Article

Steatosis Alters the Activity of Hepatocyte Membrane Transporters in Obese Rats

Catherine M. Pastor 1,2,* and Valérie Vilgrain 2,3,4

1 Department of Radiology, University Hospital of Geneva, 1205 Geneva, Switzerland
2 Centre de Recherche sur L’inflammation, Inserm, U1149, CNRS, ERL8252, F-75006 Paris, France; valerie.vilgrain@aphp.fr
3 Department of Radiology, Hôpital Beaujon, Université de Paris, 92110 Clichy, France
4 Department of Radiology, Hôpitaux Paris Nord Val de Seine (AP-HP), Université de Paris, 92110 Clichy, France
* Correspondence: catherine.pastor@unige.ch

Abstract: Fat accumulation (steatosis) in ballooned hepatocytes alters the expression of membrane transporters in Zucker fatty (fa/fa) rats. The aim of the study was to quantify the functions of these transporters and their impact on hepatocyte concentrations using a clinical hepatobiliary contrast agent (Gadobenate dimeglumine, BOPTA) for liver imaging. In isolated and perfused rat livers, we quantified BOPTA accumulation and decay profiles in fa/+ (normal) and fa/fa hepatocytes by placing a gamma counter over livers. Profiles of BOPTA accumulation and decay in hepatocytes were analysed with nonlinear regressions to characterise BOPTA influx and efflux across hepatocyte transporters. At the end of the accumulation period, BOPTA hepatocyte concentrations and influx clearances were not significantly different in fa/+ and fa/fa livers. In contrast, bile clearance was significantly lower in fatty hepatocytes while efflux clearance back to sinusoids compensated the low efflux into canaliculi. The time when BOPTA cellular efflux impacts the accumulation profile of hepatocyte concentrations was slightly delayed (2 min) by steatosis, anticipating a delayed emptying of hepatocytes. The experimental model is useful for quantifying the functions of hepatocyte transporters in liver diseases.

Keywords: hepatocyte membrane transporter; hepatocyte concentrations; transporter activity; liver imaging

1. Introduction

Nonalcoholic fatty liver disease (NAFLD) includes various dysfunctions such as increased intrahepatic triglyceride content (steatosis), inflammation, and fibrosis [1,2]. The transition from fatty livers to more severe disease (steatohepatitis) is triggered by inflammation. Cholestasis (or decreased bile flow) is a marker of severity. Significant progress has been made to decrease the burden of NAFLD, but the translation of experimental treatments to the human disease has consistently failed [3]. One of the reasons is that the mechanisms underlying the disease are numerous and complex. Nevertheless, these experimental models contribute to a better understanding of the disease.

Thus, NAFLD modifies the expression of transporters located on the basolateral and canalicular membranes of hepatocytes. Human Organic Anion Transporting Polypeptide B1/B3 (OATP1B1/B3) are not altered in steatosis, but their expression is decreased in patients with steatohepatitis [4]. In contrast, the Multidrug Resistance Associated Protein 3 (MRP3) expression is increased in fatty livers and steatohepatitis. MRP2 is upregulated in steatohepatitis [5]. The activity of these transporters determines the hepatocyte concentrations of endogenous substrates and drugs that might be unpredictable in fatty livers. Thus, patients with steatohepatitis have higher liver concentrations of mebrofenin (MEB,
Choletec®, Bracco imaging, Milan, Italy) than volunteers [6]. MEB is transported across hepatocytes via human OATPs, MRP2, and MRP3 (Figure 1A).

![Diagram of BOPTA concentrations in liver compartments](image1)

**Figure 1.** (A) Distribution of BOPTA concentrations in liver compartments: portal vein ($C_{in}$), hepatic veins ($C_{out}$), extracellular space ($C_{EC}$), hepatocytes ($C_{HC}$), and bile canaliculi ($C_{bile}$). (B) Liver concentrations of contrast agents are measured by a gamma counter placed over livers. DTPA preperfusion was used to measure $C_{EC}$ (purple symbols). DTPA distributes only into the extracellular space. DTPA concentrations were reported with BOPTA liver concentrations (***). BOPTA (black symbols) distributes into the extracellular space, hepatocytes, and bile canaliculi. (C) $^{153}$GdDTPA and $^{153}$GdBOPTA protocol of perfusion.

Experimental nonalcoholic fatty livers can be investigated in Zucker fatty (fa/fa) rats [7,8]. These rats have a mutated leptin receptor and a decreased leptin affinity. Rats develop severe obesity with insulin resistance and hepatic steatosis. The spontaneous progression of steatosis to steatohepatitis is rare and needs a second hit [9]. Heterozygous fa/+ rats serve as control animals. In biopsies, we showed previously that hepatocytes in fa/fa livers are distended by large fat droplets [10]. This fat accumulation increases the hepatocyte volume and total liver weights [11]. No inflammation or fibrosis is detected. Steatosis is associated with high triglyceride and cholesterol serum concentrations [10]. Portal pressures remain normal, but bile flow rates were significantly decreased in comparison to normal livers.

The expression of hepatocyte transporters was also investigated in fa/fa livers. According to Canet et al. [12], Oatp1a4 expression is downregulated while that of Oatp1b2 is induced in fa/fa livers. In contrast, Mrp2 and Mrp3 expression is not significantly different in fa/fa and control Sprague-Dawley livers. Another experimental study found that rats fed with a high fat diet had a decreased mRNA expression of Slco1a2, Slco1b2, Abcc2, and Abcc3 (genes coding for Oatps, Mrp2, and Mrp3) [13]. However, the protein expression of the transporters was not available.

We previously showed that fa/fa rats have a decreased bile excretion of Gadobenate dimeglumine (BOPTA, Multihance®, Bracco Imaging, Milan, Italy) [10]. BOPTA is a clinical hepatobiliary contrast agent that characterises liver parenchyma and focal lesions [14].

![Diagram of BOPTA concentrations in liver compartments](image2)

![Diagram of DTPA and BOPTA liver concentrations](image3)
BOPTA distributes into the extracellular compartment and enters into rat hepatocytes by the Organic anion transporting polypeptide transporters (Oatp1a1, Oatp1a4, and Oatp1b2) (Figure 1A) [15]. BOPTA is excreted unchanged (no metabolism) into bile canaliculi through the Multidrug resistance-associated protein 2 (Mrp2) [16]. BOPTA can also return back to sinusoids using Mrp3. BOPTA accumulation in hepatocytes is modulated by the activity of these influx and efflux transporters.

In an ex-vivo model of isolated and perfused rat liver (IPRL), we previously published the distribution of BOPTA in the extracellular space, hepatocytes, and bile canaliculi [17]. By placing a gamma counter over livers, it is possible to measure the liver concentrations of labelled BOPTA (Figure 1B). A preperfusion of labelled gadopentetate dimeglumine (DTPA) quantifies the extracellular concentrations, because it distributes exclusively into the extracellular space. Concentrations of bile canaliculi and hepatocytes were calculated. We found that the maximal BOPTA hepatocyte concentrations (obtained at the end of the perfusion period) were similar in fa/+ and fa/− livers. A compartmental pharmacokinetic modelling determined that the BOPTA influx rates were similar in both groups while the efflux rates from hepatocytes into bile and back into sinusoids were decreased.

In the present study, we reanalysed the row data of fa/+ and fa/− livers to quantify the BOPTA compartmental distribution (or respective contribution of the extracellular space, hepatocytes, and bile canaliculi to liver concentrations) and new pharmacokinetic parameters that characterise BOPTA accumulation and decay in hepatocytes.

2. Materials and Methods

2.1. Isolated and Perfused Rat Livers

Before liver isolation, Zucker fat (fa/−, n = 5) and Zucker lean (fa/+, n = 5) rats were anesthetised with pentobarbital (50 mg · kg⁻¹, ip). In these two groups, we published previous biological parameters of liver functions while the transporter functions were assessed by pharmacokinetic modelling [10]. In the present study, we returned to row data to analyse the new hepatic pharmacokinetic parameters. Rat livers were isolated, leaving the organs in the carcass. The abdominal cavity was opened and the portal vein cannulated. The hepatic artery was not perfused. The abdominal vena cava was transected, and an oxygenated Krebs–Henseleit bicarbonate (KHB) solution was pumped into the portal vein, the solution being discarded after liver distribution via a vena cava transection. The flow rate was slowly increased over 1 min up to 30 mL/min. In the second step, the chest was opened, and a cannula was inserted through the right atrium to collect solutions flowing from the hepatic veins. Finally, the abdominal inferior vena cava was ligated, allowing solutions perfused by the portal vein to be eliminated by the hepatic veins.

The perfusion system included a reservoir, a pump, a heating circulator, a bubble trap, a filter, and an oxygenator. Solutions of perfusion were equilibrated with a mixture of 95% O₂ and 5% CO₂. Livers were continuously perfused with fresh solutions using a nonrecirculating system. The common bile duct was cannulated with a PE10 catheter, and bile samples were collected every 5 min to measure the bile flow rates (Q bile, µL/min/liver) and BOPTA concentrations (C bile, µM). Samples were collected from hepatic veins every 5 min (C out, µM). C out during the BOPTA perfusion period were concentrations that did not enter into hepatocytes plus concentrations that entered into hepatocytes (via Oatps) and returned into sinusoids via Mrp3 (Figure 1A). During the rinse period, C out were concentrations leaving hepatocytes because the concentration in the portal vein (C in) was 0.

An adequate viability of livers was assessed by a steady portal pressure below 12 mmHg during the entire protocol. We previously published that a flow rate of 30 mL/min/liver maintains a normal liver O₂ consumption [18].

2.2. Perfusion of DTPA and BOPTA

Rat livers were perfused with gadopentetate dimeglumine (DTPA; Magnevist®; Bayer imaging, Berlin, Germany) and gadobenate dimeglumine (BOPTA, MultiHance®; Bracco Imaging, Milan, Italy). DTPA distributes only within sinusoids and the interstitium,
while BOPTA distributes into the extracellular space, hepatocytes, and bile canaliculi. DTPA and BOPTA labelled with $^{153}$Gd were obtained by adding $^{153}$GdCl$_3$ (1 MBq/mL) to the commercially available (0.5 M) solutions of DTPA and BOPTA. Then, $[^{153}$Gd]DTPA and $[^{153}$Gd]BOPTA were diluted in the KHB solution to obtain a 200-µM concentration. Livers were successively perfused with 200 µM $[^{153}$Gd]DTPA (10 min), KHB solution (35 min), 200 µM $[^{153}$Gd]BOPTA (30 min, accumulation or perfusion period), and KHB solution (30 min, decay or elimination period) (Figure 1C). The protocol lasted 105 min for each group.

2.3. Concentrations in Liver Compartments

To quantify the BOPTA concentrations in liver compartments, a gamma counter that collects count rates every 20 s was placed 1 cm above a right liver lobe. The counter measured the radioactivity in a region of interest that was identical in each liver. To transform count rates into BOPTA concentrations, the total liver radioactivity was measured by an activimetre at the end of each experiment and related to the last count rates. Radioactivity was corrected for decay. The gamma counter delineated a region of interest in a liver lobe from which all count rates originating from the extracellular space, hepatocytes, and bile canaliculi were divided by the liver weight to obtain liver concentrations ($C_{\text{liver}}, \mu M$). To calculate the hepatocyte concentrations, we must eliminate the BOPTA concentrations located inside the extracellular space ($C_{\text{EC}}$) and bile canaliculi ($C_{\text{BC}}$). BOPTA extracellular concentrations cannot be measured, because the substrate rapidly enters into hepatocytes (<2 min). We used DTPA that distributes only in the extracellular space to estimate the BOPTA $C_{\text{EC}}$. Assuming that DTPA $C_{\text{EC}}$ is similar to BOPTA $C_{\text{EC}}$, we can subtract BOPTA $C_{\text{liver}} - \text{DTPA } C_{\text{liver}}$ (or DTPA $C_{\text{EC}}$). The concentrations were constant during the 10-min perfusion. During BOPTA perfusion, this plateau was extended over 30 min (Figure 1B). We assumed that the concentrations inside bile canaliculi were similar to those measured in the common bile duct ($C_{\text{bile}}, \mu M$), although solute export from cholangiocytes and water transport along ductules and ducts may modify the primary bile in canaliculi. The volume ratio of the bile canaliculi and liver was previously estimated by Blouin et al. [19] at 0.43%. Concentrations in the bile canaliculi detected by the counter ($C_{\text{BC}}$) were then equal to 0.0043 · $C_{\text{bile}}$. Hepatocyte concentrations in a 78% liver volume ($C_{\text{HC}78\%}$) detected by the counter were $C_{\text{liver}} - C_{\text{EC}} - 0.0043 C_{\text{bile}}$. Indeed, Blouin et al. [19] determined previously that the volume ratio of hepatocytes to liver in the absence of fibrosis or inflammation was 78%. In situ $C_{\text{HC}100\%}$ were calculated by $(100/78) \cdot C_{\text{HC}78\%}$.

2.4. Transfer Rates and Clearances between Compartments

The BOPTA removal rate from sinusoids during the perfusion period ($v$, nmol/min) was measured by $Q_{\text{H}} \cdot (C_{\text{in}} - C_{\text{out}})$, where $Q_{\text{H}}$ was the constant liver flow rate (30 mL/min), $C_{\text{in}}$ (µM) was the constant portal concentration, and $C_{\text{out}}$ (µM) was the concentration measured in hepatic veins. The unbound fraction in the solutions was 1, because no protein was added into the solutions. Hepatic clearance ($C_{\text{H}}$, mL/min) was the ratio of $v$ and $C_{\text{in}}$ during the last min of perfusion. The BOPTA extraction ratio (ER) was $(C_{\text{in}} - C_{\text{out}})/C_{\text{in}}$. The BOPTA biliary excretion rate ($v_{\text{bile}}, \text{nmol/min}$) was $C_{\text{bile}} \cdot Q_{\text{bile}}$, where $C_{\text{bile}}$ (µM) was the concentration in the common bile duct, and $Q_{\text{bile}}$ was the bile flow rate (µL/min/liver weight). Clearance from the hepatocytes to bile canaliculi ($C_{\text{Lb}}$, mL/min) was the slope of the linear regression between $v_{\text{bile}}$ (Y-axis) and the hepatocyte concentrations ($C_{\text{HC}}$, X-axis). $C_{\text{Lb}}$ was measured during the entire protocol. During the rinse period, BOPTA concentrations leaving hepatocytes into sinusoids ($C_{\text{ef}}, \mu M$) were measured by $C_{\text{out}}$ in the absence of BOPTA entry into livers. The basolateral efflux from hepatocytes into sinusoids ($v_{\text{ef}}, \text{nmol/min}$) was $C_{\text{ef}} \cdot Q_{\text{H}}$, and the basolateral clearance ($C_{\text{Lef}}, \text{mL/min})$ was the slope of linear regression between $v_{\text{ef}}$ (Y-axis) and $C_{\text{HC}}$ (X-axis). With the assumption that $C_{\text{ef}}$ was similar during both the perfusion and rinse periods, we estimated $C_{\text{ef}}$ during the accumulation period by $(C_{\text{HC}} \cdot C_{\text{Lef}})/Q_{\text{H}}$. The hepatocyte influx rate $v_{\text{in}}$ (nmol/min) was
[C\textsuperscript{in} − (C\textsuperscript{out} − C\textsubscript{ef})] \cdot Q\textsubscript{H}, and the hepatocyte influx clearance CL\textsubscript{in} (mL/min) was \(v_{\text{in}}/C_{\text{in}}\). The CL\textsubscript{in} was measured during the last min of perfusion.

2.5. Accumulation Profile of Hepatocyte Concentrations

During the perfusion period, BOPTA hepatocyte accumulation was best described by a segmental linear regression obtained from GraphPad Prism version 8, GraphPad Software, La Jolla, CA, USA [20]. This function defines a first line \(L_1\) for a time below \(T_0\) and a second line \(L_2\) for a time higher than \(T_0\). \(T_0\) is the time when the two lines would intersect. No constraint was applied to fit the data.

2.6. Concentration Ratios (R) between Compartments

In in vivo studies, only liver-to-plasma concentration ratios are available. However, concentrations facing the sinusoidal membrane of hepatocytes where Oatps reside are the extracellular concentrations, because the space of Disse intertwines between sinusoids and the sinusoidal membrane. Thus, the hepatocyte-to-extracellular concentration ratio (\(R_{HC/EC}\)) characterises BOPTA transport across Oatps. \(R_{HC/EC}\) was measured at \(T_0\) before the BOPTA efflux out of cells. The bile-to-hepatocyte concentration ratio (\(R_{bile/HC}\)) was the slope of the relationship between \(C_{\text{bile}}\) (Y-axis) and \(C_{HC}\) (X-axis) during the entire protocol. \(R_{bile/HC}\) was independent of the BOPTA influx into hepatocytes. The hepatic vein-to-hepatocyte ratio (\(R_{HV/HC}\)) was the slope of the relationship between \(C_{\text{out}}\) (Y-axis) and \(C_{HC}\) (X-axis) during the rinse period. \(R_{HV/HC}\) was independent of the BOPTA influx into the hepatocytes. The concentrations used in these ratios were in situ concentrations. Finally, the bile-to-extracellular concentration ratio (\(R_{bile/EC}\)) assessed the ability of transporters to concentrate BOPTA from the extracellular space to bile compartments. \(R_{bile/EC}\) was measured at the end of the perfusion period, when the bile concentration was maximal.

2.7. Decay Profile of Hepatocyte Concentrations

During the decay period, the data were best described by a one phase decay (GraphPad Prism version 8, GraphPad Software, La Jolla, CA, USA) [21]. The plateau was constrained to 0, because BOPTA must leave hepatocytes. The model was defined by a rate constant of elimination (\(k_{el,HC}, \text{min}^{-1}\)). Knowing \(CL_{\text{bile+ef}}\) and \(k_{el,HC}\), we calculated the apparent hepatocyte volumes (\(V_{HC}, \text{mL}\)) as \(CL_{\text{bile+ef}}/k_{el,HC}\).

2.8. Statistics

Data were the means ± SD. The parameters obtained in normal and fatty livers were compared with a Mann–Whitney test (GraphPad Prism version 8, GraphPad Software, La Jolla, CA, USA).

3. Results

3.1. Basic Pharmacokinetic Parameters

The fatty livers were cholestatic before BOPTA perfusion. The bile flow rates (\(Q_{\text{bile}}\)) were 10.7 ± 1.0 (\(fa/+\) livers) and 8.7 ± 1.0 \(\mu\)L/min/liver (\(fa/fa\) livers, \(p = 0.03\), Figure 2A, before BOPTA perfusion, time point: 45 min). Then, BOPTA perfusion increased bile flow rates during the perfusion period (45–75 min). The area under the curve of the bile increase (\(AUC_{Q\text{bile}}\)) was not significantly different in fatty (134 ± 19 \(\mu\)L) and normal (185 ± 36 \(\mu\)L) livers (\(p = 0.06\), Figure 2B). At the end of the perfusion period, the BOPTA liver extraction ratios were low and not significantly different in the two groups (6 ± 1% in \(fa/+\) livers and 5 ± 1% in \(fa/fa\) livers, \(p = 0.17\)).
Cells 2021, 10, x FOR PEER REVIEW 6 of 13

Figure 2. (A) Bile flow rates ($Q_{\text{bile}}$) during the BOPTA perfusion and rinse periods. Livers were perfused with Krebs–Henseleit bicarbonate solution (KHB) + 200 μM [153Gd]BOPTA (45–75 min, grey area) and KHB (75–105 min). Control livers (fa/+, black circles) and fatty livers (fa/fa, open circles).

3.2. Accumulation in Liver Compartments

BOPTA accumulated in three compartments during the perfusion period (Figure 3 and Table 1). In the extracellular compartment, the concentrations ($C_{\text{EC}}$) were steady and not significantly different in the two groups. $C_{\text{EC}}$ contributed to 15 ± 7% (fa/+ livers) and 10 ± 3% (fa/fa livers) of the liver concentrations ($p = 0.22$). The maximal hepatocyte concentrations detected by the counter ($C_{\text{HC78%}}$) were also similar in lean (465 ± 61 μM) and fatty (461 ± 109 μM) livers ($p = 0.70$). In fa/+ livers, the BOPTA maximal concentrations detected by the counter in bile canaliculi ($C_{\text{BC}}$) were 54 ± 7 μM and accounted for 9 ± 1% of the liver concentrations. In fa/fa livers, BOPTA $C_{\text{BC}}$ was 37 ± 9 μM ($p = 0.02$ vs. fa/+ livers) and accounted for 7 ± 2% of $C_{\text{liver}}$ ($p = 0.10$ vs. fa/+ livers). BOPTA $C_{\text{out}}$ in hepatic veins were slightly lower than the portal vein concentrations $C_{\text{in}}$, reflecting the low liver extraction ratio of BOPTA. $C_{\text{out}}$ were not significantly different between the two groups. At the end of the accumulation period, BOPTA compartmental distribution was similar in the extracellular space and hepatocytes for both groups. However, the accumulation inside bile canaliculi was significantly lower in cholestatic livers. Nevertheless, the liver concentrations were not significant different between the two groups.

Figure 3. BOPTA compartmental distribution in fa/+ (A) and fa/fa (B) livers. Livers were perfused with Krebs–Henseleit bicarbonate solution (KHB) + 200 μM [153Gd]BOPTA (45–75 min) and KHB alone (75–105 min). Liver concentrations (black symbols) were measured by a gamma counter. Concentrations in the extracellular compartment (red symbols) were measured during the previous DTPA perfusion. Concentrations that originated from bile canaliculi (blue symbols) and from 78% hepatocytes (green symbols) were calculated.
Table 1. BOPTA concentrations at the end of the accumulation period.

| Livers   | fa/+       | fa/fa     | p     |
|----------|------------|-----------|-------|
| C_{in} (µM) | 200       | 200       |       |
| C_{out} (µM) | 189 ± 2   | 190 ± 1   | 0.19  |
| C_{EC} (µM) | 93 ± 47   | 52 ± 10   | 0.22  |
| C_{BC} (µM) | 54 ± 7    | 37 ± 9    | 0.02  |
| C_{HC78%} (µM) | 465 ± 61 | 461 ± 109 | 0.70  |
| C_{bile} (µM) | 12564 ± 1550 | 8594 ± 2208 | 0.02  |
| C_{liver} (µM) | 612 ± 79  | 550 ± 110 | 0.31  |

3.3. Accumulation Profile of BOPTA Concentrations in Hepatocytes

The accumulation profile of BOPTA concentrations in hepatocytes relied on the difference between the influx rates (v_{in}, nmol/min) and efflux rates (v_{bile+ef}, nmol/min) (Figure 4A,B, grey areas and Table 2). In both groups, v_{in} remained higher than v_{bile+ef}. At the end of the accumulation period, v_{bile+ef} was not significantly different between the two groups. However, v_{bile} was significantly lower (p = 0.02) and v_{ef} (p = 0.06) was higher in fa/fa than fa/+ livers (Table 2). Thus, in fatty livers, BOPTA efflux back into sinusoids compensates for the decreased bile excretion.

Figure 4. BOPTA hepatocyte influx rates (v_{in}, left Y-axis, black circles) and bile plus basolateral efflux rates (v_{bile+ef}, right Y-axis, open circles) during the BOPTA accumulation period from 45 to 75 min in fa/+ (A) and fa/fa (B) livers. Livers (n = 10) were perfused with Krebs–Henseleit bicarbonate solution (KHB) + 200 µM [153]Gd]BOPTA. Difference between v_{in} and v_{bile+ef} (grey area). (C) Accumulation of BOPTA hepatocyte concentrations from 45 to 75 min. Accumulation was best described by a segmental linear regression (red curve for fa/+ and blue curve for fa/fa). T₀ (red and blue arrows) were the times when BOPTA cellular efflux impacted the hepatocyte accumulation. (D). Bile concentrations measured 5 min after the start of BOPTA perfusion (50 min of the experimental protocol).
BOPTA hepatocyte accumulation was best described by a segmental linear regression (Figure 4C) [20]. This function defined a first line \( L_1 \) for a time below \( T_0 \) and a second line \( L_2 \) for a time higher than \( T_0 \), while ensuring that both lines intersected at \( T_0 \). \( T_0 \) was the time when BOPTA efflux from hepatocytes interfered with hepatocyte accumulation. \( T_0 \) occurred \( 7 \pm 1 \) min (fa/+ livers) and \( 9 \pm 2 \) min (fa/fa livers) after the start of BOPTA perfusion \( (p = 0.02) \). The \( L_1 \) slopes were \( 52 \pm 16 \) µM/min (fa/+ livers) and \( 31 \pm 4 \) µM/min (fa/fa livers, \( p = 0.01) \). These slopes characterised BOPTA influx into hepatocytes by Oatps. After \( T_0 \), the \( L_2 \) slopes were lower than the \( L_1 \) slopes (fa/+ livers: \( 11 \pm 1 \) µM/min and fa/fa livers: \( 15 \pm 4 \) µM/min, \( p = 0.06) \). Concomitant entry and efflux from the hepatocytes explained this decrease. Five minutes after the start of BOPTA perfusion, the bile concentrations were \( 1676 \pm 208 \) (fa/+ livers) and \( 443 \pm 269 \) (fa/fa livers) \( (p = 0.01, \) Figure 4D). In comparison to the lean livers, the fatty livers had a delayed \( T_0 \), decreased \( L_1 \) slopes (hepatocyte influx), similar \( L_2 \) slopes, and decreased bile concentrations.

At the end of the accumulation period, the BOPTA influx clearance (CL\(_{\text{in}}\)) was \( 2.3 \pm 0.3 \) mL/min (fa/+ rats) and \( 2.5 \pm 0.2 \) mL/min (fa/fa rats, \( p = 0.55) \). CL\(_{\text{H}}\) defined by \((C_{\text{in}} - C_{\text{out}}) \cdot Q_{\text{H}}\) was lower than CL\(_{\text{in}}\), which was defined by \([C_{\text{in}} - (C_{\text{out}} - C_{\text{d}})] \cdot Q_{\text{H}}\). CL\(_{\text{H}}\) and CL\(_{\text{in}}\) were not significantly different in the two groups.

3.4. Concentration Ratios (R) between Compartments

The concentration ratio between the hepatocytes and extracellular space (R\(_{\text{HC/EC}}\)) at \( T_0 \) was not significantly different between groups (Table 3). R\(_{\text{HC/EC}}\) was measured before BOPTA hepatocyte efflux. The concentration ratio between the bile canaliculi and hepatocytes (R\(_{\text{bile/HC}}\)) was the slope of relationship between \( C_{\text{HC}} \) (X-axis) and \( C_{\text{bile}} \) (Y-axis) during the perfusion and rinse periods. R\(_{\text{bile/HC}}\) was not significantly different between groups. In contrast, R\(_{\text{HV/HC}}\) was significant higher in fatty livers, and more BOPTA returned to sinusoids. R\(_{\text{HV/HC}}\) was the slope of the relationship between \( C_{\text{HC}} \) (X-axis) and \( C_{\text{out}} \) (Y-axis) during the rinse period (no BOPTA entry into livers). Finally, the maximal concentration ratio between the bile and extracellular space (R\(_{\text{bile/EC}}\)) quantified the ability of transporters to concentrate BOPTA from the extracellular space to bile. The ratio was measured at the end of the perfusion period and was not significantly different between groups.

### Table 3. BOPTA concentration ratios between liver compartments.

| Livers     | fa/+       | fa/fa      | \( p \) |
|------------|------------|------------|---------|
| R\(_{\text{HC/EC}}\) | 4.1 ± 1.8   | 5.5 ± 1.3  | 0.15    |
| R\(_{\text{bile/HC}}\) | 20.2 ± 3.2  | 18.6 ± 5.8 | 0.84    |
| R\(_{\text{HV/HC}}\) | 0.0068 ± 0.0018 | 0.0135 ± 0.0030 | 0.01 |
| R\(_{\text{bile/EC}}\) | 162 ± 69 | 109 ± 54 | 0.15 |
3.5. BOPTA Elimination from Hepatocytes

We used two ways to assess the BOPTA efflux from hepatocytes: BOPTA hepatocyte concentration decay and BOPTA recovery in sinusoids (CL_{ef}) plus bile canaliculi (CL_{bile}). During the rinse period, hepatocyte concentration decay was best described by a one phase decay (Figure 5A and Table 4). The model was described by a rate constant of elimination (k_{el,HC}) that was significantly decreased in fatty livers (Table 4). CL_{bile} and CL_{ef} were defined by the linear regression between v_{bile} or v_{ef} (Y-axis) and the hepatocyte concentrations (C_{HC, X-axis}) (Figure 6). CL_{bile} was significantly lower in fa/fa livers (0.26 ± 0.11 mL_{HC}/min) than in fa/+ livers (0.46 ± 0.09 mL_{HC}/min, p = 0.02, Table 2). In contrast, CL_{ef} was significantly higher in fa/fa livers (0.41 ± 0.09 mL_{HC}/min) than in fa/+ livers (0.20 ± 0.05 mL_{HC}/min, p = 0.01). In fatty livers, high CL_{ef} compensated the low CL_{bile}, and CL_{bile+ef} was similar in both groups (Table 2). The ratios between CL_{ef} and CL_{bile+ef} were 62 ± 13% (fa/fa livers) and 31 ± 7% (fa/+ livers) (p = 0.01). Knowing CL_{bile+ef} and k_{el,HC}, we calculated the apparent hepatocyte volume (V_{HC}, ml) as CL_{bile+ef}/k_{el,HC}. The V_{HC} were significantly higher in fa/fa livers.

![Figure 5](image_url)

**Figure 5.** BOPTA hepatocyte (A) and bile (B) concentrations during the rinse period (75–105 min) when the livers were perfused only with the Krebs–Henseleit bicarbonate solution. Hepatocyte decay was best described by a one phase decay (red curve for fa/+ and blue curve for fa/fa).

**Table 4.** Parameters of BOPTA hepatocyte decay.

| Livers | fa/+     | fa/fa    | p  |
|--------|----------|----------|----|
| Y_{0} (µM) | 587 ± 81 | 598 ± 137 | 1.00 |
| k_{el,HC} (min⁻¹) | 0.08 ± 0.02 | 0.03 ± 0.01 | 0.01 |
| T_{1/2} (min) | 9 ± 2 | 23 ± 4 | 0.01 |
| V_{HC} (ml) | 9 ± 2 | 21 ± 2 | 0.01 |
| 78% of liver weight (ml) | 7 ± 1 | 11 ± 2 | 0.01 |

Hepatocyte concentrations at the beginning of the rinse period (Y_{0}). Decay rate constant (k_{el,HC}). Time when Y equals Y_{0}/2 (T_{1/2}). Apparent hepatocyte volume (V_{HC}).
4. Discussion

BOPTA is a clinical hepatobiliary contrast agent that characterises liver disease and focal lesions in liver magnetic resonance imaging [14]. BOPTA distributes within the extracellular compartment and enters into rat hepatocytes by Organic anion transporting polypeptide transporters (Oatp1a1, Oatp1a4, and Oatp1b2) [15]. The contrast agent is excreted unchanged (no metabolism) into bile canaliculi through Mrp2 [16]. BOPTA can also return back to sinusoids using Mrp3. BOPTA hepatocyte accumulation is modulated by the activity of the three transporters. In this study, we showed that BOPTA has a low liver extraction ratio (<7%) that is not modified by steatosis. BOPTA transport by Mrp2 is associated with water transport (choleresis). We showed that a BOPTA-induced bile flow increase is not significantly different in fa/fa and fa/+ livers. However, a single low value of AUCQ\textsubscript{bile} in the fa/+ group might hide the statistical difference between the groups. The water transport across the canalicular membrane may be linked to BOPTA crossing via Mrp2, which triggers the trafficking and insertion of aquaporin-8-containing vesicles into the canalicular membrane [22,23]. BOPTA-induced choleresis is concentration-dependent and inhibited in livers lacking Mrp2 [16]. Similar results were published with benzylpenicillin [24].

BOPTA distributes into the extracellular space before reaching Oatps. In accordance with the absence of fibrosis in fatty livers [10], steatosis does not modify the extracellular concentrations that represent 13% of the liver concentrations in both groups. The maximal hepatocyte concentrations are also similar, as are the liver concentrations. Consequently, liver imaging following BOPTA injection would not detect differences between the two groups. However, BOPTA accumulation profiles are modified in steatosis. BOPTA accumulation into hepatocytes is best described by segmental linear regression [20]. This function

Figure 6. BOPTA clearances from the hepatocytes to bile canaliculi (CL\textsubscript{bile}) were determined by the slope of linear regression between \(v\textsubscript{bile} \) (Y-axis) and \(C\textsubscript{HC} \) (X-axis) in fa/+ (A) and fa/fa (B) livers. The BOPTA basolateral efflux clearance (CL\textsubscript{ef}) was measured during the rinse period by the slope of linear regression between \(v\textsubscript{ef} \) (Y-axis) and \(C\textsubscript{HC} \) (X-axis) in fa/+ (C) and fa/fa (D) livers. Linear regressions in normal (A and C, red) and fatty (B and D, blue) livers.
defines a first line \( L_1 \) for a time before \( T_0 \) (before BOPTA efflux from hepatocytes). \( T_0 \) is slightly delayed in fatty hepatocytes, anticipating a delay in cellular emptying. Steatosis decreases the \( L_1 \) slope or BOPTA entry into fatty hepatocytes. After \( T_0 \), \( L_2 \) slopes are much lower than \( L_1 \) slopes, because BOPTA efflux from hepatocytes counteracts the influx. Steatosis does not modify the \( L_2 \) slopes.

Another parameter of BOPTA influx into hepatocytes is the concentration ratio between the hepatocytes and extracellular space (\( R_{HC/EC} \)) measured at \( T_0 \) before BOPTA cellular efflux. Steatosis does not modify this ratio. BOPTA influx clearance (\( CL_{in} \)) and hepatic clearance (\( CL_{H} \)) are also similar at the end of BOPTA perfusion, confirming that steatosis does not substantially modify the Oatp activity. Geier et al. [11] found a decreased expression of Oatp1a1 but not Oatp1a4 in \( fa/fa \) livers. The transport activities were not investigated in this study. The reason why the \( L_1 \) slope is lower in \( fa/fa \) than \( fa/+ \) hepatocytes might be explained by the larger volume of fatty hepatocytes, the \( L_1 \) slope being expressed in \( \mu M/min \) or \( nmol/(mL \cdot min) \). With similar \( v_{in} \) in both groups, the \( L_1 \) slope will be lower in a larger volume of fatty hepatocytes.

The BOPTA accumulation in bile (\( C_{bile} \)) is significantly lower during the accumulation period in \( fa/fa \) livers, and the \( C_{bile} \) decrease during the rinse period is slower in fatty livers. Accordingly, the bile clearance (\( CL_{bile} \)) is lower in fatty livers, showing a decreased activity of Mrp2 associated with steatosis. The low activity of Mrp2 can be attributed to a lower expression [11,25]. In contrast, efflux clearance back into sinusoids (\( CL_{ef} \)) is significantly higher in fatty livers, reflecting an increased activity of Mrp3 that compensates the decreased activity of Mrp2. Thus, the sum of both efflux clearances (\( CL_{bile+ef} \)) is not significantly different between both groups. Such compensation was previously published [26]. Nevertheless, the Mrp3 expression is not increased in \( fa/fa \) livers [11,25], suggesting that a dysregulation might promote BOPTA transport without changing the transporter expression. In contrast, the human MRP3 expression is increased in fatty livers and steatohepatitis [4]. Another way to describe BOPTA hepatocyte emptying is to compare the rate constant of elimination (\( k_{el,HC} \)) obtained with the one phase decay equation. The \( k_{el,HC} \) that quantifies BOPTA efflux into bile canaliculi and sinusoids is lower in the presence of steatosis. Knowing \( CL_{bile+ef} \) and \( k_{el,HC} \), we can calculate the apparent hepatocyte volume where BOPTA distributes (\( V_{HC} \)). The \( V_{HC} \) is significantly higher in \( fa/fa \) livers. However, these apparent hepatocyte volumes overestimate the hepatocyte volume calculated by 0.78 \cdot liver weights.

When we previously determined the transporter activities of Oatps, Mrp2, and Mrp3 by pharmacokinetic modelling, we found that the BOPTA hepatocyte influx in \( fa/fa \) rats was not significantly different from \( fa/+ \) livers [10]. In contrast, the bile and sinusoidal effluxes were both significantly decreased. The pharmacokinetic modelling did not evidence that a high efflux into sinusoids can compensate for a decreased efflux into bile canaliculi. The direct quantification of transporter activities we proposed in the present study is innovative and might validate new modelling approaches. The experimental model is useful to quantify the functions of hepatocyte transporters in liver diseases.

The isolated and perfused rat liver is a convenient model, because the experimental conditions are well-controlled and simplified. The structure of the liver is maintained. However, livers are perfused only through the portal vein, avoiding the complexity of a dual-input entry but changing the physiology of liver perfusion. To simplify the protocol, we did not add proteins in the KHB solution, and BOPTA was free to enter into hepatocytes. The gamma counter placed over the liver detects the concentrations of imaging substrates every 20 s, avoiding the collection of serial biopsy samples that damage rat livers. An important assumption is that the region of interest measured by the counter is representative of the entire liver. Moreover, the hepatocyte concentrations we obtained averaged out the concentrations of numerous hepatocytes, knowing that Oatps are mainly expressed in perivenous hepatocytes where BOPTA is likely to enter. Additionally, extrapolation of the results to other species and humans must be cautious, because the expressions of membrane transporters can differ.
Author Contributions: Conceptualization, C.M.P. and V.V.; methodology, C.M.P.; formal analysis, C.M.P.; writing—original draft preparation, C.M.P.; writing—review and editing, V.V.; supervision, V.V.; and funding acquisition, C.M.P. All authors have read and agreed to the published version of the manuscript.

Funding: This research was funded by the Swiss National Science Foundation, grant number 310030-126030, and the APC was funded by the Swiss National Science Foundation.

Institutional Review Board Statement: This protocol was carried out in accordance with the Swiss Guidelines for the Care and Use of Laboratory Animals and was approved by the local animal welfare committee and the veterinary office in Geneva, Switzerland (No. 1006.3384.2).

Informed Consent Statement: Not applicable.

Data Availability Statement: The data presented in this study are available on request from the corresponding author.

Conflicts of Interest: The authors declare no conflict of interest. The funders had no role in the design of the study; in the collection, analyses, or interpretation of the data; in the writing of the manuscript; or in the decision to publish the results.

References
1. Fabbrini, E.; Sullivan, S.; Klein, S. Obesity and nonalcoholic fatty liver disease: Biochemical, metabolic, and clinical implications. Hepatology 2010, 51, 679–689. [CrossRef]
2. Geier, A.; Tiniakos, D.; Denk, H.; Trauner, M. From the origin of NASH to the future of metabolic fatty liver disease. Gut 2021, 70, 1570–1579. [CrossRef] [PubMed]
3. Tarantino, G.; Citro, V.; Capone, D. Nonalcoholic fatty liver disease: A challenge from mechanisms to therapy. J. Clin. Med. 2019, 9, 15. [CrossRef] [PubMed]
4. Vildhede, A.; Kimoto, E.; Pelis, R.M.; Rodrigues, A.D.; Varma, M.V.S. Quantitative proteomics and mechanistic modeling of transporter-mediated disposition in nonalcoholic fatty liver disease. Clin. Pharmacol. Ther. 2020, 107, 1128–1137. [CrossRef] [PubMed]
5. Dzierlenga, A.L.; Cherrington, N.J. Misregulation of membrane trafficking processes in human nonalcoholic steatohepatitis. J. Biochem. Mol. Toxicol. 2018, 32, e22035. [CrossRef]
6. Ali, I.; Slizgi, J.R.; Kaullen, J.D.; Ivanovic, M.; Niemi, M.; Stewart, P.W.; Barritt, A.S.; Brouwer, K.L.R. Transporter-mediated alterations in patients with NASH increase systemic and hepatic exposure to an OATP and MRP2 substrate. Clin. Pharmacol. Ther. 2018, 104, 749–756. [CrossRef]
7. Geier, A.; Tiniakos, D.; Denk, H.; Trauner, M. From the origin of NASH to the future of metabolic fatty liver disease. Gut 2021, 70, 1570–1579. [CrossRef] [PubMed]
8. Sanderson, S.; Ramalho, L.; Augusto, M.J.; da Silva, D.M.; Ramalho, F.S. Nonalcoholic steatohepatitis: A search for factual animal models. Biomed. Res. Int. 2015, 2015, 574832. [CrossRef]
9. Kucera, O.; Cervinkova, Z. Experimental models of non-alcoholic fatty liver disease in rats. World J. Gastroenterol. 2014, 20, 8364–8376. [CrossRef]
10. Pastor, C.M.; Wissmeyer, M.; Millet, P. Concentrations of Gd-BOPTA in cholestatic fatty rat livers: Role of transport functions through membrane proteins. Contrast Media Mol. Imaging 2013, 8, 147–156. [CrossRef]
11. Geier, A.; Dietrich, C.G.; Grote, T.; Beuers, U.; Prüfer, T.; Fraunberger, P.; Matern, S.; Gartung, C.; Gerbes, A.L.; Bilzer, M. Characterization of organic anion transporter regulation, glutathione metabolism and bile formation in the obese Zucker rat. J. Hepatol. 2005, 43, 1021–1030. [CrossRef]
12. Canet, M.J.; Cherrington, N.J. Drug disposition alterations in liver disease: Extrahepatic effects in cholestasis and nonalcoholic steatohepatitis. Expert Opin. Drug Metab. Toxicol. 2014, 10, 1209–1219. [CrossRef]
13. Zhang, L.; Xu, P.; Cheng, Y.; Wang, P.; Ma, X.; Liu, M.; Wang, X.; Xu, F. Diet-induced obese alters the expression and function of hepatic drug-metabolizing enzymes and transporters in rats. Biochem. Pharmacol. 2019, 164, 368–376. [CrossRef]
14. Vilgrain, V.; Van Beers, B.E.; Pastor, C.M. Insights into the diagnosis of hepatocellular carcinomas with hepatobiliary MRI. J. Hepatol. 2016, 64, 708–716. [CrossRef] [PubMed]
15. Planchamp, C.; Hadengue, A.; Steiger, B.; Bourquin, J.; Vonlaufen, A.; Frossard, J.L.; Quadri, R.; Becker, C.D.; Pastor, C.M. Function of both sinusoidal and canalicular transporters controls the concentration of organic anions within hepatocytes. Mol. Pharmacol. 2007, 71, 1089–1097. [CrossRef] [PubMed]
16. Millet, P.; Moulin, M.; Steiger, B.; Daali, Y.; Pastor, C.M. How organic anions accumulate in hepatocytes lacking Mrp2: Evidence in perfused rat livers. Drug Metab. Dispos. 2019, 47, 412–418. [CrossRef] [PubMed]
18. Pastor, C.M.; Morel, D.R.; Billiar, T.R. Oxygen supply dependence of urea production in the isolated perfused rat liver. *Am. J. Respir. Crit. Care Med.* **1998**, *157*, 796–802. [CrossRef] [PubMed]

19. Blouin, A.; Bolender, R.P.; Weibel, E.R. Distribution of organelles and membranes between hepatocytes and nonhepatocytes in the rat liver parenchyma. *A stereological study. J. Cell Biol.* **1977**, *72*, 441–455. [CrossRef]

20. Segmental Linear Regression. *GraphPad Software, La Jolla California, USA*. 2021. Available online: [https://www.graphpad.com/guides/prism/latest/curve-fitting/reg_segmental_linear_regression.htm](https://www.graphpad.com/guides/prism/latest/curve-fitting/reg_segmental_linear_regression.htm) (accessed on 15 September 2021).

21. One Phase Decay. *GraphPad Software, La Jolla California, USA*. 2021. Available online: [https://www.graphpad.com/guides/prism/latest/curve-fitting/reg_exponential_decay_1phase.htm](https://www.graphpad.com/guides/prism/latest/curve-fitting/reg_exponential_decay_1phase.htm) (accessed on 15 September 2021).

22. Marrone, J.; Danielli, M.; Gaspari, C.I.; Capiglioni, A.M.; Marinelli, R.A. Aquaporin gene transfer for hepatocellular cholestasis. *Biochimie* **2021**, In press. [CrossRef]

23. Marinelli, R.A.; Vore, M.; Javitt, N.B. Hepatic bile formation: Canalicular osmolarity and paracellular and transcellular water flow. *J. Pharmacol. Exp. Ther.* **2019**, *371*, 713–717. [CrossRef]

24. Ito, K.; Koresawa, T.; Nakano, K.; Horie, T. Mrp2 is involved in benzylpenicillin-induced choleresis. *Am. J. Physiol. Gastrointest. Liver Physiol.* **2004**, *287*, G42–G49. [CrossRef] [PubMed]

25. Pizarro, M.; Balasubramaniyan, N.; Solis, N.; Solar, A.; Duarte, I.; Miquel, J.F.; Suchy, F.J.; Trauner, M.; Accatino, L.; Ananthanarayan, M.; et al. Bile secretory function in the obese Zucker rat: Evidence of cholestasis and altered canalicular transport function. *Gut* **2004**, *53*, 1837–1843. [CrossRef] [PubMed]

26. Lickteig, A.J.; Fisher, C.D.; Augustine, L.M.; Aleksunes, L.M.; Besselsen, D.G.; Slitt, A.L.; Manautou, J.E.; Cherrington, N.J. Efflux transporter expression and acetaminophen metabolite excretion are altered in rodent models of nonalcoholic fatty liver disease. *Drug Metab. Dispos.* **2007**, *35*, 1970–1978. [CrossRef] [PubMed]