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Mechanistic pharmacokinetic modeling for the prediction of transporter-mediated disposition in human from sandwich culture human hepatocyte data

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Abbreviations:

A, arterial; Abile, amount in bile; Acell, amount in cell; ADME, absorption, distribution, metabolism and excretion; Amedia, amount in media; BDDCS, biopharmaceutics drug disposition classification system; B:P, blood to plasma ratio; C, concentration; cell,u, cell x fu,cell; CEC, extracellular concentration; CIC, intracellular concentration; CL, clearance; Clint, intrinsic clearance; Clint,u,pass, unbound passive diffusion Clint; Clint,u,act, unbound active uptake Clint; Clint,u,bile, unbound biliary Clint; Clint,u,met, unbound metabolic Clint; Clint,u,renal, unbound renal Clint; Cm,tissue, concentration of albumin in liver relative to plasma; CYP450, cytochrome P450; fu,cell, fraction unbound in hepatocyte; fu,media, fraction unbound in the media; fu,p, fraction unbound in plasma; gu, gut; ha = hepatic artery; HLM, human liver microsomes; i.v., intravenous; Kp, tissue to plasma partition coefficient; NME, new molecular entity; KAMC, Clint,u,act/Vm; KMET, Clint,u,met/Vc; KBIL, Clint,u,bile/Vc; KPCM, Clint,u,pass/Vc; KPMC, Clint,u,pass/Vm; li, liver; media,u, media x fu,media; OATP, organic anion transporting polypeptides; PBPK, physiologically based PK; PK, pharmacokinetics; Q, blood flow; SCHH, sandwich culture human hepatocytes; SCLint,u,act, scaled unbound active uptake Clint; SCLint,u,bile, scaled unbound biliary Clint; SCLint,u,met scaled unbound metabolic Clint; SCLint,u,pass, scaled unbound distributional Clint; SD, standard deviation; sp, spleen; T, tissues; u, unbound; V, volume; v, venous; Vc, cell volume; VEC, volume of extracellular compartment; VIC, volume of intracellular compartment; Vinc, volume of the whole incubation; Vm, media volume; Vss, volume of distribution at steady state.
Abstract

With efforts to reduce CYP450-mediated clearance (CL) during the early stages of drug discovery, transporter-mediated CL mechanisms are becoming more prevalent. However, the prediction of plasma concentration-time profiles for such compounds using physiologically based pharmacokinetic (PBPK) modeling is far less established in comparison to compounds with passively mediated PK. In this study, we have assessed the predictability of human PK for seven organic anion transporting polypeptides (OATP) substrates (pravastatin, cerivastatin, bosentan, fluvastatin, rosuvastatin, valsartan and repaglinide) where clinical intravenous (i.v.) data were available. In vitro data generated from the sandwich culture human hepatocyte (SCHH) system were simultaneously fit to estimate parameters describing both uptake and biliary efflux. Use of scaled active uptake, passive distribution and biliary efflux parameters as inputs into a PBPK model resulted in the over-prediction of exposure for all seven drugs investigated, with the exception of pravastatin. Therefore, fitting of in vivo data for each individual drug in the dataset was performed to establish empirical scaling factors to accurately capture their plasma concentration-time profiles. Overall, active uptake and biliary efflux were under- and over-predicted, leading to average empirical scaling factors of 58 and 0.061, respectively; passive diffusion required no scaling factor. This study illustrates the mechanistic and model-driven application of in vitro uptake and efflux data for human PK prediction for OATP substrates. A particular advantage is the ability to capture the multiphasic plasma concentration-time profiles for such compounds using only pre-clinical data. A prediction strategy for novel OATP substrates is discussed.
Introduction

The prediction of human pharmacokinetics (PK) is pivotal to aid in the selection of new molecular entities (NMEs) with appropriate PK properties for clinical development. Physiologically based PK (PBPK) models have long provided a mechanistic framework for improved understanding and predictions of in vivo PK (Bischoff, 1975; Kawai et al., 1998). Successful predictions of human PK have been demonstrated using relevant in vitro and physicochemical data within such models (Jones et al., 2006; De Buck et al., 2007; Rostami-Hodjegan and Tucker, 2007; Jones et al., 2011). This approach has proven particularly successful for highly permeable compounds where metabolism is the predominant clearance (CL) mechanism, with negligible contribution of transporters to the overall disposition of these molecules (biopharmaceutics drug disposition classification system (BDDCS) classes 1 and 2, (Wu and Benet, 2005)). It should be emphasized that for these types of compounds, at equilibrium, the intracellular free drug concentration is expected to be equal to the free plasma concentration in the absence of CL from the tissue; therefore, the key PBPK model assumptions of flow mediated distribution and well stirred kinetics are valid. With efforts to reduce CYP450-mediated CL during drug discovery by reducing lipophilicity and increasing polarity, transporter-mediated PK is becoming more prevalent, particularly as the focus of drug discovery is moving away from the typical aminergic G-protein coupled receptors and enzyme targets to ion channels and peptidic receptors. For these more poorly permeable compounds (BDDCS classes 3 and 4, (Wu and Benet, 2005)) liver transporters may become an important determinant of disposition. The use of generic PBPK models therefore becomes limited, as hepatic uptake will lead to significant differences between the free concentrations in the hepatocyte and plasma.

A number of recent studies provide evidence that compounds with poorly predicted PK are often substrates for transporters (Soars et al., 2009; Watanabe et al., 2010). Improvement in PK prediction for such compounds requires accurate estimation of the
extent of active uptake and/or efflux in the hepatocyte. The movement of a compound across the hepatocyte cell membrane is modulated via passive diffusion and active transport, such as active uptake via organic anion transporting polypeptides (OATP). Once in the hepatocyte, substrates may be metabolized via CYP450-mediated metabolism or excreted into the bile via efflux transporters, e.g., multidrug resistance protein 2 or breast cancer resistance protein. This has resulted in the development of a number of in vitro assays with varying complexity that allow assessment of these processes either in isolation or combination (Giacomini et al., 2010). These include suspended hepatocytes (Kitamura et al., 2008; Paine et al., 2008), plated hepatocytes (Poirier et al., 2008; Yabe et al., 2011), sandwich cultured hepatocytes (Lee et al., 2010; Yan et al., 2011) and a range of transfected cell lines expressing individual transporters (Yamashiro et al., 2006; Kitamura et al., 2008).

The sandwich cultured human hepatocyte system (SCHH) involves culturing hepatocytes in a sandwich format between collagen and matrigel to allow polarisation of the cells (Liu et al., 1999; Bi et al., 2006; Lee et al., 2010). Through modulation of calcium ions, this in vitro system can be used to assess both uptake and biliary efflux (Liu et al., 1999; Bi et al., 2006). Simultaneous assessment of all the processes occurring in SCHH and mechanistic application of the data generated is currently lacking.

Mechanistic models have been used to describe in vitro uptake in suspended and plated hepatocytes (Paine et al., 2008; Poirier et al., 2008). Such in vitro data have been integrated either into semi-mechanistic or whole body PBPK models to simulate in vivo PK for OATP substrates in rat (Paine et al., 2008; Poirier et al., 2009a; Poirier et al., 2009b; Watanabe et al., 2009) and human (Poirier et al., 2009a; Watanabe et al., 2009). In most cases, successful predictions were only achieved when empirical scaling factors were incorporated.
The aim of this work was to examine the predictability of transporter mediated PK in humans using seven OATP substrates, selected based on the availability of clinical intravenous (i.v.) data. The SCHH assay was optimized to allow investigation of active/passive uptake and biliary efflux in the same hepatocyte donor and under the same experimental conditions. For each compound, in vitro SCHH data were simultaneously modeled to generate in vitro parameter estimates. In conjunction with other ADME and physicochemical properties, these parameters were then incorporated into a whole body PBPK model to assess the predictability of the clinical PK. A scaling approach is proposed and its potential application to novel compounds is discussed.
Materials and Methods

Materials. Compounds were purchased from Sequoia Research Products (Pangbourne UK), HT media, CP media, HI media and torpedo antibiotic mix were purchased from Celsis IVT (Baltimore, MD, USA), matrigel was purchased from BD BioSciences (Woburn, MA, USA), HBSS (Hanks balanced salt solution) was purchased from Invitrogen (Carlsbad, CA, USA) and all other chemicals were purchased from Sigma-Aldrich (St Louis, MO, USA).

Compound selection. Seven compounds were investigated, namely pravastatin, cerivastatin, bosentan, fluvastatin, rosuvastatin, valsartan and repaglinide. Compound selection was based on the availability of clinical i.v. data. Corresponding in vitro and physicochemical data were generated in house for these compounds using standard assays that have been described elsewhere in the literature (Allan et al., 2008).

Sandwich culture human hepatocyte (SCHH) experimental procedure. Cryopreserved human hepatocytes from donors HU4168, RTM and BD109 were purchased from CellzDirect (Pittsboro, NC, USA), Celsis IVT (Baltimore, MD, USA) and BD BioSciences (Woburn, MA, USA) respectively. These lots have been previously characterized in house and are known to have functional transport activity. The hepatocytes were cultured in a sandwich format as reported previously (Bi et al., 2006; Li et al., 2010).

Briefly, the cryopreserved hepatocytes were thawed in completed HT medium (thawing medium) and spun down at 50 x g for 3 minutes. The excess medium was removed and the hepatocyte pellet was re-suspended to 7.0 x 10⁵ cells mL in completed CP medium (plating medium). The hepatocyte suspension was then seeded onto 24-well BioCoat collagen I plates at 0.5 mL per well and cells allowed to attach overnight in a humidified incubator at 37 °C with 5% CO₂. On day 2, the excess hepatocytes were removed and the wells were washed with completed HI culture medium (incubation medium) at room temperature. Each well was then overlaid with BD Matrigel™ at a concentration of 0.25 mg/mL in ice-cold
completed HI medium. Media were replaced with completed HI medium daily until the day of the experiment.

On day 5, cells were washed twice and pre-incubated for 10 minutes at 37 °C in either: (a) HBSS buffer containing 0.1 mM rifamycin SV (inhibits a range of transporters (Vavricka et al., 2002)), (b) HBSS buffer or (c) Ca\(^{2+}\)/Mg\(^{2+}\) free HBSS containing 1 mM EGTA. Incubations were performed in 2 donors (except bosentan) and on a number of occasions. Substrates dissolved in the relevant condition buffer were added at 1 or 2 μM and incubated at 37 °C over 0.5-30 minutes; a minimum of 3 time points were taken in duplicate for each condition. Rosuvastatin was used as a positive control in all experiments. The cells were lysed with 0.5 mL methanol containing internal standard at room temperature for 20 minutes at 150 rpm. The samples were transferred to a 96 deep well plate and evaporated under 40 °C gaseous N2. The residue was reconstituted in 70% methanol and analysed using LC-MS/MS. Parallel wells of hepatocytes were lysed with RIPA buffer (TEKnova) or M-PER Mammalian Protein Extraction Reagent (Thermo Scientific (Waltham, MA)) for protein quantification by BCA Protein Assay Kit - Reducing Agent Compatible (Thermo Scientific (Waltham, MA)). Protein amounts were determined from the difference between the protein amount for each hepatocyte donor and the protein amount in blank wells, containing Matrigel™ alone.

**Bioanalysis procedure.** Analysis of 20 μL samples was carried out using HPLC (Hewlett Packard G1310 1100 Series Isocratic Pump) followed by MS/MS (MDS Sciex API 4000) using a 2 minute run time per sample. The mobile phase used to load the column (Dash HTS Hypersil Gold 20x2.1mm 5μm) was 2mM ammonium acetate in 90% methanol containing 0.027% formic acid (v/v); elution was performed at 0.7 min using a mobile phase of 2mM ammonium acetate in 10% methanol containing 0.027% formic acid (v/v). The flow rate was set at 1 mL/min. The mass:charge ratio (m/z) and collision energies (eV) for each compound were: pravastatin m/z 423→101 -40eV, cerivastatin m/z 460→356 50eV, bosentan m/z 552→202 40eV, fluvastatin m/z 412→266 1.11 25eV, rosuvastatin m/z 480→418 -25eV.
valsartan m/z 434→350 -25eV, repaglinide m/z 453→230 25eV. The internal standard used in all analyses was an in house compound (PF-05218881: m/z 688→366 negative ion mode, m/z 686→366 positive ion mode).

**In vitro data analysis.** The modeling approach used to analyse the SCHH data was analogous to the method described previously for suspended and plated hepatocytes by Paine and Poirier respectively (Paine et al., 2008; Poirier et al., 2008). To address biliary excretion, additional model terms have been proposed for the analysis of extended incubation times (Lee et al., 2010); however, these parameters cannot be estimated with the duration of experiment used here.

The model includes compartments representing the media, cell and bile environments of the experiment, with passive diffusion, active uptake and efflux processes incorporated in a mechanistic fashion, as illustrated in Figure 2. The passive diffusion component was parameterized as an unbound distribution CL (CLint,u,pass) within the model. Active uptake was parameterized in the form of an unbound uptake CL (CLint,u,act). Two further clearance mechanisms were incorporated, namely unbound biliary CL (CLint,u,bile) and unbound metabolic CL (CLint,u,met). Efflux transport by sinusoidal transporters was assumed negligible. There are studies to suggest bidirectional transport by OATPs, but these are generally based on oocyte data (Mahagita et al., 2007) and have not been considered here. It was assumed that only unbound drug is able to pass across the cell membrane, and that any binding to the cell membrane is instantaneous. The equations used in this modeling process are shown below.

\[
\frac{dA_{media}}{dt} = -KPMC \cdot A_{media,u} + KPCM \cdot A_{cell,u} - KAMC \cdot A_{media,u} + KBIL \cdot A_{cell,u}
\]

(1)

\[
\frac{dA_{cell}}{dt} = KPMC \cdot A_{media,u} - KPCM \cdot A_{cell,u} + KAMC \cdot A_{media,u} - KBIL \cdot A_{cell,u} - KMET \cdot A_{cell,u}
\]

(2)
\[
\frac{d\text{Abile}}{dt} = K_{BIL} \cdot \text{Acell}, u
\]  

(3)

where \( K_{PMC} = \text{Cl}_{int}, u, \text{pass}/\text{Vm} \); \( \text{Vm} = \text{media volume (~µL)} \); \( u = \text{unbound} \); \( K_{PCM} = \text{Cl}_{int}, u, \text{pass}/\text{Vc} \); \( \text{Vc} = \text{cell volume (~µL)} \); \( K_{AMC} = \text{Cl}_{int}, u, \text{act}/\text{Vm} \); \( K_{MET} = \text{Cl}_{int}, u, \text{met}/\text{Vc} \); \( K_{BIL} = \text{Cl}_{int}, u, \text{bile}/\text{Vc} \); \( \text{Amedia} = \text{amount in media (~pmoles)} \); \( \text{Acell} = \text{amount in cell (~pmoles)} \); \( \text{Abile} = \text{amount in bile (~pmoles)} \); \( \text{cell}, u = \text{cell} \times f_{u,\text{cell}} \); \( \text{media}, u = \text{media} \times f_{u,\text{media}} \).

\( \text{Cl}_{int} \) units were µL/min/Mcells and it was assumed based on in-house data that 1 Mcells = 1 mg protein. The volume of the whole incubation (\( V_{inc}\)) is the sum of the medium and cell volumes, \( V_m \) and \( V_c \), respectively. \( V_c \) was estimated assuming 1 Mcells is equivalent to 4 µL (Reinoso et al., 2001). The fraction unbound in the media (\( f_{u,\text{media}} \)) was assumed to equal 1, as no protein was present. The fraction unbound in the hepatocyte (\( f_{u,\text{cell}} \)) was calculated using a rearranged form of the equation reported by (Poulin and Theil, 2000), assuming the concentration of albumin in liver relative to plasma (\( C_{m,tissue} \)) is equal to 0.5. This parameter accounts for non-specific binding of the drug intracellularly within the hepatocyte and was fixed in further modeling of \textit{in vitro} data.

\[
f_{u,\text{cell}} = \frac{1}{1 + \left( \frac{1 - f_{u,\text{p}}}{f_{u,\text{p}}} \cdot C_{m,tissue} \right)}
\]  

(4)

where \( f_{u,\text{p}} \) is the fraction unbound in the plasma.

Nonzero initial conditions were set for the cell and media compartments to account for instantaneous nonspecific binding to cells and/or experimental apparatus. This amount was calculated from \( V_m \), \( V_c \) and the binding constant (\( KB \)) as described in the literature (Paine et al., 2008; Poirier et al., 2008) (Equations 5 and 6).

\[
\text{Amedia}(t = 0) = \text{Amedia} - \frac{\text{Amedia}}{\text{Vm}} \cdot KB \cdot \text{Vc}
\]  

(5)
Where significant metabolism was observed, Clint,u,met was set to the unbound Clint value determined in human liver microsomes (HLM), adjusted from μL/min/mg to μL/min/Mcells using the ratio of hepatocellularity to microsomal recovery (HLM Clint,u x microsomal recovery / hepatocellularity). The model fitting of Clint,u,pas, Clint,u,act, Clint,u,bile and KB was performed in NONMEM version VI level 1.2., NM-TRAN subroutines version III level 1.2 (Icon Development Solutions, Ellicott City, Maryland, USA, 2006) or in acsIX version 3.0.1.6 (Aegis Technologies, Huntsville, AL, USA). The HYBRID estimation method in NONMEM was employed, where first-order estimation (FO) was used to estimate all the parameters. Residual error was estimated using a proportional error model.

**In vivo simulations.** An i.v. PBPK model was used to model the in vivo situation. The PBPK model was composed of 15 compartments corresponding to the different tissues of the body, namely, adipose, bone, brain, gut, heart, kidney, liver, lung, muscle, skin, spleen, testes, rest of body which were connected by the circulating blood system (arterial and venous). Each compartment was defined by a tissue volume and a tissue blood flow rate; these physiological parameters for human have been described elsewhere (Jones et al., 2006). Each tissue was assumed to be perfusion rate limited, with the exception of liver. The liver and kidney were considered to be the only sites of elimination.

The mass balance differential equations (except for liver) used in the model have been described previously (Jones et al., 2006; Jones et al., 2011) and follow the principles shown below.

\[
\text{non-eliminating tissues: } VT \cdot \frac{dC_T}{dt} = QT \cdot C_a - QT \cdot C_vT
\]
where \( Q \) = blood flow (L/hr); \( C \) = concentration (mg/L); \( V \) = volume (L); \( T \) = tissues; \( a = \) arterial; \( v = \) venous; \( CVT = CT/Kp*B:P; \) \( Kp = \) tissue to plasma partition coefficient of the compound; \( B:P = \) blood to plasma ratio.

\[
\text{kidney: } VT \cdot \frac{dCT}{dt} = QT \cdot Ca - QT \cdot CVT - CLint,u,\text{renal} \cdot CVT,u
\]

where \( CLint,u,\text{renal} \) = the unbound renal intrinsic clearance of the compound (L/hr).

\( CLint,u,\text{renal} \) was calculated from the renal CL reported from the respective clinical study assuming well stirred conditions (Supplementary Material, Table S1).

The \( Kp \) values (for all tissues except for the liver) were estimated using tissue composition equations developed in the literature (Rodgers and Rowland, 2006). The tissue composition parameters reported by the authors were used. The main compound specific parameters required were \( \log D_{2,0} \), \( pKa \), \( B:P \) ratio and \( f_u,p \). These predictive equations account for four main processes: (1) partitioning of unionized drug into neutral lipids and neutral phospholipids, (2) dissolution of ionized and unionized drug in tissue water (3) electrostatic interactions between ionized drug and acidic phospholipids for strong ionized bases and (4) interactions with extracellular protein for neutrals, weak bases and acids. These equations assume only passive distribution and do not account for any active transport processes.

Tissue composition data and anatomical information were available for each tissue in the model. It was desirable to include the full PBPK model to get initial estimates of exposure related to safety concerns and to provide a framework for elaboration of transporters in other tissues.

The liver was modeled as a permeability-limited tissue, incorporating scaled active uptake (\( SCLint,u,\text{act} \)) and scaled passive diffusion clearances (\( SCLint,u,\text{pass} \)) of unbound drug at the sinusoidal membrane, scaled biliary clearance (\( SCLint,u,\text{bile} \)) of unbound drug at the canalicular membrane and scaled metabolic clearance (\( SCLint,u,\text{met} \)) of unbound drug.
(where appropriate). These scaled parameters (L/h) were calculated from the *in vitro* parameters \( \text{CLint, u, act, CLint, u, pass, CLint, u, bile} \) and \( \text{CLint, u, met} \) obtained in SCHH, accounting for the hepatocellularity and liver weight as described previously (Houston, 1994). The liver compartment was sub-divided into five units of extracellular and intracellular compartments, connected by blood flow in tandem (Watanabe et al., 2009), as shown in Figure 2. Watanabe and coworkers reported that for pravastatin, five sequential compartments most closely approximated the partial differential equation dispersion model, so this number of compartments was retained. Initial modeling (results not shown) used one liver tissue and liver blood. The corresponding differential equations used are shown below:

### Extracellular liver 1:

\[
\frac{\text{VEC}}{5} \frac{d\text{CEC}_1}{dt} = Q_{\text{ha}} \cdot \text{Ca} + Q_{\text{gu}} \cdot \text{Cvgu} + Q_{\text{sp}} \cdot \text{Cvsp} - Q_{\text{li}} \cdot \text{CEC}_1 - \frac{\text{SCLint, u, pass}}{5} \cdot (\text{CEC}_1 - \text{CIC}_1) - \frac{\text{SCLint, u, act}}{5} \cdot \text{CEC}_1
\]

(9)

where \( \text{CEC} \) = extracellular concentration (mg/L); \( \text{CIC} \) = intracellular concentration (mg/L); \( \text{VEC} \) = volume of extracellular compartment (L); \( \text{VIC} \) = volume of intracellular compartment (L); \( \text{ha} \) = hepatic artery; \( \text{gu} \) = gut; \( \text{sp} \) = spleen; \( \text{li} \) = liver.

### Extracellular liver 2-5:

\[
\frac{\text{VEC}}{5} \frac{d\text{CEC}_i}{dt} = Q_{\text{li}} \cdot (\text{CEC}_{i-1} - \text{CEC}_i) - \frac{\text{SCLint, u, pass}}{5} \cdot (\text{CEC}_i - \text{CIC}_i) - \frac{\text{SCLint, u, act}}{5} \cdot \text{CEC}_i
\]

(10)

### Intracellular liver 1-5:

\[
\frac{\text{VIC}}{5} \frac{d\text{CIC}_i}{dt} = \frac{\text{SCLint, u, pass}}{5} \cdot (\text{CEC}_i - \text{CIC}_i) + \frac{\text{SCLint, u, act}}{5} \cdot \text{CEC}_i - \frac{\text{SCLint, u, bile}}{5} \cdot \text{CIC}_i
\]

(11)
The model simulations were performed in Berkeley Madonna, version 8.3.9 (University of California, USA, 1996-2006).

Maximal contribution of the active process to the total uptake was estimated and expressed as the ratio of CLint,u,act and total uptake CLint (CLint,u,act + CLint,u,pass).

**In vivo fitting.** In addition to the simulations, a fitting procedure was performed. Using the observed clinical i.v. data (extracted via Digitizelt version 1.5.7), the SCLint,u,act, SCLint,u,pass and SCLint,u,bile were estimated by the PBPK model assuming that all other parameters within the model were correct and using the scaled parameter values as the initial estimates. For cerivastatin, bosentan and fluvastatin where metabolic and biliary clearance data were available, the sum of SCLint,u,bile and SCLint,u,met were fitted as SCLint,u,bile could not be uniquely identified. For repaglinide, SCLint,u,bile was assumed negligible and the predicted SCLint,u,met was fixed. The fitting procedure was performed using a proportional error model implemented within Berkeley Madonna by log transformation of the data. For the individual compounds, empirical scaling factors were calculated for each of these parameters by dividing the measured (scaled to intact liver) value by the fitted value.

The geometric mean of the empirical scaling factors across the drugs in the dataset was calculated. The i.v. PK for each compound was further simulated using the *in vitro* data together with the average empirical scaling factors within the PBPK model. The simulations were compared graphically with the observed clinical data. The predicted PK profiles with and without average empirical scaling factors were modeled in WinNonlin version 5.2 (Mountain View, CA) using non-compartmental analysis to determine volume of distribution at steady state (Vss) and CL parameters.

**Local sensitivity analyses:** Local sensitivity analyses were conducted in acsIX version 3.0.1.6 (Aegis Technologies, Huntsville, AL, USA) for each of the seven compounds to obtain
numerical estimates of the partial derivative of the model with respect to each parameter.

Each parameter was raised or lowered by 1% with respect to its value for that compound and the value of the plasma concentration was obtained at three selected times throughout the time course during simulations of the conditions used for fitting *in vivo* parameters. Sensitivity coefficients were normalized to both the parameter value and the model output value, so when the output changes by 1% for a 1% change in the input parameters, the sensitivity coefficient is 1 or -1 depending upon the direction of change. Only parameters with normalized sensitivity coefficients greater than 0.3 or less than -0.3 are reported.
Results

Physicochemical properties. A summary of the available in vitro data (excluding hepatic uptake data) and physicochemical properties for pravastatin, cerivastatin, bosentan, fluvastatin, rosuvastatin, valsartan and repaglinide is shown in Table 1. All compounds were acidic. The LogD_{7.4} measurements ranged from very hydrophilic at -0.88 for valsartan to lipophilic at 2.1 for repaglinide. B:P ratios were comparable across compounds, ranging from 0.48-0.76. For cerivastatin, bosentan, fluvastatin, valsartan and repaglinide, fu,p values were very low (< 1%), whereas fu,p values for rosuvastatin and pravastatin were higher at 9.4 and 43%, respectively. HLM Clint values were determined via substrate depletion experiments and corrected for nonspecific binding. Pravastatin, rosuvastatin and valsartan had no measurable metabolism in HLM, whereas Clint,u,met ranged from 22-128 μL/min/mg for the remaining compounds (Table 1).

Clinical data. The corresponding human PK parameters for each compound are reported in Table S1 (Supplementary Material). These data were obtained from i.v. PK studies reported in the literature (see References in Supplementary Material). CL ranged from 0.49-14mL/min/kg for valsartan and pravastatin, respectively, whereas Vss covered a 10-fold range with fluvastatin and rosuvastatin at the low and high end, respectively. Pravastatin, rosuvastatin and valsartan exhibited 29-47% contribution of renal excretion to their total CL.

In vitro data SCHH analysis. The in vitro SCHH data obtained for all seven drugs investigated are shown in Figure S1 in the Supplementary material. The SCHH data were simultaneously modeled as described in the methods section and Figure 1. The derived parameter estimates of Clint,u,act, Clint,u,pass, Clint,u,bile are shown in Table 2. This assay was performed in 2 donors (except bosentan) and on a number of occasions. For the purposes of modeling, fu,cell, which describes the free fraction in the cell and the fraction nonspecifically bound (i.e., 1 - fu,cell) was fixed to the value predicted using Equation 4. Within the fitting process,
sinusoidal efflux was assumed to be negligible. The parameter values were estimated to an acceptable precision level and diagnostic plots and visual inspection of the observed versus fitted data indicated a good model fit (plots not shown). A 60-fold range in total uptake CLint values was observed, with pravastatin showing the lowest total uptake CLint (below 2\(\mu\)L/min/Mcells) and repaglinide the highest (119\(\mu\)L/min/Mcells). Despite this, the maximal contribution of active processes to uptake CLint varied and was not necessarily correlated with the total uptake CLint. Pravastatin and valsartan both exhibited low uptake into the hepatocyte; however, the maximal active contribution was proportionally high at 95 and 78\%, respectively. In contrast, the total uptake of fluvastatin and repaglinide were much higher, with maximal active contributions of 69 and 25\% respectively; for these compounds the contribution of passive diffusion was substantial and also subject to variability in the case of repaglinide. Bosentan, cerivastatin, and rosuvastatin exhibited intermediate uptake with a range of maximal active uptake contribution of 65, 28 and 85\% respectively. CLint,u,bile of the parent compound varied for the different compounds from 0 in the case of repaglinide to 96\(\mu\)L/min/Mcells for valsartan. For all these compounds, uptake CLint (passive and active) determined in suspended hepatocytes was within 2-3 fold of the SCHH data (data not shown).

In vivo simulations and fitting. The in vitro parameters derived from the simultaneous modeling of the SCHH data and the HLM CLint,u,met were scaled to the in vivo situation to account for hepatocellularity, microsomal recovery and liver weight. Scaled SCLint,u,act, SCLint,u,pass, SCLint,u,bile and SCLint,u,met parameters are shown in Table 3 for all the drugs investigated. The Kp values, fu,p, fu,cell and B:P ratio as well as SCLint,u,active, SCLint,u,passive, SCLint,u,bile and SCLint,u,met were input into the whole body PBPK model described in the methods section and Figure 2. The clinical dose was simulated for each compound using the whole body PBPK model. The simulated versus observed profiles for each compound are shown in Figure 3. The simulation overpredicted the exposure when...
compared to the observed data, except for pravastatin and rosvastatin, suggesting underestimation of the initial distribution phase.

To rationalize the misestimation of the observed human i.v. plasma concentration time profile, the SClint,u,act, SClint,u,pass and SClint,u,bile were estimated using the model and the observed clinical i.v. data, assuming that all other parameters within the model were correct. These fitted parameters are shown in Table 3 and the profiles originating from these fitted parameters are shown in Figure 3, in parallel to the initial simulations. As can be seen in Figure 3, the fit for each compound accurately describes the observed plasma concentration-time data and corresponds to the shape of the observed profile. Goodness of fit plots are shown in Figure S2 in the Supplementary material. The fitted parameters were compared to the predicted parameters for each drug to generate an empirical scaling factor. The in vivo fitted SClint,u,act was significantly (p<0.0001) higher than the in vitro scaled value for the entire dataset, with empirical scaling factors ranging from 12-161 for rosvastatin and fluvastatin, respectively, resulting in a geometric mean empirical scaling factor of 58. Consistent with expectations for a model approximating the liver acinar gradient (5-sequential subcompartments) versus a single tissue and blood compartment, initial modeling with the single unit model required higher empirical scaling factors for fluvastatin and other compounds with high active uptake. In contrast, values were more comparable between the two models for rosvastatin and compounds with slower uptake rates, with a geometric mean empirical scaling factor of 95 (results not shown). In the majority of cases the in vivo fitted and in vitro scaled SClint,u,pass compared well with each other giving a geometric mean scaling factor of 1. The only exception was valsartan where scaling of passive permeability was required to fit the terminal phase of the profile. For each of the seven compounds, the in vivo fitted SClint,u,bile was on average 16-fold lower than the in vitro scaled value, giving empirical scaling factors below 1 (Table 3). For cerivastatin, bosentan and fluvastatin, the empirical scaling factors for SClint,u,bile could not be
estimated as these compounds also undergo measureable P450 metabolism and the model could not uniquely identify these two parameters. For this reason the two parameters were summed and fitted together. For repaglinide, there was no measurable \( \text{CL}_{\text{int,u,bile}} \). The empirical scaling factor for \( \text{CL}_{\text{int,u,bile}} \) (0.061) was therefore calculated using only 3 compounds (pravastatin, rosuvastatin and valsartan).

These average empirical scaling factors for active uptake and biliary efflux were subsequently re-applied to the seven compounds to re-simulate the clinical i.v. data. Figure 4 shows simulated plasma concentration time profiles based on the SCHH estimates for \( \text{CL}_{\text{int,u,act}}, \text{CL}_{\text{int,u,pass}} \) and \( \text{CL}_{\text{int,u,bile}} \) scaled by the average empirical scaling factors of 58, 1 and 0.061 for the corresponding processes, respectively; all other parameter inputs were as used for simulations shown in Figure 3. Use of average empirical scaling factors, compared to simulations performed without them, resulted in better agreement between the simulated profile and observed data for the majority of the compounds. The biphasic profile for bosentan was not accurately captured using the generic empirical scaling factor; the fitting procedure reduced combined metabolism and biliary efflux to a value lower than \( \text{CL}_{\text{int,u,met}} \) (55 versus 87 L/hr) (Table 3), whereas for the re-simulation the \textit{in vitro} metabolism scaled value was used.

The predicted Vss and CL parameters using different modeling scenarios are shown in Table 4. For each compound both the CL and Vss were under predicted when the \textit{in vitro} data alone were used in the PBPK model with an absolute average fold error of 7.1 and 3.0 respectively. However, when these data were corrected for the average empirical scaling factors the predicted CL and Vss parameters corresponded more accurately with the observed data with absolute average fold errors of 1.3 and 1.7, respectively.

**Local sensitivity analyses:** For each of the seven compounds, all parameters were investigated to assess their sensitivity to the model simulation of the plasma concentration
profile. This analysis included all the physiological parameters in the model as well as those specific for each compound. Very few of the physiological parameters had normalized sensitivity coefficients greater than 0.3 or less than -0.3, indicating very limited impact when these parameters were varied individually (data not shown). Physiological parameters to which multiple compounds were sensitive included body weight, cardiac output, volume of liver tissue and liver blood flow. The fu,p, fu,cell, B:P ratio, SCLint,u,act and SCLint,u,pass, were sensitive parameters, along with SCLint,u,bile, SCLint,u,met and CLint,u,renal as appropriate to each compound. The sensitive parameters showed complex changes as would be expected for the different exposure regimens and varying importance of processes throughout the duration of the observed plasma time course. Figure 5, illustrates that the early plasma time course is increasingly sensitive to liver uptake (with SCLint,u,act and SCLint,u,pass having opposite impacts), while the later time course becomes increasingly sensitive to the clearance from the liver (SCLint,u,bile and SCLint,u,met) and CLint,u,renal.
Discussion

Prediction of human PK remains an important feature of drug discovery to help select compounds with appropriate characteristics for clinical development. Physiologically based methods for human PK prediction are reasonably well established for small lipophilic compounds cleared by liver metabolism (Jones et al., 2006; De Buck et al., 2007; Rostami-Hodjegan and Tucker, 2007; Jones et al., 2011) and have recently been applied for better understanding of intestinal first-pass (Gertz et al., 2011). However, the optimization of compound properties to improve metabolic stability has led to a reduction in lipophilicity and permeability and hence a shift in QL routes from liver metabolism to transporter mediated uptake and efflux. The prediction of such processes in humans continues to be a challenge (Liu and Pang, 2006; Poirier et al., 2009a; Watanabe et al., 2009).

Using a SCHH in vitro system together with available clinical plasma concentration-time data for seven compounds, we have established a prediction approach for active liver uptake and efflux. These SCHH in vitro data were dynamically modeled as outlined in Figure 1. In contrast to previous modeling efforts (Paine et al., 2008; Poirier et al., 2008), the current model allowed estimation of biliary efflux, active and passive uptake from the same in vitro experiment through modulation of calcium ions (Liu et al., 1999; Bi et al., 2006; Lee et al., 2010; Yan et al., 2011). The in vitro mechanistic model was parameterized in such a way to separate out the intracellular binding and the active processes; hence, the fu,cell term was fixed to the predicted value. An accurate estimation of this parameter is particularly important to estimate the free concentration in the liver and influences the rate of metabolism/biliary excretion. Here fu,cell was predicted using Equation 4, however improved prediction approaches may be needed (Yabe et al., 2011). These in vitro parameters were then scaled to in vivo and were subsequently integrated into a whole body PBPK model, together with other ADME properties, to simulate the human plasma concentrations (Figure 2). Prediction accuracy of these simulations was assessed by
comparison with observed plasma concentration-time profiles reported in the literature.

The simulations in Figure 3 show that the plasma concentration time profiles were significantly overestimated, with the initial phase of the profile being significantly mis-predicted, perhaps indicating an underprediction of the initial distribution into the liver, as observed by others (Poirier et al., 2009a; Watanabe et al., 2009). Using a fitting procedure, the PBPK model together with the observed plasma concentration-time data were used to estimate the in vivo values for the SCLint,u,act, SCLint,u,pass and SCLint,u,bile that would better describe the observed data. Table 3 shows the fitted values together with the empirical scaling factors describing the relationship between the fitted and measured parameters. In general SCLint,u,act and SCLint,u,bile were under- (58-fold) and over-predicted (16-fold), respectively; no empirical scaling factor was required for passive diffusion. The values for empirical scaling factors reported here are specific to our implementation of the SCHH; other laboratories would need to re-estimate their own system specific empirical scaling factors.

Previous studies have also reported the need for empirical scaling factors for active uptake for pravastatin and valsartan to recover the human plasma concentrations (Poirier et al., 2009a; Watanabe et al., 2009). The basis for this empirical scaling factor is unclear. An evaluation of OATP expression in SCHH, showed that, while OATP1B3/ OATP2B1 expression were reduced to ~50% of that in suspension, OATP1B1 expression was increased to ~150% (Table S2, Supplementary Materials). These results are consistent with recent studies that show no effect of the culture time on uptake transporter activity in SCHH for rosvustatin (Kotani et al., 2011), which would not support the hypothesis that the expression of uptake transporters is downregulated in this system. However, the difference between OATP expression in culture and in vivo is unknown. The model fitting was performed using plasma concentration-time data alone; the lack of liver concentration data to assist in model fitting may mean that the fitted values for SCLint,u,act, SCLint,u,pass and SCLint,u,bile have not
been estimated accurately, though this would be expected to have more impact on the biliary transport value than the active uptake. However, the issue of parameter identifiability has been investigated and the combination of parameters obtained by fitting appeared unique in their ability to accurately describe the plasma concentration-time profiles of these compounds (see contour plots in Supplemental material, Figure S3) given the model structure used and the values of the other parameters. In addition, the fitting routine assumes that all other parameters within the PBPK model were correct and that all other tissues (excluding the liver) are perfusion rate limited. However, expression of OATP transporters has been reported in other tissues, e.g., kidney (Hilgendorf et al., 2007) and this could partially explain the large empirical scaling factors required when liver alone is assumed to be the main transport organ. Initial data on transporter abundance are becoming available (Schaefer et al., 2011); however, until detailed transporter expression data are reported for the liver and other tissues, this will remain a caveat of this analysis. In terms of the passive uptake into the liver, the \( \text{SCLint, u, pass} \) scaled accurately from \textit{in vitro} to \textit{in vivo}, probably due to the lack of dependence on an enzymatic or active processes that could be up- or down-regulated on culturing. This has been observed also by others in the literature in human and rat (Paine et al., 2008; Poirier et al., 2009a; Poirier et al., 2009b; Watanabe et al., 2009).

One limitation of the modeling reported here is that it does not address enterohepatic recirculation for the drugs with substantial biliary excretion, as explicitly demonstrated with rosvastatin in rats (Nezasa et al., 2002). The absence of recycling would mean that the estimated biliary efflux would represent only the net CL (i.e., excreted minus reabsorbed) resulting in the observed overprediction. A more complex model to describe the recycling of drug from bile back into the intestine is required to fully understand this empirical scaling factor and would facilitate incorporation of additional published human data for intraduodenal dosing and biliary excretion of rosvastatin (Bergman et al., 2006).
An evaluation of our SCHH assay indicated that these biliary efflux proteins are upregulated by ~3-fold during the 5 day culture period, which would also in part explain the overprediction observed (unpublished data).

The sensitivity analysis showed complex changes as would be expected for the different exposure regimens and varying importance of processes throughout the duration of the observed plasma time course, illustrating the value of fitting multiphasic i.v. plasma concentration-time data for estimation of the empirical scaling factors (to estimate misprediction) rather than relying solely on reported PK parameter values.

Although each compound showed the same trend in terms of empirical scaling factors, there was a high degree of variability in the derived values, suggesting that the de novo application of these parameters to novel compounds may be uncertain. This variability could result from several factors e.g. multiple/different transporters between compounds and genetic polymorphisms for some transporters. The purpose of this analysis was to explore the utility of using SCHH data to simulate human transporter-mediated PK. Several issues have been identified that require further investigation. Availability of in vitro transporter data in a larger number of donors in conjunction with protein expression data is required to further improve transporter IVIVE. However, Figure 4 shows that the application of the average empirical scaling factors to these compounds gives a reasonable description of the plasma concentration-time profile. In addition, the application of these average empirical scaling factors results in a good prediction accuracy for CL and Vss parameters with absolute average fold errors of 1.3 and 1.7 respectively (Table 4). However, the true test for this prediction approach would be its application to novel compounds. We have recently applied the average empirical scaling factors derived in this study for the seven literature compounds for simulation of the human PK of four novel OATP substrates entering a first in human study at Pfizer. The PBPK prediction methodology resulted in better predictions accuracy when compared to other allometric scaling and more traditional in vitro scaling
approaches (data not shown). In our study, we used the observed renal CL values determined from the clinical data. However, when conducting simulations for novel compounds, this parameter would need to be predicted from preclinical data.

In summary, this study has provided a systematic analysis of seven transporter substrates in SCHH. A mechanistic prediction methodology has been proposed for scaling of human PK for the compounds investigated using data generated in SCHH. Although this approach relies on the use of empirical scaling factors for active uptake and biliary efflux, it allows simultaneous assessment of multiple processes occurring in the hepatocytes in a mechanistic manner and improves our understanding of the relevance of these processes for hepatic disposition of drugs. The necessity for these empirical scaling factors needs to be further understood to increase confidence in the applicability of this methodology to novel compounds.
Acknowledgements

The assistance of scientists at the Hamner Institutes for Health Sciences (Research Triangle Park, NC), including Jerry Campbell and Alina Efremenko, in converting the code to acsX software and carrying out initial local sensitivity analyses is greatly appreciated.
Authorship Contribution

*Participated in research design:* Jones, Barton, Lai, El-Kattan, Fenner.

*Conducted experiments:* Lai, Bi, Kimoto, Kempshall, Tate, Fenner.

*Contributed new reagents or analytical tools:* Not Applicable.

*Performed data analysis:* Jones, Barton, Tate.

*Wrote or contributed to writing the manuscript:* Jones, Barton, Lai, Bi, Kimoto, Kempshall, Tate, El-Kattan, Houston, Galetin, Fenner.
References


Footnotes

Financial Disclosure: H.M.J., S.K., K.S.F., H.B., Y.L., Y.B., E.K. and A.E-K. were all employees of Pfizer during this research.
**Figure Legends**

Figure 1 – Schematic of the *in vitro* model (A) Conditions: 0.1mM Rifamycin SV; (B) Conditions: HBSS; (C) Conditions Ca\(^{2+}\)/Mg\(^{2+}\) free, 1mM EGTA

Figure 2 – Schematic of the *in vivo* PBPK model

Figure 3 – Simulated, fitted and observed human i.v. plasma concentration-time profiles for (A) pravastatin; (B) cerivastatin; (C) bosentan; (D) fluvastatin; (E) rosuvastatin; (F) valsartan and (G) repaglinide

*open squares represent observed data; solid line represents predicted data using the PBPK model; dashed line represents fitted data*

Figure 4 – Simulated (using the average empirical scaling factor for individual processes) and observed human i.v. plasma concentration-time profiles for (A) pravastatin; (B) cerivastatin; (C) bosentan; (D) fluvastatin; (E) rosuvastatin; (F) valsartan and (G) repaglinide

*open squares represent observed data; solid line represents predicted data using the PBPK model and the average empirical scaling factors listed in Table 3*

Figure 5 – Time dependent sensitivity analysis of the importance of CL\(_{int,u,act}\), CL\(_{int,u,pass}\), CL\(_{int,u,met}\) and CL\(_{int,u,bile}\) parameters on the plasma kinetics of (A) pravastatin; (B) cerivastatin; (C) bosentan; (D) fluvastatin; (E) rosuvastatin; (F) valsartan and (G) repaglinide
solid line represents $\text{Cl}_\text{int},u,\text{act}$; dashed line represents $\text{Cl}_\text{int},u,\text{pass}$; dotted line represents $\text{Cl}_\text{int},u,\text{met}$; dash-dot line represents $\text{Cl}_\text{int},u,\text{bile}$ and dash-dot-dot line represents renal $\text{Cl}$.
Table 1 – Physicochemical, in vitro and clinical pharmacokinetic properties for the compounds studied

<table>
<thead>
<tr>
<th>Compound</th>
<th>LogD7.4</th>
<th>Charge</th>
<th>pKa</th>
<th>B:P</th>
<th>fu,p</th>
<th>HLM CLint,um</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pravastatin</td>
<td>-0.84</td>
<td>A</td>
<td>4.6</td>
<td>0.55</td>
<td>0.43</td>
<td>0</td>
</tr>
<tr>
<td>Cerivastatin</td>
<td>1.8</td>
<td>Z</td>
<td>5.3(B); 5.0(A)</td>
<td>0.76</td>
<td>0.0048</td>
<td>29</td>
</tr>
<tr>
<td>Bosentan</td>
<td>1.3</td>
<td>A</td>
<td>5.2</td>
<td>0.48</td>
<td>0.0053</td>
<td>22</td>
</tr>
<tr>
<td>Fluvastatin</td>
<td>1.6</td>
<td>A</td>
<td>4.6</td>
<td>0.57</td>
<td>0.0041</td>
<td>76</td>
</tr>
<tr>
<td>Rosuvastatin</td>
<td>-0.33</td>
<td>A</td>
<td>4.2</td>
<td>0.56</td>
<td>0.094</td>
<td>0</td>
</tr>
<tr>
<td>Valsartan</td>
<td>-0.88</td>
<td>A</td>
<td>3.8; 4.6</td>
<td>0.55</td>
<td>0.0010</td>
<td>0</td>
</tr>
<tr>
<td>Repaglinide</td>
<td>2.1</td>
<td>Z</td>
<td>6.1(B); 4.4(A)</td>
<td>0.48</td>
<td>0.0074</td>
<td>128</td>
</tr>
</tbody>
</table>

1 A=acidic, B=basic, Z=zwitterionic; 2 (Lave et al., 1996), corrected for microsomal binding (fu = 0.87); 3 (Gertz et al., 2010)
<table>
<thead>
<tr>
<th>Compound</th>
<th>CLint,u, active (μl/min/Mcells)</th>
<th>CLint,u, passive (μl/min/Mcells)</th>
<th>CLint,u, bile (μl/min/Mcells)</th>
<th>Predicted fu,cell</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pravastatin</td>
<td>1.8</td>
<td>0.1</td>
<td>1.2</td>
<td>0.60</td>
</tr>
<tr>
<td>Cerivastin</td>
<td>9.6 (2.7)</td>
<td>25 (3.5)</td>
<td>6.2 (1.8)</td>
<td>0.0096</td>
</tr>
<tr>
<td>Bosentan</td>
<td>9.1</td>
<td>4.8</td>
<td>17</td>
<td>0.011</td>
</tr>
<tr>
<td>Fluvastatin</td>
<td>45 (21)</td>
<td>20 (9.6)</td>
<td>17</td>
<td>0.0082</td>
</tr>
<tr>
<td>Rosuvastatin</td>
<td>9.3 (2.6)</td>
<td>1.7 (0.73)</td>
<td>1.5 (0.088)</td>
<td>0.17</td>
</tr>
<tr>
<td>Valsartan</td>
<td>2.1 (0.48)</td>
<td>0.60 (0.18)</td>
<td>96</td>
<td>0.0020</td>
</tr>
<tr>
<td>Repaglinide</td>
<td>30 (16)</td>
<td>89 (75)</td>
<td>0</td>
<td>0.015</td>
</tr>
</tbody>
</table>

1Lots: Pravastatin: BD109 (n=1), HU4168 (n=1); Cerivastatin: BD109 (n=2), HU4168 (n=2); Bosentan: HU4168 (n=1); Fluvastatin: BD109 (n=2), HU4168 (n=1); Rosuvastatin: BD109 (n=6), HU4166 (n=2); Valsartan: BD109 (n=2), HU4168 (n=1); Repaglinide: HU4168 (n=3), RTM (n=2)
Table 3 – *In vitro* scaled and fitted sandwich culture human hepatocyte estimates

<table>
<thead>
<tr>
<th>Compound</th>
<th>SCLint,u,act L/hr</th>
<th>SCLint,u,pass L/hr</th>
<th>SCLint,u,bile L/hr</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pravastatin</td>
<td>19</td>
<td>406</td>
<td>21</td>
</tr>
<tr>
<td>Cerivastatin</td>
<td>102</td>
<td>12827</td>
<td>126</td>
</tr>
<tr>
<td>Bosentan</td>
<td>96</td>
<td>8489</td>
<td>89</td>
</tr>
<tr>
<td>Fluvastatin</td>
<td>475</td>
<td>76513</td>
<td>161</td>
</tr>
<tr>
<td>Rosuvastatin</td>
<td>98</td>
<td>1190</td>
<td>12</td>
</tr>
<tr>
<td>Valsartan</td>
<td>22</td>
<td>2463</td>
<td>110</td>
</tr>
<tr>
<td>Repaglinide</td>
<td>319</td>
<td>13941</td>
<td>44</td>
</tr>
<tr>
<td>Geometric mean</td>
<td>-</td>
<td>-</td>
<td>58</td>
</tr>
</tbody>
</table>

<sup>1</sup>represents the sum of SCLint,u, bile and SCLint,u,met as for these 3 compounds both CL mechanisms are occurring and they cannot be uniquely identified in the fitting process.
Table 4 – Predicted CL and Vss parameters for the seven drugs investigated using different modeling scenarios

<table>
<thead>
<tr>
<th>Compound</th>
<th>CL mL/min/kg</th>
<th>Vss L/Kg</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Clinical&lt;sup&gt;1&lt;/sup&gt;</td>
<td>In vitro scaled&lt;sup&gt;2&lt;/sup&gt;</td>
</tr>
<tr>
<td>Pravastatin</td>
<td>14</td>
<td>8.0</td>
</tr>
<tr>
<td>Cerivastatin</td>
<td>2.9</td>
<td>0.17</td>
</tr>
<tr>
<td>Bosentan</td>
<td>2.3</td>
<td>0.14</td>
</tr>
<tr>
<td>Fluvastatin</td>
<td>8.7</td>
<td>0.46</td>
</tr>
<tr>
<td>Rosuvastatin</td>
<td>11</td>
<td>4.0</td>
</tr>
<tr>
<td>Valsartan</td>
<td>0.49</td>
<td>0.15</td>
</tr>
<tr>
<td>Repaglinide</td>
<td>7.8</td>
<td>0.74</td>
</tr>
</tbody>
</table>

Absolute average
fold error

- 7.1 1.3 - 3.0 1.7

<sup>1</sup> reported i.v. CL and Vss parameters from publications reported in Supplementary Material, Table S1; <sup>2</sup> predicted i.v. CL and Vss parameters determined by non-compartmental analysis of predicted profiles.
FIGURE 1

A

Media, u, Vm ─────> Cell, u, Vc ─────> Bile

↓

CLint, u, passive

B

Media, u, Vm ─────> Cell, u, Vc ─────> Bile

↓

CLint, u, active

C

Media + Bile, u, Vm ─────> Cell, u, Vc

↓

CLint, u, passive

CLint, u, bile
FIGURE 2

QLUNG → LUNG → QLUNG

QTISSUE 1 → TISSUE 1 → QTISSUE 1

QTISSUE 2 → TISSUE 2 → QTISSUE 2

QTISSUE N → TISSUE N → QTISSUE N

QKIDNEY → KIDNEY → QKIDNEY

QLIVER → LIVER EC SPACE 5 → LIVER IC SPACE 5 → LIVER IC SPACE 4 → LIVER EC SPACE 4

QLIVER → LIVER EC SPACE 3 → LIVER IC SPACE 3 → LIVER IC SPACE 2 → LIVER EC SPACE 2

QLIVER → LIVER EC SPACE 1 → LIVER IC SPACE 1

CLint,u,kidney

QGUT

QHEPATIC ARTERY

QGUT

QSPLEEN

VENOUS BLOOD

ARTERIAL BLOOD

SCLint,b,active

SCLint,b,passive

SCLint,b,met
FIGURE 4

A. Pravastatin 9.4 mg 2min iv infusion

B. Cerivastatin 0.1 mg 1min iv infusion

C. Bosentan 10 mg 5min iv infusion

D. Fluvastatin 1.9 mg 20min iv infusion

E. Rosuvastatin 8 mg 4hr iv infusion

F. Valsartan 20 mg iv bolus

G. Repaglinide 2 mg 15min iv infusion
Supplementary Material

Journal:

Drug Metabolism and Disposition

Title of Article:

Mechanistic pharmacokinetic modeling for the prediction of transporter-mediated disposition in human from sandwich culture human hepatocyte data

Author’s names:

Hannah M Jones, Hugh A Barton, Yurong Lai, Yi-an Bi, Emi Kimoto, Sarah Kempshall, Sonya C Tate, Ayman El-Kattan, J Brian Houston, Aleksandra Galetin and Katherine S Fenner
**Figure S1:** Raw SCHH data for (A-B) pravastatin, (C-F) cerivastatin, (G) bosentan, (H-J) fluvastatin, (K-R) rosuvastatin, (S-U) valsartan and (V-X) repaglinide.

*Solid diamonds represent HBSS only, open squares represent Ca2+/Mg2+ free, solid triangles represent HBSS + Rifamycin SV*
Figure S2: Fitted versus observed human plasma concentration time data for (A) pravastatin, (B) cerivastatin, (C) bosentan, (D) fluvastatin, (E) rosuvastatin, (F) valsartan, (G) repaglinide.
Figure S3: Contour plots for pairs of fitted parameters (A-C) pravastatin, (D-F) cerivastatin, (G-I) bosentan, (J-L) fluvastatin, (M-O) rosvastatin, (P-R) valsartan and (S) repaglinide.
**Table S1** – Clinical pharmacokinetic properties and substrate specificity for the compounds studied

<table>
<thead>
<tr>
<th>Compound</th>
<th>CL, i.v. mL/min/kg</th>
<th>Vss, i.v. L/Kg</th>
<th>% renal CL</th>
<th>Substrate specificity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pravastatin¹</td>
<td>14</td>
<td>0.46</td>
<td>47</td>
<td>OATP1B1, MRP2</td>
</tr>
<tr>
<td>Cerivastatin²</td>
<td>2.9</td>
<td>0.33</td>
<td>0</td>
<td>OATP1B1, CYP3A4, 2C8</td>
</tr>
<tr>
<td>Bosentan³</td>
<td>2.3</td>
<td>0.67</td>
<td>0</td>
<td>OATP1B1, 1B3, 2B1, CYP3A4, 2C9</td>
</tr>
<tr>
<td>Fluvastatin⁴</td>
<td>8.7</td>
<td>0.16</td>
<td>0</td>
<td>OATP1B1, 2B1, 1B3, CYP2C9</td>
</tr>
<tr>
<td>Rosuvastatin⁵</td>
<td>11</td>
<td>1.7</td>
<td>30</td>
<td>OATP1B1, 1B3, 2B1, BCRP</td>
</tr>
<tr>
<td>Valsartan⁶</td>
<td>0.49</td>
<td>0.23</td>
<td>29</td>
<td>OATP1B1, 1B3</td>
</tr>
<tr>
<td>Repaglinide⁷</td>
<td>7.8</td>
<td>0.35</td>
<td>0</td>
<td>OATP1B1, CYP3A4, 2C8</td>
</tr>
</tbody>
</table>

¹ Singhvi et al., 1990; ² Muck et al., 1997; ³ Weber et al., 1996; ⁴ Lindahl et al., 1996; ⁵ Martin et al., 2003; ⁶ Flesch et al., 1997; ⁷ Hatorp et al., 1998
### Table S2: Quantification of OATP1B1, 1B3 and 2B1 in suspension hepatocytes and SCHH

<table>
<thead>
<tr>
<th></th>
<th>Suspension</th>
<th>SCHH at day 5</th>
<th>Change compared to suspension</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(fmol/µg protein)</td>
<td>%</td>
<td></td>
</tr>
<tr>
<td>OATP1B1</td>
<td>3.42±0.11</td>
<td>5.28±0.22</td>
<td>154(^1)</td>
</tr>
<tr>
<td>OATP1B3</td>
<td>1.50±0.15</td>
<td>0.88±0.11</td>
<td>59(^1)</td>
</tr>
<tr>
<td>OATP2B1</td>
<td>1.84±0.15</td>
<td>1.23±0.12</td>
<td>67(^*)</td>
</tr>
</tbody>
</table>

\(^1\) P<0.05, as compared to suspension.
Supplementary References


