An Interactive Generic Physiologically Based Pharmacokinetic (igPBPK) Modeling Platform to Predict Drug Withdrawal Intervals in Cattle and Swine: A Case Study on Flunixin, Florfenicol, and Penicillin G

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ABSTRACT

Violative chemical residues in edible tissues from food-producing animals are of global public health concern. Great efforts have been made to develop physiologically based pharmacokinetic (PBPK) models for estimating withdrawal intervals (WDIs) for extralabel prescribed drugs in food animals. Existing models are insufficient to address the food safety concern as these models are either limited to 1 specific drug or difficult to be used by non-modelers. This study aimed to develop a user-friendly generic PBPK platform that can predict tissue residues and estimate WDIs for multiple drugs including flunixin, florfenicol, and penicillin G in cattle and swine. Mechanism-based in silico methods were used to predict tissue/plasma partition coefficients and the models were calibrated and evaluated with pharmacokinetic data from Food Animal Residue Avoidance Databank (FARAD). Results showed that model predictions were, in general, within a 2-fold factor of experimental data for all 3 drugs in both species. Following extralabel administration and respective U.S. FDA-approved tolerances, predicted WDIs for both cattle and swine were close to or slightly longer than FDA-approved label withdrawal times (eg, predicted 8, 28, and 7 days vs labeled 4, 28, and 4 days for flunixin, florfenicol, and penicillin G in cattle, respectively). The final model was converted to a web-based interactive generic PBPK platform. This PBPK platform serves as a user-friendly quantitative tool for real-time predictions of WDIs for flunixin, florfenicol, and penicillin G following FDA-approved label or extralabel use in both cattle and swine, and provides a basis for extrapolating to other drugs and species.
Drug residues that exceed allowable concentrations or unsafe chemical substances in animal-derived food products are a challenge to global food safety (Baynes et al., 2016; Canton et al., 2021; Vanselow and Griffith, 2001). Generally, veterinary drugs are used to prevent or treat infectious diseases, improve feed efficiency, and enhance productivity in food-producing animals (Cully, 2014). However, these benefits are associated with the risk of drug residues above regulatory safety levels resulting in human health hazards (Durso and Cook, 2014). Violative residues within edible products or tissues of treated animals could increase the health risk to consumers or even result in the suspension of the producer’s permit or certification to affect the global food trade of the agricultural product (NRC, 1999). Violative residues can often be caused by inappropriate extralabel drug use and/or failure to observe an adequate withdrawal interval (WDI) (KuKanich et al., 2005). Therefore, in order to prevent drug residue violations and protect human food safety relative to consumption of animal-derived food products, it is important to develop a scientifically based approach to estimate tissue residues and WDIs of drugs in food-producing animals (Canton et al., 2021; Riviere et al., 2017).

A physiologically based pharmacokinetic (PBPK) model is a mechanism-based model that is capable of describing the absorption, distribution, metabolism, and excretion of chemicals in different species by incorporating physiological- and compound-specific parameters (Lin et al., 2016a). Over the past several decades, PBPK models have been widely used in many fields including animal-derived safety assessment (Henri et al., 2017; Li et al., 2019a), nanomedicine (Lin et al., 2016b; Singh et al., 2020), as well as animal and human health assessment of environmental chemicals (Chou and Lin, 2020; Lautz et al., 2020; Lin et al., 2017; Tan et al., 2018). In particular, in the field of food safety, multiple PBPK models have been developed for several veterinary drugs in different food animals by multiple research groups from different countries to determine drug WDIs based on their respective regulatory standards (Henri et al., 2017; Riad et al., 2021; Yang et al., 2019; Zhou et al., 2021). Estimation of WDIs using a scientific tool such as PBPK models may be very useful to avoid violative drug residues and keep animal-derived food products safe for human consumption when drugs are used under field conditions.

The Food Animal Residue Avoidance Databank (FARAD) program is a U.S. Congressionally authorized U.S. Department of Agriculture-supported national food safety program with the mission of helping producers and veterinarians prevent or mitigate illegal or harmful residues of drugs, pesticides, biotoxins, and other chemical agents that may contaminate foods of animal origin (Riviere et al., 2017). To achieve this mission, FARAD researchers collect and analyze pharmacokinetic data in order to develop pharmacokinetic models, including PBPK models, to help answer requests from veterinarians for WDIs for drugs prescribed extralabel in food animals. Of note, every year FARAD receives thousands of WDI requests for many different drugs administered to various food animal species that directly impact millions of food animals as well as indirectly benefitting numerous consumers (internal data from FARAD Call Centers). However, existing PBPK models are either limited to 1 specific drug or difficult to use by individuals without programming experience. PBPK models are relatively complex in nature and require numerous model parameters such as metabolic rates and tissue partition coefficients as input. Thus, it is not easy to adapt a model developed for 1 drug to use for other drugs, especially considering that different drugs often have different metabolic processes. Therefore, existing PBPK models are not sufficient to address the food safety concern and WDI requests for many drugs dosed with various therapeutic regimens in a timely manner. A user-friendly generic PBPK model platform that can be applied to multiple drugs is urgently needed to address this national and global food safety need.

Therefore, the objective of this study was to develop a web-based generic PBPK modeling platform that can be used to estimate tissue residues and WDIs of multiple drugs rapidly and easily in different food animal species. To develop and evaluate this modeling platform, we chose flunixin, florfenicol, and penicillin G as representative medications and selected cattle and swine as model species. These 3 drugs were selected as they are medications for which FARAD received the most WDI requests over the last several years (Li et al., 2017, 2019a; Yang et al., 2019). In addition, our research team previously developed PBPK models for each of these medications individually (Li et al., 2017, 2019a; Yang et al., 2019). These earlier studies provide a basis to develop the present more robust and comprehensive PBPK model and enable comparisons of simulation results between different models. In addition, new pharmacokinetic studies have been published since the development of these earlier models. In real-time residue mitigation events, it is ideal to incorporate new published datasets to improve existing PBPK models so that they are more robust than the originally developed model. Cattle and swine were selected as the model species because they are included in the FDA’s list of major food animal species in the United States. In this study, a species-specific tissue composition-based in silico model was incorporated to estimate the tissue-to-plasma partition coefficients. As such, the final PBPK model is more flexible and may be extrapolated to other food animal species. To ensure user-friendliness, the final PBPK model was converted to a web-based interactive visualization platform.

**Key words:** drug residue; Food Animal Residue Avoidance Databank (FARAD); food safety; interactive generic physiologically based pharmacokinetic (igPBPK) model; withdrawal interval (WDI).

**MATERIALS AND METHODS**

**Pharmacokinetic data for model development.** All pharmacokinetic data used in model calibration and evaluation were acquired from the FARAD Comparative Pharmacokinetic Database (http://www.farad.org) (Riviere et al., 2017) with the keywords “Flunixin,” “Florfenicol,” and “Penicillin G.” The pharmacokinetic data for cattle and swine following oral, intravenous (IV), intramuscular (IM), and subcutaneous (SC) administration were selected. Data in dairy cows and lactating sows were excluded because of the additional elimination route through milk that may result in differences in the pharmacokinetics between lactating and non-lactating animals. The pharmacokinetic data sets of flunixin, florfenicol, and penicillin G in plasma or tissues were collected from tables or digitized from figures in the literature using WebPlotDigitizer (version 4.4, https://automeris.io/WebPlotDigitizer). A summary of key information of selected pharmacokinetic studies is provided in Table 1.
# A Summary of Pharmacokinetic Studies Used for Model Calibration and Evaluation

| Reference          | Routes      | Dose (mg/kg) | Repeat dose | Species | Matrix | Compounds     | Use   |
|--------------------|-------------|--------------|-------------|---------|--------|---------------|-------|
| **Flunixin**       |             |              |             |         |        |               |       |
| Howard et al. (2014) | IV          | 3            | Single      | Swine   | P      | FLU, SOH-FLU  | Cal   |
| Paires-Garcia et al. (2013) | IV, IM, PO | 2.2          | Single      | Swine   | P      | FLU           | Cal   |
| FDA (2005)         | IM          | 2.2          | 3 days      | Swine   | L, M, K, F | FLU           | Cal   |
| Buur et al. (2006) | IV          | 2            | Single      | Swine   | P      | FLU           | Cal   |
| EMA (1999)         | IM          | 2.4          | 3 days      | Swine   | L, M, K | FLU           | Cal   |
| Bates et al. (2020) | IM          | 2.2          | Single      | Swine   | P, L, K, M, U | FLU, SOH-FLU | Cal   |
| Kittrell et al. (2020) | IM, PO | 2.2 (IM); 3.3 (PO) | Single      | Swine   | P      | FLU           | Eval  |
| FDA (1998)         | IV          | 2.2          | 3 days      | Cattle  | L, M, K, F | FLU           | Cal   |
| Shelver et al. (2013) | IV, SC   | 2.2          | Single      | Cattle  | P      | FLU, SOH-FLU  | Cal   |
| Odensvik and Johansson (1995) | IV, IM | 2.2          | Single      | Cattle  | P      | FLU           | Cal   |
| Kissell et al. (2016) | IV          | 2.2          | 3 days      | Cattle  | P, L, M, K | FLU, SOH-FLU  | Eval  |
| Jaroszewski et al. (2008) | IV         | 2.2          | 4 days      | Cattle  | P      | FLU, SOH-FLU  | Cal   |
| Kleinhans et al. (2016) | IV         | 2.2          | Single      | Cattle  | P      | FLU           | Cal   |
| Odensvik (1995)    | IV          | 2.2          | Single      | Cattle  | P      | FLU           | Cal   |
| **Florfenicol**    |             |              |             |         |        |               |       |
| Embrechts et al. (2013) | IM      | 22.5, 30, 15 | Single or 2 days | Swine   | P      | FLO           | Cal   |
| Jiang et al. (2006) | IV, IM, PO | 20           | Single      | Swine   | P      | FLO           | Cal   |
| Kim et al. (2008)  | IM          | 5, 20        | Single      | Swine   | P      | FLO           | Cal   |
| Li et al. (2002)   | IM          | 20           | Single      | Swine   | L, M, K, L, and P | FLO | Cal   |
| Liu et al. (2003)  | IV, IM, PO  | 20           | Single      | Swine   | P      | FLO           | Cal   |
| SPAHC (2002)       | PO (water)  | 20           | Single      | Swine   | L, K, M, and F | FLOA | Cal   |
| SPAHC (2006)       | PO          | 10           | 5 days      | Swine   | L, K, M, and F | FLOA | Eval  |
| Voorspoels et al. (1999) | IM, PO | 15, 15       | Single, 3 days | Swine   | P      | FLO           | Cal   |
| Zhang et al. (2016) | IM          | 20           | Single      | Swine   | P      | FLO           | Cal   |
| Lei et al. (2018)  | IM, IV      | 30           | Single      | Swine   | P      | FLO           | Eval  |
| Varma et al. (1986) | IV, PO      | 22           | Single      | Cattle  | S      | FLO           | Cal   |
| Sidhu et al. (2014) | SC         | 40           | Single      | Cattle  | S      | FLO           | Cal   |
| Lobell et al. (1994) | IV, IM | 20           | Single      | Cattle  | S      | FLO           | Cal   |
| Lacroix et al. (2011) | IM, SC | 40           | Single      | Cattle  | P      | FLO           | Cal   |
| Gilliam et al. (2008) | IV         | 2.2          | Single      | Cattle  | S      | FLO           | Cal   |
| de Craene et al. (1997) | IV       | 20           | Single      | Cattle  | P      | FLO           | Cal   |
| Croubels (2006)    | PO          | 20           | Single      | Cattle  | P      | FLO           | Cal   |
| Bretzlauff et al. (1987) | IV | 50           | Single      | Cattle  | P      | FLO           | Cal   |
| SPAHC (2008)       | SC          | 40           | Single      | Cattle  | L, K, M | FLOA         | Cal   |
| Norbrook Laboratories (2015) | SC     | 40           | Single      | Cattle  | L, M   | FLOA         | Eval  |
| Intervet Inc. (2009) | SC         | 40           | Single      | Cattle  | L      | FLOA         | Cal   |
| **Penicillin G**   |             |              |             |         |        |               |       |
| Ranheim et al. (2002) | IM, SC | 99           | Single      | Swine   | P      | PG           | Cal   |
| Korsrud et al. (1998) | IM         | 14.9, 65.4   | 3, 5 days   | Swine   | L, K, M, F, and P | PG | Cal   |
| Korsrud et al. (1998) | IM         | 14.9         | 3 days      | Swine   | K, M, and P | PG | Cal   |
| Lupton et al. (2014) | IM         | 32.5         | 3 days      | Swine   | P, M, and K | PG | Eval  |
| Li et al. (2019b)  | IM          | 6.5, 32.5    | 3 days      | Swine   | P, M, L, and K | PG | Cal   |
| Papich et al. (1993) | IM, SC | 23.7, 65.4 (IM); 65.4 (SC) | 5 days or single (IM), single (SC) | Cattle  | P      | PG           | Cal   |
| Korsrud et al. (1993) | IM         | 23.7, 65.4   | 5 days      | Cattle  | L, K, M, and P | PG | Cal   |
| Trolldenier et al. (1986) | SC       | 8.9          | Single      | Cattle  | P      | PG           | Cal   |
| Chiesa et al. (2006) | IM         | 6.9          | 3 days      | Cattle  | K      | PG           | Eval  |
| Djebala et al. (2021) | IM         | 20.8         | Single      | Cattle  | P      | PG           | Cal   |

Only concentration data above limits of quantification in selected studies were used for model calibration or evaluation.

Abbreviations: Cal, Calibration; Eval, Evaluation; F, fat; FLU, flunixin; FLO, florfenicol; FLOA, florfenicol amine; K, kidneys; NA, not available or not applicable; L, liver; M, muscle; P, plasma; PO, oral; PG, penicillin G; and SOH-FLU, 5-hydroxy flunixin.
Model structure. A generic PBPK model structure was designed for simulations of concentration versus time profiles for the 3 selected drugs. The model structure, based on previous PBPK models (Li et al., 2017, 2019a; Yang et al., 2019), was designed to include 2 submodels, which are capable of simulating the parent compounds: flunixin, flufenicol, and penicillin G, and their corresponding major metabolites (when applicable) (Figure 1). The major metabolites of flunixin and flufenicol, 5-hydroxy flunixin, and flufenicol amine, respectively, were specifically described in the metabolite submodel. Because the metabolism of penicillin G in food-producing animals is minimal, the liver metabolism of penicillin G was described by a simplified first-order model without monitoring the distribution of its specific metabolites in the model. The parent compound submodel was composed of major edible tissues including liver, kidney, muscle, fat (a.k.a. adipose tissue), and the rest of the body connected by a blood compartment representing circulating blood system (Figure 1). The administration routes of oral, IV, IM, and SC were incorporated into the parent drug submodel. IM and SC injections were described as 2-compartment injection site model (injection sites 1 and 2 were denoted as the fast absorption and slow releasing sites) with dissolution processes from site 2 to site 1 (Supplementary Equations 1–6) based on the previous studies (Lin et al., 2015, 2017; Yang et al., 2019). This approach divided the drug into dissolved moieties that are immediately available for absorption (fast) and undissolved drug acting as a depot that is released slowly (slow). Oral administration was described as a 2-compartment model consisting of stomach and intestines to simulate the absorption of drug in the stomach and then transportation to the intestinal tract via gastric emptying (Supplementary Equations 7–9). The same model structure without absorption routes and fat compartment was used in the metabolite submodel and connected with the parent drug model through hepatic metabolism. The elimination routes, including urine, biliary, and feces, were included in the parent drug submodel, while only urine and biliary excretion was considered in the metabolite submodel. Enterobiliary circulation was also considered in the model for all 3 drugs based on previous studies (Li et al., 2017; Lin et al., 2015; Yang et al., 2019). All the above-mentioned mathematical equations (Supplementary Equations 10–16) are described in detail in the Supplementary Data.

Model parameterization. Two different types of parameters were used in the generic PBPK model, including species-specific physiological parameters and chemical-specific parameters. The species-specific physiological parameters such as body weight (BW), cardiac output (QCC), fractions of blood flow to individual tissues (eg, the fraction of blood flow to liver, VLC) as well as the volume fractions of organs (eg, the volume fraction of liver, VLC) were collected from a recent comprehensive review article (Lin et al., 2020) where these parameters were collected and summarized from published experimental data. All species-specific physiological parameters are summarized in Table 2. Chemical-specific parameters consisted of protein binding parameters, absorption rate constant, elimination rate constant, metabolic rate constant, and partition coefficients. Protein binding parameters were obtained from experimental studies in cattle and swine (Adams et al., 1987; Galbraith and McKellar, 1996; Peterson, 1978) or the fitting values from previous PBPK models (Buur et al., 2006; Li et al., 2017, 2019b; Yang et al., 2019). Absorption rate, metabolic rate, urinary, fecal, and biliary elimination rate constants were collected from previous PBPK models for the 3 selected medications (Li et al., 2017, 2019b; Yang et al., 2019). Tissue-to-plasma partition coefficient parameters were predicted using mechanism-based in silico models (further described below). These parameter values were used as initial values in the model calibration and optimized with measured pharmacokinetic data (Table 1). All collected and optimized chemical-specific parameters are provided in Table 3 for cattle and Table 4 for swine.

Estimation of partition coefficients in cattle and swine. In order to estimate tissue-to-plasma partition coefficients for drugs without experimentally measured values, a compiled integrative algorithm consisting of 5 frequently used mechanistic equations (Berezhkovskiy, 2004; Poulin and Theil, 2002; Rodgers et al., 2005; Rodgers and Rowland, 2006; Schmitt, 2008) was developed and incorporated into the PBPK model. The tissue composition data in cattle and swine (Aksu et al., 2017; Haritova and Fink-Gremmels, 2010; Poulin et al., 2019; Utsey et al., 2020) and physicochemical properties (eg, logP and pKa) for flunixin, flufenicol, and penicillin G were incorporated into the model to estimate the partition coefficients for each of the tissues in each species. Then the average predicted tissue-to-plasma partition coefficients from the 5 different models were used as initial values in the model calibration by fitting to available pharmacokinetic datasets. The detailed equations for all these methods (Supplementary Equations 17–21), tissue composition data in both cattle and swine (Supplementary Tables 1 and 2), and physicochemical properties of the 3 selected drugs (Supplementary Table 3) are provided in the Supplementary Data.

Model calibration and evaluation. Following model parameterization, the generic PBPK model was further calibrated with measured pharmacokinetic raw concentration data (ie, untransformed data) using the Levenberg-Marquardt least-squares algorithm (Chou and Lin, 2019) implemented in the R package FME. Briefly, the parameters \( \theta \) were calibrated by comparing the predicted values \( f(t_j, \theta) \) with the mean measured values \( y_{ij} \) at time point \( i \) of the data set \( j \). The least-squares function between the observed and simulated values was estimated as

\[
\bar{\theta} = \arg \min \sum_{j=1}^{n_j} \frac{(f(t_j, \theta) - y_{ij})^2}{w_{ij} \times \eta_j}, \tag{1}
\]

where the \( w_{ij} \) is a weighting factor that normalizes the difference of units or magnitudes from a variety of data sources. The \( w_{ij} \) can be estimated from the standard deviation of measured values. \( \eta_j \) represents the number of data points for data set \( j \) which was used to scale the residuals to prevent abundant data set dominating the analysis. The model was fitted with pharmacokinetic data sets simultaneously and the optimized parameters \( \bar{\theta} \) were estimated when the minimum of sum of squared residuals for all data points from all data sets.

With the optimized parameter values, the evaluation of the PBPK model was conducted by comparing model simulations with independent pharmacokinetic data sets (ie, data not used in model calibration). On the basis of the PBPK modeling guidelines from World Health Organization (WHO, 2010) and Organization for Economic Co-operation and Development (OECD, 2021), the model was considered to be reasonable and acceptable when the simulation results matched the pharmacokinetic profiles and were generally within a 2-fold difference of observed values. The global evaluation of model fit between
log-transformed values of experimentally observed and model-predicted drug concentrations in plasma and tissues was further analyzed to determine the coefficient of determination ($R^2$).

Sensitivity analysis. Sensitivity analysis can be used to identify sensitive parameters that have high impacts on the model outputs (eg, selected dose metrics). In this study, a local sensitivity analysis was performed to determine which parameters were most influential on the 24-h area under curves (AUC) of liver, kidney, and muscle concentrations of flunixin, florfenicol, and penicillin G in cattle and swine following respective label use regimens. By increasing 1% of each parameter value and calculating the corresponding AUCs, normalized sensitivity coefficients (NSCs) were estimated to evaluate the relative sensitivity of each parameter on the corresponding AUC using the equation reported previously (Chou and Lin, 2021; Lin et al., 2013).

**Figure 1.** A schematic diagram of the generic PBPK model for flunixin, florfenicol, and penicillin G in cattle and swine. A metabolite submodel was included to account for the main metabolite when needed, such as 5-hydroxy flunixin for flunixin and florfenicol amine for florfenicol. Four different administration routes, including oral, IV, IM, and SC administrations are presented in the model. A 2-compartment absorption model (fast/slow) was used to describe the IM and SC injections. The parameter Frac is the fraction of the drug that is readily available for fast absorption (unitless) after IM or SC injection.

**Table 2.** Physiological Parameters Values Used in the Generic PBPK Model in Cattle and Swine

| Parameter                              | Abbreviation | Cattle       | Swine       |
|----------------------------------------|--------------|--------------|-------------|
| Cardiac output (L/h/kg)                | QCC          | 5.45 (1.47)  | 8.7 (1.62)  |
| Hematocrit                             | Htc          | 0.378 (0.046)| 0.412 (0.05)|
| Blood flow (fraction of cardiac output, unitless) |               |              |             |
| Liver                                  | QLC          | 0.44 (0.25)  | 0.273 (0.082)|
| Kidney                                 | QKC          | 0.11 (0.08)  | 0.114 (0.032)|
| Muscle                                 | QMC          | 0.28 (0.09)  | 0.342 (0.306)|
| Fat                                    | QFC          | 0.08 (0.024) | 0.128 (0.038)|
| Rest of body                           | QRestC       | 1-QLC-QKC-QMC-QFC | 1-QLC-QKC-QMC |
| Rest of body for metabolites           | QRestC1      | 1-QLC-QKC-QMC |             |
| Tissue volume (fraction of BW, unitless) |               |              |             |
| Plasma                                 | VPC          | 0.0399 (0.0068)| 0.0412 (0.0046)|
| Liver                                  | VLC          | 0.0122 (0.0018)| 0.0204 (0.0033)|
| Kidney                                 | VKC          | 0.0021 (0.0005)| 0.0037 (0.0011)|
| Muscle                                 | VMC          | 0.361 (0.1173)| 0.3632 (0.0266)|
| Fat                                    | VFC          | 0.1218 (0.0506)| 0.1544 (0.0265)|
| Rest of body                           | VRestC       | 1-VLC-VKC-VMC-VFC |            |
| Rest of body for metabolites           | VRestC1      | 1-VLC-VKC-VMC |             |

All parameter values in both cattle and swine were collected from Lin et al. (2020b), except QFC in cattle and swine and QLC in swine were collected from Li et al. (2017). The value was expressed as “mean (SD).”
The detailed equation is provided in the Supplementary Data (Supplementary Equation 22).

Establishment of a population PBPK model. Monte Carlo simulations were incorporated into the generic PBPK model to obtain population-based simulations based on repeated random sampling from the designated distribution of each parameter. Each Monte Carlo simulation contained 1000 iterations (ie, 1000 animals). For these simulations, all physiological and chemical-specific parameters were randomly sampled around mean values of the specified distributions, and their variability was defined by coefficients of variance (CVs). Based on the default assumptions reported in previous PBPK modeling studies (Henri et al., 2017; Li et al., 2017; Yang et al., 2015), log-normal distributions were assumed for all chemical-specific parameters, and the physiological parameters (except blood flow fractions) were assumed to follow normal distributions. Due to the high variability of blood flow fractions in cattle and swine collected from experimental data (Lin et al., 2020b), the log-normal distribution was used to avoid producing negative values. CV values for most of the physiological parameters including the BW, cardiac output, and tissue volume fractions of liver and kidneys, and the fractions of blood flows in liver were calculated based on the experimental data from a comprehensive review article (Lin et al., 2020b). For the other physiological parameters whose CV values were unknown, a default CV of 30% was used. For chemical-specific parameters, based on previous studies, a CV of 20% was assigned to tissue-to-plasma partition coefficients (Henri et al., 2017; Li et al., 2017; Yang et al., 2015), 30% for absorption, metabolic, and elimination rate constants (Li et al., 2018; Table 3. Chemical-Specific Parameter Values Used in the PBPK Model for Flunixin, Florfenicol, and Penicillin G and in Cattle

| Parameter                                      | Symbol        | Flunixin | Florfenicol | Penicillin G |
|------------------------------------------------|---------------|----------|-------------|--------------|
| Absorption rate constant (/h)                  |               |          |             |              |
| IM administration                              | Kim           | 1a       | 0.16b       | 0.10*        |
|                                               | Fracim        | 0.71*    | 0.65*       |              |
|                                               | Kdissim       | 0.01b    | 0.001*      |              |
| SC administration                              | Ksc           | 0.4a     | 0.12b       | 0.04*        |
|                                               | Fraccsc       | 0.56*    | 0.76*       |              |
|                                               | Kdisssc       | 0.005*   | 0.005*      |              |
| Molecular weight                               |               |          |             |              |
| Parent compounds                               |               | 296.4d   | 358.2d      | 334d         |
| Metabolites                                    |               | 312.4d   | 247.3d      |              |
| Tissue-to-plasma partition coefficient for parent compound (unitless) |
| Liver                                          | PL            | 2.19*    | 1.98*       | 1.09*        |
| Kidney                                         | PK            | 3.38*    | 0.91*       | 1.98*        |
| Muscle                                         | PM            | 0.43*    | 1.33*       | 0.20*        |
| Fat                                            | PF            | 0.56*    | 0.61*       | 0.04*        |
| Rest of body                                   | PRest         | 6.74*    | 0.12*       | 7.99*        |
| Tissue-to-plasma partition coefficient for parent metabolite (unitless) |
| Liver                                          | PL1           | 3.11f    | 7.59*       |              |
| Kidney                                         | PK1           | 4.59*    | 1.30f       |              |
| Muscle                                         | PM1           | 2.96f    | 0.90f       |              |
| Rest of body                                   | PRest1        | 8.04f    | 0.09*       |              |
| Hepatic metabolic rate constant (/h/kg)         | KmetC         | 0.005*   | 0.23*       | 0.25*        |
| Rate constant for enterohepatic circulation (/h/kg) | KehcC        | 0.012*   | 0.13*       | 0.0004*      |
| Free fraction of chemical in plasma (unitless)  |
| Parent compound                                | fr            | 0.15*    | 0.79*       | 0.66*        |
| Metabolites                                    | fr1           | 0.008*   | 0.71b       |              |
| Biliary elimination rate (L/h/kg)               | KbileC        | 0.51*    | 0.11*       | 0.63*        |
| Metabolites                                    | KbileCl       | 0.58*    | 0.16*       |              |
| Urinary elimination rate constant (L/h/kg)      | KurineC       | 0.50*    | 0.32*       | 0.825*       |
| Metabolites                                    | KurineCl      | 0.068*   | 0.002*      |              |
| Intestinal absorption rate constant (/h)        | Kabs          | 0.4a     | 1.9b        | 1.9b         |
| Fecal elimination rate constant (/h)            | Kunabs        | 0.81*    | 0.009*      | 0.51*        |

*Li et al. (2019a).  
**Yang et al. (2019).  
*Li et al. (2017).  
*PubChem.  
*Assumed to be equal to the value of parent compounds  
**Predicted value by using an in silico model (described in the Materials and Methods section)  
*Assumed same as the value of florfenicol model from the previous study (Yang et al., 2019).  
*The parameter values in bold were obtained through model fitting. Regarding the model fitting, refer to the Materials and Methods section for further information on which datasets were used to estimate values for these parameters.
Yang et al., 2015; Zeng et al., 2019), and 10% for protein binding constants (Li et al., 2017; Riad et al., 2021). Note that previous studies have shown that the CV values for plasma protein binding percentages of selected drugs were generally within 10% (Adams et al., 1987; Galbraith and McKellar, 1996; Peterson, 1978), so the use of the default CV value of 10% was conservative enough. The 2.5th and 97.5th percentiles of each parameter were calculated as the upper and lower bounds to ensure the values were biologically plausible for each of parameters. In addition, all physiological parameters were adjusted to ensure that the sum of tissue volumes or the sum of fractions of blood flows were equal to 1 after random sampling from defined distributions to avoid unbalance of the PBPK model.

Estimation of WDIs using the population PBPK model. The population PBPK model was used to account for population variability and parameter uncertainty and to generate population simulation results of flunixin, florfenicol, and penicillin G concentrations in plasma and edible tissues in diverse populations of cattle and swine following label or extralabel dosing regimens in order to calculate WDIs. In this manuscript, FDA-approved withdrawal time (a.k.a. FDA-approved withdrawal period) refers to the time needed after label administration of a drug for tissue residue concentrations to decrease below tolerances determined using the 99th percentile tolerance limit method with a 95% confidence based on FDA guidance (FDA, 2018); whereas the term WDI is used to refer to the time for tissue residue concentrations to fall below tolerances estimated using other methods, such as the present population PBPK models and it is typically used when a drug is given in an extralabel manner. PBPK model-predicted WDIs in edible tissues were determined to be the time when the 99th percentiles of the simulated drug concentrations in target tissues fell below the tolerance in the United States or maximum residue limit from the European Medicines Agency of the drug in the corresponding tissue. For penicillin G, the tolerance for all edible tissues in cattle is 0.05 ppm or $\text{g/g}$. The tolerance of penicillin G in swine is zero, so the minimum level of applicability of 0.025 $\mu\text{g/g}$ (25 ng/g) from the USDA Food Safety and Inspection Services (USDA, 2013) was used for all edible tissues in swine. The tolerance of

### Table 4. Chemical-Specific Parameter Values Used in the PBPK Model for Flunixin, Florfenicol, and Penicillin G and in Swine

| Parameter | Symbol | Flunixin | Florfenicol | Penicillin G |
|-----------|--------|----------|-------------|--------------|
| Absorption rate constant (/h) | | | | |
| IM administration | Kim | $1^a$ | 0.21 | 0.10 |
| | Fracim | | 0.56 | 0.55 |
| | Kdissim | | $0.0253^a$ | 0.007 |
| SC administration | Ksc | 0.4 | 0.126 | 0.25 |
| | Fraccsc | | 0.50 | 0.5 |
| | Kdisssc | | 0.005 | 0.005 |
| Tissue-to-plasma partition coefficient for parent compound (unitless) | | | | |
| Liver | PL | 2.30 | 0.39 | 0.07 |
| Kidney | PK | 6.99 | 3.47 | 1.43 |
| Muscle | PM | 0.31 | 1.3 | 0.08 |
| Fat | PF | 0.6 | 0.15 | 0.25 |
| Rest of body | PRest | | | 0.46 |
| Tissue-to-plasma partition coefficient for parent metabolite (unitless) | | | | |
| Liver | PL1 | | 14.7 |
| Kidney | PK1 | | 5.98 |
| Muscle | PM1 | | 1.21 |
| Rest of body | PRest1 | | 0.49 |
| Hepatic metabolic rate constant (/h/kg) | KmetC | 0.004 | 0.0075 | 0.59 |
| Rate constant for enterohepatic circulation (/h/kg) | KehcC | 0.004 | 0.17 | 0.01 |
| Free fraction of chemical in plasma (unitless) | FR | 0.13 | 0.59 | 0.96 |
| Metabolites | FR1 | | 0.78 |
| Biliary elimination rate (L/h/kg) | Kbiuc | 0.401 | 0.03 | 0.52 |
| Metabolites | Kbiuc1 | 0.034 | 5.98E-4 |
| Urinary elimination rate constant (L/h/kg) | KurineC | 0.253 | 0.43 | 1.54 |
| Metabolites | KurineC1 | 0.007 | 0.01 |
| Intestinal absorption rate constant (/h) | Kabs | 0.4 | 1.9 | 1.9 |
| Fecal elimination rate constant (/h) | Kunabs | 0.5 | 0.11 | 0.81 |

*aLi et al. (2019a).
*bLi et al. (2017).
*cYang et al. (2019).
*dPredicted value by using the in silico model (described in Materials and Methods section).
*eAssumed same as the value of florfenicol model from the previous study (Yang et al., 2019).

*The parameter values in bold were obtained through model fitting. Regarding the model fitting, refer to the Materials and Methods section for further information on which datasets were used to estimate values for these parameters.
Development of an interactive generic PBPK model web interface. An ordinary differential equation solver R package “mrgsolve” (Baron and Gastonguay, 2015) was used to solve the differential equations in the PBPK model code and was used in the final individual and population generic PBPK model. Then, the final generic PBPK model was converted to a web-based interactive generic PBPK (igPBPK) interface with the “Shiny” package in R program based on our recently published methods (Li et al., 2019a; Lin et al., 2020a; Riad et al., 2021).

Code availability and result reproducibility. All raw data that were used to calibrate and evaluate the present PBPK model and all model code, including the code files for the igPBPK web interface, the code files that were used to generate the results presented in figures, and the code files that were used to estimate tissue/plasma partition coefficients, as well as other relevant model code files along with additional instructions are available in the GitHub (https://github.com/UFPBPK/FARAD-igPBPK). These raw data and source code files will allow readers to reproduce our results and apply our model for further research.

RESULTS

Model Calibration and Evaluation

The generic PBPK model (Figure 1) was calibrated with concentrations of the 3 selected drugs and their metabolites in plasma and edible tissues following different exposure regimens corresponding to previous pharmacokinetic studies (Table 1). A global evaluation of goodness of fit was performed by comparison between model predictions and observed values for each drug (Figures 2A–C) and evaluating predicted-to-observed (P/O) ratio versus model prediction plots (Figures 2D–F), respectively. The model adequately simulated the calibration and evaluation datasets with the ranges of estimated coefficient of determination ($R^2$) of 0.77–0.80 for flunixin and 5-hydroxy flunixin, 0.72–0.77 for florfenicol and florfenicol amine, and 0.78–0.80 for penicillin G (Figures 2A–C). The percentage of the predictions within 2-fold errors was 51.3%, 68.1%, and 57.3% for flunixin and 5-hydroxy flunixin (Figure 2D), florfenicol and florfenicol amine (Figure 2E), and penicillin G (Figure 2F) in cattle and swine, respectively. The range of the percentages within 3-fold errors was 71.7%–85.2% (Figures 2D–F). These results showed that more than 50% and 70% of model predictions were within a 2-fold and 3-fold factor of measured data, respectively.

Simulated results from the calibrated PBPK model were compared with measured concentrations from individual published studies in cattle and swine administered flunixin, florfenicol, or penicillin G through oral, IV, IM, or SC routes (representative results shown in Figure 3 for penicillin G and other results are shown in Supplementary Figures 1 and 2). The time-course comparisons for penicillin G in swine and cattle for the calibration data (Figures 3A1–C3 for swine and Figures 3D5–F6 for cattle) showed that the model generally matched the kinetic profiles of penicillin G in plasma and edible tissues in both species. In addition, as shown in Figures 3A1–D4 and Figures 3H1 and 3H2 for evaluation data, model simulations for plasma, liver, kidney, and muscle of penicillin G were in agreement with most of the datasets. Overall, results from both analyses suggest that the model adequately simulates the observed data sets used for model calibration and evaluation.

Sensitivity Analysis

Thirty-eight parameters were included in the local sensitivity analysis based on the dose metrics of 24-h AUCs of liver, kidney, and muscle concentrations of flunixin, 5-hydroxy flunixin, florfenicol, florfenicol amine, and penicillin G in cattle and swine following FDA-approved label use administration (Figures 4 and Supplementary Tables 4–7). Figure 4 is a representative figure displaying the absolute percentage of NSCs based on 24-h AUCs for muscle concentrations of flunixin, florfenicol, and penicillin G in cattle (Figure 4A) and swine (Figure 4B). The complete sensitivity analysis results for all edible tissues (eg, liver, kidney, and muscle) and all drugs with their metabolites (flunixin, 5-hydroxy flunixin, florfenicol, florfenicol amine, and penicillin G) can be found in Supplementary Tables 4–7. The results showed that the AUCs of the muscle for flunixin was highly sensitive to partition coefficient of muscle (PM), biliary excretion rate constant (KbileC), and fraction of blood flow to kidney (QKC) in both cattle (Figure 4A) and swine (Figure 4B). Similar high impacts on the AUCs of penicillin G were observed for the parameters of PM and the fraction of penicillin G immediately available for absorption (Fracim) via IM injection in both cattle and swine. The fraction of florfenicol immediately available for absorption (Fracim) via IM injection, urine elimination rate constant (KurineC), and PM was highly sensitive in the cattle model (Figure 4A), while the percentage of free drug (FB), cardiac output (QCC), blood flow to the kidney (QKC), KurineC, and PM was sensitive in the swine model (Figure 4B). For cattle, the QKC, VRestC (volume fraction of rest of body), and KurineC were the common sensitive parameters on AUCs of the selected tissues across flunixin, 5-hydroxy flunixin, florfenicol, florfenicol amine, and penicillin G (Supplementary Tables 4 and 6). In the swine model, the QKC and KurineC both were sensitive parameters on AUCs of all edible tissues for all drugs (Supplementary Tables 5 and 7). In addition, the partition coefficients of liver, kidney, and muscle were highly sensitive parameters (NSC > 0.5) on the prediction of AUCs in respective tissues for either cattle or swine model (Supplementary Tables 4–7).

Monte Carlo Simulation and WDI Estimation

The population PBPK model coupled with the Monte Carlo sampling technique was used to estimate WDIs for flunixin, florfenicol, and penicillin G (Figures 5 and 6 and Table 5). All physiological and chemical-specific parameters were involved in the population analysis. Based on the simulation after FDA-approved label use in cattle (Figure 5), the concentrations of flunixin and florfenicol amine via IM injection depleted the slowest in muscle (Figures 5C and 5F), while the penicillin G concentration was decreased the slowest in liver (Figure 5G). For swine (Figure 6), the flunixin and penicillin G concentrations depleted the slowest in the kidney (Figures 6B and 6H), and the florfenicol amine concentration was decreased the slowest in the muscle (Figure 6F). The longest WDIs among tissues for flunixin in cattle is 0.125 μg/g for liver and 0.025 μg/g for muscle (FDA, 1998), and in swine is 0.03 μg/g for liver and 0.025 μg/g for muscle (FDA, 2005). Because there is no tolerance for flunixin in kidney for both cattle and swine, the tolerances for liver were used as a surrogate. The marker residue of florfenicol is the main metabolite florfenicol amine and the tolerance for florfenicol amine is 3.7 μg/g for liver and 0.3 μg/g for muscle in cattle, and 2.5 and 0.2 μg/g for liver and muscle in swine, respectively (FDA, 2020). The tolerances of florfenicol amine in liver were used for kidney due to lack of published tolerances in kidney of cattle and swine.
flunixin, florfenicol (after IM), florfenicol (after SC), and penicillin G in cattle and swine after FDA-approved label or extralabel dosing regimens were chosen to be the model-derived WDIs to compare with withdrawal times approved by regulatory agencies (Table 5).

The FDA-approved label dose regimens were obtained from the Veterinarian’s Guide to Residue Avoidance Management (VetGRAM) (Riviere et al., 2017) and are listed in Supplementary Table 8. The estimated WDIs for flunixin, florfenicol (after IM), florfenicol (after SC), and penicillin G following FDA-label approved dosing regimens were 8 (muscle), 28 (muscle), 39 (muscle), and 7 (liver) days in cattle, respectively, while they were 14 (kidney), 16 (muscle), and 11 (kidney) days for flunixin, florfenicol, and penicillin G in swine (Figures 5 and 6 and Table 5). The predicted WDI following FDA-approved label dosing regimens in cattle were equal or close to FDA-approved label withdrawal times for florfenicol after IM administration (predicted WDI: 28 vs FDA-approved withdrawal time: 28 days) and flunixin after SC administration (predicted WDI: 39 vs FDA-approved withdrawal time: 38 days), but were relatively longer for flunixin (predicted WDI: 8 vs FDA-approved withdrawal time: 4 days) and penicillin G (predicted WDI: 7 vs FDA-approved withdrawal time: 4 days). For swine, the predicted WDIs were both close to the FDA-approved withdrawal times for flunixin (predicted WDI: 14 vs FDA-approved withdrawal time: 12 days), florfenicol (predicted WDI: 16 vs FDA-approved withdrawal time: 16 days), but were somewhat longer for penicillin G (predicted WDI: 11 vs FDA-approved withdrawal time: 6 days). From a food safety perspective, estimating longer WDIs is appropriate to minimize exposure to violative residues.

The extralabel dosing regimens for the 3 selected drugs were based on the internal WDI request data from 2019 to 2021 from FARAD Regional Call Centers, and are listed in Supplementary Table 9. Following these common extralabel dosing regimens for cattle and swine, the predicted WDIs were 8 (muscle), 29 (muscle), 48 (muscle), and 43 (liver) days for flunixin, florfenicol (after IM), florfenicol (after SC), and penicillin G (Table 5). For swine, the predicted WDIs were 22 (kidney), 22 (muscle), and 25 (kidney) days, respectively (Table 5). The model-predicted WDIs in cattle and swine following extralabel dosing regimens were longer than FDA-approved label withdrawal times for all drugs (Table 5).

User-Friendly Interface Establishment

The final generic PBPK model was converted into a web-based interactive igPBPK platform using the R Shiny package in R language and published online at the link: http://pbpk.shinyapps.io/igPBPKApp. A screenshot of this interface is shown in Figure 7. Users can input parameter values for each therapeutic regimen (e.g., administration route, dosage, number of doses, and dosing interval) to the igPBPK interface to predict the concentrations of flunixin, florfenicol, and penicillin G and/or their metabolites in edible tissues. The interface will generate a detailed report of the simulation results and the predicted WDI based on the tolerance of the drug in a particular species and based on the simulated drug concentrations in the tissues for the defined therapeutic regimen. The results will be generated in real time (i.e., a few seconds). A detailed tutorial on how to launch and use this igPBPK interface is provided in the Supplementary Data file.
This study reports a new igPBPK modeling web interface that can be used to simulate tissue residues and estimate WDIs of drugs with diverse physicochemical and pharmacokinetic properties following different routes of administration (oral, IV, IM, and SC) in cattle and swine. Another novelty of this igPBPK interface is that all partition coefficients were estimated using in silico mechanistic equations. The strengths of this approach are that these parameters do not rely on animal tissue data to...
estimate their values and these mechanistic equations can be applied to other drugs. By integrating Monte Carlo simulations into the framework, the igPBPK model platform can generate health protective population-based simulation results to estimate WDIs for the 3 selected drugs following different exposure paradigms (both FDA-approved label and extralabel uses). This igPBPK interface serves as a useful tool to answer extralabel WDI inquiries in order to help ensure safety of animal-derived food products. This igPBPK model platform can be extended to other drugs in other species to help address food safety issues for the United States and other countries.

Several PBPK models for specific veterinary drugs (eg, florfenicol, monensin, penicillin G, and oxytetracycline) have been developed in recent years and used as a tool to facilitate drug WDI estimations (Henri et al., 2017; Law, 1999; Li et al., 2019a, 2019b; Riad et al., 2021; Yang et al., 2019; Zhou et al., 2021). Some of these existing models have been converted to web-based user-friendly interactive PBPK interfaces, such as oxytetracycline (Riad et al., 2021) and flunixin (Li et al., 2019a). However, existing food animal PBPK models are typically limited to 1 specific drug for 1 model, and it requires extensive data, including tissue residue depletion data to adapt each model for each drug. This approach is time- and resource-intensive and it is an impossible task to develop a new model for each of the veterinary drugs in each species because there are so many drugs in different food animal species. Additionally, when new data become available, it is difficult to formulate new models. In this study, we developed an igPBPK platform that allows users without any modeling experience to quickly implement PBPK models. By incorporating tissue-composition-based mechanistic equations into the model, the critical model parameters (ie, partition coefficients) can be predicted based on the physicochemical properties of the drug and based on the physiology of the animal. This is particularly important for drugs with limited data as it no longer requires tissue residue depletion data to estimate partition coefficients. This makes it possible to develop PBPK models for drugs with sparse data, with ready extrapolation to other drugs. In addition, compared with previous PBPK models for the 3 selected drugs, the present model is more robust as it was adapted and evaluated with additional new pharmacokinetic datasets (Bates et al., 2020; Djebala et al., 2021; Kittrell et al., 2020; Li et al., 2019b). We expect that the concept of an interactive generic PBPK model platform will accelerate the development of the next generation of PBPK models in food animals to predict tissue residues and WDIs of animal drugs, including drugs with minimal pharmacokinetic data.

The present population PBPK model can be used to estimate WDIs based on FDA-approved label and extralabel use scenarios for the 3 selected drugs in cattle and swine. Model-predicted WDI for the 3 selected drugs in cattle and swine at the FDA-approved label doses, based on respective tolerances, is close to or equal to FDA-approved label withdrawal times (Table 5). For the common extralabel use of flunixin in cattle and swine (ie, 3 repeated IM injections at the dosage of 2.2 mg/kg with a 24-h dosing interval), the predicted WDI in cattle (8 days) is 4 days more than the FDA-approved withdrawal time (4 days for 3 repeated IV injections at a dosage of 2.2 mg/kg with a 24-h dosing interval), while the predicted WDIs in swine (22 days) is much
longer than the FDA-approved withdrawal time (12 days for a single IM injection at a dosage of 2.2 mg/kg). The predicted WDIs for penicillin G in cattle and swine models following extralabel use [43 days (5 × label dose) in cattle and 25 days (5 × label dose) in swine] are both much longer than FDA-approved label withdrawal times (4 days for cattle and 6 days for swine). The results indicate that the predicted WDIs for the extralabel doses from the current model are more conservative, and thus more protective, than withdrawal times that are based on the label dose. These results highlight the importance to use a scientific-based tool to estimate WDIs for different extralabel uses in order to avoid violative tissue residues of drugs in edible tissues of food animals.

Compared with the previous penicillin G and flunixin PBPK models (Halleran et al., 2022; Li et al., 2017, 2019a, 2019b), when based on the label use, the predicted WDIs from the current
generic PBPK model in cattle (7 and 8 days for penicillin G and flunixin, respectively) and swine (11 and 14 days) were around 14%-45% different from the results in the previous models (ie, 5 and 6 days in cattle; 6 and 16 days in market-age swine). Based on the 5× label dose of penicillin G in swine, the current model-predicted WDI of 25 days was longer than ~9 days from the previous PBPK model for market-age swine (Halleran et al., 2022; Li et al., 2017) and shorter than 38 days from the previous PBPK model for heavy sows (Li et al., 2019b). The current population PBPK model was established based on not only the available datasets used in previous studies, but also new pharmacokinetic studies (Bates et al., 2020; Djebala et al., 2021; Kittrell et al., 2020; Li et al., 2019b) published after the development of the earlier PBPK models. Of note, the pharmacokinetic data for penicillin G from both market-age swine (Li et al., 2017) and heavy sows (Li et al., 2019b; Lupton et al., 2014) were incorporated into the present model to better capture the population variability. Consequently, the results from the current PBPK model for penicillin G in swine is somewhere between results from previous models for each of these production classes. Also, all physiological parameters in cattle and swine have been updated based on a recent comprehensive review article on physiological parameters for PBPK modeling in cattle and swine (Lin et al., 2020b), and variabilities of the fractional blood flows to liver and kidney in swine were larger than those in the earlier study (Li et al., 2017). Therefore, our model includes more variability and uncertainty from more comprehensive pharmacokinetic datasets and physiological parameters, resulting in a broad range of parameter values used in Monte Carlo simulations, which in turn contributes to more rigorous and conservative estimates of WDIs compared with the previous market-age swine model for penicillin G. It should be noted that these data represent penicillin G formulation with procaine only, and not longer-acting formulations complexed with procaine and benzathine.

For flunixin, commonly used extralabel administrations are 3 repeated IM injections at 20 mg/kg with 48-h intervals and 3 repeated SC injections at 40 mg/kg with 96-h intervals in cattle and 2 repeated SC injections at 40 mg/kg with a 96-h interval in swine (Supplementary Table 9). The predicted extralabel WDIs were longer than FDA-approved label withdrawal times for SC administration in cattle (eg, predicted WDI was 48 days for 3 repeated SC doses at 40 mg/kg with 96-h intervals vs FDA-approved label withdrawal time of 38 days for a single SC dose) and in swine (eg, predicted WDI was 22 days for 2 repeated SC doses at 40 mg/kg with 96-h intervals vs FDA-approved label withdrawal time of 16 days for 5 daily dose via drinking water at 100 ppm), but it was close to FDA-approved label withdrawal times in the cattle model via IM injection (eg, predicted extralabel WDI 29 days vs label withdrawal time 28 days). These results indicate that violative residues might result following extralabel SC administrations in cattle and swine if the label withdrawal times are not extended substantially.

There are several limitations to this study. First, although the present PBPK model has successfully incorporated in silico mechanistic equations to predict partition coefficients of drugs based on physicochemical properties and tissue composition, the model does not include the prediction equations to estimate other critical model parameters such as the parameters related to protein binding, hepatic or renal clearance, and metabolism (Kamiya et al., 2019; Schneckener et al., 2019). Second, the development of the current PBPK model was mostly based on available pharmacokinetic data in market-age swine and adult cattle with only a few datasets in piglets, heavy sows, and calves (Li et al., 2019b; Lupton et al., 2014; Ranheim et al., 2002; Trollødenier et al., 1986). Although the simulation results adequately correspond to pharmacokinetics of all 3 selected drugs in different age groups of selected animal species, our model is not capable of considering the differences between ages, sexes, and production classes. Additional data, especially the physiological and anatomical data in different ages and sexes, can be incorporated into the model to enhance the predictability and ensure the model is as close to reality as possible. Recently, our group has published a series of studies to establish a database of physiological parameters for developing PBPK models for drugs and environmental chemicals in different food-producing animals, including cattle and swine (Lin et al., 2020b), chickens and...
turkeys (Wang et al., 2021), and sheep and goats (Li et al., 2021). Based on this physiological parameter database, the present igPBPK modeling platform can be extended to other food animal species in the future. Third, while the CV values of chemical-specific values were based on default assumptions that are generally acceptable to be conservative in the field of PBPK modeling (Henri et al., 2017; Li et al., 2017; Riad et al., 2021; Yang et al., 2015), there are inevitably some uncertainty associated with these parameters. Additional experimental studies that directly measure the CV values of these parameters will help improve the present model. Fourth, the present study only did a local sensitivity analysis, which does not consider interactions between parameters. Global sensitivity analysis (Hsieh et al., 2018; McNally et al., 2011; Tardiveau et al., 2022) should be performed in order to assess the relative sensitivities of model parameters and their interactions in the future. Finally, the current model was developed so that it could predict WDIs for 3 different drugs when given individually, but it does not specifically predict WDIs when these 3 drugs are given concurrently because the model does not account for potential drug-drug interactions.

CONCLUSIONS

This study reports a new interactive generic PBPK modeling platform for multiple drugs in cattle and swine that can be used in real-time field exposure scenarios. This modeling platform has been implemented to build PBPK models for 3 representative drugs (flunixin, florfenicol, and penicillin G) in both species. Model simulations, in general, are in good agreement with the observed concentrations of flunixin, 5-hydroxy flunixin, florfenicol, florfenicol amine, and penicillin G residues in edible tissues of cattle and swine after different exposure regimens. Predictions of WDIs for FDA-approved label use and common extra-label uses of the 3 selected drugs using the population PBPK model with Monte Carlo simulations demonstrate the ability of the model to provide more protective WDI recommendations to help ensure safety of food products derived from animals treated with these drugs under field dosing conditions. The final generic PBPK model has been converted to a web-based igPBPK interface to provide a user-friendly platform to facilitate the application of this PBPK modeling platform for users with or without computer programming experiences. Although this model still has some limitations, the igPBPK framework represents a proof-of-concept toward the next-generation PBPK model and provides a robust foundational tool to extrapolate to other drugs and other food animal species.

SUPPLEMENTARY DATA

Supplementary data are available at Toxicological Sciences online.

DECLARATION OF CONFLICTING INTERESTS

The authors declare no conflict of interest.

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