

CDER Small Business and Industry Assistance

Webinars 2025



Model Master Files: Advancing Modeling and Simulation in Generic Drug Development and Regulatory Submissions

March 13, 2025, 1:00 - 3:00 pm EDT

1:00 - 1:05: Welcome and Overview

Kori Adair, PharmD

Pharmacist | Small Business and Industry Assistance (SBIA) | Division of Drug Information (DDI) Office of Communications (OCOMM) | Center for Drug Evaluation and Research (CDER)

1:05 - 1:15: Introduction and Overview of the Model Master File

Lanyan (Lucy) Fang, PhD

Acting Division Director | Division of Quantitative Methods and Modeling (DQMM) | Office of Research and Standards (ORS) | Office of Generic Drugs (OGD) | CDER

1:15 - 1:35: Model Master Files: How to Develop and Submit One?

Eleftheria Tsakalozou, PhD

Lead Pharmacologist DQMM | ORS | OGD | CDER

1:35 - 1:50: Cross-Comparison to Other Drug Master Files and Lessons Learned

Erin Skoda. PhD

Supervisory Chemist | Division of Product Quality Assessment XVIII (DPQA XVIII)
Office of Product Quality Assessment III (OPQA III) | Office of Pharmaceutical Quality (OPQ)

1:50 - 2:25: Discussion Panel

Moderator: Andrew Babiskin, PhD, Lead Pharmacokineticist and Acting Deputy Division Director DQMM | ORS | OGD | CDER

Panelists:

- Robert Lionberger, PhD | Director, ORS | OGD | CDER
- Lanyan (Lucy) Fang, PhD, Acting Division Director, DQMM | ORS | OGD | CDER
- Partha Roy, PhD | Director, Office of Bioequivalence (OB) | OGD | CDER
- Bhagwant Rege, PhD | Division Director, Division of Product Quality Assessment VI (DPQA VI)
 Office of Product Quality Assessment I (OPQA I) | OPQ | CDER
- Stella Grosser, PhD | Division Director, Division of Biometrics VIII (DBVIII)
 Office of Biostatistics (OB) | Office of Translational Sciences (OTS) | CDER
- Rajanikanth Madabushi, PhD | Associate Director for Guidance and Scientific Policy
 Office of Clinical Pharmacology (OCP) | OTS | CDER
- Martha Nguyen, JD | Division Director | Division of Policy Development (DPD)
 Office of Generic Drug Policy (OGDP) | OGD | CDER

2:25 - 2:55: Question and Answer Session

Moderator: Kori Adair, PharmD

Panelists:

- Eleftheria Tsakalozou, PhD | Lead Pharmacologist, DQMM | ORS | OGD | CDER
- Ross Walenga, PhD | Senior Chemical Engineer, DQMM | ORS | OGD | CDER
- Fang Wu, PhD | Senior Pharmacologist, DQMM | ORS | OGD | CDER
- Meng Hu, PhD | Lead Engineer, DQMM | ORS | OGD | CDER
- Yuqing Gong, PhD | Acting Lead Pharmacologist, DQMM | ORS | OGD | CDER
- Ethan Stier, PhD | Associate Director for Lifecycle Management, OCP | OTS | CDER
- Hao Zhu, PhD | Division Director, Division of Pharmacometrics (DPM) | OCP | OTS | CDER
- Rebecca Moody, PhD | Pharmaceutical Scientist, Office of Product Quality Assessment II (OPQA II) | OPQ

2:55 - 3:00: Closing Remarks

Robert Lionberger, PhD | Director, ORS | OGD | CDER