Effects of Korean Ginseng and Wild Simulated Cultivation Ginseng for Muscle Strength and Endurance

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Abstract - Muscle strength and endurance activities of Korean ginseng (Panax ginseng C. A. Meyer; KG) were compared with those of wild simulated cultivation ginseng (WCG) in mice. Fifty male ICR mice were divided into five groups: A (vehicle); B (WCG 100 mg/kg); C (WCG 500 mg/kg); D (KG 100 mg/kg); E (KG 500 mg/kg). Subsequently, the mice were subjected to the forced swimming test (FST) and treadmill test at the 4th and 7th weeks. The glycogen content in the muscle and blood analysis (levels of glucose, triglyceride (TG), IGF-1) were also performed immediately after the last FST and treadmill test at the 7th week. Immobility times in FST were shorter in WCG- than KG-treated groups, and the results of the treadmill tests were also significant except for KG-treated at 100 mg/kg. The glycogen content was increased in both groups with a peak at 500 mg/kg of WCG groups. Serum concentrations of TG and glucose were decreased by administration of KG and WCG and all treated groups showed increase in the level of IGF-1 in serum. These results suggest that KG and WCG suplementations are effective in escalating the muscle strength and endurance.

Key words - Wild simulated cultivation ginseng, Muscle strength, IGF-1

Introduction

It has been reported that many natural resources had an effect on the increase of exercising ability, such as muscle strength and endurance. Among them, especially, studies like the use of ginseng and red ginseng have been carried out (Alvarez et al., 2004; Lee et al., 2009). Korean ginseng (KG), the root of Panax ginseng C.A. Meyer (Araliaceae), is a popular natural health food in Asia. Many studies have been reported that KG had an effect on neuroprotection (Lee et al., 2011), regulation of body fluid and metabolism (Jia et al., 2010), increase of heart stimulation, anti-diuresis and sexual dysfunction (Becker et al., 1996), improvement of resistance on stress (Qiang et al., 2010), anti-inflammation (Cabral de Oliveira et al., 2001; Park et al., 2010), anti-oxidation (Kim et al., 1996), promotion of immune antibody production (Kim and Jung, 1987), and anti-cancer (Chang et al., 2002). It has been also known that KG is closely linked to its protective properties against free radical attack and has nutritional ergogenic aids for enhancing endurance capacity (Chen, 1996; Lee et al., 1999; Maffei Facino et al., 1999; In et al., 2006). KG administration to rats prevented myocardial ischemia-reperfusion damage induced by hyperbaric oxygen (Maffei Facino et al., 1999), and ameliorated the muscle injury (Alvarez et al., 2004).

Wild mountain ginseng which are grown naturally in the primitive forest without getting polluted by chemical fertilizer for many years are regarded as one of the most highly medical and preservative valued plants. It is believed that the slow-growing wild roots, which are harvested at an older age, absorb more curative power from the forest floor (Shin et al., 2001). The method called wild-simulated cultivation can be used to grow ginseng without fungicide sprays and needs expensive establishment costs, and we named this type of ginseng in the study, Wild-simulated Cultivation Ginseng
It has been known that distribution of ginsenoside contents to the WCG and ginseng was similar (Jeong et al., 2010). However, there has been reported that roots of WCG contains ginsenoside Rg1 and Rb1, while ginseng has not. To our knowledge, the effects of WCG as an ergogenic aid for enhancing muscle endurance capacity has not been clarified yet in animal studies. Here, we investigated the effects on muscle strength and the improvement of endurance through the eccentric exercise and measured glycogen contents in muscle and blood analyses including glucose, triglyceride (TG) and IGF-1 in both WCG and KG supplemented mice.

Materials and Methods

Preparation of extracts of Korean Ginseng and Wild simulated Cultivation Ginseng

The Korean ginseng (KG) and Wild simulated Cultivation Ginseng (WCG) were purchased from Jirisan Sansam Farming association corporation (Jeonju-si, Korea, www.jrsansam.com, Cheon-Gyu Ko). The dried roots were grounded and extracted with 70% ethanol at 70°C for 18 hours in a reflux condenser. The boiled extracts were then filtered through Whatman No. 2 filter paper and then concentrated by a vacuum evaporator (Rotary evaporator N-1000, Eyela, Japan) at 40°C and then were lyophilized to obtain dried samples.

Experimental Animals

Male ICR mice (n = 50) were purchased from Japan SLC, Inc. at 5 weeks of age, and the five animals were housed together in one cage (27 × 17 × 13 cm) in a controlled environment under a light-dark cycle (lights on at 07:00 and off at 19:00). The experimental procedures were conducted according to the Code of Ethics for Animal Experimentation of Gachon University. After a 1 week acclimation, all mice were randomly divided into five groups (10 mice/group): A (no ginseng intake and saline); B (WCG 100 mg/kg); C (WCG 500 mg/kg); D (KG 100 mg/kg); E (KG 500 mg/kg). The solutions were administered as a bolus by gavage, using a curved and ball-tipped intubation needle affixed to a syringe for 7 weeks. All groups were allowed to eat food and drink water freely and the body weights were measured at the 1st, 4th and 7th weeks. The eccentric exercise was performed on a rodent treadmill and FST test with the modification of following protocol described by Armstrong (Armstrong et al., 1983).

Forced Swimming Test (FST)

The FST was conducted for endurance enhancement ability. After swimming-acclimation twice a week, the FSTs were carried out on the 4th and 7th week after treatment. Mice were forced to swim in acryl plastic tank (70 × 70 × 60 cm) filled with 23 – 25°C water up to 80%. A mouse was regarded as immobile when it remained still in the water, making only small movements to keep its head above water. The total duration of immobility was measured for 4 minutes after 2 minutes.

Rota Rod Treadmill Test

The rota rod test was used to assess whether there were gross motor impairments in WCG or KG treated mice. Twenty-four hours before the experiment, all mice were habituated to running on a rota rod treadmill (Rota-rod, Ugobasil) at a speed of 60 rpm until they could remain there for 60 seconds without fail. After the experiment, durations of latency to fall (the running time) were measured.

Glycogen Content in Muscle

To measure the content of glycogen in skeletal muscle, mice were sacrificed by cervical dislocation and the same amounts of right femoral muscle were collected in all experimental groups. Samples of muscle (160 mg) were homogenized in 1 ml of water at 0°C. The homogenate was hydrolyzed in 100 μl of HCl 6 M, incubating at 100°C for 30 minutes. Subsequently, the samples were cooled, neutralized with 285 μl of KOH 2 M and centrifuged at 100 g for 10 minutes. The precipitates were incubated with 0.5 ml of distilled water and 1.0 ml of anthrone (final. 0.02%) at 37°C for 20 minutes, and were quantified visually at 620 nm using spectrophotometer.

Biochemical Analyses in Serum

Blood samples were collected from the infra orbital plexus of each animal and serum was separated by centrifugation. The levels of TG and glucose were measured using a Triglyceride assay kit (BioVision) and Glucose assay kit (Abcam), respec-
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tively. At the end of the experiments, all mice were sacrificed by cervical dislocation and the same amount of left femoral muscle was homogenized in 100 mM of potassium phosphate buffer (pH 7.4) to measure enzymatic activity of skeletal muscle. The activity of lactate dehydrogenase (LDH) and ATP content in muscle were measured using QuantiChrom™ Lactate Dehydrogenase Kit (BioAssay Systems) and ATP colorimetric assay kit (Abcam), respectively. The IGF-1 level in blood was measured using the Quantikine mouse / rat IGF-1 ELISA kit (R&D Systems).

Statistical Analysis
Data were presented as means ± SD. All analyses were performed using SPSS V17.0 (SPSS, Chicago, IL). Probability values of less than 0.05 were significant (*p < 0.05, **p < 0.01). Data expressed as mean ± SD of three to five independent experiments.

Results

Effect of KG and WCG on Exercise Tolerance
To measure the change of endurance, FST was carried out on the 4th and 7th weeks after oral administration. The FST was conducted as a study for evaluating the enhancement of muscle endurance ability. Although there were significant decreased immobility times in WCG and KG groups on the 4th and 7th weeks, KG treated group showed no change in immobility time at a dose of 100 mg/kg (Fig. 1A). The decreased immobility times means that WCG (100, 500 mg/kg)- and KG (500 mg/kg)-treated groups showed the improvement of exercise tolerance.

Effect of KG and WCG on Myorelaxation
To confirm the change of exercise tolerance, a treadmill test was performed on the 4th and 7th week after treatment of WCG and KG. Activities on the rotating rod treadmill reflected the levels of myorelaxation. Supplementation of WCG and KG for 4 and 7 weeks also significantly increased the running time (p < 0.01) except for KG-treated at 100 mg/kg (Fig. 1B). In the treadmill test representing the eccentric exercise, the results of WCG-and KG-treated groups are compatible with those of the FST tests.

Effect of KG and WCG on Glycogen Content in Muscle
Maintaining adequate stores to meet energy needs help to ameliorate fatigue during exercise. Carbohydrate stores in muscle and liver are important for sustained energy. It has been reported that a low or depleted glycogen stores limit exercise time and intensity and leads to decrease the time to exhaustion during physical activity (Cochran et al., 2010). Muscle glycogen becomes important as a fuel for muscular exercise as the intensity of exercise increases. Compared with the control group, in all WCG- and KG-treated groups, the...
contents of glycogen in muscle were increased significantly (Fig. 2). The highest level of glycogen in muscle was shown in WCG-treated groups at 500 mg/kg.

**Effect of KG and WCG on Levels of Glucose and TG**

In both WCG- and KG-treated groups, the levels of blood glucose were lower than that of the control group (Fig. 3A). There were significant decreases ($p < 0.05$) of blood TG levels in 500 mg/kg of WCG- and KG-treated group (100 and 500 mg/kg) (Fig. 3B), but the levels of TG in muscle did not change in both-treated groups (data not shown).

**Effect of KG and WCG on Levels of IGF-1**

IGF-1 plays a crucial role in muscle regeneration and endurance. IGF-1 stimulates both proliferation and differentiation of stem cells in an autocrine-paracrine manner, although it induces differentiation to a much greater degree (Singleton and Feldman, 2001). In all WCG- and KG-treated groups, the levels of IGF-1 in serum were significantly increased ($p < 0.05$) compared to the control group (Fig. 3C).

**Discussion**

WCG and mountain ginseng in the wild are perennial plants belonging to the genus *Panax* and have been used for

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**Fig. 2.** Effect of KG and WCG on glycogen levels in muscle. Thirty of KOH solution was added to the skeletal muscles and samples were incubated at 100°C for 30 minutes. After incubation, samples were mixed with 95% of ethanol and centrifuged for 15 minutes at 4,000 × g. The precipitates were incubated with 0.5 ml of distilled water and 1.0 ml of anthrone (final. 0.02%) at 37°C for 20 minutes, and were quantified visually at 620 nm using spectrophotometer.

**Fig. 3.** (A) Effect of KG and WCG on glucose levels in serum. The levels of glucose were measured using Glucose assay kit. (B) Effect of KG and WCG on TG level in serum. The levels of triglyceride (TG) were measured using Triglyceride assay kit (BioVision). (C) Effect of KG and WCG on IGF-1 concentration in serum. The IGF-1 level in blood was measured using the Quantikine mouse/mouse IGF-1 ELISA kit (R&D Systems). * Significantly different from the control at $p < 0.05$. 

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typical energy-supplement herbs in Chinese and oriental medicine. In this study, we investigated the comparative effects of WCG and KG on the strength and the endurance of muscles. WCG-treated groups showed more improved muscle strength and endurance than those of KG-treated groups.

On the 4th and 7th week, all WCG-treated groups and 500 mg/kg of the KG-treated group significantly increased the treadmill running time and significantly decreased the immobility time in the FST test. These results suggested that WCG- and KG-treated groups showed the improvement of muscle endurance and WCG has a higher effect on the muscle endurance at a lower concentration comparing with those of KG. Generally, the muscle endurance means the recovery of muscle contraction through the inhibition of fatigue materials (e.g. inorganic phosphate and lactic acid in muscle accumulation from glycogen) and the rapid restoration of energy-metabolic materials like ATP and creatine phosphate (Wei et al., 1992).

After the final FST at the 7th week, the contents of glycogen in all WCG- and KG-treated groups were increased significantly compared to the control group. Glycogen is an important energy source during prolonged strenuous exercise and exercise must be abandoned if glycogen stores are depleted. Glycogen storage in the body is limited (300-500 g) (Beelen et al., 2010). As exercise continues, stored glycogen in the skeletal muscle and liver becomes depleted rapidly and exercise performed at 65-75% of maximal oxygen consumption can be maintained only for 90-120 minutes. Therefore, amelioration of glycogen depletion during prolonged strenuous exercise can be a major determinant of endurance and this accumulation of the glycogen in the muscle by KG and WCG might increase the promotion of endurance and exercise efficiency (Hawley and Burke, 2010). It has been also reported that after exhaustive exercise skeletal muscle glycogen levels of ginseng saponin treated rats were slightly higher than those of saline treated rats (Alvarez et al., 2004). These results suggested the possibility that the lowered consumption of glycogen might be due to the promotion of muscle endurance by WCG and KG supplementations.

Eccentric exercise (EC) produces high muscular tension when the muscles are stretched and is associated to delayed onset muscle soreness provoking ultra-structural and metabolic changes in the muscle cells. The injuries are subcellular and generally in small areas of the muscular fiber. Therefore, they suffer a transient reduction in strength, subsequently muscular pain and finally an inflammation process that causes further tissue deterioration. EC is also responsible for the low levels of glycogen and induces transient insulin resistance in healthy individuals (King et al., 1993; Kirwan et al., 1992).

We also measured the serum analysis such as the levels of TG, glucose and IGF-1 were performed immediately after the last FST and treadmill test at the 7th week. The TG levels in serum were decreased significantly in 500 mg/kg of WCG- and KG-treated groups. However, the TG level in muscle and the activity of LDH and ATP content in muscle were not changed (data not shown). In the glycolysis, NAD+ is needed for the ATP production and NAD+ supply by LDH is needed for serial ATP production in anaerobic exercise which resulted in lactic acid formation. These results suggest that WCG and KG could induce the conversion to aerobic exercise from anaerobic exercise forms the lactic acid. The ATP content in muscle is decreased rapidly in continuous exercise because the energy generated by degradation of ATP is used in the muscle contraction during the exercise. Finally, the glycogen is exhausted and the anaerobic energy metabolism in the muscle is activated which resulted in lactic acid accumulation (Paik et al., 1997). These metabolic materials induce the fatigue after the muscle exercise. With this reason, we analyzed the ATP content in the muscle after WCG and ginseng treatment for weeks, but WCG- and KG-treated mice did not show any changes that the ATP content in the muscle (data not shown). Therefore, these results suggest that WCG and KG were used for the ATP effectively to inhibit the rapid decrease of ATP. Overall glucose levels in serum showed the low pattern in all WCG- and KG-treated groups. These results suggest that WCG and KG might enhance the consumption of glucose in serum during FST and treadmill tests.

IGF-1 is the polypeptide consisting of 70 kinds of amino acids and is produced mainly in the liver and bone. It has been reported that IGF-1 stimulated the protein synthesis, cell differentiation, growth and regeneration of skeletal muscle and protected the neuronal cell (Singleton and Feldman, 2001). Also, it has been reported that the declines of the muscle strength and exercise ability were closely related with the decrease of IGF-1 concentration in the body and the
absence of IGF-1 causes the lack of growth, the retardation of spiritual development and the insensibility of sensory nerves (Laron, 2001; Nilsson et al., 1986). In this study, the levels of blood IGF-1 were significantly increased in all treated groups and these results suggested that WCG and KG stimulated the synthesis of IGF-1 resulting to the alleviation of stamina reduction and the improvement of the muscle endurance. The beneficial effect of WCG can be attributed to the activity of ginsenoside Rg1 and Rb1.

Due to the relatively high cost of WCG used in this study, we were unable to examine with greater parameters. Additional analyses are required to detect a clear statistical difference regarding the parameters between WCG and KG. In conclusion, we verified that WCG and KG had effects on the muscle strength and improvement of endurance.

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Literature Cited

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