Effect of Cordyceps sinensis and exercise training on hepatic damage in sepsis
摘要

冬蟲夏草是應用於治療多種發炎性疾病的中草藥，有一些研究文獻指出運動訓練可以降低敗血症帶來的傷害。本篇的目的是研究冬蟲夏草合併運動訓練對於敗血症引起之肝臟傷害的影響。實驗使用三十二隻小鼠(C57BL/6 mice)，隨機分成八組，其中四組小鼠沒有進行運動訓練，小鼠管餵冬蟲夏草的劑量分成: 0mg(saline)、20mg、40mg，在運動訓練結束後使用脂多糖(LPS)/半乳糖胺(D-GalN)誘導敗血症，取出血液與肝臟測量蘇木紫-伊紅染色(Hematoxylin-Eosin Y staining)、天冬氨酸轉氨酶(aspartate aminotransferase, AST)、腫瘤壞死因子-α(TNF-α)、介白素-10(IL-10)、超氧化物歧化酶(superoxide dismutase, SOD)、一氧化氮(nitric oxide, NO)、肝臟細胞凋亡表現蛋白、TUNEL陽性細胞。實驗結果表示單獨進行運動訓練增加了IL-10、SOD並降低AST、TNF-α、NO，減少肝臟凋亡表現蛋白與TUNEL陽性細胞。只給予高劑量冬蟲夏草可以增加IL-10、SOD並降低AST、TNF-α、NO，減少肝臟凋亡表現蛋白與TUNEL陽性細胞。高劑量冬蟲夏草合併運動訓練降低了保護效果，增加AST、TNF-α、NO產生，減少IL-10、SOD並活化肝臟凋亡表現蛋白與增加TUNEL陽性細胞產生。總結是進行運動訓練或是高劑量蟲草可以降低敗血症引起的肝臟傷害，而高劑量冬蟲夏草合併運動訓練則降低了保護效果。
Abstract

*Cordyceps sinensis* (*C.sinensis*) has been considered to be an herbal medicine used in various inflammatory diseases, and some studies show the exercise training can decrease septic responses. The purpose of this study was evaluated whether *C.sinensis* and exercise training could reduce liver damage in septic mice synergistically. Tumor necrosis factor – alpha (TNF-α), interleukin-10 (IL-10), aspartate aminotransferase (AST), superoxide dismutase(SOD), nitric oxide(NO), apoptotic proteins analyzed by Western Blotting, TUNEL assay and H&E stain were tested in thirty two male C57BL/6 mice which were randomly assigned into control (without lipopolysaccharide (LPS) combined with D-galactosamine(D-GalN) infusion), saline, CS20mg and CS40mg with or without pre-condition swimming exercise training. Exercise treatment alone can increase LPS/ D-GalN - induced IL-10 production and decrease TNF-α level and apoptosis in liver. High dose CS treatment alone can also increase the levels of IL-10 and decrease the level of TNF-α, AST and hepatic apoptosis. High dose *C.sinensis* treatment combined with exercise training attenuated the protective effect, increased the liver damage and apoptosis responses. In summary, exercise training alone or *C.sinensis* treatment alone decreased injury induced by sepsis. Exercise training combined *C.sinensis* treatment decreased the protective effect and induced liver damage in sepsis.

Key word: TNF-α, Lipopolysaccharide, D-galactosamine, Hepatic injury
CONTEXT

Effect of Cordyceps sinensis and exercise training on hepatic damage in sepsis

Introduction

Materials and Methods

Animal groups

Cordyceps sinensis treatment

Precondition exercise training

Sepsis-induced protocols

Hematoxylin-Eosin Y Staining

Biochemistry study

Enzyme-linked immunosorbent assay (ELISAs)

Measurement of nitric oxide (NO)

Superoxide dismutases (SOD) assay

Western blot analysis

Terminal deoxynucleotidyl transferase-mediated dUTP nick end labeling (TUNEL)

Statistical analysis

Results

Discussion

References

Figures
FIGURE CONTENTS

Fig 1. Effects of *C. sinensis* and exercise training on H&E stain .....34
Fig 2. Effects of *C. sinensis* and exercise training on AST (Aspartate Aminotransferase) value .............................................35
Fig 3. Effects of *C. sinensis* and exercise training on serum TNF-α concentration .........................................................36
Fig 4. Effects of *C. sinensis* and exercise training on serum IL-10 concentration ..............................................................37
Fig 5. Effects of *C. sinensis* and exercise training on NO production of three hours serum ..................................................38
Fig 6. Effects of *C. sinensis* and exercise training on NO production of six hours serum .....................................................39
Fig 7. Effects of *C. sinensis* and exercise training on TNF-α concentration of liver tissue ....................................................40
Fig 8. Effects of *C. sinensis* and exercise training on IL-10 concentration of liver tissue ....................................................41
Fig 9. Effects of *C. sinensis* and exercise training on NO production of liver tissue ..........................................................42
Fig 10. Effects of *C. sinensis* and exercise training on SOD levels .....43
Fig 11. Effects of *C. sinensis* and exercise training on ERK (P44/42) protein level of liver tissue ..........................................44
Fig 12. Effects of *C. sinensis* and exercise training on p38 protein level of liver tissue ..........................................................46
Fig 13. Effects of *C. sinensis* and exercise training on cleaved-caspase 3 protein level of liver tissue ....................................48
Fig 14. Effects of *C. sinensis* and exercise training on cleaved-caspase 6 protein level of liver tissue…………………………………………..................50

Fig 15. Effects of *C. sinensis* and exercise training on cleaved-PARP protein level of liver tissue……………………………………………………52

Fig 16. Effects of *C. sinensis* and exercise training on hepatic TUNEL (terminal deoxynucleotidyl transferase (TdT)-mediated dUTP nick end labeling) staining …………………………………………………………... 54
Effect of Cordyceps sinensis and exercise training on hepatic damage in sepsis
INTRODUCTION

1. Sepsis

Sepsis is a severe inflammatory disorder, and serious clinical problem with high mortality may lead to severe shock and multiple organ failure. The interaction between infecting microorganism and host immune, inflammation, and coagulation responses results in sepsis. It is defined as the systemic response to infection, with the most common cause being a contamination of the blood with bacteria [37].

The infection activates various mechanisms to cause tissue damage, including cytokines production that initiates the inflammatory response and changes in the systemic circulation and coagulation cascade. An unregulated response to multiple organ failure trigger by sepsis is characterized by liver, cardiovascular, pulmonary, renal and gastrointestinal dysfunction. In particular, the severe liver failure after sepsis is associated with a high mortality rate [9].

Lipopolysaccharide (LPS) has been used for induction of endotoxemia in animal experimentation and is responsible for the multiple organ dysfunctions induced by septic shock. LPS from gram negative bacteria bound with toll-like receptors (TLRs) can arrange other signaling molecules to release cytokines [25]. This process involves the activation of mitogen activated protein kinase (MAPK) [31]. Cytokines then cause oxidant stress production and tissue injury eventually. The pro-inflammatory cytokines can decrease systemic vascular resistance, resulting in profound hypotension by inducing the expression of inducible nitric oxide synthase (iNOS) and augmenting the production of nitric
oxide (NO) [39][46]. D-galactosamine (D-GalN) is a hepatotoxic agent, which increases the susceptibility of mice to LPS-induced shock by impairing liver metabolism [28].

2. *Sepsis-induced liver dysfunction*

Liver appears to be a key organ in sepsis, and portal blood flow is a subject to vasoconstriction and bacterial translocation during sepsis. The liver is made mainly of hepatocytes, Kupffer cells, and endothelial sinusoidal cells. All of these cell types are involved in the metabolic, anti-infectious and immune responses through multiple cell interactions. Kupffer cells exposed to endotoxin *in vitro* produce large amounts of tumor necrosis factor-α (TNF-α). TNF-α is also involved in induction of the nitric oxide synthase gene (*iNOS*) transcription, accounting for nitric oxide hepatic production [9].

Sepsis-induced liver dysfunction is usually attributed to systemic disturbances, endotoxin, the activation of inflammatory cytokines, and mediators. The overproduction of locally secreted and circulating mediators, such as reactive oxygen species, proinflammatory cytokines, and nitric oxide formation leading to tissue damage and cell death which contributed to liver injury [9].

3. *Role of pro-inflammatory cytokines in sepsis*

Tumor necrosis factor- alpha (TNF-α) is recognized as an important mediator in the development of sepsis and in the systemic inflammatory response syndrome (SIRS) [32]. While Toll like receptor-4 (TLR-4) recognizes the LPS of gram-negative bacteria, binding of TLRs stimulates intracellular signaling which increase the secretion of proinflammatory molecules such as TNF-α and interleukin-1β, as well as
antiinflammatory cytokines such as interleukin-10 [37].

Macrophages represent a major defense system against attack of the host by microorganisms and release a vast group of mediators including cytokines, such as TNF-α and IL-1-β, arachidonic acid metabolites, and NO that amplify the inflammatory response. TNF-α has been proposed as a therapeutic target because it plays a key role in arranging the cytokine cascade in many inflammatory diseases and it is a “master-regulator” of inflammatory cytokine production [25][32]. Proinflammatory cytokines up-regulate adhesion molecules in neutrophils and endothelial cells. Activated neutrophils exterminate microorganisms and damage endothelium by releasing mediators that increase vascular permeability. In addition, activated endothelial cells release nitric oxide, a potent vasodilator that acts as a key mediator of septic shock [3][12].

4. Role of apoptosis in sepsis

Apoptosis, commonly termed programmed cell death, can proceed by the death receptor or mitochondrial-mediated pathways that ultimately converge to activate executioner caspases including caspase-3 and caspase-6 that mediate the systematic demolition of the cell [7]. Apoptotic cell death plays an important role in LPS-related tissue injuries in clinical systemic inflammatory response syndrome, septic shock, and multiple organ failure [13]. The inflammatory and metabolic changes develop in septic patients and change the various organs into pathophysiology.

Endotoxin and various ROS elicit DNA strand breaks, which in turn activate Poly (ADP-ribose) polymerase (PARP). The role for PARP in DNA repair suggests that PARP activation serves to help rescue damaged
cells and this can occur following mild DNA damage. However, the opposite is true in excessive DNA damage. Cleavage of PARP might occur when cell is no longer able to repair its DNA and it ensures commitment to apoptosis. The excess NO production in Septic shock also activates PARP and damages DNA [34].

5. Role of nitric oxide (NO) in sepsis

Nitric oxide (NO) is involved in smooth muscle relaxation, and increased NO release contributes to sepsis-induced hypotension. NO is produced by three different NO synthases (NOS); neuronal NOS (nNOS), inducible NOS (iNOS) and endothelial NOS (eNOS). Under normal circumstances, iNOS activity stays in low levels, but its synthesis is rapidly stimulated by a wide range of inflammatory agents with endotoxin [5].

Nitric oxide (NO) is an intracellular messenger because it can diffuse through most cells and tissues with consumption or direct reaction [1]. NO is generated from L-arginine and iNOS produces large amounts of NO over prolonged periods of time. The NO metabolites nitrite and nitrate, which is the indicators of NO production, raise progressively in various animal shock models. The enhanced formation of NO contributes to the hypotension caused by endotoxin and TNF-α. NO may also directly contribute to tissue and organ injury by its direct, peroxynitrite-mediated cytotoxic effects [5][6][12].

6. The effect of Cordyceps sinensis in sepsis

Cordyceps fungi, such as Cordyceps sinensis (C. sinensis), is a parasitic complex of fungus and caterpillar, which has been utilized as a traditional medicine and used for medicinal purposes for centuries
particularly in Asia. *C. sinensis* has many bioactive components, such as 3’-deoxyadenosine (cordycepin), cordycepic acid, and Cordyceps polysaccharides [45].

*C. sinensis* shows various biological effects, e.g. down-regulation of inflammation-related and apoptotic genes [40], anti-tumor activity [8], and suppression of autoimmune diseases and allergy with their immunomodulating effects [45]. *C. sinensis* has also been used to treat liver, inflammatory and autoimmune diseases [22][23][24]. *C. sinensis* acts to accelerate macrophage phagocytosis, enhances the immune function and reverse the developing process of liver fibrosis [24].

The anti-inflammatory effect of *C. sinensis* might result from the inhibition of TNF-α [35][43]. It has been shown that *Cordyceps pruinosa* methanol extract suppresses inflammation through suppression of NF-κB-dependent inflammatory gene expression, which suggesting that the *Cordyceps pruinosa* extract may be beneficial for treatment of endotoxin shock or sepsis [15]. However, the mechanisms of *C. sinensis* contribute to the beneficial effect for hepatic dysfunction induced by sepsis is unknown.

**7. The effects of exercise in sepsis**

In general, regular exercise can enhance immunological and antioxidant functions [27]. The mechanisms of exercise-associated immune changes are multifactorial and include multiple neuroendocrinological factors. Alterations in metabolism and metabolic factors contribute to exercise-associated changes in immune function. Exercise has important modulatory effects on immunocyte dynamics and possibly on immune function [2]. These effects are mediated by
exercise-induced release of proinflammatory cytokines, classical stress hormones, and hemodynamic effects leading to cell redistribution [33].

Chen et al. [4] suggested that exercise training attenuates septic responses and protects organs from damage in sepsis. The adaptations promoted by swimming training seem to be beneficial, counteracting the cardiovascular abnormalities and pulmonary edema seen in septicemia induced by LPS [26]. The appropriate exercise in humans increases the plasma cytokine IL-6 and suppresses the endotoxemia-induced elevation in TNF-α [41]. They conclude that exercise exerts anti-inflammatory effects. Because the anti-inflammatory effect of C. sinensis and exercise training attenuates septic responses and protects organs from damage, the present study was designed to test that C. sinensis combined with precondition exercise training can attenuate hepatic injury in sepsis.
MATERIALS AND METHODS

Animal groups
Thirty-two male C57BL/6 mice (weighing 20-25g) were obtained from the National Laboratory Animal Center (NLAC), Taiwan. Mice were randomly assigned into control (without LPS combined with D-galactosamine (D-GalN) infusion), saline, CS20mg and CS40mg with or without pre-condition swimming exercise training.

Cordyceps sinensis treatment
The extract of C. sinensis is purchased from TCM Biotech International Corp., Taiwan. Mice were treated with saline or the extract of C. sinensis (20 mg/kg or 40 mg/kg) by oral gavage 4 hours before the precondition swimming exercise training.

Precondition exercise training
Four hours after C. sinensis treatment, mice were treated with precondition exercise training. The exercise protocol is swimming exercise for 30 minutes/day, 5days/week for 4 weeks in 26-28°C water.

Sepsis-induced protocols
At the end of treatment, mice were induced sepsis by LPS with D-GalN through intraperitoneally injection (LPS: 10μg/kg, Escherichia coli, B4, 10; D-GalN: 800 mg/kg, Sigma-Aldrich, St. Louis, MO, USA.).

Hematoxylin-Eosin Y staining
The livers were excised and soaked in formalin and processed to paraffin blocks. Slides were prepared by deparaffinization and rehydration. They were passed through a series of graded alcohols (100%, 95%, and 75%), for 15 minutes each. The slides were then dyed with
hematoxylin and eosin for 1 minute. After being gently rinsed with water, slides were then soaked with 85% alcohol, two changes of 100% alcohol for 15 minutes each. At the end, the slides were soaked in two changes of Xylene. Photomicrographs were obtained using Zeiss AxioImager M1 microscope (Carl Zeiss MicroImaging GmbH, Jena, Germany).

**Biochemistry study**

Blood samples were collected at 6 hours after LPS/D-GalN infusion, and immediately centrifuged at 5000g for 10 min. Plasma was stored at 4°C for biochemical examination after collection. Plasma samples were diluted 1:10 with distilled water before measurements. Aspartate aminotransferase (AST) were evaluated of liver failure.

**Enzyme-linked immunosorbent assay (ELISAs)**

TNF-α and IL-10 was measured using antibody enzyme-linked immunosorbent assays (ELISAs). The experiment was repeated with blood collected at 3 and 6 hours for cytokine analysis by intracardiac puncture and immediately allowed to centrifugation for 10 min at 5000g. The liver tissue was collected for analysis at 6 hours after LPS/D-GalN treatment. Liver tissue was homogenate and allowed to centrifugation for 20 min at 5000g. The supernatant was stored at −80°C for determination of IL-10 and TNF-α concentration using commercially available enzyme-linked immunosorbent assay kits (R&D Systems, Minneapolis, MN, USA. Duo Set ELISA development kit) according to the manufacturer’s instructions. The optical density was measured at 450 nm wavelengths by Multiskan EX Microplate Photometer (Thermos, Waltham, MA, USA).
**Measurement of nitric oxide (NO)**

The NO concentration in serum and liver were measured after the Griess reaction by incubating 20μL sample with 20μL of Griess reagent (Sigma-Aldrich, St. Louis, MO, USA) and added 160μL deionized water at room temperature for 20 minutes. The absorbance was measured at 550 nm and nitrite concentration was calculated by comparison with a standard solution of known sodium nitrite concentrations.

**Superoxide dismutases (SOD) assay**

The enzyme activity of SOD was measured in 10μL of liver sample using kit from Cayman Chemical (Ann Arbor, MI, USA). The method utilizes tetrazolium salt to quantifying superoxide radicals generated by xanthine oxidase and hypoxanthine. The standard curve was generated using quality controlled SOD standard and the optical density was measured at 450 nm wavelengths.

**Western Blot analysis**

Mice liver tissue was harvested after 6 hours following LPS/ D-GalN treatment. The tissue were homogenized and lysed for 30 min in lysis buffer (homogenize tissue sample in 1:20 (w/v) of tissue to T-PER Reagent (Thermos, Waltham, MA, USA).

Samples were then centrifuged and supernatant was collected. Protein quantification is measured by Bradford method. Western blotting is analysis for cleaved caspase-3, cleaved caspase-6, cleaved PARP (Poly (ADP-ribose) polymerase), p38 MAPKs and ERK (p44/42) protein expression (Cell Signaling Technology, Danvers, MA, USA). 150 μg proteins were separated by 12% sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) with a constant voltage 80V and
transferred onto polyvinylidene difluoride (PVDF) membranes (Bio-Rad Laboratories, CA, USA) with a transfer apparatus (Bio-Rad Laboratories, CA, USA).

The membranes were blocked in 0.05% TBS-Tween (TBST) containing 5% milk and then incubated with the primary antibody (1:1,000) in TBST with 5% skim milk at 4 °C overnight. The primary antibody was detected by incubation with horseradish peroxidase-coupled second antibody (1:2,000) at room temperature for 2 h. The chemiluminescence detection was performed by using Western Chemiluminescence HRP substrate (Millipore, Billerica, MA, USA). The blots were then stripped and incubated with anti β-actin antibody (diluted 1:10000; Sigma-Aldrich, St. Louis, MO, USA) to ensure equal loading. Quantitative analysis of bands carried out using Gel-PRO analyzer Imaging System.

**Terminal deoxynucleotidyl transferase (TdT)-mediated dUTP nick end labeling (TUNEL)**

ApopTag Peroxidase Kits (Millipore, Billerica, MA, USA) was used to label apoptotic cells. Paraffin slides were deparaffinized and pretreated with diluted Proteinase K. After washed the specimen in 2 changes of PBS, the specimen was put into 3.0% hydrogen peroxide in PBS for 5 minutes and rinsed twice with PBS for 5 minutes each time. The specimen was applied 75 μL/ 5cm2 of Equilibration buffer immediately and incubated for 10 seconds at room temperature. Next, the specimen was applied on TdT Enzyme and incubated in a humidified chamber at 37°C for 1 hour.

After an hour, the specimen was applied Stop/Wash buffer and
incubated for 10 minutes. After washed the specimen in 3 changes of PBS for 1 minute each wash, the slides were applied room temperature anti-digoxignenin conjugate and incubated in a humidified chamber for 30 minutes. The specimen was developed color in peroxidase substrate after wash in PBS and counter stained specimen in 0.5% (w: v) methyl green for 10 minutes. The specimen was dehydrated by moving the slide through three changes of xylene for 2 minutes each wash and mount under a glass coverslip in a mounting medium.

Photomicrographs were obtained using Zeiss AxioImager M1 microscope (Carl Zeiss MicroImaging GmbH, Jena, Germany). All counts were performed by at least two independent individuals in a blinded manner.

**Statistical analysis**

The data are presented as mean ± SEM. Statistical evaluation was performed by one way-ANOVA and Bonferroni’s post hoc test (Systat Software Inc., SigmaPlot, San Jose, CA, USA). *p*<0.05 was considered to be statistically significant.
RESULTS

Effects of C. sinensis and exercise training on H&E stain in LPS/ D-GalN-treated mice

Light microscopy showed no infiltration or sequestration of neutrophils in liver in control group (Fig. 1), whereas more overt sequestration of neutrophils and coagulative necrosis tissues were found in liver in saline without exercise group. The neutrophil infiltrations and liver necrosis were significantly reduced in 40mg C. sinensis without exercise group (n =4). Administration of exercise training only decreased LPS/ D-GalN-induced neutrophil infiltrations and liver necrosis compared with non-exercise group (n=4). Administration of 20 mg C. sinensis combined with exercise training decreased LPS/ D-GalN-induced neutrophil infiltrations and liver necrosis (n=4). However, 40 mg C. sinensis combined with exercise training significantly increased neutrophil infiltrations and liver necrosis (n=4) compared to 40 mg C. sinensis without exercise group.

Effects of C. sinensis and exercise training on AST (Aspartate Aminotransferase) value in LPS/ D-GalN- treated mice

AST expression in serum was measured at 6 hours after LPS/ D-GalN challenge to examine the effect of C. sinensis and exercise training (Figure 2). AST concentration in LPS/ D-GalN- treated groups was significantly increased than control group (p < 0.05), and significantly decreased with 40mg C. sinensis in non-exercise group (p < 0.05). Administration of exercise training only strongly decreased LPS/ D-GalN- induced AST elevation compared with non-exercise group (p <
Administration of 20 mg *C. sinensis* only did not decreased LPS/ D-GalN -induced AST elevation and there was no difference in 20 mg *C. sinensis* combined with exercise training group. However, 40 mg *C. sinensis* combined with exercise training significantly increased AST concentration compared to 40 mg *C. sinensis* without exercise group (*p* < 0.05).

**Effects of *C. sinensis* and exercise training on serum TNF-α concentration in LPS/ D-GalN -treated mice**

To examine the effect of *C. sinensis* and exercise training on TNF-α level in serum, three hours serum was measured by ELISA (Figure 3). Serum TNF-α concentration at 3 hours after LPS/ D-GalN challenge were significantly increased (*p* < 0.05), and increased dose of *C. sinensis* significantly decreased serum TNF-α in non-exercise group (all *p* < 0.05). Serum TNF-α concentration was significantly decreased in saline with exercise group (*p* < 0.05). Administration of 20 mg *C. sinensis* only decreased LPS/ D-GalN-induced TNF-α elevation (*p* < 0.05) and there is no difference in 20 mg *C. sinensis* combined with exercise training group. Administration of 40 mg *C. sinensis* only strongly decreased LPS/ D-GalN-induced TNF-α elevation (*p* < 0.05); however, 40 mg *C. sinensis* combined with exercise training increased TNF-α concentration compared to 40 mg *C. sinensis* without exercise group (*p* < 0.05).

**Effects of *C. sinensis* and exercise training on serum IL-10 concentration in LPS/ D-GalN -treated mice**

To examine the effect of *C. sinensis* and exercise training on IL-10 production of serum, six hours serum of mice was measured by ELISA (Figure 4). Serum IL-10 concentration at 6 hours after LPS/ D-GalN
challenge were significantly increased ($p < 0.05$), increased dose of *C. sinensis* significantly increased serum IL-10 in non-exercise group (all $p < 0.05$). Serum IL-10 concentrations at 6 hours after LPS/ D-GalN challenge were significantly increased in saline with exercise group ($p < 0.05$). Administration of 20 mg *C. sinensis* only increased LPS/ D-GalN-induced IL-10 elevation ($p < 0.05$) and there is no difference in 20 mg *C. sinensis* combined with exercise training group. IL-10 concentration after LPS/ D-GalN challenge were significantly decreased in 40 mg *C. sinensis* of exercise group compared with saline of exercise group ($p < 0.05$). Administration of 40 mg *C. sinensis* only strongly increased LPS/ D-GalN-induced IL-10 elevation ($p < 0.05$); however, 40 mg *C. sinensis* combined with exercise training decreased IL-10 concentration compared to 40 mg *C. sinensis* without exercise group ($p < 0.05$).

**Effects of *C. sinensis* and exercise training on NO production of three hours serum in LPS/ D-GalN-treated mice**

To examine the effect of *C. sinensis* and exercise training on NO production, three hours serum was measured by Griess Reaction (Figure 5). Serum nitrite concentration at 3 hours after LPS/ D-GalN challenge were significantly increased ($p < 0.05$), increased dose of *C. sinensis* significantly decreased serum nitrite in non-exercise group (all $p < 0.05$). Serum nitrite concentrations at 3 hours after LPS/ D-GalN challenge were significantly decreased in saline with exercise group ($p < 0.05$). Administration of 20 mg *C. sinensis* only decreased LPS/ D-GalN-induced nitrite elevation ($p < 0.05$) and there is no difference in 20 mg *C. sinensis* combined with exercise training group ($p < 0.05$).
Serum nitrite concentrations after LPS/ D-GalN challenge were significantly increased in 40 mg *C. sinensis* of exercise group compared with saline of exercise group (*p* < 0.05). Administration of 40 mg *C. sinensis* only strongly decreased LPS/ D-GalN-induced nitrite elevation (*p* < 0.05). However, 40 mg *C. sinensis* combined with exercise training increased nitrite concentration compared to 40 mg *C. sinensis* without exercise group (*p* <0.05).

**Effects of *C. sinensis* and exercise training on NO production of six hours serum in LPS/ D-GalN -treated mice**

To examine the effect of *C. sinensis* and exercise training on NO production, six hours serum was measured by Griess Reaction (Figure 6). Serum nitrite concentration at 6 hours after LPS/ D-GalN challenge were significantly increased (*p* < 0.05), increased dose of *C. sinensis* significantly decreased serum nitrite in non-exercise group (all *p* < 0.05). Serum nitrite concentrations at 6 hours after LPS/ D-GalN challenge were significantly decreased in saline with exercise group (*p* < 0.05). Administration of 20 mg *C. sinensis* only decreased LPS/ D-GalN-induced nitrite elevation (*p* < 0.05) and there is no difference in 20 mg *C. sinensis* combined with exercise training group. Serum nitrite concentrations after LPS/ D-GalN challenge were significantly increased in 40 mg *C. sinensis* of exercise group compared with saline of exercise group (*p* < 0.05). Administration of 40 mg *C. sinensis* only strongly decreased LPS/ D-GalN-induced nitrite elevation (*p* < 0.05); however, 40 mg *C. sinensis* combined with exercise training increased nitrite concentration (*p* < 0.05).
Effects of *C. sinensis* and exercise training on TNF-α concentration of liver tissue in LPS/ D-GalN -treated mice

To examine the effect of *C. sinensis* and exercise training on TNF-α production induced by LPS/D-GalN treatment, the excised liver of mice was measured by ELISA (Figure 7). Liver TNF-α concentration after LPS/ D-GalN challenge were significantly increased ($p < 0.05$), increased dose of *C. sinensis* significantly decreased liver TNF-α in non-exercise group (all $p < 0.05$). TNF-α concentration of liver after LPS/ D-GalN challenge were significantly decreased in saline with exercise group ($p < 0.05$). Administration of 20 mg *C. sinensis* only decreased LPS/ D-GalN-induced TNF-α elevation ($p < 0.05$) but there is no difference in 20 mg *C. sinensis* combined with exercise training group. TNF-α concentration of liver after LPS/ D-GalN challenge were significantly increased in 20 mg *C. sinensis* and 40mg *C. sinensis* of exercise group compared with saline of exercise group ($p < 0.05$). Administration of 40 mg *C. sinensis* only strongly decreased LPS/ D-GalN-induced TNF-α elevation ($p < 0.05$); however, 40 mg *C. sinensis* combined with exercise training increased TNF-α concentration ($p < 0.05$).

Effects of *C. sinensis* and exercise training on IL-10 concentration of liver tissue in LPS/ D-GalN -treated mice

To examine the effect of *C. sinensis* and exercise training on IL-10 production of liver was measured by ELISA (Figure 8). Liver IL-10 concentration after LPS/ D-GalN challenge were significantly increased ($p < 0.05$), increased dose of *C. sinensis* significantly increased liver IL-10 in non-exercise group (all $p < 0.05$). Liver IL-10 concentrations after LPS/ D-GalN challenge were significantly increased in saline with
exercise group ($p < 0.05$). Administration of 20 mg \textit{C. sinensis} only increased LPS/ D-GalN-induced IL-10 elevation ($p < 0.05$) and there is no difference in 20 mg \textit{C. sinensis} combined with exercise training group. IL-10 concentration of liver after LPS/ D-GalN challenge were significantly decreased in 20 mg \textit{C. sinensis} and 40 mg \textit{C. sinensis} of exercise group compared with saline of exercise group ($p < 0.05$). Administration of 40 mg \textit{C. sinensis} only strongly increased LPS/ D-GalN-induced IL-10 elevation ($p < 0.05$); however, 40 mg \textit{C. sinensis} combined with exercise training decreased IL-10 concentration compared to 40 mg \textit{C. sinensis} without exercise group ($p < 0.05$).

\textbf{Effects of \textit{C. sinensis} and exercise training on NO production of liver tissue}

To examine the effect of \textit{C. sinensis} and exercise training on NO production, liver tissue was measured by Griess Reaction (Figure 9). Liver nitrite concentration after LPS/ D-GalN challenge were significantly increased ($p < 0.05$), increased dose of \textit{C. sinensis} significantly decreased liver nitrite in non-exercise group (all $p < 0.05$). Liver nitrite concentrations after LPS/ D-GalN challenge were significantly decreased in saline with exercise group ($p < 0.05$). Administration of 20 mg \textit{C. sinensis} only decreased LPS/ D-GalN-induced nitrite elevation ($p < 0.05$) and there is no difference in 20 mg \textit{C. sinensis} combined with exercise training group. Liver nitrite concentrations after LPS/ D-GalN challenge were significantly increased in 40 mg \textit{C. sinensis} of exercise group compared with saline of exercise group ($p < 0.05$). Administration of 40 mg \textit{C. sinensis} only strongly decreased LPS/ D-GalN-induced nitrite elevation ($p < 0.05$); however, 40
mg *C. sinensis* combined with exercise training increased nitrite concentration (*p* < 0.05).

**Effects of *C. sinensis* and exercise training on SOD (superoxide dismutases) levels in LPS/ D-GalN-treated mice**

To examine the effect of *C. sinensis* and exercise training on SOD production, liver homogenate was measured by ELISA (Figure 10). SOD concentration after LPS/ D-GalN challenge were significantly decreased (*p* < 0.05), increased dose of *C. sinensis* significantly increased SOD level in non-exercise group (all *p* < 0.05). SOD concentrations after LPS/ D-GalN challenge were significantly increased in saline with exercise group (*p* < 0.05). Administration of 20 mg *C. sinensis* only increased LPS/ D-GalN-induced SOD elevation (*p* < 0.05) and there is no difference in 20 mg *C. sinensis* combined with exercise training group. Administration of 40 mg *C. sinensis* only strongly increased LPS/ D-GalN -induced SOD elevation (*p* < 0.05); however, 40 mg *C. sinensis* combined with exercise training decreased SOD concentration (*p* < 0.05).

**Effects of *C. sinensis* and exercise training on ERK (P44/42) protein level of liver tissue**

The effect of *C. sinensis* and exercise training on ERK pathway was measured by Western Blotting. Liver tissue was harvested and homogenate was analyzed after 6 hour of LPS/ D-GalN infusion (Figure 11). Treatment of 40mg *C. sinensis* significantly increased ERK protein in non-exercise group (*p* < 0.05). The phosphorylation of ERK after LPS/ D-GalN challenge increased in saline with exercise group (*p* < 0.05). the phosphorylation of ERK after LPS/ D-GalN challenge were no different between 20 mg *C. sinensis* without exercise group and 20 mg *C. sinensis*
with exercise group. Administration of 40 mg *C. sinensis* only strongly increased LPS/ D-GalN -induced ERK phosphorylation (*p* < 0.05); however, the phosphorylation of ERK after LPS/ D-GalN challenge were decreased in 40 mg *C. sinensis* combined with exercise group (*p* < 0.05).

**Effects of *C. sinensis* and exercise training on p38 protein level of liver tissue**

To examine the effect of *C. sinensis* and exercise training on activation p38 pathway, the liver tissue were analyzed by Western Blotting (Figure 12). Increased dose of *C. sinensis* significantly decreased p38 protein in non-exercise group (all *p* < 0.05) but the phosphorylation of p38 after LPS/ D-GalN challenge was no different in saline with exercise group. The phosphorylation of p38 after LPS/ D-GalN challenge increased in 20 mg *C. sinensis* and 40 mg *C. sinensis* with exercise group compared with saline with exercise group (*p* < 0.05). Administration of 20 mg *C. sinensis* only decreased LPS/ D-GalN-induced p38 elevation (*p* < 0.05). However, 20 mg *C. sinensis* combined with exercise training increased LPS/ D-GalN-induced p38 activation (*p* < 0.05). Administration of 40 mg *C. sinensis* only strongly decreased LPS/ D-GalN-induced p38 phosphorylation (*p* < 0.05); however, 40 mg *C. sinensis* combined with exercise training increased p38 activation (*p* < 0.05).

**Effects of *C. sinensis* and exercise training on cleaved-caspase 3 protein level of liver tissue**

The effect of *C. sinensis* and exercise training on cleaved-caspase 3 pathway was measured by Western Blotting. Liver tissue was harvested and homogenate was analyzed after 6 hours of LPS/ D-GalN infusion (Figure 13). Cleaved-caspase 3 protein levels after LPS/ D-GalN
challenge were significantly increased \((p < 0.05)\) and 40mg \(C. \textit{sinensis}\) treatment significantly decreased cleaved-caspase 3 protein activation in non-exercise group (all \(p < 0.05\)). There was no difference between 20 mg \(C. \textit{sinensis}\) alone treatment and 20 mg \(C. \textit{sinensis}\) combined with exercise training group. Administration of 40 mg \(C. \textit{sinensis}\) only strongly decreased LPS/ D-GalN -induced cleaved-caspase 3 activation \((p < 0.05)\); however, 40 mg \(C. \textit{sinensis}\) combined with exercise training increased cleaved-caspase 3 activation \((p < 0.05)\).

**Effects of \(C. \textit{sinensis}\) and exercise training on cleaved-caspase 6 protein level of liver tissue**

The effect of \(C. \textit{sinensis}\) and exercise training on cleaved-caspase 6 pathway was measured by Western Blotting. Liver tissue was harvested and homogenate was analyzed after 6 hours of LPS/ D-GalN infusion (Figure 14). Cleaved-caspase 6 protein level after LPS/ D-GalN challenge were significantly increased \((p < 0.05)\). 40mg of \(C. \textit{sinensis}\) significantly decreased cleaved-caspase 6 protein activation in non-exercise group \((p < 0.05)\). Cleaved-caspase 6 protein after LPS/ D-GalN challenge were significantly decreased activation in saline with exercise group \((p < 0.05)\). There was no difference between 20 mg \(C. \textit{sinensis}\) alone treatment and 20 mg \(C. \textit{sinensis}\) combined with exercise training group. Cleaved-caspase 6 protein after LPS/ D-GalN challenge were significantly increased activation in 20mg \(C. \textit{sinensis}\) and 40 mg \(C. \textit{sinensis}\) with exercise group compared with saline with exercise group \((p < 0.05)\). Administration of 40 mg \(C. \textit{sinensis}\) only strongly decreased LPS/ D-GalN -induced cleaved-caspase 6 activation \((p < 0.05)\); however, 40 mg \(C. \textit{sinensis}\) combined with exercise training increased
cleaved-caspase 6 activation ($p < 0.05$).

**Effects of C. sinensis and exercise training on cleaved-PARP (Poly ADP ribose polymerase) protein level of liver tissue**

The effect of C. sinensis and exercise training on cleaved-PARP pathway was measured by Western Blotting. Liver tissue was harvested and homogenate was analyzed after 6 hours of LPS/ D-GalN infusion (Figure 15). Cleaved-PARP protein level after LPS/ D-GalN challenge were significantly increased ($p < 0.05$) and 40mg C. sinensis treatment significantly decreased cleaved-PARP activation in non-exercise group (all $p < 0.05$). Cleaved-PARP protein after LPS/ D-GalN challenge were significantly decreased in saline with exercise group ($p < 0.05$). There was no difference between 20 mg C. sinensis alone treatment and 20 mg C. sinensis combined with exercise training group. Cleaved-PARP protein after LPS/ D-GalN challenge were significantly increased activation in 20 mg C. sinensis and 40 mg C. sinensis of exercise group compared with saline of exercise group ($p < 0.05$). Administration of 40 mg C. sinensis only strongly decreased LPS/ D-GalN -induced cleaved-PARP protein ($p < 0.05$); however, 40 mg C. sinensis combined with exercise training increased cleaved-PARP elevation ($p < 0.05$).

**Effects of C. sinensis and exercise training on hepatic TUNEL (terminal deoxynucleotidyl transferase (TdT)-mediated dUTP nick end labeling) staining**

To examine the effect of C. sinensis and exercise training on hepatic cell apoptosis induced by LPS/ D-GalN treatment, the liver tissue were measured by TUNEL staining (Figure 16). TUNEL positive cells after LPS/ D-GalN challenge were significantly increased ($p < 0.05$) and
increased dose of *C. sinensis* significantly decreased TUNEL positive cells in non-exercise group (all *p* < 0.05). TUNEL positive cells after LPS/ D-GalN challenge were significantly decreased in saline with exercise group (*p* < 0.05). Administration of 20 mg *C. sinensis* only decreased LPS/ D-GalN-induced TUNEL positive cells elevation (*p* < 0.05) and there is no difference in 20 mg *C. sinensis* combined with exercise training group. TUNEL positive cells after LPS/ D-GalN challenge were significantly increased in 40 mg *C. sinensis* of exercise group compared with saline of exercise group (*p* < 0.05). Administration of 40 mg *C. sinensis* only strongly decreased LPS/ D-GalN-induced TUNEL positive cells (*p* < 0.05); however, 40 mg *C. sinensis* combined with exercise training increased TUNEL positive cells (*p* < 0.05).
DISCUSSION

The main findings can be summarized as follows: (1) High dose *C. sinensis* treatment alone decreased the liver damage induced by sepsis. The AST value, TNF-α, and nitrite concentration in serum and liver were significantly decreased after LPS/ D-GalN infusion. The activation of apoptotic proteins including caspase-3, caspase-6, and PARP in liver tissue were significantly decreased. SOD activity and IL-10 level in serum and liver was also increased. (2) Preconditioning exercise training could decrease the damage induced by sepsis. The AST value, TNF-α level, nitrite concentration in serum and liver were significantly decreased. The phosphorylation of p38 protein and apoptotic proteins including cleaved-caspase 6 and cleaved-PARP levels in liver were significantly decreased after LPS/ D-GalN infusion. SOD activity and IL-10 level in serum and liver was significantly increased after LPS/ D-GalN infusion. (3) High dose *C. sinensis* treatment with preconditioning exercise training could not decrease the damage induced by sepsis. The AST value, TNF-α level, nitrite concentration in serum and liver were significantly increased. The phosphorylation of p38 protein and apoptotic proteins in liver were significantly increased after LPS/ D-GalN infusion. SOD activity and IL-10 level in serum and liver was significantly decreased after LPS/ D-GalN infusion.

Endotoxin and TNF-α cytokine contribute to the enhanced formation of NO. NO may cause tissue and organ injury by its cytotoxic effects. The overproduction of proinflammatory cytokines and NO lead to tissue damage and cell death which caused liver injury. In this study, *C. sinensis*
suppresses the expression of TNF-α and NO formation in liver of LPS/D-GalN stimulated mice. It implicated that *C. sinensis* decreased the tissue damage induced by sepsis through decreasing TNF-α production and NO formation. *C. sinensis* treatment with exercise training group increased the formation of NO and NO overproduction lead to the damage induced by sepsis.

*Cordyceps* species increases the activity of antioxidant enzymes such as superoxide dismutase in the liver [29]. Our results showing that *C. sinensis* treatment alone and precondition exercise training alone decreased the damage induced by sepsis through decreasing NO production and increasing SOD activity.

Our results showing that precondition exercise training may decrease the tissue damage induced by sepsis through decreasing TNF-α production, increasing IL-10 production, decreasing oxidative stress, further reducing hepatic apoptosis.

It is generally known that adequate and appropriate exercise is essential to maintain health and release work pressure. Some studies have demonstrated that regular exercise can enhance immunological and antioxidant functions [4][14][26]. Regular exercise induces the anti-inflammatory actions. During exercise, IL-6 is produced by muscle fibres. IL-6 stimulates the appearance in the circulation of other anti-inflammatory cytokines such as IL-10 and inhibits the production of the pro-inflammatory cytokine TNF-α [33].

Interestingly, exercise decreased the protective effect of high dose *C. sinensis* on liver damage induced by sepsis. AST value in mice received 40 mg *C. sinensis* alone were significantly decreased than saline of
non-exercise group. However, AST level in 40 mg *C. sinensis* with exercise increased significantly. The cleaved-caspase-3, 6 and cleaved-PARP protein levels in liver tissue of 40 mg *C. sinensis* with exercise group were also markedly increased. Because the TNF-α level in serum of 40 mg *C. sinensis* with exercise group was significantly increased than 40 mg *C. sinensis* alone.

The data suggested that higher NO production by exercise on high dose *C. sinensis* in liver might due to TNF-α induction. The NO production and SOD depletion was significantly increased in 40 mg *C. sinensis* with exercise group than 40 mg *C. sinensis* alone.

The controversial results of exercise combine *C. sinensis* treatment showed that physical exercise is new host factor with impact on hepatic drug metabolism. Moderate to heavy physical exercise for a few hours reduces liver blood flow leading to a decreased elimination of drugs exhibiting flow-limited metabolism [11]. The moderate physical exercise can prolong the half-life of indocyanine green for a few minutes, and the amount of prolongation increases with the duration of physical exercise. A reduction of liver blood flow amounting to 60% has been reported [36]. The physical exercise might change the metabolism of *C. sinensis* in liver and may lead to changes in drug efficacy and drug toxicity.

In summary, high dose *C. sinensis* treatment alone and precondition exercise training alone reduced the tissue damage induced by sepsis. The reverse effect of exercise on high dose *C. sinensis* in liver might due to high expression of TNF-α, NO accumulation, and caspases activation.
REFERENCES


Fig 1.

Effects of *C. sinensis* and exercise training on H&E stain in LPS/ D-GalN -treated mice (Magnifications: 400x). Mice were received *C. sinensis* (20 or 40 mg/kg /day) 4 hours before exercise and treated with precondition exercise training (NE: non-exercise, E: exercise) before administration of LPS (10 μg/kg) / D-GalN (800 mg/kg, intraperitoneally (i.p.)). Areas of necrosis are shown by arrows.
Effects of *C. sinensis* and exercise training on AST (Aspartate Aminotransferase) value in LPS/ D-GalN- treated mice. Mice were received *C. sinensis* (20 or 40 mg/kg /day) 4 hours before exercise and treated with precondition exercise training (NE: non-exercise, E: exercise) before administration of LPS (10 μg/kg)/ D-GalN (800 mg/kg, i.p.). The differences among those treatments with different letters were statistically significant (p<0.05). ✽ means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in NE group. + means CS20+LPS and CS40+LPS group compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in E group. & means the NE group compared with E group.
Fig 3.

Effects of *C. sinensis* and exercise training on serum TNF-α concentration in LPS/ D-GalN -treated mice. Mice were received *C. sinensis* (20 or 40 mg/kg/day) 4 hours before exercise and treated with precondition exercise training (NE: non-exercise, E: exercise) before administration of LPS (10 g/kg)/ D-GalN (800 mg/kg, i.p.). The differences among those treatments with different letters were statistically significant (p < 0.05). * means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in NE group. + means CS20+LPS and CS40+LPS group compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in E group. & means the NE group compared with E group.
Effects of *C. sinensis* and exercise training on serum IL-10 concentration in LPS/ D-GalN -treated mice. Mice were received *C. sinensis* (20 or 40 mg/kg /day) 4 hours before exercise and treated with precondition exercise training (NE: non-exercise, E: exercise) before administration of LPS (10 μg/kg)/ D-GalN (800 mg/kg, i.p.). The differences among those treatments with different letters were statistically significant (p < 0.05).

* means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in NE group. + means CS20+LPS and CS40+LPS group compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in E group. & means the NE group compared with E group. ^ means CS20+LPS and CS40+LPS group compared with Saline+LPS group in E group.
Fig 5.

Effects of *C. sinensis* and exercise training on NO production of three hours serum in LPS/ D-GalN-treated mice. Mice were received *C. sinensis* (20 or 40 mg/kg /day) 4 hours before exercise and treated with precondition exercise training (NE: non-exercise, E: exercise) before administration of LPS (10 µg/kg)/ D-GalN (800 mg/kg, i.p.). Serum NO which was measured by Griess Reaction was determined at 3 hours after LPS/D-GalN injection. Data are mean values ± SE (n=4). The differences among those treatments with different letters were statistically significant ($p < 0.05$). ✽ means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in NE group. + means CS20+LPS and CS40+LPS group compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in E group. & means the NE group compared with E group. ^ means CS20+LPS and CS40+LPS group compared with Saline+LPS group in E group.
Effects of *C. sinensis* and exercise training on NO production of six hours serum in LPS/D-GalN-treated mice. Mice were received *C. sinensis* (20 or 40 mg/kg/day) 4 hours before exercise and treated with precondition exercise training (NE: non-exercise, E: exercise) before administration of LPS (10 μg/kg)/D-GalN (800 mg/kg, i.p.). Serum NO which was measured by Griess Reaction was determined at 6 hours after LPS/D-GalN injection. Data are mean values ± SE (n=4). The differences among those treatments with different letters were statistically significant (p < 0.05). ✭ means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in NE group. + means CS20+LPS and CS40+LPS group compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in E group. & means the NE group compared with E group. ^ means CS20+LPS and CS40+LPS group compared with Saline+LPS group in E group.
Effects of *C. sinensis* and exercise training on TNF-α concentration of liver tissue in LPS/ D-GalN -treated mice. Mice were received *C. sinensis* (20 or 40 mg/kg /day) 4 hours before exercise and treated with precondition exercise training (NE: non-exercise, E: exercise) before administration of LPS (10 μg/kg)/ D-GalN (800 mg/kg, i.p.). The differences among those treatments with different letters were statistically significant (p < 0.05). ✽ means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in NE group. + means CS20+LPS and CS40+LPS group compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in E group. & means the NE group compared with E group. ^ means CS20+LPS and CS40+LPS group compared with Saline+LPS group in E group.
Fig 8.

Effects of *C. sinensis* and exercise training on IL-10 concentration of liver tissue in LPS/ D-GalN -treated mice. Mice were received *C. sinensis* (20 or 40 mg/kg /day) 4 hours before exercise and treated with precondition exercise training (NE: non-exercise, E: exercise) before administration of LPS (10 μg/kg)/ D-GalN (800 mg/kg, i.p.). The differences among those treatments with different letters were statistically significant (p < 0.05). ✽ means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in NE group. + means CS20+LPS and CS40+LPS group compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in E group. & means the NE group compared with E group. ^ means CS20+LPS and CS40+LPS group compared with Saline+LPS group in E group.
Effects of *C. sinensis* and exercise training on nitrite oxide production of liver in LPS/ D-GalN-treated mice. Mice were received *C. sinensis* (20 or 40 mg/kg/day) 4 hours before exercise and treated with precondition exercise training (NE: non-exercise, E: exercise) before administration of LPS (10 μg/kg)/ D-GalN (800 mg/kg, i.p.). NO concentration of liver tissue were determined at 6 hours after LPS/ D-GalN injection and measured by Griess Reaction. Data are mean values ± SE (n=4). The differences among those treatments with different letters were statistically significant (*p* < 0.05). ♠ means Saline+LPS, CS20+LPS and CS40+LPS compared with control group in NE group. + means CS20+LPS and CS40+LPS compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS compared with control group in E group. & means the NE group compared with E group. † means CS20+LPS and CS40+LPS group compared with Saline+LPS group in E group.
Effects of *C. sinensis* and exercise training on SOD (superoxide dismutases) levels in LPS/ D-GalN-treated mice. Mice were received *C.sinensis* (20 or 40 mg/kg /day) 4 hours before exercise and treated with precondition exercise training (NE: non-exercise, E: exercise) before administration of LPS (10 μg/kg)/ D-GalN (800 mg/kg, i.p.). SOD concentrations of liver tissue were determined at 6 hours after LPS/ D-GalN injection. Data are mean values ± SE (n=4). The differences among those treatments with different letters were statistically significant (p < 0.05). ✽ means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in NE group. + means CS20+LPS and CS40+LPS group compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS group compared with Saline+LPS group in E group. & means the NE group compared with E group. ^ means CS20+LPS and CS40+LPS group compared with Saline+LPS group in E group.
Fig 11. Effects of *C. sinensis* and exercise training on ERK (P44/42) protein level in LPS/D-GalN -treated mice.

(A) The representative protein products of phospho-ERK (p44/42, extracellular signal-regulated kinases) and total ERK extracted from liver in normal and LPS/ D-GalN-treated mice. Mice were received *C. sinensis* (20 or 40 mg/kg /day) 4 hours before exercise. Mice were received precondition exercise training (30 mins/day, 5 days/week; non-exercise (NE), exercise (E)) before administration of LPS (10μg/kg)/ D-GalN (800mg/kg, i.p.). Proteins were measured by Western Blotting analysis.
β-actin was used as an internal control. (B) Bars represent the protein quantification in phospho-ERK / total ERK and mean values ± SE (n=3 in each group). The differences among those treatments with different letters were statistically significant (p < 0.05). ✻ means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in NE group. + means CS20+LPS and CS40+LPS group compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in E group. & means the NE group compared with E group. ^ means CS20+LPS and CS40+LPS group compared with Saline+LPS group in E group.
Fig 12. Effects of *C. sinensis* and exercise training on p38 mitogen-activated protein kinase (MAPK) protein level in LPS/ D-GalN-treated mice

(A) The representative protein products of phospho-p38MAPK (Mitogen-activated protein kinase) and total p38 MAPK extracted from liver in normal and LPS/ D-GalN-treated mice. Mice were received *C. sinensis* (20 or 40 mg/kg/day) 4 hours before exercise. Mice were received precondition exercise training (30 mins/day, 5 days/week; non-exercise (NE), exercise (E)) before administration of LPS (10μg/kg)/
D-GalN (800mg/kg, i.p.). Proteins were measured by Western Blotting analysis. β-actin was used as an internal control. (B) Bars represent the protein quantification in phospho-p38 MAPK / total p38 MAPK and mean values ± SE (n=3 in each group). The differences among those treatments with different letters were statistically significant (p < 0.05). ✷ means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in NE group. + means CS20+LPS and CS40+LPS group compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in E group. & means the NE group compared with E group. ^ means means CS20+LPS and CS40+LPS group compared with Saline+LPS group in E group.
Fig 13. Effects of *C. sinensis* and exercise training on cleaved-caspase 3 protein level in LPS/ D-GalN -treated mice.

(A) The representative protein products of cleaved caspase-3 and β-actin extracted from liver in normal and LPS/ D-GalN- treated mice. Mice were received *C. sinensis* (20 or 40 mg/kg /day) 4 hours before exercise and treated with precondition exercise training (NE: non-exercise, E: exercise) before administration of LPS (10μg/kg)/ D-GalN (800mg/kg, i.p.). Proteins were measured by Western Blotting analysis. β-actin was used as an internal control. (B) Bars represent the protein quantification
in cleaved caspase-3 / β-actin and mean values ± SE (n=3 in each group). The differences among those treatments with different letters were statistically significant ($p < 0.05$). ✻ means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in NE group. + means CS20+LPS and CS40+LPS group compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in E group. & means the NE group compared with E group. ^ means means CS20+LPS and CS40+LPS group compared with Saline+LPS group in E group.
Fig 14. Effects of *C. sinensis* and exercise training on cleaved-caspase 6 protein level in LPS/ D-GalN -treated mice.

(A) The representative protein products of cleaved caspase-6 and β-actin extracted from liver in normal and LPS/ D-GalN-treated mice. Mice were received *C.sinensis* (20 or 40 mg/kg/day) 4 hours before exercise and treated with precondition exercise training (NE: non-exercise, E: exercise) before administration of LPS (10μg/kg)/ D-GalN (800mg/kg, i.p.). Proteins were measured by Western Blotting analysis. β-actin was used as an internal control. (B) Bars represent the protein quantification in cleaved caspase-6 / β-actin and mean values ± SE (n=3 in each group).
differences among those treatments with different letters were statistically significant \( (p < 0.05) \). ✽ means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in NE group. + means CS20+LPS and CS40+LPS group compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in E group. & means the NE group compared with E group. ^ means means CS20+LPS and CS40+LPS group compared with Saline+LPS group in E group.
Fig 15. Effects of *C. sinensis* and exercise training on cleaved-PARP (Poly ADP ribose polymerase) protein level in LPS/ D-GalN-treated mice.

(A) The representative protein products of cleaved-PARP (Poly (ADP-ribose) polymerase) and β-actin extracted from liver in normal and LPS/ D-GalN-treated mice. Mice were received *C.sinensis* (20 or 40 mg/kg /day) 4 hours before exercise and treated with precondition exercise training (NE: non-exercise, E: exercise) before administration of LPS (10μg/kg)/D-GalN (800mg/kg, intraperitoneally). Proteins were measured by Western Blotting analysis. β-actin was used as an internal
(B) Bars represent the protein quantification in cleaved- PARP/β-actin and mean values ± SE (n=3 in each group). The differences among those treatments with different letters were statistically significant (p < 0.05). ✽ means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in NE group. + means CS20+LPS and CS40+LPS group compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in E group. & means the NE group compared with E group. ^ means means CS20+LPS and CS40+LPS group compared with Saline+LPS group in E group.
Fig 16. Effects of *C. sinensis* and exercise training on terminal deoxynucleotidyl transferase (TdT)-mediated dUTP nick end labeling (TUNEL) staining in LPS/ D-GalN-treated mice.

(A) Effects of *C. sinensis* and precondition exercise training on TUNEL staining in normal and LPS/ D-GalN-treated mice. Mice were received *C. sinensis* (20 or 40 mg/kg/day) 4 hours before exercise and treated with precondition exercise training (NE: non-exercise, E: exercise) before...
administration of LPS (10µg/kg)/ D-GaIN (800mg/kg, i.p.). (B) Bars represent the TUNEL positive cells and mean values ± SE (n=3 in each group). The differences among those treatments with different letters were statistically significant (p < 0.05). ✽ means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in NE group. + means CS20+LPS and CS40+LPS group compared with Saline+LPS group in NE group. # means Saline+LPS, CS20+LPS and CS40+LPS group compared with control group in E group. & means the NE group compared with E group. ^ means CS20+LPS and CS40+LPS group compared with Saline+LPS group in E group.