

In silico Drug Discovery Workshop

hosted by the NCATS Assay Guidance Manual Program

Wednesday, October 23, 2024 - Thursday, October 24, 2024 ~ Virtual (All times are in ET)

AGENDA: Day 1

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11:00 AM	Introductory Comments Joni Rutter, National Center for Advancing Translational Sciences (NCATS), National Institutes of Health (NIH) Kimberly Sciarretta, Biomedical Advanced Research and Development Authority (BARDA)	
11:20 AM	Overview of Workshop Objectives Sarine Markossian and Alexey Zakharov, NCATS, NIH	
11:30 AM	Keynote: Computer-Aided Drug Discovery Then and Now Martha S. Head (Marti), Amgen Inc.	
12:15 PM	Lunch	
12:45 PM	Session 1: Data Chair: Sarine Ma 12:45 PM 1:15 PM	asourcing for in silico drug discovery arkossian, NCATS 15 Years of ChEMBL: Challenges & Opportunities Barbara Zdrazil, ChEMBL Protein Data Bank: From Two Epidemics to the Global Pandemic to mRNA Vaccines and Paxlovid Stephen K. Burley, Rutgers
1:45 PM	Break	
2:00 PM		Tropsha, University of North Carolina Chapel Hill Cartography-Guided Exploration of (Ultra)Large Chemical Spaces Alexandre Varnek, The University of Strasbourg Virtual Screening (TBD) Gisbert Schneider, ETH Zürich Synthesis-Aware Library Design and Optimal Experimental Design Connor W. Coley, Massachusetts Institute of Technology (MIT) Physics-Based Modeling (TBD) John D. Chodera, Memorial Sloan-Kettering Cancer Center
4:00 PM	Break	
4:15 PM	Session 3: Appl Chair: Morgan S 4:15 PM 4:45 PM	lications of in silico methodologies in drug discovery herer, BARDA Applying Artificial Intelligence in a Small Drug Discovery Company Sean Ekins, Collaborations Pharmaceuticals, Inc. TBD Victoria Feher, Novartis Institutes for BioMedical Research (NIBR)



Summary of Day 1 and Adjourn

5:15 PM

AGENDA: Day 2

11:00 AM Session 3: Applications of in silico methodologies in drug discovery (continued)

Chair: Morgan Sherer, BARDA

11:00 AM Industrializing Early Drug Discovery at Scale

Marissa Saunders, Recursion Pharmaceuticals

11:30 AM **TBD**

Alex Zhavoronkov, Insilico Medicine

12:00 PM Lunch

12:30 PM Session 4: Emerging trends in in silico drug discovery

Chair: Alexey Zakharov, NCATS

12:30 PM Understanding the Chameleonicity and Permeability of PROTACs Using Molecular

Dynamics Simulations, Markov Models, and Deep Learning

Bryn Taylor, Johnson & Johnson Innovative Medicine

1:00 PM Data Generation in Support of Computational Drug Discovery

Cheryl H Arrowsmith, University of Toronto

1:30 PM Human-Al-Robot Collaboration to Accelerate Materials and Molecular Discovery

Milad Abolhasani, North Carolina State University

2:00 PM Protein Therapeutics Design (TBD)

David Baker, University of Washington

2:30 PM Break

3:00 PM Panel Discussion: Bridging the gaps in translation

Chair and moderator: Rommie Amaro, University of California San Diego (USCD)

Panelists: Pat Walters, Relay Therapeutics; Russ B Altman, Stanford University; Christopher Austin, GSK; Martha S.

Head (Marti), Amgen Inc.

3:00 PM Introduction to the Session and Challenges Ahead

Rommie Amaro, UCSD

3:20 PM Dangers of Poor Data (TBD)

Pat Walters, Relay Therapeutics

3:40 PM Paying Attention to Attention in Al Generative Models

Russ B Altman, Stanford University

4:00 PM Team Environment for Better Translation (TBD)

Christopher Austin, GSK

4:20 PM Discussion and Q&A

5:00 PM Closing Session: Summary of Discussions and Perspectives on the Challenges Ahead

Alexey Zakharov, NCATS; Alexander Tropsha, UNC Chapel Hill; Rommie Amaro, USCD

5:15 PM Closing Statement and Adjourn