Physiologically Based Pharmacokinetic Models for Prediction of Complex CYP2C8 and OATP1B1 (SLCO1B1) Drug–Drug–Gene Interactions: A Modeling Network of Gemfibrozil, Repaglinide, Pioglitazone, Rifampicin, Clarithromycin and Itraconazole

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Abstract
Background Drug–drug interactions (DDIs) and drug–gene interactions (DGIs) pose a serious health risk that can be avoided by dose adaptation. These interactions are investigated in strictly controlled setups, quantifying the effect of one perpetrator drug or polymorphism at a time, but in real life patients frequently take more than two medications and are very heterogenous regarding their genetic background.

Objectives The first objective of this study was to provide whole-body physiologically based pharmacokinetic (PBPK) models of important cytochrome P450 (CYP) 2C8 perpetrator and victim drugs, built and evaluated for DDI and DGI studies. The second objective was to apply these models to describe complex interactions with more than two interacting partners.

Methods PBPK models of the CYP2C8 and organic-anion-transporting polypeptide (OATP) 1B1 perpetrator drug gemfibrozil (parent–metabolite model) and the CYP2C8 victim drugs repaglinide (also an OATP1B1 substrate) and pioglitazone were developed using a total of 103 clinical studies. For evaluation, these models were applied to predict 34 different DDI studies, establishing a CYP2C8 and OATP1B1 PBPK DDI modeling network.

Results The newly developed models show a good performance, accurately describing plasma concentration–time profiles, area under the plasma concentration–time curve (AUC) and maximum plasma concentration (Cmax) values, DDI studies as well as DGI studies. All 34 of the modeled DDI AUC ratios (AUC during DDI/AUC control) and DDI Cmax ratios (Cmax during DDI/Cmax control) are within twofold of the observed values.

Conclusions Whole-body PBPK models of gemfibrozil, repaglinide, and pioglitazone have been built and qualified for DDI and DGI prediction. PBPK modeling is applicable to investigate complex interactions between multiple drugs and genetic polymorphisms.

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1 Introduction

From epidemiological data, it is estimated that 5–20% of adverse drug events resulting in hospital admission are caused by drug–drug interactions (DDIs), with an especially high risk for elderly patients due to polypharmacy [1]. Indeed, data show that in the USA, 67% of the adults older than 62 years take more than five medications. As a result, about one in six older adults may be at risk for a major DDI [2] resulting in decreased efficacy, increased risk for adverse drug reactions, and increased healthcare costs. A second important aspect is that genetic polymorphisms in drug transporters or metabolizing enzymes may result in drug–gene interactions (DGIs). Similarly to DDIs, these DGIs can result in significantly altered drug exposure. In
Whole-body physiologically based pharmacokinetic (PBPK) models of gemfibrozil, repaglinide, and pioglitazone have been successfully built and evaluated for the prediction of drug–drug interactions (DDIs). The final models integrate state-of-the-art knowledge on the absorption, distribution, metabolism, and excretion of the three drugs with insights gained during the PBPK model development.

These models were applied to predict DDIs in a network of cytochrome P450 (CYP) 2C8 perpetrator and victim drugs and to describe and predict drug–gene interactions (DGIs) caused by genetic polymorphisms of CYP2C8 and solute carrier organic anion transporter family member (SLCO) 1B1 (organic-anion-transporting polypeptide [OATP] 1B1) Furthermore, the existence of physicochemical DDIs was proposed: coadministration of poorly soluble drugs such as itraconazole and pioglitazone might further decrease their solubility in the gut, leading to decreased absorption and lower drug exposure.

This study demonstrates the applicability of PBPK to investigate the DDI or DGI potential of drugs, predict complex interaction scenarios (e.g., drug–drug–drug–gene interactions), and develop potential dose adaptations for patients.

Key Points

- Whole-body physiologically based pharmacokinetic (PBPK) models of gemfibrozil, repaglinide, and pioglitazone have been successfully built and evaluated for the prediction of drug–drug interactions (DDIs).
- The final models integrate state-of-the-art knowledge on the absorption, distribution, metabolism, and excretion of the three drugs with insights gained during the PBPK model development.
- These models were applied to predict DDIs in a network of cytochrome P450 (CYP) 2C8 perpetrator and victim drugs and to describe and predict drug–gene interactions (DGIs) caused by genetic polymorphisms of CYP2C8 and solute carrier organic anion transporter family member (SLCO) 1B1 (organic-anion-transporting polypeptide [OATP] 1B1).
- The existence of physicochemical DDIs was proposed: coadministration of poorly soluble drugs such as itraconazole and pioglitazone might further decrease their solubility in the gut, leading to decreased absorption and lower drug exposure.

This study demonstrates the applicability of PBPK to investigate the DDI or DGI potential of drugs, predict complex interaction scenarios (e.g., drug–drug–drug–gene interactions), and develop potential dose adaptations for patients.

Current clinical practice, DDIs and DGIs are considered separate entities. However, they are interconnected and ignoring drug–drug–gene interactions (DDGIs) can jeopardize patient safety. Ideally, guidelines on how to manage DDIs and DDGIs should be based on results from clinical trials.

However, in reality, most DDGIs cannot be investigated in clinical trials for many reasons, including ethical and feasibility restrictions due to their complexity. Usually, classic DDI studies are performed as typical phase I studies in healthy volunteers using so-called index substances to characterize a certain DDI potential. The study participants are mostly young, healthy, take only two drugs at the same time, and are genetically homogenous, and, consequently, they do not mimic real-life multimorbid patients with polypharmacy and genetic polymorphisms [3]. Thus, there is a translational challenge to assess and manage complex multifactorial DDGIs in real-life patients.

One possibility to loosen this Gordian knot might be the application of whole-body physiologically based pharmacokinetic (PBPK) modeling. PBPK models are increasingly used to evaluate the effects of patient factors on drug exposure [4] and they are excellent tools to predict the DDGI potential of drugs in silico and allow development of alternative dosing regimens for patients. The interest in PBPK modeling is continuously rising in academia and the pharmaceutical industry. Regulatory agencies (European Medicines Agency [EMA], U.S. Food and Drug Administration [FDA]) recommend PBPK modeling for the assessment of DDI potential, the development of alternative dosing regimens, and, in some cases, even to waive clinical studies [5, 6]. To project the reality of patients, complex DDI networks and thoroughly developed PBPK models are required. Even though many perpetrator and victim drug models have been developed and published so far [7], there is still a need for further models and more comprehensive DDI networks.

The main focus of the presented work is the description of cytochrome P450 (CYP) 2C8- and organic-anion-transporting polypeptide (OATP) 1B1-based DDIs, using PBPK models of the perpetrator drug gemfibrozil (strong CYP2C8 index inhibitor and inhibitor of OATP1B1) and of the two victim drugs repaglinide (sensitive CYP2C8 index substrate and substrate of OATP1B1) and pioglitazone (moderate sensitive CYP2C8 substrate) [6, 8]. Gemfibrozil, repaglinide, and pioglitazone are all recommended by the FDA for use in clinical DDI studies [8]. As clinically relevant genetic polymorphisms of CYP2C8 and solute carrier organic anion transporter family member (SLCO) 1B1 (OATP1B1) are reported to impact the pharmacokinetics of repaglinide and pioglitazone, the effects of the CYP2C8*3 and SLCO1B1 521T>C alleles were considered and implemented into the respective models.

The aims of this study were (1) to develop a PBPK DDI network [9] for CYP2C8 and thereby to expand the library of publicly available models for DDI prediction with verified whole-body PBPK models of gemfibrozil, repaglinide, and pioglitazone, and to apply these models to (2) describe and predict DDIs including complex DDIs with more than two drugs, (3) describe and predict DGIs with two or more drugs, and (4) exemplarily develop victim drug dose adaptations for patients with genetic polymorphisms and coadministration of two perpetrator drugs. The supplementary document to this paper (Electronic Supplementary Material [ESM]) is compiled as a transparent and comprehensive documentation and reference manual, providing detailed information on all PBPK models and modeled DDI studies. Model files are freely available in the Open Systems Pharmacology (OSP) repository (https://www.open-systems-pharmacology.org).

2 Methods

2.1 Software

PBPK models were developed using PK-Sim® and MoBi® modeling software (version 7.3.0, part of the OSP suite). Parameter optimizations (Monte-Carlo-algorithm) and sensitivity analyses were performed with PK-Sim®. Clinical study
2.2 Physiologically Based Pharmacokinetic (PBPK) Model Building

PBPK model building was started with an extensive literature search to gain information on the physicochemical properties and absorption, distribution, metabolism, and excretion (ADME) processes of the drugs of interest as well as to obtain clinical studies (healthy individuals) of intravenous and oral administration in single and multiple doses. The plasma concentration–time profiles of the clinical studies were digitized and divided into an internal dataset used for model building and parameter optimization, and an external dataset used for model evaluation. Parameters that could not be informed from the literature were optimized, fitting the model simultaneously to the observed data of all studies assigned to the internal dataset.

The mean individuals used to simulate the different studies were modeled according to the respective study reports, with corresponding age, weight, height, sex, and ethnicity. If no information on these demographics was found, a 30-year-old male European was assumed, with the mean weight and height characteristics given in the PK-Sim® database. The creation of virtual populations to compare predicted and observed population plasma concentration–time profiles is described in the ESM (Sect. 3.9).

Protein expression of enzymes and transporters was implemented according to the literature, using the PK-Sim® database [10]. For more details, see ESM Table S3.9.1.

2.3 PBPK Model Evaluation

The PBPK models were evaluated by comparison of predicted population plasma concentration–time profiles to observed data. The observed data were mostly reported as arithmetic or geometric mean plasma concentration–time profiles with standard deviations. To compare the variability of predicted to observed profiles, 68% population prediction intervals were plotted, as this interval corresponds to the range of ± 1 standard deviation around the mean if normal distribution is assumed.

Furthermore, the predicted plasma concentration values were compared to their respective observed values in goodness-of-fit plots and the model performance was evaluated by comparison of predicted to observed area under the plasma concentration–time curve (AUC), maximum plasma concentration ($C_{\text{max}}$), apparent oral clearance ($\text{CL/F}$), and half-life values. For model evaluation, all AUC values were calculated from time zero to infinity ($\text{AUC}_\infty$).

As quantitative measures of the model performance, mean relative deviations (MRDs) of the predicted plasma concentrations (see ESM Eq. 3.1) and geometric mean fold errors (GMFEs) of the $\text{AUC}_\infty$, $C_{\text{max}}$, $\text{CL/F}$, and half-life values (see ESM Eq. 3.2) were calculated.

2.4 Drug–Drug Interaction (DDI) Network Modeling

In addition to the evaluation methods described in Sect. 2.3, a CYP2C8 DDI network was built to evaluate the DDI performance of the developed models (Fig. 1). Gemfibrozil with its metabolite gemfibrozil 1-O-β-glucuronide was used as the CYP2C8 and OATP1B1 inhibitor, repaglinide as the CYP2C8 and OATP1B1 victim, and pioglitazone as the CYP2C8 victim drug. In addition, repaglinide is also a substrate of CYP3A4 and OATP1B3, adding DDI potential during coadministration of CYP3A4 and OATP1B3 perpetrators, such as itraconazole, clarithromycin, and rifampicin. Rifampicin was used as the inducer and competitive inhibitor of CYP2C8, CYP3A4, OATP1B1, and OATP1B3, interacting with repaglinide and pioglitazone. Interaction processes are modeled using the equations given in the ESM (Sect. 1).

2.5 DDI Network Evaluation

The quality of the DDI modeling was evaluated by comparison of predicted to observed victim drug plasma concentration–time profiles when administered alone and during coadministration, DDI AUC ratios (Eq. 1) and DDI $C_{\text{max}}$ ratios (Eq. 2). For DDI evaluation, all AUC values were calculated from time zero to the time of the last concentration measurement ($\text{AUC}_{\text{last}}$).

$$\text{DDI AUC ratio} = \frac{\text{AUC}_{\text{last}} \text{ victim drug during perpetrator coadministration}}{\text{AUC}_{\text{last}} \text{ victim drug control}} \quad (1)$$

$$\text{DDI } C_{\text{max}} \text{ ratio} = \frac{C_{\text{max}} \text{ victim drug during perpetrator coadministration}}{C_{\text{max}} \text{ victim drug control}} \quad (2)$$

To assess the DDI modeling performance, the GMFEs of the predicted DDI AUC ratios and DDI $C_{\text{max}}$ ratios were calculated according to ESM Eq. 3.2.

2.6 Sensitivity Analysis

Sensitivity analyses were performed on the gemfibrozil, repaglinide, and pioglitazone models to investigate the impact of single model parameters on the predicted AUC at steady state given the highest recommended dose. Parameters were included into the analysis if they have been optimized, if

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they could have a strong influence on the pharmacokinetics due to their use in calculation of permeabilities or partition coefficients (e.g., fraction unbound), or if they had significant impact in formerly developed models (e.g., blood/plasma concentration ratio).

Sensitivity is calculated as the ratio of the relative change of the simulated AUC to the relative variation of the tested parameter around the parameter value used in the model, according to Eq. 3:

$$S = \frac{\Delta \text{AUC}}{\text{AUC}} \times \frac{p}{\Delta p}$$

where $S$ is the sensitivity of the AUC to the tested model parameter, $\Delta \text{AUC}$ is the change of the AUC, AUC is the AUC calculated with the original model parameter value, $p$ is the original model parameter value, and $\Delta p$ is the change of the tested model parameter value.

Sensitivity analyses were performed using a relative perturbation of 100%. The threshold value for sensitivity was set at 0.5. A sensitivity value of + 0.5 indicates that a 100% increase of the model parameter value causes a 50% increase of the predicted AUC. In addition, the parameters were varied within a 0.03- to 30-fold range and the resulting fold changes of AUC were investigated in spider plots.

Physiologically based pharmacokinetic models of itraconazole, rifampicin and clarithromycin were adopted from Hanke et al. [9]. Metabolism and transport are illustrated as black arrows. Solid red lines indicate reversible inhibition processes, dashed bold red lines indicate mechanism-based inactivation. Dashed violet lines indicate interaction processes by rifampicin consisting of inhibition as well as induction processes. The postulated physicochemical interactions are shown as dotted black lines.

Fig. 1 The developed drug–drug interaction (DDI) network with gemfibrozil and gemfibrozil 1-O-β-glucuronide as cytochrome P450 (CYP) 2C8 and organic-anion-transporting polypeptide (OATP) 1B1; itraconazole as CYP3A4, OATP1B1, and OATP1B3; rifampicin as CYP2C8, CYP3A4, OATP1B1, and OATP1B3; and clarithromycin as CYP3A4, OATP1B1, and OATP1B3 perpetrator drugs (upper part); and repaglinide as CYP2C8, CYP3A4, OATP1B1, and OATP1B3 and pioglitazone as CYP2C8 victim drugs (lower part).

3 Results

3.1 PBPK Model Building and Evaluation

Whole-body PBPK models of gemfibrozil with gemfibrozil 1-O-β-glucuronide (parent–metabolite model), repaglinide, and pioglitazone have been successfully developed. A total number of 103 studies showing plasma concentration–time profiles were used for model building and evaluation; all of these are presented in the ESM.

A detailed description of each model, including Tables listing the drug-dependent parameters, is given in ESM Sects. 3.3 (gemfibrozil and gemfibrozil 1-O-β-glucuronide model), 3.4 (repaglinide model), and 3.5 (pioglitazone model). System-dependent parameters were taken directly from the PK-Sim® database or, if not available, they were gathered from literature, as summarized in ESM Table S3.9.1. The good model performance for both internal and external datasets is demonstrated by comparison of population predicted to observed plasma concentration–time profiles in Fig. 2 (representative studies for each compound) and in ESM Figs. S3.3.1, S3.3.2, S3.3.3, S3.3.4, S3.4.1, S3.4.2, S3.4.3, S3.4.4, S3.5.1, and S3.5.2 (all studies, semilogarithmic and linear plots). Goodness-of-fit plots...
are presented in ESM Figs. S3.3.5, S3.4.5, and S3.5.3 and MRD values for all studies are given in ESM Tables S3.3.3, S3.4.3, and S3.5.3. Correlation of predicted to observed AUC and C\text{max} values is presented in ESM Figs. S3.3.6, S3.4.6, and S3.5.4, and the corresponding values are given in ESM Tables S3.3.4, S3.4.4, and S3.5.4, including calculated model GMFE values. Correlation of predicted to observed CL/F and half-life values is shown in ESM Figs. S3.3.7, S3.4.7, and S3.5.5 and the corresponding values are given in ESM Tables S3.3.5, S3.4.5, and S3.5.5. The results of sensitivity analyses are described in ESM Sects. 3.3, 3.4, and 3.5 and Figs. S3.3.8, S3.3.9, S3.3.10, S3.4.8, S3.4.9, S3.5.6, and S3.5.7.

For the development of the gemfibrozil parent–metabolite PBPK model, plasma concentration–time profiles of 23 studies (oral administration), including ten studies showing gemfibrozil 1-O-β-glucuronide concentrations (after oral administration of gemfibrozil) were used. Gemfibrozil 1-O-β-glucuronide was incorporated into the model because it is the major circulating metabolite of gemfibrozil and a mechanism-based inactivator of CYP2C8 [11], contributing significantly to the strong CYP2C8 inhibition by gemfibrozil. The gemfibrozil model applies an unspecified active uptake of gemfibrozil into hepatocytes [12], metabolism by uridine 5’-diphospho-glucuronosyltransferase (UGT) 2B7 to form gemfibrozil 1-O-β-glucuronide and glomerular filtration. The metabolite model applies a hepatic uptake transport by OATP1B1 [12], an efflux transport into bile via multidrug resistance-associated protein (MRP) 2 and glomerular filtration.

For the development of the repaglinide model, plasma concentration–time profiles of 56 studies (intravenous and oral administration) as well as the fraction metabolized via CYP2C8 information were used. The model applies hepatic uptake via OATP1B1 and OATP1B3, metabolism by CYP2C8 and CYP3A4, and glomerular filtration. An unbound Michaelis-Menten constant (K_M) value determined for repaglinide uptake into untransfected primary human hepatocytes was used as the K_M value for the OATP1B1 and OATP1B3 transport processes [13]. For the metabolism via CYP2C8, a fraction metabolized of 89% was assumed, according to a clinical DDI study with gemfibrozil [14]. The remaining repaglinide is metabolized via CYP3A4 [15]. The model slightly overpredicts the low repaglinide plasma concentrations at later times after dosing (ESM Figs. S3.4.1, S3.4.2, and S3.4.5). This could be due to a misspecification of the CYP2C8 or OATP1B1/1B3 K_M values taken from the literature, or to an unknown mechanism that impacts the pharmacokinetics of repaglinide but is missing in the model. This limitation of the model affects the prediction of low repaglinide plasma concentrations, but C\text{max} and AUC values are well-predicted (ESM Fig S3.4.6). For the investigation of DGIs, the model adequately describes the pharmacokinetics of repaglinide in carriers of the CYP2C8*3, SLCO1B1 521C and SLCO1B1 –11187A alleles. The implementation of polymorphic CYP2C8 and OATP1B1 is described in ESM Sects. 2.2, 2.3, and 3.4.

For the development of the pioglitazone model, plasma concentration–time profiles of 13 studies (oral administration), one study describing the fraction of pioglitazone excreted to urine, as well as the fraction metabolized via CYP2C8 information were used. Pioglitazone is reported to be predominantly metabolized by CYP2C8 [16], with no consistent information on the identity of other metabolic enzymes involved. Therefore, the model applies metabolism by CYP2C8, an unspecified hepatic clearance and glomerular filtration. For the metabolism via CYP2C8, a fraction metabolized of 70–75% was assumed [17]. For the investigation of DGIs, the model adequately describes the pharmacokinetics of pioglitazone in carriers of the CYP2C8*3 allele. The implementation of polymorphic CYP2C8 is described in ESM Sects. 2.2 and 3.5.

### 3.2 DDI Network Modeling

For the DDI network modeling, a total number of 34 DDI studies were available in literature and used to evaluate the modeled interactions (Fig. 1). Thereof, 23 studies describe the gemfibrozil–repaglinide DDI, four the gemfibrozil–pioglitazone DDI, and one study each the itraconazole–repaglinide DDI, the gemfibrozil–itraconazole–pioglitazone DDI, the gemfibrozil–itraconazole–repaglinide DDI, the geometric–itraconazole–pioglitazone DDI, the rifampicin–repaglinide DDI, the rifampicin–pioglitazone DDI, and the clarithromycin–repaglinide DDI. The previously developed models of itraconazole, rifampicin, and clarithromycin [9] were used without changes other than the addition of interaction parameters to model the CYP2C8 and OATP1B1/1B3 DDIs. A full description of the DDI modeling is given in ESM Sect. 4, including all interaction parameters, administration protocols, and study population demographics.

All DDI victim drug plasma concentration–time profiles are well-predicted using interaction parameters taken from the literature (listed in ESM Tables S3.3.2, S3.6.1, S3.7.1, and S3.8.1 summarizing the drug-dependent parameters of the perpetrator drugs), except for the hydroxy-itraconazole OATP1B1 and OATP1B3 inhibitory constant (K_i) values, which had to be optimized. To describe the DDIs with itraconazole, the solubility value of the previously developed itraconazole model was adjusted (see ESM Figs. S3.6.1 and S3.6.2). Itraconazole is a poorly soluble compound (8.0 mg/L in fasted state simulated intestinal fluid [18]), leading to variable absorption and, therefore, to large inter-individual differences in itraconazole plasma concentration–time profiles. In the gemfibrozil–itraconazole–pioglitazone interaction study [19], itraconazole concentration–time
profiles before and during gemfibrozil coadministration are reported, in addition to the DDI victim drug concentrations. For itraconazole and hydroxy-itraconazole, a reduction of the AUCs is described if gemfibrozil is added to the itraconazole–pioglitazone DDI. Possible explanations for these findings proposed by Jaakkola et al. [19] and Niemi et al. [20] are the displacement of itraconazole from plasma proteins or a reduction of itraconazole bioavailability by
gemfibrozil. Both hypotheses were tested with the models by either increasing the fraction unbound of itraconazole during the DDI or by reducing its solubility (to describe reduced bioavailability). The observed plasma concentration–time profiles of itraconazole and hydroxy-itraconazole could not be described properly by a change in fraction unbound, but the change of itraconazole solubility resulted in a good description of the observed data. To accurately model the observed itraconazole plasma concentrations of the gemfibrozil–itraconazole–pioglitazone interaction study [19], the solubility of itraconazole (capsule formulation, fasted state) was optimized to 14.5 mg/L at pH 6.5 in the absence of gemfibrozil and to 0.69 mg/L at pH 6.5 in the presence of gemfibrozil (shown in ESM Sect. 3.6), indicating a physicochemical DDI between gemfibrozil and itraconazole.

The second assumed physicochemical DDI affects pioglitazone during coadministration of both gemfibrozil and itraconazole (see Figs. 3c and ESM Fig. S4.7.1). Like itraconazole, pioglitazone is poorly soluble, with a reported solubility of 16.8 mg/L at pH 6.5 [21]. To accurately describe the pioglitazone plasma concentrations during

![Fig. 3 Pioglitazone plasma concentration–time profiles during the gemfibrozil–pioglitazone (a), itraconazole–pioglitazone (b), gemfibrozil–itraconazole–pioglitazone (c), and rifampicin–pioglitazone (d) drug–drug interaction (DDI). Observed data are shown as triangles ± standard deviation (dark blue: control, light blue: with perpetrator drug) [19, 38]. Pioglitazone population simulation arithmetic means are shown as lines (dark blue: control, light blue: with perpetrator drug), the dashed line (e) shows the prediction of the gemfibrozil–itraconazole–pioglitazone without pioglitazone solubility adjustment. The shaded areas represent the respective 68% population prediction intervals. Detailed information about dosing regimens and study populations is given in Electronic Supplementary Material (ESM) Tables S4.3.1, S4.5.1, S4.7.1, and S4.9.1. Predicted and observed DDI area under the plasma concentration–time curve (AUC) ratios and DDI maximum plasma concentration ($C_{\text{max}}$) ratios are compared in ESM Tables S4.3.2, S4.5.2, S4.7.2, and S4.9.2. b.i.d., twice daily, conc, concentration, po, oral, q.d., once daily, s.d., single dose.](https://example.com)
gemfibrozil–itraconazole–pioglitazone coadministration, pioglitazone solubility was adjusted to 1.59 mg/L at pH 6.5 (shown in ESM Sect. 4.7).

The good DDI performance of the models is demonstrated in predicted compared to observed victim drug plasma concentration–time profiles before and during
perpetrator drug coadministration in Figs. 3 and 4 (representative studies for each DDI) and in ESM Figs. S4.2.1, S4.2.2, S4.2.3, S4.2.4, S4.2.5, S4.2.6, S4.3.1, S4.3.2, S4.4.1, S4.5.1, S4.6.1, S4.7.1, S4.8.1, S4.9.1, and S4.10.1 (all studies, semilogarithmic and linear plots). For the gemfibrozil–repaglinide DDI, the time-dependency (Fig. 4 and ESM Figs. S4.2.3 and S4.2.4) and dose-dependency (ESM Figs. S4.2.5 and S4.2.6) of the interaction were also modeled and compared to clinical data.

As further evaluation of the performance of the entire DDI network, the correlation of predicted to observed DDI AUC ratios and DDI $C_{\text{max}}$ ratios of all modeled DDI studies is shown in Fig. 5. The corresponding values are listed in ESM Tables S4.2.2, S4.3.2, S4.4.2, S4.5.2, S4.6.2, S4.7.2, S4.8.2, S4.9.2, and S4.10.2, including GMFE values calculated for each perpetrator–victim drug combination.

### 3.3 Dose Adaptation Considerations

To show the utility of the models to individualize and improve drug therapy, dose adaptations for different DGI scenarios were calculated. Exemplarily, victim drug plasma concentration–time profiles in subjects with polymorphisms, and with polymorphisms during perpetrator drug coadministration, were simulated and compared to those of a CYP2C8 and SLCO1B1 wild-type individual without DDI (control). Then, potential dose adaptations were calculated, aiming to match control exposure (i.e., AUC in steady state). Figure 6 shows the different simulated scenarios, the extrapolated iso-exposure doses (% of control), simulations of plasma concentration–time profiles with the same dose for all individuals, and simulations of plasma concentration–time profiles with the extrapolated adjusted doses.

As expected, the worst-case scenario is the repaglinide administration to a patient carrying the CYP2C8 wild-type sequence and simultaneously homozygously the SLCO1B1 521C allele who is co-medicated with gemfibrozil and itraconazole, leading to a 49-fold increase in the repaglinide steady state AUC. According to the simulations, a repaglinide dose reduction by nearly 98% would be necessary to produce the same drug exposure in this individual as in the control person given the normal dose. A dose adjustment to simultaneously match both AUC and $C_{\text{max}}$ values to the control profiles was not possible, as the drug half-life is changed due to the polymorphisms and the DDIs. The safety and efficacy of increased or decreased $C_{\text{max}}$ values or the drug half-life following dose adaptations that were calculated to match the AUC in steady state cannot be predicted from the models.

### 4 Discussion

In this study, whole-body PBPK models of gemfibrozil (parent–metabolite model of gemfibrozil and gemfibrozil 1-O-β-glucuronide), repaglinide, and pioglitazone for the investigation and prediction of DDIs and DGIs have been successfully built and evaluated. All models reliably describe and predict plasma concentration–time profiles over a broad dosing range and for single- and multiple-dose administration. Their good performance has been demonstrated by (1) comparison of predicted to observed plasma concentration–time profiles; (2) goodness-of-fit plots; (3) the calculation of MRD values; (4) the comparison of predicted to observed AUC and $C_{\text{max}}$ values; (5) the calculation of model GMFEs; and (6) their good predictions within a CYP2C8 PBPK DDI modeling network.

The itraconazole, rifampicin, and clarithromycin PBPK models applied in this study have been evaluated in a previously described CYP3A4 DDI network [9, 22]. This network has now been expanded with the presented gemfibrozil, repaglinide, and pioglitazone models and their interactions via CYP2C8, OATP1B1, OATP1B3, and CYP3A4. To use the previously established itraconazole, rifampicin, and clarithromycin models for CYP2C8 and OATP1B1/B3 DDI modeling, interaction constants describing their induction and inhibition of CYP2C8, OATP1B1, and OATP1B3 have been added, but otherwise no drug- or system-dependent parameters have been changed. To model the induction of CYP2C8 by rifampicin, the maximum effect value ($E_{\text{max}}$ CYP2C8 = 3.2, ESM Table S3.7.1) was taken from literature [23]. Concerning the induction of OATP1B1 and OATP1B3 by rifampicin, only a modest and variable induction was found in primary human hepatocytes [24, 25]. The $E_{\text{max}}$ OATP1B1 = 0.38 (± 1.38-fold) identified during the rifampicin model development [9] was assumed to be adequate to model the induction of both OATP1B1 and OATP1B3. Except for the hydroxy-itraconazole OATP1B1 and OATP1B3 $K_i$ values (optimized, see ESM Sect. 4.4), all interaction parameter values were taken from previous experimental or modeling studies.
During the modeling of the complex DDIs with simultaneous administration of the two perpetrator drugs gemfibrozil and itraconazole, the clinical data show effects that cannot be explained with the expected CYP enzyme or drug transporter inhibition [19]. First, a reduction of itraconazole and hydroxy-itraconazole AUCs was described.

**Fig. 5** Correlation of predicted and observed drug–drug interaction (DDI) area under the plasma concentration–time curve (AUC) ratios and DDI maximum plasma concentration (C_{max}) ratios of all studies. The upper panel illustrates DDI AUC ratios (a) and DDI C_{max} ratios (b) of the gemfibrozil–repaglinide, itraconazole–repaglinide, gemfibrozil–itraconazole–repaglinide, rifampicin–repaglinide, or clarithromycin–repaglinide DDIs. The lower panel illustrates DDI AUC ratios (c) and DDI C_{max} ratios (d) of the gemfibrozil–pioglitazone, itraconazole–pioglitazone, gemfibrozil–itraconazole–pioglitazone, or rifampicin–pioglitazone DDIs. The colors represent different perpetrator drugs and the symbols the victim drugs repaglinide (dots) and pioglitazone (triangles). The straight black line marks the line of identity. Light grey lines indicate 0.8- to 1.25-fold and dark grey lines indicate 0.5- to 2-fold acceptance limits. The curved black lines show the prediction success limits suggested by Guest et al. [41]. Detailed information about dosing regimens and study populations is given in Electronic Supplementary Material (ESM) Tables S4.2.1, S4.3.1, S4.4.1, S4.5.1, S4.6.1, S4.7.1, S4.8.1, S4.9.1, and S4.10.1. The plotted DDI AUC ratios and C_{max} ratios are listed in ESM Tables S4.2.2, S4.3.2, S4.4.2, S4.5.2, S4.6.2, S4.7.2, S4.8.2, S4.9.2, and S4.10.2.
A PBPK Modeling Network of CYP2C8 Perpetrator and Victim Drugs

During coadministration of itraconazole and pioglitazone if gemfibrozil is added to this regimen as the third drug [19]. As itraconazole and hydroxy-itraconazole are both solely metabolized by CYP3A4 and gemfibrozil is not reported to affect CYP3A4, we propose the possibility of a physicochemical DDI between gemfibrozil and itraconazole. Simultaneous administration of gemfibrozil might further decrease the poor solubility of itraconazole, causing a decrease in absorption and thereby a decrease of the plasma concentrations of itraconazole and hydroxy-itraconazole. To test this hypothesis with the PBPK models, a reduction of itraconazole solubility during gemfibrozil coadministration was performed (from 14.5 to 0.69 mg/L), leading to a good description of the itraconazole exposure of this study (ESM Figs. S3.6.1 and S3.6.2). The phenomenon of decreased itraconazole plasma concentrations during co-treatment with gemfibrozil has been discussed by Jaakkola et al. [19] and Niemi et al. [20]. They proposed a displacement of itraconazole from plasma proteins by gemfibrozil as a possible explanation, which was also tested with our models but did not describe the observed effects, or an effect of gemfibrozil on itraconazole bioavailability, which is in full agreement with our hypothesis of reduced itraconazole solubility. This example demonstrates the great value of perpetrator drug plasma concentration assessment in clinical DDI studies.

Furthermore, a reduction of the pioglitazone AUC was observed during coadministration of gemfibrozil and pioglitazone if itraconazole was added to this regimen as the third drug [19]. As itraconazole alone has no effect on the pioglitazone pharmacokinetics (ESM Fig. S4.5.1), the effect of the combination of gemfibrozil and itraconazole on the pioglitazone plasma concentrations is expected to be similar to the effect of gemfibrozil alone (2.5-fold increase of AUC, no change of $C_{\text{max}}$). Surprisingly, during the combination of all three drugs, the pioglitazone AUC is increased only 2.2-fold and the $C_{\text{max}}$ is reduced to 75% of the $C_{\text{max}}$ in control conditions. To explain the considerable decrease of the pioglitazone $C_{\text{max}}$, we again postulate a physicochemical DDI decreasing the solubility of pioglitazone and thereby its absorption and plasma concentrations. To test this hypothesis applying the PBPK models, a reduction of pioglitazone solubility during gemfibrozil plus itraconazole coadministration was performed (from 16.8 to 1.59 mg/L), leading to a good description of the pioglitazone exposure during this complex DDI (ESM Fig. S4.7.1). These two examples illustrate that PBPK modeling is a valuable tool to develop and test hypotheses for unexpected clinical findings. Furthermore, they certainly raised our awareness of the possibility of solubility interactions in the gut, given the great number of poorly soluble drugs. These solubility interactions could be mediated either by the coadministered drugs themselves or by coadministered solubilizing agents that are used in the marketed formulations of Biopharmaceutics Classification System (BCS) class II drugs. However, these hypotheses need to be further investigated experimentally to verify or reject...
them. Additionally, inhibition of transporters or enzymes by excipients has been described in literature [26]. A change in bioavailability due to such an interaction cannot be excluded.

In addition to prediction of DDIs, the presented PBPK models of repaglinide and pioglitazone adequately describe the effects of the CYP2C8*3 and SLC01B1 521C alleles on repaglinide pharmacokinetics and the effect of the CYP2C8*3 allele on pioglitazone pharmacokinetics. Considering the 13% frequency of the CYP2C8*3 allele [27] and 14.3% frequency of the SLC01B1 521T>C SNP in Caucasians [28], these polymorphisms are clinically relevant. With our models, a dose adjustment in case of complex DDGs and drug–drug–drug–gene interactions was exemplarily performed to show how PBPK modeling could support drug therapy and labeling for complex scenarios.

5 Conclusion

In summary, comprehensive whole-body PBPK models of gemfibrozil (parent–metabolite model), repaglinide, and pioglitazone have been carefully built and evaluated in a CYP2C8 DDI network. The network described in this study represents an extension of a previously developed network of CYP3A4-interacting drugs [9], and the new models were challenged and verified in DDI predictions with different perpetrator and victim drugs, causing different kinds of interactions such as competitive inhibition, mechanism-based inactivation, and induction. Furthermore, the presented models are able to describe DGIs of repaglinide and pioglitazone, complex DDIs during coadministration of more than two interacting drugs, and were used to postulate DDIs on a physicochemical level. All models are transparently documented and model files are available in the OSP repository. They can be applied to investigate the DDI or DGI potential of drugs, inform the design of clinical trials, or simulate complex interactions (e.g., drug–drug–drug–gene interactions).

Compliance with Ethical Standards

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Conflict of Interest Sebastian Frechen, Thomas Eissing, and Thomas Wendt are employees and potential shareholders of Bayer AG. No potential conflicts of interest were disclosed by the other authors (Denise Türk, Nina Hanke, Sarah Wolf, Matthias Schwab, and Thorsten Lehr).

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