The Effects of Either Chrysin or Moderate Exercise on Inflammasome and Thermogenic Markers in High Fat Fed Mice

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The purpose of this study was to investigate the effects of either chrysin or exercise on the inflammasome and thermogenic markers in the livers of high-fat fed mice. C57BL/6 mice were randomly assigned to four groups: normal diet control (NC; n=5), high-fat diet control (HC; n=5), high-fat diet with chrysin (Hch; n=5), and high-fat diet with moderate exercise (HME; n=5). The mice were fed a high-fat diet (60% of calories from fat) or normal diet (18% of calories from fat). Chrysin was supplemented orally as 50mg/kg/day dissolved in a 0.1ml solution of dimethyl sulfoxide. The exercised mice ran on a treadmill at 12-20 m/min for 30-60 min/day, 5 times/week, for 16 weeks. After the intervention, the epididymal fat and liver weights were significantly decreased in the HME group compared with HC and Hch groups. The adipocyte size was effectively decreased in the Hch and HME groups compared with the HC group. The inflammasome markers NLRP3, IL-1β, and caspase1 were significantly decreased in the Hch and HME groups compared with the HC group. The thermogenic markers PGC-1α and BMP7 were significantly lower in the HC than in the NC group. However, the HME group showed an increase in the thermogenic markers. In conclusion, chrysin and moderate exercise have positive effects on obese metabolic complications induced by high-fat diets by reducing inflammasome genes. However, chrysin supplementation had no effect on thermogenic gene expression. Moderate exercise would therefore seem to be more effective in controlling obesity-induced metabolic deregulation.

Key words: Browning marker, chrysin, exercise, inflammasome, obese

Introduction

Chronic inflammation state is a pivotal mark in obesity. In obesity, inflammatory cytokines released from adipose tissue reach out the liver and can directly interfered liver functions [30]. Recently, interleukin-1 (IL-1) is one of the regulators in inflammation-related diseases, and the IL-1 secretion is called an inflammasome [19]. Among inflammasomes, the most studied in the area of metabolic disorder is the NOD-like receptor family, pyrin domain containing 3 (NLRP3) inflammasome consisting of NLRP3, apoptosis-associated speck-like protein containing a caspase activation recruitment domain (ASC) and caspase 1 (CASP1) [5]. Previous report has demonstrated that NLRP3 inflammasome may be responsible for the progression of metabolic disorders. Moreover, NLRP3 inflammasome is implicated in obesity, hepatic inflammation [32]. Vandanmagar et al. [30] have suggested that obese caused the inflammasome marker increased significantly in liver. Alleviating hepatic inflammation through regular exercise, and diet intervention may help treat obesity. Moderate exercise can prevent against of chronic inflammation [18] and reduces state of fibrosis as a marker of hepatic injury [19]. Our previous study has been shown the positive effect of treadmill exercise and resveratrol supplementation on inflammatory cytokines, mitochondrial biogenesis, adipogenesis in adipose tissue of obese animal [9, 11, 15]. Also, recent study reported that resveratrol reduced activation of NLRP3 inflammasome in the subcutaneous fat cell [16]. Many researchers use variety way to treat obese induced metabolic complication such as functional nutrition diet. Nutritional approaches to reduce hepatic inflammation using functional foods, including blueberry and resveratrol, have been introduced [3]. However the underlying mechanisms have not been clearly identified. Chrysin, (5, 7-dihydroxyflavone), is a natural flavonoids present in many plant extracts, honey and propolis [10]. Several, studies have
shown that chrysin has multiple biological activities, such as anti-inflammation, anti-oxidation [2, 3, 8]. Although most previous studies reported the gainful effects of chrysin and exercise on high-fat-induced metabolic disturbances, there were few results, comparing the effectiveness among them. Therefore, the purpose of this study was to compare the effectiveness either chrysin or moderate exercise training on inflammasome, and browning markers in liver of high fat diet mice.

**Material and Methods**

**Animals and diet**

Male C57BL/6 (Central Experiment Animal, Korea, N=20) mice were housed in cages in a standard experimental laboratory, at temperature 22±2°C, with 60±5% of humidity. After a one-week acclimatization period, the mice were fed a high fat diet (60% of calories from fat, 20% from carbohydrate, 20% from protein, Orient Bio Inc., #D12492) or a normal diet (18% calories from fat, 58% from carbohydrate, 24% from protein, Orient Bio Inc., #2018) ad for 16 weeks. The mice were divided into 4 groups: normal diet control (NC; n=5), high fat diet control (HC; n=5), high fat with chrysin (Hch; n=5), high fat with moderate exercise (HME; n=5). Liver tissues were dissected, weighed and immediately frozen. All experiments were approved by the Animal Care and Use Committee at the Chungnam National University (CNU-00494).

**Exercise protocol and chrysin supplements**

In the early phase of moderate exercise, all the mice of exercise ran at 8-10 m/min for 10 minutes on the automatically operated rodent treadmill. After that, the speed and time of exercise were gradually increased up to 10-16 m/min for 40-60 minutes. The volume of this exercise training protocol represents 60-75% of the maximum oxygen consumption of the rodents as moderate intensity exercise program [23]. A foam sponge was placed in the back of each treadmill lane to prevent the animal from being injured. Electrical shock was not used during the treadmill run. The control mice were exposed to environmental stress as well as noise from the treadmill. Chrysin supplemented mice were orally gavaged with chrysin (50 mg/kg/day) dissolved in a 0.1ml solution of Dimethyl Sulfoxide [10]. The other group received the vehicle. Each treatment was administrated once a day, five times a week, for 16 weeks before tissue collection.

**Histology of adipose tissue size**

White adipose tissue samples were taken from characteristic depots identified. Flash-frozen adipose tissues were fixed in 4% buffered parafomaldehyde and sections were stained with hematoxylin and eosin (H&E). To measure adipocyte cross-sectional area, four microscope fields in H&E-stained adipose tissue from 5 mice per group were digitally recorded. Images were analyzed using NIS-Elements imaging software (Nikon Instruments Inc., Melville, NY).

**Gene expression**

Total RNA was extracted from 20 mg liver tissue homogenate using 1ml Trizol reagent (Ambion, Carlsbad, CA, USA). The total RNA concentration was calculated by measuring the absorbance at 260 nm and 280 nm using an ultraviolet spectrophotometer. For cDNA synthesis, Maxi RT PreMix kit (iNtRON, Korea) was used according to the manufacturer’s instructions. The PCR was set using the following program: 95°C for 2 min, 95°C for 30 sec 38-40 cycle, the appropriate annealing temperature between 55-57°C for 30 sec and 72°C for 2 min. After loaded onto 1% of the agarose gel containing ethidium bromide then measured the PCR band density with Image Lab 4.0 (Bio-Rad, USA). The PCR primer sequences for each studied gene are shown in Table 1.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward</th>
<th>Reverse</th>
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<tbody>
<tr>
<td>NLRP</td>
<td>GCTCCAACCACCTTCTGAC</td>
<td>AGTTACACTGTGGTGCCTTT</td>
</tr>
<tr>
<td>ASC</td>
<td>CAGGTATTGGCCATCACTCAGT</td>
<td>TCCCATAGGTAAGGACACATAAATAA</td>
</tr>
<tr>
<td>Caspase1</td>
<td>GCAAAGAGGAAGCCATTTATCA</td>
<td>GCTTTGCTCATAGCAGTAAAT</td>
</tr>
<tr>
<td>II-1β</td>
<td>TCACAGGCAGAGCCACAAT</td>
<td>GAAACAGTGCAGCCATAC</td>
</tr>
<tr>
<td>PGC-1α</td>
<td>CGCTTTGCTGCTTCTTGA</td>
<td>AATTCGCTTGATGTTGATCTAGGT</td>
</tr>
<tr>
<td>BMP7</td>
<td>CCATCCTACGCCCACCACAT</td>
<td>GCCAAAACTGCCCATAC</td>
</tr>
<tr>
<td>UCP-1</td>
<td>TGGAGGTGTGGGACGAGTGT</td>
<td>GGAGATGAGAATGGGTCTACT</td>
</tr>
<tr>
<td>β-actin</td>
<td>TCACCCACACTGTGCCCTACGA</td>
<td>CAGCGAAACCCGCTATTGCCAATGG</td>
</tr>
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Statistical Analysis

Statistical analysis of the data was performed by SPSS Version 22.0 using one-way ANOVA with LSD post-hoc tests. Statistical significance was defined as \( \alpha = .05 \).

Results

The change of white adipose tissue and liver weight

Fig. 1 shows the epididymal fat and liver weight of high-fat fed mice. The epididymal fat weight was significantly increased in HD and Hch group compared with NC group \( (p<0.01) \). However, the weight was most effectively decreased in HME group compared with HD and Hch group \( (p<0.05) \). The liver weight was significantly increased in HD group compared with NC group \( (p<0.05) \). However, the weight was effectively decreased in HME and Hch group compared with HD group \( (p<0.05) \).

The change of fat cell size in white adipose tissue

Fig. 2 shows the fat cell size of high-fat fed mice. The epididymal fat cell size was significantly increased in HD group compared with NC group \( (p<0.05) \). However, epididymal fat cell size was significantly decreased in Hch and HME group compared with HD group \( (p<0.05) \).

The change of inflammasome genes in liver

Fig. 3 shows the effect of chrysin or exercise inflammasome genes in liver.

The NLRP3, caspase1, and IL-1\( \beta \) mRNA expression of HC group significantly increased compared to NC group. However, the NLRP3, caspase1, and IL-1\( \beta \) mRNA expression of Hch and HME group were significantly reduced compared with HC group \( (p<0.05) \). The ASC level was no significant difference among all groups.

The change of thermogenic genes in liver

Fig. 4 shows the effect of chrysin or exercise thermogenic genes in liver.

The PGC-1\( \alpha \) mRNA expression was significantly different between NC and HC group, but, HME group was significantly increased in those. The BMP7 mRNA expression was also significantly decreased in HC and Hch group compared with NC group. However, HME group was effectively increased for BMP7 mRNA expression. The UCP-1 mRNA expression was no significant difference among all groups.

Discussion

It is well known that moderate exercise has beneficial effects on the metabolic disorder [4]. Thus, physical activity has been recommended in the treatment of obesity [6]. Also, recent studies demonstrated that Chrysin has many functional roles, such as anti-inflammatory and antioxidant [8]. Nevertheless, information concerning the effects of chrysin in animal models is rather scant. Thus, it was meaningful to compare the effectiveness of moderate exercise and chrysin supplementation on inflammasome and thermogenic genes in high fat fed mice.

After 16 weeks of treatments, the epididymal fat, liver weight, and adipocyte size of HC group were significantly increased in comparison with NC group (Fig. 1, Fig. 2). But, HME group was significantly decreased in that’s comparison with HC group. These research results are consistent with the previous studies that the aerobic exercise allowed to pos-
Fig. 2. The change of adipocyte size of high-fat fed mice. Values are expressed as means ± SEM. NC: normal diet control; HC: high fat diet control; Hch: high fat diet with chrysin; HME: high fat diet with moderate exercise. *p<0.05 difference from NC group, #p<0.05 difference from HC group.

itively reduce the body and fat weight by increasing the energy consumption [9, 11]. It is suggesting that the moderate exercise can suppress the obesity induced weight gain.

In this study, we found that liver weight and adipocyte size of Hch group were effectively reduced comparison with HC group. But, no significant effect on fat pad weight comparison with HC group. The previous study reported that resveratrol supplements significantly reduced the epididymal fat pad weight [1]. In contrast, in another study this effect was not observed [1, 22]. Thus, in anti-agent studies, the results were still controversial and depended on the dosage and duration of the supplement. Thus, further studies are needed in the future.

The NLRP3 inflammasome is a key regulator linking metabolic dysfunction [5]. Once the NLRP3 inflammasome is activated, activated caspase1 cleaves a variety of protein precursors [21]. We found that induction of high-fat fed induced NLRP3, IL-1β, and Caspase-1 expression in liver. On the contrary to this, NLRP3, IL-1β, and Caspase-1 mRNA expressions were significantly decreased in Hch and HME group comparison with HC group. These results have suggested that the chrysin supplements or moderate exercise have the function to suppress the expression of inflamma-
some related genes in hepatocytes. It seems that the effec-

Fig 3. The change of inflammasome markers of high-fat fed mice. (A) NLRP3; (B) ASC; (C) Caspase1; (D) IL-1β. Values are expressed as means ± SEM. NC: normal diet control; HC: high fat diet control; Hch: high fat diet with chrysin; HME: high fat diet with moderate exercise. *p<0.05 difference from NC group, #p<0.05 difference from HC group.
tiveness on the regulation of inflammasome molecules depends on whether there is a significant liver weight loss after intervention. The future study measuring the tissue using the inhibitors of NLRP3 can determine the direct roles of inflammasome in obesity-associated metabolic dysfunction.

Peroxisome proliferator activated receptor gamma co-activator 1α (PGC-1α), have been identified as the main regulators of mitochondrial biogenesis and thermogenesis [7]. Also, BMP7 inhibits fibrosis in animal models of chronic liver disease [17]. Previous experiment results indicated that BMP7 promotes hepatocyte regeneration and inhibits liver fibrosis [20].

In our results, the PGC-1α and BMP7 were significantly decreased in high-fat fed group compared to the normal diet group. However, PGC-1α and BMP7 were significantly increased only in moderate exercise group.

Regarding the improve effect of moderate exercise on PGC-1α and BMP7 activation in high-fat fed, we suggest several pathways. First, in the exercise-induced danger signals, such as free fatty acids, cholesterol, etc., that promotes the NLRP inflammasome activation [32]. In our results, demonstrated that moderate exercise reduce inflammasome related gene in liver of high-fat fed mice (Fig. 3). Second, previous study suggested that aerobic exercise potently stimulates fatty-acid oxidation in variety tissue by inhibiting the activity of lipid synthesis related markers [15]. Although, our study did not measure this metabolite factors in this study. It will be of interest for further researchers to include. In conclusion, chrysin and moderate exercise have a positive effect on high-fat-diet-induced obese metabolic complication by reducing inflammasome genes. However, there was no effect of chrysin supplementation on the thermogenic gene expression. It seems that moderate exercise is more effective to obese-induced metabolic deregulation.

Acknowledgment

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References


초록: 고지방식이 동물의 간 조직에서 크리신 투여 또는 중강도 운동이 Inflammasome과 열 발생 유전자 발현에 미치는 효과
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본 연구 목적은 고지방식이 동물의 간 조직에서 크리신 투여 또는 중강도 운동이 Inflammasome과 thermo-genesis 유전자 발현의 차이를 규명하고자 시도되었다. 본 연구를 위해 정상식이군, 고지방식이군, 고지방식이+크리신 투여군, 고지방식이+중강도 운동군으로 분류한 후, 크리신 투여군은 16주간 50 mg/kg 농도로 투여하였으며, 운동군은 최대산소섭취량의 60-75%의 중강도 운동으로 실시되었다. 연구결과 크리신 그리고 중강도 운동군은 지방조직, 간조직 무게 그리고 지방세포 크기의 변화를 고지방식이 군과 비교하여 유의하게 감소하였다. Inflammasome 유전자 발현은 크리신 투여군 그리고 중강도 운동군에서 NLRP3, ASC, Caspase1 mRNA 발현이 고지방식이 군과 비교하여 유의하게 감소하였다. 열발생마커로 알려진 PGC-1a, BMP7 mRNA 발현은 중강도 운동군에서만 고지방식이 군과 비교하여 유의하게 증가하였다. 결론적으로 중강도 운동은 고지방식이 동물에서 지방무게, Inflammasome, 그리고 열발생 유전자 발현의 차이를 비교하여 유의하게 감소하였고, 본 연구에서는 크리신 투여 및 중강도 운동이 고지방식이 동물에서 지방조직 무게, Inflammasome, 그리고 열발생 유전자 발현의 차이를 비교하여 유의하게 감소하였고, 본 연구에서는 크리신 투여 및 중강도 운동이 고지방식이 동물에서 지방조직 무게, Inflammasome, 그리고 열발생 유전자 발현의 차이를 비교하여 유의하게 감소하였고, 본 연구에서는 크리신 투여 및 중강도 운동이 고지방식이 동물에서 지방조직 무게, Inflammasome, 그리고 열발생 유전자 발현의 차이를 비교하여 유의하게 감소하였고, 본 연구에서는 크리신 투여 및 중강도 운동이 고지방식이 동물에서 지방조직 무게, Inflammasome, 그리고 열발생 유전자 발현의 차이를 비교하여 유의하게 감소하였고, 본 연구에서는 크리신 투여 및 중강도 운동이 고지방식이 동물에서 지방조직 무게, Inflammasome, 그리고 열발생 유전자 발현의 차이를 비교하여 유의하게 감소하였고, 본 연구에서는 크리신 투여 및 중강도 운동이 고지방식이 동물에서 지방조직 무게, Inflammasome, 그리고 열발생 유전자 발현의 차이를 비교하여 유의하게 감소하였고, 본 연구에서는 크리신 투여 및 중강도 운동이 고지방식이 동물에서 지방조직 무게, Inflammasome, 그리고 열발생 유전자 발현의 차이를 비교하여 유의하게 감소하였고, 본 연구에서는 크리신 투여 및 중강도 운동이 고지방식이 동물에서 지방조직 무게, Inflammasome, 그리고 열발생 유전자 발현의 차이를 비교하여 유의하게 감소하였고, 본 연구에서는 크리신 투여 및 중강도 운동이 고지방식이 동물에서 지방조직 무게, Inflammasome, 그리고 열발생 유전자 발현의 차이를 비교하여 유의하게 감소하였다. 본 연구에서는 크리신의 비만억제 효과를 규명하기 위해 투여농도 및 기간을 고려하여 다양한 연구가 진행되어야 할 것이다.