Bio-catalyzer α+β No.11, Bio-catalyzer 2B, BIO No.66 Quenched Free Radicals and Inhibited Lipid Peroxidation in Iron-induced Epileptic Focus in Rats

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INTRODUCTION

The biochemical pathogenesis of post-traumatic epilepsy implicates the most potentially damaging hydroxyl radicals (·OH) which initiate lipid peroxidation in neuronal membrane, edema, cavity necrosis in tissues, and convulsive seizures in iron-induced epileptogenic foci in rats (1-3). The search for effective health foods and other non-drugs with minimal side effects for prevention and treatment of post-traumatic epilepsy has received much attention.

Recently, we have demonstrated that a commercially available health food called Bio-catalyzer α+β No.11 (Bio-normalizer) and its by-product Bio-catalyzer 2B are ·OH scavengers. The former was found to inhibit the formation of thiobarbituric acid reactive substances, a measure of lipid peroxidation in iron-induced epileptic foci in rats (4).

Through electron spin resonance spectrometry, we analyzed the scavenging action of a new health food called BIO No.66. The anti-epileptic potentials of Bio-catalyzer 2B and BIO No.66 were also tested using the same experimental animal model.

MATERIALS AND METHODS

Chemicals Hypoxanthine, 1,1-diphenyl-2-picrylhydrazyl (DPPH), diethylenetriaminepentaacetic acid (DETAPAC) were purchased from Sigma Chemical Co., St. Louis, MO. Xanthine oxidase and 5,5-dimethyl-1-pyrrrole-N-oxide (DMPO) were from Boehringer Gmbh and Daiichi Pure Chemicals Ltd., Tokyo, respectively. Other chemicals and reagents were of the highest grade available from commercial suppliers.
Samples Bio-catalyzer α-β No.11 (Bio-normalizer) and its by-product Bio-catalyzer 2B are white, sweet, granular, natural health food whereas BIO No.66 is a light brown, powder health food. These are products of Sun-O International Inc., (Gifu, Japan) from fermented Carica papaya Linn., Pennisetum pupureum Schum., Sechium edule Swartz, glucose and yeasts.

Animals Male adult Sprague-Dawley rats from Clea Japan, Inc., Tokyo weighing 240-260 g were used in this study.

Radical Analysis Details of the analysis of DPPH, hydroxyl and superoxide radicals by ESR spectrometry (JES-FE1XG, JEOL, Tokyo) and ESR parameters are given in a previous report (4). Hydrogen- and carbon-centered radicals were analyzed as follows: To 0.5 ml of rat cerebral cortex homogenate (0.05% w/v), 0.45 ml of Tris/HCl buffer, pH 7.4, and 0.5 ml of equimolar (2 mM) mixture of ferrous chloride and ascorbic acid was added the sample, mixed for 10 s and incubated at 37°C for 15 min. To the incubated medium was added 20 µl of DMPO, mixed for 10 s, then placed in an ESR flat cell (JEOL, Tokyo) and analyzed exactly after 50 s.

Iron-induced epileptic foci of rats Ferric chloride-induced epileptic foci of rats was performed following the procedure of Willmore et al. (5). Briefly, the left cerebral cortex of the control animals were subpially injected with the same volume of saline in place of ferric chloride which pH was adjusted to that of the iron solution. Rats were administered orally with 1 g/kg body weight of Bio-catalyzer 2B or BIO No.66 30 min prior to cortical injection of 5 µl of 100 mM ferric chloride for 5 min thru a needle inserted into the left cortex to a depth of 1-2 mm. Rats were sacrificed by cervical decapitation and the epileptogenic focal area was rapidly excised over an ice plate following the method of Glowinski and Iversen (6) and kept at -80°C until TBARS analysis. TBARS were assayed as described by Ohkawa (7).

Statistical Analysis was performed using Student's t-test.

RESULTS AND DISCUSSION

While 50 mg/ml each of Bio-catalyzer and Bio-catalyzer 2B, and BIO No.66 showed weak scavenging action on DPPH and superoxide radicals with activity ranging from 5-10%, these 3 products exhibited stronger quenching action on ·OH. BIO No.66 appears to be the most potent among the 3, requiring only 25 mg/ml to quench about 95% of the hydroxyl radicals
Fig. 1. Scavenging effect of Bio-catalyzer α·P No.11 (□) and Bio-catalyzer 2B (■) (graph A) and BIO No.66 (graph B) on hydroxyl radicals generated from Fenton reagents. The values are the means of (n = 6-8) ± SEM.

(4.4 x 10^4 spins/ml) (Fig. 1B) as compared to 50 mg/ml required by Bio-catalyzer and Bio-catalyzer 2B (Fig. 1A). At 350 mg/ml, BIO No.66 also scavenged 98% of the hydrogen radicals (7.8 x 10^4 spins/ml) and 72.3% of the carbon-centered radicals (1.2 x 10^4 spins/ml) generated in the brain homogenate induced with iron and ascorbic acid. BIO No.66 thus, appears to have wider-ranged scavenging action (Fig. 2).

The radical scavenging property of the 3 products may be attributed to papaya—the base plant material used in their preparation, and other constituents such as yeast and glucose. At 50-250 mg/ml, papaya (a Philippine medicinal plant) quenched DPPH, ·OH and superoxide radicals (unpublished data). Yeast Saccharomyces cerevisiae also quenched DPPH, hydroxyl, superoxide, hydrogen- and carbon-centered radicals (8). Glucose scavenged ·OH at 0.045 to 45.45 mg/ml. Other plant materials used may also contain natural antioxidants.

Figure 3 shows the inhibitory action of Bio-catalyzer 2B and BIO No.66 in vivo lipid peroxidation. The TBARS level was decreased in the left or ipsilateral sensorimotor cortex of rats orally administered with 1 g/kg body weight of the products whereas, the TBARS concentration remained high in the untreated iron-induced rat sensorimotor cortex. Saline-injected rat group contains appreciable TBARS. Similar preventive action on TBARS formation was reported on Bio-catalyzer (4).

The treatment by alpha tocopherol and selenium prevented epileptiform discharges in iron-induced rats (9-10). Superoxide, peroxyl radicals, organic radicals and ·OH have been reported to be liberated in the neuronal membrane, where they initiate peroxidative damage and cause experimental post-traumatic epilepsy in rats. Corollary with these reports, our
Fig. 2. Scavenging effect of 350 mg/ml of BIO No.66 on hydrogen- and carbon-centered radicals. The values are means of 4-5 determinations.

Fig. 3. Inhibitory effect of Bio-catalyzer 2B and BIO No.66 on TBARS level in the iron-induced epileptic foci of rats. The values are means (n= 7-8) ± SEM.

- p < 0.001 vs. control grp.; p < 0.001 vs. iron-injected.

Data suggest that Bio-catalyzer, Bio-catalyzer 2B and BIO No.66 may act as natural antioxidative health foods by scavenging the •OH and providing a protective barrier against these radicals.

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