Therapeutic effect of hepatocyte growth factor-secreting mesenchymal stem cells in a rat model of liver fibrosis

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Bone marrow-derived mesenchymal stromal cells (MSCs) have been reported to be beneficial for the treatment of liver fibrosis. Here, we investigated the use of genetically engineered MSCs that overexpress hepatocyte growth factor (HGF) as a means to improve their therapeutic effect in liver fibrosis. Liver fibrosis was induced by intraperitoneal injection of dimethylnitrosamine. HGF-secreting MSCs (MSCs/HGF) were prepared by transducing MSCs with an adenovirus carrying HGF-encoding cDNA. MSCs or MSCs/HGF were injected directly into the spleen of fibrotic rats. Tissue fibrosis was assessed by histological analysis 12 days after stem cell injection. Although treatment with MSCs reduced fibrosis, treatment with MSCs/HGF produced a more significant reduction and was associated with elevated HGF levels in the portal vein. Collagen levels in the liver extract were decreased after MSC/HGF therapy, suggesting recovery from fibrosis. Furthermore, liver function was improved in animals receiving MSCs/HGF, indicating that MSC/HGF therapy resulted not only in reduction of liver fibrosis but also in improvement of hepatocyte function. Assessment of cell and biochemical parameters revealed that mRNA levels of the fibrogenic cytokines PDGF-bb and TGF-β1 were significantly decreased after MSC/HGF therapy. Subsequent to the decrease in collagen, expression of matrix metalloprotease-9 (MMP-9), MMP-13, MMP-14 and urokinase-type plasminogen activator was augmented following MSC/HGF, whereas tissue inhibitor of metalloprotease-1 (TIMP-1) expression was reduced. In conclusion, therapy with MSCs/HGF resulted in an improved therapeutic effect compared with MSCs alone, probably because of the anti-fibrotic activity of HGF. Thus, MSC/HGF represents a promising approach toward a cell therapy for liver fibrosis.

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INTRODUCTION

Chronic liver damage leads to liver fibrosis, which is characterized by the hyper-accumulation of extracellular matrix (ECM) proteins including collagen,¹ which produces a fibrous scar that distorts the hepatic architecture and causes liver cell dysfunction. Currently, there is no effective therapy for liver fibrosis, and liver transplantation, which is a major surgery and is limited by an inadequate supply of transplantable livers, is the only option for end-stage liver failure.

Mesenchymal stromal cells (MSCs) were recently found to provide effective therapy in animal models of liver fibrosis and cirrhosis. Sakaida et al.² and Fang et al.³ showed that transplantation of bone marrow cells or systemic infusion of FLK1(+) MSCs, respectively, reduced carbon tetrachloride (CCL4)-induced liver fibrosis in mice. A therapeutic effect of transplanting hepatocyte growth factor (HGF)-treated bone marrow MSCs was also documented in CCL4-injured rats.⁴ MSCs were shown to differentiate into hepatocytes in vitro⁵⁻⁷ and in vivo.⁸ Collectively, these results establish a therapeutic effect of MSCs in liver fibrosis. However, the therapeutic efficacy of MSC transplantation was not significant in some reports. Carvalho et al.⁹ reported that bone marrow-derived MSCs were unable to reduce liver fibrosis, whereas treatment with MSCs/HGF resulted in an improved therapeutic effect compared with MSCs alone, probably because of the anti-fibrotic activity of HGF. Thus, MSC/HGF represents a promising approach toward a cell therapy for liver fibrosis.
fibrosis or improve liver function in a rat model of severe chronic liver injury, suggesting that the therapeutic efficacy of MSCs needed to be improved.

The mode of therapeutic action of MSCs in liver disease was postulated to include differentiation into hepatocytes or fusion with endogenous hepatocytes. However, considering their low frequency of transdifferentiation or fusion, as well as the low numbers of transplanted MSCs, it is unlikely that MSC-derived hepatocytes could provide sufficient replacement of lost liver function. Moreover, MSCs do not only differentiate into hepatocytes or endothelial cells; a large fraction differentiates into fibrogenic myofibroblasts, leading to adverse effects.

In addition to their differentiation potential, MSCs have the ability to supply beneficial cytokines. MSC-derived interleukin-10 and tumor necrosis factor-α inhibit the proliferation of Ito cells, and MSC-derived HGF was shown to induce the apoptosis of Ito cells in a co-culture system. Cytokines are produced in ample quantities by MSCs, thus supporting the view that cytokine production may be the major mode of therapeutic action of MSCs in liver fibrosis. This insight led us to postulate that augmentation of the production of therapeutic cytokines by MSCs would enhance their therapeutic efficacy in liver fibrosis.

HGF is a well-known anti-fibrotic cytokine, and delivery of the HGF gene or protein attenuates liver fibrosis in numerous in vivo models. The anti-fibrotic effect of HGF is thought to be achieved through attenuation of fibrogenic cytokine expression (transforming growth factor beta 1 (TGF-β1) and platelet-derived growth factor-bb (PDGF-bb)), and through inhibition of the proliferation and activation of Ito cells, the major ECM producer in the liver. In addition, HGF inhibits the cell death of normal hepatocytes. The effectiveness and broad mode of action of this cytokine prompted us to choose HGF for testing the concept of enhancing cytokine production by MSCs used for cell therapy of liver fibrosis. In this report, we investigated the therapeutic efficacy of human MSCs ectopically overexpressing HGF (MSC/HGF) in a rat model of liver fibrosis induced by dimethylnitrosamine (DMN) injection.

MATERIALS AND METHODS

Cell culture and gene transduction

Human MSCs were isolated from bone marrow aspirate and expanded in culture as described previously. The adenoviral vector encoding human HGF (Ad-HGF) was kindly provided by Dr SJ Kim (ViroMed, Seoul, Korea). For adenoviral transduction, MSCs were exposed to fresh medium containing Ad-HGF (multiplicity of infection = 200) for 1 h. The medium was then removed, and the cells were washed once with DMEM and re-cultured in normal medium for 24 h, after which cell transplantation was performed. Aliquots of transduced MSCs were maintained in culture for the determination of HGF production by transduced MSCs. The medium was sampled at 2, 4, 6, 8 and 10 days after transduction and analyzed for HGF using a human HGF-specific ELISA kit (Institute of Immunology, Tokyo, Japan).

Liver fibrosis animal model and cell transplantation

Five-week-old, specific-pathogen-free, male Sprague-Dawley (SD) rats were obtained from Central Laboratory Animal Inc. (Seoul, Korea) and maintained in an air-conditioned animal house (22°C, 55% humidity, and 12:12 h daylight/darkness cycles) in accordance with institutional guidelines. To induce liver fibrosis, rats (n = 65) were conditioned by intraperitoneal injection with 1% (V/V) DMN (Sigma-Aldrich, St. Louis, MO, USA) diluted in saline at a dose of 1 ml per kg of body weight, on 3 consecutive days each week for 4 consecutive weeks; control rats (n = 10) were mock-conditioned by injection with saline. Liver biopsies were performed at 4 days after the last DMN administration for collagen determination. Collagen was determined by Sirius Red assay using Sircol Collagen Assay DYE reagent (0.1% Sirius Red dye solution; Biocolor Ltd., Newtownabbey, UK) according to the manufacturer’s guidelines. Briefly, extracts of biopsy samples were prepared by sonication in 0.5 M acetic acid, and the extracts were incubated with Sirius Red for 30 min. The absorbance of Sirius Red dye bound to collagen was measured at 550 nm to provide a quantitative indicator of the amount of collagen in the liver. Rats with a collagen level within one standard deviation from the average value were randomly divided into three groups and subjected to cell transplantation. DMN-conditioned rats were injected with saline or with MSCs or MSCs/HGF (1 × 10⁶ cells per rat). Saline-conditioned rats were injected with saline. The final groups were as follows (conditioning + therapeutic treatment): saline + saline, n = 7; DMN + saline, n = 10; DMN + MSCs, n = 10; DMN + MSC/HGF, n = 11.

Analysis of liver tissue specimens and blood samples

To directly compare liver biopsy samples with liver tissue obtained at killing, rats were numbered before biopsy and tracked individually until killing. Rats were killed 12 days after transplantation. For serum biochemical analysis, blood samples were collected from the portal vein and heart. Immediately after blood collection, alkaline phosphatase (ALP), alanine transaminase, aspartate transaminase, total bilirubin, direct bilirubin, albumin and γ-glutamyl transpeptidase concentrations were measured using a Toshiba 200FR automatic chemistry analyzer (Toshiba, Tokyo, Japan).

Liver tissue was excised from a distal site of the left lobe for Sirius red, reverse transcription PCR (RT–PCR) and western blot analysis. The remaining tissue was fixed and processed for histological and immunohistochemical analysis.

Histological and immunohistochemical analysis

Liver tissues were removed, post-fixed in 4% PFA overnight, and then embedded in paraffin. For determination of liver fibrosis, 5-μm-thick paraffin sections were deparaffinized and stained with Picro-Sirius red. The ratio of the Sirius red-positive area to the total area was examined using the ImageJ 1.3s software (National Institutes of Health, Bethesda, MD).

For detection of transplanted cells and HGF expression, deparaffinized liver sections were boiled in citrate buffer (pH 6.0) for 10 min. After blocking in 3% normal serum, the sections were incubated with anti-human HGF antibody (R&D Systems, Minneapolis, MN, USA) and human mitochondria-specific antibody (hMt; Chemicon, Temecula, CA, USA) at 4°C overnight. Sections were then incubated with AlexaFluor 488– or 594-conjugated anti-IgG secondary antibodies (Life Technologies, Grand Island, NY, USA) and counterstained with bis-benzamide (Molecular Probes, Eugene, OR, USA) to visualize cell nuclei.
nuclei. To detect Ito cells, sections were immunostained with anti-desmin antibody (DAKO, Tokyo, Japan) or anti-α-smooth muscle actin antibody (α-SMA; DAKO) and visualized using an ABC kit (Vector Laboratories, Burlingame, CA, USA) according to the manufacturer’s instructions. Immunostained images were acquired using a Zeiss Axiophot microscope (Carl Zeiss, Jena, Germany).

Hydroxyproline content assay
The hydroxyproline content of the liver tissue was determined as described previously.23 Briefly, the samples were homogenized with a Polytron homogenizer (Kinematica Co., Luzern, Switzerland), followed by hydrolysis in 6 N HCl at 120 °C for 3 h, and then dried in a rotary evaporator. Hydrolyzed samples were resuspended in buffer (3.325% (W/V) citric acid 1H₂O, 0.08% (V/V) glacial acetic acid, 8% (W/V) sodium acetate 3H₂O, and 2.275% (W/V) NaOH, 20% (V/V) 1-propanol) and then incubated with chloramine T solution (50 μM) for 20 min. The samples were then subjected to reaction with aldehyde-perchloric acid reagent (15% (W/V) p-dimethylaminobenzaldehyde, 60% (V/V) n-propanol, and 26% (V/V) perchloric acid) at 60 °C for 15 min. The absorbance of each sample was read at 550 nm.

Western blotting
Liver tissues were lysed in RIPA buffer (20 mM Tris, pH 7.4; 150 mM NaCl; 1% Triton X-100; and 0.1% SDS) containing protease inhibitor (10 μg ml⁻¹ leupeptin, 10 μg ml⁻¹ aprotinin and 0.2 mM EDTA). Each liver lysate (50 μg per sample) was resolved by 8 or 10% SDS-PAGE and the proteins were transferred onto a nitrocellulose membrane. Immunoblotting was performed with antibodies against α-SMA (DAKO), desmin (DAKO), matrix metalloprotease-9 (MMP-9) (Biodesign, Kennebunk, ME, USA), or urokinase-type plasminogen activator (uPA) (Santa Cruz, Delaware Avenue, CA, USA); proteins were detected by enhanced chemical luminescence after incubation with appropriate horseradish peroxidase-conjugated secondary antibodies (Amersham, Arlington Height, IL, USA). The relative signal intensity of each lane was calculated using a GS-800 Calibrated Densitometer (Bio-Rad Laboratories, Hercules, CA, USA).

Reverse transcription-PCR
Total RNA from each sample was extracted using the Trizol Reagent (Life Technologies), and 3 μg of RNA was used as the template for cDNA synthesis using AMV reverse transcriptase (TaKaRa Bio Inc., Shiga, Japan), according to the manufacturer’s instructions. PCR amplification was subsequently performed using specific primers (summarized in Supplementary Table 1). PCR products were separated by 1.5% agarose gel electrophoresis, and the relative signal intensity of each lane was calculated using a GS-800 Calibrated Densitometer (Bio-Rad, Hercules, CA, USA). All PCRs were normalized to expression levels of GAPDH used as an internal standard.
Statistical analysis
Statistical analysis was performed using the SPSS 12.0.1. software (SPSS Inc., Chicago, IL, USA). All results are expressed as the mean ± s.d. Comparisons between groups were performed using PROC ANOVA for one-way analysis and Scheffe’s multiple-comparison procedure. \( p < 0.05 \) was considered to be significant.

RESULTS
Quantification and functional assay of secreted HGF in vitro
The amount of HGF produced by MSCs transduced with Ad-HGF was assessed by ELISA and was \( \sim 160 \) times higher than the amount endogenously secreted by MSCs (557.86 vs 3.49 ng ml\(^{-1}\), from \( 5 \times 10^5 \) cells cultured for 4 days, Figure 1a). HGF production peaked at 2–4 days after transduction and was maintained for at least 10 days (Figure 1b). The biological activity of HGF produced by MSCs/HGF was confirmed by phosphorylation of c-Met, the receptor for HGF (Figure 1c), and by scattering of MDCK\(_2\) cells (data not shown), a characteristic function of HGF in these cells.\(^{24}\)

Enhanced resolution of liver fibrosis by transplantation of MSCs/HGF
We adopted the DMN-induced rat liver fibrosis model in this study rather than the CCl\(_4\)-induced model because CCl\(_4\)-induced liver fibrosis resolves spontaneously.\(^{25}\) To address the therapeutic effect rather than the preventive effect, we injected saline, MSCs, or MSCs/HGF into the spleens of the rats once on the ninth day after the final DMN injection (4 days after biopsy; Figure 2a). The average collagen content (Figure 2c) and average body weight (Figure 2e) of all DMN-treated animals were similar at the time of cell transplantation, indicating that the extent of fibrosis was not different before transplantation. The rats were killed at 12 days after cell transplantation, and fibrosis

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**Figure 2** Assessment of liver fibrosis in MSC/HGF-transplanted rats and controls. (a) Diagram of the treatment protocol. (b) Extracellular deposition of collagen fibers stained with Sirius Red. Scale bar = 0.5 mm. (c) Quantification of collagen by image analysis. (d, e) Collagen content and body weight quantified before (white bar) and 12 days after transplantation (black bar). (f) Hydroxyproline content. Normal rats (N), control fibrotic animals (S), fibrotic animals transplanted with MSCs (M) or MSCs/HGF (H). Data represent mean ± s.d. for each group (*\( p < 0.05 \), **\( p < 0.01 \), ***\( p < 0.005 \)).
was evaluated. Liver fibrosis was evident based on the formation of fibrotic septa joining the central area (Figure 2b, right upper panel). Quantitative measurement of collagen in liver tissue by Sirius red assay also revealed increased collagen content (Figure 2d, normal, 0.506 ± 0.019 vs saline/DMN, 0.748 ± 0.018, P < 0.001), which was consistent with the histological changes. The increased collagen content of the rat livers was sustained during the entire experimental period after completion of DMN conditioning, indicating that the fibrosis in this experimental system was stable (Figure 2d, liver biopsy, 0.748 ± 0.018 vs killing, 0.751 ± 0.035 in the saline/DMN group).

The collagen-stained area in liver sections was significantly decreased by MSC transplantation and was further reduced by MSC/HGF transplantation (Figure 2c, saline, 19.2 ± 4.90% vs MSCs, 13.14 ± 3.88% vs MSCs/HGF, 9.0 ± 3.09%; saline vs MSCs P < 0.05; saline vs MSCs/HGF P < 0.001; MSCs vs MSCs/HGF P < 0.05). Tissue hydroxyproline levels, another indicator of tissue collagen content, showed a similar pattern (Figure 2f, saline, 218.7 ± 26.8 μg/g, MSCs, 134.2 ± 24.3 μg/g; MSCs/HGF, 105.2 ± 12.6 μg/g; saline vs MSCs P < 0.001, MSCs vs MSCs/HGF P < 0.05). Surprisingly, when we compared the tissues at the time of biopsy with tissues at the time of killing, the amount of collagen was decreased in the MSC group and in the MSC/HGF group, whereas the amount of collagen in the control and saline/DMN groups did not change (Figures 2c and d), suggesting that resolution of fibrosis was occurring. We also evaluated changes in body weight as a parameter reflecting the general condition of the rats. Body weight was significantly increased only in the MSC/HGF group (P < 0.05, Figure 2e). These data clearly demonstrate that MSCs/HGF were significantly more effective than MSCs alone as a therapeutic for rat liver fibrosis.

Functional recovery from DMN-induced liver damage by transplantation of MSCs/HGF

We next evaluated the effects of cell transplantation on the extent of liver injury and liver function. Serum transaminase levels, which increase when the liver is damaged, were significantly attenuated in the MSC/HGF group (Figures 3a and b), as were ALP levels, which increase when biliary epithelial cells are damaged (Figure 3c), compared with the MSC group and saline control group, demonstrating that MSCs/HGF were more effective in preventing liver damage associated with fibrosis. Similarly, the serum ammonia level, which is an indicator of urea cycle function in hepatocytes, was normalized in the MSC/HGF group to a greater extent than in the MSC group (Figure 3d). In addition, the serum albumin level, which is an indicator of liver function, was slightly increased in the MSC/HGF group but not in the MSC or saline control group (P < 0.05; Figure 3e). Taken together, our data demonstrate that MSCs/HGF are more beneficial than MSCs alone with respect to the attenuation of liver damage and recovery of liver function.

Detection of transplanted cells and determination of human HGF levels in vivo

Our model system was based on the xenotransplantation of human MSCs into rats in the absence of immunosuppression. Thus, we verified the presence of transplanted human cells and the production of human HGF at the time of killing. Because we injected the cells into the spleen, we assumed that the transplanted cells would act directly in the liver after migration or would act indirectly by secreting HGF into the bloodstream, which would eventually reach the liver via the portal vein. The human HGF level in the portal vein was measured by human HGF-specific ELISA. As shown in Figure 4a, human HGF was detected in eight of nine rats of

![Figure 3](image-url) Examination of serum parameters. (a, b) Transaminases (ALT and AST), (c) alkaline phosphatase (ALP), (d) ammonia and (e) albumin. Normal rats, N; fibrotic control animals, S; fibrotic animals transplanted with MSCs, M, or MSCs/HGF, H. Data represent the mean ± s.d. for each group (*P < 0.05, **P < 0.01, ***P < 0.005).
the MSC/HGF group, and in only one of seven rats in the MSC group. In addition, the average level of HGF in the portal vein was significantly higher in the MSC/HGF group than in the MSC group (MSC, 0.01 ± 0.018 ng ml⁻¹ vs MSC/HGF, 0.19 ± 0.154 ng ml⁻¹), indicating that a considerable amount of human HGF produced by cells transplanted into the spleen was delivered to the liver during the experimental period.

We next examined the presence of transplanted cells in the liver and assessed their function in terms of HGF production. Expression of human HGF mRNA was detected by semiquantitative RT-PCR in the liver tissue of six of nine rats transplanted with MSCs/HGF (Figure 4b). In contrast, human HGF mRNA was detected in only one of eight rats transplanted with MSCs. This difference can likely be attributed to the difference in HGF expression level between MSCs and MSCs/HGF. Therefore, the presence of the transplanted cells in the liver tissue was directly evaluated by immunohistochemistry using antibodies against human mitochondria-specific antigen or human HGF (Figure 4c). Although the background fluorescence signal was somewhat high, we could clearly recognize double-positive cells (human mitochondrial antigen and human HGF) in tissue sections from both MSC and MSC/HGF groups. The intensity of staining for human HGF was clearly higher in the MSC/HGF group than in the MSC group. It was remarkable that the xenotransplanted human MSCs...
survived for 12 days and produced human HGF without immunosuppression.

Decreased expression of profibrogenic cytokines and inactivation of Ito cells by MSCs/HGF

We next tried to elucidate the mechanistic basis for the enhanced therapeutic effect of transplanted MSCs/HGF. Because HGF was reported to downregulate TGF-β1 and PDGF-bb, we assessed the expression of TGF-β1 and PDGF-bb mRNA in the liver (Figure 5a). Although mRNA levels varied among animals, expression of TGF-β1 and PDGF-bb was significantly increased in fibrotic livers, and this effect was dramatically ameliorated by MSCs/HGF but not by MSCs (Figures 5b and c).

HGF has been reported to directly affect the survival and activation of rat Ito cells in vivo. Therefore, we assessed the effect of MSCs/HGF and MSCs on Ito cells by measuring α-SMA, which is a marker of activated Ito cells, and desmin, which is a marker of Ito cells irrespective of their activation state. Immunohistochemistry revealed that α-SMA and desmin expression decreased more in the MSC/HGF group than in the MSC group (Figure 6a). Western blotting of pooled liver lysates from each group further confirmed this observation (Figure 6b). In addition, the calculated expression of α-SMA and desmin protein for each experimental animal was similar to that in pooled samples (Figure 6c). Based on these results, we conclude that MSCs/HGF decreased both the number and the activation of Ito cells to a greater extent than MSCs, most probably as a result of the known inhibitory activity of HGF on the survival and activation of Ito cells.

Effect on ECM remodeling molecules in fibrosis resolution

The decreased collagen content in the liver after MSC/HGF transplantation suggested that there was an active remodeling of ECM components in the liver following MSC/HGF transplantation. Because the amount of ECM depends on the balance between production and degradation, we investigated the expression of Type 1 collagen, MMPs and tissue inhibitor of metalloproteases (TIMPs) in our system. The mRNA level of Type 1 collagen, a representative ECM molecule, was significantly decreased in the MSC/HGF group (Figure 7a).

MMPs and TIMPs are key regulators of ECM remodeling and many ECM glycoproteins, such as fibronectin and laminins, are degraded by MMP-2 and MMP-9, whereas MMP-2 activity is regulated by TIMP-2 and MMP-14 via a
tri-molecular interaction. In the present study, the expression of TIMP-2 and MMP-2 mRNA did not change (Figures 7c and d), whereas the reduction in MMP-14 mRNA caused by DMN administration was reversed only in the MSC/HGF group (Figure 7g). In contrast, MMP-13, which degrades interstitial collagen and is expressed in the recovery phase of liver fibrosis, was upregulated in both MSC and MSC/HGF groups (Figure 7f). TIMP-1 was downregulated only in the MSC/HGF group (Figure 7b). uPA generates plasmin, which then activates MMP-9. Protein and mRNA expression of MMP-9 (Figures 7e, h and i) and protein expression of uPA (Figures 7h and j) were increased in both MSC and MSC/HGF groups.

**DISCUSSION**

In this study, we found that MSCs that overexpress HGF have an enhanced therapeutic effect compared with non-manipulated MSCs in a rat liver fibrosis model. Recent studies have shown that the potential of MSCs to differentiate into liver parenchymal cells is not sufficient to fully resolve liver fibrosis. Furthermore, di Bonzo et al. suggested that MSCs are a two-edged sword in hepatic regenerative medicine, as they can differentiate into either hepatocytes or profibrogenic myofibroblasts. Similarly, Aswa et al. suggested inhibiting migration of bone marrow cells to the liver as a strategy for prevention of biliary fibrosis. Nonetheless, several animal and clinical studies have reported beneficial effects of MSCs in the treatment of liver fibrosis, suggesting modes of action distinct from differentiation into parenchymal cells, such as alteration of environmental conditions in the injured tissue through production of soluble factors. Given that MSCs have been demonstrated to produce a wide array of trophic and growth factors both in vivo and in vitro, a reasonable explanation for the functional benefit derived from MSCs may be that they produce organotrophic factors that protect cells from damage or activate endogenous restorative mechanisms within the injured tissue.

The clinical transplantation of organs and cells is limited practically by the availability of donors. Organs and cells to be transplanted must not only be available in sufficient quantity but must also be immunologically compatible with the recipient. Allotransplantation requires immunosuppression in addition to matching of the donor and the recipient by tissue typing. One noteworthy aspect of the present study was that although this study involved xenotransplantation, we did not use immunosuppression. MSCs are reported to escape immune recognition and to inhibit immune responses through inhibition of the activation and differentiation of immune cells such as antigen-presenting cells, T cells, B cells and natural killer cells. Moreover, HGF is known to induce immunological tolerance and has been reported to suppress T-cell proliferation and dendrite cell antigen presentation, reduces acute and chronic allograft rejection, ameliorates the progression of experimental autoimmune myocarditis, and attenuates allergic airway inflammation. In the current study, the two immune modulatory factors, MSCs and HGF,
appear to have played important roles in maintaining a significant population of transplanted cells during the treatment period in the absence of additional immunosuppression. MSCs/HGF may therefore be suitable for treatment of liver fibrosis irrespective of the immunological phenotype of the patient.

Frequently, chronic diseases, including liver fibrosis, are not diagnosed until the condition has reached an advanced stage. Therefore, therapeutic drugs, rather than preventive drugs, are promising and practical from a clinical perspective. In this experiment, MSCs or MSCs/HGF as therapeutic candidates were transplanted after liver fibrosis had been induced by DMN administration. We adopted the DMN model because it has been found to be relatively stable but may progress after administration of DMN ceases. In the current study, we initiated cell therapy after liver fibrosis was established and stabilized, as indicated by the comparable collagen content in the liver at biopsy and at killing among control animals. Taken together, the results indicate that MSCs/HGF may be a favorable therapeutic modality for treatment of liver fibrosis.

In this study, HGF was delivered by MSCs, but the results were comparable to data from studies in which HGF protein was injected directly. Our data show that MSC/HGF treatment affected most relevant parameters in producing an anti-fibrotic effect. We found several strong correlations among parameters, including, for example, HGF level vs MMP-13 level (R = 0.77). These correlations provide clues regarding possible causal relationships between parameters that need further investigation.

In summary, the present study shows that cell therapy with MSCs/HGF significantly promotes liver function and attenuates liver fibrosis. These effects are superior to treatment with MSCs alone. The enhanced therapeutic effects of MSCs/HGF were accompanied by decreased levels of the fibrogenic cytokines TGF-β1 and PDGFB-bb, which may in turn be responsible for the observed reduction in the number and activation of Ito cells. Transplantation with MSCs/HGF also resulted in a decrease in MMP expression and an increase in TIMP expression. Cell therapy with MSCs/HGF may be a new and effective tool for the treatment of fibrotic liver disease.

CONFLICT OF INTEREST
The authors declare no conflict of interest.

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