Cellular organization of normal mouse liver: a histological, quantitative immunocytochemical, and fine structural analysis

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Abstract The cellular organization of normal mouse liver was studied using light and electron microscopy and quantitative immunocytochemical techniques. The general histological organization of the mouse liver is similar to livers of other mammalian species, with a lobular organization based on the distributions of portal areas and central venules. The parenchymal hepatocytes were detected with immunocytochemical techniques to recognize albumin or biotin containing cells. The macrophage Kupffer cells were identified with F4-80 immunocytochemistry, Ito stellate cells were identified with GFAP immunocytochemistry, and endothelial cells were labeled with the CD-34 antibody. Kupffer cells were labeled with intravascularly administered fluorescently labeled latex microspheres of both large (0.5 μm) and small (0.03 μm) diameters, while endothelial cells were labeled only with small diameter microspheres. Neither hepatocytes nor Ito stellate cells were labeled by intravascularly administered latex microspheres. The principal fine structural features of hepatocytes and non-parenchymal cells of mouse liver are similar to those reported for rat. Counts of immunocytochemically labeled cells with stained nuclei indicated that hepatocytes constituted approximately 52% of all labeled cells, Kupffer cells about 18%, Ito cells about 8%, and endothelial cells about 22% of all labeled cells. Approximately, 35% of the hepatocytes contained two nuclei; none of the Kupffer or Ito cells were double nucleated. The presence of canaliculi and a bile duct system appear similar to that reported for other species. The cellular organization of the mouse liver is quite similar to that of other mammalian species, confirming that the mouse presents a useful animal model for studies of liver structure and function.

Keywords Albumin · F4/80 · GFAP · Hepatocyte · Immunocytochemistry

Introduction

The important roles performed by the liver, not only in the storage and release of nutrients but also in the neutralization and elimination of a variety of toxic substances, have prompted investigations of its cellular constituents and organization. Some of these studies have been carried out in human liver, but the importance of having an experimental model system has prompted several investigations of liver organization in laboratory mammals, primarily rats (Blouin et al. 1977; Bouwens et al. 1992; Gard et al. 1985; Geerts 2001; Kawada 1997; Leo and Lieber 1983; Marcos et al. 2004; Sigal et al. 1999; Sleyster and Knoek 1982; Wake et al. 1989; Widmann et al. 1972; Wisse 1970, 1972, 1974; Yokota and Fahimi 1981). In the species studied thus far, investigations have demonstrated that the liver is comprised of several different populations of cells. These include the major parenchymal cells, the hepatocytes (Blouin et al. 1977; David 1985; Jones and Mills 1974; Loud 1968; Yokota and Fahimi 1981), and a variety of non-parenchymal resident cells including a population of macrophages termed Kupffer cells (Bouwens et al. 1986; Naito et al. 1997, 2004; Wake et al. 1989; Widmann et al. 1972; Wisse 1974), fat storing cells termed Ito or stellate...
cells (Fahimi 1982; Ito 1973; Kawada 1997; Sato et al. 2003; Senoo 2004), and endothelial cells (Smedsrod et al. 1994; Stöhr et al. 1978; Widmann et al. 1972; Wisse 1972).

In recent years, the use of mice, and particularly genetically engineered mice, in research laboratories has increased markedly. Several studies have used mice in addressing particular questions in liver structure and function (e.g., Bartók et al. 1983; Longmuir et al. 2006; Naito et al. 1997; Robertson et al. 2008; Sigal et al. 1999; Stöhr et al. 1978; Yamada et al. 1990). Many of these studies assume similarities between the mouse and rat liver, but the putative similarities have not, to our knowledge, been demonstrated. Indeed, notable cases of differences between rat and mouse liver have been reported (Stöhr et al. 1978). Because of the important role that will be played by mice in future studies of liver function, we believe it imperative to establish the baseline of normal cellular composition, to serve as a reference for these future studies. The present paper reports the results of an empirical study of normal mouse liver, with emphasis on identifying the constituent populations of cells in the liver by using immunocytochemical markers, determining essential fine structural features of these cells, and determining the relative numbers of the different immunocytochemically identified populations of cells.

Materials and methods

Materials

Chemical supplies were purchased from Sigma Aldrich (St. Louis, MO) unless specified otherwise.

Animals

All animal work was reviewed and approved by the University of California, Irvine, Institutional Animal Care and Use Committee (IACUC) prior to conducting the experiments. Adult (2–4 mo) BALB/c or ICR female mice, obtained from Charles River (Wilmington, CA), body weight approximately 20–25 g, were used for these experiments. Mice were provided standard laboratory mouse food ad libitum, and were housed in a satellite vivarium with standard 12 h light, 12 h dark cycles.

Tissue preparation

For studies of normal structure, 27 mice were deeply anesthetized with sodium pentobarbital (50 mg/kg i.p.). Mice were perfused through the heart or the portal vein with 10 ml saline, using a pump at 5 ml/min, to clear the liver of blood, then followed with cold 4% paraformaldehyde in sodium phosphate buffer, pH 7.4 for approximately 15 min. Providing the perfusion was successful and the liver cleared of blood, no differences were detected in morphological features between the two routes of perfusion.

The liver lobes were removed, cut into 2–3 mm blocks, and fixed in 4% paraformaldehyde for 1–18 h before being placed in 30% sucrose for cryoprotection. Blocks of liver tissue were frozen in −20°C 2-methylbutane in preparation for sectioning with a cryostat. Frozen liver sections were cut on a Reichert-Jung 1800 cryostat at 10–12 μm, mounted directly on super frost slides (Fisher Scientific, Pittsburgh, PA), and air dried for 10–30 min before processing for immunocytochemistry, lectin binding, or hematoxylin and eosin (H&E) staining.

Latex microsphere and liposome injections

Twelve mice were lightly anesthetized with Ketamine–xylazine (100 mg/kg Ketamine; 5 mg/kg xylazine). Six of these mice received injections into the tail vein of 25–100 μl of a saline solution containing Fluorospheres (fluorescently labeled latex microspheres; 2.5%; Molecular Probes—Invitrogen, Carlsbad CA). Fluorospheres with mean diameters of 0.5 μm or of 0.03 μm were used, injected either separately or combined as a cocktail comprising equal volumes of the two stock suspensions. Fluorospheres of different sizes carried different fluorophores, either rhodamine or fluorescein. Following post-injection survival periods of 1–5 h, animals were deeply anesthetized with sodium pentobarbital and perfused through the heart as described above.

The other six of these mice received injections of liposomes into the tail vein. Liposomes were prepared as described previously (Longmuir et al. 2006; Robertson et al. 2008) and contained a liver targeting peptide sequence derived from the circumsporozoite protein of Plasmodium berghei. The liposomes were made from a mPEG5000 conjugate of 22:1-PE, prepared by reacting 10 μmol di22:1-PE with 20 μmol of mPEG5000 carboxylic acid N-hydroxysuccinimidy l ester (#2M4M0H01, Nektar, Huntsville, AL) and 100 μmol of triethylamine in dichloromethane. An addition of 1.2-dipalmitoyl-sn-3-phosphatidylethanolamine, labeled with 1.4 nm gold particles obtained from Nanoprobes, Inc., Yaphank, NY, was used to enable localization of the liposome fragments using electron microscopy.

Immunocytochemistry

Serial sections of liver were cut on the cryostat and collected on Superfrost/Plus coated slides (Fisher Scientific, Pittsburgh, PA) and processed for immunocytochemistry for albumin, biotin, F 4/80, glial fibrillary acidic protein
(GFAP), or the endothelial cell marker CD-34. Slides with tissue sections were rinsed in Tris buffer three times and blocked for 1 h in 3% normal goat serum (Invitrogen, Carlsbad CA) (for albumin, F4/80, CD-34) or 3% normal rabbit serum (for biotin). Each primary antibody was tested parametrically, in dilutions of Tris buffer in blocking solution, to determine the optimal antibody concentration to be used. The albumin antibody (Bethyl Labs; Montgomery TX) labeled with FITC was used at 1:500; the biotin antibody (Abcam; Cambridge MA) labeled with FITC was used at 1:500; the macrophage (Kupffer cell) antibody F/4/80 (Serotec, Raleigh, NC) was used at 1:1,000; the GFAP antibody (Dako; Carpinteria, CA) was used at 1:3,000; and the CD-34 antibody (Vector Labs) was used at 1:100. Sections were exposed to primary antibodies at room temperature and in the dark, overnight (16–18 h). The following day, slides were rinsed in Tris buffer three times. The sections for F4/80 then were incubated for 2 h with either Alexa 488 or Alexa 546 goat anti-rat IgG at 1:1,000 (Invitrogen; Carlsbad CA), sections for GFAP were incubated with either Alexa 488 or Alexa 546 goat anti-rabbit, at 1:1,000 and sections for CD-34 were incubated with Alexa 488 goat anti-mouse at 1:1,000. Following incubation, slides were rinsed with Tris buffer and coverslipped with Vectashield anti-fade fluorescent mounting medium with DAPI; DAPI served as a fluorescent (ultraviolet, UV) stain for cell nuclei.

Lectin binding

Cryostat cut sections were mounted on slides and rinsed in Tris buffer. Tomato lectin (Vector labs; Burlingame, CA) labeled with FITC was diluted (1:5,000) in Tris buffer and sections were incubated overnight in the dark. Slides were rinsed in Tris buffer and coverslipped with Vectashield anti-fade fluorescent mounting medium with DAPI; DAPI served as a fluorescent (ultraviolet, UV) stain for cell nuclei.

Hematoxylin and eosin

Slide mounted 8–12 um cryostat sections were dehydrated through absolute alcohol and rehydrated to water. Slides were placed in hematoxylin stain for 4 min, rinsed in water, differentiated in 70% alcohol and stained in 0.01% eosin Y for 2 s, rinsed in 95% ethanol, dehydrated with absolute ethanol and cleared in xylenes for 15 min before coverslipping.

Electron microscopy

Liver tissue from nine animals was used for electron microscopic studies. Six of these animals had received intravenous injections of liposomes, as described above. Following perfusion with saline followed by 4% paraformaldehyde, tissue was post-fixed in 4% paraformaldehyde and 0.5% glutaraldehyde. Tissue blocks were cut on a Vibratome at 50 µm. Some sections were processed using a gold-enhancement kit (Goldenhance-EM; Nanoprobes Inc., Yaphank, NY) following the protocols described by the manufacturer. Other sections were processed for immunocytochemistry using the F4-80 or GFAP antibodies. Sections were then placed in 1% osmium tetroxide for 30 min, embedded in epoxy, cut at 800 nm, and stained with uranyl acetate and lead citrate to enhance contrast. These thin sections were inspected on a Phillips EM-10 electron microscope equipped with a Gatan digital camera.

Dissociated cell staining

Mice aged 30 days to adult were deeply anesthetized with sodium pentobarbital and perfused through the portal vein with 25 ml cold 0.1 M EDTA in calcium free DMEM followed by 25 ml warm (37°C) 0.05% collagenase II in DMEM with calcium. Liver pieces were removed, minced and incubated with the collagenase medium at 37°C for 15 min. Tissue was centrifuged at 1,500 rpm for 5 min, the collagenase medium was removed, and the cells re-suspended in DMEM. The tissue was dissociated and strained through a 100 µm mesh to eliminate clumps of cells. The cells were then stained with trypan blue for 2 min before photographing and counting using a bright field microscope.

Image collection and processing

Slides were examined using bright field illumination (H&E) or fluorescence illumination with rhodamine, fluorescein, or ultraviolet (DAPI) filter cubes. Digital images were captured using a Nikon DS 5M digital camera and imported into Adobe Photoshop. When creating photographic plates for illustrations, brightness and contrast were adjusted for uniformity within a plate; no other alterations of images were done.

Quantitative assessments

Relative numbers of each of the different populations of cells were estimated by counting the number of nuclei of immunocytochemically labeled cells and correcting the raw counts by using the method of Abercrombie (1946). Sections of 12 µm thickness, immunocytochemically labeled with a cell specific antibody and with DAPI nuclear stain, were examined. Digital images were collected using a 40× lens and imported into Photoshop. These images included an area of 46,800 µm² (260 µm × 180 µm). Cells that displayed clear immunocytochemical labeling along with distinct DAPI positive nuclei were identified and their
Positions marked on an overlaid sheet in Photoshop. Diameters of DAPI stained nuclei were measured using the Nikon DS-5M software for two point distances or from Photoshop images, using a reticule. The number of labeled cells (defined as DAPI stained nucleus amid immunocytochemically labeled cytoplasm) in each section was adjusted by the formula presented by Abercrombie:

\[ P = A \frac{M}{L + M} \]

in which \( P \) is the calculated average number of nuclei per region, \( A \) the crude count of number of nuclei of labeled cells per section, \( M \) the tissue section thickness, and \( L \) the average diameter of nuclei.

Numbers of immunocytochemically and DAPI labeled cells were determined for more localized regions of the liver, using the liver acinus scheme of Rappaport et al. (1954). Three zones of approximate equivalent areas were identified in a liver acinus, including zone 1 close to the portal area, zone 3, around the central venule, and a zone 2 situated between zones 1 and 3. Photoshop images were analyzed by placing a graticule, measuring 7,500 \( \mu m^2 \) (50 \( \mu m \times 150 \mu m \)), over each of the zones and determining the numbers of labeled cells in each area.

**Results**

Histological organization of normal mouse liver

Sections of mouse liver stained with a standard hematoxylin and eosin (H&E) procedure reveal patterns of cellular labeling similar to what has been reported for other mammalian species. Portal areas, containing elements of the hepatic triad, that is one or more small branches of the portal vein, a branch of the hepatic artery, and a small bile duct, along with lymphatic vessels and a very small amount of connective tissue, could be identified (Fig. 1a, c). The presence of portal areas along with central venules provided evidence of a lobular structure to the mouse liver.
Liver cells are arranged in plates or cords, and are seen radiating from the regions of central venules (Fig. 1a, b, d) and extending to the portal areas. The plates or cords of cells are separated by sinusoidal capillaries (Fig. 1g).

Tissue processed with H&E (or with the fluorescent DAPI stain, see below) reveals distinctly stained nuclei of varied shapes. Most common are cells having one, or sometimes two, large round to slightly oval nuclei. Also detected in the H&E stained, or DAPI labeled, sections are other cells with smaller and darkly staining oval or oblong nuclei.

Immunocytochemical techniques were used to identify different populations of cells within the liver tissue. As shown in Fig. 1d, cells containing the large round nuclei were labeled by an antibody to albumin. Albumin immunoreactivity appeared as small puncta of intense fluorescence within the cytoplasm of virtually all of these cells (Fig. 1d), although the numbers of puncta as well as their apparent intensity displayed some variation between cells. Because of their immunoreactivity to albumin antibodies, these cells with large round nuclei were deemed to be hepatocytes (Yokota and Fahimi 1981).

Figure 1e shows immunoreactivity to the F4-80 antibody, which is known to label macrophages in other organs and labels some of the cells containing dense oblong nuclei in liver sections. The F4-80 positive cells were distributed throughout the liver lobules but were always associated with a sinusoidal capillary. Within the liver lobules, F4-80 positive cells were more frequently encountered in regions close to the portal area (region 1 of Rappaport et al. 1954) and were more sparse in regions closer to the central venule (region 3). Because their morphological features and F4-80 immunoreactivity are similar to macrophages described in other species, these F4-80 positive cells were deemed to be Kupffer cells (Austyn and Gordon 1981; von Kupffer 1898).

Cells immunoreactive for glial fibrillary acidic protein (GFAP) also contained small oblong nuclei and also were distributed throughout the liver (Fig. 1f). The GFAP positive cells were encountered less frequently than were the F4-80 positive cells. These cells were deemed to be the stellate cells of Ito (Gard et al. 1985; Ito 1973; Neubauer et al. 1996).

Endothelial cells were identified by a combination of CD-34 immunocytochemistry and DAPI labeling of nuclei. As shown in Fig. 2a and b, the CD-34 immunoreactivity appears as a cell surface marker and clearly outlines sinusoidal capillaries in liver tissue. The CD-34 immunoreactivity is associated with small and oval or oblong DAPI stained nuclei (arrows) and clearly are distinct from the large round nuclei that are characteristic of the parenchymal hepatocytes. Figure 2c–e presents a case in which a mouse received an intravascular injection of small (0.03 μm) rhodamine labeled latex microspheres, and the tissue later sectioned and processed for CD-34 immunoreactivity. A comparison of Fig. 2c and d demonstrates that these smaller latex microspheres appear to label intensely the endothelial cells. Co-labeling of endothelial cells by green Alexa 488 CD-34 immunoreactivity, red rhodamine labeled latex microspheres, and blue DAPI labeled nuclei is illustrated in the higher magnification photomicrograph of Fig. 2f.

Further immunocytochemical characterization of liver cells

The issue of whether the immunocytochemical markers resulted in overlapping populations of cells was addressed. As was shown in Fig. 1, hepatocytes are labeled by immunoreactivity for albumin. Figure 3a and b show two views of the same section, with rhodamine optics to show immunoreactivity for albumin (Fig. 3a) and under fluorescein optics to show immunoreactivity for biotin (Fig. 3b). Similar to the pattern for albumin, biotin labeling appeared punctate and also appeared to label most cells containing large round nuclei (Fig. 3b). Figure 3c presents a merger of the images in 3a and 3b, along with an ultraviolet image showing the DAPI labeled blue nuclei, and demonstrates that virtually all cuboidal cells with large round nuclei are labeled with antibodies to albumin and also to biotin. These markers are found within the same cells, although the intracellular distribution of green biotin puncta was slightly different from the intracellular distribution of red albumin related puncta.

Albumin immunoreactivity was not found to co-express with immunoreactivity to F4-80, GFAP or CD-34

Photomicrographs in Fig. 3d–f demonstrate that F4-80 labeled cells are a separate population from the GFAP labeled cells. The GFAP antibody labels a population of cells with relatively small and oblong nuclei, and with slender and widely branching cellular processes (Fig. 3d). As noted above, these are the stellate cells identified by Ito (Fahimi 1982; Gard et al. 1985; Geerts et al. 1990; Ito 1973; Kawada 1997; Neubauer et al. 1996). The F4-80 antibody labels a population of cells (Bouwens et al. 1986; Naito et al. 1997; Widmann et al. 1972) that also displays small oblong nuclei, but these cells display widely branching but somewhat broader dendritic processes (Fig. 3d). As noted above, these are the stellate cells identified by Ito (Fahimi 1982; Gard et al. 1985; Geerts et al. 1990; Ito 1973; Kawada 1997; Neubauer et al. 1996). The F4-80 antibody labels a population of cells (Bouwens et al. 1986; Naito et al. 1997; Widmann et al. 1972) that also displays small oblong nuclei, but these cells display widely branching but somewhat broader dendritic processes (Fig. 3d). The photomicrographs of Fig. 3d and e are of the same section, processed for each antibody and using secondary antibodies with different fluorescent labels. When the two images are combined (Fig. 3f) it can be seen that the two labels are associated with distinctly different cells.

The issue of the distinct populations of cells labeled by F4-80 or by GFAP was further explored in animals that received intravascular injections of relatively large (0.5 μm) fluorescently labeled latex microspheres. These
larger microspheres do not label endothelial cells, as do the smaller (0.03 μm) microspheres as shown in Fig. 1, but do label macrophages throughout the body. In sections from the livers of these animals, the rhodamine (red) labeled microspheres could be seen contained within cell bodies. These sections were processed for immunocytochemistry. Figure 3g demonstrates that virtually all cells containing red microspheres were labeled with the F4-80 antibody, although not every F4-80 positive cell contained visible microspheres. In contrast, microspheres were not associated with GFAP labeled cells (Fig. 3h).

Double nucleated cells

The issue of determining whether these cells were double nucleated was addressed by studying immunocyto-chemically identified cell types along with DAPI staining of cell nuclei.

Figure 4a presents a merged image of a section taken using the fluorescein filter set to show tomato lectin binding, which labels the extracellular glycoprotein layer surrounding virtually all liver cells (McMillan et al. 1988), and the UV filter set to reveal DAPI labeling of nuclei. The white arrows indicate several figures that appear to be double nucleated cells. Figure 4b shows a similar section, but this section was also processed for immunocytochemical labeling of albumin in red. The white arrows indicate structures, bordered by tomato lectin, containing albumin, and also containing two DAPI stained nuclei. Analysis of 1869 albumin positive cells reveals the presence of double nuclei detected in the plane of the section of 455 of these cells, or approximately 24%. Nuclei of albumin or biotin
positive cells were round, with an average diameter of 9.8 μm. In general, the nuclei of single nucleated cells appeared to have slightly greater diameters than did the nuclei of double nucleated cells, although this was not a statistically significant difference.

In other cases (n = 4 animals), livers were prepared for dissociated cell suspensions and stained with Trypan blue. Figure 4c presents an example of several liver cells, some of which appear to contain only one nucleus (black arrows) while others contained two nuclei (white arrows). Counts of these preparations reveal that approximately 35% of the dissociated cells (259 of 743) are double nucleated. Further, evidence of double labeled hepatocytes was also obtained from electron microscopic studies of liver cells, as shown in Fig. 4d.

The possibility of other cell types containing double nuclei was examined using cell specific immunocytochemistry. Figure 4e presents an example of a merged image showing F4-80 positive cells with DAPI labeled nuclei. The nuclei of F4-80 positive cells typically were ovoid in shape, with a long axis that averaged 7.9 μm and a short axis of 4.5 μm. Of 355 F4-80 cells examined, there were no instances of cells with two nuclei. Similarly, Fig. 4f presents a merged image showing GFAP positive cells...
cells. Nuclei of GFAP positive cells also were ovoid or oblong in shape, with a long axis that averaged 6.3 μm and a short axis of 4.1 μm. Of the 147 GFAP positive cells that were examined, none showed evidence of more than one nucleus. Studies of CD-34 labeled endothelial cells with DAPI stained nuclei revealed that nuclei of these cells also displayed oblong shapes, with a long axis that averaged 7.2 μm and a short axis of 3.9 μm. The CD-34 positive cells showed no clear evidence of double labeled cells. However, the borders between adjacent endothelial cells were often difficult to discern, and so it is difficult to conclude that no double labeled endothelial cells exist.

Relative numbers of immunocytochemically identified cells

Numbers of immunocytochemically identified cells were determined for sets of 12 μm thick sections, viewed with a 40× lens, in an area of 46,800 μm² (260 μm × 180 μm), and the raw numbers corrected by the method of Abercrombie (1946). As shown in Fig. 5, an average of 33.4 cells with albumin positive cytoplasm and a DAPI labeled nucleus was calculated for this field region. The F4-80 positive cells comprised a smaller population, with an average of 12.6 cells per field. The GFAP positive cells were the least frequent population, with an average of 7.1 cells in each field. Endothelial cells, identified by a combination of DAPI labeled nuclei and the CD-34 antibody, were frequently encountered, with an average of 21.8 cells in each field. Wandering or temporarily ‘fixed’ leukocytes, including the ‘pit’ cells, were not included in this analysis. Because the Abercrombie (1946) correction factor was employed, it is possible to compare the numbers of cells labeled by each of these methods. Thus, the albumin positive cells constituted about 52% of all immunocytochemically labeled cells, the endothelial cells about 22% of
labeled cells, F4-80 cells were about 18% of all labeled cells, and the GFAP positive cells constituted about 8% of all labeled cells.

The above analysis was a summary of numbers of positively labeled cells across all fields of the liver. Other investigators studying liver from other mammalian species have noted that the distributions of some populations of cells vary between the three ‘zones’ that comprise the liver acinus (Rappaport et al. 1954). These include a region surrounding the portal area (see Fig. 1a) termed zone 1, a region surrounding the central vein, termed zone 3, and a region intermediate between the two, termed zone 2. Numbers of positive cells were counted in 7,500 \( \mu \text{m}^2 \) (50 \( \mu \text{m} \times 150 \mu \text{m} \)) areas in each of these zones. Numbers of albumin positive cells were remarkably consistent in different zones of the liver lobule. Numbers of F4-80 positive cells displayed variation between different regions of the liver, with more cells observed in zone 1 (4.5 per 7,500 \( \mu \text{m}^2 \)) than in zone 2 (3.9/7,500 \( \mu \text{m}^2 \)) or zone 3 (3.7/7,500 \( \mu \text{m}^2 \)). Numbers of GFAP positive cells were quite variable, with no consistent difference observed between the different zones. Numbers of CD-34 positive endothelial cells did not vary between regions.

Fine structural features of liver tissue

An analysis of the fine structural features of hepatocytes and non-parenchymal cells was undertaken using electron microscopic techniques. Figure 6a presents a low magnification electron micrograph of an hepatocyte, showing a typical large round cell nucleus along with cytoplasmic organelles including both smooth (SER) and rough (RER) endoplasmic reticulum and many mitochondria (M). Dense glycogen granules appear distributed throughout the cytoplasm. The basal surface of the hepatocyte, neighboring the sinusoidal capillary space, displays a profuse elaboration of microvilli in the space of Disse (D). The space of Disse is situated just under the endothelial layer of the sinusoidal capillary. Figure 6b presents a higher magnification view of the space of Disse, and also shows a fragment of an endothelial cell with a fenestration (F) in the endothelial lining.

Adjacent hepatocytes are separated by a narrow intercellular space with occasional intercellular junctional complexes. The intercellular space widens to form canaliculi (Fig. 6c), and occluding junctional complexes adjacent to the canaliculi apparently separate the lumen of the canaliculi from the normal intercellular space, as shown in Fig. 6c. Golgi apparatus was seen occasionally in the cytoplasm adjacent to the canaliculi.

Non-parenchymal cells were also detected with electron microscopic methods, as illustrated in Fig. 7. Kupffer cells were encountered frequently, either situated upon underlying endothelial cells or as part of the lining of the sinusoidal capillaries. Figure 7a presents an electron micrograph of tissue that had been processed immunocytochemically for F4-80. The immunoreactivity is visible as dark staining on the plasmalemma, which displays folds and pseudopodia. Kupffer cells display large numbers of lysomes, several of which can be seen in the cell in Fig. 7a. Kupffer cells are also shown in Fig. 7d, where adjacent membranes between Kupffer and endothelial cells display no specialized contacts, and in Fig. 7G, where uptake of gold labeled liposomes is demonstrated by the many gold particles in the Kupffer cell cytoplasm, but not in the nucleus.

Ito stellate cells could be identified by the presence of prominent intracellular lipid droplets and filamentous material, as illustrated in Fig. 7b. These stellate cells were situated between endothelial cells and hepatocytes or were intercalated between hepatocytes (as in Fig. 7b).

Endothelial cells were identified by their elongated and flattened nuclei, and by the presence of fenestrations (Fig. 7c) in the cytoplasmic processes that form much of the lining of the sinusoidal capillaries. These fenestrations have diameters of approximately 125–175 nm. Fenestrations did not include membranous diaphragms over the openings; only sparse evidence of a very rudimentary basal lamina was seen associated with endothelial cells. At sites where cell processes from apparent adjacent endothelial cells meet, specialized adhesive intercellular junctional complexes are seen (Fig. 7c). Adhesive contacts are not seen between endothelial cells and hepatocytes or any other non-parenchymal cell. Finally, pit cells are occasionally seen, as illustrated in Fig. 7f.

![Fig. 5 Histograms summarizing results of counts of immunocytochemically identified cells with DAPI stained nuclei. The Y axis shows numbers of cells detected in a region with an area of 48,000 \( \mu \text{m}^2 \), and thickness of 12 \( \mu \text{m} \). Calculated numbers were adjusted using the method of Abercrombie (1946). Data are mean ± SEM.](image-url)
Discussion

Overview

The data presented here provide an overview of the histological and fine structural organization of the mature mouse liver. It is clear that the fundamental features of the mouse liver are remarkably similar to the features of liver from other mammalian species. On the one hand, the clear similarities may detract from the noteworthiness of this manuscript; the mouse liver displays the same cellular and subcellular constituents as do the livers of other mammalian species. On the other hand, however, the data presented in this manuscript are vitally important for the interpretation and understanding of other results, using either wild type or genetically engineered mice, and provide a solid reference point against which other data can be interpreted.

Although the very sparse connective tissue within the mouse liver makes recognition of its lobular structure (e.g., Rappaport et al. 1954) a challenge, the major features of the lobules, including the positions of portal areas and central venules, are recognizable (Fig. 1a, c). The types of cells that comprise the mouse liver are similar to those that have been described in other species. The prominent cell type is the hepatocyte, characterized by the presence of intracellular albumin (Bemuau et al. 1985) or biotin. Also found are phagocytic Kupffer cells (von Kupffer 1898), labeled with the F4-80 antibody (Naito et al. 1997), the numbers of which are approximately 35–40% of the number of hepatocytes. Also studied were the Ito stellate cells (Gard et al. 1985; Ito 1973; Neubauer et al. 1996), whose numbers are about 8–10% of the number of hepatocytes. As with any organ, endothelial cells form much of the lining of the sinusoidal capillaries. Although the thin squamous endothelial cells do not contribute a great deal to the volume of tissue in the liver, the number of nuclei is approximately 22% of all liver nuclei and approximately 40% of the number of hepatocytes. The relative proportions of these cells are similar to figures reported for other species (Bouwens et al. 1986; Geerts 2001; Marcos et al. 2004; Naito et al. 1997; Sato et al. 2003). Further, the biliary system of the mouse liver appears similar to that of other mammalian species, with the presence of canaliculi between adjacent hepatocytes and small bile ductules that form part of portal areas. Thus, although the present investigation did not reveal novel features of liver organization in the mouse, the value of this study is that it establishes baseline data for the mouse and demonstrates the similarities of the histological organization of the mouse liver and of other mammalian species. These data underscore the value of the mouse model for laboratory studies of liver structure and function.

Liver cell types

Parenchymal cells in the liver are the hepatocytes, while non-parenchymal cells include the Kupffer cells, Ito

Fig. 6 Electron micrographs of liver tissue. a Relatively low magnification electron micrograph showing a portion of an hepatocyte. A large round nucleus of the hepatocyte (HN) is seen in the lower part of the figure. Many profiles of rough endoplasmic reticulum (RER) can be seen throughout the cytoplasm, often intermixed with mitochondria (M). The space of Disse (D) can be seen at the basal surface, just below processes of endothelial cells that border the lumen (Lu). b Relatively high magnification electron micrograph shows a fenestration (F) in the endothelial cell (E) lining, over the space of Disse (D). c Higher magnification electron micrograph illustrating a canaliculus, and adjacent occluding junctions (arrows)
stellate cells, wandering leukocytes (including a population of lymphocytes, termed the ‘pit’ cells (Bouwens et al. 1992), and endothelial cells.

Different cell types in the liver were identified based upon their morphological features and by their immunoreactivity. Hepatocytes have long been recognized in H&E stained material as round or cuboidal cells, and recent studies have demonstrated that they contain mRNA for expressing the albumin gene and also are immunoreactive for albumin (Bernuau et al. 1985; Yokota and Fahimi 1981). The present results from mouse are fully compatible with these previous reports. The present results indicate that albumin positive hepatocytes are encountered approximately three times more frequently than are Kupffer cells, and about six to seven times more frequently than Ito stellate cells.

Further, the present studies demonstrate that hepatocytes also contain biotin. Biotin appears not to be synthesized by hepatocytes but rather is taken up from food or produced by intestinal flora (Bowman and Russell 2006), and is stored in the liver to support its role in metabolism. The presence of biotin within hepatocytes clearly is important for considerations of its metabolic role, but may also have importance in issues of immunocytochemical methodology. Immunocytochemical studies that use avidin–biotin binding as a step in localizing some antigen of interest may
yield false positive results due to avidin binding to endogenous biotin rather than to biotinylated secondary antibodies (Ramos-Vara 2005).

The fine structural features of mouse hepatocytes are similar to those reported for other species. Large round nuclei, and occasionally two nuclei, are prominent. Much of the cytoplasm is occupied by mitochondria, both smooth and rough endoplasmic reticulum, and scattered glycogen particles. The basolateral portion of the hepatocyte, which faces the sinusoidal capillaries, displays a rich microvillous elaboration of the plasmalemma; this microvillous border occupies much of the space of Disse. As in other mammalian species, the apical portion of hepatocytes is associated with canaliculi. Canaliculi are formed by a widening of the intercellular space, although the canicular lumens are set off from the intercellular space by occluding junctions. Small profiles of Golgi apparatus were seen occasionally in the apical cytoplasm close to canaliculi, but we did not detect secretory vesicles associated with canaliculi. The canaliculi are believed to form an anastomosing system, which lead eventually through ducts of Hering into small bile ductules in the portal areas. These small bile ducts are a consistent feature in the portal areas.

Monocyte derived macrophages are found in virtually every organ and tissue of the body, and comprise the diffuse reticulo-endothelial system (Aschoff 1924; Furth et al. 1972). In the liver these macrophages are termed Kupffer cells (von Kupffer 1898). Although originally, these phagocytic cells were likely confused with the stellate cells that would later be identified by Ito (1973), later studies demonstrated that Kupffer cells can be identified by their ability to phagocytose tracer substances, including carbon, India ink, or latex microspheres, and also by their immunoreactivity to the F4-80 antibody. The present studies demonstrated the presence of F4-80 positive Kupffer cells in mouse liver, and electron microscopic studies demonstrated this antibody labels a cell surface marker. Further, the results from studies of double labeling with fluorescently labeled latex microspheres and also from immunocytochemistry have demonstrated conclusively that the Kupffer cells are a population of cells distinct from the Ito stellate (fat storing) cells.

The use of latex microspheres of different diameters was useful in demonstrating that Kupffer cells could be labeled specifically with larger (0.5 μm) microspheres, while smaller microspheres (0.03 μm) labeled both Kupffer cells and endothelial cells, as demonstrated previously (Wake et al. 1989).

Other investigators (Bouwens et al. 1986, 1992; Sleyster and Knook 1982) have noted that Kupffer cells are not distributed homogenously in the liver and appear to show some variation in regard to their phagocytic activity. These authors (Sleyster and Knook 1982; Bouwens et al. 1992; Bouwens et al. 1986) have reported that Kupffer cells are more frequently encountered and also are larger in regions around the portal areas than around the central venules. The present data corroborate this finding in the mouse, although the regional differences in the mouse liver appear not as great as the regional differences reported for rat liver.

Ito stellate cells are fat storing cells of the liver, and in the present studies these were identified by immunoreactivity to glial fibrillary acidic protein (GFAP) (Gard et al. 1985; Neubauer et al. 1996). Stellate cells are identifiable by their fine structural features of prominent intracellular lipid droplets and by cytoplasmic filamentous material. The intracellular filamentous material likely forms the basis of their immunoreactivity to GFAP and to desmin (Leo and Lieber 1983; Yokoi et al. 1984; de Bleser et al. 1991). Quantitative estimates (Leo and Lieber 1983; Yokoi et al. 1984; de Bleser et al. 1991) indicated that numbers of Ito stellate cells in rat liver were about 10–12% of hepatocytes, a fraction similar to the present results in mouse. Further, de Bleser et al. (1991) reported that stellate cells were found more frequently in peri-portal areas than in peri-central areas. The present study noted that stellate cells were not distributed homogenously throughout the liver, but a consistent pattern between peri-portal and peri-central regions was not detected.

Endothelial cells are an important cell type in any organ, and certainly so in the liver. Liver endothelial cells are specialized, with the presence of fenestrations that appear aggregated into groups that form ‘sieve plates’ (Wisse 1972). The very sparse nature of a basal lamina beneath the endothelial cells, along with the absence of diaphragmatic coverings of the fenestrations, allows for apparent relatively free movement of small molecules (less than 125 nm diameter) between the space of Disse and the capillary lumen. Wisse has demonstrated both small bristle-coated micropinocytotic vesicles and large smooth macropinocytotic vesicles in the endothelial cells. Small latex microspheres, diameter of 30 nm, are not detected in hepatocytes after intravascular injection, although they do appear to label endothelial cells. This suggests that the latex microspheres are taken up by endothelial vesicles, but either do not reach the space of Disse or are not taken up by the hepatocyte microvillous border in the space of Disse.

Double nucleated cells

The present analysis indicated that a considerable portion of hepatocytes are double nucleated. The percentage gleaned from studies of tissue slices is likely to be an underestimate, however, because although two nuclei in the X or Y planes would be detected easily, if a second nucleus was obscured in the Z plane or found within an
adjacent section, it would be missed by the techniques employed in this study. Another approach taken in the present studies was to dissociate liver cells, and use the Trypan blue stain to detect double nucleated cells. This analysis yielded higher numbers, of approximately 35% double nucleated cells. These results are in agreement with other recent studies (Wheatley 1972; Gupta 2000) that have reported that the population of hepatocytes includes many double nucleated cells. The functional significance of double nucleated cells, however, is not clear. No evidence was revealed to indicate that any other cell type within the liver included double nucleated cells. Analysis of tissue sections immunolabeled with F4-80 for Kupffer cells or with GFAP for Ito stellate cells revealed no evidence of more than one DAPI stained nucleus. Kupffer and stellate cells probably were not detected in the dissociated cell studies, as different steps in the cell dissociation process need to be taken to include these cell populations (Smetsrod et al. 1985; Riccalton-Banks et al. 2003). The thin squamous endothelial cells display extensive spread of cytoplasm and membranes, making the detection of two nuclei within one cell a very difficult endeavor. We are not aware of any examples of double nucleated endothelial cells, and believe they are not likely to exist.

Conclusion

Genetically engineered mice will play a very important role in future studies of liver function, and so it is vitally important to have baseline reference information on the cellular makeup of normal mouse liver. The present paper, using histological, quantitative immunocytochemical, and fine structural analyses, demonstrates that the cellular organization of the mouse liver is quite similar to that of other mammalian species, confirming that the mouse presents a useful animal model for studies of liver structure and function.

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