

# Use of Artificial Intelligence to Facilitate the Development and Regulatory Assessment of Complex Generic Drugs

**SBIA 2021: Advancing Generic Drug Development: Translating Science to Approval**  
**Day 2, Session 1: Cutting Edge Science**

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CDER | U.S. FDA

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# Disclaimer

- This presentation reflects the views of the author and should not be construed to represent FDA's views or policies.

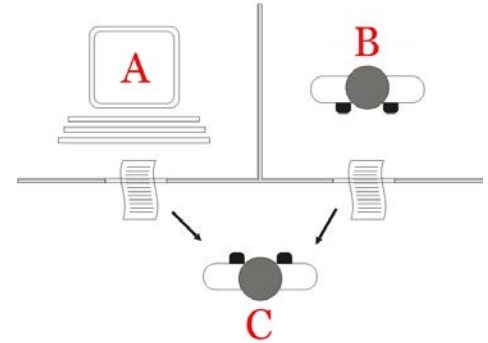
# Outline

- Artificial Intelligence (AI) is everywhere
- AI offers opportunities to facilitate generic drug development and regulatory assessment
- DQMM's efforts to answer the opportunity call
- Actions from other stakeholders
- Challenges and lessons learned
- Takeaways

# AI is everywhere



- What is AI?
  - *Turing Test* proposed by **Alan Turing** in 1950
  - “*It is the science and engineering of making intelligent machines, especially intelligent computer programs.*” by **John McCarthy** in 2004



# AI is everywhere – continued

- The thriving AI community should thank the advances in information technologies and powerful chips
  - Big data
  - Data analytics tools (e.g., machine learning and natural language processing)
  - HPC (High-performance computing) / Cloud computing
- AI is transforming many areas of everyday life
  - Smartphone (e.g., face recognition)
  - Autopilot
  - Chatbot
  - Personalized recommendation
  - ATM (automated teller machine)

# AI offers opportunities to facilitate generic drug development and regulatory assessment

- Automating labor-intensive tasks
  - Enhanced efficiency (e.g., saving time)
  - Improved consistency (e.g., reducing human error)
  - High-quality deliverables
- Exploiting advanced data analytics methods
  - Promoting business intelligence
  - Supporting regulatory assessment

# DQMM's efforts to answer the opportunity call



## Automating labor-intensive tasks

Development of:

**BEAM tool  
(bioequivalence  
assessment mate)**

data analytics tool  
to facilitate  
product-specific  
guidance (PSG)  
development

## Adopting advanced data analytics methods

Promoting business  
intelligence based on machine  
learning methodologies

Supporting regulatory  
assessment

**Prediction of abbreviated new  
drug application (ANDA)  
submission**

Equivalence assessment  
for complex particle size  
distribution based on the  
earth mover's distance

Heterogeneous treatment  
effect analysis

Multivariate analysis  
method to facilitate  
active pharmaceutical  
ingredient sameness  
assessment

Modeling the process of ANDA  
assessment



Example Case

# **Predictive Analysis of First ANDA Submission for New Chemical Entities (NCEs) Based on Machine Learning Methodology**





# Project background

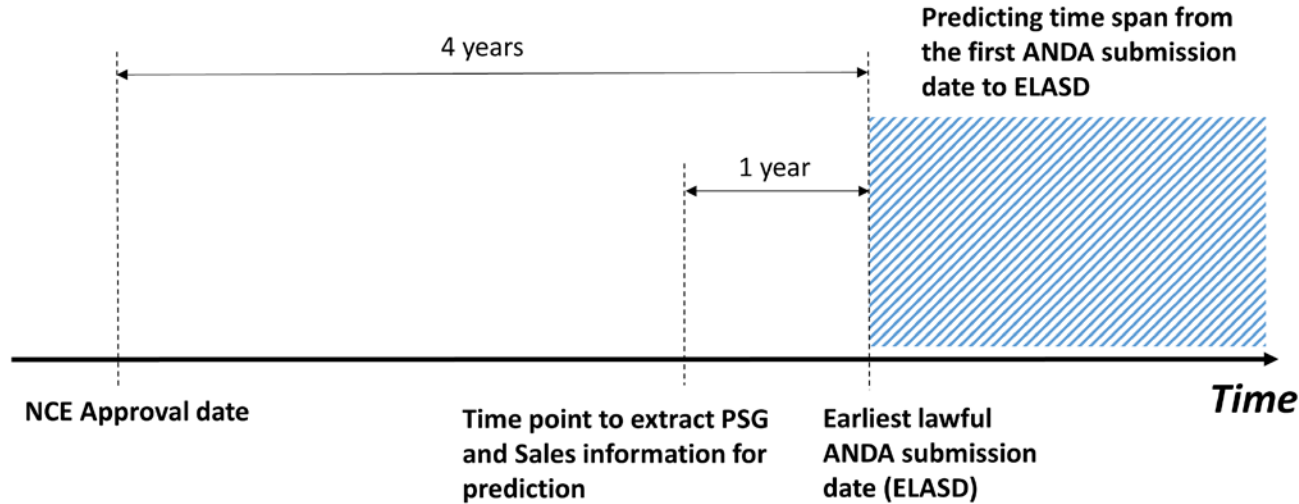
- An approved ANDA is required to ensure that generic drugs are available to facilitate drug availability and accessibility to U.S. public
- Under GDUFA II, the ANDA assessment process involves multiple offices in FDA, multiple disciplines, and tight turnaround times
- Predictive analysis of ANDA submission will critically inform resource allocation and workload management
  - Prioritizing product-specific guidance (PSG) development and research efforts
  - Optimizing resource allocation (e.g., pre-ANDA interactions)
- Study scope: Time to first submission for ANDA referencing NCE



# Collecting data for prediction

- **Drug product information**
  - Complex API
  - Complex Dosage Form
  - Complex Delivery Route
  - Complex Drug-Device Combination
  - Abuse Deterrent Formulation
  - Oral Modified Release
  - Anatomical Therapeutic Chemical (ATC)
  - Acute/Chronic Disease
- **Regulatory information**
  - NDA Approval Date
  - NCE Exclusivity Expiration Date
  - Patent Expiration Date
  - First ANDA Submission Date
  - First PSG Publication Date
  - Risk Evaluation and Mitigation Strategies (REMS)
- **Pharmacoeconomic information**
  - Drug sales from 2011 to 2017

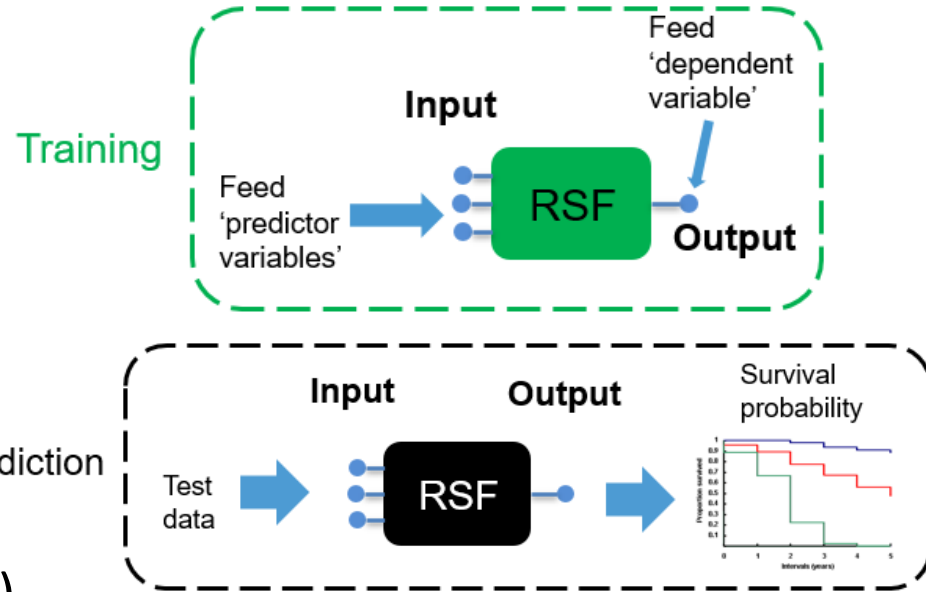
# Data model



*Time span from the first ANDA submission date to ELASD*  $\sim f(\text{Drug Information, Regulatory Information, Pharmacoeconomic Information})$

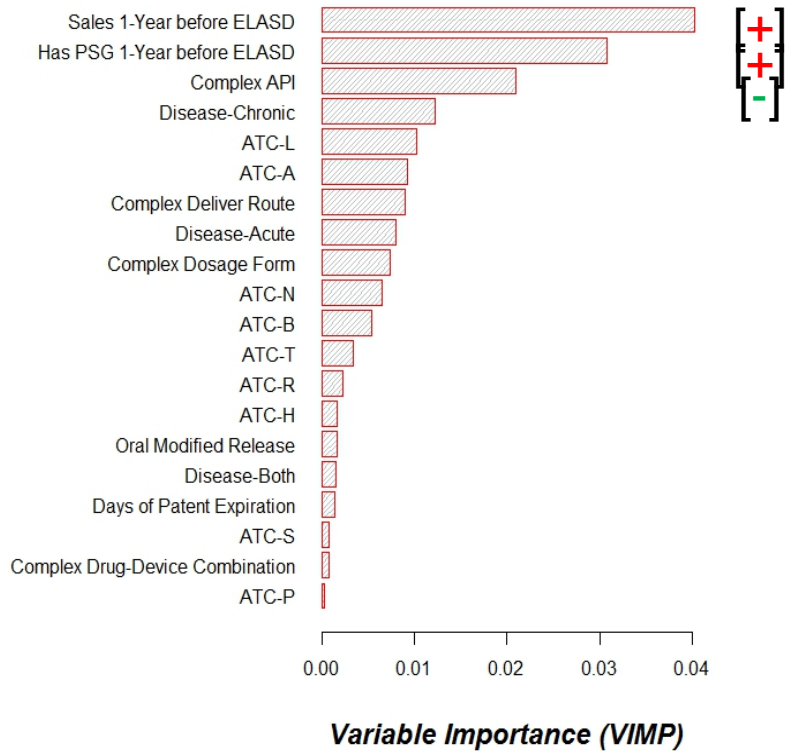
# Method

- Machine-learning based survival method
  - Random Survival Forest (RSF)
  - Data adaptive (no model assumptions)
  - Capable for large-feature problem
- Predictive Performance Evaluated by the Concordance Index (C-index)
  - C-index = 1 Perfect prediction
  - C-index = 0.5 Random guess

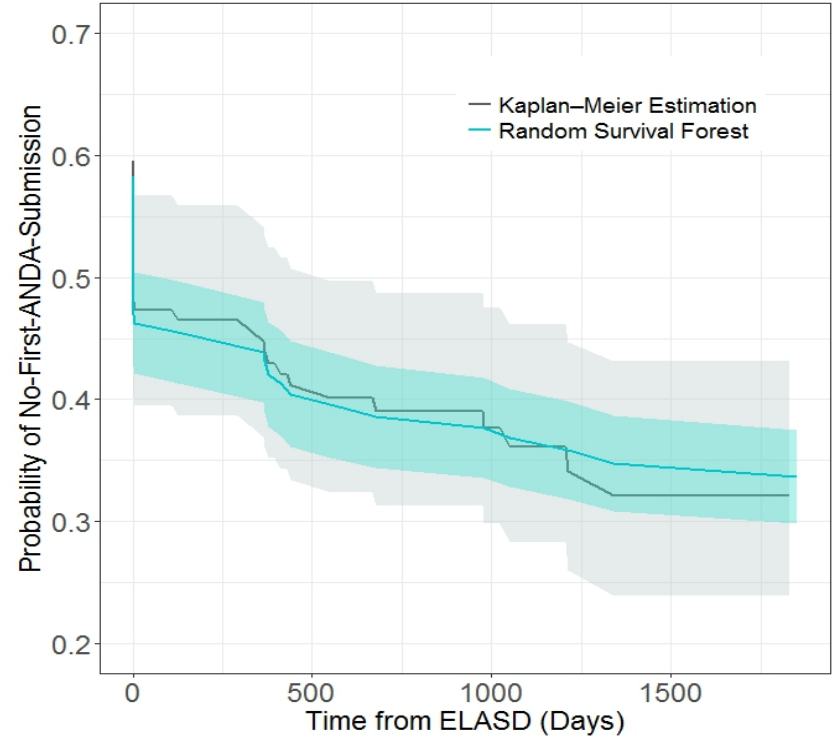


# Results

## Identification of important variables



## Prediction based on leave-one-out method (C-index = 0.767)



## **Actions from other stakeholders:**

A brief summary of the breakout session (1C) on AI in the Generic Drug Science and Research Initiatives Public Workshop (2021)

<https://sbiaevents.com/grs-2021/>



# Session Design

**Sub Session 1C:** Exploring opportunities and challenges for utilizing artificial intelligence (e.g., machine learning and natural language processing) to support generic drug development and application assessment

3:30 PM – 3:40 PM

***Artificial Intelligence in Pharmaceuticals***

**Defang Ouyang, PhD**

Assistant Professor, Univ. of Macau

3:40 PM – 3:50 PM

***Artificial Intelligence in Generic Drug Development – Experience and Opportunities***

**Jerneja Opara, PhD**

Leading Scientist, Sandoz Pharm.

3:50 PM – 4:00 PM

***Improving Generic Drugs and Streamlining Their Approval Through Artificial Intelligence***

**Charlie DiLiberti, PhD**

President, Montclair Bioequivalence Services, LLC

4:00 PM – 4:30 PM

Panel Discussion (Sub-Session 1C)

*Moderator:*

**Meng Hu, PhD**

Team Lead, DQMM, ORS, OGD, FDA

*Panelists:*

**Defang Ouyang, PhD**

Assistant Prof., Univ. of Macau

**Jerneja Opara, PhD**

Leading Scientist, Sandoz Pharm.

**Charlie DiLiberti, PhD**

President, Montclair Bioequivalence Services, LLC

**Stella Grosser, PhD**

Director, DB-VIII, Office of Biostatistics, OTS, FDA

**Liang Zhao, PhD**

Director, DQMM, ORS, OGD, FDA

**Robert Lionberger, PhD**

Director, ORS, OGD, FDA

**Donald Mager, PhD**

Prof. and Vice Chair, Department of Pharmaceutical Sciences, SUNY

**Robert Bies, PhD**

Associate Prof., Department of Pharmaceutical Sciences, SUNY

# Presentations in the session

- *AI in pharmaceuticals*
  - Challenges for formulation development and values provided by AI
  - Case study: predicting physical stability, in vitro dissolution and in vivo performance of solid dispersion
- *AI in generic drug development - experience and opportunities*
  - AI models were used to predict BE outcomes (e.g., Cmax T/R ratio)
  - Discussion on other opportunities of AI models in generic drug development
- *Improving generic drugs and streamlining their approval through AI*
  - Needs to develop regulatory framework for AI
  - Potential applications of AI to the generic drug development





# Highlights from the panel discussion

- Priority order of potential applications, the low-hanging fruits (e.g., handling outliers)
- Feasibilities from the technical perspective
  - Combination of AI models and conventional tools (e.g., Physiologically based pharmacokinetic (PBPK) models)
  - Integrating domain knowledge
  - Leveraging AI to utilize data from different resources
- Transparency / interpretability of AI models
- Business model of collaborations (e.g., between industries and the FDA)

# Challenges and Lessons Learned

- Availability of reliable data
- Incorporation of domain knowledge
- Enabling complex projection between complex datasets
- Transparency/interpretability - turning the black box into a gray (or even transparent) box
- Extrapolation ability of AI models
- Lack of guidelines for regulatory purpose
- Teamwork and collaboration



# Takeaways

- AI technologies:
  - bring opportunities to advance development and regulatory assessment of generic drugs.
  - have been applied to facilitate BE assessment, PSG development, business intelligence and regulatory assessment, etc.
  - will play more important role in providing high-quality generic drugs for U.S. public as more challenges get addressed.



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