A novel cardiovascular systems model to quantify drugs effects on the inter-relationship between contractility and other hemodynamic variables

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Abstract
The use of systems-based pharmacological modeling approaches to characterize mode-of-action and concentration-effect relationships for drugs on specific hemodynamic variables has been demonstrated. Here, we (i) expand a previously developed hemodynamic system model through integration of cardiac output (CO) with contractility (CTR) using pressure-volume loop theory, and (ii) evaluate the contribution of CO data for identification of system-specific parameters, using atenolol as proof-of-concept drug. Previously collected experimental data was used to develop the systems model, and included measurements for heart rate (HR), CO, mean arterial pressure (MAP), and CTR after administration of atenolol (0.3–30 mg/kg) from three in vivo telemetry studies in conscious Beagle dogs. The developed cardiovascular (CVS)-contractility systems model adequately described the effect of atenolol on HR, CO, dP/dtmax, and MAP dynamics and allowed identification of both system- and drug-specific parameters with good precision. Model parameters were structurally identifiable, and the true mode of action can be identified properly. Omission of CO data did not lead to a significant change in parameter estimates compared to a model that included CO data. The newly developed CVS-contractility systems model characterizes short-term drug effects on CTR, CO, and other hemodynamic variables in an integrated and quantitative manner. When the baseline value of total peripheral resistance is predefined, CO data was not required to identify drug- and system-specific parameters. Confirmation of the consistency of system-specific parameters via inclusion of data for additional drugs and species is warranted. Ultimately, the developed model has the potential to be of relevance to support translational CVS safety studies.
INTRODUCTION

Cardiovascular (CVS) safety issues are among the major causes of safety-related attrition during drug development.\textsuperscript{1–3} In this context, most attention has been given to potentially fatal arrhythmias arising from QT-prolongation, but non-QT drug effects on hemodynamic end points, such as blood pressure or cardiac contractility, also represent serious CVS safety issues.\textsuperscript{4–9} Preclinical in vivo telemetry studies are often conducted to identify both QT- and non-QT effects, in which an ascending dose or placebo is administered to animals while measuring the time course of changes in heart rate (HR), blood pressure, cardiac output (CO), and/or contractility (CTR; e.g., dP/dt\text{max}).

The interpretation of preclinical data from CVS safety studies, designed to determine the drug mode-of-action and concentration-effect relationship for specific hemodynamic variables, can be challenging due to underlying complex homeostasis and feedback inter-relationships. Quantitative CVS systems models to characterize the dynamics and pharmacokinetic/pharmacodynamic (PK/PD) relationships of hemodynamic variables after drug administration have previously been proposed to address this challenge.\textsuperscript{10,11} A hemodynamic systems model was established for eight compounds with diverse modes-of-action in rats, here referred to as the “Snelder model.”\textsuperscript{12,13} This model characterized drug effects on the inter-relationship among HR, stroke volume (SV), total peripheral resistance (TPR), CO, and mean arterial pressure (MAP).

Importantly, the Snelder model allows for separation of both drug- and system-specific (e.g., drug-independent) parameters. For identification of the system-specific parameters, invasive and challenging CO measurements were required, which limits the integration of this model into a translational modeling platform, which would be an ultimate goal. Alternatively, measures for CTR, such as left ventricular (LV) dP/dt\text{max} could replace CO, as it can be measured more easily using telemetry.\textsuperscript{14} The dP/dt\text{max} is determined by myocardial CTR and the loading conditions on the ventricle, and has been reported to be significantly correlated with SV and HR.\textsuperscript{15} Based on this, and expanding on the work from Snelder et al., an adapted model was recently published using data from telemetry studies in dogs, which replaced CO measurements by CTR (dP/dt\text{max}) measurements.\textsuperscript{16} This model can be used to characterize novel compounds’ effects on the CVS system in dogs. However, in the absence of a mechanistic basis for the inter-relationship among CO, SV, and CTR, the predictive value for other species could be limited.

LV pressure and volume have long been studied with pressure-volume (PV) loop theory. A PV loop can be generated by plotting multiple measurements of LV pressure against LV volume in a complete cardiac cycle. In this

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**Study Highlights**

**WHAT IS THE CURRENT KNOWLEDGE ON THE TOPIC?**

Hemodynamic systems models in rat and dog have been established to characterize drug effects on the inter-relationship between key hemodynamic biomarkers, such as blood pressure, heart rate, and cardiac output (CO). However, to date, no models exist that integrate contractility with hemodynamic variables in a systems manner.

**WHAT QUESTION DID THIS STUDY ADDRESS?**

Can the inter-relationship between contractility and other hemodynamic variable be described in a systems manner? Can contractility readouts (dP/dt\text{max}) replace CO readouts to identify drug- and system-specific parameters?

**WHAT DOES THIS STUDY ADD TO OUR KNOWLEDGE?**

A novel cardiovascular (CVS) systems model was developed based on pressure-volume loop theory using atenolol as proof-of-concept drug. Both systems- and drug-specific parameters of this CVS-contractility (CTR) model could be identified without CO data. Hence, the absence of CO data does not limit the application of this model to data from other species.

**HOW MIGHT THIS CHANGE DRUG DISCOVERY, DEVELOPMENT, AND/OR THERAPEUTICS?**

It is expected that the incorporation of pressure-volume loop theory will provide the mechanistic basis to apply the developed CVS-CTR model for inter-species scaling. It has the potential to be integrated into a translational modeling platform to support CVS safety evaluations.
theory, the changes in the end-systolic pressure-volume relationship reflect inotropic changes and the influence of vascular loading on systolic pressure and flow generation. This provides insight in the myocardial CTR and loading conditions of the heart for the hemodynamic modeling. As the PV loop theory captures the physiological principles of the cardiac cycle, it is uniquely suited as a basis for integration of CTR in the Snelder model in a system-based manner.

Here, we describe the development of a novel CVS-CTR systems model, which integrates dP/dt_max with CO, and other hemodynamic biomarkers (including HR and MAP) based on the principles from PV loop theory, using data from multiple dog telemetry studies for the selective β1-blocker atenolol as proof-of-concept. We characterized the model through identifiability analyses with respect to drug- and system-specific parameters, performed external validation studies, and investigated if dP/dt_max can replace CO for the identification of system-specific parameters.

**METHODS**

**Experimental data**

We obtained data from three in vivo telemetry studies conducted at or on behalf of different pharmaceutical companies in conscious Beagle dogs, which were administered increasing doses of atenolol (0.3–30 mg/kg) or placebo with washout periods in between treatments (Table 1). Each study was conducted according to company-specific standard procedures. In each study, several CVS hemodynamic variables were measured over time for a period of 6 to 24 h. In this analysis, we only used time course measurements for MAP, HR, CO, and LV dP/dt_max. Data from studies 1 and 2 were used for model development, whereas data from study 3 were used for external validation. The data from studies 1 and 3 were obtained after administration of control compounds as part of earlier drug development programs, and have not been previously published. The animals and experimental procedures are described for each respective study in the Supplementary Materials.

**Pharmacokinetic model**

Because PK was not measured in the experimental studies, we utilized a previously published PK model to predict drug plasma concentrations over time. The model was a three-compartment model with first-order absorption and elimination, and the following PK parameters: $K_a = 1.13 \, h^{-1}$, $CL = 3.35 \, L/(kg*h)$, $V_1 = 4.05 \, L/kg$, $Q_2 = 8.85 \, L/(kg*h)$, $V_3 = 3.74 \, L/kg$, $Q_3 = 5.73 \, L/(kg*h)$,
$V_1 = 11.9$ L/kg, and $F_1 = 0.783$. The predicted concentrations in the central compartment were directly linked to the drug effects.

**Relationship between hemodynamic variables**

We defined relationships between the hemodynamic variables used in the systems modeling framework as follows: MAP was described as a product of CO and TPR based on Ohm’s law, and CO was the product of HR and SV (Equation 1)

\[
\text{MAP} = \text{CO} \cdot \text{TPR} \\
\text{CO} = \text{HR} \cdot \text{SV}
\]

SV was defined further as:

\[
\text{SV} = \text{EDV} \cdot \text{ESV}
\]

To describe the inter-relationship between SV and CTR, we considered the LV PV loop theory. There are two key linear relationships in PV loops (Figure 1). Line $E_A$ (Figure 1) represents the afterload relationship with the slope as arterial elastance ($E_A$) (Equation 3)

\[
\text{Line } E_A: P_A = (\text{EDV} - \text{ESV}) \cdot E_A
\]

where $E_A$ is the product of HR and TPR (Equation 4).

\[
E_A = \text{HR} \cdot \text{TPR}
\]

Line ESPVR (Figure 1) represents the end-systolic pressure-volume relationship, which depends on CTR, and is defined as follows in relation to CTR:

\[
\text{Line ESPVR: } P_A = (\text{ESV} - V_0) \cdot \text{CTR}
\]

Here, $V_0$ is the x-axis intercept of ESPVR, or the minimum volume required to get pressure in the LV. Equations 3 and 5 were solved to obtain the following relationship for ESV (Equation 6):

\[
\text{ESV} = \frac{\text{EDV} \cdot E_A + V_0 \cdot \text{CTR}}{E_A + \text{CTR}}
\]

$dP/dt_{\text{max}}$ is used in this study as indicator of myocardial CTR, which depends on end diastolic volume (EDV; preload) and duration of isovolumetric contraction (TIC). Therefore, we defined the variable CTR measurements (CTRM), which reflects for $dP/dt_{\text{max}}$ measurements (Equation 7), as follows:

\[
\text{CTRM} = \text{CTR} \cdot \frac{\text{EDV}}{\text{TIC}}
\]

For a typical young Beagle dog (27 months) a TIC of 0.256 s was reported.

**Model structure**

The CVS system is regulated via various control mechanisms, such as the baroreflex system and the renin-angiotensin system. To account for these negative feedback mechanisms, and to allow for the inclusion of drug effects into the model, the dynamics of HR, EDV, TPR, and CTR were described by a set of four differential equations, which were linked via negative feedback through MAP (Equation 8), similar to the Snelder model.

\[
\frac{d\text{HR}}{dt} = k_{\text{inHR}} \cdot (1 - \text{FB} \cdot \text{MAP}) - k_{out\text{HR}} \cdot \text{HR}
\]

\[
\frac{d\text{EDV}}{dt} = k_{\text{inEDV}} - k_{out\text{EDV}} \cdot \text{EDV}
\]
\[
\frac{dTPR}{dt} = k_{inTPR} \cdot (1 - FB \cdot MAP) - k_{outTPR} \cdot TPR \tag{8}
\]
\[
\frac{dCTR}{dt} = k_{inCTR} \cdot (1 - FB \cdot MAP) - k_{outCTR} \cdot CTR
\]

Here, FB is the feedback effect parameter through MAP to all the three primary parameters. The system was forced to be in steady-state, or dynamic equilibrium, before drug administration, which means HR, EDV, TPR, and CTR do not change over time and equal their baseline (BSL) values. Therefore, the production rates of HR, EDV, TPR, and CTR can be expressed in terms of baseline and \( k_{out} \) as follows:

\[
k_{inHR} = \frac{k_{outHR} \cdot BSL_{HR}}{1 - FB \cdot BSL_{MAP}}
\]
\[
k_{inEDV} = k_{outEDV} \cdot BSL_{EDV}
\]
\[
k_{inTPR} = \frac{k_{outTPR} \cdot BSL_{TPR}}{1 - FB \cdot BSL_{MAP}}
\]
\[
k_{inCTR} = \frac{k_{outCTR} \cdot BSL_{CTR}}{1 - FB \cdot BSL_{MAP}}
\]

### Circadian rhythms in hemodynamic variables

Three cosine functions were defined as follows to model the circadian rhythm in HR, TPR, and CTR (CRHR, CRTPR, and CRICTR) influencing \( k_{inHR} \), \( k_{inTPR} \), and \( k_{inCTR} \), respectively:

\[
CR_{HR}(t) = amp_{HR} \cdot \cos \left(2\pi \cdot \frac{t + t_{hor_{HR}}}{24}\right)
\]
\[
CR_{CTR}(t) = amp_{CTR} \cdot \cos \left(2\pi \cdot \frac{t + t_{hor_{CTR}}}{24}\right)
\]
\[
CR_{TPR}(t) = amp_{TPR} \cdot \cos \left(2\pi \cdot \frac{t + t_{hor_{TPR}}}{8}\right)
\]

Here, amp refers to amplitude and hor is the horizontal displacement. We tested periods of 8, 12, and 24 h, and identified the optimal period for each variable. The system was forced to be in oscillating steady-state at the start of pharmacological intervention by shifting the observations 1 week (determined empirically) in time (i.e., the system was initialized at 0 h and the pharmacological interventions started at 168 h).

### Modelling of the concentration-effect relationship

Concentration-effect relationships were evaluated using maximum effect (\( E_{max} \)) models.

\[
E_{max} \text{ model: } EFF(C) = \frac{E_{max} \cdot C}{EC_{50} + C} \tag{11}
\]

Here, EFF represents the drug effect at concentration C, \( E_{max} \) is the maximum effect constant, and \( EC_{50} \) is the concentration constant at which half of the maximum effect is achieved. Based on literature values for the binding affinity (\( K_D \)) of atenolol to the \( \beta_1 \) and \( \beta_2 \)-adrenoceptors from Baker et al.,\textsuperscript{23} we fixed the \( EC_{50} \) for the effects on HR and CTR, which are mediated through the \( \beta_1 \)-adrenoceptors, to 58.3 ng/ml and the \( EC_{50} \) for the effect on TPR, which is mediated through the \( \beta_2 \)-adrenoceptor, to 272.5 ng/ml.

### Incorporation of drug effects in the systems model

The complete model consisted of a combination of Equations 1–11 and the following differential equations:

\[
\frac{dHR}{dt} = k_{inHR} \cdot (1 - FB \cdot MAP) \cdot (1 - EFF_{HR}) \cdot (1 + CR_{HR}) - k_{outHR} \cdot HR
\]
\[
\frac{dEDV}{dt} = k_{inEDV} - k_{outEDV} \cdot EDV
\]
\[
\frac{dTPR}{dt} = k_{inTPR} \cdot (1 - FB \cdot MAP) \cdot (1 - EFF_{TPR}) \cdot (1 + CR_{TPR}) - k_{outTPR} \cdot TPR
\]
\[
\frac{dCTR}{dt} = k_{inCTR} \cdot (1 - FB \cdot MAP) \cdot (1 - EFF_{CTR}) \cdot (1 + CR_{CTR}) - k_{outCTR} \cdot CTR
\]

### Structural identifiability analysis

A structural identifiability analysis was performed to verify whether the model parameters could be estimated from experimental data. Structural identifiability of the model was evaluated using the MATLAB toolbox GenSSI 2.0, which is a MATLAB toolbox together with MATLAB (version R2020a).\textsuperscript{24} We implemented simultaneously two different drug effects, namely, EFFHR and EFFCTR, to influence the production rate \( k_{in} \) of HR and CTR, respectively. At the same time, we applied three different circadian rhythms (i.e., CRHR, CRTPR, CRICTR), to influence...
the production rate of HR, TPR, and CTR, respectively. We verified the structural identifiability of the model for two different sets of observable outputs, namely HR, CTRM, CO, and MAP and HR, CTRM, and MAP.

Model estimation and development

The data from studies 1 and 2 were simultaneously analyzed using a nonlinear mixed-effects modeling approach implemented in NONMEM (version 7.4.3, Icon Development Solutions, Ellicott, MD) with Perl speak to NONMEM toolkit (PsN, version 4.8.1, Uppsala University, Sweden). First order linear conditional estimation method with interaction was used for model estimation. Interindividual variability (IIV) on BSLHR, BSLTPR and BSLCTRM were evaluated using log-normal distributions. Exponential, additional, proportional, and combined error models were evaluated for residual error variability. A decrease of 3.84 (corresponding to \( p < 0.05 \) in a chi-squared distribution with degree of freedom = 1) in the objective function value (OFV) by adding an additional parameter between nested models was considered statistically significant. The baseline parameters for HR, CTR, and TPR were estimated together with \( V_0 \), while ensuring that these parameters remain positive during the estimation.

Relevance of the CO data

To investigate the importance of CO data in the estimation, the final model was fitted to a dataset without CO readouts. In the absence of remaining informative data, the circadian rhythm (CRTPR) and IIV on BSLTPR were omitted from the model and BSLTPR was fixed to 0.0743 mmHg*ml/min, which was the estimate in the final model. Fixing this parameter was required to avoid over-parameterization.

Model evaluation

Visual predictive checks (VPC) with 1000 samples and goodness-of-fit plots (GOF) were conducted to evaluate the predictive performance of the final model. External validation consisted of comparing the model prediction (95% prediction interval using the final model) with hemodynamic observations from study 3.

External validation

The external validation was conducted based on the data from study 3. A VPC was performed to predict the effect of atenolol on the hemodynamic markers as detailed here above. All the system-specific parameters were assumed the same as the final estimates from study 1.

Identification of the right site of action of compounds given the data

Stochastic simulations and re-estimation (SSE) using PsN toolkit in conjunction with NONMEM were conducted to investigate if the model can be used to identify the mode of action of new drugs. We simulated three types of models with drug effects on HR, CTR, or TPR, defined as the original models in this analysis. Subsequently, we re-estimated the parameters on the simulated datasets using models with drug effects on HR, CTR, or TPR and no drug effect, defined as the alternative models in this analysis. The simulation scenarios included three combinations of measurements at rich timepoints for (i) HR, dPdt\(_{\text{max}}\), CO, and MAP; (ii) HR, dPdt\(_{\text{max}}\), and MAP; and (iii) HR and MAP. The parameter values used in the simulation steps were fixed to the final estimates from previous model fitting. The difference in OFV (dOFV) between drug effect models and no drug effect models was calculated to compare the performance of the models with different sites of action.

Simulations to illustrate the properties of the CVS system

To illustrate the properties of the system for drugs with a primary effect on HR, CTR, or TPR, simulations were performed using the final model (Equations 1–11). The values of system-specific parameters in the simulation were fixed to the final estimates obtained from the final model, and hypotheses of both negative and positive effects (reflecting different mechanisms of action [MOA]) were tested.

RESULTS

Structural identifiability analysis

The structural identifiability analysis showed that the proposed model is at least locally identifiable for both sets of observable outputs (HR, CTRM, CO, and MAP and HR, CTRM, and MAP), indicating the possibility of estimating parameter values from experimental data. The difference between the two sets of observable variables is that when CO is measured all baseline parameters (i.e., BSLCTRM, BSLHR, BSLTPR, and \( V_0 \)) are globally identifiable indicating that they can be uniquely determined, whereas when no
CO data are available BSL_{TPR} and V_0 are only locally identifiable indicating that more than one solution is possible.

**Model development and evaluation**

Data from studies 1 and 2 were adequately described by the final CVS-CTR model (Figure 2 and Equations 1–11) as can be seen in the VPC (Figure 3) and GOF plots (Figures S1–S4). All parameter could be estimated precisely, with relative standard errors less than 50% (Table 2). The concentration-effect relationships were best described by E_{max} models. The EC_{50} for HR and CTR was fixed to the reported K_D of 58.3 ng/ml for binding of atenolol to the β1 receptor. E_{max} for the effect on HR and CTR was estimated to be 0.415 and 0.422, respectively. An E_{max} model was added for a positive effect on TPR with the EC_{50} fixed to 272.5 ng/ml. However, as the final estimate of E_{max} for TPR approached zero, and the OFV did not decrease significantly (3.84, p < 0.05), this additional drug effect was rejected. Overall, the final model was structured with inhibiting effects on both HR and CTR.

The rate constants K_{outHR}, K_{outCTR}, K_{outTPR}, and K_{outEDV} were initially estimated separately and did not differ significantly. As the OFV did not increase significantly (p < 0.05) after estimating the same K_{out} for all four biomarkers, these four parameters were lumped into one parameter (K_{out}). Because CO was not measured in study 2, the typical values of BSL_{TPR} were assumed valid for both studies. The other baseline parameters differed between the two studies, with a 3.12% higher BSL_{HR} and 8.42% higher V_0 for study 1 as compared to study 2, whereas BSL_{CTRm} for study 1 was 55.94% higher than for study 2. BSL_{SV}, BSL_{CO}, and BSL_{MAP} were derived from these parameters, resulting in higher BSL_{SV}, BSL_{CO}, and BSL_{MAP} for study 1 as compared with study 2. The amplitude parameters am_{HR}, am_{CTR}, and am_{TPR}, could not be distinguished and were estimated to be 0.0931 for study 1 and 0.168 for study 2.

The IIV was estimated on the baseline parameters BSL_{HR}, V_0, BSL_{CTRm}, and BSL_{TPR}, and covariance was estimated between ETA_{BSL_{HR}} and ETA_{V_0}, and between ETA_{BSL_{CTRm}} and ETA_{BSL_{TPR}}. The residual errors on HR and dP/dt_{max} were best described by exponential residual error models with lower RSEs compared to additive, proportional, and combined residual error models. The residual error of TPR was very small and, therefore, it was fixed to 0.

**Evaluation of the relevance of the CO data**

Fitting the CVS-CTR model to a dataset from which the CO data were omitted did not result in a significant change in parameter estimates; almost all estimates were within the 95% confidence interval (CI) of the parameters of the final model fitted to the complete dataset including CO data (Table 2). The only parameter outside the 95%
CI, IIV on BSL on HR, was also close to the lower boundary of 95% CI and would not significantly influence the model prediction.

**External validation**

The external validation for study 3 (Figure S5) showed that in general the validatory data were reasonably well-predicted. However, the model slightly underpredicted the effect on dP/dt\text{max} in the 3 mg/kg dose group, possibly reflecting interstudy variability.

**Identification of the right site of action of compounds given the data**

In the SSE analysis, the models with the “correct” MOA showed the highest reduction in OFV for all three simulation scenarios (Figure 4). With decreasing numbers of observations, the difference of “correct” and “incorrect” models in OFV gradually shrank. Especially, when re-estimating the parameters on the HR/MAP data from the model with the original effect on CTR, the dOFV for “incorrect” and “correct” models was quite close.

**Simulations to illustrate the properties of the CVS system**

In the simulations of drug effects with different MOAs, there were noticeable differences between the signature profiles of HR, dP/dt\text{max}, CO, TPR, and MAP. As shown in Figure 5, inhibition of HR, CTR, or TPR always lead to a drop in MAP indicating that the magnitude of negative feedback from MAP is lower than the primary effect. Similarly, the positive effects on HR, CTR, or TPR resulted in an increase in MAP, as shown in Figure S6. The site of action of a compound with unknown MOA can be derived from the combination of the direction of the effects on MAP, HR, and CTR.

**DISCUSSION**

We describe the development of a novel CVS-CTR systems model to characterize drug effects on the inter-relationship among CTR and CO, MAP, HR, TPR, SV, ESV, and EDV. The proposed model was based on a previously developed CVS model by Snelder et al.,\textsuperscript{12,13} which captures the inter-relationship between CO, MAP, HR, TPR, and SV. To date, no mechanistic models exist that integrate CTR with the other mentioned hemodynamic variables, except for a model that was recently published by Venkatasubramanian et al.\textsuperscript{16} This model describes SV as a function of CTR and HR, to account for the effect of lowering SV due to shorter filling time with increased HR during diastole, and was successfully applied to characterize drug effects on the hemodynamic system in dogs. Although the relationship between SV, CTR, and HR has a mechanistic basis, the proposed model is not truly mechanistic, as it does not account for the effect of afterload/TPR on the ability of the ventricle to eject blood, altering ESV and SV. PV-loop theory does cover these aspects.\textsuperscript{17} Hence, we based our model on this theory. Better capturing the physiology of the CVS system via PV-loop theory allows for improved separation of system-specific and drug-specific parameters, which is crucial to understand variation between species.\textsuperscript{26} Therefore, the developed CVS-CTR model is the first systems model that integrates CTR with other key hemodynamic variables. By fitting the CVS-CTR systems model to a dataset that did not include CO data, we showed that CTR data can replace CO data. The absence of CO data does not limit the application of this model to data from other species, which facilitates the development of a translational modeling platform.

The developed CVS-CTR model adequately described the effect of atenolol on the inter-relationship between CTR and the other hemodynamic variables. Atenolol was selected as a proof-of-concept compound because it is widely used, and its hemodynamic effects are well-investigated. To identify the system-specific parameters in the best possible way, we fixed the EC\textsubscript{50} for the effects of atenolol on HR and CTR to literature (K\textsubscript{D}) values. Although the EC\textsubscript{50} is a lumped parameter for binding to the receptor and signal transduction, we assumed that the EC\textsubscript{50} could be fixed to the reported K\textsubscript{D}. To evaluate if these K\textsubscript{D} values were covered by the exposures in the dog studies we simulated the atenolol PK profiles following single oral doses of 0.3, 1, 3, 10, or 30 mg/kg atenolol (Figure S7). Simulated concentrations after doses of 0.3 and 1 mg/kg did not reach the K\textsubscript{D} of β\textsubscript{1}-adrenoceptors, which is in line with the weak drug effect observed in these low-dose groups. Moreover, at high doses of 10 and 30 mg/kg,
only peak concentrations were above the $K_D$ for binding to β2-adrenoceptors, which is in line with the finding that adding a positive drug effect on TPR did not significantly improve the description of the data. Overall, our assumed

\[ EC_{50} \]

seems justified, as the CVS effects of atenolol could be adequately described by a combination of primary inhibitory effects on both HR and CTR, which corresponds well with its high affinity for β1-adrenoceptors. By comparing

### TABLE 2

The system- and drug-specific parameter values from the CVS-contraction model fitted to a dataset including CO data (final model) and a dataset from which the CO data were omitted (model without CO data)

| Parameter | Final model | Model without CO data |
|-----------|-------------|-----------------------|
|           | Value (RSE) | 95% CI                | Value (RSE) |
| System-specific parameters |            |                       |             |
| BSLHR\(_{1}\) (beats/min\(^{-1}\)) | Baseline of HR for study 1 | 79.4 (5.78%) | 70.4–88.3 | 78.4 (4.69%) |
| $V_0$ (ml) | $V_0$ for study 1 | 9.92 (9.81%) | 8.02–11.8 | 10.1 (11.0%) |
| BSLCTRM\(_{1}\) (mmHg/s) | Baseline of dP/dt\(_{\text{max}}\) for study 1 | 3777 (7.59%) | 3215–4338 | 3652 (9.04%) |
| BSLHR\(_{2}\) (beats/min\(^{-1}\)) | Baseline of HR for study 2 | 77.0 (2.63%) | 73.0–81.0 | 76.4 (2.73%) |
| $V_0$ (ml) | $V_0$ for study 2 | 9.15 (9.62%) | 7.42–10.9 | 8.55 (4.76%) |
| BSLCTRM\(_{2}\) (mmHg/s) | Baseline of dP/dt\(_{\text{max}}\) for study 2 | 2422 (7.50%) | 2067–2779 | 2480 (7.70%) |
| BSLTPR (mmHg*min/ml) | Baseline of TPR | 0.0743 (4.55%) | 0.0677–0.0810 | 0.0743 FIX |
| BSEDM (ml) | Baseline of EDV | 31.13 | FIX | 31.13 FIX |
| $K_{\text{out}}$ (h\(^{-1}\)) | Degradation rate | 0.830 (21.8%) | 0.475–1.18 | 1.07 (22.9%) |
| FB (mmHg\(^{-1}\)) | Feedback effect | 0.00558 (12.5%) | 0.00422–0.00694 | 0.00589 (27.9%) |
| Circadian Rhythm |            |                       |             |
| Amp\(_{1}\)\(^{b}\) | Amplitude for study 1 | 0.0931 (41.3%) | 0.0178–0.168 | 0.0775 (48.1%) |
| HorHR\(_{1}\) | Horizontal displacement of HR | 7.86 (28.7%) | 3.43–12.3 | 6.33 (17.8%) |
| HorCTR\(_{1}\) | Horizontal displacement of CTR | 9.82 (18.9%) | 6.19–13.5 | 8.57 (16.6%) |
| Amp\(_{2}\) | Amplitude for study 2 | 0.168 (13.6%) | 0.123–0.213 | 0.172 (22.6%) |
| HorHR\(_{2}\) | Horizontal displacement of HR | 19.4 (1.60%) | 18.8–20.1 | 19.5 (1.99%) |
| HorCTR\(_{2}\) | Horizontal displacement of CTR | 21.8 (1.84%) | 21.0–22.6 | 21.8 (2.84%) |
| HorTPR | Horizontal displacement of TPR | 6.33 (1.96%) | 6.09–6.58 | 0 FIX |
| Drug-specific parameters |            |                       |             |
| $EC_{50\text{HR}}$ and $EC_{50\text{CTR}}$ (ng/ml) | $EC_{50\text{HR}}$ and $EC_{50\text{CTR}}$ | 58.3 | FIX | FIX |
| $E_{\text{max\text{HR}}}$ | $E_{\text{max\text{HR}}}$ | 0.415 (11.6%) | 0.321–0.510 | 0.402 (18.0%) |
| $E_{\text{max\text{CTR}}}$ | $E_{\text{max\text{CTR}}}$ | 0.422 (9.56%) | 0.343–0.501 | 0.421 (14.8%) |
| Interindividual variability |            |                       |             |
| BSLHR (CV\%) | BSLHR (CV\%) | 6.08 (24.2%) | 4.41–7.39 | 3.89 (45.0%) |
| BSLCTRM (CV\%) | BSLCTRM (CV\%) | 16.58 (34.2%) | 9.47–21.52 | 16.99 (33.5%) |
| BSLTPR (CV\%) | BSLTPR (CV\%) | 6.48 (29.6%) | 4.2–8.15 | 0 FIX |
| BSLCTRM X BSLTPR (CV\%) | BSLCTRM X BSLTPR (CV\%) | 8.88 (30.9%) | 5.57–11.26 | 0 FIX |
| Residual variability |            |                       |             |
| Res. Error\(_{\text{HR}}\) (CV\%) | Res. Error\(_{\text{HR}}\) (CV\%) | 11.53 (12.7%) | 9.98–12.9 | 11.37 (11.9%) |
| Res. Error\(_{\text{dP/dt\text{max}}}\) (CV\%) | Res. Error\(_{\text{dP/dt\text{max}}}\) (CV\%) | 10.43 (13.6%) | 8.93–11.74 | 10.26 (13.6%) |

Abbreviations: CI, confidence interval; CO, cardiac output; CVS, cardiovascular; $EC_{50}$, half-maximal effective concentration; EDV, end diastolic volume; $E_{\text{max}}$, maximum effect; HR, heart rate; RSE, relative standard error; TPR, total peripheral resistance.

\(^{a}K_{\text{out\text{HR}}} = K_{\text{out\text{CTR}}} = K_{\text{out\text{TPR}}} = K_{\text{out\text{EDV}}} = K_{\text{out}}.\)

\(^{b}Amp_{\text{HR}} = Amp_{\text{CTR}} = Amp_{\text{TPR}} = Amp_{\text{Amp}}_{\text{Amp}}_{\text{1}}\) and $Amp_{\text{2}}$ are the amplitudes of circadian rhythm for studies 1 and 2 respectively, and for TPR the Amp and Hor were assumed to be the same as in study 1.
the value of amplitude and $E_{\text{max}}$ parameters, it was found that the magnitude of circadian rhythm is approximately three to four times lower than the maximum drug effect. In addition, in the current analysis we have fixed $K_{\text{outCTR}}$, $K_{\text{outTPR}}$, and $K_{\text{outEDV}}$ to the same value as $K_{\text{outHR}}$, as these parameters could not be estimated independently. Identical $K_{\text{out}}$ values could be biologically plausible because all variables studied are influenced by feedback through the baroreflex system. To confirm this, our model should be fitted to data for additional drugs with diverse mode of action in the future.

In our experimental data, a drop was observed in both HR and CTR after administration of atenolol. Therefore, we assumed that the primary drug effect is on both HR and CTR, which is in line with the MOA of $\beta$1-blockers. This assumption was challenged by forward inclusion and backward exclusion during the model estimation. An increase in HR leads to an increase in the force of contraction generated by the myocardial cells through the Bowditch effect, which indicates that CTR can be regulated by HR. This phenomenon is associated with calcium ion handling and mishandling in cardiac cells. However, this pathway was not included in our current model structure as it could not be distinguished from the effect atenolol on both HR and CTR. In future research, we will consider to introduce this regulation on CTR by HR in the model, if a similar behavior pattern of HR and CTR is observed for other compounds with different MOAs as well.

The developed model incorporates EDV as a key variable. As the available data did not contain any information to estimate baseline EDV, this parameter had to be fixed to a reported literature value to avoid over-parameterization. Importantly, although EDV does not currently impact model predictions, its inclusion was considered necessary to allow future application, for example, compounds with a primary drug effect on EDV. Dynamics of EDV may be further evaluated in future work if data for drugs with a known effect on SV would be available, and the necessity to estimate a different $K_{\text{out}}$ for EDV could then be investigated.

In the MOA-identification analyses, the site of action of new compounds could be identified correctly with three combinations of observations: (1) HR, CO, $dP/dt_{\text{max}}$ and MAP; (2) HR, $dP/dt_{\text{max}}$ and MAP; and (3) HR and MAP. This implies that CO readouts are not needed for identification of the site of action of compounds in the SSE analyses. In the simulations to illustrate the system properties after simulating drug effects on HR, CTR, and TPR, there are noticeable differences in the behavior of HR, $dP/dt_{\text{max}}$, TPR, CO, and MAP (Figure 5 and Figure S5). This indicates that the site of action of new compounds can be distinguished by comparing the direction of effect on HR, $dP/dt_{\text{max}}$ and MAP. Because the negative feedback effect is always lower than the
primary effect, the primary site of action is the biomarker with the same direction of effect as the effect on MAP. Thus, we anticipate that the model can be used to identify the site of new compounds with a single site of action, for which the mechanism underlying CVS effect is unclear. For more complex MOAs with more than one primary site of action, different assumptions will need to be evaluated.

CONCLUSION

We successfully developed a novel CVS-CTR model to characterize drug effects on the inter-relationship between CTR and other key hemodynamic variables based on pressure-volume loop theory. Through fitting our model to data from telemetry studies of experimental compounds, our model can support the identification and quantification of the mode of action on key hemodynamic end points. We expect that the incorporation of PV-loop theory will provide mechanistic basis that could be of relevance when applying this model for interspecies scaling. Key hurdles in this extrapolation may, for instance, be differences in homeostatic feedback mechanisms and set points. Future studies may focus on applying the developed model to additional compounds in different species to confirm consistency of system-specific parameters and scalability toward other species, and to potentially study how disease state or exercise will influence hemodynamic responses.

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CONFLICT OF INTEREST

The authors declared no competing interests for this work.
AUTHOR CONTRIBUTIONS
Y.F., H.T., M.M.S., J.G.C.H., and N.S. designed and performed the research and analyzed the data. Y.F., H.T., M.M.S., E.R., T.A.C., S.B., D.L., P.H.G., J.G.C.H., and N.S. wrote the manuscript.

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