Antioxidative and Neuroprotective Activities of the Pre-Germinated Brown Rice Extract

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ABSTRACT

We evaluated the effects of pre-germinated brown rice extract (PGBR ex) with enhanced levels of GABA on proliferation and apoptosis of neuronal SK-N-SH cells line. Firstly, we used HPLC methods to study the level of \( \gamma \)-aminobutyric acid (GABA) in all rice extracts. We found that the concentration of GABA in the PGBR ex were 3 and 8 times higher than the GABA concentration in non-germinated brown rice (BR ex) and white rice (WR ex) compared with the standard GABA respectively. Next we study the protective effects of brown rice extract by investigating various methods, we found that the effects of dose-dependent study by treated with PGBR ex, BR ex and WR ex at \( 0 - 4000 \mu g/ml \). The data from MTT assay showed that the higher concentration of all rice extracts were not induced toxicity to SK-N-SH cells. To test the protective effect by study the viability of SK-N-SH cells. These results showed that PGBR ex and BR ex can protect cells by significantly increase cells survival up to 29.3% ± 0.01% and 13.4% ± 0.07% (\( p < 0.05 \)) but not WR ex comparable with 150 \( \mu M \) H2O2 alone which caused cells death > 56.9% ± 0.02 % (\( p < 0.05 \)), compared with untreated cells (control). Next study we test the effect of cells apoptotic by ROS assay and DNA fragmentation. The results showed that PGBR ex were definitely decrease the amount of ROS formation and had a little of DNA ladders comparable with condition that induced by 150 \( \mu M \) H2O2. Our data indicating that PGBR ex with enhanced levels of GABA effectively inhibit SK-N-SH cells proliferation and apoptosis. These present results suggest that intake of PGBR and BR instead of WR is effective to protect cell proliferation and apoptosis which may be useful nutritional to prevent neuronal cells from neurodegenerative disease.

Keywords: Pre-Germinated Brown Rice; SK-N-SH Cells; Antioxidant; GABA; Apoptosis; DNA Fragmentation

1. Introduction

Rice is a major cereal food and is a dietary staple world wide, especially in Asian countries. Rice seeds and rice germ contain fiber and several kinds of antioxidants, such as ferulic acid, phytic acid, tocopherols, and oryzanols. Brown rice is a rice seed from which only the hull is removed. Recently, we found that pre-germinated brown rice contains a much higher concentration of essential amino acids, such as lysine, isoleucine, methionine, than conventional brown rice, and over 13 times the amount of \( \gamma \)-aminobutyric acid (GABA) [1,2]. Pre-germinated brown rice (PGBR) is brown rice, which has been soaked in water for up to a day and had a germ of approximately 1 mm long. During germination, nutrients in the brown rice change drastically. Nutrients that increase in content include \( \gamma \)-aminobutyric acid (GABA), dietary fiber, inositol, ferulic acid, phytic acid, tocotrienols, magnesium, potassium, zinc, \( \gamma \)-oryzanol, and prolylendopeptidase inhibitor. According to Kenichi, germinated brown rice contained more total ferulic acid (126%), total dietary fiber (145%), soluble dietary fiber (120%) and insoluble dietary fiber (150%) compared to the brown rice [3].

Free radicals have been found to be crucial because they can cause several severe diseases such as cancer, cardiovascular and cell degeneration [4]. This damage results from the imbalance between antioxidants and free radicals in the body [5]. Thus efforts are being expended in the search for substances that can prevent and inhibit the activity of free radicals. An important source of antioxidants is daily vegetables and fruits [6]. Oxidative stress induced cell damage has been shown to be involved in neurodegenerative diseases such as Alzheimer’s disease, Parkinson’s disease, and stroke [7]. The damage is mediated by reactive oxygen species (ROS), mainly superoxide anion (\( O_2^- \)) and hydrogen peroxide (\( H_2O_2 \)). Accumulation of ROS in neuronal cells results in lipid peroxidation, protein and DNA damage, and finally cell death[8]. Sev-
eral studies revealed that ROS can be inhibited by antioxidants [9].

The neuroprotective effects of various natural extracts has been reported in the literature [10]. The water extract of *Curcuma longa* reduces rat pheochromocytoma PC12 cell death induced by pyrogallol and H$_2$O$_2$ [11]. The water-soluble extracts of the seed of *Celastrus paniculatus* have neuroprotective effects against glutamate-induced toxicity in embryonic rat forebrain neuronal cells [12]. Relevant studies associated with rice include the observations that aqueous-ethanol extracts of rice bran exhibited antioxidative properties feruloyl-myoinositol present in rice bran inhibited phorbolester-induced superoxide anion generation in HL-60 cells [13] the cyaniding 3-O-β-D-glucoside isolated from pigmented rice suppressed reactive oxygen species in an in vitro assay [18] and protocatechuic acid methyl ester isolated from black rice inhibited the enzymatic activity of tyrosinase. Recently, Choto-san was shown to act as an antioxidant and neuroprotective agent against oxidative damage in NG108-15 cells [19]. Among them, rice bran is a by-product of the rice milling process and contains various antioxidant factors showing beneficial effects on human health. As well known antioxidants in rice bran, tocopherols, tocotrienols, oryzanols (ferulate esters of tri-terpene alcohols) are isolated from fat-soluble extracts of rice bran against oxygen radical-related chronic diseases.

2. Materials and Methods

2.1. Cells Culture

The human neuroblastoma SK-N-SH cells line was obtained from the ATCC (Rockville, MD, USA) and maintained in minimum essential medium (Life Technology, Inc.) containing 2 mM L-glutamine in a humidified incubator at 37°C and 5% CO$_2$. The medium was supplemented with 10% fetal bovine serum, 1 mM sodium pyruvate, 0.1 mM minimal essential medium nonessential amino acids (Life Technology, Inc.), 100 units/ml penicillin, and 100 mg/ml streptomycin (Biofluids, Rockville, MD). Cells were plated at 5 × 10$^5$ cells/ml were cultured in 96 well plates (for Reactive Oxygen Species (ROS) formation and for cell viability (MTT assay). Cell were plated at 1 × 10$^6$ cells/ml on 6 well plates for DNA fragmentation then maintained in serum free optimal minimal modified Eagles medium (MEM) supplemented with Fetal bovine serum (Invitrogen, Carlsbad, CA). Cultures were maintained for 24 h before treatments.

2.2. Preparation of Brown Rice Extracts

Pre-germinated brown rice (*Oryza sativa* L.) was supplied by Innofood (Thailand) Co., Ltd. in Pratumthani and was germinated by soaking in the following solutions at 25°C - 26°C in the dark for 72 h: was air dried, frozen in liquid nitrogen pre-germinated brown rice extracts were prepared as previously described [10] Briefly, the pre-germinated brown rice (PGBR), brown rice (BR), polish rice or white rice (WR) were ground with a mortar and pestle, and then added with distilled water mix with Vortex for 10 min and then kept in water bath at 70°C for 30 min. The samples were centrifuged at 15,000 rpm at 4°C for 30 min, and the supernatants were collected, passed through filters with 0.45 μm pores, and used as extracts.

2.3. Effect of Brown Rice Extracts on Cell Viabilities of SK-N-SH Cell Lines

To determine the influence of brown rice extract on cell viability which induced by 150 μM H$_2$O$_2$. Cells were plated 5 × 10$^5$ cells/ml in 96-well plates (Falcon, Germany) in medium supplemented with 10% fetal bovine serum and penicillin/streptomycin. After additions by treated with increasing concentrations of brown rice extract (0 - 4000 μg/ml) at 37°C for 3 h and further added with 150 μM H$_2$O$_2$ in culture incubation at 37°C for 24 h as indicated. Viabilities of the cells were assay by the MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide] method, MTT solution was added at a final concentration of 0.5 mg/ml at 37°C for 3 h incubation. Then solubilize solution (DMSO) was then added, the plates were shaken for 10 min and absorbance was read at 570 nm in a microplate reader.

2.4. Assay for Reactive Oxygen Species Formation

Detection of intracellular reactive oxygen species (ROS). The production of intracellular reactive oxygen species
was estimated by using a fluorescent probe, 2′,7′-dichloro-
fluorescein diacetate (DCFH-DA) [21]. DCFH-DA is trans-
ported across the cell membrane and hydrolyzed by in-
tracellular esterases to form nonfluorescent 2′,7′-dichlo-
rofluorescein (DCFH), which is then rapidly converted to
highly fluorescent 2′,7′-dichlorofluorescein (DCF) in the
presence of reactive oxygen species. The DCF fluo-
rescence intensity is believed to be parallel to the amount of
reactive oxygen species formed intracellular. To study
the free radical scavenging effect of Brown rice extracts
(rice extract at final concentration 2000 μg/ml) was added
3 h before added with 150 μM H_2O_2 to the culture and
incubated at 37°C for 24 h. To determine the amounts of
ROS induced by H_2O_2, DCFH-DA (50 μM final con-
centration in DMSO) was added to the cells culture and then
incubated at 37°C for 2 h. The production of reactive oxygen
species was measured immediately by microplate reader
using excitation and emission at 485 nm and a 530 nm.

2.5. Effect of Brown Rice Extracts on Apoptosis
and DNA Fragmentation by DNA Extraction
and Agarose Gel Electrophoresis

Cultured cells were prepared at 1 × 10^6 cells/well in 6
well plates. Rice extracts were added into the cultures at
2000 μg/ml and cultured at 37°C for 3 h and further
added with 150 μM H_2O_2 in culture incubation at 37°C
for 24 h then cells were collected. Genomic DNA was
extracted using the Apoptotic DNA Ladder Kit, (Roche
Applied Science, Mannheim, Germany) with slightly
modification. In brief, cells were washed twice with ice
cold 1 × PBS, then 200 μl binding/lysis buffer was
added and mixed immediately. After holding for 10 min at
22°C, 100 μL of 100% isopropanol was added and the
solution was vortexed for 10 sec. The lysate was run
through the column then washed twice with washing buffer.
DNA was eluted with 200 μL of pre-warmed (70°C) elu-
tion buffer and concentrated with a speedvac. Extracted
DNA was subjected to gel electrophoresis, and the image
was captured with GelDoc™ EQ (Bio-Rad Laboratories,
Ltd. Hercules, CA).

2.6. Statistical Analysis

The results were expressed as mean ± SEM of triplicate
assays. The statistical comparison between control and
treated experimental groups were carried out using Stud-
ent’s t-test. P-value less than 0.05 were considered to be
significantly different.

3. Results and Discussion

3.1. The Cytotoxicity of Pre-Germinated Brown
Rice Extract on SK-N-SH Cells

First, we attempted to establish the doses dependent of
the brown rice extract, the cytotoxicity of brown rice
extract on human neuronal SK-N-SH cells was observed.
PGBR ex, BR ex and WR ex were applied in culture by
increasing concentrations 0 - 4000 μg/ml to SK-N-SH
cells for 24 h. We found that the data of cells survival
showed that std. GABA up to 96.4 ± 0.052 (p > 0.48),
PGBR ex 98.9 ± 0.032 (p > 0.30), BR ex 96.3 ± 0.039 (p
> 0.39) and WR ex 95.3 ± 0.037 (p > 0.25), respectively.
The data of viability cells had no significantly decrease
cell survivals which had no toxicity affect on human
neuronal SK-N-SH cells (Figure 1). However, treated
with 150 μM H_2O_2 alone caused cells death > 56.9% ±
0.02% (p < 0.05) compared with control indicating at this
concentration H_2O_2 had toxicity to SK-N-SH cells. (data
not shown) Similar to Chan-Ho Oh and Suk-Heung Oh
had tested the effect of the brown rice extracts on the
viability of Hela cells, by treated cells with the brown
rice ex extracts 2000 μg/ml in cultured, the viability of
the cells was assayed. Their results had shown that
brown rice extracts also had no effects on the retardation
of HeLa cell proliferation.

3.2. PGBR Ex Inhibit the Neuronal SK-N-SH
Cell Death Induced by H_2O_2

Next we tested the protective effects of the brown rice
extracts on the viability of SK-N-SH cells, cells were pre-
treated with the PGBR ex, BR ex and WR ex at 2000 μg/ml
and then treated with 150 μM H_2O_2. These results showed
that PGBR and BR ex can protect cells. The cells survival
significantly increase up to 29.3% ± 0.01% and 13.4% ±
0.07% (p < 0.05), but not WR ex comparable with 150
μM H_2O_2 treated alone which caused cells death >56.9 ±
0.02% compared with control (Figure 2). PGBR ex con-
tains approximately 13 times of GABA and the amount of

![Figure 1. Effect of brown rice extract on the viability of neuronal SK-N-SH cells. Cells (5 × 10^6 cells/ml) were incubated with increasing concentrations of brown rice extracts; PGBR, BR and WR at 0 - 4000 μg/ml. Cultures were incubated at 37°C for 24 h. The cytotoxicity of brown rice extract was measured using MTT assay, as described in Materials and methods. Results are expressed as percentages of the control value (untreated cells) and data shown are means ±S.E.M of three replicate experiments.](https://example.com/figure1.png)
roxidative damage [23]. Ferulic acid, a representative antioxidant, inhibits Aβ-induced neurotoxicity. In vitro studies have shown that β-tocopherol, a representative antioxidant, inhibits Aβ-induced neuronal cell death and lipid peroxidation [22]. Ferulic acid also possesses free radical scavenging activity and reduces peroxidative stress is one of the mechanisms of Aβ-induced neurotoxicity. In vitro studies have shown that β-tocopherol, a representative antioxidant, inhibits Aβ-induced neuronal cell death and lipid peroxidation [22]. Ferulic acid also possesses free radical scavenging activity and reduces peroxidative damage [23].

3.3. PGBR Ex Inhibits the Apoptosis-Associated DNA Fragmentation Induced by H2O2

DNA fragmentation is a marker of late stage of apoptosis. To verify the possible involvement of apoptosis in the 150 μM H2O2-induced death of neuronal SK-N-SH cells. We investigated its inhibitory effect of PGBR ex on the apoptosis induced by 150 μM H2O2 by observing DNA fragmentation levels in SK-N-SH cells. This figure demonstrates internucleosomal DNA degradation from gel electrophoresis. As a control, we used DNA isolated from untreated cells and no DNA ladder similar results was shown when treated with PGBR ex (Figure 3, lanes 2 & 3). Fragmented DNA was observed when cells were treated with 150 μM H2O2 resulted in the characteristic apoptotic DNA ladder. DNA fragments smeared the whole lane (Figure 3, lanes 4). These fragmentation patterns are entirely consistent with the molecular weight patterns expected to result from internucleosomal DNA cleavage [24]. But DNA fragments was decreased when co-treated with PGBR ex and 150 μM H2O2 (Figure 3, lanes 5). These DNA fragmentation patterns are entirely consistent with the molecular weight patterns expected to result from internucleosomal DNA cleavage. DNA fragmentations are indicative of early and late stage of apoptosis, respectively. From this results shown that the PGBR ex in our present study showed the anti-apoptotic effect on SK-N-SH cells.

3.4. PGBR Ex Inhibits H2O2 Induced ROS Generation in Neuronal SK-N-SH Cells

Intracellular ROS have been implicated with DNA fragmentation and apoptosis [25]. The intracellular radical scavenging activity of a given substance can be evaluated by DCFHDA assay. To determine whether PGBR ex inhibits the ROS generation induced by 150 μM H2O2 in neuronal SK-N-SH cells, and to determine whether the protection of SK-N-SH neuronal cells from apoptosis is accompanied by free radicals scavenging by the PGBR ex, changes in the concentrations of ROS in whole cell suspensions were analyzed over 2 h period after treatment with PGBR extract and/or 150 μM H2O2. SK-N-SH cells showed relatively low levels of basal fluorescence but when treated with 150 μM/ml H2O2 for 2 h, a marked increase in fluorescence was observed. However, this increase in fluorescence by 150 μM H2O2 rapidly returned to the control level in PGBR ex pretreated (2000 μg/ml) to the cells (Figure 4).

4. Conclusions

The concentrations of GABA in the PGBR ex, BR ex and WR ex were compared with standard pure glutamic acid. The concentration of GABA in the PGBR ex (1200 nmol/ml) was 3 times higher than the GABA concentration in the BR ex (400 nmol/ml). The concentration of GABA in the PGBR ex was 8 times higher than the GABA concentration in the WR ex (150 nmol/ml). In this study, we evaluated the effects of PGBR extracts with enhance levels of GABA against 150 μM H2O2 induced cytotoxicity, oxidative stress and apoptosis were compared with those of BR ex, and WR ex on neuronal...
SK-N-SH cells. To our knowledge, this is the first report of the protective effects of PGBR on human neuronal SK-N-SH cells.

Natural antioxidants that can neutralize ROS include cysteine, reduced glutathione, polyphenolic compounds, carotenoids, ascorbic acid (vitamin C), β-tocopherol (vitamin E) and indole carbinds. Some researchers have suggested that germination may bring about changes in nutrients and physiologically active substances. During the germination of wheat [26] and Panigum edule Reinit [27] and rice seed vitamin C, vitamin E, ferulic acid, and total phenolic acid contents, alanine and β-aminoxybutyrate have been reported to increase significantly [28]. Upon malting of finger millet, changes in both free and bound phenolic acid contents were observed and these reflected their antioxidant properties. We speculate that, during germination, as seed moisture increases, the seed coat may be a method to improve health-related benefits.

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