## Title page

# In vitro – in vivo extrapolation of OATP1B-mediated drug-drug interactions in cynomolgus monkey

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## Running title page

Running title: IVIVE of OATP DDIs in cynomolgus monkey

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#### List of non-standard abbreviations

ABT, 1-aminobenzotriazole; AUC, area under the plasma concentration-time curve; AUCR, ratio of the area under the plasma concentration-time curve; AUCpo, area under the plasma concentration-time curve of oral statin; AUC<sub>last</sub>, area under the plasma concentration-time curve from time 0 to the last time point;  $AUC_{0-\infty}$ , area under the plasma concentration-time curve from time 0 to infinity; BCA, bicinchoninic acid; BCRP, breast cancer resistance protein; CL<sub>active</sub>, active uptake clearance; CL<sub>diff</sub>, passive diffusion clearance; CL<sub>hepatic</sub>, hepatic clearance; CL<sub>iv</sub>, intravenous clearance; C<sub>max</sub>, maximum plasma concentration; CL<sub>repal</sub>, renal clearance; DDI(s), drug-drug interaction(s); DPBS, Dulbecco's Phosphate Buffered Saline; ECCS, extended clearance classification system; F, oral bioavailability; Fa, fraction of drug absorbed; Fg, fraction of drug escaping intestinal extraction; Fh, fraction of drug escaping hepatic extraction; fu,cell, fraction of unbound drug in the cell; fu,med, nonspecific binding in the media; gmfe, geometric mean fold error; HEK293, human embryonic kidney 293 cells; IC<sub>50</sub>, inhibitory constant (concentration at which 50% of total inhibitory effect is observed); IVIVE, in vitro-in vivo extrapolation; K<sub>i</sub>, reversible inhibition constant; K<sub>m</sub>, affinity constant; LC-MS/MS, liquid chromatography-tandem mass spectrometry; MRP2, multidrug resistance protein 2; NTCP, sodium taurocholate co-transporting polypeptide; OATP, organic anion transporting polypeptide; Q<sub>h</sub>, hepatic blood flow; t<sub>1/2</sub>, half-life; Vd<sub>ss</sub>, volume of distribution; V<sub>max</sub>, maximum uptake rate.

#### **Abstract**

Hepatic organic anion transporting polypeptides (OATP) 1B1 and 1B3 are clinically relevant transporters associated with significant drug-drug interactions (DDIs) and safety concerns. Given that OATP1Bs in cynomolgus monkey share >90% degree of gene and amino acid sequence identity with human orthologs, we evaluated the in vitro-in vivo translation of OATP1B-mediated DDI risk using this preclinical model. In vitro studies using cynomolgus monkey hepatocytes showed active uptake K<sub>m</sub> values of 2.0 and 3.9 μM for OATP1B probe substrates, pitavastatin and rosuvastatin, respectively. Rifampicin inhibited pitavastatin and rosuvastatin active uptake in monkey hepatocytes with IC<sub>50</sub> values of 3.0 and 0.54 µM, respectively, following pre-incubation with the inhibitor. Intravenous pharmacokinetics of <sup>2</sup>H<sub>4</sub>-pitavastatin and <sup>2</sup>H<sub>6</sub>rosuvastatin (0.2 mg/kg) and the oral pharmacokinetics of cold probes (2 mg/kg) were studied in cynomolgus monkeys (n=4) without or with co-administration of single oral ascending doses of rifampicin (1, 3, 10 and 30 mg/kg). A rifampicin dose-dependent reduction in intravenous clearance of statins was observed. Additionally, oral pitavastatin and rosuvastatin plasma exposure increased up to 19- and 15fold at the highest dose of rifampicin, respectively. Use of IC<sub>50</sub> obtained following 1h pre-incubation with rifampicin (0.54 µM) predicted correctly the change in mean intravenous clearance and oral exposure of statins as a function of mean unbound C<sub>max</sub> of rifampicin. This study demonstrates quantitative translation of in vitro OATP1B IC50 to predict DDIs using cynomolgus monkey as a preclinical model and provides further confidence in application of in vitro hepatocyte data for the prediction of clinical OATP1B-mediated DDIs.

#### Introduction

Drug-drug interactions (DDIs) involving hepatic organic anion transporting polypeptides (OATPs) are widely recognized as clinically important due to potential serious adverse events associated with them (Giacomini et al., 2010; Yoshida et al., 2012; El-Kattan et al., 2016; Galetin et al., 2017). Therefore, there is a strong need to quantitatively predict OATP-mediated DDIs early in candidate identification and drug development. Despite tremendous strides in establishing *in vitro* tools for assessing transporter role, confidence in quantitative prediction of transporter-mediated DDIs using *in vitro* data is arguably still low-to-moderate (Zamek-Gliszczynski et al., 2013; Jones et al., 2015; Yoshida et al., 2017).

Recently, cynomolgus monkey has been increasingly evaluated as a potential animal model for the assessment of OATP1B-mediated DDIs (Shen et al., 2013; Takahashi et al., 2013; Chu et al., 2015; Shen et al., 2015; Takahashi et al., 2016) due to >90% degree of gene and amino acid sequence identity between cynomolgus monkey and human orthologs for OATP1B uptake transporters (Ebeling et al., 2011; Shen et al., 2013; Takahashi et al., 2013). In addition to clinical drug probes, increasing evaluation of endogenous biomarkers for OATP1B DDIs (e.g., coproporhyrin I, bile acids) has been reported in this preclinical species (Chu et al., 2015; Watanabe et al., 2015; Shen et al., 2016; Thakare et al., 2017). DDIs reported in cynomolgus monkey for statins including rosuvastatin, pitavastatin and atorvastatin showed a good agreement in DDI classification (strong/moderate) relative to those observed in humans (Table S1, Supplementary Material). However, direct comparison of hepatic transporter-mediated DDIs between humans and monkeys may be challenging due to species differences in dosing regimen, transporter expression levels, activities and/or mechanisms involved in drug clearance. For instance, cynomolgus monkey liver expression of OATP1B1 and OAT1B3 was shown to be ~6- to 13-fold higher compared to human liver, respectively (Wang et al., 2015). In the case of OATP2B1 and sodium taurocholate co-transporting polypeptide (NTCP), the protein levels were either ~6-fold greater in human liver or comparable between the two species, respectively (Wang et al., 2015). Nevertheless, an in vitro-in vivo extrapolation (IVIVE) strategy using either static or physiologically-based pharmacokinetic (PBPK) modelling approaches accounting for the species differences could be employed for predicting clinical DDI risk.

Discrepancy in the translation of inhibitory potency (IC<sub>50</sub> or K<sub>i</sub>) to in vivo was reported for OATP inhibitors

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(Varma et al., 2012; Li et al., 2014; Yoshikado et al., 2017). For example, rifampicin *in vitro* K<sub>i</sub> values were shown to be several fold higher than the *in vivo* K<sub>i</sub> estimated from the clinical DDI data using mechanistic modelling (Varma et al., 2014; Barnett et al., 2017; Yoshikado et al., 2017). In addition, a number of studies demonstrated potentiation of OATP1B inhibition following pre-incubation with the inhibitor, trend particularly evident for cyclosporine (Amundsen et al., 2010; Gertz et al., 2013; Izumi et al., 2015; Takahashi et al., 2016; Pahwa et al., 2017) and incorporated in the recent FDA DDI guidance document (FDA, 2017). While the clinical DDIs with cyclosporine are well recovered with the IC<sub>50</sub> obtained following pre-incubation (Gertz et al., 2013), there is limited understanding of the significance of pre-incubation in predicting DDIs of other inhibitor drugs. Additionally, recent studies reported substrate-dependence in the *in vitro* inhibition data for OATP1B inhibitors (Noé et al., 2007; Izumi et al., 2013). Due to the gaps in the available clinical DDI data, understanding of the translation of pre-incubation and substrate-dependent inhibition *in vitro* phenomena is still ambiguous.

Using cynomolgus monkey as a preclinical model, our goals in the current study are (i) to evaluate the predictability of OATP1B-mediated DDIs from *in vitro* inhibition data, and (ii) to understand the *in vivo* relevance of the effect of pre-incubation and substrate-dependency in rifampicin inhibition potential measured *in vitro*. In this study, the *in vitro* inhibition of transporter-mediated uptake of rosuvastatin and pitavastatin was investigated after incubation of cynomolgus monkey primary hepatocytes either with buffer or OATP inhibitors (rifampicin, cyclosporine and rifamycin SV). Prior to inhibition studies, an uptake kinetic characterization was performed for these two statins. Pitavastatin and rosuvastatin represent extended clearance classification system (ECCS) class 1B and class 3B, respectively, where OATP-mediated hepatic uptake is the rate-determining step in the systemic clearance in human (Varma et al., 2015; Varma et al., 2017). To allow separate evaluation of inhibitory effect of hepatic versus intestinal disposition, pharmacokinetics of both statins were studied following simultaneous intravenous (stable-labelled) and oral (cold) administration to cynomolgus monkey (n=4 animals) and over a wide rifampicin dose range (1-30 mg/kg). Finally, the extrapolation of rifampicin *in vitro* inhibition potency data obtained

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using monkey hepatocytes to the *in vivo* changes in systemic clearance and plasma (i.v./oral) exposure of statins was evaluated.

#### **Materials and Methods**

Chemicals and Reagents. 3'-Phosphoadenosine-5'-phosphosulfate (PAPS), simvastatin, rifamycin SV, rifampicin, 1-aminobenzotriazole (ABT), cyclosporine A, naloxone, and tolbutamide were purchased from Sigma-Aldrich (St Louis, MO, USA). Deuterium labeled rifampicin (<sup>2</sup>H<sub>8</sub>-rifampicin) was obtained from ALSACHIM (Illkirch, Graffenstaden, France). Pitavastatin and rosuvastatin were purchased from Sequoia Research Products Ltd. (Oxford, UK). Deuterium labeled pitavastatin (<sup>2</sup>H<sub>4</sub>-pitavastatin) and rosuvastatin (<sup>2</sup>H<sub>6</sub>-rosuvastatin) were purchased from Clearsynth (Ontario, Canada). Atorvastatin was purchased from Toronto Research Chemicals (Toronto, Canada). InVitroGro CP hepatocyte medium and Torpedo antibiotic mix were purchased from InVitro GmbH (Frankfurt, Germany). Collagen I coated 24-well plates were obtained from VWR International (Leicestershire, UK). Cryopreserved cynomolgus monkey hepatocytes (female, pooled lot 10353012) were purchased from *In vitro* ADMET Laboratories, LLC (Columbia, Maryland). Bicinchoninic acid (BCA) protein assay kit was purchased from Life Technologies Ltd (Paisley, UK). Dulbecco's Phosphate Buffered Saline (DPBS) was obtained from Life Technologies Ltd (Paisley, UK). Acetonitrile, water, ammonium hydroxide were obtained from Fisher Scientific (Fair Lawn, New Jersey). Methanol was purchased from Fisher Scientific (Fair Lawn, New Jersey) and VWR International (Leicestershire, UK).

In vitro transport studies using monkey hepatocytes. Cryopreserved cynomolgus monkey hepatocytes were thawed in pre-warmed InVitroGRO CP medium supplemented with torpedo antibiotic mix (2.2% v/v) according to the protocol from InVitro GmbH and cell viability was determined by Trypan Blue exclusion method. Hepatocyte suspension was diluted to 0.7 x 10<sup>6</sup> cells/mL with the prepared InVitroGRO CP medium and hepatocytes were seeded into collagen I-coated 24-well plates at a density of 350,000 cells per well. Cells were cultured for 4h at 37°C and 5% CO<sub>2</sub> in an incubator to allow attachment to the collagen. Cell confluency and monolayer formation was visually assessed before each experiment. Both the hepatic transporter uptake and its inhibition were investigated in plated cynomolgus monkey hepatocytes 4h post seeding using rosuvastatin and pitavastatin as probe substrates. Uptake was measured over a range of concentrations (0.1-0.3-1-3-10-30-100 μM) for 30, 60, 90 and 120 s at 37°C in triplicate to determine uptake kinetics, as described previously (Ménochet et al., 2012).

Inhibition of rosuvastatin and pitavastatin hepatic uptake by monkey hepatocytes was assessed in triplicate using the OATP inhibitor rifampicin (0.01-100 µM). In addition, inhibition with cyclosporine, (0.01-6 μM), and rifamycin SV (0.01-1000 μM) was investigated. Cyclosporine was included as a strong OATP inhibitor with evidence of pre-incubation effect in human in vitro systems (Gertz et al., 2013; Izumi et al., 2015), whereas rifamycin SV was considered as dual inhibitor of OATPs and NTCP (Bi et al., 2017). Rosuvastatin and pitavastatin concentrations used in the inhibition studies were 1 and 0.3 µM, respectively. The medium was removed after plating and cell monolayers were rinsed twice with prewarmed DPBS. Effect of pre-incubation on the inhibition of hepatic uptake transporters in cynomolgus monkey hepatocytes was investigated by the addition of an inhibitor solution (400 µL) on the cell monolayer for 1h. In control group, pre-incubation was performed with DPBS containing 1 mM ABT. Following pre-incubation, cell monolayers were co-incubated with pre-warmed DPBS containing the inhibitor and the probe substrate for 3 min. This incubation was stopped by the removal of the medium and rinsing of the cell monolayers three times with 800 µL of ice cold DPBS. The cell monolayers were lysed in 200 µL ice cold deionised water for subsequent liquid chromatography-tandem mass spectrometry (LC-MS/MS) analysis. The in vitro sample preparation and LC-MS/MS analysis are described in Supplementary Material, Section S1. The LC-MS/MS conditions for each individual drug and their corresponding internal standards are detailed in Supplementary Material, Table S2.

In vitro data analysis. Determination of the uptake kinetic parameters including the affinity constant  $K_m$  (µM), the maximum uptake rate  $V_{max}$  (pmol/min/ $10^6$  cells), passive diffusion clearance  $CL_{diff}$  (µL/min/ $10^6$  cells) and fraction unbound in the cell  $f_{u,cell}$  for rosuvastatin and pitavastatin was performed using the two-compartment mechanistic model in Matlab 7.14 (2012) (The MathWorks,Inc., Natick, MA), as described previously (Ménochet et al., 2012). The active uptake clearance  $CL_{active}$  (µL/min/ $10^6$  cells) was estimated from the  $V_{max}$  to  $K_m$  ratio. Parameter estimates were corrected for nonspecific binding in the media ( $f_{u,med}$ ) which was calculated from the slope of the linear regression of the unbound substrate concentration extrapolated at t=0 vs the initial media concentration curve. The cellular concentrations were normalized for protein content as measured by the BCA protein assay kit at the end of incubation. Hepatocyte volume was set to 3.9  $\mu$ L/ $10^6$  cells as in rat (Reinoso et al., 2001); conversion of monkey hepatocyte data

expressed per mg protein to M cells was based on the 1:1 relationship between mg protein and M cells (in house data).

The data on substrate uptake (expressed as % control) at each inhibitor concentration were used to estimate the half-maximal inhibitory concentration  $IC_{50}$  ( $\mu$ M) of the inhibitors used. The analysis was performed in GraFit<sup>TM</sup> v6.0 (Erithacus Software Ltd, Horley, UK) by fitting a nonlinear least squares regression model as shown in Eq. 1 to the experimental data:

Substrate uptake (% control) = 
$$\frac{Max - Min}{1 + \left(\frac{I}{IC_{50}}\right)^s} + Min$$
 (1)

where Max and Min represent the fitted maximum and minimum uninhibited uptake, respectively, I is the inhibitor concentration, corrected for  $f_{u,med}$  (0.85, 0.7 and 0.95 for rifampicin, cyclosporine and rifamycin SV, respectively), and s is the slope factor. To increase the precision of the  $IC_{50}$  estimates, the minimum uptake was fixed to the experimental data. In all cases, the experimental data for minimum uptake was within 25% of the estimated value. Statistical analysis of the pre-incubation effect on  $IC_{50}$  was performed using a paired t-test where p<0.05 was considered as statistically significant.

In vivo studies in cynomolgus monkeys. All procedures performed on these animals were in accordance with regulations and established guidelines and were reviewed and approved by Pfizer Institutional Animal Care and Use Committee, and were conducted at Pfizer Groton (Connecticut, US). Male Cynomolgus macaque Mauritian monkeys (approximately 6 to 8.5 years of age) were used for these studies. A cross-over study design was employed, where the same four animals were dosed over a series of five studies, following a minimum one-week wash-out period between each study. One exception was the 3 mg/kg rifampicin dose group, where one of four monkeys was dosed only in that single study. Animals were provided a normal food schedule the day before the study (meals at 8:00 am and 11:00 am, with one treat daily) and were allowed free access to water. Animals were housed in metabolism cages during sample collection. On the day of the study, monkeys were fed at approximately 1h and 3h post-dose and allowed water ad libitum. Rifampicin was administered via oral gavage at 0 (blank vehicle), 1, 3, 10, and 30 mg/kg, at a dose volume of 2 ml/kg in a 0.5% (w/v) methylcellulose (in water) suspension. Rifampicin administration was immediately followed by oral doses of pitavastatin and rosuvastatin at a

dose of 2 mg/kg. Approximately one hour and 15 minutes following the oral rifampicin administration, <sup>2</sup>H<sub>4</sub>pitavastatin and <sup>2</sup>H<sub>6</sub>-rosuvastatin were administered via intravenous (i.v.) bolus (cephalic vein), at dose of 0.2 mg/kg, in a dosing volume of 0.2 ml/kg; 2% dimethyl sulfoxide (DMSO) (v/v) and 98% of TRISbuffered saline (pH ~7.7). All i.v. formulations were sterile filtered prior to administration. Serial blood samples were collected via the femoral vein into K<sub>2</sub>EDTA tubes prior to dosing and then at 0.083, 0.25, 0.5, 0.75, 1, 2, 3, 5, 6, and 24 hours post i.v. dosing. Blood samples were stored on wet ice prior to being centrifuged to obtain plasma (3000 RPM, 10 minutes at 4°C; Jouan BR4i refrigerated centrifuge). Urine was also collected on wet ice, pre-dose and at intervals of 0 to 6 hours and 6 to 24 hours post-dose. Due to the potential instability of rifampicin and possible inter-conversion of lactone and acid forms of pitavastatin or rosuvastatin, each plasma and urine sample was equally divided into two aliquots prior to being stored frozen. The first aliquot was untreated matrix, while the second aliquot was added to an equal volume of 0.1 M sodium acetate buffer (pH 4). All urine and plasma samples, treated and untreated, were kept cold during collection, after which they were stored frozen at -20°C. The analysis of <sup>2</sup>H<sub>4</sub>pitavastatin, <sup>2</sup>H<sub>6</sub>-rosuvastatin, pitavastatin, rosuvastatin and rifampicin in plasma samples by LC-MS/MS was described in Supplementary Material, Section 2. The analytes were monitored using multiple reaction monitoring with settings listed in Supplementary Material, Table S3.

Pharmacokinetic analysis and DDI predictions. Analyst 1.4.2 software (SCIEX, Framingham, MA) was used for LC-MS peak integration of plasma and urine samples. Raw data were imported into Watson LIMS<sup>TM</sup> version 7.4 (Thermo Fisher Scientific Inc, Waltham, MA) for standard curve regression and non-compartmental pharmacokinetic parameter calculations – area under the plasma concentration-time curve (AUC), maximum plasma concentration ( $C_{max}$ ), intravenous clearance ( $CL_{iv}$ ), volume of distribution ( $Vd_{ss}$ ) and half-life ( $t_{1/2}$ ). Other parameters were subsequently calculated on the basis of pharmacokinetic first principles. Oral bioavailability (F) of statins was described by Eq. 2:

$$F = F_a \cdot F_g \cdot F_h$$
 (2)

where  $F_a$ ,  $F_g$  and  $F_h$  represent fraction of drug absorbed, fraction of drug escaping intestinal and hepatic extraction, respectively. The  $F_h$  was described by Eq. 3.

$$F_h = 1 - \frac{CL_{hepatic}}{Q_h}$$
 (3)

where  $CL_{hepatic}$  represents hepatic blood clearance (plasma  $CL_{hepatic}$ /blood-to-plasma ratio), and  $Q_h$  is hepatic blood flow in cynomolgus monkey (44 mL/min/kg) (Hosea et al., 2009). Measured blood-to-plasma ratios of rosuvastatin and pitavastatin in cynomolgus monkey were 0.55 and 0.58, respectively.

The ratio of the area under the plasma concentration-time curve (AUCR) of oral statins in the presence (AUC'<sub>po</sub>) and absence (AUC<sub>po</sub>) of rifampicin was predicted based on Eq 4.

$$AUCR = \frac{AUC'_{po}}{AUC_{po}} = \frac{F_{a'}}{F_{a}} \cdot \frac{F_{g'}}{F_{g}} \cdot \frac{F_{h'}}{F_{h}} \cdot \frac{(CL_{\dot{N}})}{(CL_{\dot{N}}')} \tag{4}$$

where ' indicates parameters in the presence of rifampicin. For the prediction purposes, it was assumed that rifampicin has no impact on  $F_a$  and  $F_g$  of both statins, and therefore  $F_a$ ' and  $F_g$ ' were the same as in the control phase.  $F_h$ ' was estimated from the hepatic blood clearance in the presence of rifampicin ( $CL_{hepatic}$ ') as shown in Eq. 5.

$$F_h' = 1 - \frac{CL_{hepatic'}}{Q_h}$$
 (5)

The CL<sub>iv</sub> of statins is the sum of hepatic and renal clearance and assuming uptake is the rate-determining step for hepatic clearance it can be expressed as shown in Eq. 6:

$$CL_{iv} = CL_{hepatic} + CL_{renal} = CL_{active} + CL_{diff} + CL_{renal}$$
 (6)

where CL<sub>renal</sub>, CL<sub>active</sub> and CL<sub>diff</sub> represent renal, sinusoidal active uptake and passive diffusion clearance, respectively.

In vivo  $IC_{50}$  values were estimated using the Eq. 7 using unbound  $C_{max}$  of rifampicin ( $[C_{u,max}]$ ) as independent variable and intravenous clearance of statins as the depending variable.

$$CL_{iv}' = \left[ \frac{CL_{active}}{1 + \frac{[C_{u,max}]}{IC_{50}}} + CL_{diff}' \right] + CL_{renal}'$$
 (7)

CL<sub>diff</sub> was obtained from the data in the presence of the highest rifampicin dose assuming complete inhibition of active hepatic uptake (Eq. 8). It was assumed that rifampicin had no impact on the passive

diffusion and renal clearance of both statins, and therefore  $CL_{diff}$  and  $CL_{renal}$  were the same as  $CL_{diff}$  and  $CL_{renal}$ .

$$CL_{diff} = CL_{iv(+30\,mg/kg\,rifampicir)} - CL_{renal} \tag{8}$$

Geometric mean fold error (gmfe) was calculated to assess the bias of the predicted rosuvastatin and pitavastatin AUCR across rifampicin dose range, as shown in Eq. 9 (Gertz et al., 2010).

$$gmfe = 10^{\frac{1}{N} \sum \left| log \frac{predicted}{observed} \right|}$$
 (9)

where N represents the number of observations.

#### Results

*In vitro* uptake kinetics of rosuvastatin and pitavastatin in monkey hepatocytes. Uptake of rosuvastatin and pitavastatin (0.1 to 100 μM) was investigated after incubation with plated cynomolgus monkey hepatocytes and the kinetic parameters were estimated using the mechanistic two-compartment model. The measured f<sub>u,med</sub> was 0.83 and 0.99 for rosuvastatin and pitavastatin, respectively, which was used for correction of the initial media concentrations of both drugs. Kinetic profiles of both drugs in monkey hepatocytes demonstrated a saturable and nonsaturable uptake phase (Figure 1, Table 1). The K<sub>m</sub> of rosuvastatin and pitavastatin were 3.29 and 1.99 μM, respectively. Although pitavastatin active uptake clearance was greater than for rosuvastatin (CL<sub>active</sub> of 109 μL/min/10<sup>6</sup> cells *vs.* 98.3 μL/min/10<sup>6</sup> cells), the contribution of active process to total uptake was approximately 96% in case of rosuvastatin versus 80% for pitavastatin. The CL<sub>diff</sub> was approximately 6-fold higher for pitavastatin (26.5 μL/min/10<sup>6</sup> cells) than in the case of rosuvastatin (4.27 μL/min/10<sup>6</sup> cells). Furthermore, the extent of intracellular binding differed between the two probes, as f<sub>u,cell</sub> was 0.25 and 0.024 for rosuvastatin and pitavastatin, respectively.

In vitro uptake inhibition potency of rifampicin and cyclosporine in monkey hepatocytes. Both rifampicin and cyclosporine inhibited OATP-mediated rosuvastatin and pitavastatin uptake in a concentration-dependent manner (Figure 2). For rosuvastatin, up to 88% inhibition was observed after pre-incubation with rifampicin (Figure 2A). The uninhibited uptake of rosuvastatin (12%) was in close agreement with the contribution of passive diffusion (4%) estimated from the uptake kinetic study in monkey hepatocytes. The  $IC_{50}$  of rifampicin obtained after pre-incubation with buffer (1.14  $\mu$ M) was reduced by approximately 2-fold after pre-incubation with rifampicin (0.54  $\mu$ M) (Table 2). This marginal (2-fold) effect of pre-incubation on rifampicin potency seen with rosuvastatin reflects similar findings reported for this inhibitor in human OATP1B1 and OATP1B3 transfected HEK293 cells with estradiol-17 $\beta$ -glucuronide as a probe (2- and 3-fold increase in potency, details in Supplementary Table S4). Cyclosporine showed similar extent of maximal inhibition of rosuvastatin uptake in monkey hepatocytes at the highest concentration (Figure 2C). However, the increase in cyclosporine inhibition potency following pre-incubation was more pronounced (7-fold shift) (Table 2).

In the case of pitavastatin, maximal 55% inhibition of uptake was observed at the highest rifampicin concentration and after pre-incubation with inhibitor (Figure 2B). There was no significant change in the IC<sub>50</sub> value after pre-incubation with rifampicin relative to pre-incubation with buffer (Table 2). This is in agreement with the data reported in HEK293 cells expressing cynomolgus monkey OATP1B1/1B3 using pitavastatin as a probe, where pre-incubation with inhibitor resulted in no significant change in rifampicin IC<sub>50</sub> (Supplementary Table S4). Discrepancy between the uninhibited uptake (45%) and contribution of passive diffusion (20%) estimated from uptake kinetic data was apparent in case of pitavastatin. Cyclosporine also showed incomplete inhibition of pitavastatin uptake (Figure 2D), but in contrast to rifampicin its inhibitory potency increased 3.7-fold following pre-incubation with the inhibitor (Table 2). Additionally, reduction in the steepness of the IC<sub>50</sub> curves was observed after pre-incubation in particular with cyclosporine (Figure 2C and D). DDI studies with rifamycin SV were also performed to elucidate the possible involvement of NTCP in pitavastatin uptake, which would not be inhibited by rifampicin. The maximum inhibition of pitavastatin uptake increased to 64% after pre-incubation with rifamycin SV, suggesting some contribution of NTCP in pitavastatin uptake in monkey hepatocytes (Figure S1, Supplementary Material). No effect of pre-incubation on rifamycin SV inhibition potency was observed in plated monkey hepatocytes (Table 2). Assuming almost complete inhibition of transporter-mediated uptake of pitavastatin at 1 mM rifamycin SV (Thakare et al., 2017), this uninhibited uptake should represent the contribution of passive diffusion to total cellular uptake, which in this case was higher compared with estimates obtained by mechanistic modelling of kinetic data done over short incubation times. Such incomplete inhibition of uptake for pitavastatin has previously been reported in monkey hepatocytes using rifampicin and cyclosporine (Takahashi et al., 2013) and in human hepatocytes using rifampicin (Pahwa et al., 2017) and may reflect the combination of passive diffusion and involvement of transporter uptake not inhibited by these inhibitors.

Dose-dependent effect of rifampicin on i.v. and oral pharmacokinetics of rosuvastatin and pitavastatin in monkeys. Intravenous pharmacokinetics of stable-labeled rosuvastatin and pitavastatin and the oral pharmacokinetics of cold rosuvastatin and pitavastatin were measured in cynomolgus monkeys after single oral ascending does of rifampicin (1 mg, 3 mg, 10 mg and 30 mg), and compared

with the vehicle only dosing (control). The plasma concentration-time profiles and the corresponding pharmacokinetic parameters are shown in Figure 3 and 4 and Table 3. Rosuvastatin i.v. clearance, when dosed along with 30 mg/kg rifampicin, decreased to almost 50% of control, while there was no significant change in the volume of distribution. This is in agreement with the previously reported marginal decrease in Vd<sub>ss</sub> of i.v. rosuvastatin when given with a lower rifampicin dose in cynomolgus monkey (Chu et al., 2015). Oral AUC and C<sub>max</sub> of rosuvastatin were increased up to 15-fold by the highest dose of rifampicin. In case of pitavastatin, reduction in i.v. clearance was more prominent (18.0 mL/min/kg versus 4.3 mL/min/kg), and the volume of distribution was reduced from 1.8 L/kg (control) to 0.55 L/kg (30 mg/kg rifampicin group). The decrease in intravenous clearance was accompanied by a significant decrease in Vd<sub>ss</sub>, consistent with a previous report (Takahashi et al., 2013), and therefore, t<sub>1/2</sub> was not altered significantly by rifampicin. Rifampicin dose-dependent increase in oral pitavastatin plasma exposure was noted with AUC change of up to ~19-fold and C<sub>max</sub> increase of up to ~12-fold at the highest dose tested (30 mg/kg) compared to the control group.

The lactone forms of orally administered pitavastatin and rosuvastatin were also measured in plasma samples. The plasma  $AUC_{0-\infty}$  of the lactone forms in the absence and presence of rifampicin and the lactone-acid ratios are reported in the Supplementary Table S5 and S6, respectively. No significant increase in the overall lactone plasma  $AUC_{0-\infty}$  was observed for rosuvastatin (Supplementary Table S5). In the case of pitavastatin lactone, a significant increase in its plasma  $AUC_{0-\infty}$  was observed at 3 mg/kg rifampicin. No clear dose dependent trend in lactone  $AUC_{0-\infty}$  of both statins was apparent. A significant increase in lactone-acid  $AUC_{0-\infty}$  ratios was observed for rosuvastatin only at 1 mg/kg rifampicin relative to control, whereas no significant changes were seen for pitavastatin at any rifampicin dose (Supplementary Table S6). There was an apparent decrease in lactone-acid  $AUC_{0-\infty}$  ratios for both statins particularly at the highest rifampicin doses. Previously a decrease and no change in lactone-acid ratio of pitavastatin (in human) and rosuvastatin (in cynomolgus monkey), respectively was observed in the presence of rifampicin (Prueksaritanont et al., 2014; Chu et al., 2015). Given that the AUC ratio of a metabolite to parent reflects both the formation and subsequent elimination of the metabolite (Houston, 1981), such behavior is not unexpected.

IVIVE of OATPs inhibition. In vivo inhibitory potency values estimated using unbound C<sub>max</sub> of rifampicin were  $0.99 \pm 3.8 \,\mu\text{M}$  and  $0.22 \pm 0.14 \,\mu\text{M}$  against  $\text{CL}_{\text{iv}}$  of rosuvastatin and pitavastatin, respectively (Figure 6, Table 2). In vitro IC<sub>50</sub> (0.54 μM), obtained in monkey hepatocytes following rifampicin pre-incubation and using rosuvastatin as probe substrate, described reasonably well rifampicin dose-dependent inhibition of rosuvastatin CLiv. In contrast, in vitro IC50 obtained using pitavastatin as probe substrate did not recover the in vivo inhibition activity of rifampicin against pitavastatin CLiv. However, in vitro IC50 generated using rosuvastatin described well the in vivo data of pitavastatin. Subsequently, in vitro IC50 value of 0.54  $\mu$ M and inhibitor unbound  $C_{max}$  were employed to predict the rifampicin dose-dependent change in AUC of both statins dosed either intravenously or orally (Figure 7). The prediction bias calculated from the gmfe was <1.6-fold for both statins, with 78 and 88% of the predicted AUCR values within 2-fold of the observed data for rosuvastatin and pitavastatin, respectively. An apparent underprediction was noted for the rosuvastatin oral AUC ratio at higher doses of rifampicin. This is likely due to increased oral absorption as a result of inhibition of rosuvastatin intestinal efflux, which was not captured in the current prediction model (see Methods). To note, exploration of IVIVE using IC $_{50}$  of 0.42 µM obtained in HEK293 cells expressing cOATP1B1 (lowest value compared to those obtained for other OATPs) and using rosuvastatin as an OATP probe (Shen et al., 2013) resulted in either no or marginal improvement in the prediction bias, highlighting the physiological relevance of monkey hepatocytes for comparison with the inhibition parameters obtained from the *in vivo* data in a top down manner.

The  $F_aF_g$  of rosuvastatin estimated from the intravenous and oral data of each study arm showed on average a 178% (2.8-fold) increase in rosuvastatin  $F_aF_g$  in monkeys receiving 30 mg/kg rifampicin relative to the control arm; this trend was not evident for pitavastatin. Correction of the average predicted AUCR values shown in Figure 7A by this increase in  $F_aF_g$  reduced the underprediction of the magnitude of rosuvastatin DDI at the highest dose of rifampicin (data not shown).

**Discussion** 

In comparison to the increased success of quantitative prediction of CYP-mediated DDIs, the confidence in the successful prediction of OATP-mediated DDIs using *in vitro* data is still low (Gertz et al., 2013; Prueksaritanont et al., 2014; Jones et al., 2015; Bi et al., 2017; Yoshida et al., 2017). Therefore, the objective of our study was to evaluate the IVIVE of OATP inhibition potential using cynomolgus monkey as a preclinical model and to gain confidence in mechanistic translational approach to predict clinical DDIs mediated by these transporters. Furthermore, it was aimed at improving our understanding of the utility of the cynomolgus monkey as a model for human to drive evaluation of clearance mechanisms (rate-determining step) and DDI risk. Collective results suggest that quantitative OATP1B DDI predictions can be made from *in vitro* IC<sub>50</sub> measured in primary hepatocytes using multiple probe substrates following pre-incubation with the inhibitor.

In the current study, both rosuvastatin and pitavastatin showed a high affinity for uptake transport in monkey hepatocytes (Table 1). Rosuvastatin CL<sub>active</sub> was approximately 4-fold greater in the current monkey donor investigated compared to the previous data (Shen et al., 2013), possibly reflecting donor differences in the transporter activity, as the estimated K<sub>m</sub> values were comparable. Although CL<sub>diff</sub> was about 6-fold greater in the donor used in the current study, contribution of the passive process to the overall uptake of rosuvastatin (4.2%) was in agreement with previous reports in monkey hepatocytes (Shen et al., 2013).

In this study, the effect of pre-incubation on the inhibition of OATP-mediated rosuvastatin and pitavastatin uptake in monkey hepatocytes was investigated using prototypical inhibitors rifampicin and cyclosporine. In contrast to the marginal decrease in rifampicin  $IC_{50}$  after pre-incubation, more pronounced increase in cyclosporine potency (up to 7-fold) was seen regardless of the substrate probe used. This clear pre-incubation effect on cyclosporine inhibition potency demonstrated here in monkey hepatocytes is in agreement with previous literature reports on this inhibitor. But the reported magnitude of shift in cyclosporine potency varied between *in vitro* systems and probes used (Table S4, Supplementary Material), with up to 22- and 23-fold increase in cyclosporine potency noted after pre-incubation in human

and monkey OATP1B1 transfected HEK293 cells, respectively, highlighting that the effect of preincubation on the potency of OATP inhibitors is dependent not only on the substrate used but also on the cellular system investigated.

Incomplete inhibition of uptake was observed for pitavastatin in the current study. As a number of reports demonstrated the contribution of sodium-dependent NTCP to the uptake of pitavastatin and rosuvastatin in human (Bi et al., 2013; Bi et al., 2017) and monkey hepatocytes (Thakare et al., 2017), pitavastatin uptake was further evaluated in the presence of rifamycin SV. Although rifamycin SV increased the maximal inhibition of pitavastatin uptake in monkey hepatocytes, incomplete inhibition of uptake was still evident, highlighting potentially lower transporter activity and/or higher passive contribution in the pooled donor investigated.

The quantitative translation of interaction noted in monkeys to humans may not be straightforward due to possible species difference in the multiple mechanisms involved in the clearance. However, several recent reports suggested good agreement between cynomolgus monkey and human in the magnitude of DDIs with rifampicin as an OATP inhibitor (Shen et al., 2013; Watanabe et al., 2015). In this study, rifampicin showed a dose-dependent effect with no further change in statins pharmacokinetics between rifampicin doses of 10 and 30 mg/kg in monkeys. The unbound C<sub>max</sub> of rifampicin achieved (~2 to 7 μM) at these doses are comparable to the unbound C<sub>max</sub> in human following single 600 mg dose (Varma et al., 2012; Prueksaritanont et al., 2014; Yoshikado et al., 2017). However, the magnitude of change in the oral AUC for both statins observed in the cynomolgus monkeys (~15 to 20-fold, Table 3) is much higher than noted in humans (AUC ratio ~5) (Prueksaritanont et al., 2014). In contrast, an earlier study showed ~3fold increase in oral rosuvastatin AUC in cynomolgus monkey with rifampicin dose of 15 mg/kg orally (Shen et al., 2013). While the reasons for such a difference are not apparent, a relatively smaller magnitude of rosuvastatin DDI in human may be attributed to the contribution of hepatic OATP2B1 which is not affected by rifampicin at in vivo relevant plasma concentrations (OATP2B1 in vitro IC<sub>50</sub>>60 µM). However, the contribution of OATP2B1 to rosuvastatin hepatic uptake in monkey is likely to be minor because of its relatively low expression in cynomolgus liver (Wang et al., 2015). In addition to OATPs,

both rosuvastatin and pitavastatin are substrates of efflux transporters such as multidrug resistance

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protein 2 (MRP2) and breast cancer resistance protein (BCRP) (Prueksaritanont et al., 2014; Lee et al., 2015). As rifampicin is also a substrate of OATP1Bs (Yamaguchi et al., 2011), its inhibitory effect on the biliary efflux of rosuvastatin and pitavastatin was considered. Previously, Chu et al. (2015) showed relatively weak inhibition of cMRP2 by rifampicin in HEK293 cells (IC $_{50}$  of 118  $\mu$ M). Assuming the reported unbound liver-to-plasma concentration ratio (Kp $_{uu}$ ) of 3.3 for rifampicin in humans (Chu et al., 2015) is comparable to that in cynomolgus monkey, the IC $_{50}$  for cMRP2 is much higher than the estimated rifampicin unbound liver concentrations (approximately 23  $\mu$ M at the highest rifampicin dose used), suggesting that the increase in plasma exposure of both statins is unlikely to be caused by MRP2 inhibition. In case of potential MRP2 inhibition, expectation is that liver AUC will change, consistent with an understanding of the rate-determining processes that affect hepatic exposure of these drugs (Tsamandouras et al., 2015).

The increase in  $F_aF_g$  of rosuvastatin seen in this study is most likely attributed to the effect of rifampicin on intestinal BCRP. Rosuvastatin absorption in humans is approximately 50% of the oral dose (Martin et al., 2003), whereas only ~20% was noted in monkeys (control arm). This difference in  $F_aF_g$  values suggests species differences in the expression of the intestinal transporters including BCRP. This is supported by previous reports of generally greater mRNA expression of intestinal efflux transporters in cynomolgus monkey relative to human (60-fold difference in the case of BCRP) (Takahashi et al., 2008). While, we cannot assess the differential effects of OATP2B1 uptake versus BCRP efflux definitively, increased absorption following rifampicin administration implies that BCRP efflux plays a prominent role in the oral absorption of rosuvastatin in monkey. In contrast, pitavastatin is a highly permeable drug and the  $F_aF_g$  (~0.7 in control arm) is not expected to be limited by intestinal efflux (EI-Kattan et al., 2016). Taken together, the current monkey study suggests potential contribution of intestinal efflux inhibition to the rifampicin-rosuvastatin DDIs.

The rich dataset of rifampicin dose-dependent effect on two statins simultaneously dosed i.v. and orally was leveraged to investigate the IVIVE of OATP1B-mediated DDIs. The *in vivo* IC<sub>50</sub> values, estimated

using intravenous clearance of rosuvastatin/pitavastatin and unbound  $C_{max}$  of rifampicin, were within 2.5-fold of the *in vitro* IC<sub>50</sub> (0.54  $\mu$ M) obtained following pre-incubation with rifampicin and using rosuvastatin as probe substrate in primary monkey hepatocytes. This *in vitro* IC<sub>50</sub> predicted the AUCR of both statins after i.v. and oral dosing with high accuracy (minimum 78% within 2-fold of the observed data), when employing the static mechanistic model (Eq. 7). In contrast, the *in vitro* IC<sub>50</sub> obtained under all other experimental conditions (Table 2) considerably underpredicted the fold change in the AUC of statins investigated. Consistent with the IVIVE noted here in monkey, Varma et al. demonstrated that similar inhibition potency (0.5  $\mu$ M), obtained using OATP1B1-transfected cells, correctly predicted rifampicin clinical DDIs when using a static mechanistic model (>85% within 2-fold of observed AUC ratios, n=22) (Varma et al., 2014). Overall, the current monkey study validates mechanistic translational framework to predict OATP1B-mediated DDIs in human.

Substrate-dependent inhibition was argued as a potential cause for concern in prospective prediction of OATP1B-mediated interactions (Noé et al., 2007; Izumi et al., 2013; Zamek-Gliszczynski et al., 2013). We report almost 6-fold difference in rifampicin IC50 measured with pre-incubation when using rosuvastatin (0.54 μM) vs. pitavastatin (3.0 μM) (Table 2). Interestingly, the change in i.v. clearance and oral exposure of both rosuvastatin and pitavastatin are recovered well only by the rifampicin in vitro IC50 measured against rosuvastatin (Figure 6 and 7). Based on these findings, it can be inferred that, (i) substratedependent inhibitory potency noted in vitro may not translate to differences in vivo, (ii) IC50 measured following pre-incubation with inhibitor is needed for quantitative prediction of OATP-mediated DDIs, and (iii) in vitro inhibition studies should consider more than one probe substrates and the most potent measurement should be employed for reliable prospective predictions. Of note, plasma coproporphyrins (I and III) have recently been described as selective OATP1B biomarkers in both cynomolgus monkey and human (Shen et al., 2016; Barnett et al., 2017). Therefore, the coordinated use of the cynomologus monkey (pre-human dosing), with measurements of changes in biomarker exposure in Phase 1, could be leveraged to quickly discharge OATP1B DDI risk. Given that current regulatory agency-driven OATP1B DDI risk analyses are conservative, and present a relatively high false positive rate (~30%) (Vaidyanathan et al., 2016), consideration of such approaches is warranted.

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In summary, our studies using the cynomolgus monkey as a model demonstrated a rational translation of *in vitro* inhibition potency in predicting OATP1B DDIs. Rifampicin *in vitro* inhibition potency measured using monkey hepatocytes was influenced by a pre-incubation step and/or the probe substrate employed. However, *in vivo* IC<sub>50</sub> values obtained from the change in rosuvastatin or pitavastatin i.v. clearance as a function of rifampicin unbound C<sub>max</sub> were not statistically different. Additionally, the most potent IC<sub>50</sub> obtained under pre-incubation conditions provided good quantitative predictions of fold-changes in plasma AUC for both statins following oral administration. Collectively, these findings suggest the need for employing multiple probe substrates and a pre-incubation step when conducting *in vitro* inhibition studies. Finally, this study emphasizes further the utility of cynomolgus monkey as a preclinical model in supporting the early assessment of OATP-mediated DDI risk.

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**Authorship Contribution** 

Participated in research design: Ufuk, Kosa, Gao, Bi, Rodrigues, Tremaine, Varma, Houston, and Galetin

Conducted experiments: Ufuk, Kosa, Gao, Bi, Modi, Gates

Contributed new reagents or analytical tools: Gao

Performed data analysis: Ufuk, Kosa, Varma, Houston, and Galetin

Wrote or contributed to the writing of the manuscript: Ufuk, Kosa, Gao, Modi, Gates, Rodrigues,

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# Footnote

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**Figure Legends** 

Figure 1. Uptake kinetic profiles of rosuvastatin (A) and pitavastatin (B) measured in cryopreserved

cynomolgus monkey hepatocytes plated for 4h. Symbols represent the observed total uptake. Solid,

dashed and dotted lines represent the predicted total, active and passive diffusion related uptake,

respectively.

Figure 2. Rifampicin (A-B) and cyclosporine (C-D) concentration-dependent inhibition of rosuvastatin (A-

C) and pitavastatin (B-D) as OATP substrates in cryopreserved cynomolgus monkey hepatocytes plated

for 4h. Pre-incubation with either buffer or inhibitor (rifampicin 0.01-100 μM, cyclosporine 0.01-6 μM) was

performed for 1h prior to co-incubation with the inhibitor and the probe substrates. Data represent mean ±

SD of at least triplicate measurements.

Figure 3. Effect of single ascending oral doses of rifampicin on the intravenous and oral

pharmacokinetics of rosuvastatin in cynomolgus monkey. Plasma concentration-time profiles of i.v. <sup>2</sup>H<sub>6</sub>-

rosuvastatin (A) and oral cold rosuvastatin (B), and the estimated pharmacokinetics (C) are depicted.

Pharmacokientics parameters were estimated with each monkey serving as its own control. Data

represent mean and s.d. (n=4). One-way ANOVA with Dunnett's multiple comparisons test was employed

to test significance with \*p<0.05 and \*\*p<0.01.

Figure 4. Effect of single ascending oral doses of rifampicin on the intravenous and oral

pharmacokinetics of pitavastatin in cynomolgus monkey. Plasma concentration-time profiles of i.v. <sup>2</sup>H<sub>4</sub>-

pitavastatin (A) and oral cold rosuvastatin (B), and the estimated pharmacokinetics (C) are depicted.

Pharmacokientics parameters were estimated with each monkey serving as its own control. Data

represent mean and s.d. (n=4). One-way ANOVA with Dunnett's multiple comparisons test was employed

to test significance with \*p<0.05 and \*\*p<0.01.

Figure 5. Rifampicin mean unbound plasma concentration-time profiles at different doses following oral

administration in cynomolgus monkey. Horizontal lines represent in vitro IC50 values estimated using

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rosuvastatin and pitavastatin as probe substrates following pre-incubation with buffer or rifampicin (Table

2).

Figure 6. IVIVE of hepatic uptake inhibition by rifampicin in cynomolgus monkey. Plots depict rifampicin

plasma concentration-dependent inhibition of intravenous clearance of rosuvastatin (A) and pitavastatin

(B). Data were fitted to Eq. 7 to estimate in vivo  $IC_{50}$  with each statin (red curves with shaded area – 95%

CI). Additionally, change in intravenous clearance was predicted using in vitro IC50 obtained in monkey

hepatocytes following pre-incubation with rifampicin with rosuvastatin as a probe (green curve). Vertical

lines represent in vivo and in vitro IC<sub>50</sub> values (Table 2).

Figure 7. Predicted versus observed interactions between rifampicin and rosuvastatin (A) or pitavastatin

(B), when administered with single ascending oral doses of rifampicin. Change in exposure (AUC ratio) of

intravenous (open points) and oral (closed points) statins was predicted based on eqs. 4-7 using the

measured unbound C<sub>max</sub> of rifampicin and in vitro IC<sub>50</sub> of 0.54 μM. Each monkey served as its own control

in calculating the AUCR. Solid and dotted lines represent the line of unity and 2-fold error, respectively.

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Table 1

Uptake kinetic parameters of rosuvastatin and pitavastatin estimated in plated cynomolgus monkey hepatocytes using a mechanistic two-compartment model

Drug	K <sub>m</sub>	$V_{max}$	$\mathbf{f}_{u,cell}$	$CL_{diff}$	CL <sub>active</sub>	
	μM	pmol/min/10 <sup>6</sup> cells		μL/min/10 <sup>6</sup> cells		
Rosuvastatin	3.29 ± 0.45	$323 \pm 44$	0.25 ± 0.08	4.27 ± 1.07	98.3	
Pitavastatin	1.99 ± 0.31	217 ± 34	0.025 ± 0.009	26.50 ± 2.74	109	

Table 2

In vitro  $IC_{50}$  of rifampicin on the OATP-mediated uptake of rosuvastatin and pitavastatin with or without pre-incubation in cryopreserved plated cynomolgus monkey hepatocytes; and the in vivo  $IC_{50}$  estimated based on change in intravenous clearance for these two statins

Substrate	Inhibitor	In vitro	In vivo IC <sub>50</sub> (μM)	
		Pre-incubation	Pre-incubation with	
		with buffer	inhibitor	
Rosuvastatin	Rifampicin	$1.14 \pm 0.34^{**}$	0.54 ± 0.13	$0.99 \pm 3.80$
	Cyclosporine	0.72 ± 0.04**	0.10 ± 0.04 <sup></sup>	-
Pitavastatin	Rifampicin	3.80 ± 1.82	2.98 ± 0.78	0.22 ± 0.14
	Cyclosporine	0.78 ± 0.14	0.21 ± 0.06	-
	Rifamycin SV	6.17 ± 0.96	4.30 ± 1.28	-

Data represent mean  $\pm$  SD of at least triplicate measurements. "Significant difference between incubation conditions by paired t-test ( $p \le 0.01$ )

Table 3

Summary of pharmacokinetic estimates of pitavastatin and rosuvastatin in the presence of single ascending doses of rifampicin in cynomolgus monkey

monkey

	Control	+Rifampicin 1mg/kg	+Rifampicin 3mg/kg	+Rifampicin 10mg/kg	+Rifampicin 30mg/kg
	Mean ± SD	Mean ± SD	Mean ± SD	Mean ± SD ू	Mean ± SD
Pitavastatin				rnals	
CL (mL/min/kg)	$18.0 \pm 4.6$	15.0 ± 2.5	$7.5 \pm 2.0$	4.9 ± 1.3	4.3 ± 1.6
Fold change		0.83	0.42**	$4.9 \pm 1.3$ org at ASPET Journals $0.27^{**}$ 0.52 ± 0.16 $0.29^{**}$ 3.8 ± 0.7	0.24**
Vd <sub>ss</sub> (L/kg)	1.79 ± 0.72	$1.97 \pm 0.50$	$0.47 \pm 0.20$	0.52 ± 0.16	$0.55 \pm 0.20$
Fold change		1.10	0.26**	0.29** Jour	0.30**
t <sub>1/2</sub> (h)	$5.9 \pm 0.9$	7.2 ± 1.7	2.9 ± 1.9	$3.8 \pm 0.7$ mals	$4.5 \pm 0.6$
Fold change		1.21	0.50**	0.63 on March 4070 ± 4357	0.76
Oral AUC <sub>last</sub> (ng.h/mL)	299 ± 146	515 ± 417	2690 ± 2710	4070 ± 4357 Erg	5808 ± 4579
Fold change		1.72	8.99**	13.60**	19.41**
Oral C <sub>max</sub> (ng/mL)	225 ± 200	520 ± 822	2730 ± 2040	1920 ± 2950 224	2780 ± 3166
Fold change		2.31	12.13**	8.53**	12.36**
Rosuvastatin					
CL (mL/min/kg)	$27.5 \pm 7.8$	$23.6 \pm 2.4$	$20.0 \pm 2.1$	15.6 ± 1.9	$13.4 \pm 3.7$
Fold change		0.86	0.73	0.57**	0.49**
Vd <sub>ss</sub> (L/kg)	$0.67 \pm 0.41$	$0.61 \pm 0.06$	$0.64 \pm 0.16$	$0.71 \pm 0.08$	$0.79 \pm 0.16$
Fold change		0.91	0.96	1.05	1.17
t <sub>1/2</sub> (h)	$2.5 \pm 0.7$	$2.5 \pm 0.8$	$2.1 \pm 0.3$	$3.1 \pm 0.1$	$4.8 \pm 2.0$
Fold change		1.01	0.86	1.23	1.93*
Oral AUC <sub>last</sub> (ng.h/mL)	49 ± 12	$63 \pm 28$	$232 \pm 43$	467 ± 267	744 ± 512

Fold chan	ge	1.28	4.68**	9.41** Do	15.01**
Oral C <sub>max</sub> (ng/mL)	15 ± 11	24 ± 29	122 ± 46	150 ± 100 wnlo	227 ± 169
Fold change		1.54	7.72**	9.49** aded	14.37**
				from	
Rifampicin				jpet	
Oral AUC <sub>last</sub> (ng.h/mL)		985 ± 396	11300 ± 3260	70100 ± 10800 g	$240000 \pm 73900$
Dose-normalized change		-	3.8	7.1 tjour	8.1
related to 1mg/kg				nals.	
Oral C <sub>max</sub> (ng/mL)		176 ± 99	2050 ± 245	7980 ± 2480	24200 ± 14100
Dose-normalized change		-	3.8	4.5 AS	4.6
related to 1 mg/kg				SPET (	

Data represent mean and standard deviation (N=4). One-way ANOVA with Dunnett's multiple comparisons test was employed to test significance with \*p<0.05 and \*\*p<0.01.

On March 20, 2024

Figure 1

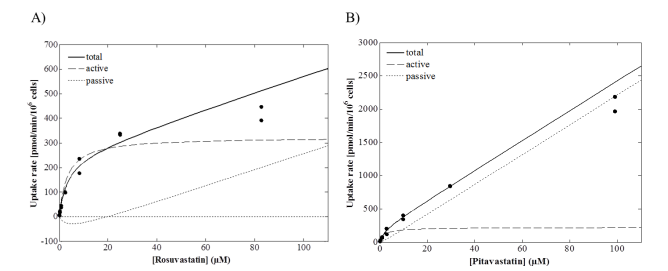


Figure 2

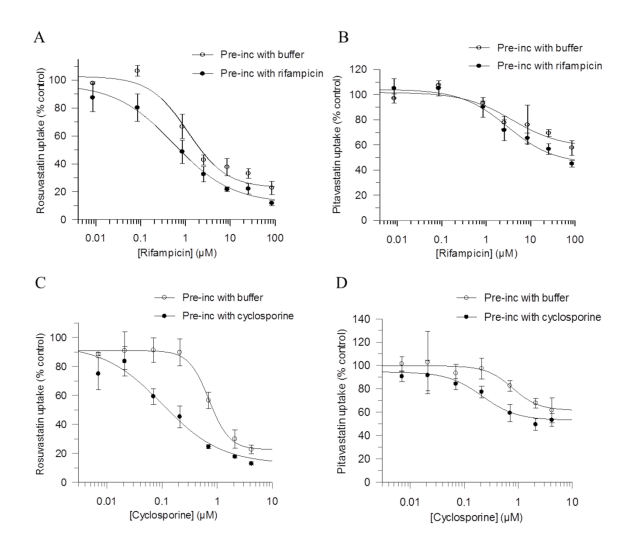


Figure 3

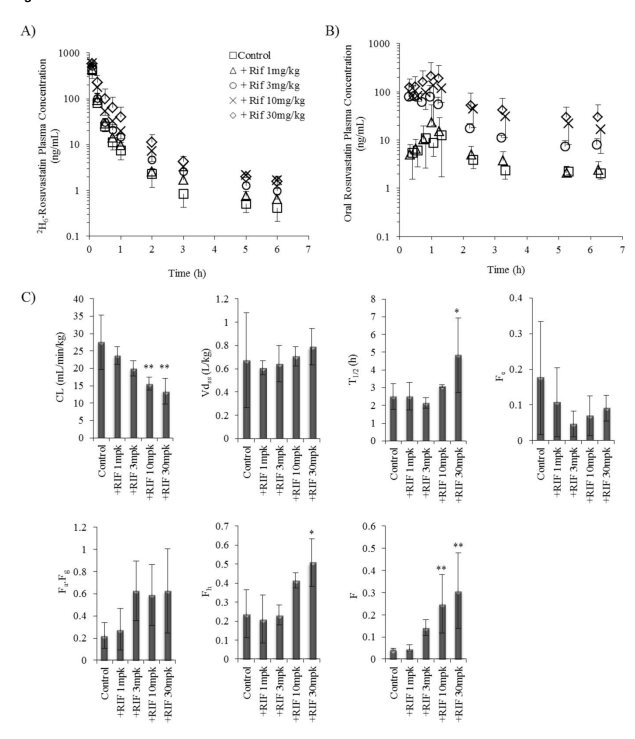


Figure 4

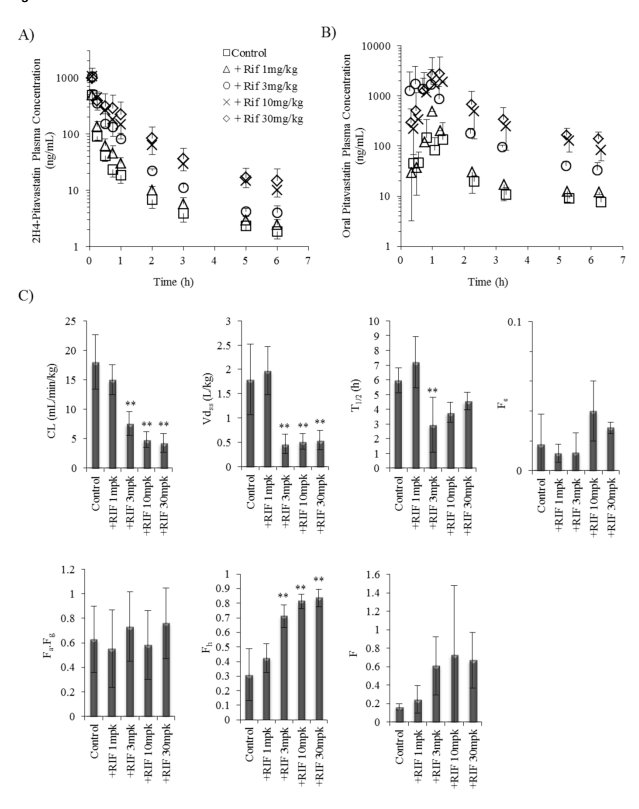


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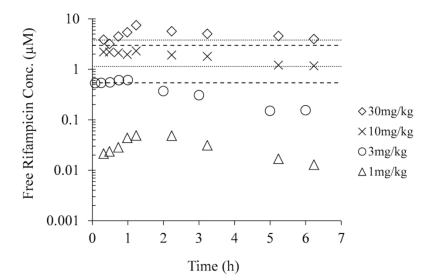


Figure 6

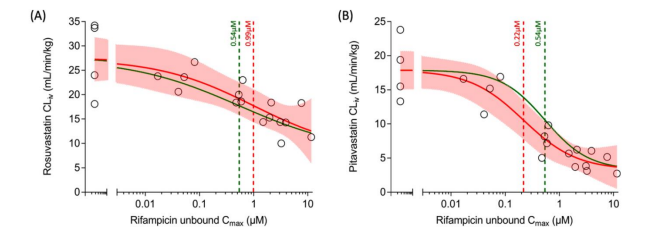


Figure 7

