

DDI Management in Oncology Drug Development

Ping Zhao, PhD, Integrated Development - Quantitative Sciences

FDA/ISOP Workshop on Quantitative Methods in Dosage Optimization of Oncology Products. Oct 16, 2023

BILL & MELINDA GATES foundation

Draft Panel Questions

Can we use MIDD to assess the doses needed in acute (short-term) DDI vs DDI effects at steady state?

Can we use MIDD to help design better dosing regimens for time-dependent inhibitors over the course of treatment?

Can we use MIDD to simulate/explore better dosing regimens when DDI cannot be avoided (or when we might want to leverage this phenomenon)?

Can we use MIDD to evaluate dosing in clinical scenario that are infeasible to conduct studies?

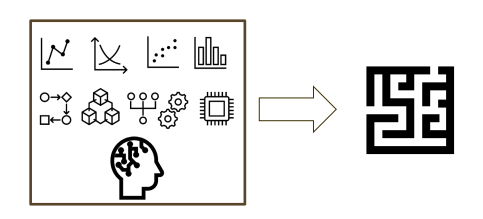
Outline

Questions	Answers	Rationale
Does technology allow us to answer DDI questions?	Yes	FDA's MIDD definition: "exposure-based, biological, and statistical models derived from preclinical and clinical data sources" Availability and industrialization of user-friendly computational tools
Are we able to address these DDI questions for all drugs with certainty now?	No	Lack of required data to derive the model, both drug or biology /physiology relevant Evolution of science and technology
Are we able to address these DDI questions for all drugs with certainty in the future?	Likely	More efficient knowledge integration Disappearance of barrier between modelers and non- modelers

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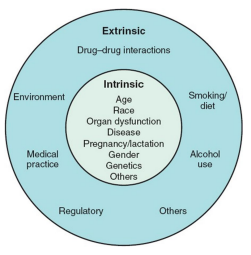
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A Complex Problem May Require Complex System to Address



Complex System

Complex Problem



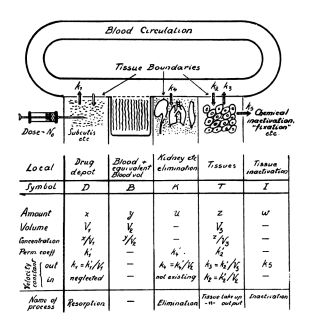
e.g., individualization of medications

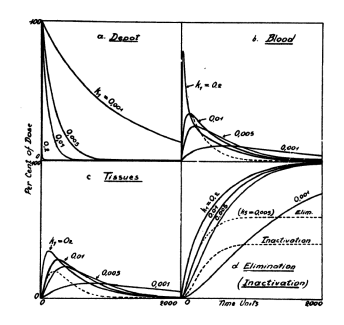
Huang and Temple, Clin Pharmacol Ther, 2008

FDA's MIDD definition: "...exposure-based, biological, and statistical models derived from preclinical and clinical data sources"

PDUFA Commitment Letters

Pharmacology Is Complex



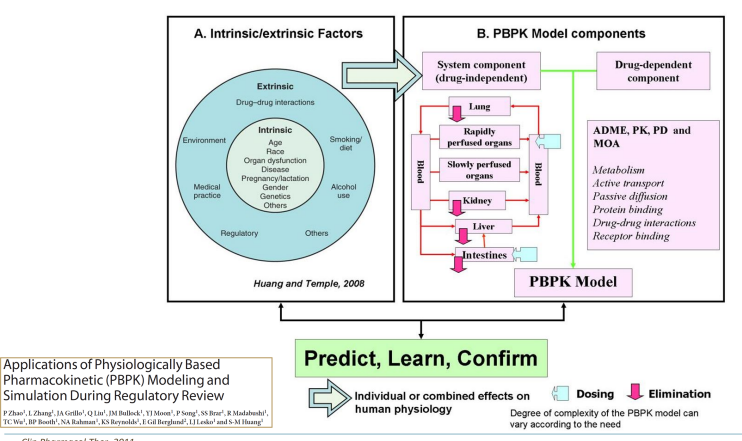


Teorell, Arch Intern Pharmacodyn, 1937

Pharmacology is the science that studies how drugs and medications work in the body, as well as their effects on the body and how they can be used to treat various medical conditions

ChatGPT description to a lay person

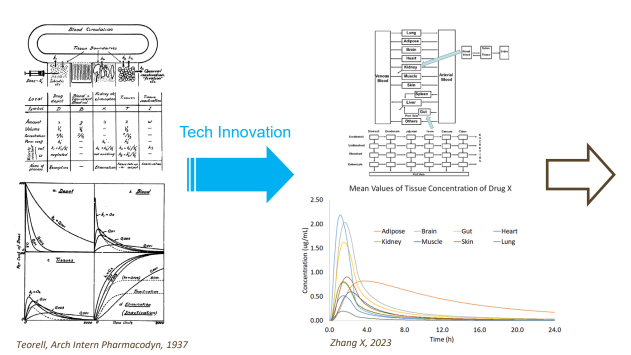
Physiologically-based Pharmacokinetic Models (PBPK)



Clin Pharmacol Ther, 2011

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|Technology Innovation Realizes A Doctor's Dream



PBPK Industrialization

- · Routine Applications
- Efficient Analysis
- Efficient Communications
- Powerful Knowledge Integration

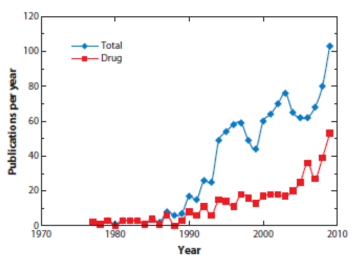
Why Software Is Eating the World

by Marc Andreessen

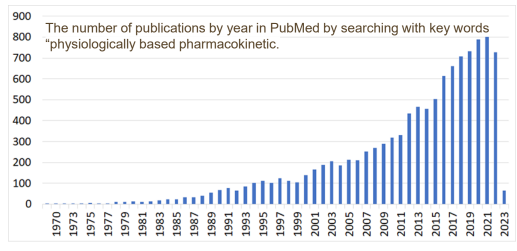
Industrialization: Common Drivers

Cost Reduction Technology Standardization Efficiency Scale
Productivity Automation
Volume Competition Unmet Needs Market Demand

PBPK Industrialization: Need, Demand, Efficiency...



Rowland M et al. Ann Rev Pharmacol Toxicol, 2010



Zhang X, 2023

Industrialization: Routine Use in R&D and Registration

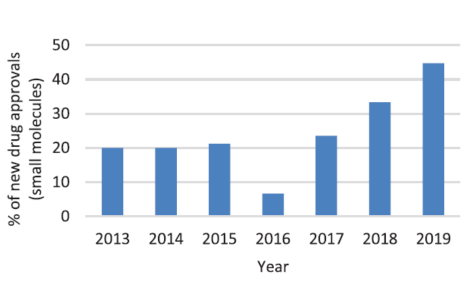


Figure 2. Percentage of new drug approvals containing physiologically based pharmacokinetics (2013-2019).

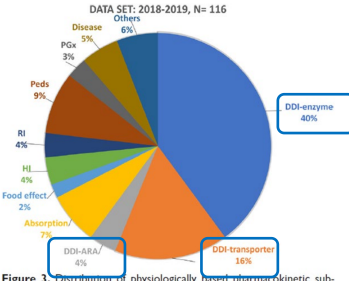


Figure 3. Distribution of physiologically based pharmacokinetic submissions by application areas (2018-2019). DDI-ARA, acid-reducing agent-mediated drug-drug interaction; DDI-enzyme, enzyme-mediated drug-drug interaction; DDI-transporter, transporter-mediated drug-drug interaction; HI, hepatic impairment; peds, pediatrics; PGx, pharmacogenomics; RI, renal impairment.

Prominent use-case is management of drug-drug interactions (DDIs)

Industrialization of PBPK in Oncology

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- Multiple Regulatory Authorities
- >100 new drugs, 325 label (simulation results in lieu of clinical studies)

Courtesy, M Jamei

Updated July. 2023

Learning from 2023 FDA Approvals

As of Sep 22, there are 36 NDAs and BLAs with FDA review published

Submissions including PBPK	15 (12 NDAs and 3 BLAs)
Oncology/Hematology	8
DDI	13 Five submissions included multiple applications (DDI as both victim and perpetrator, hepatic and renal impairment)
Label impact	Yes: 8 No, not obvious or not specified: 7
Methods	SimCYP®: 11 Gastroplus®: 1 Not specified: 3

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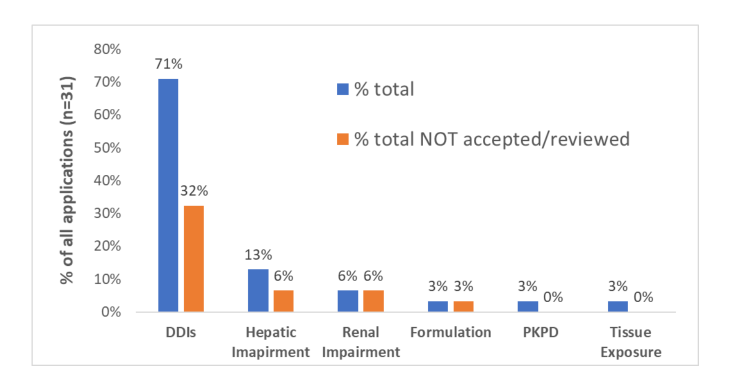
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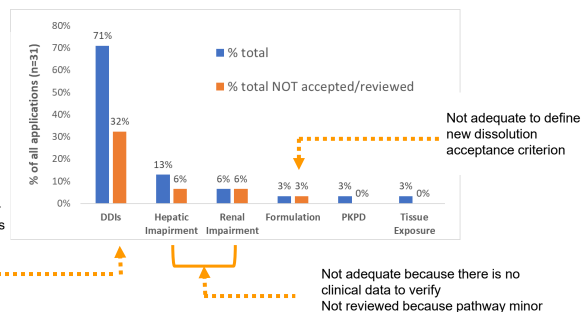
| 45% of PBPK Applications Not Accepted or Reviewed



Reasons for Not Being Accepted/Reviewed

Not adequate

- CYP contribution uncertain (NDA216718)
- Perpetrator (efavirenz) model not adequate but supportive (NDA217639)
- Rifampin's combined CYP3A induction and OATP1B1/NTCP inhibition not delineated (NDA216386)
- Model for mild CYP3A inducer; effect on CYP1A2 substrate (NDA217759)
- Underpredicted effect of rifampin; pathway not clear for CYP2C8/2C19 to predict effect of these inhibitors and effect of efavirenz (215559)
- Inadequate to predict effects of cytokine change on CYPs (BLA 761342 and 761309)



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Pragmatism - Living with Uncertainties

Jesduvroqas a CYP2C8 substrate (NDA216951):

DRUG INTERACTIONS-----

- Moderate CYP2C8 Inhibitors: Reduce starting dose. (7.1)
- CYP2C8 Inducers: Monitor hemoglobin and adjust the dose of JESDUVROQ as appropriate. (7.2)

Daprodustat AUC and C_{max} are expected to increase at least 4-fold and 3-fold, respectively, following concomitant administration of daprodustat with clopidogrel 75 mg once daily (moderate CYP2C8 inhibitor).

'However, the effects of clopidogrel on daprodustat <u>may be underestimated because</u> the clopidogrel PBPK models used by the Applicant were not reliably validated and these clopidogrel models significantly underpredicted the clinically observed interactions between clopidogrel and repaglinide (see Section 14.6 for details). This suggests that the effect of clopidogrel on daprodustat exposure <u>is expected to be no less than the model-predicted effects</u>.' NDA 216951

"All models are wrong, some are useful" G. Box

Re-usable Models – Efficiency, Generalizability, Predictability

Software library models or sponsors' models based on publications

- Sohonos (NDA 215559): 'library files sim-ketoconazole-400 mg QD, SV-rifampin-MD, SV-Efavirenz, SVErythromycin_EC, SV-Fluconazole, and SV-Fluoxetine were used for DDI simulations without any modification unless otherwise noted' (NDA 215559)
- Jesduvroq (NDA216951): 'The default PBPK models of gemfibrozil and trimethoprim in Simcyp were used for DDI prediction. The PBPK models of clopidogrel and its glucuronide were developed based on published data (Tornio et al. 2014; Shebley et al. 2017; Varma et al. 2019). Simcyp built-in models of repaglinide and the PBPK models of pioglitazone and montelukast built by the Applicant based on were used to qualify the ability of the clopidogrel PBPK model to simulate DDIs with CYP2C8 substrates as well as a CYP3A4 substrate. No modifications were made to the original models'

| More Models - Knowledge and Experience

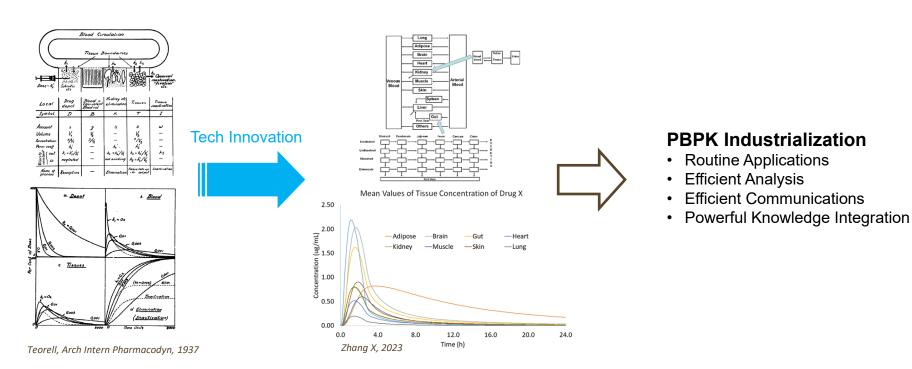
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Updated July. 2023

Shift in Mindset of Pharmacology Models



"My Model? Your model? His model? Her model? Whose model???"

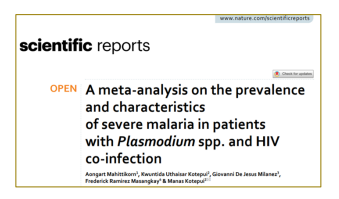
Rostami-Hodjegan, 2018

DDIs in Low Income Settings Are More Common

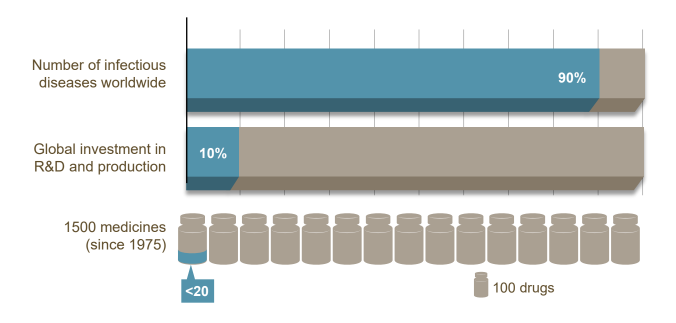
"It is estimated that one-third of the 40 million people living with HIV/AIDS worldwide are co-infected with TB. People with HIV are up to 50 times more likely to develop TB in a given year than HIV-negative people."

https://www.who.int/3by5/TBfactsheet.pdf

"The odds of SM (severe malaria) were significantly higher in co-infected patients than in Plasmodium mono-infected patients (OR <u>2.41</u>; 95% CI 1.43–4.08; I 2 = 85%; P= 0.001)...."

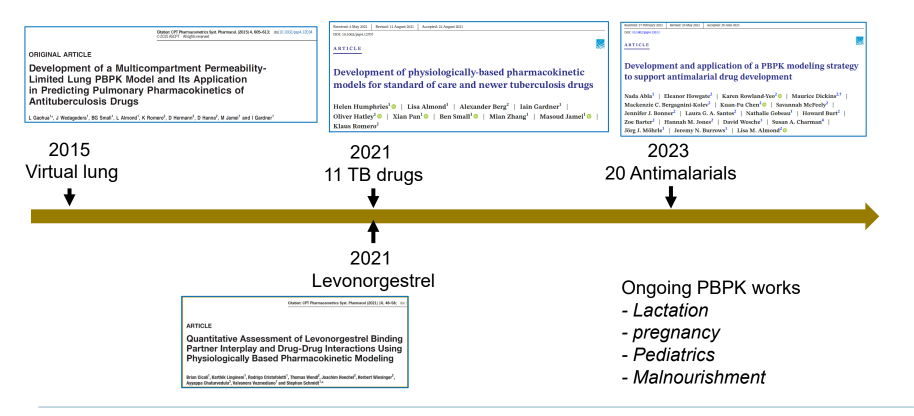


Developing Countries Under-represented in Pharma R&D

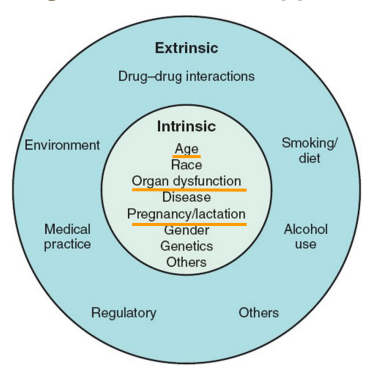


Developing countries

Collaborations to Manage Complex DDI and Comorbidity



Stagnation of PBPK Applications



Huang and Temple, Clin Pharmacol Ther, 2008

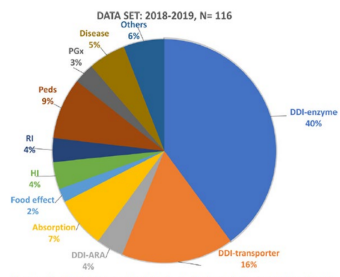


Figure 3. Distribution of physiologically based pharmacokinetic submissions by application areas (2018-2019). DDI-ARA, acid-reducing agent-mediated drug-drug interaction; DDI-enzyme, enzyme-mediated drug-drug interaction; DDI-transporter, transporter-mediated drug-drug interaction; HI, hepatic impairment; peds, pediatrics; PGx, pharmacogenomics; RI, renal impairment.

Zhang et al, J Clin Pharmacol, 2021

Bigger needs vs fewer or no applications!

Removing Barrier between Modelers and Non-modelers

Demystifying physiologically based pharmacokinetic modelling among non-modelers towards model-informed medicine use in under-served populations

Jolien Freriksen¹, Joyce van der Heijden¹, Marika de Hoop-Sommen¹, Trevor Johnson², Karen R Yeo², Essam Kerwash³, Susan Cole³, Janet Nooney², Rick Greupink¹, Ping Zhao ⁴, Saskia de Wildt^{1,5}

Workshop	Date, location and duration	Attendees	Populations discussed
Part of the Radboud Summer School "Introduction to Pharmacokinetic and Pharmacodynamic Analysis"	July 2021 Online Half day	26 delegates ^a (global) 5 tutors	Pediatric and pregnant population
Pre-congress course of the 19th European Society for Developmental Perinatal and Paediatric Pharmacology (ESDPPP) meeting	June 2022 In-person Liverpool (UK) One day	37 delegates (mostly European) 11 tutors	Pediatric and pregnant population
Part of the Radboud Summer School "Introduction to Pharmacokinetic and Pharmacodynamic Analysis"	July 2022 Online Half day	29 delegates ^a (global) 5 tutors	Pediatric and pregnant population
Satellite Session of the American Society for Clinical Pharmacology & Therapeutics (ASCPT) meeting	September 2022 Online One day	45 delegates (global) 14 tutors	Pediatric and pregnant population
Medicines and Healthcare products Regulatory Agency (MHRA) workshop	October 2022 In-person London (UK) One day + two online sessions	32 delegates ^b (mostly UK) 7 tutors	Pregnant population

Gates Open Research, 2023

¹ Radboud University, Nijmegen, Gelderland, The Netherlands

² Certara UK Limited, Sheffield, UK

³ Medicines and Healthcare Products Regulatory Agency, London, UK

⁴ Bill & Melinda Gates Foundation, Seattle, Washington, USA

⁵ Erasmus MC-Sophia Children's Hospital, Rotterdam, The Netherlands

Summary

- Technology innovation allows efficient and comprehensive evaluation of DDI scenarios
- Fit-for-purpose, practical analyses can support dosing decisions
- Knowledge-base continues to grow, enabling efficient learning across applications
- Technology innovation can further remove the barriers for many to understand and apply MIDD

| Acknowledgements

Former colleagues at US FDA

Current collaborators in global health