Intravital multiphoton microscopy can model uptake and excretion of fluorescein in hepatic ischemia-reperfusion injury

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Abstract. The liver is important in the biotransformation of various drugs, where hepatic transporters facilitate uptake and excretion. Ischemia-reperfusion (I/R) injury is a common occurrence in liver surgery, and the developing oxidative stress can lead to graft failure. We used intravital multiphoton tomography, with fluorescence lifetime imaging, to characterize metabolic damage associated with hepatic I/R injury and to model the distribution of fluorescein as a measure of liver function. In addition to measuring a significant increase in serum alanine transaminase levels, characteristic of hepatic I/R injury, a decrease in the averaged weighted lifetime of reduced nicotinamide adenine dinucleotide phosphate was observed, which can be attributed to a changed metabolic redox state of the hepatocytes. I/R injury was associated with delayed uptake and excretion of fluorescein and elevated area-under-the-curve within the hepatocytes compared to sham (i.e., untreated control) as visualized and modeled using images recorded by intravital multiphoton tomography. High-performance liquid chromatography analysis showed no differences in plasma or bile concentrations of fluorescein. Finally, altered fluorescein distribution was associated with acute changes in the expression of liver transport proteins. In summary, multiphoton intravital imaging is an effective approach to measure liver function and is more sensitive in contrasting the impact of I/R injury than measuring plasma and bile concentrations of fluorescein. © 2013 Society of Photo-Optical Instrumentation Engineers (SPIE) [DOI: 10.1117/1.JBO.18.10.101306]

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1 Introduction

A growing interest in intravital imaging has evolved due to limitations of conventional techniques in measuring dynamic real-time events. Intravital microscopy provides information about molecular and cellular processes with high spatial and temporal resolution in vivo. Imaging of live animal organs has been going on for 80 years, but has largely increased in the last decade with the improvement of conventional microscopes, especially nonlinear excitation, which has allowed deep tissue imaging.

Multiphoton tomography (MPT) is the most popular application of fluorescence microscopy for imaging thick tissue and live animals, producing high-resolution three-dimensional images. It involves the excitation of two (or more) photons of lesser energy normally produced by single photon absorption of higher energy. MPT is particularly well suited for in vivo studies of drug distribution in biological tissues, as it reduces tissue damage and enhances image quality due to reduced photobleaching and increased penetration depth compared to conventional fluorescence microscopy. Fluorescence lifetime imaging microscopy (FLIM) in combination with MPT can identify fluorescent probes with overlapping spectral properties and measure lifetime specific for the probe and the surrounding environment. For example, autofluorescent nicotinamide adenine dinucleotide (NADH) is a direct measure of cellular metabolic state and can be used to study cellular damage. The free and bound forms of NADH are involved in the anaerobic and aerobic production of adenosine triphosphate (ATP), respectively. Accordingly, a change in the free/bound ratio gives insight into the metabolic state of a cell. In a previous study, our group showed a decreased metabolic activity as shown by an increased ratio of free/bound NADH in ischemia-reperfusion (I/R) injury in the liver.

I/R injury is a common phenomenon in liver transplantation and resection, with morbidity and mortality as complications. During liver transplantation, ischemic damage occurs when blood flow is cut off to the donating liver, and the injury is further intensified when the liver is transplanted into the recipient. The reason for reperfusion injury is the reintroduction of oxygenated blood, which results in oxidative stress, causing increased microvascular permeability, intracellular ATP loss, inflammatory cell infiltration, and, potentially, cell death.
As a consequence, oxidative stress may lead to graft failure and rejection of the liver. Cellular damage occurring in I/R injury have been shown to affect membrane proteins, which could affect distribution of drugs in the liver.

Hepatic transporters are membrane proteins located on the sinusoidal, lateral, or canalicular membranes, facilitating uptake and excretion of many endogenous and exogenous substances. Fluorescein is a fluorescent probe and a substrate for the uptake transporters organic anion transport polypeptides (OATPs). It is metabolized to fluorescein monoglucuronide. Both fluorescein and fluorescein monoglucuronide are excreted into the bile by multidrug resistance-associated protein 2 (MRP2). It is not known if they are substrates for MRP3, an export transporter located on the lateral membrane of the cell, transporting substrate back to the blood (Fig. 1). However, other members of the fluorescein family, such as 5(6)-carboxy-2'-7'-dichloro-fluorescein and 5(6)-carboxyfluorescein, are secreted into the blood by MRP3; thus, it is likely that fluorescein and fluorescein monoglucuronide also interact with this transporter.

The aim of this study was to investigate the transport of fluorescein in the liver following I/R injury by intravitral multiphoton tomography, correlating these results with changes in biliary excretion, plasma concentrations, and expression of the transporters.

2 Materials and Methods

2.1 Chemicals and Reagents

Sodium fluorescein and β-glucuronidase were purchased from Sigma Aldrich (St. Louis, MO). Ilium xylazil and ketamine hydrochloride were obtained from Bayer Australia (Pymble NSW, Australia) and Parnell Laboratories, Australia, respectively. Reagents used for western blot analysis were purchased from Bio-rad (Protein std precision plus BIO-Rad, IMMuno-blot membrane PVDF, 10× TGS Buffer, 10× TG Buffer, 30% Acrylamide/Bis Solution, 29:1, TEMED and Ammonium Persulfate); Thermo Fisher (Supersensitive West Femto Maxi Sensitivity, Pierce BCA Protein Assay Kit, Dental Developer replenisher, Dental fixer replenisher and T-mat g film); and Whatman (Whatman filter paper). MRP2 monoclonal [M2 III-6] antibody (catalogue number AB3373) was purchased from Sapphire Bioscience (Sydney NSW, Australia); OATP2 (catalogue number AB3572P) was purchased from Merck Millipore (Kilsyth VIC, Australia); and rabbit antibody MRP3 from Sigma Aldrich. The reagents used in real-time polymerase chain reaction (RT-PCR) were obtained from Invitrogen to Life Technologies Australia (Mulgrave, Victoria, Australia) (Sybr master mix) and Sigma Aldrich [quantitative PCR (qPCR) primers].

2.2 Animals

Male Wistar rats, weighing 320 ± 20 g, purchased from the University of Queensland Biological Resources, were used in all experiments. All studies were approved by the Animal Ethics Committee at the University of Queensland. Rats were housed at the University of Queensland in Pharmacy Australia Centre of Excellence, where the temperature was maintained at 20 ± 1°C and humidity at 60 to 75%, with artificial light for 12 h (7 a.m. to 7 p.m.) daily. All animals had unlimited access to food and water.

2.3 Surgical Procedures and I/R Model

Rats were anaesthetized initially by an intraperitoneal injection of ketamine hydrochloride (80 mg/kg) and xylazine (10 mg/kg). Anesthesia was maintained throughout the experiment by administering ketamine (2.2 mg/100g) and xylazine (0.25 mg/100g). Body temperature was controlled by placing rats on a heating pad set to 37°C. Ischemia was induced locally in approximately 70% of the liver, by clamping the portal vein, hepatic artery, and bile duct supplying the median and left lobes using a microvascular clamp. After 60 min, the clamp was removed to allow reperfusion of the liver for 4 h and rats were allowed to recover during this time under close monitoring. Sham rats (i.e., untreated controls) underwent the same procedure without clamping the vessels. After 3 h of reperfusion, rats were again anaesthetized. The jugular vein and carotid artery were cleared of surrounding tissue and cannulated using polyethylene (PE)-tubing for administration of sodium fluorescein (intravenous bolus 10 mg/kg at 4 h of reperfusion) and blood collection. A midline laparotomy was performed to expose the liver and to cannulate the bile duct, which was cleared of surrounding fat tissue and cannulated using PE-tubing. The left lobe was slightly raised above the intraperitoneal cavity to allow imaging of fluorescein distribution. The liver and intraperitoneal cavity were kept moist by continuous administration of 0.9% saline throughout the experiment.

2.4 Tissue Collection

Blood (0.2 mL) was collected during the surgery from the inferior vena cava using a 30 gauge needle and plasma concentration of alanine transaminase (ALT) was measured using a Hitachi 747 analyzer (Hitachi Ltd., Tokyo, Japan). Blood for fluorescein determination was collected at 5, 10, 30, 60, 120, and 180 min from the carotid artery. Bile was collected in pre-weighted tubes every 10 min for 1 h and every hour after that for 3 h. At the end of the experiment, the left and median lobes were excised and portions were snap frozen in liquid nitrogen and stored at −70°C for later analysis with western blot and RT-PCR.

2.5 Multiphoton Intravital Imaging

MPT images were recorded using a DermalInspect system (JenLab GmbH, Jena, Germany) equipped with an ultrashort (85 fs pulse width) pulsed mode-locked 80-MHz Titanium: Sapphire MaiTai laser (Spectra Physics, 25 Mountain View,
USA). The excitation wavelength was set to 740 nm for liver autofluorescence and 920 nm for fluorescein fluorescence, with an emission signal range of 350 to 650 nm established through the use of a BG39 bandpass filter. Images were recorded with 10× (low magnification) or 40× (high magnification) objectives. The laser power was set to 15 or 20 mW and the acquisition time for obtaining the images was 13.4 or 7.4 s per frame for high- and low-magnification imaging, respectively. Each image was 179 × 179 μm wide at a resolution of 512 × 512 pixels. The left lobe of the liver was placed on a metal plate, attached to an adjustable stand that could be elevated or lowered as required. The plate was slightly raised above the intraperitoneal cavity to reduce movement in the organs underneath. Images were recorded before injection of fluorescein and then continuously throughout the experiment up to 180 min.

FLIM data were collected with a time-correlated single-photon counting (TCSPC) SPC-830 detector (Becker & Hickl GmbH, Berlin, Germany) integrated into the MPT system. The TCSPC module assembles a photon distribution of the scan area across the x and y coordinates taken from four photon counters, with only three used for this study. The module determines the time of arrival for each photon detected within the fluorescence decay. Fluorescence emission was spatially resolved between three linearly arranged photon counters through the use of three dichroic filters in the beam path, spectrally dividing the emission light into three channels for each photon detected within the fluorescence decay. Fluorescence emission was spectrally resolved using a standard curve prepared in homogenized liver. The area under the curve (AUC) was calculated as

\[ AUC = \alpha_1 + \alpha_2 \]

where \( \alpha_1 \) and \( \alpha_2 \) are amplitudes, respectively. Each FLIM image was collected at an exposure of 13.4 s and acquisition image size of 214 × 214 μm.

2.6 Image Processing

All image processing was done in ImageJ 1.44p (National Institutes of Health, USA). To trace the distribution of fluorescein in blood and hepatocytes, these areas were selected in the images recorded by low magnification. For analysis of fluorescein distribution in bile, this area was selected in images recorded by high magnification. The profile of fluorescein concentration (in sinusoid or hepatocyte) versus time was fitted by an empirical triexponential model.

\[ C = A_1 e^{-B_1 t} + A_2 e^{-B_2 t} + A_3 e^{-B_3 t}, \]

where \( C \) is the concentration of \( F, t \) is the time, \( A_1, A_2, A_3, B_1, B_2, B_3 \) are model parameters, and \( t/2 \) was calculated as 0.693/B3. The area under the curve (AUC) was calculated as

\[ AUC = \frac{A_1}{B_1} + \frac{A_2}{B_2} + \frac{A_3}{B_3}. \]

Model fitting was performed using software SCIENTIST® (Micromath Scientist, Salt Lake City, UT).

2.7 FLIM Data Analysis

FLIM data analysis was performed using SPCIImage 3.9.7 software (Becker & Hickl GmbH). FLIM data is composed of an array of pixels containing several time channels distributed across the fluorescence decay curve. The decay curve represents a sum of multiple exponentials, or components, as each pixel contains an overlay of fluorescence from several endogenous fluorophores at various conformations. For the data we obtained, a double-exponential decay model function was used.

\[ f(t) = \alpha_1 e^{-t/\tau_1} + \alpha_2 e^{-t/\tau_2} \quad \text{with} \quad \alpha_1 + \alpha_2 = 1. \]

The fitted decay curve establishes two lifetimes: the short (\( \tau_1 \)) and long (\( \tau_2 \)) fluorescence decay lifetimes (ps) and with corresponding relative amplitude coefficients \( \alpha_1 \) and \( \alpha_2 \) (%), respectively.

The instrument response function (IRF) was calibrated using a surose crystal standard first analysed by FLIM (Ajax Finechem Pty Ltd, Sydney, NSW, Australia) was used. Each individual FLIM image is convoluted with IRF(\( t \)) in the model function, \( f(t) \), to calculate the function \( F(t) \). The function below corresponds to the shape of the signal if the measured optical signal was identical to the model function \( f(t) \).

\[ F(t) = f(t) \ast IRF(t); * \text{represents the convolution symbol}. \]

Using the \( F(t) \) function, the measured data points are obtained and analyzed to determine the average lifetime and relative contributions of each endogenous molecule.

2.8 High-Performance Liquid Chromatography

The concentration of fluorescein and fluorescein monoglucuronide in each collected bile sample was determined by high-performance liquid chromatography (HPLC, Shimadzu, Kyoto, Japan) using 1 mL/min flow rate with a C18 column [Agilent HC-C18 (2) 4.6 × 150 mm, 5 μm] that included a security guard column (Phenomenex C18 4 × 3 mm). Fluorescence excitation and emission wavelengths were 488 and 515 nm, respectively, with a retention time of 5 min. The mobile phase consisted of 50 mM NaH2PO4 and methanol (2:3).

Analysis of fluorescein involved diluting 5 μl of bile to 500 μl with 0.2 M acetate buffer. For analysis of fluorescein monoglucuronide (FG), a 5-μl bile sample was diluted in 50 μl β-glucuronidase (10 U/100 μl). All samples were then incubated for 1 h at 37°C in a water bath. Fluorescein samples were further diluted in deionized water (1:10), whereas FG samples were first diluted in an acetate buffer (1:20) and then in water (1:10). Samples (10 μl) were injected into the HPLC system. Bile was collected in preweighted tubes, enabling calculation of the excretion rates for fluorescein and fluorescein monoglucuronide.

2.9 RT-PCR

RNA was extracted from 100 to 150 mg liver tissue using TRIzol Reagent as instructed by the manufacturer. DNA contaminations were removed by incubation with RNase-free DNase-1. First strand cDNA was synthesized from 1 μg total RNA using SuperScript III reverse transcriptase (Invitrogen) and oligo dT as instructed by the manufacturer.

cDNA was diluted (1:20) in water, and 2 μl of the sample was added to each well in duplicates. A mixture containing 5 μl platinum SYBR® Green qPCR, 0.2 μl of forward and 0.2 μl of reverse primer and 2.6 μl water, was added to each well, after
which the plate was centrifuged to mix the solutions. RT-PCR was read using a Roche LightCycler480.

### 2.10 Western Blot

Liver slices were homogenized in protein lysis buffer solution (1:20, weight:volume), after which triton (10 µL/ml buffer) was added and stored on ice. The solution was centrifuged at 13,000 rpm at 4°C for 20 min and the supernatant was collected. The supernatant, equivalent to 30 or 50 µg of protein, was mixed with loading buffer and heated for 5 min at 100°C. Acrylamide resolving gel (10%) and 4% stacking gel, both with sodium dodecyl sulfate, were prepared with tris-glycine as a running buffer. Electrophoresis was run at 150 V for 45 min, after which the gel was transferred to membranes at 100 V for 1 h at 4°C. The membranes were blocked in 10% skim milk in blocking buffer. Electrophoresis was run at 150 V for 45 min, after which it was washed in TBS-T. Membranes were incubated with primary antibody (1:500 for OATP2 and MRP2 and 1:1000 for MRP3) in blocking buffer for 1.5 h, after which it was washed in TBS-T. Membranes were incubated with the secondary antibody (1:100 K for OATP2, 1:300 K for MRP3, and 1:20 K for MRP2) for 1.5 h, followed by washing in TBS-T first and then TBS. The membrane was developed onto film using a luminescence kit (SuperSignal® West Femto) with a reaction time of 5 min. β-actin was used as a loading control (1:1000 for primary and 1:200 K for secondary antibody). Protein bands were quantified using ImageJ and adjusted with density of β-actin.

### 2.11 Statistical Analysis

The data are expressed as mean ± standard error of the mean (n = 3 to 7). A student’s t-test was used for statistical comparison of serum ALT levels, fluorescence lifetime data, and pharmacokinetic parameters between sham and I/R injury groups. To analyze statistical differences in mRNA or protein levels between sham and I/R injury groups at 4 and 24 h, a one-way analysis of variance was used with a post hoc Newman-Keuls test. Statistical analysis was done using GraphPad Prism v5.03 (GraphPad Software Inc., La Jolla, California) with a p value less than 0.05 considered statistically significant.

### 3 Results

#### 3.1 I/R-Induced Elevation in Serum ALT Levels

Figure 2 demonstrates extensive liver damage as indicated by significantly elevated (p < 0.05; n = 4) serum ALT levels in rats subjected to I/R injury (1200 ± 260 U/L) compared to sham (54 ± 7 U/L).

#### 3.2 Intravital FLIM Imaging of the Rat Liver after Sham or I/R Injury

Figure 3(a) shows representative intravital FLIM images (pseudocolored according to average weighted lifetime $\tau_{av}$), across three spectral channels (i.e., 350 to 450, 450 to 515, and 515 to 620 nm), of hepatocytes from within the rat liver following sham or I/R injury. The data demonstrate that in the second spectral channel [450 to 515 nm; NAD(P)H/FAD], $\tau_{av}$ appears to be smaller in the rat liver after I/R injury (blue) relative to sham (yellow-green). This observation is also seen in Fig. 3(b), indicated by the left-shift of the $\tau_{av}$ histogram derived from the FLIM image. The average $\tau_{av}$ [450 to 515 nm; NAD(P)H/FAD] of hepatocytes from I/R-treated rats (1575.3 ± 41.9 ps) was significantly lower ($p < 0.001; n = 6$) than the sham control (1225.4 ± 16.3 ps) [Fig. 3(c)]. Further examination revealed that the fast decay lifetime component ($\tau_1$) within hepatocytes was significantly smaller ($p < 0.01$) in I/R-treated rats (1054.5 ± 31.4 ps) compared to the sham control (932.4 ± 17.4 ps) [Fig. 3(d)]. The tendency for the long decay lifetime component ($\tau_2$) in I/R-treated rats to be smaller than the sham control was not statistically significant. In addition, while the $\tau_{av}$ of I/R-treated rats, in the 350 to 450 nm [NAD(P)H] spectral channel appeared smaller than the sham control [Fig. 3(b)], this difference was not statistically significant.

Similar to Fig. 3, Fig. 4 shows representative redox FLIM images of hepatocytes from I/R and sham control rats. The images were pseudocolored according to the ratio of amplitude coefficients $\alpha_1$ and $\alpha_2$. Figure 4(a) demonstrates that in the 350 to 450 nm [NAD(P)H] spectral channel, the redox ratio ($\alpha_1/\alpha_2$) appears higher in hepatocytes after I/R injury (red) relative to sham (blue-green). This tendency is reflected in the $\alpha_1/\alpha_2$ ratio histograms in Fig. 4(b). However, the average $\alpha_1/\alpha_2$ ratio in I/R-treated rats (2.663 ± 0.05) is not significantly higher ($p = 0.187$) than the sham control (2.501 ± 0.12) [Fig. 4(c)].

#### 3.3 Fluorescein Uptake, Distribution, and Clearance from the Rat Liver after I/R Injury

Figure 5 shows representative fluorescence intravital images of the rat liver, from sham and I/R injury groups, over time after injection of fluorescein. Representative intravital images, taken at x10 and x40 magnifications [Fig. 5(a)], demonstrated elevated hepatocyte autofluorescence [two photon excitation ($2P_{exc}$): 740 nm] in the sham control rats compared to I/R injury group. Figure 5(b) and 5(c) shows representative images of fluorescein ($2P_{exc}$: 920 nm) uptake, distribution, and clearance within the liver of sham and I/R injury rats after injection at selected time points over a 180-min time-course. Fluorescein was detected within minutes after injection of fluorescein in sham and I/R injury rats. Fluorescein appears to be taken up and cleared from the liver faster in the sham group compared to the I/R injury group, which displayed delayed uptake and elevated retention within the liver over time [Fig. 5(b) and 5(c)].

Figure 6(a) to 6(c) show the changes in fluorescein concentration within the sinusoids, hepatocytes, and bile, respectively, of sham and I/R injury rats after fluorescein injections. It appears that the fluorescein clearance rate from the sinusoids...
was slower for I/R injury, as indicated by a shallower slope [Fig. 6(a)], which was statistically significant ($p < 0.05$) at 10 to 25 min. Similarly, the excretion rate from the hepatocytes also appeared slower in the I/R group [Fig. 6(b)].

Figure 7(a) and 7(b) shows the AUC value and elimination half-life, respectively, for fluorescein in the sinusoids and hepatocytes, calculated by fitting data from Fig. 6(a) and 6(b) to a triple exponential equation. The data show a significantly increased ($p < 0.05$) AUC in the hepatocytes ($5100 \pm 340 \mu g \cdot min / mL$) for the I/R injury group compared to the sham group ($2800 \pm 640 \mu g \cdot min / mL$). The AUC in the sinusoids and the half-life in the hepatocytes tended to be higher for the I/R injury group ($2030 \pm 510 \mu g \cdot min / mL$ and $430 \pm 61$ min, respectively) relative to the sham group ($1200 \pm 390 \mu g \cdot min / mL$ and $250 \pm 55$ min, respectively), but these differences were not statistically significant.

**Fig. 3** I/R injury associated changes in the average weighted fluorescence lifetime ($\tau_m$) within the liver in vivo. (a) Representative $\tau_m$ intravital images of sham and I/R injury in the liver. FLIM data were measured within three spectral ranges [pseudocolor range (blue-to-red) indicated in parenthesis]: 350 to 450 (750 to 1500 ps), 450 to 515 (1000 to 1750 ps), and 515 to 620 nm (500 to 1750 ps) to capture NAD(P)H, NAD(P)H/FAD, and FAD/fluorescein, respectively. Each image is $214 \times 214 \times 1 \mu m^3$. (b) $\tau_m$ lifetime histograms of sham and I/R injury liver NAD(P)H, NAD(P)H/FAD, and FAD/fluorescein, respectively. The histograms represent the average pixel intensity for each $\tau_m$ lifetime. (c) The corresponding average $\tau_m$ lifetimes from the three spectral channels are charted for the sham and I/R injury groups. (d) The average short ($\tau_1$) and long ($\tau_2$) decay lifetimes obtained from the NAD(P)H/FAD (450 to 515 nm) channel for the sham and I/R injury groups. Values are presented as the mean±SEM ($n = 6$), * $p < 0.01$ or ** $p < 0.001$ significantly different from sham.
3.4 Concentration of Fluorescein in Plasma and Excretion of Fluorescein and Metabolites in Bile

Figure 8(a) demonstrates that the plasma concentration of fluorescein, measured by HPLC, was similar in sham and I/R injury groups at 4 h of reperfusion. The biliary excretion rate of fluorescein [Fig. 8(b)] showed a slight tendency to be lower in the I/R injury group relative to the sham control group, though these differences were not statistically significant. In contrast, the biliary excretion rate of fluorescein monoglucuronide for sham and I/R injury groups was comparable over time [Fig. 8(c)].

3.5 mRNA and Protein Expression of OATP2, MRP2, and MRP3

Figure 9 shows the mRNA and protein expression of OATP2, MRP2, and MRP3 within excised liver tissue from sham and I/R injury groups. mRNA expression of the transporter gene MRP2 tended to be higher in the I/R injury group compared to the sham group after 4 h of reperfusion, respectively, while OATP2 and MRP3 transporter expression appeared to decrease after 4 and 24 h of reperfusion, respectively [Fig. 9(a)]. However, none of these differences were statistically significant. Similar to mRNA expression, OATP2 protein levels for the I/R injury group were similar to those of the sham group after 4 h of reperfusion [Fig. 9(b)].
The injury group appeared to be higher than the sham group after 4 h of reperfusion [Fig. 9(b) and 9(c)]. MRP2 protein levels were comparable between sham and I/R injury groups at both time points. After 4 h of reperfusion, MRP3 levels in the I/R injury group were significantly lower than the sham [Fig. 9(c)]. At 24 h, MRP3 protein levels were comparable to the sham group.

### 4 Discussion

In this study, we used intravital imaging of the rat liver to further characterize hepatic metabolic damage associated with I/R injury. I/R injury was confirmed by measuring elevated serum ALT levels and changes in the metabolic and redox state of hepatocytes by FLIM. Intravital imaging of fluorescein uptake,
distribution, and clearance was a more sensitive indicator of impaired liver function for I/R injury than measuring fluorescein levels in plasma or biliary excretion. Changes in uptake and clearance were also associated with acute changes in liver transport protein expression.

As with previous studies showing an increased ALT in response to I/R injury, the present model for I/R injury also resulted in a significant elevation of serum ALT (Fig. 2), indicative of liver damage.

We have previously described differences in the redox state of hepatocytes, using fluorescence lifetime imaging, between the sham and I/R damaged rat liver. Specifically, the amplitude coefficients ratio ($\alpha_1/\alpha_2$) of the short ($\tau_1$) and long ($\tau_2$) decay lifetimes increased over 4 h after I/R injury. The ratio is an indicator of the redox state of the cells, representing the proportion of free and protein-bound NAD(P)H within the cell. While we observed a tendency for the redox ratio to be elevated, this difference was not statistically significant (Fig. 4).

In this study, we further characterized changes in the metabolic state of hepatocytes, after I/R injury, identifying a decrease in average weighted lifetime ($\tau_m$), specifically due to a decrease in the $\tau_1$ lifetime (Fig. 3). This lifetime difference was observed in both the 350 to 450 and 450 to 515 nm spectral channels, both of which encompass NAD(P)H autofluorescence, though this was statistically significant only in the latter. The lack of any change in the fluorescence lifetime in the 515 to 620 nm spectral channel, which predominantly isolates the FAD autofluorescence, further supports the conclusion that NAD(P)H is responsible for the lifetime changes observed in the 450 to 515 nm channel. Moreover, as no lifetime change was simultaneously observed for FAD, this suggests that an environmental factor (e.g., pH) may not be responsible for the decrease in the $\tau_1$ lifetime (Fig. 3) of which encompass NAD(P)H autofluorescence, though this was statistically significant only in the latter. The lack of any change in the fluorescence lifetime in the 515 to 620 nm spectral channel, which predominantly isolates the FAD autofluorescence, further supports the conclusion that NAD(P)H is responsible for the lifetime changes observed in the 450 to 515 nm channel. Moreover, as no lifetime change was simultaneously observed for FAD, this suggests that an environmental factor (e.g., pH) may not be responsible for the decrease in the $\tau_1$ lifetime (Fig. 3) of which encompass NAD(P)H autofluorescence, though this was statistically significant only in the latter. The lack of any change in the fluorescence lifetime in the 515 to 620 nm spectral channel, which predominantly isolates the FAD autofluorescence, further supports the conclusion that NAD(P)H is responsible for the lifetime changes observed in the 450 to 515 nm channel. Moreover, as no lifetime change was simultaneously observed for FAD, this suggests that an environmental factor (e.g., pH) may not be responsible for the decrease in the $\tau_1$ lifetime (Fig. 3) of which encompass NAD(P)H autofluorescence, though this was statistically significant only in the latter. The lack of any change in the fluorescence lifetime in the 515 to 620 nm spectral channel, which predominantly isolates the FAD autofluorescence, further supports the conclusion that NAD(P)H is responsible for the lifetime changes observed in the 450 to 515 nm channel. Moreover, as no lifetime change was simultaneously observed for FAD, this suggests that an environmental factor (e.g., pH) may not be responsible for the decrease in the $\tau_1$ lifetime (Fig. 3).

Fig. 6 Change in fluorescein concentration over time within the sinusoid (a), hepatocytes (b), and bile (c) of sham and I/R injury groups. Data represent the mean±SEM ($\text{n}=3$). Significant differences are indicated with an asterisk ($p < 0.05$).

Fig. 7 Pharmacokinetic parameters of fluorescein clearance within the hepatocytes of sham and I/R injury groups. Fluorescein accumulation (area under the curve) (a) and elimination half-life (b) were calculated from fluorescein fluorescence intensity changes within the hepatocytes. Data represent the mean±SEM ($\text{n}=3$). Significant differences are indicated with an asterisk ($p < 0.05$).

While
these processes also involve oxidative damage, the outcome is ultimately cell death and a breakdown of internal compartmentalization within the cell, which may alter protein recruitment of NAD(P)H and thus result in the lifetime changes we observe. Further studies are required to identify the protein-binding associated changes for NAD(P)H in response to I/R injury and cell death. To our knowledge, we are the first to associate hepatic I/R injury with specific lifetime changes in NAD(P)H, in addition to the free/bound redox ratio association we have published earlier.

Our results demonstrate a delayed appearance and clearance of fluorescein within the sinusoid of liver with I/R injury (Fig. 5). Our mathematical model showed an increased elimination half-life and accumulation of fluorescein in the hepatocytes in the I/R injury group compared to the sham group, although only hepatocyte fluorescein accumulation was significantly elevated (Fig. 7). The delayed uptake of fluorescein following I/R injury is consistent with a previous study that showed delayed uptake and clearance of indocyanine green (ICG) in the rabbit liver following I/R injury. Another recent study also found a reduced uptake and clearance of perfused rhodamine 123 in the rat liver after I/R injury. To our knowledge, we are the first to use intravitral imaging to visualize and model the uptake and clearance of a fluorescent compound to assess liver function following I/R injury.

Delayed ICG uptake by the rabbit liver after I/R injury was associated with impaired hepatic microcirculation. In addition, another study reports that I/R injury is associated with a reduction in sinusoid density within the liver, which is believed to impair hepatic microcirculation. Therefore delayed fluorescein uptake may be due to a decrease in sinusoid diameter that impairs blood flow and thus availability of the dye to the hepatocytes.

Interestingly, the differences in uptake and clearance of fluorescein measured by intravitral imaging were not observed in plasma or bile using traditional HPLC to assay fluorescein or fluorescein monoglucuronide (Fig. 8). However, the model of both fluorescein and fluorescein monoglucuronide within the rat liver of sham and I/R injury groups (4 and 24 h of reperfusion). Data represent the mean ± SEM (n = 4). Significant differences are indicated with an asterisk (*p < 0.05).

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compared to the sham group (Fig. 9). As MRP3 is responsible for transporting fluorescein from the hepatocytes into the sinusoids, this result may explain the increased AUC accumulation of fluorescein within the hepatocytes observed in Fig. 7.

The expression of hepatic transporters is reported to be altered as a result of I/R injury. For example, the expressions of the bile salt export pump, MRP2, and OATP(s) were all reported to be decreased after 24 h of reperfusion in rats, while multidrug resistance protein 1B was increased.22–24 In nonalcoholic fatty liver disease, expression of MRP2 has been reported to be upregulated,41 whereas it was relatively well preserved in cholestasis and primary biliary cirrhosis.42,43

Interestingly, mRNA and protein levels of the OATP2 transporter appeared contradictory after 4 h of reperfusion [Fig. 7(a) and (7c)]. While the differences in OATP2 expression between sham and I/R injury groups are not statistically significant, the trend may suggest an initial upregulation of OATP2 expression, as reflected by elevated protein levels after 4 h of reperfusion, to augment impaired fluorescein uptake into the hepatocytes from the sinusoids due to I/R damage. A consequence of increased levels of transporters may be that they facilitate liver regeneration and possibly increase bile secretion per weight of functional liver and provide hepatic resistance against cytokine-metabolic stress.46 Decreased mRNA transcripts of OATP2 in the I/R injury group after 4 h may be indicative of a negative feedback loop that has not yet affected protein levels at this time point. The comparable mRNA and protein levels of OATP2 between sham and I/R injury groups after 24 h appears to support this hypothesis. Additional studies measuring changes in transcription and protein levels real-time are necessary to elucidate this possibility.

Our study has demonstrated that MPT imaging and the kinetics of fluorescein uptake and/or clearance may be used to measure liver function after transplantation, in order to assess damage from I/R injury. This approach may be used to assess liver damage intraoperatively by directly imaging selected lobes of the transplanted or damaged liver. The advantage of this approach is that potential damage could be determined in a few hours, rather than up to 18 to 24 h post-transplantation using plasma-ICG clearance,45 to allow for clinical intervention prior to the completion of surgery. MPT imaging equipment and the use of microendoscopic objectives are improving the size and portability as demonstrated for dermatological imaging46 and small animal surgery,47 suggesting current equipment can be used for assessing liver function in vivo as demonstrated by our study.

In conclusion, intravital MPT is a simple and effective optical technique to study drug distribution in normal and diseased liver. It has proven to be a more sensitive alternative to bile and plasma concentration measurements. Furthermore, we have provided a simple mathematical model in studying drug uptake and excretion using intravital MPT.

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