

Physiologically-based pharmacokinetic modeling and simulation approaches: Best practices for regulatory applications related to locally-acting generic drugs

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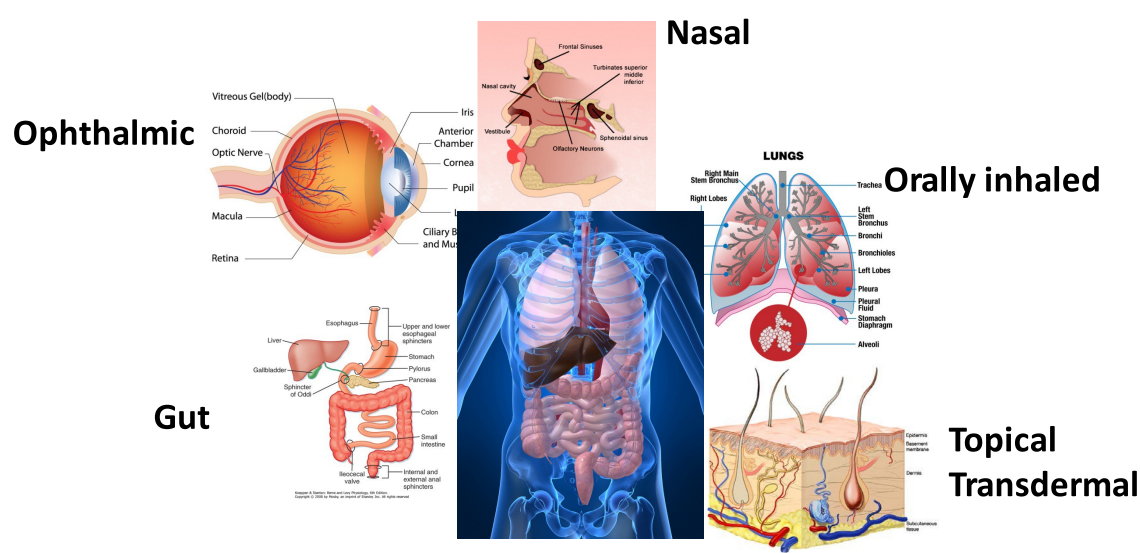
Overview



- Bioequivalence (BE) of locally-acting drug products
- Physiologically-based pharmacokinetic (PBPK) modeling Best practices on:
 - \circ Model development
 - \circ Model performance assessment
 - o Virtual bioequivalence (VBE) studies
 - \odot Reporting and documentation
- Case example: approved Abbreviated New Drug Application (ANDA) for a complex topical drug product

Locally-acting drug products

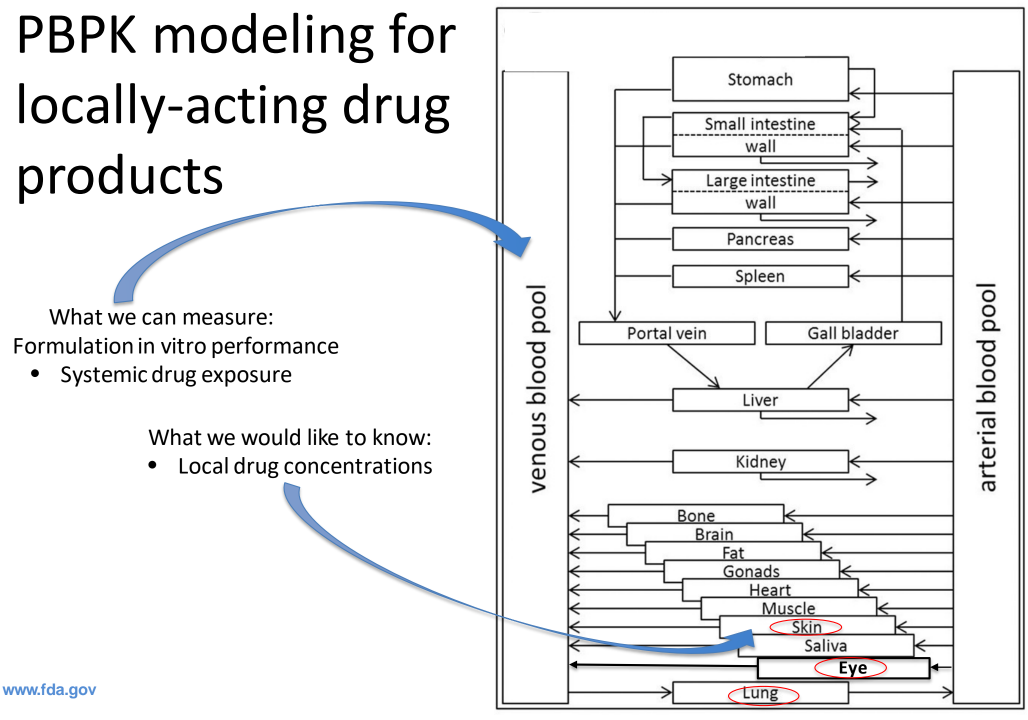




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products

Applications of PBPK modeling



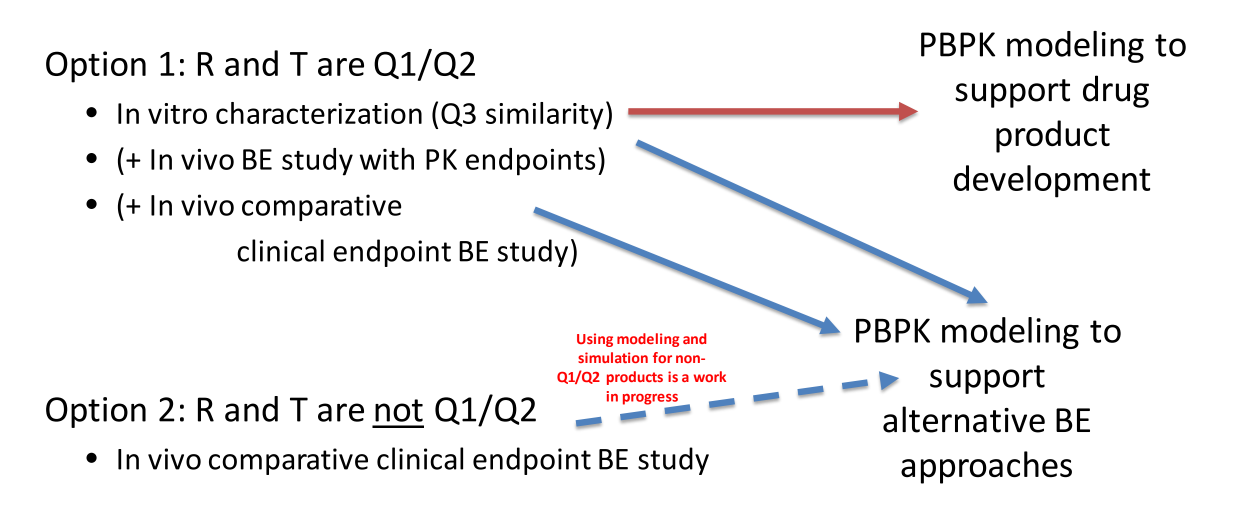
New investigational agents:

- Candidate selection in preclinical phase
- Animal-to-human extrapolation studies
- Drug-drug interaction studies
- Early formulation selection studies
- Assess disease impact
- Dose adjustment for specific populations (organ impairment, pediatric population, etc.)

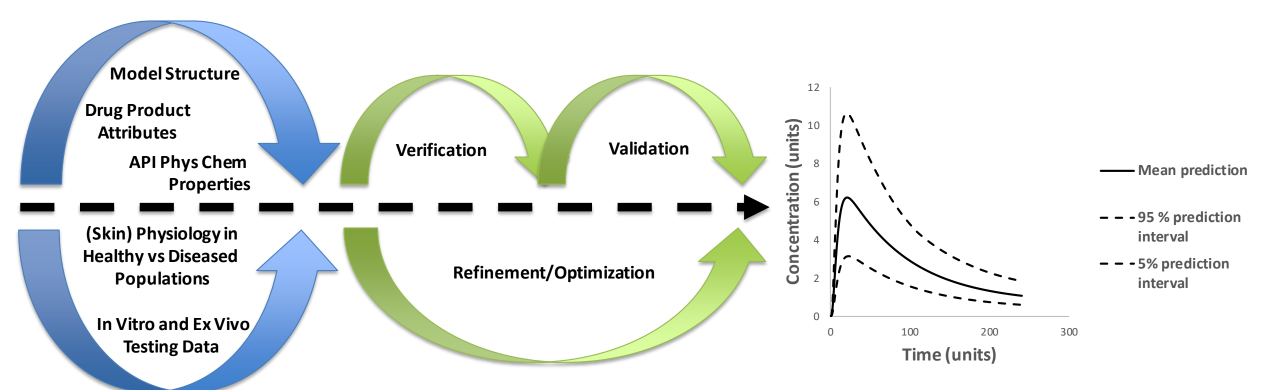
Generic drug products:

- Product-specific guidance (PSG) development
 - Alcohol dose dumping
 - Risk assessment for change in drug release mechanism
- Alternative approaches for demonstrating bioequivalence (BE)
 - o In vitro testing in lieu of in vivo BE studies
 - \circ Locally-acting drug products
- Extrapolate BE assessments in subpopulations
 - Disease
 - Age
- Drug product development
 - "Safe space" for critical attributes of drug products (dissolution specifications)

PSGs for complex locally-acting drug products



PBPK modeling for locally-acting drug products



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Best practices: internal reporting and documentation



• Modeling analysis plan:

 Objectives, model input (assumptions) and reference sources for in vivo/in vitro data used

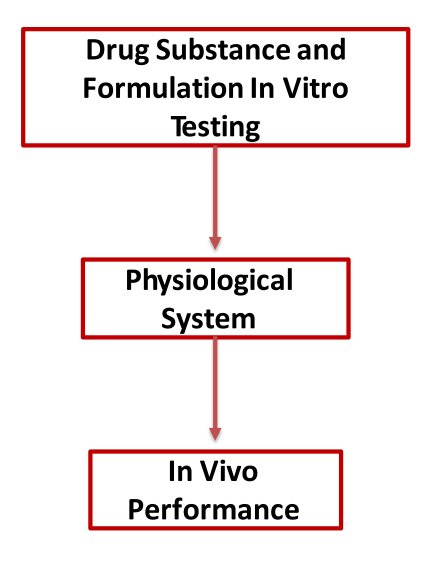
o "living document"

• Modeling analysis report:

 \odot Technical aspects of the modeling task

o Conclusions that may inform decisions should be clearly presented and discussed

Best practices: model development



- Model input (data)

 Good quality
 Low uncertainty
 Experimentally derived
 Biologically plausible
- Model structure justified and supported by data or sensitivity analysis
- Parameter/population variability
- Parameter correlations
- Disease state impacting BE assessment outcome



Best practices: model development

Model verification

- Model components:
 - Differential equations
 - \circ Code
 - \odot Engine for executing applications
- Biological plausibility:

• Physiology (route of administration and site of action)

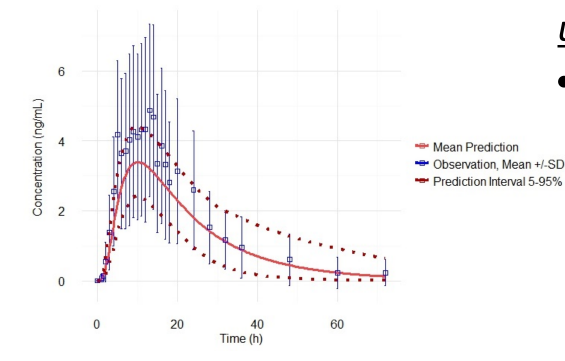
- Orug substance ADME properties
- Drug product attributes



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"Model verification should provide sufficient information to clearly demonstrate that the proposed PBPK model is <u>appropriate for the modeling</u> <u>purpose or question asked for the particular drug product and study</u> <u>population</u> and is robust enough to <u>respond to perturbations</u> in uncertain parameters"*





Model validation <u>for purpose (intended</u> <u>use) for drug product of interest</u>

- Observed data on systemic and local exposure:
 - \circ Good quality
 - \circ Clinically relevant scenarios
 - Drug product-specific
 - o Variability
- Acceptance criteria:
 - Regulatory impact of the decision

 \odot Established a priori and used consistently 12

Oxybutynin chloride extended release tablet (DITROPAN XL[®]), 15 mg given orally at fasting state*

* Tsakalozou et al. AAPS, November 2016, Denver CO.

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Platform performance assessment:

- Models developed under the same principles as the drug product of interest
 - o APIs of similar physicochemical properties and ADME characteristics
 - Dosage forms of variable compositional complexity for the same route of administration
 - \circ Not used for model development
- Models developed to capture physiology variables and their interplay with product attributes:
 - o Disease
 - \circ Patient population



Platform performance assessment

• Ocular PBPK:

• Validate model in animal species - interspecies extrapolation

In vitro or ex vivo data (site of action)

o In vivo systemic and local exposure data in humans

• Inhalation/nasal PBPK:

In vitro or ex vivo data (site of action)

o In vivo systemic and local exposure data in humans

• Dermal PBPK:

In vitro or ex vivo data (site of action)

 \odot In vivo systemic and local exposure data in humans



Considerations and challenges:

- Data availability for model validation, for establishing in vivo-in vitro relationships, for describing drug product-physiology interplay
- Systemic exposure does not reflect local concentrations
- Model validated only for the range of conditions of the dataset used

Best practices: model refinement



"middle-out" modeling approach involving parameter optimization Knowledge gaps?

- Fit-for-purpose model
- Model structural and numerical identifiability
- Parameter collinearity
- Additional model verification necessary

Best practices: model application



Virtual bioequivalence assessments

 Leverage models developed for R and T product to generate population predictions

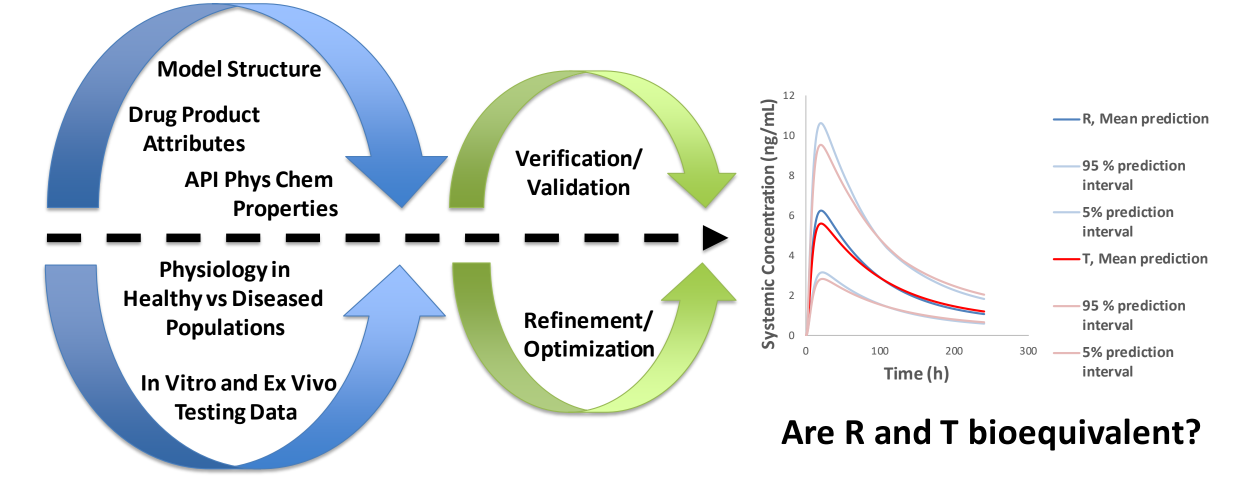
Identification of clinically relevant drug product attributes (sensitivity analysis)

• Considerations:

 Sources of variability incorporated into the model realistic population predictions

o Sample size impacted by highly variable pharmacokinetic behavior





Best practices: regulatory submission



Physiologically Based Pharmacokinetic Analyses — Format and Content Guidance for Industry August 2018 Clinical Pharmacology

III.	FORMAT AND CONTENT	.2
А.	Executive Summary	.2
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C.	Materials and Methods	.2
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	. Model Verification and Modification Model Application Discussion	

13 December 2018 EMA/CHMP/458101/2016 Committee for Medicinal Products for Human Use (CHMP)

Guideline on the reporting of physiologically based pharmacokinetic (PBPK) modelling and simulation

4. Reporting of PBPK modelling and simulation4
4.1. Objective and regulatory purpose4
4.2. Background information4
4.3. Qualification
4.4. Model parameters6
4.4.1. Assumptions
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4.5. Model development
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4.7.2. Evaluation of the predictive performance of the drug model
4.8. Results
4.9. Discussion of the regulatory application9

Best practices: regulatory submission

Highlights:

• Modeling and Simulation report:

o Model development and performance assessment are documented in detail

o Model assumptions and conclusions are data- and science-supported

- Model limitations are clearly stated and considered for simulations and outcome reporting
- Executable final model files, model input and output files and their description
- Submitted material updated as a result of interactions with the Agency

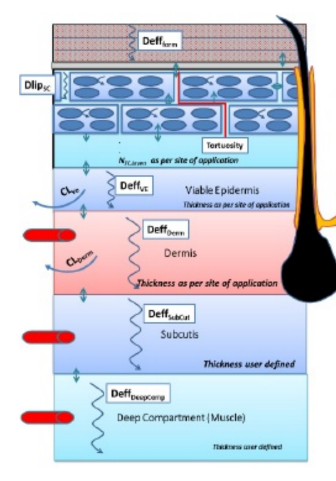
Take home messages



- PBPK models for locally-acting products can be used to support:
 - Development of a drug product prior approval
 - o Alternative BE approaches
- Model development is an intense and resource-demanding process due to:

 Complexity of the models and the drug products (remote target site)
 Limitations in data availability in model development and validation
- PBPK modeling supporting an ANDA: early interaction between industry and regulatory agency should be initiated - pre-ANDA meeting request program, GDUFA II

Dermal PBPK model supporting ANDA 211253 approval



Formulation (Gel, cream, lotions, poste, patch, ointments, etc.) Stratum Corneum (SC) * Define cell shape and size

- Cell membrane permeability
- Keratin bonding kinetics
 Tortuosity and diffusivity
- Hair follicle density and size
- Viable Epidermis (VE) Thickness, diffusivity Metabolism

Dermis

Thickness, diffusivity
 Metabolism, blood flow

Subcutis

- Thickness, diffusivity
- Blood flow

Deep Tissue

- Thickness, diffusivity
- Blood flow

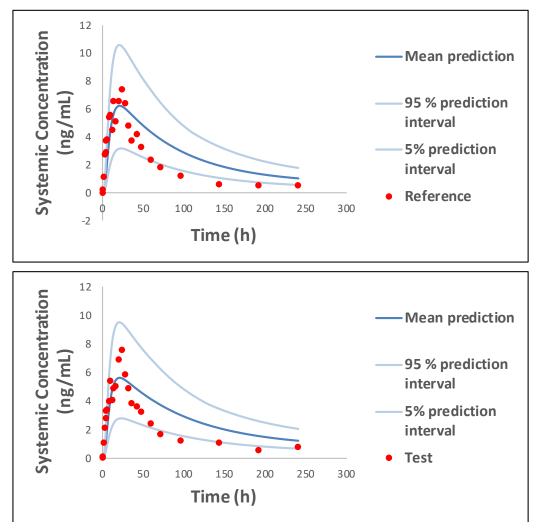
- Diclofenac sodium topical gel, 1%
- Alternative BE approach for a Q1/Q2/Q3 formulation: dermal PBPK model in lieu of an in vivo comparative clinical endpoint BE study
- Model development:
 - o API physicochemical properties
 - o API ADME properties
 - Formulation attributes for R and T drug products (viscosity, globule size, pH)

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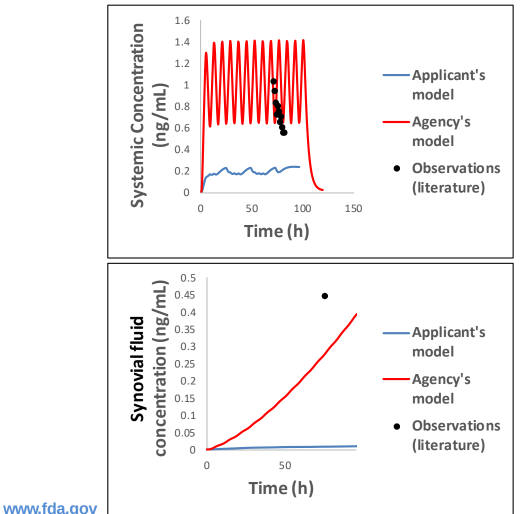
Dermal PBPK model supporting ANDA approval



- Platform performance assessment:
 - \circ >10 PBPK models for TDS and topical products
 - Multiple doses/product strengths and dosing regiments
 - \circ Satisfactory model performance
- Model performance assessment for diclofenac sodium topical gel, 1%:
 - Literature and application data on doses, product strengths, dosing regiments, routes of administration and local/systemic exposure data
 - $\,\circ\,$ Formulation attributes for R and T
 - \circ Good predictions of systemic exposure

R: Reference, T: Test, TDS: Transdermal Delivery System

Dermal PBPK model supporting ANDA approval



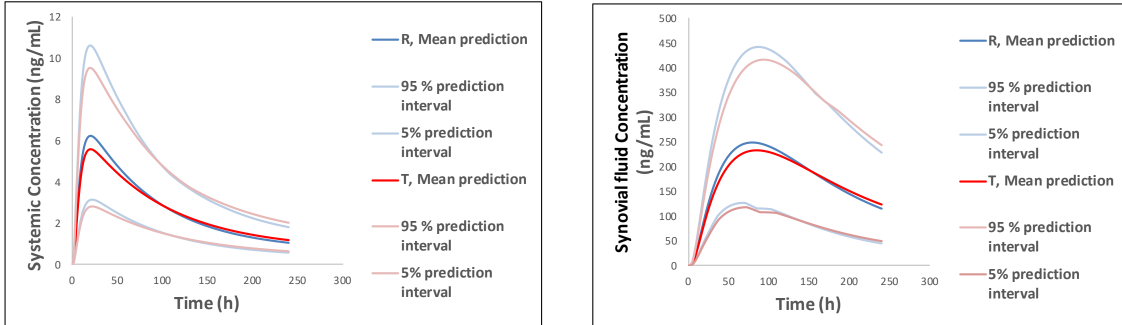
- Refined model to improve synovial fluid exposure predictions (by the Agency)
 - $\circ\,$ Protein binding in all skin layers
 - $\,\circ\,$ Drug product attributes updated
 - Partition coefficients modified leveraging observed local drug amounts

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Dermal PBPK model supporting ANDA approval

Conducted virtual BE assessments on predicted systemic and local exposure data

✓ R and T drug products were found bioequivalent





Conclusions



- PBPK models can be used to inform product development decisions and support alternative BE approaches for generic locally-acting drug products.
- Applicants are encouraged to follow best practices when developing PBPK models for generic locally-acting drug products as these are communicated by the Agency in guidances and other public forums.
- Applicants are encouraged to engage with the Agency early in their product development program by making use of the pre-ANDA meeting request program (GDUFA II) – case example of the approved ANDA for a complex topical drug product.



Acknowledgments

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Generic Drug User Fee Amendments: Regulatory

Grant/Contract	Institute	Grant or Contract No.
An integrated multiscale-multiphysics modeling framework for evaluation of generic ophthalmic drug products	CFD Research Corporation	HHSF223201810151C
GastroPlus OCAT model extension and validation	SimulationsPlus, Inc	HHSF223201810255P
A Multiscale Computational Framework for Bioequivalence of Orally Inhaled Drugs	CFD Research Corporation	HHS223201810182C
Characterization of key system parameters of mechanistic dermal PBPK models in various skin diseases and performance verification of the model using observed local and systemic concentrations	Simcyp, Ltd	1U01FD006521
Assessment of Transdermal Drug Product Quality and Performance Attributes via Enhanced Virtual Bioequivalence Simulations	SimulationsPlus, Inc	1U01FD006526
Formulation drug product quality attributes in dermal physiologically-based pharmacokinetic models for topical dermatological drug products and transdermal delivery systems	University of Queensland	1U01FD006522
PBPK and Population Modeling Seamlessly Linked to Clinical Trial Simulation in an Open-Source Software Platform	Children's Hospital of Los Angeles	1U01FD006549



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