 4.16%.

Evaluation of DNA protection against hydroxyl radicals. The order of the level of the ratio of DNA protection against hydroxyl radicals is shown in Fig. 5. The puffer fish sauce showed the highest ratio of DNA protection, 68.9%. Based on the data in the figure, the ratio of DNA protection of salmon fish sauce, dark color soy sauce, light color soy sauce, puffer fish sauce, squid fish sauce (Ishiru) and colorless soy sauce showed high hydroxyl radical scavenging capabilities. The IC$_{50}$ values of these sauces were 0.09, 0.14, 0.18, 0.20, 0.26 and 0.28%, respectively. On the other hand, sandfish fish sauce (Shottsuru) showed weak antioxidative activity compared with the sauces described above; the IC$_{50}$ value of this fish sauce was 4.16%.
sauce, squid fish sauce (Ishiru), dark color soy sauce and light color soy sauce was 67.0, 60.1, 49.7, 34.1, 28.2, and -4.4%, respectively. Puffer, salmon and sandfish fish sauces showed high ratios of DNA protection against hydroxyl radicals.

Relationship between the ratio of DNA protection and the IC50 value of hydroxyl radical scavenging. The relationship between the ratio of DNA protection and the IC50 value of hydroxyl radical scavenging was examined using correlation coefficient (r), regression line and coefficient of determination (R2) (Fig. 6). As a result, the equation of the regression line was y = 5.0124x + 39.525, the correlation coefficient was 0.2861 and the coefficient of determination was 0.0819. Therefore, we judged that a mutual relation was absent at 5% (p=0.05) of the level of significance.

Discussion

Apurinic/apyrimidinic (AP) sites are the most common DNA lesions which are spontaneously created by the release of deaminated bases under various physiological conditions (13). Oxidative attack by hydroxyl radicals on the deoxyribose moiety produces a multiplicity of modifications in DNA, namely it leads to the release of free bases from DNA, generating strand breaks with various sugar modifications and simple AP sites. In fact, AP sites are one of the major types of damage generated by hydroxyl radical species. It has been estimated that endogenous reactive oxygen species can result of damage generated by hydroxyl radical species. It has been reported that edarabone (3-methyl-1-phenyl-pyrazolin-5-one) reacted with the aldehyde group of AP sites, and damage to these biotin-tagged sites can be detected by an ELISA-like assay (20,21).

The AP site DNA method has been used in various research fields, namely DNA damage and its repair in bacteria, yeast or cell culture, and the protection against acute renal damage using rats and postischemic brain injury (22-26). For example, Monti et al reported the influence of base excision repair defects on the processing of Me-lex (a methyl sulfonate ester appended to a neutral N-methylpyrrole-carboxamide-based dipeptide) lesions on plasmid DNA that harbors the p53 cDNA as a target in yeast using the AP site SA DNA method (22). Also, using this method, Satoh et al reported that edarabone (3-methyl-1-phenyl-pyrazolin-5-one) as a novel free radical scavenger protected against cisplatin-induced acute renal damage in male Wistar rats (25). In addition, it was reported, again using this method, that abasic sites, hallmarks of oxidative DNA damage, were significantly increased in DNA from the ischemic brain of folate-deficient animals at early time points after middle cerebral artery occlusion (26).

For the first time, Makino et al tried the AP site DNA method in the field of food antioxidants in 2001 (19). We further developed this method in order to research the antioxidative activity of fish and soy sauce seasonings. In this study, we found that puffer fish sauce and other fish and soy sauces exhibited a high ratio of DNA protection against hydroxyl radicals. However, no relation between the protection against DNA damage by hydroxyl radicals and the hydroxyl radical scavenging capability of fish and soy sauces was observed. Some mechanism, which mediates or depresses the reactivity of the hydroxyl radicals to DNA molecules, may be present. The mechanism seems to affect the efficiency of the antioxidative activity of the sauce products. Therefore, we conclude that the antioxidative activity of food cannot be evaluated only from the viewpoint of radical scavenging capability.

In the future, it is essential that we isolate and research the substances from antioxidative foods which protect DNA against hydroxyl radical damage.

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