Application of a Physiologically Based Pharmacokinetic Model to Assess Propofol Hepatic and Renal Glucuronidation in Isolation: Utility of In Vitro and In Vivo Data

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ABSTRACT
A physiologically based pharmacokinetic (PBPK) modeling approach was used to assess the prediction accuracy of propofol hepatic and extrahepatic metabolic clearance and to address previously reported underprediction of in vivo clearance based on static in vitro–in vivo extrapolation methods. The predictive capacity of propofol intrinsic clearance data (Cl\text{int}) obtained in human hepatocytes and liver and kidney microsomes was assessed using the PBPK model developed in MATLAB software. Microsomal data obtained by both substrate depletion and metabolite formation methods and in the presence of 2% bovine serum albumin were considered in the analysis. Incorporation of hepatic and renal in vitro metabolic clearance in the PBPK model resulted in underprediction of propofol clearance regardless of the source of in vitro data; the predicted value did not exceed 35% of the observed clearance. Subsequently, propofol clinical data from three dose levels in intact patients and anhepatic subjects were used for the optimization of hepatic and renal Cl\text{int} in a simultaneous fitting routine. Optimization process highlighted that renal glucuronidation clearance was underpredicted to a greater extent than liver clearance, requiring empirical scaling factors of 17 and 9, respectively. The use of optimized clearance parameters predicted hepatic and renal extraction ratios within 20% of the observed values, reported in an additional independent clinical study. This study highlights the complexity involved in assessing the contribution of extrahepatic clearance mechanisms and illustrates the application of PBPK modeling, in conjunction with clinical data, to assess prediction of clearance from in vitro data for each tissue individually.

Introduction
Propofol is a probe substrate for UGT1A9 and is also cleared by cytochrome P450 (P450) enzymes, primarily via CYP2B6 and to a minor extent by CYP2C9, CYP1A2, and CYP3A4 (Guitton et al., 1998; Court et al., 2001; Oda et al., 2001; Court, 2005). Propofol has not been reported to be a substrate for transporters and undergoes minimal renal excretion (Vree et al., 1987; Simons et al., 1988; Verolli et al., 1992), and therefore represents a good candidate for exploring the prediction of clearance due to metabolism alone. However, pronounced underprediction of in vivo clearance has been observed for propofol using static in vitro–in vivo extrapolation (IVIVE) techniques (Killford et al., 2009; Gill et al., 2012). Use of inappropriate in vitro systems, exclusion of extrahepatic metabolism, inadequacy of scaling factors, and/or models applied to the in vitro data may all contribute to this underprediction trend. Both in vivo and in vitro data indicate that the kidneys play an important role in the glucuronidation of certain drugs, including morphine and propofol (Mazoit et al., 1990; Pichette and du Souich, 1996; Soars et al., 2002; Takizawa et al., 2005a; Gill et al., 2012). Similarly, there is extensive evidence that both P450 and conjugation metabolism in the small intestine represent important contributors to drug clearance (Galetin et al., 2008; Cubitt et al., 2009, 2011; Gertz et al., 2010). We previously showed that the inclusion of renal metabolic clearance data in IVIVE improved prediction of glucuronidation clearance; however, underprediction was still apparent for certain drugs, including propofol (Gill et al., 2012).

Recently, there has been an increased use of dynamic modeling techniques such as physiologically based pharmacokinetic (PBPK) models to predict drug exposure and clearance (Rowland et al., 2011; Huang and Rowland, 2012). The application of PBPK models by the pharmaceutical industry and regulatory bodies together with some of the limitations of this approach have been highlighted recently (Poulin et al., 2011; Zhao et al., 2011; Huang and Rowland, 2012; Jones et al., 2012). A variety of physiologic and drug-specific parameters, including in vitro and in vivo clearance and tissue binding data, can...
be incorporated to optimize these models and improve prediction of in vivo pharmacokinetics (Nestorov, 2007; Huang and Rowland, 2012). The relative accuracy of predictions of hepatic versus extrahepatic clearance from in vitro data has not previously been assessed as the availability of in vivo data to allow such analysis is very limited. However, the use of a dynamic approach such as PBPK modeling in conjunction with suitable in vivo data would be expected to improve the understanding of the predictive capacity of in vitro clearance data for different tissues in isolation.

Previous PBPK modeling for propofol has not incorporated in vitro hepatic clearance data and has ignored the contribution of renal metabolic clearance (Levitt and Schneider, 2005; Upton and Ludbrook, 2005). In vivo propofol concentration-time data for subjects during liver transplantation have been reported in the literature (Veroli et al., 1992); under these conditions, the clearance observed is solely mediated by extrahepatic metabolism. Such in vivo data used in conjunction with mechanistic PBPK models allow independent assessment of the predictive capacity of in vitro clearance data for hepatic and renal metabolism.

The aim of this study was to apply a PBPK model to assess the prediction of propofol systemic clearance using in vitro metabolism data from the kidney and liver. Propofol blood concentration-time profiles and systemic clearance after intravenous dosing were predicted using intrinsic clearance (CLint) data obtained in different in vitro systems (microsomes and hepatocytes); the predictive ability of hepatic in vitro CLint data from different sources was assessed. The rich in vivo data available for propofol were used to optimize the PBPK model and to bridge the gap in the IVIVE of propofol clearance. In vivo data from anhepatic patients allowed the analysis of the prediction success of renal glucuronidation clearance in isolation from the liver; in conjunction with data from intact patients, these data were used to refine the prediction of renal and hepatic clearance.

Materials and Methods

Chemicals. Propofol, alamethicin (from Trichoderma viride), UDP-glucuronic acid, EDTA, bovine serum albumin (BSA), saccharic acid lactone, MgCl₂, NAD⁺, isocitric acid, and isocitric acid dehydrogenase were purchased from Sigma-Aldrich (Gillingham, UK). Propofol glucuronide was purchased from Toronto Research Chemicals (North York, ON, Canada). All other reagents were of the highest grade available.

Source of the Microsomes. BD UltraPool human liver microsomes (HLMs) were purchased from BD Gentest (Woburn, MA). HLMs were pooled from 150 Caucasian donors, 59% of whom were female, with a mean age of 53 years (range, 18–79 years). Pooled human kidney and intestinal microsomes (HKMs and HIMs, respectively) were purchased from Xenotech (Tebu-Bio Ltd., Peterborough, UK). HKMs were pooled from eight donors, 88% of whom were Caucasian and 50% were female, and a mean age of 61 years (range, 46–69 years). HIMs were pooled from 13 donors, 92% of whom were Caucasian and 46% were female, with a mean age of 40 years (range, 18–55 years). Microsomes were stored at –80°C. Activity for pooled HLMs was reported as 0.054 nmol/min per milligram protein for CYP2B6 and 2.00 nmol/min per milligram protein for UGT1A9, using specific probe substrates. For HKM and HIM pools, glucuronidation capacity was characterized for 4-methylumbelliferone (a substrate of multiple uridine diphosphate glucuronosyltransferases [UGTs]), with reported activity of 125 and 7.22 nmol/min per milligram protein, respectively.

Experimental Conditions for Microsomal P450 Depletion Assays. Incubations were performed in duplicate (Eppendorf vials (Eppendorf, Hamburg, Germany) using a substrate depletion approach. A final substrate concentration of 3 μM was used, which is >3-fold below the reported Kᵣ (Michaelis-Menten constant) value in HLMs (Guitton et al., 1998; Court et al., 2001; Al-Jahdari et al., 2006), ensuring suitable conditions to determine the intrinsic clearance of propofol. Propofol was preincubated at a microsomal protein concentration of 0.5 mg/ml in 0.1 M phosphate buffer, pH 7.4, for 5 minutes at 37°C shaken at 900g using an Eppendorf thermomixer (Kilford et al., 2009). The reaction was initiated by addition of the NADPH regenerating system, containing 1 mM NADPH, 7.5 mM isocitric acid, 1 mM isocitric acid dehydrogenase, and 10 mM MgCl₂ in a final incubation volume of 1 ml (Kilford et al., 2009). Concentration of the organic solvent used (methanol) was 0.5% v/v of the incubation media. The total length of the incubation was 45 minutes. To terminate the reaction, 100-μl samples of the incubation were removed at each time point and added to an equal volume of ice-cold methanol containing 1 μM tolbutamide as the internal standard. Samples were kept at –20°C for at least 1 hour and then centrifuged (MSE Mistral 3000i centrifuge; MSE, London, UK) at 4°C and 2500g for 30 minutes. An aliquot of the supernatant (30 μl) was analyzed by liquid chromatography–tandem mass spectrometry (LC-MS/MS) for propofol concentration. Experiments were repeated on three separate occasions. No P450 clearance was observed in HKMs and HIMs in the absence of BSA.

Experiments performed in the presence of BSA used the same methodology, with the exception of the addition of 2% BSA to the incubation. Samples of 100 μl of the incubation were removed at each time point and added to double volume of ice-cold acetone/95%/tolbutamide containing 1 μM tolbutamide to terminate the reaction. Samples were refrigerated for at least 10 minutes and then centrifuged (Eppendorf Mini Spin) at 13,400g and room temperature for 5 minutes prior to analysis by LC-MS/MS for propofol concentration. Similarly to data in the absence of BSA, no P450 clearance was observed in HKMs and HIMs in the presence of BSA. Nonenzymatic depletion of propofol was monitored and clearance estimates were corrected for the observed nonenzymatic loss.

Experimental Conditions for Propofol Glucuronide Formation in Microsomes. Incubations were performed in duplicate using Eppendorf vials. Initial experiments showed propofol glucuronide formation was linear up to 10 min at protein concentrations ≤0.5 mg/ml in all tissue samples, both in the presence and absence of 2% BSA. The time period and protein concentration for each tissue were selected to ensure that metabolite formation was within the linear range. Substrate concentrations ranging from 0.5 to 400 μM were used to assess the kinetic parameters Kₘ and Vₘₕ (maximum rate of reaction), with the exception of HIMs in which a concentration range of 2.5 to 600 μM was employed. Microsomal protein concentrations for experiments performed in the absence of BSA were 0.2, 0.1, and 0.4 mg/ml for HLMs, HKMs, and HIMs, respectively. Microsomes were treated with alamethicin (50 μg/mg protein) for 15 minutes on ice as reported previously (Fishier et al., 2000; Cubitt et al., 2009; Kilford et al., 2009). Activated microsomes were preincubated with propofol and 0.1 M phosphate buffer, pH 7.1, containing 3.45 mM MgCl₂, 1.15 mM EDTA, and 15 μM saccharic acid lactone, for 5 minutes at 37°C shaken at 900g (Fishier et al., 2001; Cubitt et al., 2009; Kilford et al., 2009). The reaction was initiated after the addition of UDP-glucuronic acid (5 mM in incubation) to give a final incubation volume of 0.1 ml. Organic solvent (methanol) made up 1% v/v of the incubation media. Control incubations were performed with no cofactor present to account for any potential cofactor independent formation of the metabolite over the incubation time. In the absence of BSA, the reaction was terminated after 10 minutes by addition of an equal volume ice-cold methanol containing 1 μM of the internal standard tolbutamide. Samples with starting propofol concentrations in excess of 10 μM were diluted in ice-cold blank 1:1 matrix:methanol to give a final propofol concentration of ≤10 μM. Samples were kept at –20°C for at least 1 hour and then centrifuged (MSE Mistral 3000i centrifuge) at 4°C and 2500g for 30 minutes. An aliquot of the supernatant (20 μl) was analyzed by LC-MS/MS for propofol glucuronide concentration.

Incubation conditions for experiments including BSA were comparable with those without BSA, with the exception of the addition of 2% BSA. Optimal conditions of 0.1 mg/ml microsomal protein for HLMs and HKMs or 0.2 mg/ml for HIMs were employed for the kinetic assessment in the presence of BSA. The reaction was terminated after 10 minutes by addition of double volume ice-cold acetone/tolbutamide containing 1 μM of the internal standard tolbutamide. Samples with starting propofol concentrations in excess of 10 μM were diluted in ice-cold blank 1:2 matrix:acetone to give a final propofol concentration of ≤10 μM. Samples including BSA were refrigerated for at least 10 minutes and then centrifuged (Eppendorf Mini Spin) at 13,400g and room temperature for 5 minutes. An aliquot of the supernatant (20 μl) was analyzed by LC-MS/MS for propofol glucuronide concentration. Experiments were repeated on three separate occasions.
Propofol intrinsic clearance due to glucuronidation (CL_{int,UGT}) data in HLMs, HKMs, and HIMs obtained by the depletion method and corresponding incubation details were reported previously (Gill et al., 2012). The batch of HLMs and HKMs used were the same as those used herein for determination of intrinsic clearance due to P450s (CL_{int,P450}) and metabolite formation CL_{int,UGT}. A different pool of HIMs was used due to the limited supply of the batch used previously; the donor demographics and activity characterization were comparable between the two HIM pools.

In addition to microsomal data, use of in-house unbound CL_{int} data obtained in hepatocytes at a single propofol concentration of 5 μM was also investigated. CL_{int} data were generated by the substrate depletion approach using pooled cryopreserved human hepatocytes from 10 donors purchased from In Vitro Technologies (Baltimore, MD). Experiments were performed in the absence and presence of nonspecific P450 inhibitor 1-aminobenzotriazole (ABT) at 2.5 mM to determine total CL_{int} (combined CL_{int,P450} and CL_{int,UGT}) and CL_{int,UGT} respectively.

LC-MS/MS. Propofol and propofol glucuronide were analyzed on a Waters 2790 (Waters, Milford, MA) or an Agilent 1100 high-performance liquid chromatography system (Agilent Technologies, Santa Clara, CA), respectively, with a Micromass Quattro Ultima (Waters) triple quadrupole mass spectrometer in negative mode. Source temperature was 125°C, desolvation temperature was 350°C, and the desolvation gas flow rate was 600 l/h. Cone gas flow rate was 150 l/h for propofol and 50 l/h for propofol glucuronide. The capillary voltage was 3.25 kV. Propofol and propofol glucuronide were analyzed using single ion recording due to their limited fragmentation. The transitions for propofol, propofol glucuronide, and tolbutamide were 176.90, 353.55, and 269.00 m/z, respectively. Analytes were separated using a Luna C18 (3 μm, 50 × 4.6 mm) column (Phenomenex, Macclesfield, UK). Four mobile phases were used, with varying gradients for each compound: 1) 90% water, 10% methanol, and 0.05% formic acid; 2) 10% water, 90% methanol, and 0.05% formic acid; 3) 90% water, 10% methanol, and 1 mM ammonium acetate; and 4) 10% water, 90% methanol, and 1 mM ammonium acetate. The flow rate was 1 ml/min, splitting to 0.25 ml/min prior to entry into the mass spectrometer. Cone voltage was set to 60 V for propofol and 75 V for tolbutamide, and the corresponding retention times were 3.40 and 2.80 minutes. When analyzing for the metabolite, the cone voltage was set to 30 and 45 V for propofol glucuronide and tolbutamide, with retention times of 3.40 and 3.20 minutes, respectively. The lower limit of quantification for propofol and its glucuronide metabolite were 0.039 and 0.020 μM.

Data Analysis. The mean propofol concentrations of the duplicate samples from the P450 depletion assays at each time point were analyzed using GraFit 5 (Erithacus Software, Horley, UK) to determine the elimination rate constant (k) by fitting a single exponential equation to the data. This rate constant was used to calculate the CL_{int,P450} (Eq. 1). The nonenzymatic loss was taken into account when analyzing the experimental data. The CL_{int,P450} values were calculated for nonspecific binding (CL_{int,P450}/f_{u,inc}), using the experimentally determined fraction unbound in the microsomal incubation (f_{u,inc}) in the presence and absence of 2% BSA reported previously (Gill et al., 2012), to generate the unbound intrinsic clearance (μl/min per milligram protein). The unbound CL_{int,P450} values have been reported.

\[
CL_{int,P450} = \frac{k \times \text{volume of incubation}}{\text{amount of microsomal protein in assay}}
\]

In vitro microsomal CL_{int} data from the various assays were scaled with the microsomal protein yields: 40, 12.8, and 20.6 mg protein per gram tissue were used for hepatic, renal, and intestinal data, respectively (Al-Jahdari et al., 2006; Barter et al., 2007; Cubitt et al., 2009). Hepatocyte unbound CL_{int} data were scaled with a hepatocellularity value of 120 × 10^6 cells/g tissue (Brown et al., 2007). The scaled CL_{int} data were compared between the in vitro systems.

Development of the Propofol PBPK Model. A previously reported in-house whole-body PBPK model (Gertz et al., 2011) was adapted to predict propofol concentration-time profiles and pharmacokinetics. The resultant model contained 14 organ compartments connected by arterial and venous blood supplies (Fig. 1). An additional compartment representing the rest of the body was included, which accounted for <5% of the total body weight and <8% of the total blood flow. Propofol plasma binding, blood to plasma partition coefficient, and renal excretion of unchanged drug were collated from the literature, as reported previously (Gill et al., 2012). Tissue to plasma concentration ratios (K_{p,t}) were predicted using the Rodgers and Rowland (2006) method, and are detailed in Fig. 1 and Supplemental Table 1. Tissue blood flow and volume were collated from the literature (ICRP, 2002) (Fig. 1; Supplemental Table 1).

All tissues were assumed to be well stirred compartments where unbound tissue concentration is at equilibrium with the unbound concentration in the emerging blood (Pang and Rowland, 1977). Eq. 3 was used for noneliminating organs (Nestorov, 2003).

\[
\frac{dC_T}{dt} = \frac{Q_T \left( C_{h,T} - C_T \right)}{V_T} (3)
\]

where C_T, Q_T, C_{h,T}, and V_T represent tissue concentration, tissue blood flow, unbound concentration in arterial blood, tissue blood concentration ratio, and tissue volume.

The liver, kidney, and enterocytes were considered to potentially contribute to systemic drug clearance. The liver and kidney were separated into cellular tissue and blood compartments to allow assessment of the extraction ratio across each tissue. Eq. 4 represents the tissue compartment and Eq. 5 represents the blood compartment for the kidney and the liver.

\[
\frac{dC_{T,T}}{dt} = \frac{Q_T \left( C_{h,T} - C_{T,T} \right) - \frac{f_{u,b}}{K_{b,T}} \cdot C_{T,T} \cdot \left( CL_{int,UGT} + CL_{int,P450} \right)}{V_{T,T}} (4)
\]

where C_{T,T}, Q_T, C_{h,T}, f_{u,b}, V_{T,T}, and V_{T,T} represent tissue cell concentration, permeability surface area (set to >10,000 × tissue blood flow to ensure perfusion limited kinetics), concentration in blood residing in the tissue, blood binding, volume of tissue cells, and volume of blood residing in tissue. Microsomal CL_{int,UGT} and CL_{int,P450} were scaled with the microsomal recovery and mass of the relevant tissue, as described above. Hepatocyte CL_{int} (combined CL_{int,UGT} and CL_{int,P450}) data were scaled with the hepatocellularity value and the mass of the liver. Kidney and liver blood volumes were set to 100 ml (ICRP, 2002). The portal vein concentration represents the differential of emergent blood concentrations from the large and small intestine, enterocytes, stomach, spleen, and pancreas.

The extraction ratio for the enterocytes was not calculated and therefore this tissue was defined as a single compartment (Eq. 6).

\[
\frac{dC_T}{dt} = \frac{Q_T \left( C_{h,T} - C_T \right) - \frac{f_{u,b}}{K_{b,T}} \cdot C_T \cdot \left( CL_{int,UGT} + CL_{int,P450} \right)}{V_T} (5)
\]

The blood flow to the small intestine represents approximately 10% of the cardiac output (ICRP, 2002) and the enterocytic blood flow represents approximately 50% of the small intestinal blood flow (Gertz et al., 2011). The
rate equations were solved in MATLAB software (version 7.12.0; MathWorks, Natick, MA) using the ODE15s solver. The dose recovery over time was assessed using mass balance equations.

Assumptions Applied to the Propofol PBPK Model. The following assumptions were made: 1) there is no active uptake or efflux of propofol in any tissue and therefore tissue distribution/elimination is perfusion rate limited; this assumption seems justifiable because, together with the high lipophilicity of propofol (Reiner et al., 2009), there is a lack of in vitro or in vivo data suggesting that propofol is a substrate of transporters; 2) propofol does not affect the cardiac output (Grounds et al., 1985; Price et al., 1992); and 3) the volume of distribution at steady state (Vss) for the anhepatic patients was not expected to be markedly different from that in healthy subjects. This is supported by a number of studies where changes in Vss for patients with varying grades of liver cirrhosis were not apparent in comparison with patients with healthy livers (mean values ranging from 202 to 637 l in subjects with mild liver cirrhosis) (Servin et al., 1988a, b, 1990). Similarly, no changes in propofol Vss during chronic hepatic cirrhosis were stated in the AstraZeneca propofol product monograph (http://www.astrazeneca.ca/documents/Product-Portfolio/DIPRIVAN_PM_en.pdf). In addition, no change in blood binding of propofol was observed in patients with mild liver cirrhosis (Servin et al., 1990).

Validation of Propofol PBPK Model. The PBPK model was validated using data from two clinical studies, covering a propofol dose range of 2–18 mg/kg (Gepts et al., 1987; Doenicke et al., 1997). The study details and subject demographics for these studies are presented in Table 1. The cardiac output was corrected for the mean age of each dose group (Brown et al., 1997). One of the studies (Gepts et al., 1987) included elderly subjects but no alterations were made to the tissue volumes for this population. Plasma binding was assumed to be equivalent in young and elderly subjects for which evidence could be found in the literature (Kirkpatrick et al., 1988). Minimal age and sex effects have been observed for in vitro microsomal UGT activity for a range of probe substrates including propofol (Court, 2010); therefore, no alterations were made to microsomal protein yields for the different populations.

Vss for propofol is highly variable in the literature, with mean values reported in 14 studies collated from the literature (including 194 subjects) ranging from 121 to 722 l; no trends were observed regarding sex, age, dose level, length of infusion, or disease status (details provided in Supplemental Fig. 1). The Vss data considered for this analysis were taken from studies in which the propofol blood concentrations were available over at least 8 hours, which is considered adequate to describe its pharmacokinetics. Where necessary, the predicted Kp values were optimized using a universal scalar for all tissues to ensure that the reported Vss was recovered correctly for each dose group used in the model validation and for the prediction of propofol blood concentration-time profiles.

Contradicting reports exist in the literature over the potential propofol metabolism and/or sequestration in the lung (Dawidowicz et al., 2000; He et al., 2000; Hiraoka et al., 2005; Takizawa et al., 2005a). Reduction in propofol concentration across the lung in one study was proportional to the formation of the quinol metabolite and assessment of the blood concentration data suggested a lung extraction ratio of 0.40 (Dawidowicz et al., 2000). Assumption of that extent of lung extraction would predict clearance in excess of the observed values for the in vivo studies used in the current analysis. In contrast, the majority of other available data proposed that the reduction in propofol concentration across the lung was not due to metabolism but binding, with subsequent slow release from the lung (He et al., 2000; Hiraoka et al., 2005; Levitt and Schnider, 2005; Takizawa et al., 2005a; Upton and Ludbrook, 2005). Previous studies showed that sequestration of propofol was not apparent after intravenous infusion dosing, particularly for patients aged >35 years (He et al., 2000; Levitt and Schnider, 2005). The majority of the in vivo data used in the
TABLE 1
Data are presented as mean (± S.D.) unless otherwise specified.

<table>
<thead>
<tr>
<th>Study</th>
<th>Status of Patients</th>
<th>Dose Type</th>
<th>Dose Level</th>
<th>Subjects</th>
<th>Body Weight</th>
<th>Age</th>
<th>Male</th>
</tr>
</thead>
<tbody>
<tr>
<td>Doenicke et al. (1997)</td>
<td>Intact</td>
<td>Intravenous bolus</td>
<td>2</td>
<td>12</td>
<td>ND</td>
<td>24–42</td>
<td>100</td>
</tr>
<tr>
<td>Gepts et al. (1987)</td>
<td>Intact</td>
<td>Intravenous infusion over 2 h</td>
<td>6</td>
<td>6</td>
<td>75.0 (7.2)</td>
<td>60.3 (4.9)</td>
<td>100</td>
</tr>
<tr>
<td>Veroli et al. (1992)</td>
<td>Anhepatic</td>
<td>Intravenous bolus</td>
<td>0.5</td>
<td>10</td>
<td>60.0 (7.0)</td>
<td>38.0 (ND)</td>
<td>ND</td>
</tr>
</tbody>
</table>

ND, not detailed; PBPK, physiologically based pharmacokinetic.
* Mean age not detailed; age range of subjects presented instead.

current analysis were reported in subjects of similar or greater age after an infusion dose, which may explain the lack of evidence for sequestration in the lung. With the lack of suitable data to define association and dissociation rate constants for potential propofol binding in the lung, a truly physiologic model to account for this process could not have been included within the current PBPK model and therefore no adjustment for lung sequestration was employed. Accumulation within lysosomes, one potential route of sequestration within tissues such as lungs, is not anticipated for propofol due to its physicochemical properties. The potential for propofol sequestration in other tissues or via other mechanisms could not be investigated, although this cannot be ruled out.

In Vitro Clearance Data Used as Inputs for the PBPK Model to Predict in Vivo Clearance of Propofol. In vitro CLint,P450 and CLint,UGT data for HLMs, HKMs, and HIMs were used within the PBPK model to assess the success of in vivo clearance predictions. The prediction success of systemic clearance was investigated using CLint,UGT data from substrate depletion assays, obtained previously from Gill et al. (2012), and metabolite formation assays (current study). CLint,P450 data were obtained using the substrate depletion approach (detailed herein). Clearance prediction success for in-house data determined via the substrate depletion method in hepatocytes (computed CLint,P450 and CLint,UGT) in conjunction with CLint,UGT data from HKMs and HIMs from either the substrate depletion or metabolite formation methods was also investigated. Details of the in vitro data used for prediction of in vivo clearance are given in Table 2.

In addition, the in vitro data were used in the PBPK model to predict hepatic and renal extraction ratios (EQh and EQr, respectively) at steady state. The predicted extraction ratios were compared with values reported in a clinical study (Hiraoka et al., 2005), which was independent to the clinical studies used for model validation. The reported EQh (0.93) and EQr (0.69) were obtained from blood concentration measurements in the renal vein, hepatic vein, and radicle artery at steady state. The extraction ratios for each tissue (EQt) were calculated by Eq. 7.

$$ET = \frac{C_{in} - C_{T,B}}{C_{in}}$$  

where C_{in} represents the concentration in blood entering the tissue.

Optimization of Predictions of Renal and Hepatic Clearance from in Vitro Data. Initial analysis indicated underprediction of hepatic and renal CLint regardless of the in vitro system used. Therefore, optimization was performed to bridge the gap in IVIVE and determine the empirical scaling factors required for the in vitro data to accurately recover in vivo clearance, by fitting the PBPK model to the in vivo propofol blood concentration-time profiles. Data from three dose levels (6–18 mg/kg) in intact patients (Gepts et al., 1987) and from anephric patients (Veroli et al., 1992) (who received a 0.5 mg/kg dose) were fitted simultaneously in MatLab using a nonlinear least-squares regression analysis (lsqnonlin function). For the anephric patients, the PBPK model was adapted removing the liver compartment to reflect the in vivo situation. Data from the anephric patients were used to delineate the contribution of the renal CLint,UGT and to refine the PBPK model with respect to renal metabolism, whereas data from intact patients provided input for optimization of the hepatic CLint and Kp. Propofol concentration-time profiles in the anephric patients were only available for 1 hour; therefore, Vss for these patients could not be determined from the profile. The data from intact patients were used to inform the optimization of the Vss for both intact and anephric patients in the simultaneous fitting routine. Optimization of hepatic and renal CLint in the PBPK model allowed the assessment of the degree of underprediction of in vitro CLint for each tissue individually.

In the in vitro CLint data for the intestinal microsomes were low in comparison with that in the kidney and the liver. Considering their lower tissue mass and blood flow, the enterocytes were not expected to contribute extensively to the metabolism of propofol. There is also a lack of suitable in vivo data for assessment of the predictive capacity of the intestinal in vitro CLint,UGT data. For these reasons, the optimization was performed for the kidney and liver metabolism only.

Results
Propofol P450 Depletion Assays in Human Hepatic, Renal, and Intestinal Microsomes. No P450-mediated clearance was observed in renal or intestinal microsomes; however, in the liver, CLint,P450 in the presence of BSA was 2.2-fold higher than CLint,UGT obtained via substrate depletion (Table 2). BSA increased propofol CLint,P450 by 2-fold in HLMs (Table 2), whereas a 3-fold change in CLint,UGT in the presence of BSA was reported previously (Gill et al., 2012). Depletion profiles in HLMs over time are shown in the Supplemental Fig. 2.

Propofol Glucuronide Formation Assays in Human Hepatic, Renal, and Intestinal Microsomes. Propofol Kmax and Vmax were determined in hepatic, renal, and intestinal microsomes both in the presence and absence of BSA; corresponding kinetic profiles are shown in Fig. 2. Use of a protein concentration of 0.1 mg/ml for HLMs and HKMs and HIMs in combination with 2% BSA had no impact on protein P450-mediated metabolism. There is also a lack of suitable in vivo data for assessment of the predictive capacity of the intestinal in vitro CLint,UGT data. For these reasons, the optimization was performed for the kidney and liver metabolism only.
Comparison of CLint Data Derived Using Different in Vitro Systems.

In the absence of BSA, propofol scaled CLint,UGT estimates derived from metabolite formation assays in HLMs and HKMs were lower than those from substrate depletion assays (0.6 versus 2.7 ml/min per gram tissue and 0.7 versus 1.2 ml/min per gram tissue, respectively) (Table 2). However, in the presence of 2% BSA, CLint,UGT data were comparable from both assays (Table 2). Consequently, the increase in propofol CLint,UGT upon inclusion of BSA in HLMs and HKMs was greater for the metabolite formation data in comparison with the substrate depletion data. For HIMs, CLint,UGT estimates obtained by metabolite formation were lower both in the presence (9-fold) and absence (22-fold) of BSA compared with those obtained by depletion (Table 2). Combined CL_{int,P450} and CL_{int,UGT} data for hepatic microsomes (26 and 29 ml/min per gram tissue using the depletion or formation assays in the presence of BSA, respectively) gave higher total CLint per gram of liver than estimates obtained from hepatocytes (18 ml/min per gram tissue).

The in vitro fm,UGT for HLM CLint data derived using substrate depletion were 0.23 and 0.31 in the absence and presence of 2% BSA, respectively. When using HLM CLint,UGT data derived via metabolite formation in the presence of BSA, the resulting fm,UGT (0.37) was similar to using HLM CLint,UGT data obtained by substrate depletion. Use of metabolite formation CLint,UGT data obtained in the absence of BSA reduced the fm,UGT estimate. The fm,UGT determined in hepatocytes using the substrate depletion approach in the presence and absence of ABT (0.43) was similar to that determined from HLMs. The in vitro fm,UGT for HKMs and HIMs were 1.0 due to the lack of P450 clearance observed in these tissues.

Prediction of Propofol Clearance and Blood Concentration-Time Profiles Using Data from Different in Vitro Systems. Blood concentration-time profiles in vivo and clearance were predicted using various combinations of in vitro CL_{int,UGT} and CL_{int,P450} data from the different systems used; a summary of the prediction accuracy for each system is shown in Table 3 (full details for individual dose groups are

### Table 2

<table>
<thead>
<tr>
<th>Assay Type</th>
<th>Parameter</th>
<th>Without BSA</th>
<th>With 2% BSA</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>HLMs</td>
<td>HKMs</td>
<td>HIMs</td>
</tr>
<tr>
<td>Metabolite formation</td>
<td>V_{max} (pmol/min per milligram protein)</td>
<td>1460 (244)</td>
<td>5220 (726)</td>
</tr>
<tr>
<td></td>
<td>K_{m} (µM)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>CL_{int,UGT} (ml/min per gram tissue)</td>
<td>107 (39.9)</td>
<td>91.0 (12.8)</td>
</tr>
<tr>
<td></td>
<td>CL_{int,P450} (ml/min per gram tissue)</td>
<td>0.594 (0.215)</td>
<td>0.735 (0.0247)</td>
</tr>
<tr>
<td></td>
<td>CL_{int,UGT} (ml/min per gram tissue)</td>
<td>2.71 (0.18)</td>
<td>1.17 (0.41)</td>
</tr>
<tr>
<td></td>
<td>CL_{int,P450} &amp; UGT (ml/min per gram tissue)</td>
<td>9.12 (2.03)</td>
<td>ND</td>
</tr>
</tbody>
</table>

BSA, bovine serum albumin; HIM, human intestinal microsomes; HKM, human kidney microsomes; HLM, human liver microsomes; (—), not applicable; ND, no depletion observed; NT, not tested; PBPK, physiologically based pharmacokinetic.

**Fig. 2.** Formation rate plots for propofol glucuronide as a function of substrate concentration in alamethicin activated human hepatic, renal, and intestinal microsomes in the presence and absence of 2% BSA. Data represent the mean of three experiments, each performed in duplicate. Error bars represent the standard deviation. (A) Data generated in the absence of BSA. (B) Data generated in the presence of BSA. △, ○, and □ represent propofol glucuronide formation in hepatic, renal, and intestinal microsomes, respectively.
presented in Supplemental Table 2). The predicted concentration-time profiles for all dose groups are shown in the Supplemental Figs. 4 and 5.

Use of the PBPK model resulted in underestimation of clearance regardless of the source of the in vitro data. Predicted clearance did not exceed 35% of the observed values and recovery of the blood concentration-time profiles was poor when using CL\text{int} data from all in vitro systems (Table 3). As expected from comparison of the in vitro data, no difference was observed in the predicted in vivo clearance from microsomal CL\text{int,UGT} data derived by either the substrate depletion or metabolite formation methods in the presence of BSA (Table 3). The lower in vitro CL\text{int} obtained in hepatocytes or microsomes in the absence of BSA reduced the clearance prediction accuracy even further. Therefore, the microsomal CL\text{int} data obtained using the substrate depletion approach (in the presence of BSA) were used for further analysis and optimization of the PBPK model; a representative predicted blood concentration-time profile obtained using these data is shown in Fig. 3. When these in vitro data were used in the PBPK model, renal glucuronidation clearance contributed 11% to the predicted systemic clearance, which is much lower than the approximately 30% contribution that has been reported in vivo to the predicted systemic clearance, which is much lower than the approximately 30% contribution that has been reported in vivo (Hiraoka et al., Takizawa et al., 2005).

With microsomal substrate depletion clearance data as inputs, the PBPK model was used to predict the extraction ratio due to metabolic clearance for the kidney and the liver at steady state. These data were compared with corresponding in vivo values detailed in a clinical clearance for the kidney and the liver at steady state. These data were compared with corresponding in vivo values detailed in a clinical clearance for the kidney and the liver at steady state. These data were compared with corresponding in vivo values detailed in a clinical study (Hiraoka et al., 2005). Using in vitro data derived in the presence of BSA, an underprediction of hepatic extraction was observed (predicted E\text{H} was 0.39, which represented 42% of the observed value). A more pronounced underprediction was found for E\text{R} with the predicted value (0.07) being only 10% of the observed value.

**Optimization of the Model for Prediction of Renal and Hepatic Metabolic Clearance.** Considering the underprediction observed, in vitro clearance parameters were optimized within the PBPK model by performing simultaneous fitting of the concentration-time data from intact and anhepatic patients (Table 4); in vitro data from microsomal substrate depletion assays in the presence of BSA were used as the initial estimates of CL\text{int}. Simultaneous optimization of in vitro hepatic and renal CL\text{int} in the PBPK model showed that renal glucuronidation clearance was underpredicted to a greater extent than hepatic clearance, resulting in an empirical scaling factor of 17 versus 9 required in the case of liver (Table 4). Coefficients of variation for the hepatic and renal CL\text{int} empirical scaling factors were 59 and 39%, respectively (Table 4). Use of the empirical scaling factors solely for in vitro kidney CL\text{int,UGT} data, in conjunction with the nonoptimized liver CL\text{int}, resulted in predicted clearance within 2-fold (59% on average) of observed values. Representative blood concentration-time profiles predicted using either nonoptimized in vitro data or optimized renal and hepatic clearance (in vitro hepatic and renal CL\text{int} data with the empirical scaling factors) are shown in Fig. 3. A high degree of correlation was observed for this dataset (R\textsuperscript{2} = 0.92, n = 94 data points) with 99% of the propofol blood concentrations predicted within 2-fold of the line of unity across the dose range (Fig. 4). Use of the optimized in vitro clearance data in the PBPK model predicted an overall E\text{R,UGT} value of 0.53, in agreement with in vivo estimates of approximately 0.6 (Favetta et al., 2002). In addition, predicted E\text{H} and E\text{R} at steady state based on the use of the optimized scalars for in vitro hepatic and renal CL\text{int} were within 20% of the observed values [91 and 82% of observed values, respectively; data from an additional independent study by Hiraoka et al. (2005)].

**Discussion**

This is the first study to assess the prediction accuracy of renal versus hepatic clearance from CL\text{int} data determined in different in vitro systems by applying a PBPK modeling approach. Propofol was used as an example drug and clinical data reported in intact and anhepatic patients were used to refine the predictions of clearance from in vitro data and to optimize the PBPK model for prediction of propofol blood concentration-time profiles.

**In Vitro Characterization of Propofol Metabolism in Microsomes.** No P450 clearance of propofol was detectable in HKMs, in agreement with studies reporting low P450 mRNA levels in the kidney (Nishimura and Naito, 2006; Bieche et al., 2007). The effect of albumin on CYP2B6 (the main enzyme for propofol P450 metabolism) has not been investigated to date; this enzyme is not reported to be involved in free fatty acid clearance and a marked impact of albumin is not anticipated, in agreement with the minimal increase (2-fold) in CL\text{int,P450} observed in HLMs upon inclusion of BSA. Similar to previous reports, propofol liver CL\text{int,P450} was greater than CL\text{int,UGT} (3-fold) in the absence of BSA with the extent of this difference being reduced upon inclusion of BSA (Al-Jahdari et al., 2006; Kilford et al., 2009).

This study shows that the trend of increased CL\text{int,UGT} upon inclusion of albumin is consistent in hepatic, renal, and intestinal microsomes; however, the extent differs between the tissues. Similarly to our findings, previous reports of the effect of albumin on UGT1A9 and 2B7 substrates in HLMs also showed a decrease in K\text{m} with minimal impact on V\text{max} (Rowland et al., 2007, 2008, 2009). This is considered to be due to the sequestration of inhibitory free fatty acids released during microsomal incubations, allowing estimation of the true K\text{m} (Rowland et al., 2007, 2008). Propofol kinetic parameters obtained in HLMs, HKMs, and HIMs were comparable with previously published data, where available (Supplemental Table 3 and Supplemental references). In the presence of BSA, HLM and HKM CL\text{int,UGT} values were similar from both metabolite formation and substrate depletion assays. In this study, a decrease in the propofol

<table>
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<tr>
<th>TABLE 3</th>
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<tr>
<td>Accuracy of propofol in vivo clearance predicted using CL\text{int,P450} and CL\text{int,UGT} data derived from different in vitro systems</td>
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<tr>
<td>Data are shown as mean (± S.D.) predicted CL/observed CL (%). Data for individual dose groups are detailed in Supplemental Table 2.</td>
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<td>In Vitro Systems Used</td>
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<tr>
<td>Substrate depletion</td>
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<tr>
<td>Metabolite formation</td>
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* Substrate depletion approach used to determine hepatocyte CL\text{int} data.
model that allows incorporation of metabolism in extrahepatic tissues is required to adequately assess the prediction of \( f_{\text{m,UGT}} \) from in vitro CLint data derived in both hepatic and extrahepatic microsomes. However, inclusion of such data in the PBPK model still resulted in an underprediction of \( f_{\text{m,UGT}} \) (0.39), with the predicted value being comparable with that derived from HLM data alone.

**Prediction of Propofol Renal Glucuronidation and Hepatic Metabolism Using In Vitro Clearance Data from Different Systems and Optimization Using In Vivo Data.** This study represents the first report where a PBPK model has been used to predict propofol renal metabolism from in vitro clearance data, in contrast to previous propofol PBPK modeling efforts that have mainly focused on accommodating potential lung sequestration (Levitt and Schnider, 2005; Upton and Ludbrook, 2005). Inclusion of BSA in the microsomal assays improved the prediction of propofol in vivo clearance using the PBPK model, consistent with the trend reported previously for UGT1A9 substrates using static IVIVE methods (Rowland et al., 2008; Kiford et al., 2009; Gill et al., 2012). However, the underprediction trend was still apparent regardless of whether hepatocyte or microsomal data in the presence of 2% BSA were used as an input for hepatic CLint: predicted clearance was <35% of the observed value. Therefore, clinical data from three dose levels in intact patients and data reported for subjects during the anhepatic phase of liver transplantation were used to optimize the in vitro clearance parameters. Use of clinical data in conjunction with the developed PBPK model allowed differentiation between underprediction of hepatic and renal CLint and highlighted more pronounced underestimation of propofol renal CLint,UGT in comparison with hepatic clearance (17-fold versus 9-fold). Prior to optimization, predicted extraction ratios were 0.07 and 0.42 for kidney and liver, compared with 0.43–0.87 and 0.76–0.98 estimated in vivo, respectively (Hiraoka et al., 2005; Takizawa et al., 2005a, 2005b). Following optimization of renal in vitro glucuronidation data, the predicted in vivo clearance was within 2-fold of the observed values and the corresponding extraction ratios and \( f_{\text{m,UGT}} \) values were in good agreement with the in vivo data (from independent studies than used for model development). The intestinal metabolism was not investigated in the PBPK model due to the anticipated low contribution of this tissue to its systemic clearance. This assumption is supported by UGT and P450 expression data for the intestine, showing low levels of UGT1A9 and CYP2B6 in comparison with the liver and kidney (Bièche et al., 2007; Court et al., 2012; Harbort et al., 2012). Due to the lack of suitable in vivo data, we could not assess the prediction accuracy for in vitro CLint,UGT data for the intestine or potential for propofol sequestration in either lungs or other tissues, although this cannot be ruled out.

Reported UGT mRNA data showed regional differences in the kidney (Gaganis et al., 2007; Lash et al., 2008). In addition, glucuronidation capacity has been found to differ between the sections

\[
V_{\text{max}} \text{ was observed at microsomal protein concentrations <0.1 mg/ml and in the presence of BSA, in agreement with a previous report (Walsky et al., 2012). Similar effects were observed at both 1 and 2% BSA, suggesting that the effect could not be simply rationalized by the differential concentration ratio of microsomal protein to BSA. Current findings are in contrast to an increase in } V_{\text{max}} \text{ reported for other UGT1A9 substrates in the presence of BSA and at low microsomal protein concentrations (Manevski et al., 2011, 2012). However, differences in assay conditions in the published studies prevent direct comparison with the findings herein; further investigation is required to confirm whether these effects on } V_{\text{max}} \text{ are directly associated with the use of BSA at very low protein concentrations. Current findings highlight the importance of careful selection of the assay conditions when determining glucuronidation kinetic parameters in the presence of BSA, particularly when using microsomal protein concentrations of <0.1 mg/ml. The highest in vitro } f_{\text{m,UGT}} \text{ estimate was obtained from in-house hepatocyte data (0.43), whereas the } f_{\text{m,UGT}} \text{ calculated from HLM in vitro data derived in the presence of BSA was <0.4. These in vitro estimates ignore the contribution of extrahepatic metabolism and therefore it was not surprising that the fraction was lower than the value of 0.6 reported in vivo (Favetta et al., 2002). A mechanistic approach to confirm whether these effects on } f_{\text{m,UGT}} \text{ require additional investigation.}
\]

**Fig. 3.** Predicted blood concentration-time profiles for propofol using optimized liver and kidney in vitro CLint data derived from microsomes in the presence of BSA. ○ represent mean ± S.D. observed blood concentration-time data for the 18 mg/kg dose level from Gepts et al. (1987). Black line represents predicted concentrations using in vitro CLint data from the substrate depletion microsomal assays (in the presence of BSA) without optimization; red line represents predicted concentrations using optimized kidney CLint,UGT data; and green line represents predicted concentrations using optimized CLint data for both liver and kidney.

\[
\text{TABLE 4}
\]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Initial Parameter Estimate</th>
<th>Model Fitted Parameter Estimate</th>
<th>Model Fitted Scalar</th>
<th>CV% for Fitted Scalar</th>
</tr>
</thead>
<tbody>
<tr>
<td>( K_p )</td>
<td>0.786</td>
<td>13.6</td>
<td>0.54</td>
<td>9.5</td>
</tr>
<tr>
<td>Renal CLint,UGT (l/h per gram tissue)</td>
<td>1.57</td>
<td>14.5</td>
<td>9.21</td>
<td>59</td>
</tr>
</tbody>
</table>

CV, coefficient of variation.

* Parameters estimated by simultaneous fitting of in vivo concentration-time data from intact and anhepatic patients, as detailed in the Materials and Methods.


* Initial CLint,UGT data taken from microsomal substrate depletion assays in the presence of BSA. Full details are presented in Table 2.

* Combined CLint,P450 and CLint,UGT data.
of the kidney in laboratory animals, with the highest activity observed in the proximal tubular cells of the cortex (Cojocel et al., 1983; Hjelle et al., 1986). However, data on regional differences in glucuronidation activity in human kidney tissue are very limited. Two studies have shown up to 3-fold higher glucuronidation clearance for naproxen and morphine (mainly cleared by UGT2B7) in cortical compared with medullary microsomes (Yue et al., 1988; Gaganis et al., 2007). In contrast, frusemide (primarily cleared by UGT1A9) glucuronidation clearance was comparable in microsomes from human kidney cortex and medulla (Kerdpin et al., 2008). Although these data may suggest that there are no regional differences in glucuronidation clearance via UGT1A9 in the kidney, these findings should be considered with caution because potential differences in microsomal recovery depending on the region of the kidney were not considered. The information on the proportion of the cortex and medulla used for the preparation of the microsomes employed in this study was not available; similarly, the region of the kidney used in the study reporting the kidney microsomal recovery (Al-Jahdari et al., 2006) is unknown, all of which may affect the glucuronidation activity and contribute to the particularly poor prediction of renal glucuronidation observed with the PBPK model prior to optimization of the in vitro data.

Although we have successfully developed a PBPK model for the prediction of propofol clearance and exposure from in vitro CLint data, we could not assess the use of our empirical scaling factors with other drugs. The availability of blood concentration-time data identifying drug metabolism across the kidney is extremely limited and hinders the assessment of this model with a wider range of compounds. In addition to the factors discussed above, poor prediction of renal glucuronidation may also suggest that well stirred assumptions, which are adequate for many tissues in PBPK modeling, are not appropriate for the kidney and more complex models may need to be considered. This will be particularly relevant for drugs that are also substrates of the renal transporters (Giacomini et al., 2010). Unlike the liver, the kidney is not a homogeneous tissue and consists of distinct regions with varying blood flows. However, in vivo and in vitro data required for development and validation of more complex kidney models, such as absolute abundance data for the UGTs and transporters in the various regions of the kidney, are currently lacking in the literature.

In conclusion, this study has shown consistent increases in propofol clearance estimates in hepatic, renal, and intestinal microsomes due to the “albumin effect” and highlighted the importance of careful selection of assay conditions when using BSA at very low microsomal protein concentrations. The analysis provides an example of the application of clinical data to refine the developed PBPK model and assess the predictive capacity of in vitro data for propofol renal and hepatic metabolic clearance in isolation. This study highlighted a more pronounced underprediction of renal glucuronidation than hepatic metabolism; further assessment of the contribution of extrahepatic microsomal mechanisms and the adequacy of currently available scaling factors and models for renal metabolism is required.

Acknowledgments

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Authorship Contributions

Participated in research design: Gill, Gertz, Houston, Galetin.
Conducted experiments: Gill.
Performed data analysis: Gill.
Wrote or contributed to the writing of the manuscript: Gill, Gertz, Houston, Galetin.

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Applying a PBPK Model to Predict Renal Metabolism


Giacomini KM, Huang SM and Rowland M (2012) The role of physiologically based pharmacokinetic modeling


