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Title: Physiologically Based Pharmacokinetic Modeling of Vitamin D₃ and Metabolites in Vitamin D Insufficient Patients

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VitD₃, Vitamin D₃; 25D₃, 25(OH)D₃; 1,25D₃, 1,25(OH)₂D₃; 24,25D₃, 24,25(OH)₂D₃

3 Abstract

A physiologically based pharmacokinetic (PBPK) model of vitamin D₃ and metabolites [25(OH)D₃, 1,25(OH)₂D₃, and 24,25(OH)₂D₃] is presented. In this study, patients with 25(OH)D₃ plasma concentrations below 30 ng/ml were studied after a single dose of 5,000 I.U. (125 µg) cholecalciferol, provided with 5,000 I.U. daily cholecalciferol supplementation until vitamin D replete (25(OH)D₃ plasma concentrations above 30 ng/ml), and had serial plasma samples were collected at each phase for 14 days. Total concentrations of vitamin D₃ and metabolites were measured by ultra-high performance liquid chromatography tandem mass spectrometry. A nine-compartment PBPK model was built using MATLAB to represent the triphasic study nature (insufficient, replenishing, sufficient). Stimulatory and inhibitory effect of 1,25(OH)₂D₃ were incorporated by fold-changes in the primary metabolic enzymes CYP27B1 and CYP24A1, respectively. Incorporation of dynamic adipose partition coefficients for vitamin D₃ and 25(OH)D₃ and variable enzymatic reactions aided in model fitting. Measures of model predictions agreed well with data from metabolites, with 97%, 88%, and 98% of the data for 25(OH)D₃, 24,25(OH)₂D₃, and 1,25(OH)₂D₃, respectively, within 2-fold of unity (*fold error* values between 0.5 and 2.0). Bootstrapping was performed and optimized parameters were reported with 95% confidence intervals. This PBPK model could be a useful tool for understanding the connections between vitamin D and its metabolites under a variety of clinical situations.

4 Significance Statement

This study developed a physiologically based pharmacokinetic (PBPK) model of vitamin D₃ and metabolites for patients moving from an insufficient to a repleted state over a period of 16 weeks.

5 Visual Abstract

Not required

6 Introduction

Vitamin D is a fat-soluble, prohormone that plays an essential role in regulating calcium and phosphorus to maintain musculoskeletal health. The main source of vitamin D is through endogenous production of cholecalciferol (VitD₃) in the skin upon ultraviolet B exposure from the sun. The active form of VitD₃, 1,25(OH)₂D₃ (1,25D₃) has been found to exhibit many pleiotropic actions beyond calcium and phosphorus homeostasis (e.g. musculoskeletal health), that are often referred to as “non-classical” actions. Additionally, 1,25D₃ controls over 200 genes, including those responsible for the regulation of hormone secretion, immune function, and cell proliferation and differentiation [Holick, 2006].

VitD insufficiency is a global health issue and it has been estimated that up to 80% of men and women in the United States of America (USA), Canada, and Europe meet this classification [Ganji et al., 2012; van Schoor and Lips, 2018]. VitD status is based on concentrations of 25-hydroxyvitamin D, 25(OH)D (25D), given that it is the major circulating form and it is readily assayed in hospital laboratories. There is a lack of consensus on the serum concentrations of 25D considered to be adequate, but it is commonly agreed that 25D concentrations should be above 20 ng/ml (1 ng/ml = 2.5 nmol/l) [Ganji et al., 2012, van Schoor and Lips, 2018]. The Institute of Medicine (IOM) defines deficiency as 25D < 12 ng/ml and recommends a target serum concentration of 20 ng/ml [Ross, 2011]. The Endocrine Society (ENDO) defines VitD deficiency as 25D < 20 ng/ml and recommends a target concentration of 30 ng/ml [Holick et al., 2011]. However, other leading experts define VitD insufficiency as 25D concentrations between 20 and 30 ng/ml and deficiency as concentrations < 20 ng/ml [Cianferotti and Marcocci, 2012; Dawson-Hughes et al., 2005; Holick, 2007; Holick et al., 2011]. While many people with VitD insufficiency take supplements, treatment recommendations are not consistent. The IOM recommended a daily intake of 600 I.U. VitD for children, adolescents, and adults, and 800 I.U. for adults over the age of 70 to maintain 25D concentrations of 20 ng/ml [Jernigan and Andress, 2003]. ENDO recommended 1500-2000 I.U./day to prevent or treat VitD insufficiency and achieve 25D concentrations above 30 ng/ml, with a preferred range of 40-60 ng/ml [Holick et al., 2011]. While these are the common guidelines in the U.S., recommendations in other

regions of the world differ based on many population-specific factors such as sunlight exposure, skin pigmentation, clothing, and dietary practices [Haq et al., 2018; Munns et al., 2016; Pérez-López et al., 2012; Płudowski et al., 2013; Rizzoli et al., 2013; Society, 2012].

Activation of VitD occurs through two sequential hydroxylation steps to generate the metabolically active metabolite 1,25D. Vitamin D is hydroxylated in the liver primarily by the cytochrome P450 (CYP) enzyme CYP2R1, but also through pathways with CYP27A1 [Sakaki et al., 2005] and CYP3A4 [Battault et al., 2012; Wang et al., 2013], to form 25D, the major circulating metabolite. 25D can then be hydroxylated in the kidney by either CYP27B1 to the active metabolite, 1,25D, or by CYP24A1 to the inactive metabolite 24,25-dihydroxyvitamin D (24,25D). Phase 2 metabolism pathways have also been described for VitD but appear to be of lesser importance [Gao et al., 2017; Wang et al., 2014; Wong et al., 2018]. 1,25D is tightly regulated in a feedback loop with CYP27B1 and CYP24A1; high concentrations of 1,25D will respectively suppress and stimulate expressions of CYP27B1 and CYP24A1.

Despite VitD being used clinically for decades, there are a limited number of published studies that include assessments of the parent VitD compound and metabolites using pharmacokinetic or physiologically based pharmacokinetic (PBPK) approaches [Armas et al., 2004; Bailie and Johnson, 2002; Benaboud et al., 2013; Fassio et al., 2020; Hsu et al., 2021; Ilahi et al., 2008; Jetter et al., 2014; Kimura et al., 1991; Levine and Song, 1996; Meekins et al., 2014; Ocampo-Pelland et al., 2016, 2017; Ramakrishnan et al., 2016; Roth et al., 2012]. The current study sought to develop a PBPK model of VitD₃ and metabolites in VitD insufficient patients in the U.S. who were treated with moderate (5000 I.U.) daily doses of VitD₃ for up to 16 weeks to achieve replacement as defined by 25D \geq 30 ng/ml. Through establishing a model that might align well with a triphasic clinical experience of a patient—an initial visit for establishing VitD concentrations, time to achieve repletion on a dosing regimen, and a follow-up visit—this study aimed to better understand the longitudinal effects of daily supplementation of moderate doses of VitD₃.

7 Materials and Methods

7.1 Design Overview

The parent study from which the data was obtained was carried out in accordance with the Declaration of Helsinki. All subjects provided informed consent and the protocols were approved by the Institutional Review Boards at the University of Colorado and the University of Pittsburgh. The parent study (NCT02360644) was registered under ClinicalTrials.gov and can be referenced for additional details.

7.1.1 Study Participants

Healthy subjects from the University of Colorado and University of Pittsburgh with VitD insufficiency defined in the study as total 25D below 30 ng/ml and not receiving VitD replacement therapy were recruited. Other eligibility criteria included age 18-75 years, predicted compliance with study visits, not pregnant or lactating, no changes in medications within 4 weeks, no predisposition to hypercalcemia, and hemoglobin ≥ 10 g/dL. Exclusion criteria were active autoimmune disease, active or recent infections requiring antimicrobial treatment and hepatic insufficiency. Note that while the parent clinical study focused on patients with chronic kidney disease, this study utilized data from a control group of healthy study participants from the parent clinical study. Baseline characteristics for study participants ($n=11$) can be found in Table 1.

7.1.2 Clinical Study Design

Vitamin D insufficient subjects were admitted to the Clinical and Translational Research Centers (CTRC) at the University of Colorado or University of Pittsburgh for a 12 h stay, followed by visits at 24, 48, 168, and 336 h. Participants were required to be fasting at the beginning of the study. Any prescribed medications were withheld for the first two hours of the study. Subjects were given a single 5,000 I.U. (125 μ g) oral dose of VitD₃ (Jarrow Formulas, Los Angeles, CA). Serial blood samples (7.5 ml) were collected at baseline and at 0.5, 1, 2, 4, 8, 12, 24, 48, 168, and 336 h into heparinized vacutainers. Blood samples were centrifuged immediately following collection for 10 min at 3000 xG at 4°C and plasma samples were stored at -80°C until analysis. After the 336 h

blood draw, participants were given up to 16 weeks of daily supplementation of 5000 I.U. VitD₃ to attain replete concentrations (≥ 30 ng/ml). At the repleted study phase, participants were given a final dose of 5,000 I.U. VitD₃ followed by collection of blood samples as previously described for up to 336 h. The total study reflected three phases—the insufficient phase, the replenishing phase, and the repleted phase. Data are available for subjects in the insufficient and repleted phases.

7.1.3 Analytical Assay

Total concentrations of VitD₃ and metabolites [25D₃, 1,25D₃, and 24,25D₃] were determined by a novel ultra-high performance liquid chromatography-tandem mass spectrometry (UHPLC-MS/MS) assay capable of detecting all four analytes simultaneously as previously described [Stubbs et al., 2014] with minor modifications. UHPLC was performed with a Waters Acquity UPLC I-class (Waters, Milford, MA, USA), which includes a sample manager and a binary solvent manager. Briefly, 500 μ l samples were precipitated with acetonitrile, extracted with methyl tert butyl ether, then derivatized with 4-phenyl-1,2,4-triazoline-3,5dione. Separation of derivatized VitD analytes was achieved using a Waters Acquity BEH C18 column (150 mm \times 2.1 mm, 1.7 μ m particles) with a gradient elution of water with 0.1% formic acid and acetonitrile. The flow rate was 500 μ l/min and the total run time was 8 min. Detection of analytes was achieved using positive atmospheric pressure chemical ionization and selected reaction monitoring on a TSQ Quantum Ultra triple quadrupole mass spectrometer (Thermo Scientific, San Jose, CA). Standards and quality control samples were constructed using blank human serum. Standard curve ranges were 0.10-15 ng/ml for VitD₃ and 24,25D₃, 0.0100-0.500 ng/ml for 1,25D₃, and 1.0-100 ng/ml for 25D₃. Mean correlation coefficients were ≥ 0.994 for all calibration curves. The within-run and between-run accuracy and precision percentage coefficient of variation were $< 10.6\%$ for all analytes.

7.2 Physiologically Based Pharmacokinetic Model Development

A nine-compartment PBPK model (adipose tissue, brain, heart, kidney, liver, rapidly perfused tissue, slowly perfused tissue, and plasma) was developed for VitD₃ and its metabolites using MATLAB (version R2020a, The

Mathworks Inc, Natick, MA). A schematic of the overall PBPK model for each compound is shown in Figure 1 with the connections between each metabolite PBPK model shown in Figure 2.

Physiological parameters were obtained from a previous publication [Davies and Morris, 1993]. The slowly perfused compartment was composed of skin and muscle. The rapidly perfused compartment served as a mass balance compartment and comprised all tissue not previously named. Fractional blood flow rate for the slowly perfused compartment was equal to the sum of the rates of its components. The fractional blood flow rate of the rapidly perfused compartment was equal to the remaining fractional blood flow rates. Following the methods of Ramakrishnan et al. [2016], this study assumed no binding to red blood cells and scaled the blood flow to tissues by multiplying the blood flow Q_B and the blood volume V_B by BP , the blood:plasma ratio, to model the rate of flow of plasma, as shown in Equations (1) and (2):

$$V_P = BP \cdot V_B \quad (1)$$

$$Q_P = BP \cdot Q_B \quad (2)$$

The PBPK model was comprised of conventional mass balance equations assuming well-stirred distribution into each model compartment, with detailed equations for the PBPK model found in the Appendix (Equations (A1) to (A14)). Calculations for relevant partition coefficients and physiological values used in this study are found in the supplemental materials (Supp. Table S1 and S2).

7.2.1 Metabolic Routes

The metabolic cascade for VitD₃ considered in this study consists of three CYP enzymes: CYP2R1, CYP27B1, and CYP24A1. Although the CYP enzyme CYP2R1, a microsomal enzyme found mainly in the liver, is primarily responsible for the 25-hydroxylation of VitD₃ to 25D₃ following Michaelis-Menten kinetics [Cheng et al., 2004; Shinkyō et al., 2004], there are several other enzymatic pathways that may be involved in the 25-hydroxylation pathway including CYP27A1 [Sakaki et al., 2005], CYP2J2 and CYP3A4 [Battault et al., 2012; Wang et al.,

2013]. This study chose to focus on the actions of CYP2R1, with the assumption that V_{max} may vary depending on levels of sufficiency [Abramson, 1986]. The variable rate of V_{max} is given by Equation (3):

$$V_{max} = \frac{V'}{1 + \left(\frac{[D]}{D_{50}}\right)^{h_V}} \quad (3)$$

where V' is the maximum value of V_{max} , $[D]$ is the total liver concentration of VitD₃, D_{50} is the concentration at which 50% of the inhibition occurs, and h_V is the Hill coefficient for this function. The conversion of VitD₃ to 25D₃ by CYP2R1 is given by standard Michaelis-Menten kinetics (Equation (4)) [Ocampo-Pelland et al., 2016]:

$$CL_D = \frac{V_{max} * [D]}{K_m + [D]} \quad (4)$$

Endogenous production of VitD₃ through incidental exposure to sunlight and additional dietary sources are incorporated into the model using Equation (4). This follows the assumption that the input rate for VitD₃ is equal to the output rate of VitD₃ to 25D₃ when there is no external supplementation at low baseline concentrations of VitD₃ [Ocampo-Pelland et al., 2016]. Following the approach of Ramakrishnan et al. [2016], the baseline concentration of VitD₃ and its metabolites in tissue were calculated using the relationship between measured total plasma concentrations and partition coefficients as $C_{T,base} = P_{T,p} \cdot C_{plasma,base}$, where $C_{T,base}$ is the baseline concentration in tissue T , $P_{T,p}$ is the calculated tissue:plasma partition coefficient, and $C_{plasma,base}$ is the baseline concentration of the compound in the plasma compartment.

The metabolite 25D₃ is further converted to either 1,25D₃ by the 1 α -hydroxylase CYP27B1, or deactivated to the metabolite 24,25D₃ by the 24-hydroxylase CYP24A1. Both CYP27B1 and CYP24A1 are renal mitochondrial enzymes which have the capability to demonstrate Michaelis-Menten kinetics [Inouye and Sakaki, 2001; Sakaki et al., 2005]. However, since observed concentrations of 25D₃ range roughly 20-50 times smaller than literature values of K_m for CYP27B1 [Inouye and Sakaki, 2001] and 200-400 times smaller than literature values of K_m for CYP24A1 [Sakaki et al., 2005], this study assumes linear first-order kinetics for CYP27B1 and CYP24A1 acting on 25D₃.

Supplementation doses of VitD₃ during the replenishment period were incorporated into the model through pulsing the initial condition for A_{gut} , the amount of oral VitD₃ effectively introduced in the intestine en route to the liver. During the initial and final portions of the study where no daily supplements were given, the only source of input to the model is through the endogenous production of VitD₃ throughout sunlight and diet. For the purposes of continuity of the model, all participants were assumed to have 16 weeks of supplementation to achieve sufficient concentrations of 25D₃.

7.2.2 Dynamic Adipose Partition Coefficients

Previous publications have suggested the possibility that partition coefficients in tissues, in particular the kidney, may change with levels of sufficiency [Quach et al., 2015] and the adipose tissue [Sawyer et al., 2017]. We chose to incorporate dynamic partitioning for the adipose tissue compartment for VitD₃ and 25D₃ similar to [Sawyer et al., 2017] using the following equations (Equations (5) and (6)):

$$AdiPC_D(t) = \frac{Adi_{D,max}}{1 + \left(\frac{Adi_{D,50}}{[D]}\right)^{h_D}} \quad (5)$$

$$AdiPC_{25D}(t) = \frac{Adi_{25,max}}{1 + \left(\frac{[25D]}{Adi_{25D,50}}\right)^{h_{25}}} \quad (6)$$

Here, $AdiPC_{\Gamma}(t)$ are the adipose partition coefficients at any point in t related directly to the total plasma concentration of either VitD₃ or 25D₃ at time t (indicated by $[D]$ and $[25D]$, respectively), $Adi_{\Gamma,50}$ is the concentration of VitD₃ or 25D₃ at which 50% of the inhibition occurs, and h_{Γ} is the Hill coefficient for these functions.

7.3 Stimulatory and inhibitory effects of 1,25D₃

Circulating and tissue concentrations of 1,25D₃ are tightly regulated through its synthesis by CYP27B1 and degradation by CYP24A1. CYP27B1 expression is tightly regulated by parathyroid hormone and 1,25D₃, where

high levels of 1,25D₃ suppress expression of CYP27B1 [Schuster, 2011]. Conversely, the expression of CYP24A1 is enhanced by the presence of 1,25D₃; upregulation of CYP24A1 by 1,25D₃ serves as feedback control to reduce concentrations of 1,25D₃ [Ramakrishnan et al., 2016]. To incorporate the regulatory effects of 1,25D₃ plasma concentrations, this study follows the approach previously outlined [Ramakrishnan et al., 2016] and adjusts the fold change (FC; ratio of changed/basal mRNA levels) of CYP27B1 and CYP24A1 in proportion to total 1,25D₃ plasma concentrations calculated from the partition coefficient in the relevant tissues. The rate of CYP27B1 fold change was expressed as shown in Equation (7):

$$\frac{dCYP27B1_{FC}}{dt} = k_{in,27} \cdot \left(1 - \frac{I_{max} [1,25D_K]^{\gamma_2}}{IC_{50}^{\gamma_2} + [1,25D_K]^{\gamma_2}} \right) - k_{out,27} \cdot CYP27B1_{FC} \quad (7)$$

with I_{max} the maximum inhibitory effect, IC_{50} the total plasma concentration of 1,25D₃ in the kidney to achieve 50% of I_{max} , γ_2 is the Hill coefficient for CYP27B1 in the kidney, and $[1,25D_K]$ is the total concentration of 1,25D₃ in the kidney [Ramakrishnan et al., 2016]. For CYP24A1, Equation (8) gives the fold change in tissue T , for T liver, kidney, intestine, or brain:

$$\frac{dCYP24A1_{FC,T}}{dt} = k_{in,24,T} \left(1 + \frac{E_{max,T} [1,25D_T]^{\gamma_{1,T}}}{EC_{50,T}^{\gamma_{1,T}} + [1,25D_T]^{\gamma_{1,T}}} \right) - k_{out,24,T} \cdot CYP24A1_{FC,T} \quad (8)$$

where $E_{max,T}$ is the maximum stimulatory effect in tissue T , $EC_{50,T}$ is the total plasma concentration of 1,25D₃ in the tissue to achieve 50% of E_{max} , $\gamma_{1,T}$ is the Hill coefficient for CYP24A1 in the tissue, and $[1,25D_T]$ is the total concentration of 1,25D₃ in the tissue [Ramakrishnan et al., 2016]. Following the methods of [Noh et al., 2020, Ramakrishnan et al., 2016], $k_{in,[]}$ and $k_{out,[]}$ are presumed to have the same value but different (appropriate) units for the differential equation.

7.4 Model simulations and parameter estimations

This study follows the approach outlined by McNally et al. [2011] to explore the global sensitivity of the kinetic parameters for this PBPK model. The model equations were coded in MATLAB (version R2020a, The Mathworks Inc, Natick, MA) and sensitivity analysis was conducted using the SAFE Toolbox [Pianosi et al.,

2015]. For the initial analysis, Morris' Method was used to distinguish between the set of influential and noninfluential parameters through ranking [Morris, 1991]. Parameters in this study were considered influential if the normalized mean sensitivity measure obtained using Morris' Method was greater than 0.1 [Hsieh et al., 2018]. Using this method, the set of influential parameters were chosen for optimization and all other parameters were held at available literature values. If a parameter did not have an available literature value, it was also included in the set chosen for optimization.

This set of parameters was optimized using nonlinear least squares where the cost of fit was calculated using the least squares difference between the observed and predicted model data. To avoid unfairly skewing the model, each compound's data were transformed by subtracting the minimum observed value and dividing by the interquartile range of the observed data for that compound. Data were further weighted by the count of the number of available points for optimization in each phase; this ensured that data below the analytical level of quantification were not considered in and did not negatively affect the optimization routine. This was present primarily in the insufficient VitD₃ data and comprised approximately 15% of the overall data points (3% of points without the insufficient VitD₃ data) used for optimization. The lack of data due to patient VitD₃ concentrations below the limit of quantification poses some issue for initialization of the model for this compartment; however, the set of influential parameters span the entire timespan of the model, so we used visual inspection of the curve of the model prediction to assess fit in this area for this compartment. The model was initialized to the mean of the first two time points ($t = 0$ and $t = 0.5$ hours).

To generate 95% confidence intervals for the optimized parameters, a bootstrapping method was used after initial optimization of the model. For this method, parameters identified as candidates for optimization were allowed to vary two-fold from their optimized values, which were based on available literature values, or the best result from multiple iterative guesses. Bootstrapping was performed by systematically running the optimization solver over randomly sampled parameters within these constraints, holding non-optimized parameters constant at literature values. The resulting parameter output range for each optimized parameter was then subject to standard

bootstrapping techniques and the 95% confidence intervals were found for each parameter. To generate a 95% confidence interval around the optimized model, parameters were randomly sampled in a normal distribution from a two-fold range of their optimized value for 50 samples, the model run over these parameter sets, and then the 95th percentile of the total model outputs at every half-hour mark was used to generate the total band for the duration of the time-course data.

7.5 Assessment of Prediction Accuracy

The accuracy of the PBPK predictions was evaluated on predicted plasma concentration fit to observations for the three measured compounds. The predicted plasma concentration was formulated using the set of optimized parameters developed from the bootstrapping results. The goodness of prediction for each compound was based on the average fold error (*AFE*), root mean square error (*RMSE*), and the normalized root mean square error (*NRMSE*) [Sheiner and Beal, 1981]. The *fold error* [Equation (9)], *AFE* [Equation (10)], *RMSE*, [Equation (11)], and *NRMSE* [Equation (12)] were calculated as follows:

$$\text{fold error} = \frac{\text{Predicted}}{\text{Observed}} \quad (9)$$

$$\text{AFE} = \exp\left(\frac{\sum \ln(\text{fold error})}{n}\right) \quad (10)$$

$$\text{RMSE} = \sqrt{\frac{\sum (\text{Pred} - \text{Obs})^2}{n}} \quad (11)$$

$$\text{NRMSE} = \frac{\text{RMSE}}{\text{IQR}(\text{Obs})} \quad (12)$$

where *Pred* is the predicted value, *Obs* is the observed value, *IQR(Obs)* is the interquartile range of the observed values, and *n* is the number of observed samples.

8 Results

A basic PBPK approach was applied to VitD₃ and metabolites using intrinsic clearance parameters determined from the published literature and physiological distribution parameters (Figure 1). Data from eleven subjects (demographics given in Table 1) were used to fit the model; two subjects withdrew from the study before the conclusion of the experiment.

Due to limited VitD₃ time course data, the nearly instantaneous conversion of VitD₃ to 25D₃ in VitD deficient individuals [Heaney et al., 2008], and the adipose tissue as a storage compartment that may contribute to circulating levels of VitD₃ [Best et al., 2020], our model was insufficient at predicting available VitD₃ time course data (results not shown). However, clinically, VitD₃ levels are not measured and not likely to be important since therapy decisions are based on 25D₃ levels. All optimized and literature model parameters are reported in Table 2. Bootstrapping was performed and optimized parameters are reported with 95% confidence intervals.

The observed and final model predicted concentrations for 25D₃, 24,25D₃ and 1,25D₃ during the insufficient and repleted periods are shown in Figure 3 (a-c), respectively, with a 95% confidence interval band around the optimized parameter model as described in section 7.5. Detailed model predictions for the insufficient and repleted periods are shown as insets in Figure 3. The wavy line in the center of the plots indicate a break in the supplementation period for better clarity on the insufficient and repleted period.

Predictive performance of the PBPK model for the insufficient and repleted periods are shown in Figure 4. The average fold error (*AFE*) for the VitD₃ metabolites ranges between 1.01 and 1.05 and the normalized root mean square error (*NRMSE*) ranges between 0.26 and 0.38. *AFE* values close to one and *NRMSE* close to zero are indicative of better model fits to data. The correlation of determination (R^2) for each metabolite is 0.77, 0.67, and 0.63, respectively, and 0.92 when considering all data points together. When considering the *fold error* values for the data, the model performed well with 97%, 88%, and 98% of the data for 25D₃, 24,25D₃, and 1,25D₃,

respectively, within 2-fold of unity (*fold error* values between 0.5 and 2.0). When considering a tighter range of fold error (1.5-fold of unity, with *fold error* values between 0.67 and 1.5), the model performs adequately with nearly 80% of 25D₃ and 1,25D₃ captured within a 1.5-*fold error* of unity. 24,25D₃ data was captured less well, with a 1.5-*fold error* value of 63%.

9 Discussion

Vitamin D insufficiency is highly prevalent in the community, afflicting up to 80% of men and women in the USA, Canada, and Europe [Ganji et al., 2012; van Schoor and Lips, 2018]. While the definitions of VitD insufficiency (and deficiency) are not standardized across medical organizations, targeted concentrations are generally in the 20 ng/ml to 40 ng/ml range for 25D, the primary metabolite used for classification [Ganji et al., 2012; Holick et al., 2011; Ross et al., 2011; van Schoor and Lips, 2018]. While various treatment recommendations have also been proposed according to the IOM [Jernigan and Andress, 2003] and ENDO [Holick et al., 2011], there is variability in success of achieving target concentrations of 25D₃ in patients. There is currently not a clinically established approach to enable prediction of plasma 25D concentrations that might result from a given treatment regimen for a given patient. There is further complication since the plasma concentration of 25D will be impacted by the function of numerous CYP enzymes through activation and deactivation pathways, as well as through sunlight exposure, skin pigment, diet, and clothing. PBPK modeling has a potential to predict expected plasma concentrations of 25D and subsequent metabolites after administration of oral VitD therapy. The current study reports the development of a PBPK model of VitD₃ and metabolites in VitD insufficient subjects in the U.S. who were treated with moderate doses (5,000 I.U.) of daily VitD₃ for up to 16 weeks to achieve replacement as defined by 25D \geq 30 ng/ml. The comprehensive PBPK model for VitD₃ incorporated dynamic adipose:plasma partition coefficients, atypical kinetics for CYP2R1 using a variable V_{max} based on the total liver concentration of VitD₃, and inhibitory and stimulatory effects of 1,25D₃ for CYP27B1 and CYP24A1. The resultant model sufficiently predicted the concentrations of VitD₃ metabolites well in healthy subjects throughout a multiphase study that incorporated an insufficient phase and replete phase. The model informs an understanding of the disposition of VitD₃ metabolites and could be utilized to predict plasma concentrations that might result from a given dosing regimen of VitD₃ in human patients.

The developed PBPK model for VitD₃ and its metabolites consisted of nine compartments (adipose tissue, brain, heart, intestines, kidney, liver, rapidly perfused tissue, slowly perfused tissue, and plasma). Partition coefficients

were either selected from the literature or calculated as described in the Supplemental Material. Although a previous publication in mice suggested that partition coefficients may differ in states of vitamin D deficiency [Quach et al. 2015], our simulations showed less than a 2% change in liver partition coefficients (as defined by the ratio of liver concentration to plasma concentration) between insufficient and sufficient states for VitD₃ and metabolites, and less than 1% for kidney partition coefficients (data not shown). However, we did find a difference in model outcomes with the inclusion of variable partitioning in the adipose tissue compartment. As subjects in our study were initially at levels of insufficiency, there is evidence to suggest that normal daily inputs and endogenous production of vitamin D is not sufficient for accumulation in tissues [Heaney et al., 2009]; however, at higher levels of sufficiency, the body is able to store VitD₃ and 25D₃ in adipose tissue for use at a later time [Abbas, 2017; Heaney et al., 2009; Mawer et al., 1972]. The addition of a variable adipose partition coefficient for VitD and 25D, as described in section 7.2.2 and discussed in [Sawyer et al., 2017] led to enhanced fits for the model. In addition, our model agrees well with previously published human studies without K_p values defined [Holick et al., 2008, figure 2B].

As VitD₃ is acquired endogenously through the skin from the effects of ultraviolet-B light on 7-dehydrocholesterol (7-DHC) and dietary sources including dairy and fish, these contribute to baseline concentrations of VitD₃ in the plasma. The PBPK model incorporated endogenous VitD₃ levels with an assumption of output to 25D₃. After oral ingestion and absorption of supplemental VitD₃, the free fraction is taken up in the liver where it is hydroxylated by 25-hydroxylase to form calcifediol or 25D₃. Subsequently, 25D₃ is hydroxylated by the kidneys to form calcitriol or 1,25D₃, which is considered the most active metabolite of VitD₃. The current PBPK model used linear first order kinetics for CYP27B1 and CYP24A1 based on the significantly lower 25D₃ concentrations vs. published K_m values for the respective enzymes. Supplemental dosing of cholecalciferol was incorporated into the model through pulsing the amount introduced into the intestines.

Circulating and tissue concentrations of 1,25D₃ are tightly regulated through the synthesis by CYP27B1 and degradation by CYP24A1. While CYP27B1 expression is tightly regulated by the binding of 1,25D₃ and

parathyroid hormone [Schuster, 2011], the expression of CYP24A1 is enhanced by the presence of 1,25D₃; upregulation of CYP24A1 by 1,25D₃ serves as a feedback control to reduce concentrations of 1,25D₃ [Ramakrishnan et al., 2016]. The current PBPK model incorporated the regulatory effects of 1,25D₃ plasma concentrations by adjusting the fold change of CYP27B1 and CYP24A1 in proportion to 1,25D₃ plasma concentrations in the liver, kidney, intestine, and brain, as previously identified [Ramakrishnan et al., 2016]. While the fold-change data was simulated and although the dosing regimen and the supplementation in [Ramakrishnan et al., 2016] is different than our dosing regimen and supplementation, we observed the same general behavior for the fold change of each enzyme (results not shown), namely that as the level of 1,25D₃ increases, the fold change of CYP27B1 decreases and the levels of CYP24A1 increase in each compartment. When the supplementation was removed at the repleted portion of the study, we observed an expected increase and decrease, respectively, in the fold changes of the CYP24A1 and CYP27B1 enzymes comparative to the concentration of 1,25D₃ in the appropriate compartments. This behavior follows the expected directions for CYP27B1 and CYP24A1 given the critical concentration of 1,25D₃ governing these physiological effects on these enzymes.

Recent publications have discussed the advancement of atypical Michaelis-Menten kinetics for a variety of CYP enzymes, including several enzymes involved in the secondary metabolism cascade for vitamin D. This includes CYP3A4 [Arendse et al, 2013] and CYP2J2 [Leow and Chan, 2019; Leow et al., 2021]. In particular, CYP2J2 shares 72.5% sequence similarity with CYP2R1, so it is not unreasonable to assume that CYP2R1 may also exhibit atypical Michaelis-Menten kinetics at varying sufficiency levels. To model this, we chose to use a modification on substrate inhibition kinetics, as shown in Equation (3). This led to strong improvement of model predictions, particularly in latter time points (data not shown).

Since the metabolites versus the parent VitD₃ compound are measured clinically to assist clinicians in determining whether patients are VitD₃ insufficient and/or whether they require changes to dosing regimens, we performed some simulations of predicted concentrations with common regimens. A simulation using the current model was

performed inputting a dose of 1000 I.U./daily, which is consistent with dosing recommendation according to the IOM. Our model predicted 25D₃ plasma concentrations well (Figure 5) and is consistent with a previous study with this dosing regimen [Holick et al., 2008, fig. 2B]. Note subjects in [Holick et al., 2008] began the study with sufficient levels of VitD; we hypothesis this explains the slight underprediction of our model to participant data at later time points as the ratio of VitD₃ and metabolites may not remain the same across levels of sufficiency for initialization of the model.

Overall, the developed PBPK model of VitD₃ that incorporated data from insufficient participants into a repleted phase after daily administration of 5000 I.U. VitD₃ for 12-16 weeks led to acceptable and satisfactory predictions in healthy human subjects. When considering the *fold error* values for the data, the model performed well with 97%, 88%, and 98% of the data for 25D₃, 24,25D₃, and 1,25D₃, respectively, within 2-fold of unity (*fold error* values between 0.5 and 2.0). Due to limited VitD₃ time course data, the nearly instantaneous conversion of VitD₃ to 25D₃ in VitD deficient individuals [Heaney et al., 2008], and the adipose tissue as a storage compartment that may contribute to circulating levels of VitD₃ [Best et al., 2020], our model was insufficient at predicting available VitD₃ time course data (results not shown). However, this may not be overly relevant since VitD₃ levels are not measured or used clinically. An additional limitation to this model is the lack of available experimental data for comparison of atypical kinetic parameters for CYP2R1; however, the inclusion of the atypical kinetics greatly improved our model behavior and is within the realm of biological possibility. The model serves to inform an understanding of VitD₃ and metabolite disposition and could be utilized to predict 25D₃ plasma concentrations that might result from a given dosing regimen in human patients. A beneficial future addition to this study would include incorporation of patient-specific data throughout the supplementation phase to better inform the model predictions. This model has applications in studying the effects of repletion schemes for populations of patients with impaired enzymatic abilities, such as chronic kidney disease, an important diseased population we are currently evaluating.

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11 Authorship Contributions

Participated in research design: Joy and Nolin

Conducted experiments: Joy, Nolin, Tuey, and West

Performed data analysis: Sawyer

Wrote or contributed to the writing of the manuscript: Sawyer, Tuey, Joy, and Nolin

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13 Footnotes

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- b.) Sawyer CW, Tuey SM, West III RA, Nolin TD, Joy MS. A Triphasic Physiologically Based Pharmacokinetic Model of Vitamin D3 and Metabolites in Vitamin D Insufficient Patients. Presented at *BAMM! Biology and Medicine through Mathematics*, Richmond, VA, May 19, 2022.
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14 Figure Legends

Figure 1: General physiologically based pharmacokinetic model diagram for VitD₃ and metabolites. Symbols are defined as the following: Q is the plasma flow rate, C_{ap} is the arterial concentration of the compound; $C_{vp:T} = A_T / (V_T * P_T)$ is the apparent concentration of the compound in tissue T defined as the amount of compound divided by the volume of the tissue times the tissue:plasma partition coefficient; k_a is the absorption rate from the gut.

Figure 2: Network diagram for the VitD₃ metabolic cascade model. The network contains compounds with measured concentrations and black arrows define conversion steps with kinetic equations as defined in Table 2. Note that concentrations of 1,24,25(OH)₃D₃ were not measured in this study.

Figure 3: Predicted PBPK plasma concentrations for VitD₃ metabolites (25D₃, 24,25D₃, and 1,25D₃). Box plots indicate the middle 50% of the data with the median data indicated by a line, and the shading indicates the 95% confidence interval of the model predictions. Insets indicate enhanced view of data for predicted PBPK plasma concentrations for VitD₃ metabolites (25D₃, 24,25D₃, and 1,25D₃), clipping out the supplementation period from days 20 through 92 of the experiment, where the wavy line in the center indicates the clipped portion of the model during the supplementation period where no data were observed.

- (a) 25D₃ serum levels (nmol L⁻¹)
- (b) 24,25D₃ serum levels (nmol L⁻¹)
- (c) 1,25D₃ serum levels (nmol L⁻¹)

Figure 4: Observed plasma concentrations versus predicted PBPK plasma concentrations in vitamin and three metabolites. Red x-marks indicate observations from the deficient period, blue circles indicate observations from the replete period, the dotted line represents the identity line and the green lines represent the 2-fold interval.

Figure 5: Simulation of daily 1000 I.U. dosing regimen. Solid line indicates model predictions, data taken from [Holick et al., 2008] are mean ± SEM.

15 Tables

Table 1: Baseline characteristics of study participants ($n = 11$)

Study Site	University of Pittsburgh	9 (82%)
	University of Colorado	2 (18%)
Gender	Female	6 (55%)
	Male	5 (45%)
Race	White	6 (55%)
	Black	5 (45%)
Ethnicity	Non-Hispanic	11 (100%)
Age (years)		57 (11.5)
Weight (kg)		89.2 (42.6)
BMI (kg/m ²)		30.5 (13.2)
eGFR (ml/min/1.73 m ²)		92 (10.5)
Serum Albumin (g/dl)		4.3 (0.2)
Serum VDBP (µg/ml)		294 (88.7)

Data are presented as median (IQR) or number (%).

Abbreviations: eGFR – estimated glomerular filtration rate; VDBP – vitamin D binding protein

Table 2: Kinetic parameters for the PBPK model. Optimized parameters are given with bootstrapped 95% confidence intervals and all other parameters are held to available literature values or estimates as notated. Note the 95% CIs are not necessarily symmetric about the optimized parameter due to the method of sampling. See the *Model simulations and parameter estimations* section for details.

Parameter	Description	Value	Units	Reference
K_m	Michaelis-Menten constant for CYP2R1	5.91 ^a (5.09, 5.95)	nmol L ⁻¹	Optimized
D_{base}	Endogenous baseline concentration of VitD ₃	2.05 ^a (2.00, 2.06)	nmol·L ⁻¹	Optimized
k_a	Absorption constant	0.32	h ⁻¹	[Ocampo-Pelland et al., 2016]
R_{27B1}^{25}	Synthesis of 25D ₃ to 1,25D ₃ by CYP27B1	17.7 ^b (15.85, 18.97)	L·hr ⁻¹	Optimized
f_{ub}^{25}	Percent free unbound 25D ₃	0.03	--	[Bikle et al., 1986]
Synthesis of 24,25D ₃ from 25D ₃ by CYP24A1 in tissue				
$R_{24A1,L}^{25}$	Liver	10.2 ^b (8.45, 10.8)	L·hr ⁻¹	Optimized
$R_{24A1,K}^{25}$	Kidney	2.43 ^b (2.06, 2.68)	L·hr ⁻¹	Optimized
$R_{24A1,Br}^{25}$	Brain	13.3 ^b (12.2, 13.9)	L·hr ⁻¹	Optimized
$R_{24A1,I}^{25}$	Intestine	15.4 ^b (13.2, 16.8)	L·hr ⁻¹	Optimized
Clearance of unbound (free) 1,25D ₃ by CYP24A1 in tissue T , $CL_{free,int,T}^{125}$				
$CL_{free,int,L}^{125}$	Liver	0.0057	mL·hr ⁻¹ ·g ⁻¹	[Noh et al., 2020]

$CL_{free,int,K}^{125}$	Kidney	0.066 ^a (0.065, 0.082)	$\text{mL}\cdot\text{hr}^{-1}\cdot\text{g}^{-1}$	Optimized
$CL_{free,int,Br}^{125}$	Brain	0.0083	$\text{mL}\cdot\text{hr}^{-1}\cdot\text{g}^{-1}$	[Noh et al., 2020]
$CL_{free,int,I}^{125}$	Intestine	0.0028	$\text{mL}\cdot\text{hr}^{-1}\cdot\text{g}^{-1}$	[Noh et al., 2020]
$CL_{free,int,K}^{2425}$	Clearance of unbound (free) 24,25D	990 ^b (946, 1068)	$\text{pmol}\cdot\text{hr}^{-1}$	Optimized
I_{max}	Maximal inhibitory effect of CYP27B1	1	fold change	[Noh et al., 2020]
IC_{50}	Total 1,25D ₃ concentration when reaching 50% I_{max}	672 ^a (680,1068)	pmol L^{-1}	Optimized
Maximal stimulatory effect of CYP24A1 in tissue T , $E_{max,T}$				
$E_{max,L}$	Liver	122 ^a (88.7, 131.6)	fold change	Optimized
$E_{max,K}$	Kidney	25 ^a (23.9, 28.0)	fold change	Optimized
$E_{max,Br}$	Brain	22 ^a (21.5, 23.0)	fold change	Optimized
$E_{max,I}$	Intestine	675	fold change	[Noh et al., 2020]
Total tissue 1,25D ₃ concentration when reaching 50% E_{max} , $EC_{50,T}$				
$EC_{50,L}$	Liver	2600	pmol L^{-1}	[Noh et al., 2020]
$EC_{50,K}$	Kidney	143	pmol L^{-1}	[Noh et al., 2020]
$EC_{50,Br}$	Brain	97 ^a (74.5, 97.5)	pmol L^{-1}	Optimized
$EC_{50,I}$	Intestine	2640	pmol L^{-1}	[Noh et al., 2020]
Hill coefficient of CYP24A1 in tissue T , $\gamma_{1,T}$				

$\gamma_{1,L}$	Liver	3.05 ^a (2.99, 3.47)	--	Optimized
$\gamma_{1,K}$	Kidney	3.59	--	[Noh et al., 2020]
$\gamma_{1,Br}$	Brain	1.12 ^c (0.90, 1.15)	--	Optimized
$\gamma_{1,I}$	Intestine	4.16	--	[Noh et al., 2020]
γ_2	Hill coefficient of CYP27B1	2.7	--	[Noh et al., 2020]
$k_{out,27}$	Turnover rate constant of CYP27B1	0.245	h ⁻¹	[Noh et al., 2020]
Turnover rate constant of CYP24A1 in tissue T , $k_{out}^{24,T}$				
$k_{out,24,L}$	Liver	0.47	h ⁻¹	[Noh et al., 2020]
$k_{out,24,K}$	Kidney	0.28	h ⁻¹	[Noh et al., 2020]
$k_{out,24,Br}$	Brain	0.86	h ⁻¹	[Noh et al., 2020]
$k_{out,24,I}$	Intestine	0.047	h ⁻¹	[Noh et al., 2020]
Atypical Kinetic Parameters for V_{max}				
V'	Maximum value of V_{max} for CYP2R1	5.76 ^b (5.73, 5.76)	L hr ⁻¹	Optimized
D_{50}	Concentration at which 50% of V' occurs	8.35 ^b (8.34, 8.35)	nmol L ⁻¹	Optimized
h_V	Hill constant for variable V_{max}	2.27 ^b (2.26, 2.27)	--	Optimized
Dynamic Adipose partition coefficient parameters				
$Adi_{D,max}$	Maximum adipose PC for VitD ₃	9.9 ^b (9.78, 10.4)	--	Optimized
$Adi_{25,max}$	Maximum adipose PC for 25D ₃	1.64 ^b (0.90, 1.73)	--	Optimized

$Adi_{D,50}$	Concentration at which 50% of $Adi_{D,max}$ occurs	8.96 ^b (4.93, 11.98)	nmol L ⁻¹	Optimized
$Adi_{25D,50}$	Concentration at which 50% of $Adi_{25D,max}$ occurs	117 ^b (61.8, 121)	nmol L ⁻¹	Optimized
h_D	Hill constant for $AdiPC_D$	2.12 ^b (2.10, 2.30)	--	Optimized
h_{25}	Hill constant for $AdiPC_{25}$	0.96 ^b (0.86, 1.01)	--	Optimized

Abbreviations: D₃, VitD₃; 25D₃, 25(OH)D₃; 1,25D₃, 1,25(OH)₂D₃; 24,25D₃, 24,25(OH)₂D₃

^aConfidence interval does not contain literature values

^bNo literature values available for comparison

^cConfidence interval contains literature value

16 Figures

Attached separately

18 Appendix

A.1 PBPK Model Equations

The parameters in the PBPK model equations refer to tissue (T), arterial blood flow (Q), compound amount (A), concentration (C), volume (V), and tissue:plasma partition coefficients ($P_{T:p}$). The concentration in each tissue is given by Equation (A1), where the tissue:plasma partition coefficients are given in Supp. Table S2.

$$C_T = A_T / (V_T \cdot P_{T:p}) \quad (\text{A1})$$

For compartments that have the same equation form across all compounds, general mass balance differential equations are given in Equations (A2) and (A3) based on Figure 1 in the main text. Definitions and values for parameter values for Equations (A2) through (A14) are given in Table 2 and Supp. Tables S1 and S2.

Non-eliminating tissue in the set Z (adipose, heart, rapidly perfused, slowly perfused)

$$V_Z \cdot dC_Z/dt = BP \cdot Q_Z \cdot (C_p - C_{p:Z}) \quad (\text{A2})$$

Plasma:

$$V_p \cdot dC_p/dt = \sum_T [BP \cdot Q_T \cdot C_{p:T}] - BP \cdot Q_{CO} \cdot C_p \quad (\text{A3})$$

Equations (A4) through (A7) describe the mass balance equations for the remaining four compartments (brain [Br], intestines [I], kidney [K], liver [L]) for VitD₃ and each metabolite, where additional components are described in Equations (A8) through (A14).

VitD₃ equations:

$$V_{Br} \cdot dC_{Br}/dt = BP \cdot Q_{Br} \cdot (C_p - C_{p:Br}) \quad (\text{A4a})$$

$$V_I \cdot dC_I/dt = BP \cdot Q_I \cdot (C_p - C_{p:I}) + k_a \cdot A_{gut} \quad (\text{A4b})$$

$$V_K \cdot dC_K/dt = BP \cdot Q_K \cdot (C_p - C_{p:K}) \quad (\text{A4c})$$

$$V_L \cdot dC_L/dt = BP \cdot Q_L \cdot (C_p - C_{p:L}) + ENDOG - R25D \quad (\text{A4d})$$

25D₃ equations:

$$V_{Br} \cdot dC_{Br}/dt = BP \cdot Q_{Br} \cdot (C_p - C_{p:Br}) - R2425D_{Br} \quad (\text{A5a})$$

$$V_I \cdot dC_I/dt = BP \cdot Q_I \cdot (C_p - C_{p:I}) - R2425D_I \quad (\text{A5b})$$

$$V_K \cdot dC_K/dt = BP \cdot Q_K \cdot (C_p - C_{p:K}) - R125D - R2425D_K \quad (\text{A5c})$$

$$V_L \cdot dC_L/dt = BP \cdot Q_L \cdot (C_p - C_{p:L}) + R25D_3 - R2425D_L \quad (A5d)$$

24,25D₃ equations:

$$V_{Br} \cdot dC_{Br}/dt = BP \cdot Q_{Br} \cdot (C_p - C_{p:Br}) + R2425D_{Br} \quad (A6a)$$

$$V_I \cdot dC_I/dt = BP \cdot Q_I \cdot (C_p - C_{p:I}) + R2425D_I \quad (A6b)$$

$$V_K \cdot dC_K/dt = BP \cdot Q_K \cdot (C_p - C_{p:K}) + R2425D_K - CL2425D_{Br} \quad (A6c)$$

$$V_L \cdot dC_L/dt = BP \cdot Q_L \cdot (C_p - C_{p:L}) + R2425D_L \quad (A6d)$$

1,25D₃ equations:

$$V_{Br} \cdot dC_{Br}/dt = BP \cdot Q_{Br} \cdot (C_p - C_{p:Br}) - CL125D_{Br} \quad (A7a)$$

$$V_I \cdot dC_I/dt = BP \cdot Q_I \cdot (C_p - C_{p:I}) - CL125D_I \quad (A7b)$$

$$V_K \cdot dC_K/dt = BP \cdot Q_K \cdot (C_p - C_{p:K}) + R125D - CL125D_K \quad (A7c)$$

$$V_L \cdot dC_L/dt = BP \cdot Q_L \cdot (C_p - C_{p:L}) - CL125D_L \quad (A7d)$$

Additional differential equations for the PBPK model are given in Equations (A8) to (A14), where concentrations in tissue T are indicated by $[M_T]$ where M is VitD₃ or a metabolite:

Absorption of VitD₃:

$$dA_{gut}/dt = -k_a \cdot A_{gut} \quad (A8)$$

Endogenous production of VitD₃:

$$ENDO_G = \frac{V_{max} \cdot [D_{base}]}{K_m + [D_{base}]} * [D_{base}] \quad (A9)$$

Conversion of VitD₃ to 25D₃ in the liver (L):

$$R25D = \frac{V_{max} \cdot [D_L]}{K_m + [D_L]} * [D_L] \quad (A10)$$

Conversion of 25D₃ to 1,25D₃ in kidney by CYP27B1:

$$R125D = R_{27B1}^{25} \cdot [25D_K] \cdot f_{ub}^{25} \cdot CYP27B1_{FC} \quad (A11)$$

Conversion of 25D₃ to 24,25D₃ in tissue T by CYP24A1:

$$R2425D_T = R_{24A1,T}^{25} \cdot [25D_T] \cdot f_{ub}^{25} \cdot CYP24A1_{FC,T} \quad (A12)$$

Clearance of 1,25D₃ in tissue T by CYP24A1:

$$CL125D_T = CL_{free,int,T}^{125} \cdot [1,25D_T] \cdot CYP24A1_{FC,T} \quad (A13)$$

Clearance of 24,25D₃ in the kidney:

$$CL_{2425D} = CL_{free,int,K}^{2425} \cdot [24,25D_K] \quad (A14)$$

The initial conditions for the model were generated by taking the mean of the data points to generate an initial estimate for the median plasma concentration.

Figure 1

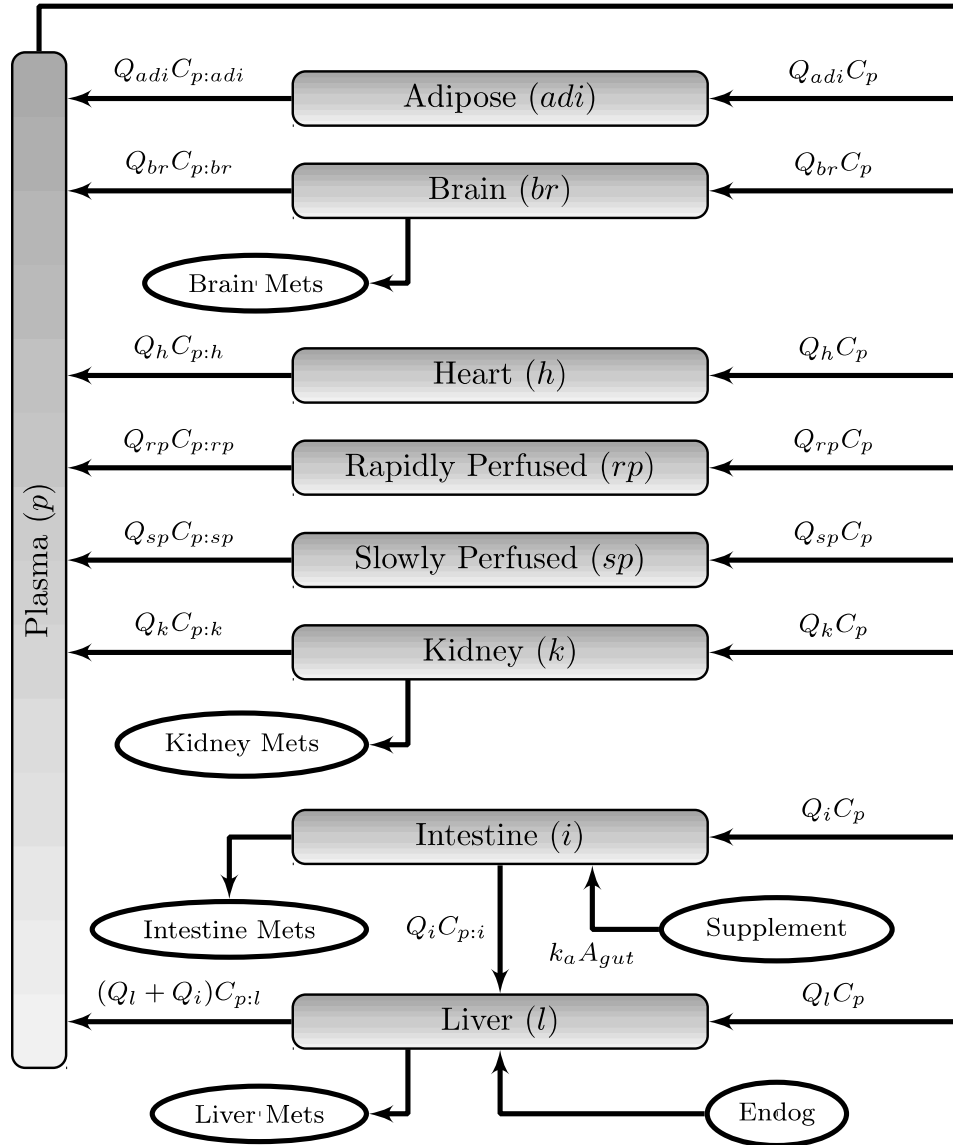


Figure 2

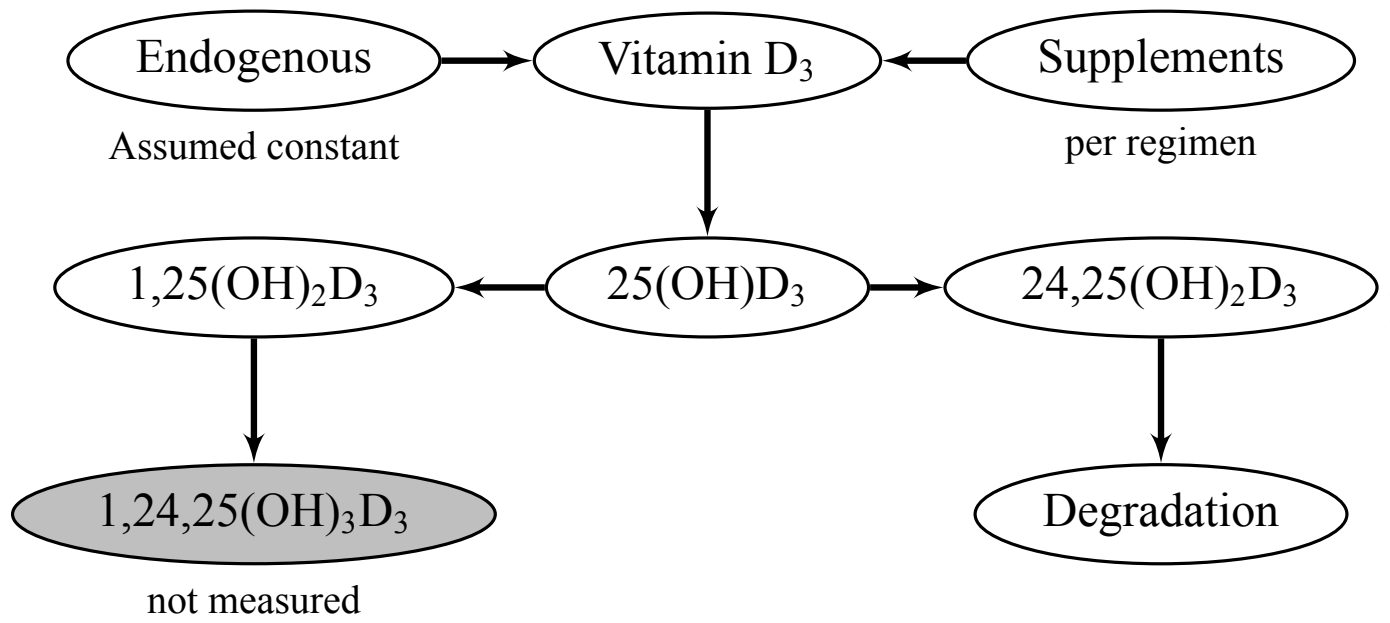


Figure 3

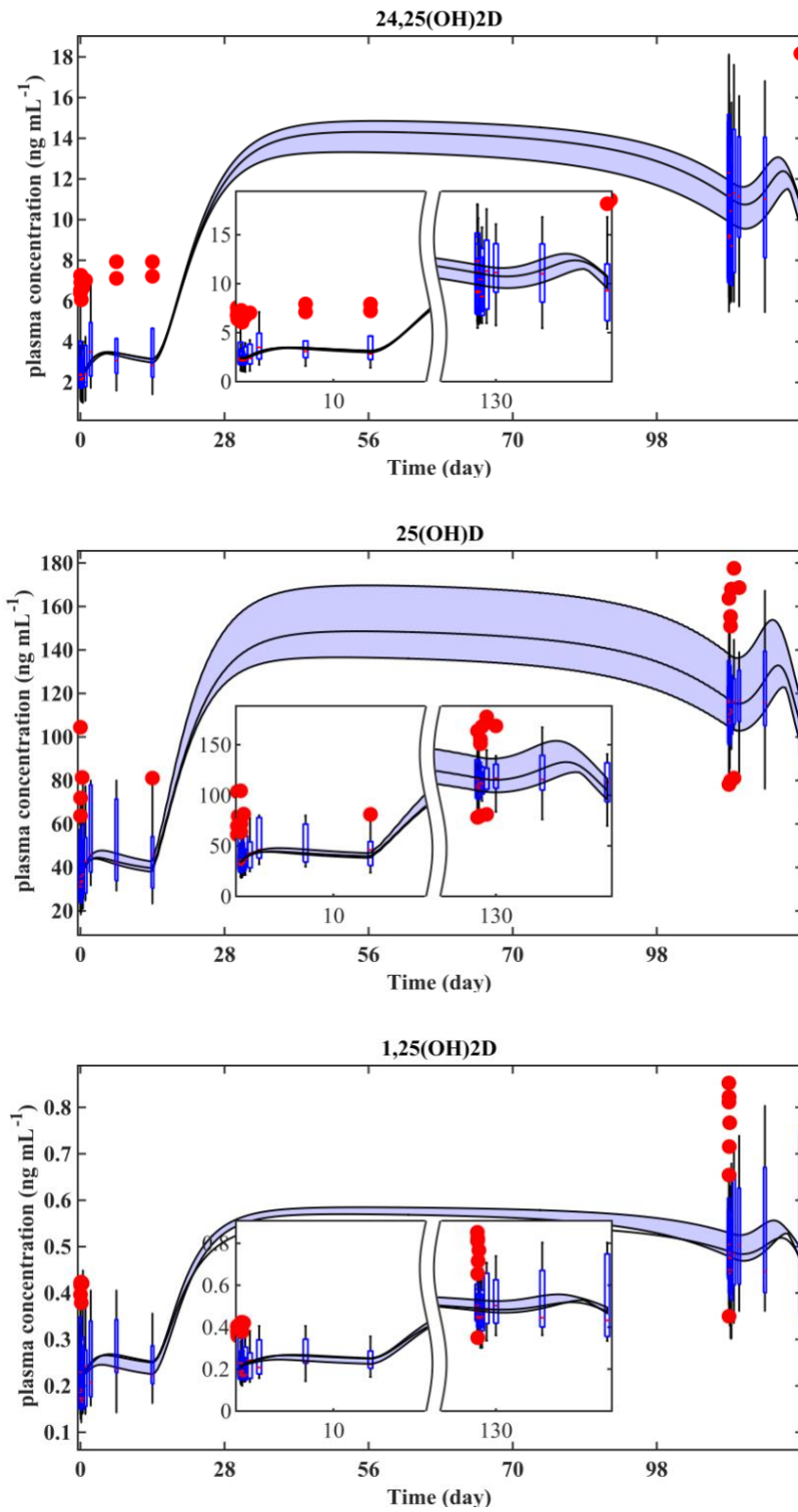


Figure 4

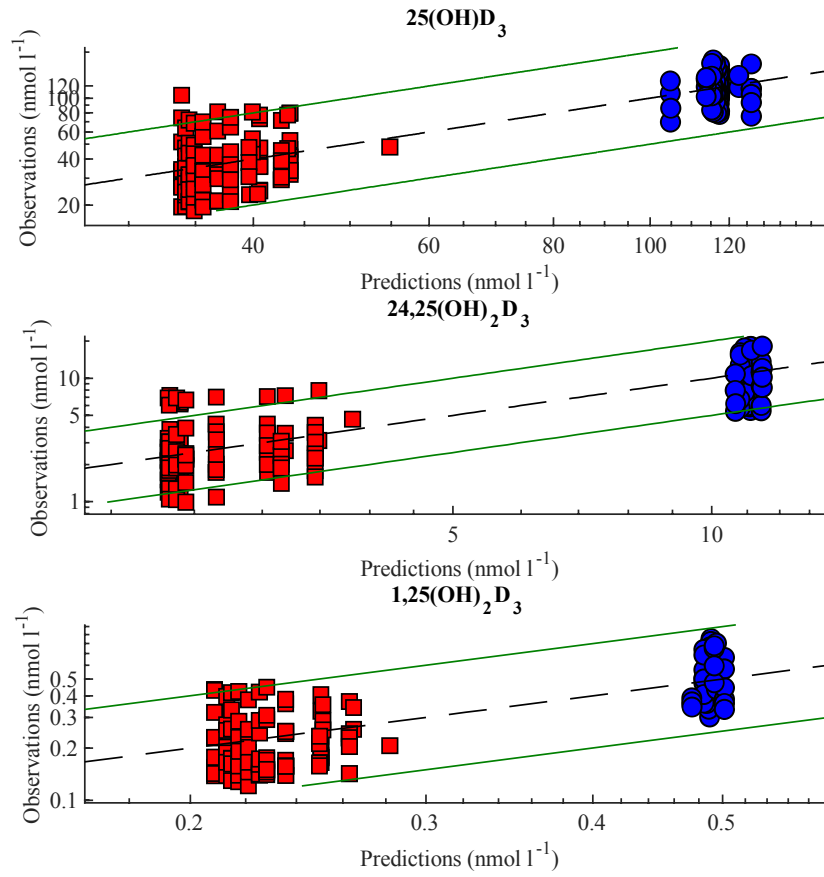
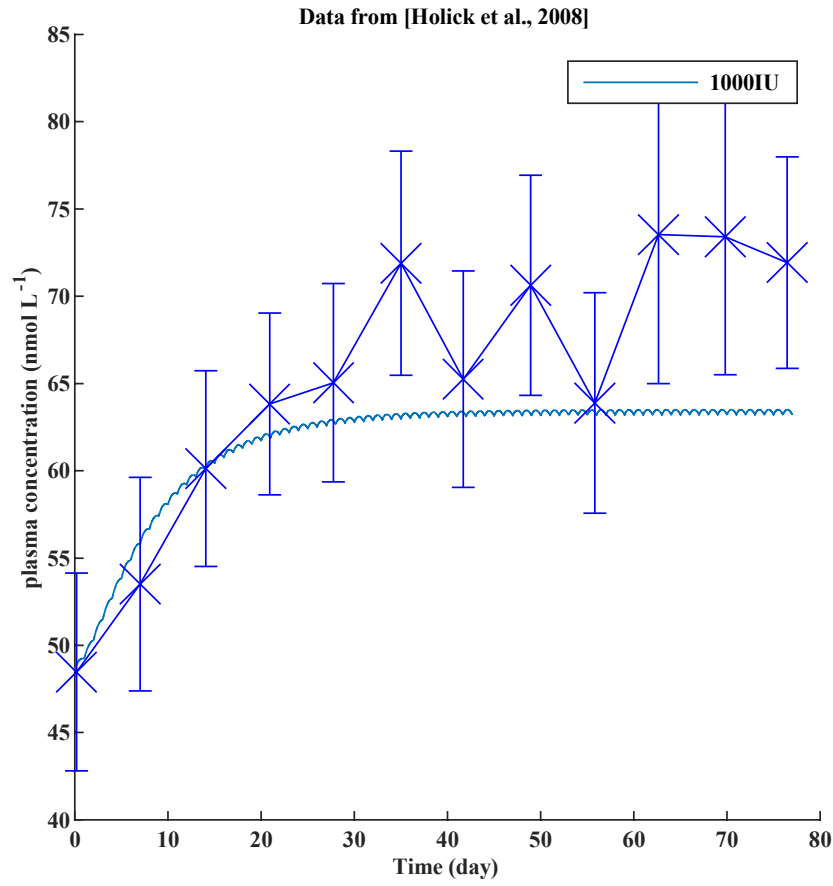


Figure 5



```
% Supplemental Material: Drug Met Disp, DMD-AR-000609
% Title: Physiologically Based Pharmacokinetic Modeling of Vitamin D3 and Metabolites in
Vitamin D Insufficient Patients
% Authors: Colton W. Sawyer, Stacey M. Tuey, Raymond E. West III, Thomas D. Nolin,
Melanie S. Joy
% Code title: DynamAdi
```

```
function [AdiPC] = DynamAdi(K, venconc)
% K = Kinetic values structure
% Adiconc = concentration of venous in VitD and 25D
% AdiPC = adjusted adipose PC for VitD and 25D

Adimax = [K.Adimax K.Adimax25] ;
hAdi = [K.hAdi K.hAdi25];
Adi50 = [K.Adi50 K.Adi50_25];

AdiPC(1) = Adimax(1) ./ (1 + ( Adi50(1)./venconc(1)).^hAdi(1));
AdiPC(2) = Adimax(2) ./ (1 + ( venconc(2)./Adi50(2)).^hAdi(2));

end
```

```

%
% Supplemental Material: Drug Met Disp, DMD-AR-000609
% Title: Physiologically Based Pharmacokinetic Modeling of Vitamin D3 and Metabolites in
Vitamin D Insufficient Patients
% Authors: Colton W. Sawyer, Stacey M. Tuey, Raymond E. West III, Thomas D. Nolin,
Melanie S. Joy
% Code title: main

```

```

function [serum, A, tout] = main(T, g, varargin)

```

```

% -----
% in:
% T = parameter table
% g = group of interest
% varargin = {doseIU, suppweeks}
% out:
% serum = {D, 25, 125, 2425}
% A = Amt (nmol where appropriate)
% tout = time (days)
% -----

```

```

if nargin==4
    doseIU = varargin{1};
    suppweeks = varargin{2};
else
    doseIU = 5000;
    suppweeks = 16;
end

```

```

WD = 384.64; %weight of vitamin D in g/mol = Da
doseug = doseIU/40; %IU -> ug
dosenmol = doseug*1e3/WD; %ug -> nmol

```

```

[P] = initconcs(T,g); %subfunction to get baseline concentration
%initial conditions for vitD
InitAmts = [P.AmtD; P.Amt25; P.Amt125; P.Amt2425];
initbase = [1; ones(4,1)]; %baseline fold: 27B1; 24A1 for 4 tissues
initrates = zeros(11,1);

```

```

T.Pars = P; %update with initial concentrations.

```

```

t1 = 2*7;
t2 = suppweeks*7;
t3 = 2*7;

```

```

daycount = 1:sum([t1,t2,t3]); %16 week trial\

```

```

tvec = 24.*[0:daycount(end)]; %time markers (hours)

```

```

dailyintake = 0; %average dietary supplement
dailydose = [zeros(t1,1); dosenmol*ones(t2,1); zeros(t3,1);0];
dailydose = dailydose + dailyintake*ones(size(dailydose));
initdose = [dosenmol; zeros(t1-1,1); zeros(t2,1); dosenmol; ...
    zeros(t3-1,1);0];

```

```

intake = initdose + dailydose;

```

```

initamt = [InitAmts(:); %nmol [D, 25, 125, 2425]
    initbase(:); %baseline fold change
    inirates(:); %synthesis and elimination rate tracking
    initdose(1)]; %initial gut dose

```

```

%%%%%%%%%%

```

```

refine = 4;

```



```

options = odeset('Refine',refine);

tout = tvec(1);
A = initamt';

%these will be called in multiple times, reduce number of passes
P = T.Pars;
Kin = T.Kinetics(1,:);
Kstruct = table2struct(Kin);

for i = 1:length(tvec)-1 %counts the number of terminal events
    %use vectorized PBPK file with the structure for kinetics
    helper = @(t, amt) PBPK(t,amt,T,Kstruct);
    [t,Aout] = ode15s(helper,[tvec(i) tvec(i+1)],initamt,options);
    nt = length(t);
    vec = 2:nt;
    tout = [fliplr(tout); t(vec)];
    A = [A; Aout(vec,:)]; %nmol in plasma, total amount

    % Set the new initial conditions, introducing input every 24ish hours
    initamt = Aout(nt,:);
    initamt(end) = initamt(end) + intake(i+1); %to GI reserve
end

%% now extrapolate total amount in venous plasma.

serum.time = tout(1:end)/24;

Vplas = P.Vol(9); %volume of plasma
totserum = A(:,9:9:(9*4))./(Vplas); %nmol/L in plasma

serum.vD = totserum(:,1);
serum.v25 = totserum(:,2);
serum.v125 = totserum(:,3);
serum.v24 = totserum(:,4);

end %----- End Main Function -----

%% Initial Concentrations
function [P] = initconcs(T,g)

P = T.Pars;
K = T.Kinetics(1,:);

%P = parameter table
%g = group number

[D] = readdata; % D = table of data

WD = 384.64; %weight of vitamin D in g/mol = Da
W25 = 400.64; %g/mol
W125 = 416.6; %g/mol
W24 = 416.64; %g/mol
MolWeights = [WD; W25; W24; W125];
CF = 1e3./MolWeights; %conversion from ug/L to nmol/L

%now take the average of the first and second time points because there is
%a dip between these points for all cases (0-30min)
mpvD = nanmean(D.medVitD3{g}(1:2)).*CF(1); % nmol/L
mpv25 = nanmean(D.med25{g}(1:2)).*CF(2); %
mpv2425 = nanmean(D.med2425{g}(1:2)).*CF(3); %
mpv125 = nanmean(D.med125{g}(1:2)).*CF(4); %

```

```

%% now use ideas of Ramakrishnan2016 (A2): C_(T,base)/C_(Plas,base) = K_T
%% % % => C_(T,base) = K_T .*C_(Plas,base)
P.ConcD = P.PC_D.*mpvD; %this will be used later in ENDOG in the PBPK file
P.Conc25 = P.PC_25.*mpv25;
P.Conc125 = P.PC_125.*mpv125;
P.Conc2425 = P.PC_2425.*mpv2425;
%
% allow for dynamic adipose partition coefficient based on total 25D conc
% in venous
if K.AdiPC
    venconc = [P.ConcD(end) P.Conc25(end)];
    AdiPC = DynamAdi(K,venconc); %pass in venous concentrations
    P.ConcD(5) = mpvD*AdiPC(1);
    P.Conc25(5) = mpv25*AdiPC(2);
end

P.AmtD = P.ConcD.*P.Vol; % = nmol/L * L = nmol for amt in
P.Amt25 = P.Conc25.*P.Vol;
P.Amt125 = P.Conc125.*P.Vol;
P.Amt2425 = P.Conc2425.*P.Vol;

end

```

```

% Supplemental Material: Drug Met Disp, DMD-AR-000609
% Title: Physiologically Based Pharmacokinetic Modeling of Vitamin D3 and Metabolites in
Vitamin D Insufficient Patients
% Authors: Colton W. Sawyer, Stacey M. Tuey, Raymond E. West III, Thomas D. Nolin,
Melanie S. Joy
% Code title: Pars

```

```
function [T] = Pars
```

```
%output: T = structure of parameters for model, fields:
```

```

%       T.Pars: Physio table
%       T.Kinetics: Kinetic parameter table
%       T.Misc: Misc structure
%       T.C: Conversion structure
%
%-----

```

```
%% Table setup
```

```
% method of plasma rate flow taken from Ramakrishnan2016
```

```

VarNames = {'Vol', 'Rate', 'PC_D', 'PC_25', 'PC_125', 'PC_2425'};
Doublecell = cellfun(@char, repmat({'double'},1,length(VarNames)), 'Un', 0);

```

```
VarTypes = [Doublecell(:)];
```

```
compt = {'Liv', 'Kid', 'Brain', 'Int', 'Adi', 'RP', 'SP', 'Heart', 'Plasma'};
```

```
TblSz = [length(compt), length(VarNames)];
```

```

P = table('Size', TblSz, 'VariableNames', VarNames, 'VariableTypes', ...
         VarTypes, 'RowNames', compt);
%-----

```

```
%% physio parameters (Jones 2013)
```

```
BW = 70; %in kg
```

```
CO = 5.600; %L/min %Davies1993; %15*(BW)^(0.74); %L/hr (Brown 1997)
```

```
BP = .55; % blood:plasma ratio = (1-.45)
```

```
%%%%%%%%%% VOLUMES in L (assuming BW = 70kg) (Davies1993)
```

```
VPlasma = 3.000; %
```

```
VL = 1.690; %Liver
```

```
VK = 0.280; %Kidney
```

```
VBr = 1.450; %brain
```

```
VHt = 0.310; %heart
```

```
% VB = 14.3; %bone ----- density = 1.92 g/cm^3
```

```
VSP = 42.800; %Muscle + skin
```

```
VAdi = 10.000; %adipose ----- density = 0.916 g/cm^3
```

```
VI = 0.637; %Intestine --- 0.91%
```

```
%Brown, Delp, Lindstedt, Rhomberg, & Beliles, 1997 in Noh2020
```

```
VRP = BW - (VPlasma + VL + VK + VBr + VHt + VSP + VAdi + VI); %RP tissue
```

```
P.Vol = [VL; VK; VBr; VI; VAdi; VRP; VSP; VHt; VPlasma]; %this is in L=kg
```

```
%%%% BLOOD FLOW RATES in L/min, adjusted to plasma flow later (Davies1993)
```

```
qL = 0.300; %liver % this is the hepatic flow rate
```

```
qK = 1.240; %kidney
```

```
qBr = 0.700; %brain
```

```
qHt = 0.240; %heart
```

```
qSP = 1.050; %muscle + skin
```

```
qAdi = 0.260; %adipose
```

```
qI = 1.150; %intestine + spleen --- portal vein
```

```
qRP = CO - (qL + qK + qBr + qHt + qSP + qAdi + qI); %rapidly perfused
```

```
qPlasma = CO; %L/min blood flow - will convert to plasma flow rate next
```

```
%plasma flow rates since D binds only in plasma with VDP (vitamin d binding
```

```

%protein) Ramakrishnan2016
% converted to L/hr %% plasma flow rate
P.Rate = [qL; qK; qBr; qI; qAdi; qRP; qSP; qHT; qPlasma].*(60.*BP);

%-----
%% tissue:plasma PCs -- function below
[P,M] = ParCoeffs(P);
%[L;K;Br;Int]
mousewts = [1.75; 0.32; 0.506; 0.36]*1e-3; %mL -> L mouse vol Noh2020
humanwts = P.Vol(1:4); %L human tissue volume
C.manmouse = humanwts./mousewts;
%-----
%% kinetics table

VarNames = {'Kinetic', 'LB', 'UB'};
Doublecell = cellfun(@char, repmat({'double'},1,length(VarNames)), 'Un', 0);

VarTypes = Doublecell(:);

kins = {'VmaxD', 'KmD', ...%clearance of D to 25(OH)D - Ocampo-Pelland2016
'R27B1', ... %ratio for CYP27B1
'Rsyn24L', 'Rsyn24K', 'Rsyn24Br', 'Rsyn24I', ...%rate of synthesis for 24,25 from 25 by 24A1
'fkCL2425', ... %free fraction clearance of 24,25 from kidney
'IC50', 'EC50L', 'EC50K', 'EC50Br', 'EC50I', ... %(Ramakrishnan2016)
'Imax', 'EmaxL', 'EmaxK', 'EmaxBr', 'EmaxI', ... %(Ramakrishnan2016)
'g1L', 'g1K', 'g1Br', 'g1I', ... %gamma 1 (Ramakrishnan2016, Noh2020)%
'kout24L', 'kout24K', 'kout24Br', 'kout24I', ... %kin for 24A1 (Ramakrishnan2016)
'fLCL125L', 'fKCL125K', 'fBrCL125Br', 'fICL125I', ... %CL_int,met,T for 24A1 (Ramakrishnan2016)
'g2', ...%kin27', .
'kout27', ... %CYP27B1 (Ramakrishnan2016, Noh2020)
'Dbase', ...%(Ocampo-Pelland2016)
'ka', ... %endogenous base, gut absorb (Ocampo-Pelland2016)
'maxVmax', 'hVmax', 'D50', 'VarVmax', ...% variable vmax values
'Adimax', 'hAdi', 'Adi50', ...% variable adipose PC
'Adimax25', 'hAdi25', 'Adi50_25', 'AdiPC' ...% variable adipose PC
};

%
TblSz = [length(kins),length(VarNames)];

K = table('Size', TblSz, 'VariableNames', VarNames, 'VariableTypes', ...
VarTypes, 'RowNames', kins);

%%
K.Kinetic = [1-2
1.62; %VmaxD %Ocampo-Pelland2016
5.90; %KmD
%% 3
17.7; %R27B1 %endog synth of 1,25 from 25D;
%% 4-7
%% rate of synthesis of 24,25 from 25 by 24A1
1.20; %Rsyn24L
2.43; %Rsyn24K
13.3; %Rsyn24Br
15.4; %Rsyn24I
%% 8
%% free fraction clearance of 24,25 from kidney
990; %fkCL2425; assume units are pmol/hr
%%
%%Inhib/Stim Ramakrishnan2016 fitted values (TM) model
%% UNITS pmol/kg tissue
%% 9-13

```

```

673; %IC50
2600; %EC50L
143;%113; %EC50K
97;%235; %EC50Br
2640; %EC50I
%% Imax and Emax %Noh2020 combined doses
%% 14-18
1; %Imax %upper bound = 1
122; %EmaxL
25; %EmaxK
22; %EmaxBr
675; %EmaxI
%%
%% 19-22
%% Hill constant for indirect response function of 24A1
3.05; %g1L %Ramakrishnan2016 TM model
3.59; %g1K %Ramakrishnan2016
1.12; %g1Br %Ramakrishnan2016
4.16; %g1I %Ramakrishnan2016
%% turnover rates of CYP24A1
%% 23-26
0.047; %kout24L
0.28; %kout24K
0.86; %kout24Br
0.047; %kout24I
%%
%%-----
%% free fraction CL_int,met,T (Noh2020)-- units mL/hr/g tissue
%% tissue - will adjust in PBPK code to account for man
%% vs mouse tissue
%% 27-30
0.00057; %fLCL125L
0.066; %fKCL125K
0.0083; %fBrCL125Br
0.0028; %fICL125I
%% -----
%%
%% 31-32
%% parameters for CYP27B1
2.7; %g2 , %Ramakrishnan2016 TM model
0.245; %kout27b1
%% 33-34
%% Dbase and gut absorption (Ocampo-Pelland2016)
2.05; %Dbase, nmol/L
0.32; %gut absorption 1/hr
%% 35-38
%% Variable Vmax
5.76; %max Vmax (in def state)
2.27; % exponential coeff for dynamic kid PC
8.34;%; %50% concentration of D nmol/L
1;%Vmax switch
%% 39-41
%% variable adipose PC D
9.9; %adimax
2.12; %hadi
8.97; %Adi50 D (nmol/L)
%% 42-45
%% variable adipose PC 25
1.64; %adimax25
.96; %hadi25
117; %Adi50 25 (nmol/L)
1;%adipose switch
];

```

```

%% -----
K = rows2vars(K);

K.Properties.RowNames = K{:,1};
K(:,1) = []; %delete first row which is unnecessary now

K{2,:} = .5*K{1,:};
K{3,:} = 2*K{1,:};
    K{[2 3],14} = 1; %Imax
    K{[2 3],38} = 1; %variable vmax switch
    K{[2 3],45} = 1; %variable adiPC switch

%-----
%% Now package everything into a structure
T.Pars = P;
T.Kinetics = K;
T.Misc = M;
T.C = C;
T.C.optvec = [1:13, 15:37, 39:44]; %leave out imax, vmax switch
T.C.defopt = [3:8 35:37 39:44];
end

%% -- ParCoeffs function -----
function [P,M] = ParCoeffs(P)

% P = parameter table; will update with partition coefficients
% M = structure with misc data

%% Fraction unbound in plasma
%build table M for miscellaneous data here
%convert everything to mol/L
WD = 384.64; %weight of vitamin D in g/mol = Da
W25 = 400.64; %g/mol
W125 = 416.6; %g/mol
W24 = 416.64; %g/mol

MolWeights = [WD; W25; W125; W24];
freefrac = [0.07; 0.03; 0.43; 0.03]*1e-2;

M.fup = freefrac;
M.MolWeights = MolWeights;

%-----
%% Partition coefficients
% %
% compt = {'Liv','Kid','Brain','Int','Adi','RP','SP','Heart', 'Plasma'};
%
% now use comparison of concentrations of D and 25D in different
% compartments
%kid, liv, brain, intestines ratios from Ramakrishnan2016, assume
% heart=liver
BlumSerummol = 7.8;%nmol/kg Blum et al 2008. Endocrine (33)
BlumSerumug = BlumSerummol*WD/1e3; %ug/kg

%data taken from Jakobsen2007, Br J Nutr (98) in ug/kg
P.PC_D([1:4 8]) = (2.67/BlumSerumug)*[1 2 1/3 8/3 1]; %[0.8895]
P.PC_D(5) = (7.47/BlumSerumug); %[2.4885]; adipose
P.PC_D(6) = (1.11/BlumSerumug); %[0.3698]; rapidly (lean)
P.PC_D(7) = (2.99/BlumSerumug); %[0.9961]; Slowly perfused (skin)

JSerum25D = 18.1; %ug/kg
P.PC_25([1:4 8]) = (3.95/JSerum25D) * [1 2 1/3 8/3 1]; %[0.2182]
P.PC_25(5) = (7.47/JSerum25D); % [0.4127]; adipose

```

```
P.PC_25(6) = (0.89/JSerum25D);% [0.0492]; rapidly (lean)
P.PC_25(7) = (3.18/JSerum25D);%[0.1757]; Slowly perfused (skin)

%[L,K,Br,I], liv = heart
P.PC_125([1:4 8]) = [0.15; 0.34; 0.05; 0.4; 0.15]; %Ramakrishnan2016
P.PC_125(5) = P.PC_25(5)./P.PC_25(7) *0.335; %ratio of 25D for adipose/SP
P.PC_125(6) = P.PC_25(6)./P.PC_25(2) *0.34; %assume RP = ratio 25D RP/kid
P.PC_125(7) = 0.335; %SP = peripheral compartment in Noh2020

P.PC_2425 = P.PC_125; %comparable molecular weights

%set plasma PCs = 1 because ratio of plasma/plasma = 1
P.PC_D(9) = 1;
P.PC_25(9) = 1;
P.PC_125(9) = 1;
P.PC_2425(9) = 1;

end
```

```

% Supplemental Material: Drug Met Disp, DMD-AR-000609
% Title: Physiologically Based Pharmacokinetic Modeling of Vitamin D3 and Metabolites in
Vitamin D Insufficient Patients
% Authors: Colton W. Sawyer, Stacey M. Tuey, Raymond E. West III, Thomas D. Nolin,
Melanie S. Joy
% Code title: PBPK

```

```

function [dA] = PBPK(t,A,T,K)

```

```

% A = amt vector
% T = paramter structure
% K = Kinetic structure
% dA = diff eq vector (see below)

```

```

P = T.Pars;
M = T.Misc;

```

```

% differential equations have form:

```

```

% dA = [ vit D (11)
%       25(OH)D (11)
%       1,25(OH)2D (11)
%       24,25(OH)2D (11)
%       CYP27B1 fold change (1)
%       CYP24A1 fold change (4)
% %     Vmax 2R1 eqn (1)
%       synthesis (3)
%       synthesis via 24A1 (4)
%       elimination via 24A1 (4)
%       entry (1)
%     ];
% total equations = 62

```

```

%% Separate out the amounts into compounds

```

```

A_D = A([1:9]+0*9); %ug
A_25 = A([1:9]+1*9);
A_125 = A([1:9]+2*9);
A_2425 = A([1:9]+3*9);

```

```

FC27 = A(4*9+1); %37
FC24 = A([4*9+2]:[4*9+5]); %38-41

```

```

Agut = A(end);

```

```

% free, unbound portion
fup25 = M.fup(2);

```

```

%% build the concentrations

```

```

% C_{vT} = C_T/(Kp) %Kp is the tissue:plasma PC
% Ramakrishnan2016
C_D = (A_D./P.Vol) ./ (P.PC_D); %nmol/L
C_25 = (A_25./P.Vol) ./ (P.PC_25);
C_125 = (A_125./P.Vol) ./ (P.PC_125);
C_2425 = (A_2425./P.Vol) ./ (P.PC_2425);

```

```

%
if K.AdiPC % allow for dynamic adipose partition coefficient based on total 25 conc in
venous

```

```

    venconc = [C_D(end) C_25(end)];
    AdiPC = DynamAdi(K,venconc); %pass in venous concentrations
    C_D(5) = A_D(5)./(P.Vol(5).*AdiPC(1)); %adjust adipose D partition coefficient
    C_25(5) = A_25(5)./(P.Vol(5).*AdiPC(2)); %adjust adipose 25 partition coefficient

```

```

end

```



```

ClivD = C_D(1); % liver D
C25 = C_25(1:4);
C125 = C_125(1:4); %total concentration
Kkid2425 = C_2425(2); %kidney 24,25
R27B = K.R27B1; %L/hr

%rate of synthesis of 24,25 from 25 by 24A1
Rsyn24 = [K.Rsyn24L; K.Rsyn24K; K.Rsyn24Br; K.Rsyn24I];

fkCL2425 = K.fkCL2425; %free fraction clearance of 24,25 from kidney (nmol/hr);

kout27 = K.kout27;
kout24 = [K.kout24L; K.kout24K; K.kout24Br; K.kout24I];
kin27 = kout27;
kin24 = kout24;

%gamma 1 terms for CYP24A1 Ramakrishnan2016
g1 = [K.g1L; K.g1K; K.g1Br; K.g1I];
g2 = K.g2; %hill coefficient

%free fraction unbound clearance terms %Noh2020 (mL/hr in mouse tissue -> L/hr in human
tissue)
freeCLT = [K.fLCL125L; K.fKCL125K; K.fBrCL125Br; K.fICL125I].*P.Vol(1:4); %scaled
according to tissue weight

CFpmolnmol = 1e-3; %conversion factor from pmol/L to nmol/L; 1 pmol = 1e-3 nmol

IC50 = K.IC50.*CFpmolnmol; %CONCENTRATION OF 1,25, now in nmol/L
Imax = K.Imax; %max effect - no units

EC50 = [K.EC50L; K.EC50K; K.EC50Br; K.EC50I].*CFpmolnmol; % now in concentration of 1,25
(nmol/L)
Emax = [K.EmaxL; K.EmaxK; K.EmaxBr; K.EmaxI]; %max effect, no units
%%% -----

ka = K.ka; %absorption into the gut; 1/hr

%%%
%% BASIC EQUATIONS
% rates are all plasma flow rates (already multiplied by BP)

Qlivtot = P.Rate(1) + P.Rate(4); %hepatic artery (liver) plus portal vein (intestine)

%vit D
dAD(1) = P.Rate(1).*C_D(9) + P.Rate(4).*C_D(4) - Qlivtot.*C_D(1); %liver with portal vein
dAD(2:8) = P.Rate(2:8).*(C_D(9) - C_D(2:8)); % main tissues
dAD(9) = Qlivtot.*C_D(1) + sum(P.Rate([2:3 5:8]).*C_D([2:3 5:8])) - C_D(9).*P.Rate(9); %
plasma not including intestine

%25(OH)D
dA25(1) = P.Rate(1).*C_25(9) + P.Rate(4).*C_25(4) - Qlivtot.*C_25(1); %liver with portal
vein
dA25(2:8) = P.Rate(2:8).*(C_25(9) - C_25(2:8)); % main tissues
dA25(9) = Qlivtot.*C_25(1) + sum(P.Rate([2:3 5:8]).*C_25([2:3 5:8])) - C_25(9).*P.Rate
(9); %plasma not including intestine

%1,25(OH)2D
dA125(1) = P.Rate(1).*C_125(9) + P.Rate(4).*C_125(4) - Qlivtot.*C_125(1); %liver with
portal vein

```

```

dA125(2:8) = P.Rate(2:8).*(C_125(9) - C_125(2:8)); % main tissues
dA125(9) = Qlivtot.*C_125(1) + sum(P.Rate([2:3 5:8]).*C_125([2:3 5:8])) - C_125(9).*P.
Rate(9); %plasma not including intestine

%24,25(OH)2D
dA2425(1) = P.Rate(1).*C_2425(9) + P.Rate(4).*C_2425(4) - Qlivtot.*C_2425(1); %liver with
portal vein
dA2425(2:8) = P.Rate(2:8).*(C_2425(9) - C_2425(2:8)); % main tissues
dA2425(9) = Qlivtot.*C_2425(1) + sum(P.Rate([2:3 5:8]).*C_2425([2:3 5:8])) - C_2425(9).
*P.Rate(9); %plasma not including intestine

%%%%%%%%%%%%%%
%% now the fold change terms

%Ramakrishnan2016
%27B1 terms
ICterm = 1 - (Imax.*C_125(2).^g2)./(IC50.^g2 + C_125(2).^g2);
dFC27 = kin27.*ICterm - kout27.*FC27;

%24A1 terms
ECterm = 1 + (Emax.*C125.^g1)./(EC50.^g1 + C125.^g1);
dFC24 = kin24.*ECterm - kout24.*FC24;

%DE fold change terms
dFC = [dFC27; dFC24];
dFC = dFC(:);
%% Now the metabolism terms and other rates
%units are nmol/hr
Gut = ka*Agut; %absorption from the gut based on gut amount

%% This section from Ocampo-Pelland2016

if K.VarVmax %variable Vmax
    [VmaxD] = VaribVmax(K, ClivD); %function below
else
    VmaxD = K.VmaxD; %nmol/h; Ocampo-Pelland2016
end

KmD = K.KmD; %nmol/L; %Ocampo-Pelland2016
Dbase = K.Dbase;

ENDOG = (VmaxD.*Dbase./ (KmD + Dbase))*Dbase; %endogenous synth of D in nmol/hr
CLD25 = (VmaxD.*ClivD./ (KmD + ClivD))*ClivD;

%% other synthesis terms
S_27B1 = C25(2)* R27B * FC27*fup25; %25-> 1,25 by 27B1 in kidney - first order rate
(Ocampo-Pelland2016)

S_24A1 = C25.*fup25 .*Rsyn24 .*FC24; %free amt

%% elimination terms
E_24A1 = C125 .* freeCLT .* FC24; %free amts

E_2425 = Ckid2425 * fkCL2425.*CFpmolnmol; %free amt %elimination of 24,25 from the
kidney, first order process

%% keep these for tracking purposes
dKin = [ENDOG; CLD25; S_27B1; S_24A1; E_24A1];

dKin = dKin(:);
%% Now adjust basic equations to include metabolism and other rates
% adjust D (1-9)

```

```
dAD(1) = dAD(1) + ENDOG - CLD25; %change in liver compt
dAD(4) = dAD(4) + Gut; %intestine
```

```
% adjust 25 (10-18)
```

```
dA25(1) = dA25(1) - S_24A1(1) + CLD25; %liver
dA25(2) = dA25(2) - S_27B1 - S_24A1(2); %kidney
dA25(3) = dA25(3) - S_24A1(3); %brain
dA25(4) = dA25(4) - S_24A1(4); %intestines
```

```
% adjust 1,25 (19-27)
```

```
dA125(1) = dA125(1) - E_24A1(1); %liver
dA125(2) = dA125(2) + S_27B1 - E_24A1(2); %kidney
dA125(3) = dA125(3) - E_24A1(3); %brain
dA125(4) = dA125(4) - E_24A1(4); %intestines
```

```
% adjust 24,25 (28-36)
```

```
dA2425(1) = dA2425(1) + S_24A1(1); %liver
dA2425(2) = dA2425(2) + S_24A1(2) - E_2425; %kidney
dA2425(3) = dA2425(3) + S_24A1(3); %brain
dA2425(4) = dA2425(4) + S_24A1(4); %intestines
```

```
dAmt = [dAD(:); dA25(:); dA125(:); dA2425(:)];
dAmt = dAmt(:);
```

```
%adjust gut
```

```
dA_gut = -Gut; %absorption from gut
```

```
%% now put it all together
```

```
dA = [dAmt; %1-36
      dFC; %37-41
      dKin; %42-52
      dA_gut]; %53
```

```
dA = dA(:); %ensure column vectors
```

```
end
```

```
%----- End Main Function -----
```

```
%% Variable Vmax function
```

```
function [Vmaxout] = VaribVmax(K, Dconc)
```

```
%
```

```
maxVmax = K.maxVmax;
hVmax = K.hVmax;
D50 = K.D50;
```

```
Vmaxout(1) = maxVmax./( 1 + (Dconc ./ D50).^hVmax);
```

```
end
```

Supplemental Material: *Drug Met Disp, DMD-AR-000609*

Title: Physiologically Based Pharmacokinetic Modeling of Vitamin D₃ and Metabolites in Vitamin D Insufficient Patients

Authors: Colton W. Sawyer, Stacey M. Tuey, Raymond E. West III, Thomas D. Nolin, Melanie S. Joy

Supplemental Material

S.1 Partition Coefficients

Human distribution data for vitamin D₃ and metabolites are limited [Blum et al., 2008; Didriksen et al., 2015; Mawer et al., 1972, Ramakrishnan et al., 2016]. To approximate the tissue:plasma partition coefficient (PC) for humans for VitD₃ and 25D₃, we used the ratio between tissue and plasma concentrations in pigs from [Blum et al., 2008, Jakobsen et al., 2007]. This study further assumed sufficient similarity between pigs and humans [Upton, 2008] and set the human tissue:plasma PCs equal to pig tissue:plasma PCs. These PCs were taken to be the ratio between tissue and serum concentrations. For VitD₃, as Jakobsen et al. [2007] did not measure serum VitD₃ concentrations, we used the mean serum VitD₃ concentration from [Blum et al., 2008] to find the PCs. Rapidly perfused tissue was set equal to the “lean” compartment and slowly perfused tissue was set equal to the “skin” compartment in [Jakobsen et al., 2007]. As compound amounts in the brain, intestines, or kidney were not given, this study assumed these tissues had PCs equal to the ratio of the PCs in [Ramakrishnan et al., 2016] with respect to the liver. The heart PC was set equal to the liver PC. Note the adipose tissue:plasma PC for

VitD₃ and 25D₃ was allowed to vary in this study based on the concentrations of VitD₃ and 25D₃ using Equations (5) and (6), respectively, in the main text.

For PCs for 1,25D₃ and 24,25D₃, this study used mouse PCs from [Ramakrishnan, et al., 2016] with the assumption that the heart and liver had identical PCs. The PC for the rapidly perfused compartment was set to the ratio of the PC of 25D₃ in the rapidly perfused to the kidney multiplied by the given PC in the kidney in [Ramakrishnan et al., 2016]. The adipose PC was constant and set to the ratio of the PC of 25D₃ in the adipose to the slowly perfused compartment multiplied by the given PC in the peripheral compartment in [Ramakrishnan et al., 2016]. The slowly perfused compartment for 1,25D₃ was set to the peripheral compartment. Due to the similar molecular size, PCs for 24,25D₃ were assumed equal to 1,25D₃. All PCs can be found in Supp. Table S2.

S.2 Supplemental References

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- Mawer E, Backhouse J, Holman C, Lumb G, and Stanbury S (1972) The distribution and storage of vitamin D and its metabolites in human tissue. *Clin Sci* 43:413–31.
- Ramakrishnan V, Yang QJ, Quach HP, Cao Y, Chow EC, Mager DE, and Pang KS (2016) Physiologically-based pharmacokinetic-pharmacodynamic modeling of 1 α , 25-dihydroxyvitamin D3 in mice. *Drug Metab Dispos* 44:189–208.
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S.3 Supplemental Tables

Table S1: Physiological Parameters for the PBPK model for vitamin D₃ and metabolites

Parameter	Description	Value	Units	Reference
Miscellaneous physiological parameters				
BW	Body weight ^a	70	kg	[Davies and Morris, 1993]
Q_{CO}	Cardiac Output	5.6	L·m ⁻¹	[Davies and Morris, 1993]
BP	Blood:plasma ratio	0.55 ^b	Unitless	[Ramakrishnan et al., 2016]
Volumes				
V_{adi}	Adipose	10.0	L	[Davies and Morris, 1993]
V_{br}	Brain	1.45	L	[Davies and Morris, 1993]
V_h	Heart	0.31	L	[Davies and Morris, 1993]
V_i	Intestine ^c	0.637	L	[Davies and Morris, 1993]
V_k	Kidney	0.28	L	[Davies and Morris, 1993]
V_l	Liver	1.69	L	[Davies and Morris, 1993]
V_{rp}	Rapidly perfused ^d	9.83	L	[Davies and Morris, 1993]
V_{sp}	Slowly perfused ^e	42.8	L	[Davies and Morris, 1993]
V_p	Plasma	3.0	L	[Davies and Morris, 1993]
Blood flow rates				
Q_{adi}	Adipose	8.58	L·m ⁻¹	[Davies and Morris, 1993]
Q_{br}	Brain	23.1	L·m ⁻¹	[Davies and Morris, 1993]
Q_h	Heart	7.92	L·m ⁻¹	[Davies and Morris, 1993]
Q_i	Intestine	37.95	L·m ⁻¹	[Davies and Morris, 1993]
Q_k	Kidney	40.9	L·m ⁻¹	[Davies and Morris, 1993]

Q_l	Liver ^f	9.9	$L \cdot m^{-1}$	[Davies and Morris, 1993]
Q_{rp}	Rapidly perfused	21.78	$L \cdot m^{-1}$	[Davies and Morris, 1993]
Q_{sp}	Slowly perfused	34.65	$L \cdot m^{-1}$	[Davies and Morris, 1993]

^a Reference body weight. Organ density is assumed to be 1 kg = 1 L

^b Plasma flow rates assumed a hematocrit value of 0.45 as vitamin D₃ and its metabolites are primarily bound to plasma compounds

^c Intestine were assumed to comprise 0.91% of total body weight [Brown et al., 1997]

^d Richly perfused tissues included adrenal, lungs and thyroid. This compartment served as a mass balance compartment, namely $BW - \sum_c V_c$ for other compartments c

^e Poorly perfused tissues were set to the sum of muscle and skin

^f Hepatic flow rate

Table S2: Unitless partition coefficients, $P_{T:plasma}$, for the PBPK model of VitD₃ and metabolites.

Partition coefficients were calculated as discussed in the Supplemental Materials.

Partition Coefficients	Description	D_3	$25D_3$	$1,25D_3$	$24,25D_3^a$
$P_{adi:p}$	Adipose/plasma	2.49	0.41	0.78	0.78
$P_{br:p}$	Brain/plasma	0.30	0.07	0.05	0.05
$P_{h:p}$	Heart/plasma	0.89	0.22	0.15	0.15
$P_{i:p}$	Intestine/plasma	2.37	0.58	0.4	0.4
$P_{k:p}$	Kidney/plasma	1.78	0.43	0.34	0.34
$P_{l:p}$	Liver/plasma	0.89	0.22	0.15	0.15
$P_{rp:p}$	Rapidly perfused/plasma	0.37	0.05	0.04	0.04
$P_{sp:p}$	Slowly perfused/plasma	1.00	0.18	0.34	0.34

D_3 , VitD₃; $25D_3$, 25(OH)D₃; $1,25D_3$, 1,25(OH)₂D₃; $24,25D_3$, 24,25(OH)₂D₃,

^aAssume $1,25D_3$ and $24,25D_3$ partition coefficients are identical