Glycyrrhizin alleviates non-alcoholic steatohepatitis via modulating bile acids and meta-inflammation

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Abbreviations:

Acat1, acetyl-CoA acetyltransferase 1; ACTD, actinomycin D; ALT, Alanine aminotransferase; AST, aspartate aminotransferase; ASC, apoptosis-associated speck like protein containing a caspase recruitment domain; Bsep, bile salt export pump; CA, cholic acid; CASP1, caspase 1;
Cola, collagen; DAMPs; damage-associated molecular patterns; DCA, deoxycholic acid; FACS, fluorescence-activated cell sorting; Fatp, fatty acid transport protein; FXR, farnesoid X receptor; Gapdh, glyceraldehyde 3-phosphate dehydrogenase; G-CA, glycocholic acid; GA 30, GA 30 mg/kg; GA, glycyrhrhetic acid; GL, glycyrhrizin; GL50, glycyrhrizin 50 mg/kg; HDCA, hyodeoxycholic acid; HDL, high-density lipoprotein; H&E, hematoxylin and eosin; Hmgel, 3-hydroxy-3-methylglutaryl-CoA lyase; Hmgcs, hydroxymethylglutaryl-CoA synthase; Il1b, interleukin 1b; Il6, interleukin 6; PAMPs, pathogen-associated molecular patterns; Ppara, peroxisome proliferator-activated receptor alpha; qPCR, quantitative polymerase chain reaction; SHP, small heterodimer partner; LCA, lithocholic acid; LDL, low-density lipoprotein; LPS, lipopolysaccharide; LRH, liver receptor homolog; LXR, liver X receptor; MCA, muricholic acid; MCD, methionine-choline-deficient; MCS, methionine-choline-sufficient; Mmp, matrix metalloproteinase; Myd88, myeloid differentiation primary response 88; NAFLD, non-alcoholic fatty liver disease; NAS, non-alcoholic steatosis; NASH, non-alcoholic steatohepatitis; NLRP3, NLR family, pyrin domain-containing 3; Scd1, Stearoyl-CoA desaturase 1; Srebp1, sterol regulatory element-binding transcription factor 1; TC, total cholesterol; TCM, traditional Chinese medicine; T-CA, taurocholic acid; T-CDCA, taurochenodeoxycholic acid; T-DCA, taurodeoxycholic acid; TG, triglyceride; Tgfb1, transforming growth factor beta-1; T-HDCA, taurohyodeoxycholic acid; Timp, tissue inhibitor of metalloproteinases; TLR, toll-like receptor; T-β-MCA, tauro-β-muricholic acid; TNF, tumor necrosis factor; T-UDCA, tauroursodeoxycholic acid; TUNEL, terminal deoxynucleotidyl transferase-mediated dUTP nick end labeling; UDCA, ursodeoxycholic acid.
Non-alcoholic steatohepatitis (NASH) is the progressive stage of non-alcoholic fatty liver disease that may ultimately lead to cirrhosis and liver cancer, and there are few therapeutic options for its treatment. Glycyrrhizin (GL), extracted from the traditional Chinese medicine (TCM) liquorice, has potent hepatoprotective effects in both preclinical animal models and in humans. However, little is known about its effects and mechanisms in treating NASH. To explore the effects of GL on NASH, GL or its active metabolite glycyrrhetinic acid (GA), was administered to mice treated with a methionine-choline-deficient (MCD) diet-induced NASH model, and histological and biochemical analyses used to measure the degree of lipid disruption, liver inflammation and fibrosis. GL significantly improved MCD diet-induced hepatic steatosis, inflammation and fibrosis, and inhibited activation of the NLR family pyrin domain-containing 3 (NLRP3) inflammasome. GL significantly attenuated serum bile acids accumulation in MCD diet-fed mice partially by restoring inflammation-mediated hepatic farnesoid X receptor (FXR) inhibition. In Raw 264.7 macrophage cells, both GL and GA inhibited deoxycholic acid-induced NLRP3 inflammasome-associated inflammation. Notably, both intraperitoneal injection of GL’s active metabolite GA, and oral administration of GL, prevented NASH in mice, indicating that GL may attenuate NASH via its active metabolite GA. These results reveal that GL, via restoring bile acids homeostasis and inhibiting inflammatory injury, can be a therapeutic option for treatment of NASH.
Introduction

Non-alcoholic fatty liver disease (NAFLD) affects 25% of the global adult population, and has become a leading cause of chronic liver disease in the Western countries and increasingly affects the Asian population (Younossi et al., 2016). Early stage NAFLD is asymptomatic, however, when NAFLD progresses to non-alcoholic steatohepatitis (NASH), patients have a high risk of adverse events including fibrosis, cirrhosis, and NASH-driven hepatocellular carcinoma (Wong et al., 2014; Wong et al., 2016; Zoller and Tilg, 2016). Traditional Chinese medicine (TCM) has been investigated for treating NASH (Jadeja et al., 2014; Zhang and Schuppan, 2014). Glycyrrhizin (GL) is a potent hepatoprotective constituent extracted from TCM liquorice as revealed by various animal experimental liver injury models (Li et al., 2014; Yan et al., 2016), and thus is clinically used for treating chronic liver diseases in some Asian countries (Li et al., 2014). Clinical prescription preference for GL is found mainly for treating chronic liver diseases especially for viral hepatitis in China and Japan (Li et al., 2014), while the preclinical studies of GL in treating liver diseases mostly focused on viral hepatitis and drug/toxins-induced hepatotoxicity (Li et al., 2014; Yan et al., 2016; Zhou et al., 2016). However, the effect of GL in preventing NASH is largely unknown. Until now, only glycyrrhetinic acid (GA), the metabolite of GL in vivo, was found to prevent high-fat diet-induced NAFLD in rats in vivo and free fatty acids-induced toxicity in HepG2 cells in vitro (Wu et al., 2008). Whether and how GL and GA affect NASH, the progressive stage of NAFLD, is still unknown.
Bile acids homeostasis, which is mainly modulated by FXR, is disrupted in experimental NASH mouse models and clinical human NASH patients (Arab et al., 2017). Dysregulated bile acids disruption correlates with Non-alcoholic steatosis (NAS) score in obese NAFLD (Bechmann et al., 2013), causes cholestatic liver injury (Schoemaker et al., 2003), and promotes liver carcinogenesis in NASH-driven hepatocellular carcinoma model in mice (Xie et al., 2016). Furthermore, a bile acid sequestrant was found to prevent NAFLD and NASH (Arab et al., 2017), suggesting that to suppress the accumulation of bile acids during NASH could benefit NASH treatment. On the other hand, the NLRP3 inflammasome is known to be required for hepatocyte pyroptosis, liver inflammation, and fibrosis development in NAFLD (Wree et al., 2014a; Wree et al., 2014b). Metabolic homeostasis is associated with the immune system (Osborn and Olefsky, 2012), among which several bile acids were recently demonstrated to directly activate the NLRP3 inflammasome (Hao et al., 2017), providing a link between immune dysfunction with metabolic disorder. Since bile acids accumulate in NASH livers, it is reasonable to infer that toxic bile acids induced NLRP3 inflammasome activation and could be a key factor in mediating NASH progression. Bile acids-NLRP3 inflammasome axis is a potential pharmacological target for treating NASH.

GL was demonstrated to inhibit lithocholic acid (LCA)-induced cholestatic liver injury (Wang et al., 2012), bile acid-induced cytotoxicity in rat hepatocytes (Gumpricht et al., 2005) as well as alpha-naphthyl isothiocyanate-induced liver injury and bile acids disruption (Wang et al., 2017), indicating a potential role for GL in treating cholestatic liver injury. Additionally,
glycyrrhizin is also an inhibitor of NLRP3 inflammasome activation in macrophage in vitro and NLRP3 inflammasome-associated adipose tissue inflammation in mice (Honda et al., 2014). Thus, GL might target bile acids-NLRP3 inflammasome to elicit its hepatoprotective function for decreasing NASH.

In this study, the methionine-choline deficient (MCD) diet-induced NASH mouse model was used to examine the hepatoprotective effect and potential mechanisms of GL. GL was found to potently inhibit MCD diet-induced liver lipids accumulation, inflammation, and fibrosis. GL dampened the activation of NLPR3 inflammasome and restored bile acids homeostasis. Moreover, GL also directly inhibited deoxycholic acid-induced NLRP3 inflammasome activation. GL gavage as well as GA significantly prevented MCD diet-induced liver injury, suggesting GL by oral intake might be applied in the clinic as a therapeutic option for NASH patients.
Methods and Materials

Chemicals and Reagents

GL (97%) was purchased from TCI (Shanghai, China). 18β-GA (97%), actinomycin D (ACTD), Tween 80 and lipopolysaccharides (LPS) from Escherichia coli O111:B4 were purchased from Sigma (USA). Recombinant human TNF-α was purchased from R&D (Cat. No. 210-TA, MN, USA). MCD diet and MCS diet were purchased from Trophic Animal Feed High-Tech Co., Ltd (Nantong, China) or Dyets Inc (Bethlehem, PA, USA). The In situ cell death detection kit was purchased from Roche (USA).

Experimental Animals and Treatments

6- to 8-week-old male C57BL/6 mice were obtained from Academy of Military Medical Sciences, China or the National Institutes of Health. Mice were acclimatized to the facilities for a week and then randomly divided into different groups. The animal room was maintained at 25 ± 2 °C with a 12-h light-dark cycle and 50 ± 10 % humidity. All animal studies were approved by the Animal Ethics Committee of China Pharmaceutical University and carried out in accordance with the Guide for the Care and Use of Laboratory Animals, National Institutes of Health.

To assess the influence of GL in treating NASH, mice were randomly divided into 4 groups with different treatments for 8 weeks: MCS group (saline, I.P.), MCS+GL50 group (50 mg/kg/day GL, I.P.), MCD group (saline, I.P.), and MCD+GL50 group (50 mg/kg/day GL, I.P.).
All animals were fed MCD or MCS diet for 6 weeks, and then injected with 50 mg/kg/day of GL for two consecutive weeks, while mice were fed matched MCD diet or MCS diet for the last two weeks until the end of experiments. All mice were fasted for 4 hours with free access to water before killing. For testing the effects of GA injection in preventing NASH, mice were divided into 3 groups with different treatments for 6 weeks: MCS group (MCS diet feeding, saline, I.P.), MCD group (MCD diet feeding, 5% Tween 80/saline, I.P.), and MCD+GA30 group (MCD diet feeding, 30 mg/kg/day of GA, dissolved in 5% Tween 80/saline, I.P.). Similarly, in the study for testing the effect of GL gavage, 3 groups of mice were treated accordingly: MCS group (MCS diet feeding, saline, I.P.), MCD group (MCD diet feeding, saline, P.O.), and MCD+GL50 group (MCD diet feeding, 50 mg/kg/day of GL, P.O.). All mice were fed MCD or MCS diet for 6 weeks, and injected with control vehicle or 50 mg/kg/day of GL gavage for the last 3 consecutive weeks, respectively. All mice were killed with CO₂ asphyxiation, and blood and livers were collected and stored for further use. GL was dissolved in saline and pH value adjusted to 7.0-7.5 with 1M sodium hydroxide (NaOH) solution (Sigma). GA was freshly dissolved in saline containing 5% Tween 80.

**Histology Analysis**

Formalin-fixed liver tissues were embedded in paraffin and 5 μm thick sections were cut for H&E staining, TUNEL staining, and Masson’s Trichrome Staining based on respective manual protocol. A portion of fresh livers were embedded in Tissue-Tek Optimal Cutting Temperature (O.C.T.) compound (Sakura Finetek, Torrance, USA) and then flash frozen for
Oil Red O staining. Further samples processing and analysis for Figure 1-3 was performed in the department of Pathophysiology in Affiliated Hospital of Nanjing University of Chinese Medicine (Nanjing, Jiangsu, China) and Figure 7 at Histoserv Inc. (Germantown, MD, USA).

**Targeted Bile Acids Metabolome Analysis**

The levels of main bile acid species were determined by high-performance liquid chromatography-quadrupole time-of-flight tandem mass spectrometry (UPLC-triple/TOF-MS) as described previously (Hao et al., 2017).

**In Vitro Cell Culture and Treatment**

Human HepG2 cells were obtained from the American Type Culture Collection (Manassas, VA, USA). For testing the protective effects of GL or GA on ACTD/TNF-α-induced HepG2 cell death, HepG2 cells were seeded in 96-well plates for cell viability assays or 6-well plates for FACS assays and then grown to 70%-90% confluence before use. HepG2 cells were pretreated with control DMSO, GL or GA together with 0.3 μM of ACTD for 30 min, and then treated with recombinant human TNF-α for additional 13 h or 24 h for cell viability assays with the Cell Counting Kit-8 (CCK-8, Dojindo Laboratories, Kumamoto, Japan). For FACS assays, samples were collected at 13 h after treatment and analyzed according to the manufacturer’s instruction for the Annexin V FITC apoptosis detection kit (BD Biosciences, San Diego, CA).

**Reporter Luciferase Assay**
The PGL4-Shp-TK firefly luciferase construct and human FXR expression plasmid were provided by Grace L. Guo (Rutgers University). A *Renilla* luciferase reporter gene (pRL-luciferase; Promega; Madison, WI) was used as a transfection efficiency control. The plasmids were transfected into HepG2 cells that were cultured in 10% Roswell Park Memorial Institute 1640 (RPMI 1640) medium and seeded in 24-well plates at a confluency of 70%-90%. Cells were transfected with plasmids using lipofectamine 3000 reagent (Invitrogen, Thermo Fisher Scientific, MA, USA) for 24 h, and then treated with LPS, GL or GA for another 24 h. The cells were lysed and luciferase activities measured with the Dual Luciferase Reporter Assay kit (Promega Corp., Madison, WI) and a Tecan Genios Pro luminescence plate reader (Research Triangle Park, NC).

**Serum Biochemical Analysis**

Biochemical parameters including ALT levels, AST levels, total cholesterol (TC) levels, triglyceride (TG) levels, low-density lipoprotein cholesterol (LDL-cholesterol) and high-density lipoprotein cholesterol (HDL-cholesterol) in serum, TC levels and TG levels in liver were quantified using a standard clinical automatic analyzer at Zhongda Hospital Affiliated to Southeastern University.

**Western Blot Analysis**

Liver samples were lysed in RIPA Lysis Buffer (Millipore, Temecula, California, USA) with protease and phosphatase inhibitors (Halt Protease and Phosphatase Inhibitor Cocktail, Cat #78446, Thermo Scientific, Rockford, USA), and then the protein extracts were separated
by sodium dodecyl sulfate-polyacrylamide gel electrophoresis and transferred to a polyvinylidene difluoride membrane. The membrane was incubated overnight at 4 ℃ with antibodies against mouse β-ACTIN (Cell Signaling, Danvers, MA, Cat#4970), mouse monoclonal anti-caspase 1 (Santa Cruz, Cat#sc-56036), and rabbit polyclonal anti-IL-1β (Santa Cruz, Cat#sc-7884).

**Elisa of Serum IL-1β Level in Mice**

An IL-1β ELISA kit was purchased from R&D systems (Cat#, MLB00C, MN, USA). Fifty μL of serum was used for analysis based on the manual protocol accordingly.

**Quantitative Polymerase Chain Reaction**

Total tissue RNA extraction was performed by using the RNAiso Plus reagent (Takara, Bio-Tech Co., Ltd, Dalian, China). Purified total RNA was reverse-transcribed using the Prime Script RT Reagent Kit (Takara Biotechnology Co., Ltd.). Quantitative polymerase chain reaction (qPCR) was performed by using the ABI PRISM 7000 Sequence Detection System (Applied Biosystems, Bedford, MA) and Sybr Green reagent kit (Bio-Rad Co., Ltd., Shanghai, China). Values were normalized to Gapdh. Sequences for qPCR primers could be provided upon request.

**Statistical analysis**

Data are presented as means ± SD. Statistic differences between experimental groups were determined by two-tailed Student’s t test or one-way ANOVA followed by Dunnett’s multiple
comparisons test in GraphPad Prism 7.0 as stated (GraphPad Software Inc., San Diego, CA). P value less than 0.05 was considered statistically significant.
Results

GL attenuates NASH-induced liver injury

To determine whether GL attenuates MCD diet-induced liver injury in vivo, C57BL/6 mice were treated with GL after MCD diet feeding for 6 weeks, and the GL treatment continued for the last two weeks while the mice were fed matched MCD diet or MCS diet during the injection (Figure 1A). GL treatment did not significantly affect food intake and body weight (Supplemental Figure 1, A and B) as well as liver index (Supplemental Figure 1C) both in MCS diet-fed mice and MCD diet-fed mice. H&E staining data showed that 50 mg/kg of GL sharply decreased hepatic steatosis in MCD diet-fed mice, while 50 mg/kg of GL treatment alone did not cause liver toxicity (Figure 1B). Apoptotic hepatocyte death is known to be involved in NASH (Feldstein et al., 2003), and therefore liver tissues were analyzed by TUNEL staining to measure apoptosis. GL treatment significantly reduced the positive staining when compared with control saline treatment in MCD diet-fed mice (Figure 1C). Consistently, the apoptosis marker gene mRNA, Bcl2a1c, was also significantly inhibited by GL treatment in MCD diet-fed mice (Supplemental Figure 1D). Since TNF-α is an inducer of apoptosis (Wang et al., 1996), and ACTD/TNF-α is frequently used to mimic apoptotic cell death (Leist et al., 1994), the effects of GL in alleviating ACTD/TNF-α-induced apoptosis were examined in HepG2 cells. GL directly inhibited ACTD/TNF-α-induced decrease of cell viability at both 13 h and 24 h after ACTD/TNF-α treatment (Supplemental Figure 1, E and F), and ameliorated the increase in apoptotic cell death as assessed by FACS assay (Supplemental Figure 1G), supporting that
GL could directly inhibit TNF-α-mediated apoptosis. In addition, serum ALT levels were increased in MCD diet-fed group, while they were markedly decreased in the MCD+GL50-treated group (Figure 1D). NAFLD scores among the MCS, MCD and MCD+GL50 groups by a double-blinded analysis, also showed that the MCD diet induced a significant increase of NAFLD activity, steatosis, and inflammation in livers of the MCD group, which were all markedly alleviated by GL treatment (Figure 1, E, F and G). Taken together, these results suggest that GL rescues apoptotic liver injury induced by MCD diet feeding, at least partially through directly inhibiting TNFα-induced hepatocyte apoptosis.

GL improves NASH-related liver fibrosis

Liver fibrosis is a hallmark of NAFLD progression to NASH. Masson Trichrome staining of collagen was used to measure liver fibrosis, and GL was found to significantly ameliorate NASH-associated collagen deposition induced by MCD diet feeding (Figure 2A). α-SMA is also another sensitive marker of liver fibrosis. MCD diet-induced upregulation of α-SMA expression was significantly decreased by GL treatment both at the protein level as determined by immunohistochemical staining (Figure 2B), and by mRNA analysis (Figure 2C). GL also significantly dampened the expression of fibrogenetic gene mRNAs including Tgfb1, Timp1, Timp2, Cola1, Cola2, Mmp2 and Mmp9 (Figure 2, D-J). These data demonstrate that GL significantly improves NASH-related liver fibrogenesis.

GL restores NASH-related hepatic lipids accumulation
In NASH, lipotoxicity caused by accumulating lipids in liver is a major trigger of liver toxicity (Fuchs and Sanyal, 2012). Therefore, effect of GL in MCD diet-induced lipid disruption was determined. When compared with control vehicle-treated MCD group, intraperitoneal injection of GL at 50 mg/kg significantly attenuated MCD diet-induced lipid accumulation in liver as revealed by Oil Red O staining (Figure 3A). When compared with the MCS group, liver TG and TC levels were both significantly increased in the MCD group, and both were significantly decreased by GL treatment (Figure 3, B and C). In contrast, serum TG and TC levels were significantly decreased after MCD diet feeding, which were also both normalized by GL treatment (Figure 3, D and E). Similarly, GL treatment also normalized MCD diet-induced disruption of serum HDL-cholesterol and LDL-cholesterol levels (Figure 3, F and G). These data strongly suggest that GL treatment could improve MCD diet-induced lipids disruption both in liver and serum.

Systematic lipids homeostasis is maintained by balanced mutual regulation of de novo lipids synthesis and lipids degradation by oxidation and subsequent ketogenesis (Tessari et al., 2009). To examine how GL restores MCD diet-induced dyslipidemia, lipid regulatory pathways were analyzed. When compared with MCS diet-fed mice, MCD diet-feeding significantly increased expression of Lxr and Srebp1 mRNAs and the downstream lipids synthesis gene mRNA, Fas. Although GL treatment decreased the induction of Lxr, Srebp1a and Srebp1c mRNA, it showed no significant effect on the mRNA level of Fas and significantly restored MCD diet-decreased Scd1 mRNA, both of which are encoded by two key lipogenic genes.
Moreover, GL treatment significantly inhibited MCD diet-induced mRNA upregulation of the fatty acid transport gene mRNAs Fatp3 and Fatp4, when compared with vehicle-treated MCD-fed mice (Figure 4H). MCD diet-feeding did not significantly change the expression of the ketogenesis pathway when compared with the MCS diet-fed group. In MCD diet-fed mice, GL treatment even significantly decreased Acat1, Hmgcs, Hmgcl, Ppara and Cyp4a14 mRNA levels, and showed no significant effect in other mRNAs expression involved in ketogenesis and lipids oxidation, when compared with control vehicle treatment (Figure 3, I and J). These data indicate that GL may indirectly reduce lipogenesis by suppressing the expression of fatty acid transporters and not through enhancing lipid degradation via ketogenesis and oxidation.

**GL decreases MCD-induced bile acids accumulation**

Accompanied with the dysregulated lipids metabolism, bile acids are commonly accumulated in NASH both in mice and clinical patients (Bechmann et al., 2013; Gong et al., 2016; Arab et al., 2017). Several species of bile acids may exert direct toxic effects against hepatocytes and serve as danger molecules in triggering NLRP3 inflamasome activation (Gong et al., 2016; Hao et al., 2017). Thus, the effect of GL treatment on serum bile acids was examined under conditions of experimental NASH. MCD diet feeding significantly induced accumulation of free bile acids including α-MCA, β-MCA, CA, DCA, UDCA, and HDCA, by 2.9-, 12.1-, 18.0-, 3.7-, 7.6-, and 7.2-fold, respectively, all of which were markedly reduced by GL treatment (Figure 4A). Similarly, GL treatment significantly restored MCD diet-induced
serum accumulation of taurine-conjugated bile acids, T-β-MCA, T-CA, T-DCA, T-CDCA, T-UDCA and the glycine conjugated bile acids G-CA (Figure 4B). FXR activates the expression of small heterodimer partner 1 (SHP-1) that inhibits the liver receptor homolog 1(LRH-1) and represses expression of CYP7A1, which catalyzes the rate-limiting step in bile acid biosynthesis (Goodwin et al., 2000). Since the present study showed that all tested bile acids including the main primary bile acid, CA, were decreased by GL, GL treatment was inferred to influence the de novo synthesis of bile acids in liver. Therefore, FXR-SHP-CYP7A1 signaling, the predominant pathway for bile acids synthesis, was analyzed. In MCD diet-fed mice, hepatic FXR signaling was inhibited, and Cyp7a1 mRNA was upregulated, when compared with MCS diet-fed mice. As expected, GL treatment significantly activated FXR, evidenced by upregulation of Shp mRNA and the downstream suppression of Cyp7a1 mRNA, while Fxr mRNA itself was also significantly increased by GL treatment when compared with the vehicle-treated group in the mouse NASH model (Figure 4, C, D and E). In contrast, the bile acid export pump, Bsep mRNA was not significantly influenced by GL treatment (Figure 4F), suggesting a role of GL in inhibiting de novo bile acids synthesis.

To explain how GL could activate FXR in MCD diet-fed livers, the possibility that GL could act as a FXR ligand was explored. In mouse primary hepatocytes and HepG2 cells, no significant effect of FXR activation were found by neither GL nor GA treatment (Supplemental Figure 2, A and B) as well as GL-treated normal mice in vivo (Supplemental Figure 2C). Consistently, both GL and GA failed to activate FXR, while GW4064 induced FXR reporter
luciferase by 20-fold as a positive control in HepG2 cells (Supplemental Figure 2D), excluding GL as a direct FXR activator. These data prompted us to ask whether GL activated FXR only under NASH-related pathological conditions. Since NF-κB activation, a common pathological character under NASH condition, is known to lead to FXR inactivation (Wang et al., 2008), rescue of FXR inactivation induced by NF-κB by GL was examined. LPS treatment slightly but significantly repressed FXR reporter activity, and both GL and GA treatment restored LPS-induced FXR inactivation (Supplemental Figure 2E). These data suggest that GL may restore bile acid homeostasis via attenuating the loss of FXR signaling induced by inflammatory damage.

**GL inhibits NLRP3 inflammasome activation and meta-inflammation in mice**

NLRP3 inflammasome activation is required for fibrosis development in NAFLD (Wree et al., 2014a; Wree et al., 2014b; Mridha et al., 2017). Activation of the NLRP3/inflammasome is a two-step process requiring a priming step (signal 1) induced by binding of damage-associated molecular patterns (DAMPs) or pathogen-associated molecular patterns (PAMPs) to receptors such as Toll-like receptor (TLR) to increase the transcription of NLRP3 and pro-IL-1β, followed by an activation step (signal 2) induced by the recruitment of ASC to activate the NLRP3 inflammasome to convert pro-caspase-1 into active caspase 1, which in turn processes pro-IL-1β into mature IL-1β (Guo et al., 2015). Since GL is a known inhibitor of the NLRP3 inflammasome (Honda et al., 2014), whether GL inhibited activation of the NLRP3 inflammasome in the NASH model was tested. When compared with MCS diet-fed mice, MCD
diet feeding induced a significant 2- to 4-fold increase of upstream Tlr4, Tlr9, and Myd88 mRNAs, all of which were normalized by GL treatment (Figure 5, A, B and C). Similarly, MCD diet-induced Casp1, Nlrp3, and Asc mRNAs, by 5.5-, 4.8- and 7.7-fold, respectively, which were significantly reduced by GL treatment (Figure 5, D, E and F). These data suggest that GL could dampen the signal 1 activation of NLRP3/inflammasome in MCD diet-fed mice. Western blot analysis was performed to test the protein levels of pro-CASP1, active CASP-1, pro-IL-1β and active IL-1β. MCD diet induced upregulation of signal 1 and signal 2 when compared with MCS diet treatment. GL treatment decreased the protein expression of pro-CASP1 and pro-IL1β (Figure 5G), confirming GL’s effect in inhibiting signal 1 of NLRP3 inflammasome activation. Furthermore, GL treatment was also found to inhibit the MCD diet-induced upregulation of active caspase1 (CASP1 p20) and cleaved IL-1β (Figure 5G). Additionally, GL significantly inhibited the upregulation of downstream proinflammatory cytokine mRNAs Tnfa, Il6, and Il1b (Figure 5, H, I and J) and serum release of IL-1β protein (Figure 5K), demonstrating a significantly improvement of NASH-related meta-inflammation by GL. These data demonstrate that GL treatment could dampen MCD diet-induced TLR/NLRP3 inflammasome activation and the related meta-inflammation.

Since previous reports and the present results indicate that bile acids accumulate in NASH and may be the main trigger for activation of the NLRP3 inflammasome (Arab et al., 2017; Hao et al., 2017), the possibility was explored whether GL could directly inhibit bile acid-induced NLRP3 inflammasome activation. Deoxycholic acid (DCA) was shown to induce NLRP3
inflammasome activation (Hao et al., 2017) and was thus used as a model to test GL’s effect in inhibiting bile acid-induced NLRP3 inflammasome activation. In Raw 264.7 cells, both GL and GA significantly decreased DCA-induced upregulation of Tnfa, Nlrp3, and Il1b mRNAs (Figure 6). These data demonstrate that GL as well as its metabolite GA could directly suppress DCA-induced NLRP3 inflammasome activation in vitro, which at least partially explains GL’s role in dampening MCD diet-induced NLRP3 inflammasome activation in vivo.

**GA, the active metabolite of GL, improves MCD diet-induced liver injury**

Structurally, GL is a glycoside that hydrolyzed into its metabolite GA by intestinal bacteria after oral administration (Akao et al., 1994; Takeda et al., 1996). To test whether GA influences NASH, mice were treated with the MCD diet for 6 weeks, and then either intraperitoneally injected with GA (30 mg/kg) or intragastrically administered with GL by gavage (50 mg/kg) for an additional 3 weeks (Figure 7A). GA as well as GL gavage did not significantly change the body weight, liver weight, and liver index (Supplemental Figure 3). GA was found to significantly decrease the MCD diet-induced increase of ALT and AST levels (Figure 7, B and C). Similarly, GL gavage at a dose of 50 mg/kg also significantly ameliorated MCD diet-induced increase of serum ALT levels and AST levels (Figure 7, D and E). By histological analysis, 30 mg/kg of GA ameliorated MCD diet-induced hepatic steatosis (Figure 7, F and G). Similarly, GL gavage at a dose of 50 mg/kg improved the liver lipid overloading (Figure 7, H and I). These results indicate that GA may be required for the hepatoprotective effects of GL for the treatment of NASH.
Discussion

GL and its active metabolite GA have been demonstrated to be hepatoprotective, however, they have not been examined for their activities toward NASH. The current study shows that GL, via its activate metabolite GA, could decrease MCD diet-induced NASH, as revealed by improved hepatic steatosis, inflammation and fibrosis. Mechanistically, GL may restore NASH-induced dysregulation of bile acids and lipids and the resultant meta-inflammation.

Many factors such as cholesterol crystal (Duewell et al., 2010), ceramides (Chaurasia and Summers, 2015), and secondary bile acids (Hao et al., 2017) are known to accumulate in NASH livers and could act as DAMPS to induced NLRP3 inflammasome activation. Activation of the NLRP3 inflammasome results in the release of IL-1β, together with other pro-inflammatory cytokines such as TNFα, that trigger an inflammatory cascade that ultimately promotes pathological development of NASH. Thus, accumulated DAMPs, via activating NLRP3 inflammasome and other inflammatory pathways, may represent core pathological triggers in NASH development. In the present study using the MCD diet-induced NASH model, GL was shown to inhibit NLRP3 inflammasome activation accompanied by restoring abnormal serum bile acids accumulation, and GL could normalize the dysregulated lipids and bile acids and thereby reduce meta-inflammation. Since several bile acids are known to activate NLRP3/inflammasome signaling, GL may inhibit the NLRP3 inflammasome via the regulation of FXR signaling. However, GL did not activate FXR in normal mice in vivo, hepatocytes in vitro and failed to induce FXR reporter luciferase activity, indicating that GL is not a direct...
FXR activator. FXR is known to have a reciprocal regulation with NF-κB activation (Wang et al., 2008). Under excessive inflammation, FXR signaling is reduced, resulting in the accumulation of bile acids and dysregulated lipid catabolism, which reciprocally triggers further inflammation (Bechmann et al., 2013). Although the exact mechanism warrants further clarification, the current study strongly indicates that GL and GA may target FXR mediated meta-inflammation in the treatment of NASH. The present data agree with a previous report that GL induces SHP expression and inhibits Cyp7a1 expression in LCA-induced cholestatic liver injury model (Wang et al., 2012). Additionally, because the accumulation of bile acids may directly damage hepatocytes (Gumpricht et al., 2005), the observation that GL can restore bile acids homeostasis suggests that GL may be particularly suitable for the therapy of NASH featured with cholestasis.

In addition to restoration of the lipid and bile acid homeostasis, GL could also directly inhibit DCA-induced NLRP3 inflammasome activation, suggesting that it may target multiple nodes in treating NASH. In line with the present findings, GL was previously shown to inhibit NLRP3 inflammasome activation induced by diverse DAMPs or PAMPs in bone marrow-derived macrophages (Honda et al., 2014). MCC950, a NLRP3 selective inhibitor, was recently demonstrated to improve NAFLD pathology and fibrosis (Mridha et al., 2017), suggesting that the NLRP3 inflammasome might be a potential target for NASH therapy. Moreover, it was found that MCD diet-induced steatosis sensitized cholestatic liver injury and dysregulated bile acids synthesis and transport (Lionarons et al., 2016). The present results together with these
previous findings strongly indicate that dysregulated metabolism of lipids and bile acids and the resultant meta-inflammation is a pivotal event in NASH development.

GL by the oral route shows extremely low bioavailability (approximately 1%) and most of GL by oral intake is hydrolyzed into GA in the gut by intestinal bacteria, which is then absorbed and elicits its pharmacological effects (Nose et al., 1994; Yamamura et al., 1995; Takeda et al., 1996). In line with this point, GL gavage as well as direct GA i.p. injection could both combat experimental NASH, suggesting that effect of GL against MCD diet-induced NASH is largely through its primary metabolite GA. In contrast, GL but not GA, contributes to hepatoprotective effect of GL in ameliorating acetaminophen-induced liver injury (Yan et al., 2016). Thus, it is important to note that although both GL and GA have efficacy in various types of hepatic injury, the exact effects and thereby precise clinical applications of these agents need to be carefully considered.

In summary, the herbal medicine-derived compound GL may be a therapeutic option for NASH treatment, particularly that featured with cholestasis. Mechanistically, GL and/or GA may target bile acid-mediated meta-inflammation and block the reciprocal FXR-NLRP3 inflammasome pathway. All the data obtained from present MCD diet-induced NASH model demonstrate that GL and GA may be potentially effective against NASH. However, it is important to note that the MCD-diet induced NASH model, despite the presented liver steatosis and inflammation, does not show obesity, insulin resistance nor mimic aberrant increase of serum lipids, all of which are common pathophysiological characters of human NASH.
(Hebbard and George, 2011). Thus, using long-term high-fat and/or high-sugar diet feeding model would help further confirm the translational potential of GL and GA in combating metabolic syndrome-related NASH.
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Authorship Contributions

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Wrote or contributed to the writing of the manuscript: Yan, Gonzalez, Hao.
References


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Footnotes

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Figure Legends

Figure 1. Glycyrrhizin significantly alleviates MCD diet-induced liver damage, improves TUNEL staining, and reduces serum transaminases.

(A), Mouse experiment procedure scheme. (B), H & E-staining of liver sections, original magnification: 20×; Scale bar, 50 μm. (C) TUNEL-staining analysis of paraffin-embedded livers, original magnification: 20×; Scale bar, 50 μm. (D), Serum ALT levels. (E, F and G), NAFLD scoring statistics for H&E slides. Data are presented as means ± SD. Statistic differences between experimental groups were determined by two-tailed Student’s t test. n=6-8 for each group. MCS, MCS diet-fed mice treated with saline; MCS+GL50, MCS diet-fed mice treated with 50 mg/kg of GL. MCD, MCD diet-fed mice treated with saline; MCD+GL50, MCD diet-fed mice treated with 50 mg/kg of GL. ###p<0.001 versus MCS group. **p<0.01 and ***p<0.001 versus MCD group.

Figure 2. Glycyrrhizin significantly dampens MCD-induced liver fibrogenesis.

(A and B), Masson’s Trichrome staining (A) and immunohistochemistry staining of α-SMA (B) for paraffin-embed livers, original magnification: 20X; scale bar, 50 μm. (C, D, E, and F), levels of αSma, Tgfb1, Timp1 and Timp2 mRNAs in livers. (F, G, H, I and J), Cola1, Cola2, Mmp2 and Mmp9 mRNAs in livers. Data are presented as means ± SD. Statistic differences between experimental groups were determined by two-tailed Student’s t test. n=6-8 in each group. MCS, MCS diet-fed mice treated with saline; MCD, MCD diet-fed mice treated with saline;
MCD+GL50, MCD diet-fed mice treated with 50 mg/kg of GL. ##p<0.01 and ###p<0.001 versus MCS group. *p<0.05, **p<0.01 and ***p<0.001 versus MCD group.

Figure 3. Glycyrrhizin significantly reduces MCD-induced lipids accumulation in liver via inhibiting fatty acid uptake. (A) Oil Red O-staining of liver sections, original magnification: 20×; scale bar, 50 μm. (B and C), Liver TG and TC levels. (D and E), Serum TG and TC levels. (F and G), Serum HDL-cholesterol and LDL-cholesterol levels. (H-J), Effect of GL in the mRNA expression involved in the pathway of lipogenesis (H), ketogenesis (I), and beta-oxidation (J). Data are presented as means ± SD. Statistic differences between experimental groups were determined by two-tailed Student’s t test. n=6-8 in each group. MCS, MCS diet-fed mice treated with saline; MCD, MCD diet-fed mice treated with saline; MCD+GL50, MCD diet-fed mice treated with 50 mg/kg of GL. #p<0.05; ##p<0.01 and ###p<0.001 versus MCS group. *p<0.05, **p<0.01 and ***p<0.001 versus MCD group.

Figure 4. Glycyrrhizin activates liver FXR/SHP/CYP7A1 signaling and rescues MCD-induced bile acids disruption. (A), Serum level of free bile acids, α-MCA, β-MCA, CA, DCA, UDCA and HDCA. (B), Serum level of conjugated bile acids, T-β-MCA, T-CA, T-DCA, T-CDCA, T-UDCA, T-HDCA and G-CA. (C, D, E and C), hepatic mRNA levels of Fxr, Shp, Cyp7a1 and Bsep. Data are presented as means ± SD. Statistic differences between experimental groups were determined by two-tailed Student’s t test. n=6-8 in each group. MCS, MCS diet-fed mice treated with saline; MCD, MCD diet-fed mice treated with saline;
MCD+GL50, MCD diet-fed mice treated with 50 mg/kg of GL. #p<0.05; ##p<0.01 and ###p<0.001 versus MCS group. *p<0.05, **p<0.01 and ***p<0.001 versus MCD group.

Figure 5. Glycyrrhizin significantly reduces MCD-induced TLR/NLRP3 inflammasome activation and the related meta-inflammation. (A, B and C), Tlr4, Tlr9 and Myd88 mRNA levels in livers. (D, E and F), mRNA levels of Nlrp3, Casp1 and Asc in livers. (G), Western blot analysis of pro-CASP1, cleaved CASP1, pro-IL1β, and cleaved IL1β in livers. (H, I and J), mRNA levels of proinflammatory cytokines, Tnfa, Il6, and Il1b in livers. (K), Elisa analysis of serum IL-1β level in mice. Data are presented as means ± SD. Statistic differences between experimental groups were determined by two-tailed Student’s t test. n=6-8 in each group. MCS, MCS diet-fed mice treated with saline; MCD, MCD diet-fed mice treated with saline; MCD+GL50 or MCD+GL, MCD diet-fed mice treated with 50 mg/kg of GL. #p<0.05; ##p<0.01 and ###p<0.001 versus MCS group. *p<0.05, **p<0.01 and ***p<0.001 versus MCD group.

Figure 6. Both GL and GA significantly inhibit DCA-induced Tnfa, Nlrp3, and Il1b mRNA. (A, B and C), Effect of GL in DCA-induced mRNA of Tnfa, Nlrp3, and Il1b mRNAs in Raw 264.7 cells. (D, E and F), Effect of GA in DCA-induced Tnfa, Nlrp3, and Il1b mRNAs in Raw 264.7 cells. Data are presented as means ± SD. Statistic differences determined by one-way ANOVA followed by Dunnett’s multiple comparisons test among multiple-group comparisons. n=3 per group. Raw 264.7 cells were pretreated with 0.1% DMSO, GL or GA for 30 minutes, and then treated with 200 μM of DCA for additional four hours to further perform
mRNA analysis. DMSO, 0.1% DMSO/vehicle-treated control vehicle group; DCA, 0.1% DMSO/DCA-treated group; DCA+GL50, 50 μM of GL/DCA-treated group; DCA+GL250, 250 μM of GL/DCA-treated group; DCA+GA3, 3 μM of GA/DCA-treated group; DCA+GA6, 6 μM of GA/DCA-treated group. # p<0.05; ##p<0.01 and ###p<0.001 versus DMSO group. *p<0.05, **p<0.01 and ***p<0.001 versus DCA group.

Figure 7. GL’s active metabolite, GA, significantly reduces MCD-induced liver injury. (A), Mouse experiment procedure scheme. (B and C), Serum ALT and AST levels in saline or GA-treated mice. (D and E), Serum ALT and AST levels in saline or GL gavage-treated mice. (F and G), Representative H&E staining (F) and Oil Red O staining (G) for livers from MCD group and MCD+GA30 group; (H and I), Representative H&E staining (H) and Oil Red O staining (I) for livers from MCD group and MCD+GL50 (po) group. Original magnification: 20×; Scale bar, 50 μm. Data are presented as means ± SD. Statistic differences between experimental groups were determined by two-tailed Student’s t test. n=5 in each group. MCS, MCS diet-fed mice treated with saline; MCD, MCD diet-fed mice treated with saline; MCD+GA30, MCD diet-fed mice treated with 30 mg/kg of GA by intraperitoneal injection; MCD+GL50 (p.o.), MCD diet-fed mice treated with 50 mg/kg of GL by gavage. #p<0.05; ##p<0.01 and ###p<0.001 versus MCS group. *p<0.05, **p<0.01 and ***p<0.001 versus MCD group.
Figures

Figure 1

A

Matched MCS or MCD Feeding Time (W, Week)

B
MCS
MCS+GL50

C
MCD
MCD+GL50

D

E

F

G

ALT (IU/L)

NAD Activity Score

Steatosis Score

Inflammation Score

MCS
MCD
MCD+GL50

MCS
MCD
MCD+GL50

MCS
MCD
MCD+GL50
Figure 3

A

MCS
MCD
MCD+GL50

B

Liver TG
Liver TC
Serum TG
Serum TC

C

D

E

F

G

H

I

J

Relative mRNA

Acath
Hmgcs
Hmads
Bdel

Lxr
Srebp1a
Srebp2
Fas
Scdl
Fatp1
Fatp2
Fatp3
Fatp4
Fatp5

Ppara
Cyp26a1
Cdd1
Acot1
Mkksa

# P < 0.05
** P < 0.01
*** P < 0.001
## P < 0.001
### P < 0.0001
Figure 4

A

B

C

D

E

F

Serum Bile Acids Level (μM)

Serum Bile Acids Level (μM)

Relative mRNA

Relative mRNA

Relative mRNA

Relative mRNA

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Figure 6
Figure 7

A

MCD Feeding Time (W, Week)

0 1 2 3 4 5 6

GA or GL

B

ALT (IU/L)

500

400

300

200

100

0

MCD

MCD+GA30

MCD+GL50 (P.O.)

MCD+GA30

MCD

MCD+GA30

MCD+GL50 (P.O.)

MCD

MCD+GA30

MCD+GL50 (P.O.)

D

ALT (IU/L)

600

500

400

300

200

100

0

MCD

MCD+GL50 (P.O.)

MCD

MCD+GL50 (P.O.)

F

MCD

MCD+GA30

F

MCD

MCD+GA30

MCD

MCD+GL50 (P.O.)

H

MCD

MCD+GA30

MCD

MCD+GL50 (P.O.)

I

MCD

MCD+GA30

MCD

MCD+GL50 (P.O.)

E

AST (IU/L)

500

400

300

200

100

0

MCD

MCD+GA30

MCD+GL50 (P.O.)

MCD+GA30

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MCD+GL50 (P.O.)

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MCD+GL50 (P.O.)

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