Overexpression of HNF4α enhances umbilical cord mesenchymal stem cell function to treat acute liver failure in mice through macrophage polarization

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Research

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Abstract

Background and aim: Advances in research on stem cell therapy provide new feasible solutions for acute liver failure (ALF) treatment. Recent studies have demonstrated that the expression of hepatocyte nuclear factor 4α (HNF4α) reset within the damaged hepatocytes can restore normal physiological function. This study aimed to determine the role of human umbilical cord mesenchymal stem cells (HuMSCs) overexpressing HNF4α in ALF treatment.

Method: HuMSCs in vitro were isolated, cultured, and reprogrammed by the lentiviral expression of HNF4α (hereinafter referred to as HuMSC-HNF4α). HuMSC-HNF4α was intraperitoneally administered into the mice immediately after exposed to D-galactosamine/lipopolysaccharide (D-galN/LPS). The liver histological and serum biochemical analyses were performed to investigate the effects of HuMSC-HNF4α in ALF. Macrophages differentiation and cytokines secretion by HuMSCs were evaluated to elucidate the underlying mechanisms.

Results: HuMSC-HNF4α had more obvious therapeutic effects on ALF compared with negative control virus-transfected the HuMSCs (HuMSC-CON). In vitro, HuMSC-HNF4α promoted the polarization of liver macrophages (Kupffer cells) to the M2 phenotype, inhibited the inflammatory response of macrophages, and reduced the levels of inflammatory factors such as TNF-α and IL-1β to reduce liver damage.

Conclusions: This study confirmed that the therapeutic effect of HuMSC-HNF4α on ALF was not the same as the previous passive support but as active intervention on excessive inflammation in the body. The findings provided new ideas for research and clinical practice in the future.

1. Introduction

Acute liver failure (ALF) is a clinical syndrome characterized by acute and severe liver damage, caused by various factors such as viral infection, toxic drugs, metabolic diseases, and alcohol-induced damage [1]. Orthotopic liver transplantation (OLT) is one of the most effective ways to treat ALF. However, this therapy is limited by the rarity of donor livers and the poor systemic condition of patients before surgery due to long waiting time [2]. Preliminary studies indicated that stem cells played a key role in treating ALF by trans-differentiating into hepatocytes to perform alternative functions, including promoting liver regeneration, reducing hepatocytes apoptosis, inhibiting liver fibrosis, and regulating the immune and inflammatory response [3-6]. Based on importance of immune responses in ALF, the immune regulation of stem cells may be the most important factor affecting the treatment of ALF [7].

Stem cells, especially mesenchymal stem cells (MSCs), are excellent genetic carriers due to their accessibility, strong differentiation potential, and proliferative capacity. Previous studies have confirmed that, the overexpression of thioredoxin-1 (Trx-1), insulin like growth factor 1 (IGF-1), hepatocyte growth factor (HGF) and other genes enhanced the protective effect of MSCs on ALF through reverse transcription and other technologies [8-10]. MSCs with the high expression of CXC chemokine receptor
type 4 (CXCR4) could improve the quality of life and prolong the survival time of patients with ALF caused by small-for-size syndrome after liver transplantation.

Hepatocyte nuclear factor 4α (HNF4α) is a nuclear transcription factor previously shown to be lowly expressed in liver diseases such as liver tumors and ALF. Also, the physiological functions of hepatocytes can be restored by resetting the normal expression of HNF4α [11, 12]. Further, the overexpression of HNF4α in MSCs though gene transfection technology promoted the differentiation of MSCs into hepatocytes [13-15]. HNF4α is closely related to ALF and has strong regulatory effects on functions of MSCs, suggesting that MSCs with the high expression of HNF4α are expected to possess potential curative effects on ALF.

Macrophages in the liver (Kupffer cells) can differentiate into M1 or M2 macrophages in response to various local microenvironments [16]. M1 macrophages mediate inflammatory damage and promote inflammation. Conversely, the M2 phenotype, which is characterized by anti-inflammatory factors, such as IL-4 and interleukin-10 (IL-10), and protective proteins (TGF-β and VEGF), inhibits inflammation and repair tissue damage [17, 18]. MSCs have been shown to promote M2 macrophage polarization and thus exert an anti-inflammatory effect by secreting IDO, PGE2, and so forth in response to environmental signals including elevated levels of IL-6 [19-21].

In this study, human umbilical cord-derived MSCs (HuMSCs) with overexpression of HNF4α (HuMSC-HNF4α) were first established. Then, their therapeutic effects on lipopolysaccharide/D-galactosamine (LPS/D-GalN)-induced ALF were observed to explore their therapeutic potential for ALF.

2. Materials And Methods

2.1. Isolation and identification of HuMSCs

This study was approved by the Institutional Review Board and Human Ethics Committee of Ren Ji Hospital, School of Medicine, Shanghai Jiao Tong University, Shanghai, China. All umbilical cords used in this study were obtained from healthy full-term cesarean section fetuses at Renji Hospital affiliated to Shanghai Jiao Tong University School of Medicine. Written consent for the use of samples for research purposes was obtained from all patients. HuMSCs were treated as previously described [14]. HuMSCs differentiated into chondrogenic (Cyagen Biosciences, Guangzhou, China), osteogenic (Cyagen Biosciences, Guangzhou, China), and adipogenic lineages (Cyagen Biosciences, Guangzhou, China), as indicated by positive type II collagen (Abcam, Cambridge, MA, USA, Cat. ab34712), Alizarin red, and Oil Red O staining, respectively, after induced with a conditioned medium for 2-3 weeks. The expression patterns of CD73, CD90, CD105, CD34, CD45, major histocompatibility complex, class II, DR (HLA-DR) were detected by flow cytometry to evaluate the phenotypes of HuMSCs.

2.2. Establishment and identification of HuMSC-HNF4α
HuMSCs with stable overexpression of HNF4α, called HuMSC-HNF4α, were constructed as previously described [14]. HuMSCs transfected with a lentiviral vector containing only green fluorescent protein (HuMSC-CON) were treated as controls. The lentiviral transduction efficiency was detected by Western blot and Real-time quantitative reverse transcription–PCR.

### 2.3. Flow cytometry

Flow cytometry (BD Sciences) was performed to characterize the phenotypes of HuMSCs and macrophages. Antibodies against the human antigens CD11b (Cat. #562721), CD68 (Cat. #565594), CD80 (Cat. #561134), CD206 (Cat. #550889), CD73 (Cat. #562430), CD90 (Cat. #563070), CD105 (Cat. #563803), CD34 (Cat. #562383), CD45 (Cat. #563792), and HLA-DR (Cat. #562804) were purchased from BD Sciences, San Diego, CA, USA. Further, antibodies against the mouse antigens CD86 (BD Sciences, San Diego, CA, USA, Cat. #553690) and CD206 (BD Sciences, San Diego, CA, USA, Cat. #565250) were used to observe the effect of in vivo on macrophages phenotype. The data were analyzed using the BD CellQuest Pro software (V1.0.2).

### 2.4. Animal

Wild-type mice (C57BL/6J) were purchased from Shanghai Slark Experimental Animal Co., Ltd., and managed by the Animal Experimental Center of Renji Hospital, Shanghai Jiaotong University (Specific Pathogen Free grade). The experimental procedure was approved by the Animal Ethics Committee of Renji Hospital, Shanghai Jiao Tong University School of Medicine. Further, 2 × 10⁶ HuMSC-HNF4α and HuMSC-CON were added to phosphate-buffered saline (PBS) solution. Mice aged 6 weeks were intraperitoneally injected with HuMSC-HNF4α cells, HuMSC-CON, or PBS (10 mL/kg). D-GalN (700 mg/kg, Sigma-Aldrich, St. Louis, MO, Cat. #G0500) and LPS (10 μg/kg, Sigma-Aldrich, St. Louis, MO, Cat. #L2630) were administered via the abdominal cavity after 24 h of pretreatment. The survival status of mice was observed and recorded 4 h after LPS/D-GalN injection for 48 h. It was recorded every half hour from 4 h to 12 h and every 6 hours between 12 h and 48 h. The mice were anesthetized with 1% sodium 3 h after intraperitoneal injection of LPS/D-GalN. Blood and livers were collected subsequently. The blood samples were centrifuged at 5000 rpm for 15 min to collect plasma. The plasma and liver tissues were frozen immediately in liquid nitrogen and stored at −80°C until analysis.

### 2.5. Real-time quantitative reverse transcription-PCR

Total RNA was isolated from cultured HuMSCs using TRIzol reagent (Invitrogen Inc, CA, USA) and reverse-transcribed into cDNA using a reverse transcription kit (Takara Bio USA, CA, USA). The PCR reaction was carried out in 20 μL of a final volume containing 0.1 μM of each forward and reverse primer, cDNA and 10 μL of SYBR Green PCR Master Mix (Takara Bio USA, CA, USA). DNA was amplified and analyzed using a StepOnePlus real-time PCR machine (Thermo-Fisher Scientific, Inc., Rochford, IL, USA).

### 2.6. Western blot analysis
To determine the expression of HNF4α, total protein was lysed with radio immunoprecipitation assay (RIPA) peptide lysis buffer (Beyotime Biotechnology, Jiangsu, China, Cat. #P0013B) containing 1% protease inhibitors (Thermo Fisher Scientific, Waltham, USA, Cat. # A32953). Western blot analysis were performed according to anti-β-actin (Santa Cruz, CA, USA, Cat. #sc-8432) and anti-HNF4α (Santa Cruz, CA, USA, Cat. #sc-374229).

2.7. Detection of serological indicators

The enzyme activities of aspartate aminotransferase (AST) and alanine transaminase (ALT) in plasma were detected with assay kits (Nanjing Jiancheng Bioengineering Institute, China). Enzyme-linked immunosorbent assay (ELISA) kits (R&D Systems, Minneapolis, MN) were used to evaluate TNF-α and IL-1β levels in plasma. HuMSC-CON and HuMSC-HNF4α were seeded in six-well plates at a density of 5 × 10^5/well. When the cells reached 70% confluence, the remaining supernatant was washed with PBS. Fetal bovine serum (FBS)-free F12 (Gibco, MA, USA) was used to continue culture routinely. After 48 h, the supernatant was collected into 15 mL centrifuge tubes and tested with ELISA kits to assess the levels of various immune factor indicators such as IL-10 and macrophage colony stimulating factor (M-CSF).

2.8. Histopathological examination

The tissue samples were cut into appropriate size and fixed with 4% paraformaldehyde for hematoxylin-eosin staining (H&E) and immunohistochemical staining. Antibodies used for immunohistochemical analysis were used as follows: F4/80 (Abcam, Cambridge, UK, Cat. #ab6640), MPO (Abcam, Cambridge, UK, Cat. #ab208670), TNF-α (Abcam, Cambridge, UK, Cat. #ab9579). Terminal deoxynucleotidyl transferase-mediated dUTP-biotin nick end labeling assay (TUNEL) was performed for apoptosis.

2.9. THP-1 cell culture and differentiation into macrophages

The 1640 complete medium (Gibco, MA, USA) with 1% double antibody and 10% FBS (Gibco, MA, USA) were used. THP-1 cells with ideal growth activity were seeded in six-well plates at a density of 1×10^6/well, and 100ng/mL phorbin 12-myristate 13-acetate (PMA, Sigma, MA, USA, Cat. #P1585) was added to the medium to induce differentiation for 24h. Then, the cells were cultured for 4 days, and the morphology of cells was observed under a light microscope. The induced macrophages were collected, and the surface antigens CD11b and CD68 were detected by flow cytometry.

THP-1 cells were induced to differentiate into macrophages and cultured in six-well plates. A 0.4-μm Transwell chamber (Corning, NY) was placed on a six-well plate. Further, 5×10^5 HuMSC-HNF4α were seeded into the chamber (control group with HuMSC-CON and negative control group without cells) and co-cultured with macrophages in 1640 medium for 24 h to construct co-culture systems. The MTT assay was used to detect the viability of macrophages. The Transwell chamber was discarded after collecting cells of the three groups. The mixed cells were cultured and stimulated in 1640 medium containing LPS (1 μg/mL) for 6 h. The negative control group cultured for another 6 h. The cell culture medium of macrophages in the three groups was collected. Immune factors secreted by macrophages, such as TNF-α,
α and IL-1β, were detected using ELISA kits. The expression patterns of macrophage CD80 and CD206 were detected by flow cytometry.

2.10. Statistical analysis

The data were expressed as mean ± standard deviation (mean ± SD). SPSS 19.0 statistical software was used for data analysis. The comparison of indicators depended on a single-factor analysis of variance. The t-test was used for comparison between groups. A P value < 0.05 indicated a statistically difference.

3. Results

3.1 Identification of HuMSCs and HuMSC-HNF4α

The cells isolated from the umbilical cord were fibroblast-like and grew in the form of fusiform strips adherently under the light microscope (Fig. 1a). The morphology of the cells changed after induction of adipogenesis, osteogenesis, and chondrogenesis. The induced cells were detected using Oil Red O staining, Alizarin red, and type II collagen, and their morphology was found to be in accordance with their differentiation performance (Fig. 1b-1d). Flow cytometry revealed that the cells overexpressed CD73, CD90, and CD105, but expressed CD34, CD45, and HLA-DR at a low level (Fig. 1e). In summary, the cells met the identification criteria of MSCs [22].

HuMSCs were transfected with either a pWIPi-Green Fluorescent Protein (GFP, HuMSC-CON) or pWIPi-HNF4α-GFP (HuMSC-HNF4α) lentiviral vector for 5-7 days. After infection, the ratio of GFP-positive cells revealed that the transfection efficiency was > 95% (Fig. 2a). Western blot analysis and real-time PCR indicated the obvious upregulation of the expression of HNF4α in HuMSC-HNF4α cells (Fig. 2b). In summary, HuMSC-HNF4α and HuMSC-CON were successfully constructed.

3.2 Protective effects of HuMSC-HNF4α in mice with ALF

A survival rate experiment was performed, liver function indexes and the levels of inflammatory factor in serum were detected, and H&E and immunohistochemical staining were performed after LPS/D-GalN injection to determine the impact of HuMSC-HNF4α on survival.

The survival curves indicated that the intraperitoneal injection of both HuMSC-HNF4α and HuMSC-CON could improve the survival of mice with ALF (P < 0.05) (Fig. 3a). However, the effect of the injection of HuMSC-HNF4α was better. In detail, the mortality in the HuMSC-HNF4α and HuMSC-CON groups after 9 h was 44% and 90%, respectively.

The plasma levels of liver enzyme indexes ALT and AST detected in the LPS/D-GalN-induced ALF mouse model were the lowest in the HuMSC-HNF4α group compared with both the HuMSC-CON and PBS groups (P < 0.01) (Fig. 3b).
The levels of pro-inflammatory cytokines in circulation were examined. TNF-α and IL-1β plasma levels were both significantly reduced in the HuMSC-HNF4α and HuMSC-CON groups after ALF. However, the levels of TNF-α and IL-1β in the HuMSC-HNF4α group were lower (Fig. 3c).

H&E staining revealed extensive congestion and necrosis of liver tissues in the PBS group. The liver tissues in the HuMSC-CON group exhibited moderate microscopic deterioration. In contrast, only mild liver injury was seen in the HuMSC-HNF4α group (Fig. 3d). HuMSC-HNF4α alleviated liver damage in mice with ALF. The number of TUNEL-positive cells in the liver tissues significantly increased in the PBS group compared with the other two groups. The mice injected with HuMSC-HNF4α showed the lowest level of apoptosis (Fig. 4a).

Next, the study investigated the hepatic infiltration of immune cells by immunohistochemical staining of MPO (a marker of neutrophils), F4/80 (a marker of macrophages), and TNF-α (a marker of liver cell death) [23-25]. Liver sections from the HuMSC-HNF4α group contained fewer F4/80-positive cells (macrophages and Kupffer cells) compared with the HuMSC-CON group. A large number of macrophages infiltrated into liver tissues of mice injected with PBS (Fig. 4b). TNF-α, secreted by activated macrophages, is the most important inflammatory factor that causes hepatocyte apoptosis and necrosis during the pathogenesis of ALF [23]. The level of TNF-α in liver tissues of mice injected with HuMSC-HNF4α was lower than that in the HuMSC-CON group (Fig. 4b). No significant difference in neutrophil infiltration in liver tissues was observed among these three groups (Fig. 4b). Taken together, these results indicated that HuMSC-HNF4α mainly acted on macrophages. Experiments were subsequently designed to explore specific mechanism of action of macrophages action in vitro.

### 3.3 HuMSC-HNF4α inhibited LPS-stimulated macrophage inflammation and promoted macrophages to polarize to M2 states

Circular and suspended THP-1 cells were cultured with PMA (100ng/ml) for 24 h; the cells grew adherently and lost the ability to proliferate subsequently. After incubation for 4 days, the cells became larger and developed extended amoeba-like pseudopodia (Fig. 5a). The expression of human macrophage markers CD11b and CD68 was significantly higher in differentiated cells compared with undifferentiated cells (Fig. 5b). In summary, the cells met the criteria of experimental macrophages [26].

Macrophages co-cultured with HuMSC-HNF4α or HuMSC-CON were stimulated with LPS for 6 h. The cultured supernatant was collected, and the levels of TNF-α and IL-1β were determined. Macrophages released a large number of inflammatory factors, such as TNF-α and IL-1β, after stimulation. Either HuMSC-HNF4α or HuMSC-CON decreased the levels of inflammatory factors secreted by macrophages. HuMSC-HNF4α inhibited the release of TNF-α more significantly ($P < 0.01$). No difference was found in the effect of inhibiting the release of IL-1β between HuMSC-HNF4α and HuMSC-CON (Fig. 5c).

Macrophages polarized towards M1 phenotype or M2 phenotype according to the local microenvironment, which played a key role in the development of ALF. M1 macrophages were characterized by the overexpression of CD80, CD86, and so forth. M2 phenotype overexpressed CD206,
CD369 (Dectin-1), and so forth [27, 28]. The expression of CD80 and CD206 was examined by flow cytometry after collecting macrophages co-cultured with HuMSC-HNF4α or HuMSC-CON cells stimulated with LPS for 6 h and macrophages without LPS (CTR). The expression of CD80 in macrophages co-cultured with HuMSC-HNF4α decreased. In contrast, the expression of CD206 increased (Fig. 5d). Overall, these results indicated that HuMSC-HNF4α promoted M2 polarization of macrophages, which was consistent with previous conclusions that HuMSC-HNF4α reduced the level of TNF-α secreted by macrophages.

3.4 Overexpression of HNF4α up-regulated the level of IL-10 and M-CSF secreted by HuMSCs

The supernatants in the HuMSC-HNF4α and HuMSC-CON groups were collected for protein chip analysis to determine the mechanism of HuMSC-HNF4α in regulating macrophage polarization (Fig. 6a). The results showed that the overexpression of HNF4α changed the levels of IL-10 and M-CSF. The protein-chip assay was performed to verify the results, and the same conclusion was obtained (Fig. 6b). The mice were pretreated with HuMSC-CON and HuMSC-HNF4α for 24 h as previously described, and an ALF model was constructed to elucidate further how HuMSC-HNF4α protected mice from ALF. The liver, spleen, lung, and small intestine of mice in these two groups were collected to detect GFP fluorescence. The results indicated no distribution of HuMSCs in these organs (Fig. 6c). The difference in serum IL-10 and M-CSF cells suggested that HuMSC-H NF4α cells reduced ALF in mice via the release of IL-10 and M-CSF, independent of their locations in the liver (Fig. 6d). The percentages of M1 and M2 phenotypes of macrophages in the liver after cell input therapy were observed by flow cytometry to determine whether the overexpression of HNF4α in HuMSCs could affect macrophage polarization in vivo (Fig. 7a). HuMSC-HNF4α promoted macrophages to polarize to M2 states in vivo. In addition, recent studies confirmed that IL-10 and M-CSF promoted the polarization of liver macrophages (Kupffer cells) toward the M2 phenotype and play a role in inhibiting inflammation and repairing damage [29, 30]. It was inferred that HNF4α might enhance the immunoregulatory function of HuMSCs by increasing the secretion of IL-10 and M-CSF.

4. Discussion

The use of genetically modified cells, with more obvious therapeutic effects, has replaced the use of existing cells to treat diseases with the rapid development of genetic engineering technology. Hence, attempts were made to use gene-regulated MSC transplantation for treating ALF and hence enhance the therapeutic effect of MSCs.

Previous studies showed that the overexpression of HNF4α in HuMSCs promoted HuMSC differentiation into hepatocytes [10, 14]. They also demonstrated that HNF4α, with strong regulatory effects on various functions of MSCs, was closely related to ALF. Thus, it was suggested that HuMSCs with the overexpression of HNF4 might possess amazing therapeutic effects on ALF.

Therefore, a series of experiments were performed to verify this hypothesis. HuMSC-HNF4α effectively improved the survival of mice with ALF, suppressed the immune responses, reduced liver damage, and
reduced necrosis and apoptosis of liver tissues in vivo. The therapeutic effect of HuMSC-HNF4α cells on ALF was to inhibit the inflammatory response of liver macrophages, reduce the infiltration of macrophages in liver tissues, and enhance the release of inflammatory factors.

Based on the importance of macrophages in the pathogenesis of ALF, an experiment was designed in this study to co-culture HuMSC-HNF4α and macrophages so as to explore the effect of HuMSC-HNF4α on the inflammatory response of macrophages in vitro. HuMSC-HNF4α inhibited the inflammatory responses caused by LPS stimulation and reduced the level of TNF-α secreted by macrophages. In vivo, experiments were designed to testify that HuMSC-HNF4α could promote macrophage polarization to the M2 phenotype by increasing the secretion of IL-10 and M-CSF, thereby reducing liver injury in mice.

HuMSC-HNF4α promoted macrophage polarization toward the M2 phenotype, suggesting that HuMSC-HNF4α changed the microenvironment of the culture medium. The culture supernatants of HuMSC-HNF4α and HuMSC-CON were collected for protein chip analysis to examine the distinguishing factors so as to find the factors secreted by HuMSC-HNF4α influencing the polarization of macrophages. More than 40 factors with significance were found (P< 0.05). The factors related to the inflammatory response of macrophage were examined to confirm that the levels of IL-10 and M-CSF secreted by HuMSC-HNF4α were significantly higher than the levels of those secreted by HuMSC-CON. IL-10 and M-CSF activated the polarization of macrophages toward the M2 phenotype, inhibited inflammation, and repaired damage [29, 30]. Thus, it was speculated that HNF4α enhanced the immune-regulatory function of HuMSCs by increasing the secretion of immune factors such as IL-10 and M-CSF.

Recent studies confirmed that immunoinflammatory responses were important in ALF. Hence, the immunomodulatory function of MSCs has become the latest focus. However, several studies have shown that hepatic macrophages, as the key cells in inflammation regulation, are central in the pathogenesis of liver injury and have been proposed as the potential targets in combatting fibrosis [31]. The last few years have revealed several key regulators of macrophage polarization, including the signal transducer and activator of transcription family of molecules such as macrophage chemoattractant CCL2 [32], peroxisome proliferator-activated receptor gamma [33], CCAAT enhancer-binding proteins (C/EBP) family, and interferon regulatory factors [27]. Recent studies have also suggested that microRNAs and long noncoding RNAs influence macrophage polarization [34]. The present study reported another regulatory mechanism of macrophage polarization. MSCs exerted immunoregulatory functions on lymphocytes, NK T cells, DC cells, and other immune effector cells in ALF caused by different etiologies. Similarly, the study also found that HuMSC-HNF4α had a stronger anti-inflammatory effect on macrophages in ALF compared with HuMSCs used alone. This finding indicated that the effects of both MSCs and HNF4α were complicated. An unknown mechanism might exist, which needs exploration.

HuMSC-HNF4α with a powerful immunomodulatory effect in the pathogenesis of ALF is expected to form a mixed system with immortalized hepatocytes to jointly transplant or construct a hybrid biological artificial liver. This process combines functional support and immune regulation to treat ALF more effectively and regarded as a future research direction. However, this study also had certain limitations.
After the protein-chip assay, the mechanism of HNF4α gene regulating the secretion of immune factors by MSCs needs to be further explored in vivo. In addition, database analysis was performed to explore the upstream transcription factors related to the differential gene expression according to the sequencing in ChIP-X Enrichment Analysis Version 3 (ChEA3). The bar chart revealed the relevance between HNF4α and the differential gene including IL-10 and M-CSF (Fig. S1). Of course, the possible mechanism needs further verification.

5. Conclusion

The present study confirmed the therapeutic effects of HuMSC-HNF4α in ALF in vivo and in vitro. It revealed that HuMSC-HNF4α cells promoted macrophages to polarize towards the M2 phenotype by secreting IL-10 and M-CSF. HuMSC-HNF4α suppressed inflammation and repaired damage, which actively interfered with excessive inflammatory reactions in the body instead of passive support. Meanwhile, the findings provided new ideas for scientific research and clinic practice.

Abbreviations

ALF, Acute liver failure; D-GalN, D-galactosamine; HNF4α, hepatocyte nuclear factor 4α; HuMSC-HNF4α, HuMSCs overexpressing HNF4α; HuMSCs, human umbilical cord mesenchymal stem cells; HuMSC-CON, HuMSCs without HNF4α; LPS, lipopolysaccharide; MSCs, mesenchymal stem cells; IL-10, interleukin-10; M-CSF, macrophage colony stimulating factor; AST, aspartate aminotransferase; ALT, alanine transaminase

Declarations

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Not applicable.

Authors’ contributions

YPY, QQZ, QX, and HHL were involved in conception and design of the study. NW and QQZ performed the in vivo experiment. YPY and QQZ performed the in vitro experiments. JC and LX performed the data analysis and interpretation. YPY, JJZ, and HLH wrote the manuscript. NW, QX, JZ, JJZ, and HLH were involved in review and editing of the manuscript. All authors commented the final version. All authors had full access to all data. All authors read and approved the final manuscript.

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Availability of data and materials

All data and materials used in this work are available on request.

Ethics approval and consent to participate

This study was approved by the Institutional Review Board and Human Ethics Committee of Ren Ji Hospital, School of Medicine, Shanghai Jiao Tong University, Shanghai, China.

Consent for publication

All authors consent to publish this manuscript.

Competing interests

The authors declare that they have no competing interests.

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