Alcoholic fatty liver disease inhibited the co-expression of Fmo5 and PPARα to activate the NF-κB signaling pathway, thereby reducing liver injury via inducing gut microbiota disturbance

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Abstract

Background
Alcohol-induced intestinal dysbiosis disrupts and inflammatory response are essential in the development of alcoholic fatty liver disease (AFLD). Here, we investigate the effects of Fmo5 on changes in enteric microbiome composition in a model of AFLD and dissect the pathogenic role of Fmo5 in AFLD-induced liver pathology.

Materials and methods
The expression profile data of GSE8006 and GSE40334 datasets were downloaded from the GEO database. Weighted gene co-expression network analysis (WGCNA) approach allowed us to investigate the AFLD-correlated module. DEGs were used to perform Gene Ontology functional and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analyses. Four PPI networks were constructed using the STRING database and visualized by Cytoscape software. Cytohubba plug-in was used to identify the hub genes. Western blot and immunohistochemistry assays were used to detect protein expressions. ELISA assay was used to detect the level of serum inflammatory cytokines. Lipid droplets in cytoplasm were observed using Oil Red O staining. Cell apoptosis was detected using TUNEL assay and flow cytometry analysis. ROS level were detected using flow cytometry analysis. The nuclear translocation of NF-κB p65 was observed using immunofluorescence staining. Co-immunoprecipitation were used to detect the co-expression of PPARα and Fmo5 in L02 cells. 16S rDNA sequencing defined the bacterial communities in mice with AFLD.

Results
Fmo5 was a key DEG and closely associated with gut microbiota and PPAR signaling pathway. Gut microbiome function in AFLD was significantly related to PPAR signaling pathway. AFLD induced shifts in various bacterial phyla in the cecum, including a reduction in Bacteroidetes and increased Firmicutes. Fmo5 and PPARα co-expression in cell and animal model with AFLD, which decreased significantly. Silencing of Fmo5 and PPARα aggravated the functions of AFLD inducing apoptosis and inflammatory response, promoting liver injury, and activating the NF-κB signaling pathway in vivo and in vitro. NF-κB inhibitor abolished the functions of silencing of Fmo5 and PPARα promoting AFLD-induced apoptosis, inflammatory response, and liver injury.

Conclusions
Our data indicated that the co-expression of Fmo5 and PPARα was involved in AFLD-related gut microbiota composition, alleviated AFLD-induced liver injury, apoptosis and inflammatory response via
inhibiting the nuclear translocation of NF-κB p65 to inhibit NF-κB signaling pathway.

**Introduction**

The toxic effect of alcohol can affect the important organs of the body, especially the liver [1]. The liver disease caused by long-term heavy drinking usually only manifests as fatty liver at first, but with the continuation of drinking, the disease can continue to develop into alcoholic hepatitis and alcoholic liver fibrosis, resulting in alcoholic cirrhosis and hepatocellular carcinoma [1, 2]. In recent years, the incidence of alcoholic liver disease (alcoholic liver disease, ALD) is also increasing year by year. AFLD is the most common and reversible stage of alcoholic liver disease [3]. When the body is stimulated by alcohol, the excessive input of lipids leads to the disturbance of lipoprotein synthesis and metabolism in the liver, and the insufficient oxidation of fatty acids, which leads to the deposition of fat in the liver, resulting in fatty liver [4]. Although a large number of studies have been done on the pathogenesis of alcoholic fatty liver at home and abroad [3], there is still no effective treatment for alcoholic fatty liver in clinic. Therefore, a systematic study of the pathogenesis of AFLD can provide a more effective treatment for alcoholic liver disease.

The intestinal fl ora is a diverse microecosystem composed of more than 2000 species of bacteria and has 150 times more genomes than the human genome [5]. Intestinal microfl ora can affect the basic functions of host such as digestion and absorption, energy metabolism and immune defense by regulating endocrine, nerve and immunity [6, 7]. The composition of intestinal microfl ora is affected by external environment, diet and host itself. The imbalance of intestinal fl ora can make the body in a state of chronic inflammation for a long time [8]. Alcoholic fatty liver is closely related to the changes of intestinal fl ora [9, 10]. Therefore, to explore the specific mechanism of inflammatory response induced by intestinal fl ora imbalance in AFLD needs to be further studied.

Flavin containing monooxygenases (FMOs) catalyze the ingredients of medicine and diet and have the unique catalytic mechanism [11, 12]. Flavin containing monooxygenase 5 (Fmo5) is a member of FMOs family, and its expression in liver tissue of aging mice is lower than that of normal mice [13]. Salvia miltiorrhiza can significantly upregulate the expression in injured liver [14]. FMO5 as a regulator of glucose homeostasis and a potential sensor of gut bacteria [15]. This shows that Fmo5 has a protective effect on the liver and was closely related to the balance of intestinal fl ora. Previous bioinformatics studies showed that Fmo5 was significantly associated with intestinal fl ora and alcoholic diet. Therefore, the role of Fmo5 in AFLD-mediated intestinal fl ora disorder and liver injury needs to be further studied.

**Materials And Methods**

**Data sources**

The NCBI-GEO database is a free and public database containing gene profiles. Four microarray datasets (GSE8006 and GSE40334) were obtained from the GEO database (https://www.ncbi.nlm.nih.gov/gds/).
The gene expression profiles of GSE8006 data set included 4 control liquid diet group samples and 4 alcohol liquid diet group samples. The gene expression profiles of GSE40334 data set included 3 colon germ free group samples and 3 colon specific pathogen free group samples.

**Identification of DEGs**

GEO2R is an interactive web tool that can compare and analyze two different groups of samples under the same experimental conditions. In this study, the selected datasets of DEGs were analyzed using GEO2R. Subsequently, DEGs met the cutoff criteria of the adjusted \( P\)-value < 0.05 and |log fold change (FC)| > 1.0 were considered DEGs.

**Functional enrichment analysis of DEGs**

GO functional analysis and KEGG pathway analysis were carried out to predict the potential functions of the DEGs by using the Database for Annotation, Visualization and Integrated Discovery (DAVID; https://david.ncifcrf.gov/; version 6.8). DEGs from GSE8006 and GSE40334 data sets were submitted to the DAVID online program. The top 10 items of the cellular component (CC), biological process (BP), and molecular function (MF) categories and KEGG pathways were then sorted and presented in the form of bubble maps. These bubble plots were drawn using the ggplot2 R package based on \( P\)-value through the statistical software R (version 3.6.1). \( P < 0.05 \) was considered statistically significant.

**PPI network construction and hub gene identification**

The Search Tool for the Retrieval of Interacting Genes database (Version 10.0, http://string-db.org) was used to predict potential interactions between gene candidates at the protein level. A combined score of >0.4 (medium confidence score) was considered significant. Additionally, Cytoscape software (Version 7.2.0, http://www.cytoscape.org/) was utilized for constructing the PPI network. CytoHubba, a Cytoscape plugin, was utilized to explore PPI network hub genes; it provides a user-friendly interface to explore important nodes in biological networks and computes using eleven methods, of which MCC has a better performance in the PPI network (cytoHubba: identifying hub objects and sub-networks from complex interactome.).

**Weighted gene co-expression network analysis (WGCNA)**

The WGCNA R package was used to construct a weighted correlation network between the prognostic mRNAs. In the network, we used the pairwise Pearson coefficient to evaluate the weighted co-expression relationship between all genes in the adjacency matrix. The soft threshold was used to ensure a scale-free network. In the unsigned co-expression network, genes with high absolute correlations were clustered into the same module. The modules were also identified by hierarchical clustering of the weighted coefficient adjacency matrix to calculate the topological overlap matrix (TOM). In addition, the topological overlap of intramodules and adjacency modules was used for selecting the functional modules.
16S rDNA sequencing

Cecal contents were collected from some animals and frozen at -80°C. DNA was extracted using Stool DNA Extraction Kit (Qiagen) according to the manufacturer's instructions. To check DNA quality and 16S content prior to sequencing, universal primers were used for SYBR Green quantitative polymerase chain reaction (qPCR) with the following extended cycling protocol: 95°C 10 min; 95°C 15 sec, 60°C 30 sec, 72°C 30 sec for 40 cycles. Sequencing was completed at the Cincinnati University Children's Hospital Medical Center's DNA Sequencing and Genotyping Facility Core (Cincinnati, OH) as described [14]. All antibiotic-treated samples failed to yield 16S rDNA sequence data; one sample each from the ethanol- and pair-fed groups was excluded based on insufficient sequence data.

UPARSE and UTAX (http://www.drive5.com/usearch/manual/cmd_utax.html) were used to generate OTU tables from 16S rDNA read data and to make taxonomic assignments. QIIME package scripts were used for calculations of α-(PD_whole_tree, chao1, observed_otus and shannon) and β-(Bray-Curtis, Un-Weighted UniFrac, and Weighted UniFrac) diversity.

Animal Experimentation

Male C57BL/6J mice (22 ± 2 g, six weeks old) and Fmo5 gene knockout (Fmo5-/-) mice (20 ± 3 g, six weeks old) were purchased from the Beijing Vital River Laboratory Animal Technology Co., Ltd. (Beijing, China) and housed individually in cages with a 12-h light/dark cycle at 23 ± 2°C with full access to chow diet and water. All experiment protocols were approved by the Institutional Animal Care and Use Committee at the Zhengzhou University.

The AFLD mouse model was established using the modified Lieber–DeCarli liquid diet (21). The 40 mice were separated into four groups, each consisting of 10 mice. The first group (control group) and the third group (Fmo5-/- group) was fed with a normal standard growth diet while the other two groups (AFLD group and Fmo5-/-_AFLD group) were fed high-fat liquid diets (35% fat, 18% protein, 47% carbohydrates, provided by TROPHIC Animal Feed High-Tech Co., Ltd., Nantong, China) for an acclimation period of two weeks acclimation. While the amount of ethanol included increased over two weeks to reach a final concentration of 20% (v/v). Body weight gain and food intake were assessed once a week. The pair-fed control group (Con) was included in this model.

After 12 weeks, the fasted mice were euthanized and blood samples were collected. The feces in the colon were also harvested into 2-ml sterile tubes to assess the gut microbiota. The tissues were immediately removed and weighed, and the liver coefficient (liver weight/body weight) was calculated. The left lobe of the liver was instantly fixed in 10% buffered formalin for histology analysis with the rest of the tissues frozen in liquid nitrogen and stored at ~80°C until further use.

Hepatic Triglyceride Staining

The liver tissue was separated and rapidly fixed in 4% neutral buffered formalin solution for 24 h and then processed for paraffin embedding. Five-micrometer-thick paraffin sections were stained with
hematoxylin and eosin (H&E) and oil-red solution. The liver steatosis status was examined under a light microscope (Olympus, Tokyo, Japan), and photographed at 200 × magnification.

**Measurement of liver function and serum pro-inflammatory cytokines**

Blood samples were obtained from the common carotid artery to detect serum alanine aminotransferase (ALT), aspartate aminotransferase (AST), tumor necrosis factor-α (TNF-α) and interleukin-6 (IL-6) when rats were sacrificed at 24 h after reperfusion. The levels of serum ALT and AST were measured using corresponding kits (Jiancheng, Nanjing, China) to assess the liver function. Serum TNF-α and IL-6 levels were measured by radioimmunoassay kits (Albert Poole Biotechnology, Beijing, China).

**Apoptosis and immunohistochemical examination of liver**

Harvested livers were placed into 10% formalin immediately after excision, and immersed for 24 h. Liver specimens were then embedded in paraffin, and sections were cut at 5 μm. Sections were stained for: (1) Apoptosis was detected by the TdT-mediated dUTP-biotin nick end labeling (TUNEL) method using an *in situ* apoptosis detection kit (Boster Biological Engineering, Wuhan, China) as previously described. The number of cells with TUNEL-positive nuclei was counted from 20 randomly selected fields at × 200 magnification per liver sample. Results were expressed as the mean number of TUNEL-positive apoptotic hepatocytes per microscopic field; and (2) Immunobiological analysis of Fmo5, caspase-3, NF-κB p65 expression in paraffin wax-embedded liver sections were performed using a standard peroxidase-antiperoxidase technique as described previously, using a Fmo5 rabbit anti-mouse polyclonal antibody (Abcam Inc, United Kingdom) or mouse monoclonal antibodies against caspase-3 and NF-κB p65 (Santa Cruz, CA, United States) at a 1:150 dilution, with a biotinylated goat anti-rabbit or goat anti-mouse antibody (Santa Cruz, CA, United States) as the secondary antibody. Brown color in the cytoplasm of the hepatocytes was evaluated as positive staining.

**Cell culture**

Human normal hepatocyte cell line L02 was purchased from the Chinese Academy of Science (Shanghai, China). Cells were cultivated in RPMI-1640 medium (HyClone, Logan, UT, USA) supplemented with 10% heat-inactivated fetal bovine serum (FBS), 100 U/ml penicillin, and 100 μg/ml streptomycin according to the manufacturer’s instructions. Cells were maintained at 37°C in a humidified atmosphere with 5% CO₂. Half of the growth medium was changed each day. L02 cells at 60-70% confluency after the fourth passage were plated for the experiments.

**Construction of AFLD cell model**

To deplete L02 cells of iron, the chelator, desferrioxamine mesylate (DFO) (100 μM, D9533; Sigma-Aldrich), was added to the cell culture medium. Iron loading of L02 cells was achieved by incubating cells in the presence of iron-dextran (0–100 μg/ml; Sigma-Aldrich). Cells were collected 24 h after treatment and at 48 h after treatment for extraction of mRNA and protein, respectively. For alcohol treatment, L02
cells were treated for 48 h with 200 nM alcohol (Sigma-Aldrich); then, cells were treated for 24 h with DFO (100 μM) in the presence of 200 mM alcohol.

**Cell viability**

The cells were seeded in 96-well plates at a density of 5 × 10^3 cells/well and cultured overnight. Subsequently, the cells were pretreated with various concentrations of ethyl alcohol (0-200 nM) for 72 h at 37°C. Cell viability was estimated using the cell counting kit-8 colorimetric assay (CCK-8 assay, Dojindo Molecular Technologies, Inc., Kumamoto, Japan). In addition, the viability was estimated using the cell counting kit-8 colorimetric assay.

**Oil Red O staining**

The Oil Red O stock solution was composed of 5 g Oil Red O in 100 ml isopropyl alcohol. Fresh Oil Red O working solution was prepared by diluting the Oil Red O stock with distilled water in a 3:2 ratio. Cells were washed twice in PBS after PA treatment, fixed with 4% formaldehyde for 30 min, and stained with Oil Red O working solution for 40 min at room temperature. Then, L02 cells were washed with 60% isopropyl alcohol once followed by washing with water twice. The nuclei of the cells were stained with Hematoxylin. The number of Oil Red O-positive cells was counted in replicates in 15 randomly selected fields at 20 × magnification under a bright-field microscope.

**Co-immunoprecipitation (Co-IP)**

For Co-IP, anti-Fmo5 was coupled with protein A-Sepharose beads (Sigma) in RIPA buffer overnight at 4°C. The immune complex was then added to cell lysate and incubated at 4°C for 2 h. Endogenous Fmo5 was immunoprecipitated using an Fmo5 antibody (Santa Cruz Biotechnology, Inc., Dallas, TX, USA). The IP samples were analyzed by western blot using the following antibodies: Fmo5 (Abcam, Cambridge, MA, USA) and PPARα (Abcam).

**Triglyceride (TG) assay**

Cells from the different groups were harvested and washed twice with PBS. The intracellular triglycerides were measured using a triglyceride assay kit according to the manufacturer’s instruction (Applygen Technologies Inc., Beijing, China). The TG concentrations were normalized to the total cell protein concentration.

**Cell transfection**

SiRNA was used to specifically target Fmo5 or PPARα. L02 cells were transiently transfected with Fmo5 siRNA or PPARα siRNA (Genesis Biotechnology, Wuhan, China) or negative control siRNA (Santa Cruz, CA, USA) according to the standard protocols. Briefly, cells were seeded at 2 × 10^5 in 12-well plates without antibiotics. Next, cells with 30–50% confluence were transfected with 50 nM siRNA or negative control using Lipofectamine 2000 (Invitrogen, USA) according to the manufacturer’s protocol.
Fmo5 expressing plasmids or PPARα expressing plasmids were obtained from Genesis Biotechnology (Wuhan, China). Briefly, cells were seeded at 2 × 10^5 in 12-well plates without antibiotics. Next, cells with 30–50% confluence were transfected with 30 nM expressing plasmids or negative control using Lipofectamine 2000 (Invitrogen, USA) according to the manufacturer’s protocol.

**Determination of apoptosis by Annexin V-FITC/PI staining**

Chondrocyte apoptosis was determined by Annexin V-FITC/PI double labeling according to the manufacturer's protocol. Briefly, after indicated cultures, chondrocytes were digested and suspended in binding buffer. Then 5 μl Annexin V and 5 μl PI solutions were added into cells and incubated for 15 min. Apoptotic rate was detected by BD FACSVerse™ flow cytometer (Becton Dickinson, Heidelberg, Germany) and analyzed by Cell Quest software (BD Biosciences). Experiments were conducted in triplicate.

**Intracellular reactive oxygen species (ROS) levels detection by flow cytometry**

Intracellular reactive oxygen species (ROS) levels were measured by ROS Assay Kit (Beyotime, Shanghai, China) with an oxidation-sensitive fluorescent probe dye, DCFH-DA (2, 7- dichlorodihydro fluorescein diacetate). In brief, cells were seeded in 6-well plates (5 × 10^3 cells per well) and incubated at 37°C, 5% CO₂ atmosphere for 24 h. Next, the cells were collected and washed with PBS, and then incubated at 37°C with DMEM containing 10 μM DCFH-DA for 30 min. After that, cells were washed three times using serum-free DMEM. The samples were finally analyzed using a FACScan flow cytometer and Flowjo 7.6.1 software.

**Immunofluorescence staining**

The cells were seeded onto 96 black well plate (5 × 10^4 cells/well) and cultured until 80% confluence. After 30 min of incubation, cells were fixed with 4% paraformaldehyde and permeabilized with 0.05% Triton X. Then, cells were incubated overnight with primary antibodies against NF-κB p65 (1:400) at 4°C. Samples were then washed with PBS, incubated for an hour with Alexa Fluor 488-conjugated goat anti-rabbit IgG (1:400), followed by DNA staining for 15 min with hoechst 33342 staining (1: 1000). The fluorescently labeled NF-κB p65 was detected by Operetta High-Content Imaging System.

**Western blot analyses of tissue and cells**

Dissected livers were snap-frozen in liquid nitrogen and protein lysates for the detection of HIF1α and HIF2α proteins were extracted with the NE-PER kit (Pierce). Primary hepatocytes and Hep3B cells were lysed in buffer of 50 mM Tris–HCl, pH 8, 150 mM NaCl, 0.5% NP-40, 1 mM PMSF and a complete protease inhibitor cocktail tablet (Roche), kept on ice for 20 min and centrifuged at 12 000 g for 20 min at 4°C. 100 μg of proteins from liver or cells were resolved by SDS-PAGE, blotted, and probed with the following primary antibodies: Fmo5, PPARα, caspase-3, NF-kb, p65, IL-6, TNF-a, and GAPDH. The secondary antibody, either anti-mouse or anti-rabbit, was conjugated to horseradish peroxidase (1:5000; Bio-Rad Laboratories). The Pierce ECL system (ThermoScientific) was used for detection.
**RNA Extraction and Real-Time PCR**

The total RNA was extracted by using a TRIzol reagent. The first-strand cDNA was synthesized from 1 μg of total RNA using the Reverse Transcription System Bestar qPCR RT Kit according to the manufacturer instruction. Real-time PCR was carried out with an ABI 7500 Real-Time PCR System (Applied Biosystems, Lincoln Centre Drive, Foster City, CA 94404, USA). Each assay was performed in triplicate. The relative amount of Fmo5 and PPARα were calculated using a $2^{-\Delta\Delta C_t}$ method and normalized using GAPDH as an internal control. The primers used in this study were shown below: for Fmo5, 5'-CCAGTTACGTGAATGATTCG-3' (forward), 5'-AGCGCGTGTGAATGCAGGCC-3' (reverse); for PPARα: 5'-CTTATAACTGCGGGAGGAC-3' (forward), 5'-TACGTGCTGAACATGAT-3' (reverse); and for GAPDH: 5'-CATCGATTAGGCATGC CGC-3' (forward), 5'-TCGTAACTAGGGCTACCGC-3' (reverse).

**Measurement of TG content in liver**

TG content in liver was determined as previously described. Briefly, 250 mg of liver sample was homogenized in 1.5 ml of methanol, and then added with 5.0 ml of MTBE and shaken at room temperature for 1 h. Subsequently, 1.25 ml of high purity water was added and mixed for 10 min, followed by centrifuging at 1,000 g for 10 min, and the upper organic layer was collected. The aqueous layer was re-extracted with MTBE/methanol/water mixture (10/3/2.5 v/v/v; 2 ml) and the combined organic layers were dried under nitrogen. The extracted dried lipids were dissolved in a mixture of triton X-114/methanol (2:1 v/v, 60 μl) and analyzed for triglyceride (L-Type TGH) using commercially available kits (Wako Diagnostics, Richmond, VA, USA).

**Statistical analysis**

Sequence data were processed using Quantitative Insights into Microbial Ecology (QIIME, version 1.9.1). The low-quality sequences, which had lengths of <150 bp and average Phred scores of <20 and contained ambiguous bases and mononucleotide repeats of >8 bp, were filtered with the following criteria. Paired-end reads were assembled using FLASH. The remaining high-quality sequences were clustered into operational taxonomic units (OTUs) at 97% sequence identity by UCLUST (Edgar 2010). OTU taxonomic classification was conducted by BLAST and the OTUs containing more than 99.999% of total sequences across all samples were reserved.

QIIME (version 1.9.1) was used to calculate all diversity indexes in these samples, and R software (version 2.15.3) was used for visual analysis. LEfSe software (LEfSe 1.0) was used for linear discriminant analysis. Principal coordinate analysis (PCoA) was performed using Bray-Curtis. The functions of gut microbiota were predicted for the CON and AFLD groups using Tax4Fun in the KEGG database.

All results are presented as means ± standard deviation. Data were analyzed with SPSS 19.0 using One-way analysis of variance (ANOVA), and, when appropriate, using a two-tailed Student’s t test between different groups. Differences among groups were evaluated for significance with the comparable
variances, followed by Tukey's and least significant difference (LSD) tests. $P < 0.05$ was considered statistically significant.

## Results

**AFLD regulates the gut microbiota composition in mice**

The microbiota can influence the development and progression of AFLD. High-throughput 16S rRNA gene sequencing produced a total of 25253 good-quality sequences from 6 samples. These were grouped into 382 OUTs based on 97% similarity (Figure 1A). 285 OTUs were identified as core bacterial OTUs (Figure 1B). We detected 7 bacterial phyla and the 60 genera among the mice. The structure and composition of gut microbiota were significantly influenced by AFLD. The results of the top 10 most abundant OTUs at all taxonomic levels in samples, as inferred by GraPhlAn, showed that *Bacteroidetes*, *Firmicutes*, and *Proteobacteria* were the most abundant phyla (Figure 1C), and *Muribaculaceae*, *Lachnospiraceae*, *Runminococcaceae*, and *Erysipelotrichaceae* were the most abundant family among the OTUs (Figure 1F). Additionally, the relative bacterial abundance between CON group and AFLD group at the phylum and family levels was shown in Figure 1D and 1E, respectively. At the phylum level, the taxonomic profiles indicated that the proportions of *Bacteroidetes* and *Proteobacteria* decreased significantly, and the abundance of *Firmicutes* increased markedly in the AFLD group. The relative abundance ration of *Firmicutes* and *Bacteroidetes* increased significantly in the AFLD group. At the family level, the taxonomic profiles indicated that the proportions of *Muribaculaceae* decreased significantly, and the abundance of *Lachnospiraceae*, *Runminococcaceae*, and *Erysipelotrichaceae* increased markedly in the AFLD group. A heat map of genera indicated that the 1 genera (*Firmicutes*) increased significantly and the 8 genera decreased significantly (Figure 1H) at the phylum level, and the 12 genera increased significantly, and the 8 genera decreased significantly (Figure 1I) at the family level. The alpha diversity (observed species, Shannon, Simpson, ACE, Chao1, and Good’s coverage) of intestinal flora was calculated between CON group and AFLD group in mice. The Shannon index was significantly lower in the AFLD group than in the CON group (PShannon = 0.04; Figure 1J). In addition, there was no difference in observed species index, Simpson index, ACE index, Chao1 index, and Good’s coverage index between the CON group and AFLD group. Multi-response permutation procedure (MRPP) testing between the CON group and AFLD group and the CON and AFLD groups yielded A > 0. The intergroup differences were larger than the intragroup differences, which further indicate that the grouping of this study was reasonable. Combined with the nonmetric multi-dimensional scaling (NMDS) plot (Figure 1K) and PCoA plots (Figure 1L), it was shown that the CON and AFLD group were significantly different. The results of the LDA Effect Size analysis show phylum, class, order, family, genus, or species with significantly different abundance between CON group and AFLD group. The results have two parts: LDA value distribution histogram and evolutionary branch diagram (phylogenetic distribution). *Bacteroidetes* is the most important phylum, class, order, family, genus, or species in the CON group, and *Firmicutes* is the most phylum, class, order, family, genus, or species in the AFLD group (Figure 2).
Prediction of gut microbiome function

The Tax4Fun analysis based on metagenomes was used to predict the function of gut microbiota that were influenced by AFLD in mice. According to the results of the predicted function, the top 20 functional information of maximum abundance was selected for each sample of the KEGG classification level, and relative functional abundance heat map was generated (Figure 3A). At level 3, the results revealed that 29 KEGG pathways changed significantly between AFLD group and CON group using Welch's test (Figure 3B). In particular, we found several interesting changes among the 29 altered KEGG pathways at level 3. PPAR signaling pathway was increased in the CON group compared to that in the AFLD group, whereas, apoptosis was increased in the AFLD group compared to that in the CON group.

Identification of DEGs and construction of PPI network in GSE8006 data set

A total of 292 DEGs (174 downregulated genes and 118 upregulated genes) in GSE8006 data set were shown in Figure 4A and 4B, and used to preform GO, and KEGG enrichment analysis and construct a PPI network using DAVID and STRING database. Our results showed that 292 DEGs were mainly enriched in three GO terms, including BP (Figure 4C-4E), CC (Figure 4F-4H) and MF (Figure 4I-4K). KEGG enrichment analysis results showed that PPAR signaling pathway was an important KEGG pathway (Figure 4L). Next, the top 10 hub genes of a PPI network (Figure 4M), including Fmo5, Cmpk2, Isg15, Ifit2, Ifit47, Usp18, Ifit3, Ifit1, Rsad2, and Irf7, were identified using cytohubba plug-in of Cytoscape software based on “MCC” method (Figure 4N).

Identification of DEGs and construction of PPI network in GSE40334 data set

A total of 246 DEGs (98 downregulated genes and 148 upregulated genes) in 40334 data set were shown in Figure 5A and 5B, and used to preform GO, and KEGG enrichment analysis and construct a PPI network using DAVID and STRING database. Our results showed that 246 DEGs were mainly enriched in three GO terms, including BP (Figure 5C-5E), CC (Figure 5F-5H) and MF (Figure 5I-5K). KEGG enrichment analysis results showed that PPAR signaling pathway was an important KEGG pathway (Figure 5L). Next, the top 10 hub genes of a PPI network (Figure 5M), including Gstm2, Mgst3, Fmo5, Cyp2c55, Cyp2c39, Gstm4, Gstm3, Gsta4, Gsta1, and Gsta2, were identified using cytohubba plug-in of Cytoscape software based on “MCC” method (Figure 5N).

WGCNA analysis of GSE8006 and GSE40334 data sets

The WGCNA was performed to analyze the GSE8006 microarray data. First, the dataset was properly normalized and filtered to reduce outliers (Figure 6A). The power value was screened out that when the value was 6, the scale independence was 0.86 (Figure 6B). Thus, this network conformed to the power-law distribution and was closer to the real biological network state. As shown in Figure 6C, the cluster dendrogram contained 31 co-expression modules. The co-expressed genes were primarily clustered in the darkmagenta module. In order to explore the function of a module, we selected the darkmagenta module for further analysis. Our results indicated that the co-expression genes in the darkmagenta module was
significantly associated with colon specific pathogen free (correlation coefficient=0.86, p=2.3e-125). Therefore, the darkmagenta module was a central module in all of the modules (Figure 6D). A total of 181 hub genes in the darkmagenta module were identified according to the absolute value of the threshold of correlation degree > 0.8. We also constructed a PPI network of the 181 hub genes using STRING database and Cytoscape was used to perform the visualization of a PPI network (Figure 6E). Next, the top 10 hub genes of a PPI network, including Fcgr1, Fmo5, C3, Slc27a2, Cybb, C1qa, Gstt1, Gstp1, Apob, and Gstk1, were identified using cytohubba plug-in of Cytoscape software based on “MCC“ method (Figure 6F).

The WGCNA was performed to analyze the GSE40334 microarray data. First, the dataset was properly normalized and filtered to reduce outliers (Figure 6G). The power value was screened out that when the value was 11, the scale independence was 0.83 (Figure 6H). Thus, this network conformed to the power-law distribution and was closer to the real biological network state. As shown in Figure 6I, the cluster dendrogram contained 111 co-expression modules. The co-expressed genes were primarily clustered in the darkturquoise module. In order to explore the function of a module, we selected the darkturquoise module for further analysis. Our results indicated that the co-expression genes in the darkturquoise module was significantly associated with alcohol liquid diet (correlation coefficient=0.75, P=9.6e-187). Therefore, the darkturquoise module was a central module in all of the modules (Figure 6J). A total of 172 hub genes in the darkturquoise module were identified according to the absolute value of the threshold of correlation degree > 0.8. We also constructed a PPI network of the 172 hub genes using STRING database and Cytoscape was used to perform the visualization of a PPI network (Figure 6K). Next, the top 10 hub genes of a PPI network, including Fmo5, Hist2hg2aa1, Wdr5, Polr2i, Cyp2u1, Hsd3b5, Cyp4a12a, Cyp2c37, Ugt2b35, and Cyp3a11, were identified using cytohubba plug-in of Cytoscape software based on “MCC“ method (Figure 6L).

**Fmo5 plays an important role in the AFLD-induced apoptosis and inflammatory response in the small intestine and liver tissues of mice and is closely related to PPARα expression and the NF-κB signaling pathway in vivo and in vitro**

Our results showed that Fmo5 was the only one overlapping gene from the hub genes of the 4 PPI network (Figure 7A). Coexpedia database indicated that there is a co-expression relationship between Fmo5 and PPARα (Figure 7B). Furthermore, the protein levels of Fmo5 and PPARα in the small intestine and liver tissues of mice with AFLD decreased significantly and the protein levels of cleaved caspase-3, IL-6, and TNF-α and the phosphorylation level of NF-κB p65 in the small intestine and liver tissues of mice with AFLD increased significantly (Figure 7C). In order to explore the function of Fmo5 in AFLD, we selected Fmo5 knockout mice for further analysis. The results showed that Fmo5 knockout aggravated the AFLD-induced hepatic injury, apoptosis and inflammatory response (Figure 7D-7F). In addition, Fmo5 knockout inhibited PPARα expression and activated the NF-κB signaling pathway using western blot and IHC assay in mice with AFLD (Figure 7G and 7H).

**In vitro**, we found that PPARα expression in Fmo5 overexpression transfected L02 cells increased significantly and in siRNA targeting Fmo5 transfected L02 cells decreased significantly by RT-PCR and
western blot assays (Figure 8A-8E). Co-IP experiments indicated that Fmo5 directly interact with PPARα in L02 cells (Figure 8F). CCK-8 results showed that ethyl alcohol inhibited cell growth in a concentration dependent manner (Figure 8G). Therefore, 200 nM was considered as the action concentration to induce AFLD in vitro. Our results showed that the accumulation of lipid droplet was observed in cytoplasm of L02 cell using Oil Red O staining (Figure 8I). The levels of TG, AST, and ALT in AFLD group increased significantly (Figure 8J-8L). It showed that AFLD cell model constructed successfully. In addition, the protein levels of Fmo5 and PPARα in AFLD group decreased significantly and the phosphorylation level of NF-κB p65 in AFLD group increased significantly (Figure 8M). Next, the functions of AFLD inhibiting cell viability, inducing cell apoptosis, and increasing the levels of TG, AST, ALT, IL-6, TNFα, and ROS were upregulated by Fmo5 inhibition, whereas, which were reversed by Fmo5 overexpression (Figure 8N-8U). In addition, inhibition of Fmo5 upregulated cleaved caspase-3 expression, and the phosphorylation level of NF-κB p65 and inhibited PPARα expression in AFLD cell model, meanwhile, Fmo5 overexpression downregulated cleaved caspase-3 expression, and the phosphorylation level of NF-κB p65 and increased PPARα expression in AFLD cell model (Figure 8V).

**Fmo5 and PPARα co-expression alleviates AFLD-induced apoptosis, inflammatory response and the nuclear translocation of NF-κB p65 in vitro**

The results that PPARα expression in PPARα overexpression transfected L02 cells increased significantly and in siRNA targeting PPARα transfected L02 cells decreased significantly by RT-PCR and western blot assays (Figure 9A). The functions of Fmo5 silencing or PPARα silencing inhibiting AFLD-induced the inhibition of cell viability, cell apoptosis, and the elevator of TG, AST, ALT, IL-6, TNFα, and ROS levels were abolished by PPARα overexpression or Fmo5 overexpression (Figure 9B-9I). Likewise, siRNA-Fmo5 and siRNA-PPARα co-transfection inhibited cell viability, induced cell apoptosis, and increased the levels of TG, AST, ALT, IL-6, TNFα, and ROS in AFLD cell module. These functions in AFLD+siFmo5+siPPARα group were significantly higher than that in AFLD+siFmo5 group. Immunofluorescence staining results proved that Fmo5 inhibition promoted AFLD-induced the nuclear translocation of NF-κB in L02 cells (Figure 9J). However, PPARα overexpression reduced AFLD-induced the nuclear translocation of NF-κB p65 in siRNA-Fmo5 transfected L02 cells, meanwhile, Fmo5 overexpression reduced AFLD-induced the nuclear translocation of NF-κB p65 in siRNA-PPARα transfected L02 cells. In addition, the functions of co-transfection of siRNA-Fmo5 and siRNA-PPARα promoting the nuclear translocation of NF-κB p65 were significantly higher than that of siRNA-Fmo5 transfection in AFLD cell model.

**Fmo5 and PPARα co-expression alleviates AFLD-induced apoptosis and inflammatory response via inhibiting the activation of NF-κB signaling pathway in vitro**

CHS-828 (NF-κB specific inhibitor), inhibited significantly the phosphorylation level of NF-κB p65 and increased significantly the protein levels of Fmo5 and PPARα (Figure 10A). CCK-8 assay showed that CHS-828 reversed the functions of the co-transfection of siRNA-Fmo5 and siRNA-PPARα promoting the inhibition of cell viability and the increased of TG, AST, ALT, IL-6, and TNFα levels in AFLD cell model (Figure 10B and 10C). Flow cytometry analysis showed that the effects co-transfection of siRNA-Fmo5
and siRNA-PPARα on promoting AFLD-induced cell apoptosis and ROS level were abolished by CHS-828 (Figure 10D and 10E).

**Discussion**

There is a close relationship between intestine and liver in anatomy and function, that is, intestinal-liver axis [16]. With the decrease of liver function, the intestinal balance is destroyed. With the decrease of liver function, the intestinal balance is destroyed [17]. The accumulation of harmful substances such as alcohol leads to liver injury and promotes the secretion of bile acid into the intestinal tract by the liver, thereby regulating the activity of various metabolic reactions in the intestinal tract [2, 4, 5, 13]. The change of the type and number of intestinal flora is an important factor leading to abnormal liver function [18]. Liver disease can also lead to overgrowth and imbalance of intestinal flora. Long-term drinking will cause intestinal flora to release endotoxin and destroy intestinal barrier, thus excessive production of inflammatory factors (such as IL-6, TNF α), and promote the occurrence and development of AFLD [19]. Inflammation increases intestinal osmotic pressure, causing endotoxin to cross the intestinal barrier and enter the body, forming a vicious circle. The result of 16S rDNA sequencing indicated that the ratio of *Firmicutes/Bacteroidetes* in gut of mice with AFLD was higher than that in gut of normal mice and the abundance of bacteria in gut of mice with AFLD was lower than that in gut of normal mice. These data showed that AFLD induced gut microbiota disturbance. Intestinal flora can directly act on intestinal epithelial cells, affect their gene expression, and then affect intestinal function and the activation of related signal pathways. Disturbance of intestinal flora can further damage the intestinal barrier and spread alcohol-related damage to liver tissue [20]. Therefore, the expression of bacterial genes related to inflammation and alcohol metabolism will change significantly in the intestines and liver of patients with AFLD. Our results showed that gut microbiome function was significantly related to PPAR signaling pathway and apoptosis. The results of bioinformatics analysis showed that Fmo5 was the only hub gene, significantly associated with intestinal microorganisms and alcoholic diet, which had a protective effect on the liver. In addition, interestingly, there is a co-expression relationship between Fmo5 and PPARα. PPARα is a key gene in PPAR signaling pathway, which plays a role in cellular lipids by regulating the expression of genes involved in lipid metabolism in the liver [21, 22]. PPARα gene knockout promotes inflammation and aggravates lipid deposition in the liver. It can be seen that Fmo5 and PPARα are beneficial to the metabolic function of the liver. Activated PPARα reduces the release of inflammatory cytokines by inhibiting NF-κB signal pathway.

NF-κB is an upstream signal molecule of many inflammatory mediators, which can promote the production of many inflammatory mediators and directly participate in acute and chronic inflammation of the liver [23]. In addition, NF-κB exerts its anti-apoptotic effect by directly activating anti-apoptotic genes [24]. Liver injury in alcoholic fatty liver is closely related to hepatocyte apoptosis, which in turn is related to the activation of NF-κB. Studies have shown that in the process of establishing the model of alcoholic fatty liver, NF-κB is consistent with the degree of steatosis in liver tissue at the gene transcriptional level. The level of NF-κB in serum of patients with alcoholic fatty liver is positively correlated with the indexes of serum liver function. NF-κB is involved in the occurrence and development
of alcoholic fatty liver [25, 26, 27]. Inflammation induces the production of free radicals or decreases the ability of the body to scavenge free radicals, thus prompting the body to release a large amount of ROS [28]. ROS can further activate inflammatory response by activating NF-κB signal pathway [29]. The results of animal experiments show that Fmo5 and PPARα expression in small intestine and liver tissues of mice with AFLD decreased significantly and the apoptosis-related protein (cleaved caspase-3), phospho-NF-κB p65, and inflammation-associated cytokines (IL-6 and TNFα) in in small intestine and liver tissues of mice with AFLD increased significantly. To sum up, AFLD induces intestinal flora imbalance, reduces the expression of intestinal-liver protection related factor (Fmo5 and PPARα), activates NF-κB signal pathway, and promotes apoptosis and inflammation. However, the anti-inflammatory and anti-apoptotic effects of Fmo5 in the liver need to be further studied.

Fmo5 gene knockout can significantly aggravate AFLD-mediated liver injury, lipid accumulation and apoptosis, and promote the release of inflammatory cytokines. In addition, PPARα expression in liver tissue of Fmo5/- mice and Fmo5/- mice with AFLD decreased significantly. Fmo5 knockout promoted the activation of NF-κB signaling pathway caused by AFLD. The above results further suggest that AFLD activates NF-κB signal pathway by inhibiting the expression of Fmo5 and PPARα, thus promoting inflammation and apoptosis and damaging liver tissue. The results of cell experiment in vitro showed that Fmo5 and PPARα were co-expressed in liver cells, and AFLD promoted the nuclear translocation of NF-κB p65 by upregulating the phosphorylation level of NF-κB p65, thus activating NF-κB signal pathway. Silencing of Fmo5 and PPARα can aggravate the role of Fmo5 silencing in promoting AFLD-induced apoptosis, inflammatory response, upregulation of ROS level and activation of NF-κB signal pathway. Rescue experiments showed that NF-κB inhibitor reversed the functions of silencing of Fmo5 and PPARα promoting cell apoptosis, and inflammatory response, increasing ROS level, and activating NF-κB signaling pathway caused by AFLD. These results suggest that the co-expression of Fmo5 and PPARα in liver tissue has a protective effect on AFLD-mediated liver injury, which is closely related to the inhibition of the activation of NF-κB signal pathway.

In conclusion, Fmo5 and PPARα are significantly downregulated in the liver tissue of AFLD and are closely related to intestinal flora and alcoholic diet. AFLD induces intestinal flora imbalance and reduces the expression of intestinal-liver protective factor α (Fmo5 and PPARα), thus activating NF-κB signal pathway, thereby promoting apoptosis and inflammation. Co-expression of Fmo5 and PPARα reduces AFLD-mediated apoptosis, inflammation and liver injury by inhibiting the activation of NF-κB signal pathway.

**Abbreviations**
Declarations

Ethics approval and consent to participate

Study were approved by the Ethics Committee of the First Affiliated Hospital of Zhengzhou University. All animal protocols were performed in accordance with the guidelines of the Institutional Animal Care and Use Committee of Zhengzhou University.

Consent for publication

Not applicable.

Availability of data and materials

The datasets generated/analyzed during the current study are available.

Competing interests

The authors declare that they have no competing interests.

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Authors’ contributions

Lingjian Kong and Jing Chen wrote the main manuscript and analyzed the data. Xiaoli Ji, Qian Qin, and Deliang Li performed the experiments. Lingjian Kong, Huiyu Yang, Dan Liu, and Meiling Sun designed the study. All authors read and approved the final manuscript.

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Figures
Figure 1

Bacterial community analysis and comparison (A) Number of Tags/or 16S OTUs amongst mice with AFLD. (B) Core bacterial operational taxonomic units (OTUs) in mice from different groups. The core community was determined based on OTUs detected in every sample. The OTUs were assigned at a 97% sequences similarity threshold, and a Venn diagram was used to summarize the number of common OTUs between the two groups. The composition and abundance distributions of each group at the phylum (C) and family (F) levels were shown using QIIME software. At the phylum (D and E) and family (G) levels, pairwise comparisons, conducted to determine the sequence amounts between two groups, were presented as pair-wise comparisons using Metastats analysis. ** represents $P < 0.01$ compared with the CON group. A heat map of genera at the phylum (H) and family (I) levels were shown. Red represents upregulation and green represents downregulation. (J) The alpha diversity (observed species, Shannon, Simpson, ACE, Chao1, and Good's coverage) of intestinal flora between CON group and AFLD group was shown. Cluster analysis result for both CON group and AFLD group according to NMDS (K) and Principal coordinate analysis (PCoA) (L). PCoA was performed using Bray-Curtis.
Figure 2

Results of LEFSE analysis. The histogram shows species whose LAD Score exceeded the default value of $|2|$. The length of the histogram represents the impact of the phylum (A), class (B), order (C), family (D), genus (E), or species (F). The different species have the same color as the group in the evolutionary branch diagram. Red represents AFLD group and green represents CON group.
Figure 3

Analysis of gut microbiome function (A) Relative abundance heat map of microbiota functions based on the KEGG database on the third level. (B) A total of 29 significantly changed the KEGG pathways in AFLD mice on the third level using Welch's test. These data were obtained by Tax4Fun.
Figure 4

Identification of DEGs and construction of PPI network in GSE8006 data set (A) The heat map for DEGs. (B) The volcano plot for DEGs. Black dots represent genes which are not differentially expressed, left dots represent the upregulated genes, and the right dots represent the downregulated genes. GO BP term was shown in (C-E). GO CC term was shown in (F-H), and GO MF term was shown in (I-K). (L) KEGG enrichment analysis was shown based on DAVID database. (M) A PPI network of DEGs was visualized by Cytoscape software. (N) The top 10 hub genes were identified by the cytohubba plug-in of Cytoscape software.
Figure 5

Identification of DEGs and construction of PPI network in GSE40334 data set (A) The heat map for DEGs. (B) The volcano plot for DEGs. Black dots represent genes which are not differentially expressed, left dots represent the upregulated genes, and the right dots represent the downregulated genes. GO BP term was shown in (C-E). GO CC term was shown in (F-H), and GO MF term was shown in (I-K). (L) KEGG enrichment analysis was shown based on DAVID database. (M) A PPI network of DEGs was visualized by Cytoscape software. (N) The top 10 hub genes were identified by the cytohubba plug-in of Cytoscape software.
Figure 6

WGCNA analysis of GSE8006 and GSE40334 data sets (A) One outlier (GSE8006) was detected by sample clustering. (B) analysis of the scale-free topology model fit index for soft threshold powers (β) and the mean connectivity for soft threshold powers. (C) A cluster dendrogram was built based on the dissimilarity of the topological overlap, which presents 31 gene co-expression modules in colon specific pathogen free group. Each designated color represents a certain gene module. (D) The module relationships and significance. (E) A PPI network of the hug genes from in the darkmagenta module was visualized by Cytoscape software. (F) The top 10 hub genes of this PPI network were identified by the cytohubba plug-in of Cytoscape software. (G) One outlier (GSE40334) was detected by sample clustering. (H) analysis of the scale-free topology model fit index for soft threshold powers (β) and the mean connectivity for soft threshold powers. (I) A cluster dendrogram was built based on the dissimilarity of the topological overlap, which presents 111 gene co-expression modules in colon specific pathogen free group. Each designated color represents a certain gene module. (K) The module relationships and significance. (K) A PPI network of the hug genes from in the darkturquoise module was visualized by Cytoscape software. (L) The top 10 hub genes of this PPI network were identified by the cytohubba plug-in of Cytoscape software.
Figure 7

Fmo5 plays an important role in the AFLD-induced apoptosis and inflammatory response in the small intestine and liver tissues of mice and is closely related to PPARα expression and the NF-κB signaling pathway in vivo (A) Venn map of the one overlapping gene from the hub genes of the above 4 PPI network. (B) The co-expression network of Fmo5 was constructed using Coexpedia database. (C) Western blot assay was used to detect the expression of Fmo5, PPARα, cleaved caspase-3, phospho-NF-κB p65, NF-κB p65, IL-6 and TNF-α in the small intestine and liver tissues between the control group and the AFLD group. (D) ELISA assay was used to detect the levels of ALT, AST, and TG in blood sample between the control group and the AFLD group. (E) ELISA assay was used to detect the levels of IL-6 and TNF-α in blood sample between the control group and the AFLD group. (F) H&E staining was used to observe the pathological change in liver tissues among the four group, Oil Red staining was used to observe the lipid deposition in liver tissues among the four group, and TUNEL was used to detect the apoptosis in liver
tissues among the four group. (G) Western blot assay was used to detect the expression of Fmo5, PPARα, phospho-NF-κB p65 and NF-κB p65 in liver tissues among the four group. (H) IHC assay was detect the expression of PPARα, cleaved caspase-3, and phospho-NF-κB p65 in liver tissues among the four group. GAPDH used as a load control. Data are presented as the mean ± standard deviation. ** P < 0.01 vs. Control group, ## P < 0.01 vs. AFLD group, and ^^ P < 0.01 vs. Fmo5/- group.
Figure 8

Fmo5 plays an important role in the AFLD-induced apoptosis and inflammatory response in the small intestine and liver tissues of mice and is closely related to PPARα expression and the NF-κB signaling pathway in vitro (A-C) Western blot assay was used to detect the protein levels of Fmo5 and PPARα in siRNA targeting Fmo5 transfected L02 cell or vector expressing Fmo5 transfected L02 cells. (D-E) RT-PCR assay was used to detect the mRNA level of Fmo5 and PPARα in siRNA targeting Fmo5 transfected L02 cell or vector expressing Fmo5 transfected L02 cells. (F) Co-immunoprecipitation experiments showed that Fmo5 directly interact with PPARα in L02 cell. (G) CCK-8 assay was used to detect the cell growth inhibition rate after L02 cells were treated with 0-200 nM of ethyl alcohol. (H) CCK-8 was used to detect the cell viability between the control and AFLD groups. AFLD cell model was shown in I. (J-L) ELISA assay was used to detect the levels of TG, AST, and ALT in L02 cells between the control and AFLD groups. (M) Western blot assay was used to detect the protein levels of Fmo5, PPARα, phospho-NF-κB p65 and NF-κB p65 in L02 cells between the control and AFLD groups. ELISA assay was used to detect TG level (N), AST level (O), ALT level (P), TNF-α level (R), and IL-6 level (S) in L02 cells among the six groups. (Q) CCK-8 assay was used to detect the cell viability among the six groups. Flow cytometry assay was used to detect cell apoptosis (T) and ROS level (U) in L02 cells among the six groups. (V) Western blot assay was used to detect the protein levels of cleaved caspase-3, phospho-NF-κB p65, NF-κB p65 and PPARα in L02 cells among the six groups. GAPDH used as a load control. Data are presented as the mean ± standard deviation. Con+siNC vs. Con+NC=ns; AFLD+siNC vs. AFLD+NC=ns. ** P < 0.01 vs. Con+siNC and Con+NC groups, ## P < 0.01 vs. AFLD+siNC and AFLD+NC groups.
Figure 9

Fmo5 and PPARα co-expression alleviates AFLD-induced apoptosis, inflammatory response and the nuclear translocation of NF-κB p65 in vitro (A) Western blot and RT-PCR assays were used to detect the protein and mRNA levels of PPARα in siRNA targeting PPARα transfected L02 cell or vector expressing PPARα transfected L02 cells. (B) CCK-8 assay was used to detect the cell viability in L02 cells among six groups. ELISA assay was used to detect TG level (C), AST level (D), ALT level (E), IL-6 level (F), and TNF-α level (G) in L02 cells among six groups. (J) Immunofluorescence staining was used to observe the nuclear translocation of NF-κB p65 in L02 cells among six groups. Flow cytometry assay was used to detect cell apoptosis (I) and ROS level (H) in L02 cells among the six groups. Data are presented as the mean ± standard deviation. ** P < 0.01 vs. NC group, ## P < 0.01 vs. siNC group, a P < 0.01 vs. Con group, b P < 0.01 vs. AFLD group, c P < 0.01 vs. AFLD+siFmo5 group, d P < 0.01 vs. AFLD+siFmo5+ PPARα group, and e P < 0.01 vs. AFLD+Fmo5+ siPPARα group,
Figure 10

Fmo5 and PPARα co-expression alleviates AFLD-induced apoptosis and inflammatory response via inhibiting the activation of NF-κB signaling pathway in vitro. (A) Western blot assay was used to detect the protein levels of Fmo5, PPARα, phospho-NF-κB p65 and NF-κB p65 in CHS-828 treated L02 cells. After CHS-828 pre-treated with L02 cells with AFLD for 4 h, siRNA-Fmo5 and siRNA-PPARα were co-transfected
into cells, the levels of TG, AST, ALT, IL-6, and TNF-α level (C) were detected by ELISA assay, cell apoptosis (D), and ROS level (E) were detected by Flow cytometry assay. Data are presented as the mean ± standard deviation. ** P < 0.01 vs. Con group, ## P < 0.01 vs. AFLD group, and ^^ P < 0.01 vs. AFLD+siFmo5+ siPPARα group.