Application of physiologically-based pharmacokinetic/pharmacodynamic models to evaluate the interaction between nifedipine and apatinib

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Aim: It has been found that the co-administration of nifedipine with apatinib could cause exposure changes of nifedipine in vivo. But, whether this pharmacokinetic drug-drug interaction (DDI) between nifedipine and apatinib could enhance the antihypertensive effect of nifedipine, causing severe changes of blood pressure was unknown. Therefore, the aim of the present study was to conduct the pharmacokinetic/pharmacodynamic (PK/PD) modelling to evaluate the effect of pharmacokinetic changes on the antihypertensive effect of nifedipine. Thus, the results could guide the co-administration of these two drugs in clinic.

Methods: A physiologically-based pharmacokinetic (PBPK) model was first developed for nifedipine. The pharmacokinetic DDI between nifedipine and apatinib was evaluated. Then the verified PBPK models were linked to a PD model for investigating whether the exposure changes of nifedipine could cause severe changes in blood pressure. Furthermore, the changes in blood pressure caused by combination with apatinib were also assessed in patients with hepatic impairment via the PBPK/PD models.

Results: The predicted area under plasma concentration-time profile (AUC), maximum concentration (Cmax), area under effect-time profile (AUE), and maximum reduction in systolic blood pressure (Rmax) are all within 0.5–2.0-fold of the observed data, indicating that the PBPK/PD models for nifedipine are successfully established. The increases of predicted AUC and Cmax of nifedipine in the presence of apatinib are 1.73 and 1.41-fold, respectively. Co-administration of nifedipine with apatinib could cause exposure changes of nifedipine in vivo. However, the predicted AUE and Rmax changes of nifedipine in the presence to the absence of apatinib in cancer patients as well as in patients with hepatic impairment are all within 1.25-fold. The results indicate that the exposure changes of nifedipine caused by combination of apatinib has little effect on the changes of systolic blood pressure both in cancer patients and patients with hepatic impairment.
Conclusion: The pharmacokinetic changes of nifedipine caused by co-administration with apatinib has little impact on the antihypertensive effect of nifedipine. Apatinib is unlikely to cause severe pharmacodynamic DDI via inhibition of CYP3A4. It is suggested that nifedipine could be used in combination with apatinib without dose adjustment in clinic.

KEYWORDS
pharmacodynamics, drug-drug interaction (DDI), physiologically-based pharmacokinetics (PBPK), nifedipine, apatinib

Introduction

Apatinib, a small-molecule tyrosine kinase inhibitor, suppresses the tumor angiogenesis by selectively inhibiting the activity of vascular epidermal growth factor receptor (VEGFR) tyrosine kinases (Tian et al., 2011; Qiu et al., 2018; Meng et al., 2020). Apatinib was approved by the National Medical Products Administration (NMPA) for patients suffering from advanced gastric adenocarcinoma or gastric esophageal junction cancer in 2014 (Roviello et al., 2016; Geng et al., 2018). In 2020, the new indications for apatinib were approved for the treatment of advanced hepatocellular carcinoma. It has been reported that apatinib showed certain benefit in the treatment of other malignant tumors, such as non-small cell lung cancer, breast cancer, and colorectal cancer (Zhang, 2015; Xue et al., 2018; Maroufi et al., 2020). Although the clinical efficacy of apatinib is agreeable, further research on the potential drug–drug interactions (DDIs) is yet needed since cancer patients usually receive combination therapy, many of which are enzymes inhibitors or inducers. It is of great significance to evaluate the potential DDI risks associated with apatinib for its reasonable administration in clinic.

The in vitro metabolism studies showed that apatinib was mainly metabolized via CYP3A4/5, and CYP2D6 (Ding et al., 2013). Research has shown that the area under plasma concentration-time profile (AUC) of apatinib was significantly affected by co-administration with itraconazole (AUC0–48h increased by 75%) or rifampin (AUC0–1 decreased by 83%) in humans (Liu et al., 2018). In our previous study, the DDI simulation showed 2 to 4-fold changes in apatinib exposures by moderate CYP3A4 inhibitors and CYP3A4 inducers (Liu et al., 2021). A moderate increase of apatinib exposure (1.25 to 2-fold) was found with strong CYP2D6 inhibitor (Liu et al., 2021). However, the detailed evaluation of apatinib as a perpetrator in CYP450-based DDIs is lacking. Apatinib was reported to exert potent inhibition on CYP3A4 and CYP2C9 with the IC50 values of 1.80, 0.83, and 0.44 μM for midazolam hydroxylation, testosterone hydroxylation, and tolbutamide hydroxylation, respectively (Zhu et al., 2020). Thus, apatinib might affect the plasma exposures of CYP3A4 and CYP2C9 substrates when using combination therapy causing clinical efficacy and safety issues.

Many cancer patients were also treated with antihypertensive agents concomitantly. The combination of calcium channel blockers (CCB) was unavoidable, among which nifedipine was one of the most commonly used due to the rapid onset of action without central nervous system depression (Haddad, 1996; Ndanusa et al., 1997). Besides, hypertension is a commonly reported adverse event for patients receiving apatinib and, therefore, co-administration of nifedipine with apatinib was common in clinical practice (Peng et al., 2018; Shao et al., 2020). The in vitro studies indicated that nifedipine was a typical substrate of CYP3A4 which was almost completely metabolized by CYP3A4 (Guengerich et al., 1986). The research conducted by Zhu et al. (2020) has confirmed that co-administration of nifedipine with apatinib significantly increased the AUC0–48h and Cmax of nifedipine by 83% and 64%, respectively. Co-administration of apatinib with nifedipine could cause exposure changes of nifedipine in vivo. But, whether this pharmacokinetic DDI might cause significant clinical effects, in other words, whether enhancing the antihypertensive effect of nifedipine leading to a risk of hypotension is still unknown (Welch and Todd, 1990). The dosing regimen of nifedipine when co-administered with apatinib might be of vital importance. That is to say, when apatinib and nifedipine are co-administered, should the nifedipine be administered in regular dose in clinic, or the dose needs to be reduced. If the dose of nifedipine needs to be reduced, what is the appropriate dose?

In the present study, physiologically-based pharmacokinetic (PBPK) modelling was applied to evaluate the impact of apatinib on the exposure of nifedipine. Meanwhile, the developed PBPK models were linked to a pharmacodynamic (PD) model to investigate whether the exposure changes of nifedipine could cause severe changes in blood pressure (Chetty et al., 2014). Furthermore, the DDIs between nifedipine and apatinib were also assessed via the PBPK/PD models to determine the reasonable combination regimen in patients with hepatic impairment. The results of the present study could clarify the potential DDI risks via inhibition on CYP3A4 by apatinib, providing data basis for guiding the reasonable application in clinic.
Materials and methods

Development and verification of nifedipine physiologically-based pharmacokinetic model

The PBPK model for nifedipine was built-in with Simcyp™ (version 16, Simcyp Limited, Sheffield, United Kingdom), a commercially available PBPK software. Nifedipine has low solubility and high permeability, which belongs to the Biopharmaceutics Classification System Class II drug (Nader et al., 2016). CYP3A4 metabolism is the predominantly eliminated pathway (Guengerich et al., 1986; Nader et al., 2016). The input parameters for nifedipine are summarized in Table 1. The physicochemical parameters such as molecular weight, oil/water partition coefficient (log P), dissociation equilibrium constants (pKa), plasma unbound fraction (fu), and blood to plasma partition coefficient (B/P) were obtained from the nifedipine compound file in Simcyp. The in vivo absorption of nifedipine controlled release (CR) tablet was described using the advanced dissolution, absorption, and metabolism (ADAM) model. The effective permeability (P_{eff}) of nifedipine in human was calculated using the permeability data of MDCK II (Polli et al., 2001). The in vivo release of nifedipine CR tablet was described by Weibull function of dissolution profiles (Doki et al., 2017). The first-order model was used to describe the in vivo absorption of nifedipine immediate release (IR) tablet. A minimal PBPK distribution model and enzyme kinetics with recombinant CYP enzyme data were used to characterize the elimination of nifedipine in the Simcyp.

The developed PBPK models for nifedipine were verified with reported clinical study on a single-dose of 60 mg nifedipine CR tablet or 20 mg nifedipine IR tablet in healthy volunteers (Holtbecker et al., 1996; Toal et al., 2012). A Sim-Healthy volunteer population with population size of 100 in Simcyp at the age of 26–65 years old were used for the verification. The proportion of female was 0.5. The predicted results were compared with the clinical observations, and the predicted accuracy was measured by calculating the fold error of the pharmacokinetic parameters (C_{max}, T_{max}, and AUC), as described in Eq. 1 (Zhang et al., 2018; Li et al., 2019; Li et al., 2021).

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\text{fold error} = \frac{\text{simulated}}{\text{observed}}
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Development of nifedipine-apatinib drug-drug interaction model

The PBPK models for apatinib were already developed and verified in the previous study (Liu et al., 2021). The predicted

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**Table 1** Summary of input data for nifedipine in Simcyp™

| Parameters                        | Value  | Source                  |
|-----------------------------------|--------|-------------------------|
| Physiochemical parameters         |        |                         |
| Molecular weight (g/mol)          | 346.3  | Simcyp built-in data    |
| log P                             | 2.69   |                         |
| Compound type                      |        | Monoprotic base         |
| pKa                               | 2.82   |                         |
| B/P                               | 0.685  |                         |
| fu                                | 0.039  |                         |
| Absorption parameters             |        |                         |
| First-order model for IR tablet   |        |                         |
| f_a                               | 1      | Simcyp built-in data    |
| k_a (1/h)                         | 4.6    |                         |
| ADAM model for CR tablet          |        |                         |
| MDCK II permeability (10^{-6} cm/s)| 61    | Polli et al. (2001)    |
| P_{eff} (10^{-4} cm/s)            | 10.5   | Simcyp calculated       |
| Disposition parameters            |        |                         |
| Minimal PBPK model                | 0.57   | Simcyp calculated       |
| V_s (L/kg)                        |        |                         |
| Elimination parameters            |        |                         |
| Enzyme                            | rCYP3A4| Simcyp built-in data    |
| V_{max} (pmol/min/pmol CYP)       | 22     |                         |
| K_{m} (μM)                        | 10.95  |                         |
| Enzyme                            | rCYP3A5| Simcyp built-in data    |
| V_{max} (pmol/min/pmol CYP)       | 3.5    |                         |
| K_{m} (μM)                        | 31.9   |                         |
mean plasma concentration-time curves of apatinib in the presence or absence of itraconazole and rifampin were matched well with the clinical observed ones. Summary of the input data are listed in Table 2.

Apatinib was reported to exert potent inhibition on CYP3A4 in a competitive way (Zhu et al., 2020). In the DDI models between nifedipine and apatinib, the Ki was set to 0.12, with the fD values at 0.65 (Bao et al., 2018; Zhu et al., 2020). The PK profiles of nifedipine in the presence of apatinib were predicted and compared with the clinical observations. The subject characteristics as well as trial design were consistent with the reported single-dose DDI studies (Zhu et al., 2020). A SimChinese volunteer population with population size of 24 in Simcyp at the age of 26–65 years old were used for the verification. The proportion of female was 0.5. Subjects either received a 30 mg of oral nifedipine CR tablet on day 1 or treated with 750 mg of oral apatinib once daily for eight consecutive days with concomitantly administration of 30 mg nifedipine CR tablet on day 6. The simulated results were compared with the observations (Zhu et al., 2020). The change of nifedipine exposure was determined by AUC ratio and Cmax ratio, as described in Eq. 2 (Zhang et al., 2018; Li et al., 2019).

### Development and verification of pharmacodynamic model for nifedipine

Studies have shown that the control of systolic blood pressure (SBP) is more important than diastolic blood pressure (DBP) in most patients with hypertension (Donnelly et al., 1994; Meredith et al., 1994). Hence, SBP was employed in the PD modelling. An ordinary Emax model (Eq. 3) was used to describe the relationship between nifedipine concentration and the change of SBP (Abernethy et al., 1990; Donnelly et al., 1993). The Emax and EC50 were obtained from literature, which were set at −30 mmHg and 12.12 ng/ml, respectively (Edgar et al., 1987; Donnelly et al., 1994; Levine et al., 2003). The developed PD model was then linked to the PBPK model to investigate the effect of

### TABLE 2 PBPK model parameters for apatinib (Liu et al., 2021).

| Parameters                 | Value                  | Unit         |
|----------------------------|------------------------|--------------|
| Molecular weight           | 397.48                 | g/mol        |
| log P                      | 3.14                   |              |
| Compound type              | Diprotic base          |              |
| pKa                        | pKa1 = 6.60 pKa2 = 5.31|              |
| B/P                        | 0.995                  |              |
| fD                         | 0.076                  |              |
| ADAM model Caco-2 permeability | 6.81 x 10^{-6}    | cm/s         |
| Peff                       | 8.00 x 10^{-4}         | cm/s         |
| Kp scalar                  | 0.7                    |              |
| Vss (L/kg)                 | 2.684                  |              |
| CYP2D6 Vmax                | 9.82                   | pmol/min/mg protein |
| Km                         | 1.41                   | μM           |
| CYP3A4 Vmax                | 39.1                   | pmol/min/mg protein |
| Km                         | 2.18                   | μM           |
| CYP3A5 Vmax                | 3.28                   | pmol/min/mg protein |
| Km                         | 1.93                   | μM           |
pharmacokinetic DDI between nifedipine and apatinib on the change of SBP. The predictive accuracy was measured by calculating the maximum reduction in SBP ($R_{\text{max}}$) and the area under the effect-time curve (AUE) (Levine et al., 2003; Mourad, 2008; Toal et al., 2012).

Application of the nifedipine physiologically-based pharmacokinetic/pharmacodynamic model

The developed PBPK/PD model was used to predict the changes in SBP caused by the exposure changes of nifedipine with the combination of apatinib. The virtual populations in Simcyp with sizes of 100 at the age of 26–65 years old were used in the simulations. The proportion of female was 0.5. Two clinical scenarios were simulated. The first scenario was designed to determine whether co-administration of apatinib with regular dose level of nifedipine could cause severe changes of SBP in cancer patients. The second scenario was designed to determine if dose adjustment of nifedipine was necessary in the combination of apatinib in patients with hepatic impairment.

Results

Verification of the physiologically-based pharmacokinetic model for nifedipine

The simulated and the observed mean plasma concentration-time curves of nifedipine CR tablet at 60 mg or IR tablet at 20 mg dose levels in healthy volunteers are depicted in Figure 1. Table 3 shows the comparison of the model predicted pharmacokinetic parameters ($C_{\text{max}}$, $T_{\text{max}}$, and AUC) to the observed data, and the calculated fold errors of $C_{\text{max}}$, $T_{\text{max}}$, and AUC of nifedipine. The predicted mean plasma concentration-time curves of nifedipine were generally consistent with the clinical observed ones. Additionally, as shown in Table 3, the $C_{\text{max}}$ and AUC values were successfully predicted within 2-fold of observed values. The results indicate a good prediction of the nifedipine PBPK model.

Evaluation of the drug–drug interactions prediction for nifedipine and apatinib

The predicted and the observed mean plasma concentration-time curves of nifedipine with apatinib in cancer patients are shown in Figure 2. Table 4 shows calculated AUC ratio as well as $C_{\text{max}}$ ratio in the presence to the absence of apatinib. The successful simulation of the pharmacokinetic DDIs between nifedipine and apatinib is obtained by the established PBPK model. The increases of the predicted AUC and $C_{\text{max}}$ of nifedipine in the presence of apatinib are 1.73 and 1.41, respectively.

Verification of pharmacodynamic model for nifedipine

When the $E_{\text{max}}$ was set to −30 mmHg, the PD model fitted best. The comparison of the predicted SBP changes to the observations for nifedipine CR tablet at 60 mg or IR tablet at 10 mg dose levels is presented in Figure 3. Results show that the PD models for nifedipine CR tablet and IR tablet are successfully developed in predicting the clinical data. The predicted and observed values of $R_{\text{max}}$ and AUE ratios are listed in Table 5. The ratios are all between 0.5 and 2-fold. The results indicate the good predictive performance of the current nifedipine PD model.
Application of the physiologically-based pharmacokinetic/pharmacodynamic model for dosing recommendations

The developed PBPK/PD model was used to predict SBP changes caused by the exposure changes of nifedipine with the combination of apatinib in cancer patients. The model-predicted SBP changes for nifedipine CR tablet at 60 mg or 30 mg dose levels and IR tablet at 30 mg dose levels with or without co-administration of apatinib are presented in Figure 4. The calculated Rmax and AUE ratio in the presence to the absence of apatinib in cancer patients are listed in Table 6. Table 7 summarizes the changes of SBP in hepatic impairments when combination of nifedipine with apatinib. The predicted Rmax and AUE changes of nifedipine in the presence to the absence of apatinib in cancer patients and in hepatic impairments are all within 1.25-fold. Results show that the exposure changes of nifedipine caused by co-administration of apatinib has little effect on the SBP, indicating that apatinib is unlikely to cause severe pharmacodynamic DDI via inhibition of CYP3A4. Thus, nifedipine could be used in combination with apatinib without dose adjustment in clinic.

Discussion

Many cancer patients are also treated with antihypertensive agents concomitantly. Thus, the combination of CCB is unavoidable. Nifedipine is a commonly used antihypertensive drug in clinical practice and is also a sensitive substrate of CYP3A4. Besides, hypertension is a commonly reported adverse event for patients who receive apatinib, and nifedipine is most frequently used for drug-induced hypertension. Clinical study has revealed that the occurrence rate of hypertension as the adverse event of apatinib reached 73.9%, which necessitated the...
Figure 3
Predicted SBP changes and clinical observations in hypertensive patients at a single dose of 60 mg nifedipine CR tablet (A), and 10 mg nifedipine IR tablet (B).

Table 5 Comparison of the predicted $R_{\text{max}}$ and AUE to the observations for nifedipine CR tablet and IR tablet.

| Dose level            | Parameters | Unit    | Predicted | Observed | Fold error |
|-----------------------|------------|---------|-----------|----------|------------|
| 60 mg nifedipine CR tablet | $R_{\text{max}}$ | mmHg    | -22.21    | -23.09   | 0.96       |
|                       | AUE        | mmHg·h  | 433.98    | 440.10   | 0.99       |
| 10 mg nifedipine IR tablet   | $R_{\text{max}}$ | mmHg    | -30.13    | -33.20   | 0.91       |
|                       | AUE        | mmHg·h  | 144.84    | 102.00   | 1.42       |
concurrent use of antihypertensive drugs (Zhu et al., 2020). Apatinib was reported to be a potent inhibitor of CYP3A4, with IC50 values of 1.80 and 0.83, and the corresponding Ki of 0.71 and 0.27 for midazolam hydroxylation and testosterone hydroxylation, respectively, indicating that apatinib might cause pharmacokinetic DDI via inhibition of CYP3A4 (Zhu et al., 2020). Therefore, it is of great significance to assess the magnitude of the DDI between nifedipine and apatinib. It has been shown that compared with the single oral administration of nifedipine, coadministration with apatinib contributed to the significant increases of AUC0–48h and Cmax of nifedipine by 83% and 64%, respectively (Zhu et al., 2020). But whether this pharmacokinetic DDI between nifedipine and apatinib could enhance the antihypertensive effect of nifedipine, causing severe changes of blood pressure is unknown, which is critical for dosing regimens.

In the development of the DDI model between apatinib and nifedipine, the type of interaction and related enzyme inhibition parameters were crucial. The types of inhibition in Simcyp included competitive inhibition and mechanism-based inhibition (time-dependent inhibition), with the key parameters of Ki for competitive inhibition, and Kapp,a as well as Kinact for mechanism-based inhibition, respectively. There were two studies regarding the kinetic parameters of enzyme inhibition by apatinib. The IC50 and IC50-shift results indicated that apatinib might not be a time-dependent inhibitor (Bao et al., 2018; Zhu et al., 2020). On the basis of the research, apatinib might inhibit CYP3A4 in a competitive way. However, it was found that the Ki values reported in these two articles were quite different. The Ki value of midazolam hydroxylation reported by Zhu et al. (2020) was 0.71 μM via competitive inhibition, while a much higher Ki value of 11.5 μM was estimated in another study, also with midazolam as a probe substrate (Bao et al., 2018). Due to the fact that the experiment conducted by Zhu et al. was not published, we couldn’t compare the difference of the experiments between the two papers. We then used these two data as the model input parameters and conducted the DDI simulations separately. The simulation results were compared with the clinical observations. It was found that the predicted value was much closer to the clinical observation value in the competitive inhibition mode, with Ki of 0.71 μM as the initial value.

PK/PD modelling can elucidate the causative relationship between drug exposure and response, providing a better understanding of the pharmacological effects that results in the observed drug effect. In the simplest form, the measured plasma concentration serves as input for the concentration-effect relationship. And the observed pharmacological effects are directly linked to the site concentration. The classic and the most common PD model under these conditions is the Emax model. Many reports have repeatedly confirmed a close and direct relationship between circulating drug concentrations and antihypertensive effect (Abernethy et al., 1990; Donnelly et al., 1993; Donnelly et al., 1994; Meredith et al., 1994). Furthermore, it has been demonstrated that the magnitude of the first-dose responsiveness correlates with the responsiveness during steady-state treatment (Donnelly et al., 1993; Donnelly et al., 1994). This correlation creates the possibility of predicting the response to long-term treatment with nifedipine in cancer patients.
individual patients (Meredith et al., 1994). Similar PKPD studies have demonstrated that the antihypertensive responses to other dihydropyridine CCBs, including amlodipine and felodipine, are directly related to the plasma concentration–time profiles for each drug (Edgar et al., 1987; Abernethy et al., 1990; Donnelly et al., 1993).

In the present study, the PBPK model for nifedipine was linked to the PD model to predict the changes in blood pressure caused by the exposure changes of nifedipine with the combination of apatinib. Therefore, the results may be helpful for the dosing regimen for combined administration. That is to say, when apatinib and nifedipine are co-administered, should the nifedipine be administered in regular dose in clinic, or the dose needs to be reduced. If the dose of nifedipine needs to be reduced, what is the appropriate dose?

The CR and IR formulations of nifedipine are most commonly used in clinic. In the study, the two formulations were both incorporated in the development of PBPK models for nifedipine. Owing to the fact that the absorption process of the CR tablet needs to be described by combining the in vitro dissolution data as well as the membrane permeability, the ADAM model was used for nifedipine CR tablet. The in vivo absorption of nifedipine IR tablet was described by the first-order model.

It has been reported that the control of SBP is more important than diastolic blood pressure (DBP) in most patients with hypertension (Levine et al., 2003; Mourad, 2008). Hence, SBP was employed in the PD modelling. An ordinary $E_{\text{max}}$ model (Eq. 3) was used to describe the relationship between nifedipine concentration and the changes in SBP. The $E_{\text{50}}$ was reported to be 12.12 ng/ml for nifedipine at the regular doses (Hirasawa et al., 1985; Levine et al., 2003; Meredith and Elliott, 2004; Niu et al., 2021). So, the $E_{\text{50}}$ value was set to 12.12 ng/ml in this study. Based on the range of $E_{\text{max}}$ values reported in the literature (Hirasawa et al., 1985; Levine et al., 2003; Meredith and Elliott, 2004; Niu et al., 2021), the PD model under different $E_{\text{max}}$ values was examined to fit the observed SBP change caused by nifedipine at therapeutics doses. It has been found that the $E_{\text{max}}$ was set to $-30 \text{ mmHg}$, the PD model fitted best. The ratios of the predicted and observed values of $R_{\text{max}}$ and AUE for nifedipine CR tablet as well as IR tablet were within 2, indicating the good predictive performance of the current nifedipine PD model.

Studies have shown that compared with the single oral administration of nifedipine, co-administration with apatinib contributed to the significant increases of AUC$_{0-48\text{h}}$ and C$_{\text{max}}$ of nifedipine by 83%, and 64%, respectively (Zhu et al., 2020). In the present study, similar statistical results are also seen based on the AUC and C$_{\text{max}}$ increases of 1.73 and 1.41, respectively, after the co-administration of apatinib. The results suggest that co-administration of apatinib could significantly increase the exposure of nifedipine. But whether this pharmacokinetic DDI between nifedipine and apatinib could enhance the antihypertensive effect of nifedipine, causing severe changes of blood pressure is unknown, which is critical for dosing regimens. Hence, the developed PBPK/PD model was used to evaluate the changes in SBP caused by the exposure changes of nifedipine with the combination of apatinib. Two clinical scenarios were simulated using Simcyp virtual populations. The first scenario was designed to determine whether co-administration

| Dose level | Parameters | Unit | Nifedipine alone | Nifedipine combined with apatinib | Ratio |
|------------|------------|------|------------------|-------------------------------|-------|
| 60 mg nifedipine CR tablet | $R_{\text{max}}$ | mmHg | -22.21 | -23.44 | 1.06 |
|  | AUE | mmHg·h | 787.55 | 916.08 | 1.16 |
| 30 mg nifedipine CR tablet | $R_{\text{max}}$ | mmHg | -18.62 | -20.30 | 1.09 |
|  | AUE | mmHg·h | 607.64 | 752.79 | 1.24 |
| 30 mg nifedipine IR tablet | $R_{\text{max}}$ | mmHg | -26.12 | -26.67 | 1.02 |
|  | AUE | mmHg·h | 206.78 | 210.03 | 1.02 |

| Hepatic impairments | Parameters | Unit | Nifedipine alone | Nifedipine combined with apatinib | Ratio |
|---------------------|------------|------|------------------|-------------------------------|-------|
| Child-pugh A | $R_{\text{max}}$ | mmHg | -21.44 | -21.64 | 1.01 |
|  | AUE | mmHg·h | 749.26 | 775.87 | 1.04 |
| Child-pugh B | $R_{\text{max}}$ | mmHg | -23.83 | -24.18 | 1.01 |
|  | AUE | mmHg·h | 909.16 | 949.93 | 1.04 |
| Child-Pugh C | $R_{\text{max}}$ | mmHg | -24.97 | -25.49 | 1.02 |
|  | AUE | mmHg·h | 998.17 | 1055.30 | 1.06 |
of apatinib with regular dose level of nifedipine could cause severe changes of SBP in cancer patients. The second scenario was to determine if dose adjustment of nifedipine was necessary in the combination of apatinib in patients with hepatic impairment. Our study shows that apatinib is unlikely to cause severe pharmacodynamic DDI via inhibition of CYP3A4. It is suggested that nifedipine could be used in combination with apatinib without dose adjustment in clinic. Further clinical studies in small sample are required to confirm the results.

However, there are some limitations in this study which warrant further discussion. First, due to lack of the observed data, the predictive performance of DDI for the nifedipine SR/CR formulations is not verified in this study. Second, although CYP3A4 might be more important in elimination of nifedipine compared to CYP3A5, few studies have suggested that CYP3A5 genotypes might explain variability in systemic exposure to nifedipine to certain extent (Haas et al., 2013). This was particularly important since the frequency of CYP3A5 expressors in Chinese population was highly relative to that in populations of European ancestry. Considering the inhibitory effect of apatinib on CYP3A4, we first conduct a DDI study on the CYP3A4. The effect of CYP3A5 is not yet evaluated. Third, a common adverse effect of apatinib was gastrointestinal disturbances, which might affect the transit time across different gastrointestinal segments, particularly for the controlled release formulation. Besides, nifedipine solubility might also be affected by variation in gastrointestinal pH. These are also not considered in the simulations. Despite the limitation existing in our refined model, we still believe that the model can give a rough understanding of the possible exposure change caused by DDI as well as the resulted pharmacodynamic changes to avoid some dangerous DDI in advance.

Conclusion

In the present study, we have investigated the DDI between nifedipine and apatinib. Results show that apatinib could increase the exposure of nifedipine. But this exposure changes has little impact on the antihypertensive effect of nifedipine. Apatinib is unlikely to cause severe pharmacodynamic DDI via inhibition of CYP3A4. It is suggested that nifedipine could be used in combination with apatinib without dose adjustment in clinic.

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Data availability statement

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding authors.

Author contributions

Participated in research design: HL, XX, and BH. Conducted experiments: HL. Performed data analysis: HL and YY. Contributed new reagents or analytic tools: CW, NG, and XW. Wrote or contributed to the writing of the article: HL, XX, and BH. Contributed to the English editing: LL.

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Conflict of interest

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