Hepatic stellate cells contribute to progenitor cells and liver regeneration

Claus Kordes,1 Iris Sawitza,1 Silke Götze,1 Diran Herebian,2 and Dieter Häussinger1

1Clinic of Gastroenterology, Hepatology and Infectious Diseases, and 2Department of General Pediatrics, Neonatology and Pediatric Cardiology, Heinrich Heine University, Düsseldorf, Germany.

Retinoid-storing hepatic stellate cells (HSCs) have recently been described as a liver-resident mesenchymal stem cell (MSC) population; however, it is not clear whether these cells contribute to liver regeneration or serve as a progenitor cell population with hepatobiliary characteristics. Here, we purified HSCs with retinoid-dependent fluorescence-activated cell sorting from eGFP-expressing rats and transplanted these GFP+ HSCs into wild-type (WT) rats that had undergone partial hepatectomy in the presence of 2-acetylaminoﬂuorene (2AAF) or retrorsine, both of which are injury models that favor stem cell–based liver repair. Transplanted HSCs contributed to liver regeneration in host animals by forming mesenchymal tissue, progenitor cells, hepatocytes, and cholangiocytes and elevated direct bilirubin levels in blood sera of GUNN rats, indicating recovery from the hepatic bilirubin–handling defect in these animals. Transplanted HSCs engrained within the bone marrow (BM) of host animals, and HSC-derived cells were isolated from BM and successfully retransplanted into new hosts with injured liver. Cultured HSCs transiently adopted an expression profile similar to that of progenitor cells during differentiation into bile acid–synthesizing and –transporting hepatocytes, suggesting that stellate cells represent a source of liver progenitor cells. This concept connects seemingly contradictory studies that favor either progenitor cells or MSCs as important players in stem cell–based liver regeneration.

Introduction

Hepatic stellate cells (HSCs) are characterized by exceptionally high amounts of retinoids in their quiescent state. These are mainly stored as retinyl palmitate in membrane-coated lipid vesicles and exhibit characteristic retinoid fluorescence after excitation by UV light. HSCs are primarily known as collagen-producing cells in the liver, which are responsible for fibrogenesis in chronic liver diseases (1). Despite intensive research for decades, HSCs have remained enigmatic (2). For instance, stellate cells express molecular markers of different germ layers, which impede a clear statement about their origin (3). The presence of glial fibrillary acidic protein (GFAP) and many other neuroectodermal proteins in stellate cells (4) led to the view that they may derive from the neural crest. This concept has been questioned by microscopic as well as fate-mapping analysis in the developing liver, which indicated that stellate cells apparently originate from mesenchymal cells located in the subependymal space of the septum transversum (5, 6). This view is supported by the expression of mesodermal marker proteins such as desmin and α-smooth muscle actin (α-SMA or actin α 2/ACTA2), especially in activated HSCs (7).

Until recently, HSCs were mainly studied with regard to their fibrogenic potential in chronic disease, whereas their identity and function in normal liver received little attention. Recent data suggest that stellate cells represent liver-resident mesenchymal stem cells (MSCs) due to their MSC-related expression profile, their potential to differentiate into adipocytes or osteocytes, and their supportive effects on extramedullary hematopoiesis (8, 9). In line with this, stellate cells can originate from the bone marrow (BM) (10, 11), where MSCs were initially discovered (12), and reside in the liver close to endothelial cells, as is the case for MSCs in other organs (13). Quiescent stellate cells are typically located between sinusoidal endothelial cells and hepatocytes in the space of Disse, which has characteristics of stem cell niches (14, 15). Moreover, activated stellate cells can develop into hepatocyte-like cells in vitro (14, 16), and fate-mapping experiments using Gfap and Acta2 promoters indicate that they contribute to liver regeneration in vivo (17–19). In contrast to this, lineage-tracing experiments with lecithin retinol acyltransferase (LRAT) and mesoderm posterior 1 homolog (MESPI) failed to demonstrate a differentiation of HSCs into epithelial cells during regeneration of the injured mouse liver (20, 21). Interestingly, LRAT and MESPI, which are suggested to be specifically expressed by HSCs or their precursors (20, 21), are expressed by embryonic and adult stem cells from mice and humans (Embryonic Stem Cell Database, http://biit.cs.ut.ee/escd/ref.s22 23). Owing to these opposing lineage-tracing studies, we performed cell transplantation experiments with HSCs in the present study to determine their contribution to liver regeneration. Transplanted stellate cells from the pancreas were recently reported to participate in liver repair through differentiation into epithelial cell lineages in a stem cell–based in vivo liver regeneration model (24). Also, MSCs from the BM and adipose tissue were shown to differentiate into liver parenchymal cells (25–27). Tissue-specific engrainment (homing), trafficking to sites of organ injury, and participation in tissue repair are important properties of stem cells that can be analyzed by cell transplantation experiments (28–31). Stem cells can be repeatedly transplanted into...
Owing to their high lipid content, HSCs formed a clear layer at the upper Nycodenz gradient after centrifugation (not shown). HSCs enriched by this technique were then sorted by retinoid-dependent fluorescence-activated cell sorting (FACS). During FACS analysis, we determined the morphology of HSCs by forward and side scatter (Figure 1A). HSCs with similar morphology (Figure 1A, gate) and characteristic retinoid (vitamin A) fluorescence after excitation at 350 nm (Figure 1B, gate) were finally collected. We found that HSCs were about 13.2 ± 0.24 μm (n = 16) in diameter immediately after sorting and exhibited typical stellate-shaped morphology, contained lipid vesicles with retinoids, and expressed eGFP as determined after excitation of living cells by blue and UV light after 2 days in culture (Figure 1C). Ectodermal and mesodermal stellate cell markers such as GFAP, nestin, and vimentin (red), along with desmin (green), as determined by immunofluorescence. Cell nuclei were stained with DAPI (blue). Scale bars: 100 μm (C); 50 μm (D–F).

Results

Characterization of isolated HSCs. HSCs from male eGFP+ Wistar rats were isolated by enzymatic perfusion of the liver and subsequently enriched by Nycodenz density gradient centrifugation.

Figure 1. Characterization of freshly isolated HSCs from eGFP+ rats sorted by retinoid-dependent FACS.

Owing to their high lipid content, HSCs formed a clear layer at the upper Nycodenz gradient after centrifugation (not shown). HSCs enriched by this technique were then sorted by retinoid-dependent fluorescence-activated cell sorting (FACS). During FACS analysis, we determined the morphology of HSCs by forward and side scatter (Figure 1A). HSCs with similar morphology (Figure 1A, gate) and characteristic retinoid (vitamin A) fluorescence after excitation at 350 nm (Figure 1B, gate) were finally collected. We found that HSCs were about 13.2 ± 0.24 μm (n = 16) in diameter immediately after sorting and exhibited typical stellate-shaped morphology, contained lipid vesicles with retinoids, and expressed eGFP as determined after excitation of living cells by blue and UV light after 2 days in culture (Figure 1C). Ectodermal and mesodermal stellate cell markers such as GFAP, nestin, vimentin, and desmin were detected by immunofluorescence in HSCs obtained by FACS (Figure 1D–F), whereas hepatobiliary marker proteins such as keratins and α-fetoprotein were undetectable (not shown). This indicated that typical HSCs were highly purified through the combination of density gradient centrifugation and retinoid-dependent FACS without contamination by other liver cells.

Transplantation of HSCs (2AAF/partial hepatectomy model). Freshly isolated HSCs from male eGFP+ rats obtained by combined density gradient centrifugation and retinoid-dependent FACS were transplanted into female wild-type (WT) rats via tail-vein injection immediately after liver injury through partial hepatectomy (PHX). The host animals were pretreated with 2AAF for 7 days in order to inhibit the proliferation of mature hepatocytes and to induce stem cell–based liver regeneration. We investigated...
The localization of transplanted eGFP+ HSCs in host livers by eGFP immunofluorescence after 7 and 14 days of liver regeneration. We found no significant eGFP staining in WT control livers (Figure 2A), whereas eGFP-expressing cells were detected mainly in the portal field of injured rat livers (2AAF/PHX model) that received eGFP+ HSCs (Figure 2, B and O). We assessed the abundance of eGFP+ male HSCs in female host livers by quantitative PCR (qPCR) analysis of male-specific sex-determining region Y
such as collagen type 1 α2 chain (Col1a2), Col4a1, or Acta2 compared with the expression levels observed in injured control livers (2AAF/PHX model), as investigated by qPCR on days 1, 7, and 14 (shown for day 14 in Figure 2F). Expression of fibrosis-associated markers transiently increased in livers of the 2AAF/PHX model during regeneration, with maximal expression around day 7 (not shown). We also observed engraftment of transplanted eGFP+ HSCs in the pancreas of WT host rats, as investigated by RT-PCR and qPCR of eGFP mRNA. The eGFP+ HSCs showed a low abundance of about 0.02% (± 0.01%; n = 4; 7 days) and 0.3% (± 0.09%; n = 8; 14 days) in the host pancreas. Pancreatic tissue from eGFP+ transgenic (100%) and WT rats (0%) were used for normalization. Although transplanted eGFP+ HSCs were below the detection limit of qPCR 1 day after transplantation (Figure 2, D and E), RT-PCR of eGFP and immunohistochemistry of eGFP using 3,3′-diaminobenzidine (DAB) revealed their presence in sinusoids.

Figure 3. Transplanted male HSCs form bile duct cells and functional hepatocytes as investigated by FISH of gender-specific chromosomes and GUNN rats (2AAF/PHX model). (A) FISH of Y chromosome (red) on a male liver section was used as a positive control. (B) Female liver without positive FISH of Y chromosome. (C) FISH of Y (red) and X (green) chromosomes in male liver. Costaining of (D) HNF4α (green) and (E) panK with FISH of Y chromosome (red) in female host livers after transplantation of male HSCs. Arrows indicate HSC-derived (D) hepatocytes and (E) bile duct cells with Y chromosome. (F) Detection of Y (red) and X (green) chromosomes in female host liver. Arrows indicate diploid and tetraploid HSC-derived hepatocytes without cell fusion. Microscopic images were taken in the portal fields of injured livers (zone 1). (G–I) HSCs were transplanted into GUNN rats that underwent PHX in the presence of 2AAF. (G) The presence of hepatocytes derived from transplanted HSCs in homozygous GUNN rats was analyzed by RT-PCR of Ugt1a1 mRNA followed by enzymatic cleavage (BstNI) of the mutation site. (H) Serum concentrations of direct (red bars) and indirect bilirubin (yellow bars) were determined after 4 weeks and compared with those detected in GUNN rats (controls), which were treated in a similar way but without transplanted HSCs, and in normal rats (n = 3–6; *P < 0.05). (I) Albumin concentrations were measured in the blood serum of normal rats as well as GUNN rats with and without transplanted HSCs (n = 3–6). Scale bars: 50 μm (A–D and F); 20 μm (E).

(Sry) DNA and eGFP mRNA (Figure 2, D and E). About 60% of the host animals that received eGFP+ HSCs exhibited measurable Sry DNA or eGFP mRNA levels. Female host livers with stellate cell engraftment showed that approximately 10% to 14% of liver cells derived from transplanted male eGFP+ HSCs after 7 and 14 days of regeneration, whereas HSC engraftment remained undetectable by qPCR in the host liver (Figure 2, D and E) and lung (not shown) 1 day after transplantation (n = 5). Conventional semiquantitative reverse transcriptase–polymerase chain reaction (RT-PCR) revealed the presence of eGFP+ HSCs in host livers 1 day after transplantation (not shown). The eGFP mRNA levels within the WT host liver varied between 6% and 22% (7–14 days) among different animals (n = 10) when qPCR was applied. The increase in Sry DNA and eGFP amounts during the first week indicated that transplanted HSCs proliferated in the host liver (Figure 2, D and E) without elevating the expression of fibrosis-associated markers such as collagen type 1α2 chain (Col1a2), Col4a1, or Acta2 compared with the expression levels observed in injured control livers (2AAF/PHX model), as investigated by qPCR on days 1, 7, and 14 (shown for day 14 in Figure 2F). Expression of fibrosis-associated markers transiently increased in livers of the 2AAF/PHX model during regeneration, with maximal expression around day 7 (not shown). We also observed engraftment of transplanted eGFP+ HSCs in the pancreas of WT host rats, as investigated by RT-PCR and qPCR of eGFP mRNA. The eGFP+ HSCs showed a low abundance of about 0.02% (± 0.01%; n = 4; 7 days) and 0.3% (± 0.09%; n = 8; 14 days) in the host pancreas. Pancreatic tissue from eGFP transgenic (100%) and WT rats (0%) were used for normalization.

Although transplanted eGFP+ HSCs were below the detection limit of qPCR 1 day after transplantation (Figure 2, D and E), RT-PCR of eGFP and immunohistochemistry of eGFP using 3,3′-diaminobenzidine (DAB) revealed their presence in sinusoids.
When HSC clones, which derived from single eGFP+ HSCs by differentiation potential within the regenerating liver (2AAF /PHX) livers (2AAF /PHX model) (Figure 2O). We found a similar differentiation (Figure 2N). We did not observe immunofluorescence of HNF4α also investigated by fast red immunohistochemistry of eGFP and 2M) zones. The differentiation of HSCs into hepatocytes was the periportal (zone 1; Figure 2L) and pericentral (zone 3; Figure 2M) of the host liver (Supplemental Figure 1, B and C). We verified the specificity of the Y chromosome probe on male (positive control) and female (negative control) liver sections (Figure 3, A and B) and by combining Y and X chromosome probes through distinct signals for each chromosome in the cell nuclei of male control liver sections (Figure 3C). After transplantation of primary HSCs from male rats sorted by FACS, Y chromosome was taken in the portal fields of injured livers (zone 1). Scale bars: 200 μm (A and C); 50 μm (B, D, G, and J-L); 10 μm (E, F, H, and I).

oids as desmin-expressing stellate cells and demonstrated their homing to the host liver (Figure 2G). We also found eGFP+ HSCs to be integrated and associated with bile ducts in the host liver 1 day after transplantation (Figure 2H). WT control livers (2AAF/PHX model) displayed no DAB staining of eGFP (Figure 2I). DAB immunohistochemistry of eGFP, in combination with immunofluorescence of the bile duct marker K19 and hepatocyte marker HNF4α, indicated that transplanted HSCs contributed to bile duct and hepatocyte formation after 14 days of liver regeneration (Figure 2, J–M). We found HSC-derived hepatocytes in both the periporal (zone 1; Figure 2L) and pericentral (zone 3; Figure 2M) regions. The differentiation of HSCs into hepatocytes was also investigated by fast red immunohistochemistry of eGFP and immunofluorescence of HNF4α (Figure 2N). We did not observe positive fast red immunohistochemistry of eGFP in WT control livers (2AAF/PHX model) (Figure 2O). We found a similar differentiation potential within the regenerating liver (2AAF/PHX) when HSC clones, which derived from single eGFP+ HSCs by clonal expansion, were transplanted. As indicated by qPCR analysis of eGFP mRNA, different HSC clones reconstituted about 16% (± 5%; range 3%–28%; n = 4) of the host liver (Supplemental Figure 1, A–C). We investigated the differentiation of male HSC clones into epithelial cells within the female host liver by fluorescence in situ hybridization (FISH) of male-specific Y chromosome. FISH analysis revealed HSC-derived Y chromosome in hepatocytes and bile duct cells when immunofluorescence of HNF4α or keratins (pan keratin, panK) was added (Supplemental Figure 1, B and C). We verified the specificity of the Y chromosome probe on male (positive control) and female (negative control) liver sections (Figure 3, A and B) and by combining Y and X chromosome probes through distinct signals for each chromosome in the cell nuclei of male control liver sections (Figure 3C). After transplantation of primary HSCs from male rats sorted by FACS, Y chromosome was detected by FISH in HNF4α+ and panK+ cells within the female host liver (2AAF/PHX model), indicating HSC-derived hepatocytes and bile duct cells within the female host liver (Figure 3, D and E). Simultaneous FISH of Y and X chromosomes displayed normal diploid and tetraploid hepatocytes that derived from transplanted male HSCs without fusion with preexisting female hepatocytes (Figure 3F). Transplanted HSCs were able to restore liver function of homozygous GUNN rats, which lack uridine 5′-diphospho-glucuronosyltransferase 1A1 (UGT1A1) activity due to mutation in exon 4. The UGT1A1 enzyme is expressed in hepatocytes and required for bilirubin glucuronidation (conjugation), which enables bilirubin excretion. Owing to Ugt1a1 deficiency, blood sera from homozygous GUNN rats exhibit high bilirubin serum levels compared with levels in normal rats. We verified the presence of Ugt1a1 transcripts without mutation in homozygous GUNN rat livers after transplantation of HSCs by enzymatic cleavage (BstNI) of Ugt1a1 PCR products, which results in additional PCR products similar to those obtained in heterozygous GUNN or normal rats. In contrast, mutated Ugt1a1 remained uncleaved and appeared as
a single PCR product at 465 bp (Figure 3G). In line with this, serum levels of conjugated (direct) bilirubin in homozygous GUNN rats increased 2-fold within 4 weeks after transplantation of HSCs derived from rats with normal \textit{Ugt1a1} when compared with serum levels in GUNN rats without transplanted HSCs (Figure 3H). This indicates partial correction of the bilirubin-handling defect in GUNN rats through differentiation of transplanted HSCs into hepatocytes, although the amounts of unconjugated (indirect) bilirubin were not yet significantly altered (Figure 3H). The albumin serum concentrations remained constant (Figure 3I).

Transplanted HSCs and ductular reactions (2AAF/PHX model). The ductular reaction is regarded as a response of stem/progenitor cells to severe liver injury and is formed by K19+ progenitor cells as well as surrounding mesenchymal cells. These ducts transiently appeared in the 2AAF/PHX model and were found in the regenerating liver from day 7 to day 14 after PHX (Figure 4). Mesenchymal marker proteins such as vimentin and desmin were highly expressed in liver tissue with ductular reactions (Figure 4, A and B). Progenitor cells that express the mesodermal protein vimentin and the epithelial marker K19 constituted the core of the ducts (Figure 4, C–F) surrounded by cells that expressed the stellate cell markers desmin (Figure 4G) and nestin (not shown). We verified the presence of vimentin in K19+ cells within these duct-like structures by immunofluorescence and ELYRA super-resolution microscopy (Zeiss) (Figure 4, E and F). Although desmin- and K19-expressing cells seem to be clearly distinct in these structures (Figure 4G), super-resolution microscopy revealed the presence of desmin residues in K19+ cells (Figure 4, H and I), indicating that intermediate states of mesenchymal and epithelial cells exist in putative progenitor cells of the liver. We found transplanted eGFP+...
HSCs in the host liver in areas with ductular reactions by day 7 of liver regeneration (2AAF/PHX model), as shown by DAB immunohistochemistry of eGFP. Simultaneous immunohistochemical staining of eGFP with the progenitor cell markers K19 and α-fetoprotein indicated that transplanted eGFP+ HSCs contributed to putative epithelial progenitors that formed duct-like structures (Figure 4, J-L). Although we found HSC-derived eGFP+ and K19+ cells among ductular cells, the majority of eGFP+ cells seemed to surround the ducts as mesenchymal tissue (Figure 4, J and K).

Retransplantation of HSCs (2AAF/PHX model). In addition to their presence in the liver, we also found engraftment of transplanted HSCs (1%-2%; n = 5; Figure 5, A and B) and HSC clones (1%; n = 5; not shown) in the BM of host rats as investigated by qPCR of Sry DNA and eGFP mRNA 7-14 days after tail-vein injection. Since cells of the BM can be gently isolated from long bones without enzymes, we used this source for retransplantation experiments with stellate cells. FACSorted eGFP+ HSCs were first stained by CellTracker Red and then transplanted into WT rats immediately after PHX in the presence of 2AAF. One day after transplantation, the host BM cells were harvested and cultured. The intense CellTracker fluorescence, as demonstrated for freshly isolated HSCs (Figure 5C), was used to find transplanted HSC-derived eGFP+ cells among WT BM cells (Figure 5, D-F), which finally converted into myofibroblast-like cells during culture (not shown). eGFP expression in HSC-derived cells was maintained during culture (Figure 5G) and used to analyze their engraftment in the liver and BM 14 days after their transplantation into new host rats (2AAF/PHX model) (Figure 5, H and I). The retransplanted eGFP+ HSC-derived cells were able to reconstitute about 17% (± 5%; n = 3) of the injured host livers as investigated by qPCR of eGFP mRNA. The percentage of eGFP+ cells varied in the different areas of the host livers (Supplemental Table 1).

Transplantation of HSCs (retrorsine/PHX model). FACSorted HSCs from eGFP+ male rats were also injected into WT female rats that underwent PHX in the presence of retrorsine, which prevents the proliferation of mature hepatocytes to facilitate engraftment of transplanted cells in the injured liver. In this model, transplanted eGFP+ HSCs formed about 6% (± 2%; range 1%-14%; n = 7) of the host liver, as investigated by qPCR of eGFP mRNA (Figure 6A). Engraftment of transplanted eGFP+ HSCs was detectable by qPCR in all host rats when the retrorsine/PHX model was applied. The eGFP+ HSCs were maintained as stellate cells in the liver sinusoids of the host liver 14 days after their transplantation (retrorsine/PHX model), as demonstrated by combined DAB immunohistochemistry of eGFP and immunofluorescence of desmin (Figure 6B). Immunohistochemistry of eGFP (fast red) and immunofluorescence of the hepatocyte marker HNF4a revealed that transplanted eGFP+ HSCs also differentiated into hepatocytes (Figure 6C). The generation of hepatocytes by male HSCs in female host liver was confirmed by FISH of Y chromosome in conjunction with HNF4a immunofluorescence (Figure 6D). We also found the Y chromosome in bile duct cells when Y chromosome probes were combined with panK immunofluorescence (Figure 6E). We found that the transplanted male HSCs were able to generate hepatocytes in the female host liver without cell fusion, as determined by FISH of Y and X chromosomes (Figure 6F).

Hepatic differentiation of MSC populations in vitro. Participation of transplanted HSCs in ductular reactions and the potential of
Figure 7. MSCs transiently acquire the gene expression of liver progenitor cells during their differentiation into hepatocytes. (A–O) Freshly isolated HSCs, clonally expanded HSCs (clones 1E7, 4E7, and 3F8), BM MSCs, and UCBSCs (clone 1G11) from rats were treated for 21 days with a hepatocyte differentiation medium that contained HGF and FGF4. Expression of (A) vimentin, (B) desmin, (C) K19, (D) Epcam, (E) α-fetoprotein, (G) Cyp7a1, (H) Hnf4α, (I) albumin, and (J) Sox9 was analyzed by qPCR. mRNA samples were taken at weekly intervals. Gene expression of the cells at the beginning of the differentiation experiments is indicated as Day 1. (K) To verify hepatic differentiation, the hepatocyte marker albumin was also measured in the culture supernatant of HSCs, HSC clones, BM MSCs, and UCBSCs by a rat-specific albumin ELISA. Hepatic function of MSC-derived hepatocyte-like cells was also determined by quantitation of the bile acids (L) cholic acid, (M) chenodeoxycholic acid, (N) ursodeoxycholic acid, and (O) ω/α-muricholic acid in culture supernatants by UHPLC-MS/MS. Bile acids were allowed to accumulate for 2 days in the culture medium before analysis. Bar values represent arithmetic means and were interpolated to indicate the trend (colored lines) for each cell group. Expression profiles, albumin synthesis, and bile acid release were determined in 3 independent experiments (n = 3).
HSCs to develop into hepatocytes (Figure 2, M and N, and Figure 3D) and cholangiocytes (Figure 2, K and L, and Figure 3E) raise the possibility that stellate cells could transiently acquire the expression pattern of liver progenitor cells during hepatic differentiation. Stellate cells were recently described as liver-resident MSCs (8) and showed an expression profile similar to that of clonally expanded HSCs, rat BM MSCs, and umbilical cord blood stem cells (UCBSCs) (Supplemental Figure 2). We treated these MSC populations with hepatocyte growth factor (HGF) and human fibroblast growth factor 4 (FGF4) for 21 days in order to monitor hepatic differentiation. The relative mRNA amounts of stellate cell or mesenchymal markers (vimentin, desmin, Lrat), indicators for progenitor cells and putative stem/progenitor cells (K19, Epcam, leucine-rich repeat-containing G protein–coupled receptor 5 [Lgr5], α-fetoprotein) as well as markers of mature hepatocytes (cytochrome P450 7A1 [Cyp7a1], Hnf4α, albumin) were analyzed by qPCR (Figure 7 and Supplemental Figure 3, A–C). The expression levels of vimentin, desmin, and Lrat steadily decreased by 20% to 30% during hepatic differentiation (Figure 7, A and B, and Supplemental Figure 3, A and B), indicating that only a subset of MSCs had differentiated in vitro. In contrast, the expression levels of K19, Epcam, Lgr5, and α-fetoprotein transiently increased and peaked on day 14 during hepatic differentiation of cultured HSCs and other MSC populations (Figure 7, C–F). The mRNA values of the hepatocyte markers Cyp7a1, Hnf4α, and albumin continuously increased until day 21 of differentiation in all MSCs (Figure 7, G–I), indicating final differentiation into the hepatic cell lineage. The mRNA values of sex-determining region Y box 9 (Sox9) decreased in primary cultured HSCs in both the growth factor–treated (Figure 7J) and control groups (Supplemental Figure 3C), whereas the Sox9 levels of HSC clones and other MSC populations remained unchanged in all experimental media (Figure 7J). This may indicate that Sox9 was apparently not involved in hepatic differentiation. Neither progenitor cells (K19, Epcam, or α-fetoprotein) nor hepatocyte markers (Cyp7a1, Hnf4α, or albumin) were detected by qPCR in the controls without growth factors over a 21-day culture, as demonstrated for albumin expression in Supplemental Figure 3D. We confirmed the appearance of hepatocyte–like cells by quantification of albumin in the culture supernatant of the MSCs using a rat-specific albumin enzyme-linked immunosorbent assay (ELISA) (Figure 7K). In contrast to HSCs, muscle fibroblasts were unable to respond by albumin mRNA expression or protein synthesis after growth factor treatment for 21 days (Supplemental Figure 3, D and E). Also, the release of bile acids such as cholic acid, chenodeoxycholic acid, ursodeoxycholic acid, and α/ω-muricholic acid into the culture supernatants indicated that all MSC populations acquired hepatocyte-specific functions during growth factor treatment (Figure 7, L–O). In the culture media of HGF- and FGF4-treated MSCs, 19 of 29 investigated bile acids were detected by ultra-high-performance liquid chromatography–tandem mass spectrometry (UHPLC-MS/MS) (Supplemental Figures 4–7), whereas bile acid synthesis was not found under control conditions without cytokines within a 21-day period (Supplemental Figure 3F and Supplemental Tables 2–5). With the exception of Sox9 expression in HSC clones — BM MSCs and UCBSCs that remained unchanged (Figure 7) — all trends in gene expression, albumin synthesis, and bile acid synthesis were significant, as determined by ANOVA (not indicated in Figure 7; see Supplemental Figure 3).

We also investigated the transient appearance of putative progenitor cells in differentiation experiments of primary cultured HSCs using combined immunofluorescence of desmin with K19, α-fetoprotein, or LGR5. Small cells with K19 filaments, α-fetoprotein, or LGR5 exhibited desmin filaments 14 days after HGF and FGF4 treatment (Figure 8, A–C). Only LGR5 was detectable under control conditions in myofibroblast–like cells, which typically develop from culture-activated stellate cells, whereas K19 and α-fetoprotein remained undetectable (Figure 8, D–F). Hepatocyte-like cells with residual desmin filaments in HSC primary cultures were identified by immunofluorescence of bile salt export pump (BSEP), sodium taurocholate cotransporting polypeptide (NTCP), and K18 21 days after we initiated the growth factor treatment (Figure 8, G–L). In control experiments without HGF and FGF4, HSC-derived myofibroblasts expressed desmin but remained negative for BSEP, NTCP, and K18 within 21 days of culture (Figure 8, M–O). In summary, we found that stellate cells and other MSCs reduced their mesodermal characteristics during differentiation into hepatocytes (endoderm), which was accompanied by a transient appearance of progenitor cell markers (Supplemental Figure 8).

Discussion

This study demonstrates for the first time to our knowledge that HSCs are transplantable, home to the liver of host animals, and participate in liver regeneration through differentiation into parenchymal and bile duct cells without cell fusion. Transplanted HSCs were able to ameliorate the glucuronidation defect in GUNN rats, as indicated by increased serum levels of conjugated bilirubin and the appearance of nonmutated UGT1A1 enzymes in the liver of these animals. This confirms the contribution of stellate cells to the formation of functional hepatocytes, although total bilirubin serum concentrations in host GUNN rats were not significantly lowered. A complete resolution of the bilirubin-handling defect may require higher numbers of transplanted HSCs and additional time for bilirubin clearance. In addition to these findings obtained with freshly isolated HSCs, single-cell clones of HSCs were established that preserved stellate cell characteristics and the ability to differentiate, indicating a self-renewal potential of HSCs. These important stem cell features further support the characterization of stellate cells as liver-resident MSCs (8, 16). In line with their characterization as MSCs, engraftment of transplanted HSCs was also found in the BM of host animals, as demonstrated in the present study. MSCs from BM or adipose tissue were reported to contribute to liver repair (25–27), which suggests that hepatic differentiation capacity is a general feature of MSCs. There is currently no consensus about a direct contribution of stem cells with mesodermal characteristics to epithelial tissue regeneration in the liver, although mesodermal proteins such as vimentin are expressed by progenitor cells along with epithelial markers (35). It is assumed that stem cells with epithelial characteristics are the source of liver progenitor cells (34), which ensure tissue repair in the event of impaired hepatocyte proliferation (32, 33). In line with this, epithelial markers such as K19 and EPCAM are expressed in ductular reactions (35), which are described as a stem/progenitor cell response (36). However, the simultaneous appearance of epithelial and mesodermal proteins in liver progenitor cells has remained
unexplained thus far. As shown in the present study, transplanted eGFP+ HSCs can contribute to the formation of K19+ and α-fetoprotein+ cells during ductular reactions, indicating that MSCs can generate progenitor cells in vivo. Furthermore, isolated HSCs cultured on collagen can form tube-like structures, as reported earlier (16). We provided an explanation of the simultaneous expression of mesenchymal and epithelial proteins in progenitor cells by time-resolved hepatic differentiation experiments. During in vitro differentiation of HSCs, HSC clones, BM MSCs, and UCBSCs, all MSC populations shifted from a mesodermal to an endodermal expression pattern during differentiation into hepatocytes. In this intermediate stage, they transiently coexpressed mesenchymal and epithelial markers, as is known to occur with progenitor cells. In addition, we found the recently described hepatic stem cell marker LGR5 (37) to be transiently upregulated during hepatic differentiation of MSCs. These findings, on the one hand, are in line with the observed contribution of transplanted HSCs to ductular reactions as well as the occurrence of mesodermal protein residues (vimentin and desmin) in K19+ progenitor cells in vivo and, on the other hand, suggest a participation of MSC populations in progenitor cell formation and liver repair. Desmin+ and nestin+ cells that surround ductular structures with liver progenitor cells in the cen-

ter (38) indicate the presence of MSCs. The present study shows that liver progenitor cells can originate from liver-resident stellate cells and other MSC populations that can potentially migrate from distant organs to the liver after severe injury in order to support regeneration. Thus, mesenchymal cells that surround ductular reactions may not only influence the differentiation of progenitor cells, as described recently (36), but may also contribute to their population. Evidence exists that hepatocyte and bile duct precursors originate from stem cells in the canals of Hering (39), and the ductular reaction emerges in the portal field. It remains an unresolved question whether MSC populations always contribute to progenitor cell formation or only under extreme conditions. Removal of HSCs from their niche in the space of Disse through enzymatic isolation and transplantation into the regenerating liver via the circulation creates an extreme situation that can promote their differentiation. Also, as shown in the present study, the injury model affects the outcome of the experiments. We found a higher engraftment of transplanted HSCs in the 2AAF/PHX model compared with that in the retrorsine/PHX model. The environment obviously controls the behavior of transplanted HSCs, which can respond in at least 3 ways: (a) HSCs remain activated and produce extracellular matrix and factors to influence neighboring cells; (b)
activated HSCs can become quiescent again; and (c) HSCs differentiate into epithelial cells. The latter 2 possibilities may explain the absence of increased Collα2, Col4α1, and Acta2 expression in the host liver despite the proliferation of transplanted HSCs. Regardless of the controversies emerging from recent lineage-tracing studies that argue for (17–19) or against (20, 21) a contribution of HSCs to liver regeneration, evidence for the characterization of HSCs as liver-resident stem cells and for their contribution to tissue repair was obtained in this study. These findings do not contradict those of previous studies demonstrating a contribution of HSCs to fibrogenesis, because chronic liver diseases certainly affect their behavior, as is also supposed for MSCs (40). With respect to mesenchymal and epithelial characteristics in progenitor cells, a recent study indicated a contribution of prominin-1/ K19^+ liver progenitor cells with mesodermal markers to fibrosis in biliary atresia (41). Future research should address whether MSC populations are connected to stem cell niches within the canals of Hering and contribute to progenitor cell formation. Monitoring of hepatic differentiation of MSC populations and mesodermal proteins in epithelial cells in ductular reactions, as demonstrated in this study, as well as recent lineage-tracing experiments with the Acta2 promoter (18, 19), support this possibility.

Methods

Cell isolation and culture. A detailed description of cell isolation and culture as well as FACS protocols is provided in the Supplemental Methods.

Liver injury and HSC transplantation. Pellets that contained 2-acetylaminofluorene (70 mg, 2AAF 14-day release; Innovative Research of America) were implanted under the neck skin of WT Wistar rats (250 g) (Heinrich Heine University Animal Facility) 7 days before PHX was performed as previously described (42, 43). eGFP^+ HSCs from Wistar-TgN(CAG-GFP)184Ys rats (Rat Resource & Research Center, University of Missouri, Columbia, Missouri, USA) were enriched by density gradient centrifugation (44), purified by retinoid-dependent FACS, and finally transplanted into WT female rats via tail-vein injections immediately after PHX. About 700,000 ± 100,000 HSCs or clonally expanded HSCs from male eGFP^+ rats were injected into each rat that underwent liver injury. HSCs were transplanted into a normal and a mutant Wistar rat strain (GUNN-UGT1A1^+/wt*; wt**; Rat Resource & Research Center) (45). Homozygous mutants lack UGT1A1 enzyme activity and exhibit elevated bilirubin serum levels. Direct (conjugated) and indirect (unconjugated) bilirubin as well as albumin concentration in the serum of rats were determined by the Central Laboratory of Heinrich Heine University 4 weeks after HSC transplantation. For retransplantation experiments, FACS-sorted eGFP^+ HSCs were stained with 10 μM CellTracker Red (C34552; Invitrogen) in suspension under serum-free conditions at 37°C for 45 minutes. After removal of excess dye by repeated centrifugation, about 100,000 double-labeled HSCs were transplanted into WT rats treated with 2AAF immediately after PHX. BM cells of the host animal were harvested 1 day after transplantation, cultured for 14 days, and then transplanted into other WT rats pretreated with 2AAF and PHX (about 10 million cells). In another injury model, female WT Wistar rats (250 g) received 2 i.p. retinorsine (R0382-100MG; Sigma-Aldrich) injections (30 mg/kg) 14 days apart. After an additional 4 weeks, PHX was performed (46), and FACS-sorted eGFP^+ HSCs from male Wistar rats were transplanted as described above.

Cell differentiation. The differentiation of isolated HSCs, HSC clones, BM MSCs, and UCBSCs was induced by 50 ng/ml FGF4 (100-31; PreproTech) and 40 ng/ml human HGF (100-39; PreproTech) dissolved in Iscove’s modified Dulbecco’s medium (12440-046; Invitrogen) supplemented with 1% insulin-transferrin-sodium selenite (ITS) (11884; Sigma-Aldrich), 1% linolic acid-albumin from bovine serum (L9530; Sigma-Aldrich), and 1% antibiotic-antimycotic solution (Invitrogen). The differentiation medium was exchanged every second to third day. Cells in control experiments were cultured in the same medium, but without FGF4 and HGF. The progression of cell differentiation was monitored at weekly intervals.

Analysis of cells and tissue sections. Isolated cells and cryosections of the liver were fixed with either ice-cold methanol for 5 minutes or 4% formalin for 10 to 20 minutes and incubated with antibodies against GFAP (MAB3402; Millipore), K18 (BM2275P; Acris), K19 (NB100-687; Novus Biologicals or 65129; ProGen Biotechnik), panK (CM162; Biocare Medical), desmin (ab8592; Abcam), vimentin (M0725; Dako), nestin (sc-33677; Santa Cruz Biotechnology Inc.), LGR5 (GTX62071; GeneTex), and HNF4α (313; Cell Signaling Technology) for immunofluorescence staining. The primary antibodies against NTCP and BSEP were provided by B. Stieger and P. Meier (University Hospital Zürich, Zürich, Switzerland). Appropriate secondary cyanine dye 3 (Cy3) or FITC-labeled antibodies (AP182C, AP162C, AP192F, and AP182F; Milipore) and DAPI (P36935, ProLonged Gold; Invitrogen) were applied to mark primary antibodies and cell nuclei, respectively. Transplanted eGFP^+ HSCs were detected by immunohistochemistry on formalin-fixed and Triton X-100–treated (0.1% in PBS, 3051.3, Carl Roth GmbH) cryosections of the liver using monoclonal (2956, D5.1 XP; Cell Signaling Technology) and polyclonal antibodies (210-PS-1GFP; Immunotech) against eGFP followed by labeling with secondary antibodies coupled to alkaline phosphatase or HRP (K5361, EnVision G/2 Double Stain System Rabbit/Mouse; Dako). Polyclonal eGFP antibodies and secondary antibodies used to detect eGFP were preincubated with liver homogenate from WT rats for at least 30 minutes to prevent unspecific binding. Fast red tablets containing levamisole for blocking endogenous alkaline phosphatase (1I496549001; Roche) were used to prepare the substrate solution for alkaline phosphatase. Cell nuclei were counterstained by diluted hematoxylin solution (1:4 in distilled water) (1.05175; Merck) on fast red–stained liver sections. The hematoxylin solution was removed by washing with distilled and tap water. eGFP^+ cell types were identified by antibodies against HNF4α, K19, or desmin and subsequently labeled by appropriate secondary antibodies coupled to fluorescent dyes. For reverse transcriptase PCR (RT-PCR), tissue samples were taken from at least 4 different liver lobes, and the first-strand cDNA was made from 1 μg total RNA per 20 μl reaction volume using the RevertAid H Minus First Strand cDNA Synthesis Kit (K1632; Thermo Scientific). PCR was performed according to standard protocols using the 2xPCR Master Mix (K0171; Thermo Scientific), 0.8 μmol/l primers (Supplemental Table 6), and 1 or 5 μl template cDNA per 25 μl reaction volume. A maximum of 20 or 35 cycles was used to amplify PCR products. The SensiMix SYBR No-ROX Kit (QT650-05; Bioline) was used for real-time qPCR according to the manufacturer’s instructions. To assess the engraftment of male HSCs after transplantation, total DNA was purified with the DNeasy Blood & Tissue Kit (69881; Qiagen). For the amplification of PCR products, 12.5 ng cDNA or 500 ng DNA and 0.6 μmol/l primers (Supplemental Table 7) were taken for each reaction. After an initial denaturation at 95°C,
the annealing was carried out at 58°C, and elongation was performed at 72°C using 40 cycles and a 7Optical Cycler (Analytik Jena AG). All samples were measured in triplicate, and hypoxanthine-guanine phosphoribosyltransferase 1 (Hprt1 mRNA of isolated cells) or β-actin (DNA of liver tissue) was used as a reference for the normalization of the results obtained by the 2^−ΔΔCt method. Rat albumin was quantified in the culture supernatant by a rat-specific albumin ELISA (E110-125; Bethyl Laboratories) according to the manufacturer’s recommendations.

For the detection of UGTT1A enzymes in the livers of normal and Gunn rats, the PCR product of Ugt1a1 mRNA was cleaved by the BstNI restriction enzyme (New England Biolabs) according to the manufacturer’s instructions and separated by agarose gels (3%). DNA probes for rat Y (IDR1070; IDLabs) and X (IDRF1067; IDLabs) chromosomes were purchased for FISH analysis after fixation of liver sections with Carnoy’s fixative for 15 minutes and tissue digestion by Pepsin (10108057001; Roche) for 5 minutes as previously described (24).

Analysis of bile acids and their conjugates. Bile acids were analyzed by UHPLC-MS/MS in culture supernatants of HGF- and FGF4-treated cells. The UHPLC-MS/MS system consisted of a UPLC-H class coupled to a Xevo TQ-S triple-quadruple mass spectrometer (Waters). Electrospray ionization was performed in the negative ionization mode. Chromatographic separation was performed on a UPLC HSS C18 column (Waters) (2.1 x 100 mm, 1.8 μm). The mobile phase consisted of water containing 0.1% formic acid and 5 mM ammonium acetate (elucent A) and acetonitrile (elucent B). Analytes were separated by a gradient elution. The injection volume was 5 μl, and the column was maintained at 40°C. Detection of the bile acids and their glycine and taurine conjugates was performed in selected reaction monitoring (SRM) mode. All standards as well as the deuterated internal standards (IS) substance (d4-CA, d4-GCA, and d4-GCDCA) were purchased from Sigma-Aldrich and Steraloids.


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Address correspondence to: Dieter Häussinger, Clinic of Gastroenterology, Hepatology and Infectious Diseases, Heinrich Heine University Düsseldorf, Moorvenstraße 5, 40225 Düsseldorf, Germany. Phone: 004.211.81.17569; E-mail: haeussin@uni-duesseldorf.de.


