Visualization of hepatobiliary excretory function by intravital multiphoton microscopy

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1 Introduction

One of the major hepatobiliary functions is the uptake, processing, and excretion of a variety of endogenous or exogenous organic anions by the hepatocytes. The anions are obtained from the sinusoids and conjugated with glutathione, glucuronate, or sulfate in hepatocytes, and then excreted by an ATP-dependent mechanism into the bile canaliculi through the apical membrane of hepatocytes. Conventional methods to investigate these processes usually involved the biochemical analyses of the contents in the liver, bile, or other body.
fluids such as blood or urine. Real-time imaging of hepatobiliary excretion has been approached on *ex-vivo* liver tissues by laser confocal microscopy. However, the *ex-vivo* model cannot be used to investigate the *in-vivo* dynamics of hepatic activities. Therefore, the establishment of an intravital imaging system aimed at following these functional processes may be of tremendous value in hepatology.

In this work, we report the design and application of an intravital hepatic imaging chamber on the mouse for the *in-vivo* investigation of the hepatobiliary excretory function of 6-CFDA.

### 2 Methods

#### 2.1 Design of the Intravital Hepatic Imaging Chamber

Our hepatic imaging chamber is composed of two doughnut-shaped titanium rings: the outer and inner lids [Figs. 1(a) and 1(b)]. The outer and inner diameters of the outer lid are 12.5 and 6.5 mm, respectively, with a thickness of 1 mm. For the inner lid, the outer and inner diameters are 11.5 and 6.5 mm, respectively, and it has a thickness of 0.8 mm. Both lids are threaded and can be fastened together by simple screwing.

#### 2.2 Installation of the Hepatic Imaging Chamber on the Mouse

C57BL/6 mice beyond 5 weeks of age were anesthetized using intraperitoneal injection of 2-2-2 tribromoethanol at a dose of 0.35 mg/g. The hairs on the abdomen were shaved and a vertical incision of the skin and peritoneum was made. To visualize the sinusoids, the inferior vena cava was exposed for the injection of 10 mg/mice of rhodamine B isothiocyanate-dextran 7000 (Sigma, Saint Louis, Missouri). The incision of the lower abdomen was sutured, leaving the upper part open to create a circled wound to position the inner lid of the hepatic imaging chamber inside the abdominal wall. Prior to installing the inner lid, an 8-mm round cover glass was adhered around the outer rim by a polyvinylacetate glue. The cover glass serves as the observation window of intravital hepatic activities. The inner lid was then sutured through the
small holes on the ring to the skin and peritoneum. To ensure tight contact of the liver surface with the cover glass and to minimize motional artifacts associated with heartbeat and respiration, the adhesive such as tissue adhesive Histoacryl (B. Braun Melsungen AG, Germany) was applied to the edge of the inner lid to adhere the liver to the imaging chamber. For gluing purposes, we tested a number of adhesives. While some such as fibrin did not glue well, others appeared to cause changes in the appearance of the liver surface not all adhesives were tested under ex-vivo conditions for liver appearance changes. Therefore, the significant criteria of glue selection rest on how well the adhesive can attach the liver to the chamber and the glue’s biocompatibility. For comparison purposes, we also acquired images of the hepatic metabolic activities of 6-CFDA using for medical device, Henkel Loctite, Rocky Hill, Connecticut and instant adhesive, Henkel Loctite. In all cases, the tissue adhesive was not applied to the imaged regions. Finally, the outer lid is attached to the inner lid by screwing [Figs. 1(c) and 1(d)].

2.3 Accommodation of the Mouse onto the Microscopic Stage

The positioning of the mouse installed with the hepatic imaging chamber and the subsequent imaging procedures are illustrated in Fig. 2. The mouse was kept in the supine position with the hepatic imaging chamber fitted into the U-shaped groove of a steel plate [Fig. 2(a)], and the plate was slid toward jaw of the mouse and lodged the liver window device to the terminus of the groove [Fig. 2(b)]. The mouse with the steel plate was then placed on the microscopic stage of an inverted microscope (TE2000, Nikon, Japan) by screwing the plate onto the microscopic stage [Fig. 2(c)]. In this manner, the mouse can be firmly attached for multiphoton imaging purposes. A hot pack between the gauze was put on top of the mouse for warming purposes [Fig. 2(d)].

2.4 Multiphoton Microscopy

The multiphoton microscope is illustrated in Fig. 2(e). A titanium-sapphire laser with 780-nm output (Tsunami, Spectra Physics, Mountain View, California) pumped by a diode-pumped solid-state laser (Millennia X, Spectra Physics) was used for excitation. The laser was scanned by an x-y mirror scanning system (Model 6220, Cambridge Technology, Cambridge, Massachusetts) and guided toward the modified inverted microscope. The laser was beam expanded and reflected into the back aperture of a long working distance (7.4 mm) objective (Plan Fluor ELWD 20×, NA 0.45, Nikon) by a primary dichroic mirror (700DCSPXRUV-3p, Chroma Technology, Rockingham, Vermont). The power at the sample was around 21 mW, and sample luminescence
was collected in the epi-illuminated or backscattering geometry. After passing through the primary dichroic mirror, the second harmonic generation (SHG) and fluorescence signals are separated into four simultaneous detection channels by secondary dichroic mirrors (435DCXR, 495DCXR, 555DCLP, Chroma Technology) and additional bandpass filters (HQ390/20, HQ460/50, HQ525/50, HQ590/80, Chroma Technology). The detection bandwidths for the SHG, blue, green, and red fluorescence are 390±10, 460±25, 525±25, and 590±40 nm, respectively. In our studies, the SHG signal was collected in the backscattering geometry. Single-photon counting photomultiplier tubes (R7400P, Hamamatsu, Japan) were used as optical detectors. Each optical scan is composed of 256×256 pixels and took approximately 4 s to complete. For image processing, we used the software of ImageJ (National Institute of Health, Bethesda, Maryland) and MetaMorph (Universal Imaging Corporation, Downingtown, Pennsylvania). Right before the visualization of hepatobiliary excretory dynamics, we would inject 50 μg/mouse of 6-carboxyfluorescein diacetate (6-CFDA, Sigma) either through the tail or jugular veins, and began to acquire the serial multiphoton images at 4-s intervals.

The procedures of the animal experiments were approved by the Institutional Laboratory Animal Care Committee of National Taiwan University, College of Medicine.

### 3 Results

Figures 3 and 4 shows the representative multiphoton images acquired using tissue adhesive Histoacryl. Depth for imaging hepatobiliary function was set at approximately 30 μm below the disappearance of the SHG signals from the capsule. Upon injection of 6-CFDA through the jugular vein, the sinusoids already demarcated by the red fluorescence. In addition, the scattered intensely fluorescent spots (5 to 15 μm) on the margins of the sinusoids were visible [Fig. 3(a)]. These structures are identified as the stellate cells that contain vitamin A emitting strong autofluorescence. Following 6-CFDA injection, green fluorescence starts to increase in the hepatic cords between the sinusoids and peaks at the 8-min point [Figs. 3(b)–3(d)]. The green fluorescence intensity then starts to decrease and almost reaches the background level at approximately 50-min post-6-CFDA injection [Fig. 3(f)].

Soon after finishing the time-course observations at approximately 50-min post-6-CFDA injection, we imaged the same microscopic field at different depths of 5, 20, 30, and 50 μm [Figs. 4(a)–4(d)]. The green fluorescence visible in the bile canaliculi at earlier time points is no longer apparent. This observation is consistent with the fact that the uptake, processing, and excretion of 6-CFDA is now complete.

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**Fig. 3** Representative intravital multiphoton hepatic images acquired using Histoacryl. (a) through (f) are the time-lapsed multiphoton images acquired approximately 30 μm below the capsule. Time in minutes indicates period post-6-CFDA injection. Small arrow in (a) indicates the presence of stellate cells (blue), while large arrow indicates sinusoid positions (red). Arrow in (b) indicates hepatocytes in cords emitting CF fluorescence (green), and arrows in (c), (d), and (e) indicate CF fluorescence in bile canaliculi.

**Fig. 4** Representative intravital multiphoton hepatic images acquired using Histoacryl. Depth-dependent multiphoton images acquired approximately 50 min after 6-CFDA injection. The stellate cell (small arrow) and sinusoid (large arrow) are clearly visible in (c).
The kinetics of the mean green fluorescence intensity of the liver is shown in Fig. 5. Also shown are the intensity profiles (6-CFDA jugular vein injection) acquired using the adhesives of 4011 and 406. As the figure demonstrates, although the intensity profiles obtained using the three adhesives show different temporal dependence, hepatic metabolism of 6-CFDA is mostly complete at the 50-min mark. Since we only applied the adhesives at the chamber edge, we feel that the effects of glue on hepatic metabolism are minimal. In addition, since the intensity profile of 406 decayed the fastest, the biocompatibility of the glues appear not to be a factor in affecting the hepatic metabolic activities of 6-CFDA. Therefore, we feel that the difference in measured intensity profiles may be due to variations in the individual mouse metabolism and the variations in 6-CFDA injection procedures.

4 Discussion

Multiphoton microscopy offers several advantages in the intravital imaging of living tissues. It is superior to videomicroscopy in its high depth-discrimination nature and also in providing simultaneous multicolor observation of tissue specimens. Multiphoton imaging is also superior to confocal microscopy due to the increased imaging penetration depth and markedly reduced phototoxicity. In addition, SHG microscopy can also be used to detect collagen fibers in the tissues without the addition of fluorophores. Intravital multiphoton microscopy has been used in the investigations of renal physiological and pathological processes, but to the best of our knowledge, it has never been applied in the hepatology field. In this work, we have successfully demonstrated the monitoring of hepatic metabolic activities by the use of multiphoton microscopy and an intravital hepatic imaging chamber.

During the course of this work, we found that the vibration of liver tissues caused by respiration and heartbeat can be minimized by gluing the edges of the inner lid of the hepatic imaging chamber to the underlying liver with tissue adhesive. The adhesive-free regions of the liver were used for imaging purposes. The preadministered rhodamine dextran stays in circulation for as long as 5 h and thus the sinusoids and venules in the liver are well demarcated by its red fluorescence. 6-CFDA can be taken up by the hepatocytes and then be hydrolyzed by esterase into carboxyfluorescein (CF), which would emit at 517 nm. After additional hepatocyte processing, CF is then excreted into bile. Our data clearly showed the sequential illumination by CF of hepatocytes and then bile canaliculi, indicating the hepatobiliary excretion of this organic anion. After the microscopic examination, the animals might, carrying the hepatic imaging chembron their abdomen, live as usual, needing no restriction on activities.

These results support the use of our methodology to be a powerful technique to investigate the hepatobiliary excretory functions in diseases, such as extrahepatic and intrahepatic cholestasis. In addition, hepatic metabolism of chemicals and drugs over an extended period of several days may be investigated with our approach. The application of imaging over a large tissue area may also open a way to observe acute or chronic liver injuries, or follow targeted cell migration and proliferation in complex liver environments.

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