# FDA-CRCG Workshop on Visionary Standards: Advancing Science and Regulation in Generic Ophthalmic Products

# Public Workshop November 19-20, 2025 Agenda

8:30 AM – 8:35 AM Welcome and CRCG Opening Remarks

Anna Schwendeman, PhD Co-Director, CRCG

8:35 AM – 8:50 AM FDA Opening Remarks

11:40 AM - 12:40 PM

Lunch Break

Markham Luke, MD, PhD Director, DTP I, ORS, OGD, FDA

## Session 1: Ophthalmic Product Standards and General Bioequivalence Considerations

This session will explore key regulatory standards, scientific principles, and methodologies that underpin the demonstration of bioequivalence in ophthalmic drug products. Presentations will cover FDA regulatory research shaping the ophthalmic landscape, current and emerging approaches for establishing bioequivalence, and case studies highlighting formulation and manufacturing challenges. Additional topics will address the influence of preservatives on ocular absorption, scientific considerations for non-Q1/Q2 formulations, and strategies for developing robust in vitro release testing methods. Together, these talks will provide a comprehensive perspective on advancing safe, effective, and high-quality ophthalmic generics that meet rigorous regulatory and patient care standards.

8:50 AM – 8:55 AM	Speaker Introductions Bin Qin, PhD	Senior Staff Fellow, DTP I, ORS, OGD, FDA
8:55 AM – 9:15 AM	A Look at the Impact of FDA Regulate Andre O'Reilly Beringhs, PhD	ory Research on the Landscape of Ophthalmology Drugs Pharmacologist, DTP I, ORS, OGD, FDA
9:15 AM – 9:35 AM	Establishing Bioequivalence for Opht Bruce Lerman, PhD	halmic Products: Current Approaches and Future Perspectives Lead Pharmacologist, DB I, OB, OGD, FDA
9:35 AM – 9:55 AM	Challenges and Opportunities of Q1/ Nemanja Aničić, PhD	<b>Q2 Pathway for Biowaiver of Ophthalmic Products</b> Senior Scientist, Pharmaceutical Development, Sandoz Development Center
9:55 AM – 10:15 AM	Preservative Effects on Ocular Absorp Arto Urtti, PhD	otion from Topically Instilled Eye Drops Professor, Biopharmaceutics, Univ. of Eastern Finland
10:15 AM – 10:30 AM	Coffee Break	
10:30 AM – 10:50 AM	Scientific Hurdles to Demonstrate Bioequivalence for Non-Q1/Q2 Ophthalmic Solutions in Generic Drug Development	
	Yoriko Harigaya, PharmD	Senior Staff Fellow, DB II, OB, OGD, FDA
10:50 AM – 11:10 AM	Seeing the Options: Challenges and S Xiaoming Xu, PhD	itrategies in Developing IVRT Methods for Ophthalmic Products Division Director, DPQRV, OPQR, OPQ, FDA
11:10 AM – 11:40 AM Moderator: Panelists:	Q&A Session with Panel Bin Qin, PhD Nemanja Aničić, PhD Andre O'Reilly Beringhs, PhD Yoriko Harigaya, PharmD Bruce Lerman, PhD Arto Urtti, PhD Xiaoming Xu, PhD	Senior Staff Fellow, DTP I, ORS, OGD, FDA Senior Scientist, Pharmaceutical Development, Sandoz Development Center Pharmacologist, DTP I, ORS, OGD, FDA Senior Staff Fellow, DB II, OB, OGD, FDA Lead Pharmacologist, DB I, OB, OGD, FDA Professor, Biopharmaceutics, Univ. of Eastern Finland Division Director, DPQRV, OPQR, OPQ, FDA

#### Session 2: Regulatory Science Applied to Ophthalmic Dispersion Products

This session will focus on regulatory science strategies for evaluating complex ophthalmic dispersions, including suspensions and ointments. Speakers will discuss pharmacokinetic (PK) and pharmacodynamic (PD) assessments, formulation challenges, and case studies illustrating both in vitro and in vivo evaluation methodologies. Presentations will highlight bioequivalence perspectives, modeling approaches, including Physiologically-Based Pharmacodynamic (PBPK/PD) and Computational Fluid Dynamics (CFD) frameworks, and the extrapolation of pharmacodynamic effects across species. The session aims to provide practical insights and scientific tools to support the development of high-quality generic ophthalmic dispersion products.

12:40 PM – 12:45 PM	Speaker Introductions Huzeyfe Yilmaz, PhD Yan Wang, PhD	Senior General Physical Scientist, DPQR II, OPQR, OPQ, FDA Deputy Director, DTP I, ORS, OGD, FDA
12:45 PM – 1:05 PM	Topical and Intracameral Pharmacol Vatsala Naageshwaran PhD	cinetics of Complex Ophthalmic Suspensions CEO, Franklin Biolabs
1:05 PM – 1:25 PM	Development and Optimization of Po Romit Jani, MS	article Size for Ophthalmic Dispersion Formulations Senior Director, Formulation R&D, Solaris Pharma
1:25 PM – 1:45 PM	In Vitro & In Vivo Assessment of Tobramycin & Dexamethasone Ophthalmic Ointments for Generic Product Equivalence	
	Xiuling Lu, PhD	Professor, School of Pharmacy, University of Connecticut
1:45 PM – 2:05 PM	Bioequivalence Perspective for Comp Hee Sun Chung, PhD	plex Ophthalmic Dispersion Products—Common Challenges and Deficiencies  Lead Pharmacologist, DB I, OB, OGD, FDA
2:05 PM – 2:25 PM	Extrapolation of PD Effects Across Sp Jessica Spires, PhD	Pecies Using Ocular PBPK Modeling Principal Scientist, Simulations Plus, Inc
2:25 PM – 2:45 PM	Ocular PBPK/PD Modeling to Explore Mingliang Tan, PhD	e In Vitro Bioequivalence Approaches for Brinzolamide Ophthalmic Suspensions Senior Pharmacokineticist, DQMM, ORS, OGD, FDA
2:45 PM – 3:05 PM	In Silico Modeling of Ophthalmic Sus Carrie German, PhD	pensions: From In Vitro to In Vivo Director, Experimental and Computational Biology, CFD Research Corp.
3:05 PM — 3:35 PM Moderators: Panelists:	Q&A Session with Panel Huzeyfe Yilmaz, PhD Yan Wang, PhD Hee Sun Chung, PhD Carrie German, PhD Romit Jani, MS Xiuling Lu, PhD Vatsala Naageshwaran, PhD Jessica Spires, PhD Mingliang Tan, PhD	Senior General Physical Scientist, DPQR II, OPQR, OPQ, FDA Deputy Director, DTP I, ORS, OGD, FDA Lead Pharmacologist, DB I, OB, OGD, FDA Director, Experimental and Computational Biology, CFD Research Corp. Senior Director, Formulation R&D, Solaris Pharma Professor, School of Pharmacy, University of Connecticut CEO, Franklin Biolabs Principal Scientist, Simulations Plus, Inc Senior Pharmacokineticist, DQMM, ORS, OGD, FDA
3:35 PM – 3:40 PM	Closing Remarks for Virtual Audience Yan Wang, PhD	Deputy Director, DTP I, ORS, OGD, FDA
3:40 PM – 4:00 PM	Coffee Break	

### Session 3: Small Working Group Sessions (In-Person Only)

These interactive, in-person discussions will provide a forum for deeper engagement with FDA speakers and other attendees on session topics from Day 1. Attendees will have the opportunity to exchange perspectives, share experiences, and explore potential regulatory and research pathways for advancing generic ophthalmic products. The small group format will encourage collaboration and practical problem-solving among FDA scientists, industry representatives, and academic experts.

4:00 PM – 5:30 PM Session 3: In Person-Only Discussion

## Day 2 November 20

9:00 AM – 9:10 AM **Day 1 Summary** 

Yan Wang, PhD Deputy Director, DTP I, ORS, OGD, FDA

## Session 4: Regulatory Science Applied to Ophthalmic Insertable and Implantable Products

This session will examine the regulatory science guiding the development, evaluation, and approval of ophthalmic inserts and implants. Topics will include performance characterization of intracanalicular and intravitreal products, advanced analytical methods for polymer-based delivery systems, and reverse-engineering studies that inform generic development. Presentations will also cover novel ocular delivery platforms, strategies for overcoming development and regulatory challenges, and key quality considerations. Attendees will gain insights into how rigorous evaluation and innovative research ensure safe, effective, and long-acting ophthalmic therapies.

9:10 AM – 9:15 AM	Speaker Introductions William Smith, PhD	Research Scientist, DPQR V, OPQR, OPQ, FDA
9:15 AM – 9:35 AM	Product Evaluation and Performance Michael VandenBerg, PhD	Characterization of Dexamethasone Intracanalicular Inserts Chemical Engineer, DPQR VI, OPQR, OPQ, FDA
9:35 AM – 9:55 AM	Analytical Methods for Mini-Size Col Kinam Park, PhD	mplex Long-Acting Injectable Formulations Professor, Purdue University; President, Akina Inc.
9:55 AM – 10:15 AM	Reverse Engineering of Dexamethas Coleman Johnson, PhD	one Intravitreal Implants Postdoctoral Research Fellow, OPQR, OPQ, FDA
10:15 AM – 10:30 AM	Coffee Break	
10:30 AM – 10:50 AM	Image-Based In Vitro In Silico Release Predication Software & Reusable Database for Insertable & Implantable Products	
	Andrew Clark, PhD	Senior Director of Research and Strategy, digiM Solution LLC
10:50 AM – 11:10 AM	Drug Product Quality Considerations for Dexamethasone Intravitreal Implant	
	Megha Barot, PhD	Pharmaceutical Scientist, DPQA IV, OPQA I, OPQ, FDA
11:10 AM – 11:40 AM	Q&A Session with Panel	
Moderator:	William Smith, PhD	Research Scientist, DPQR V, OPQR, OPQ, FDA
Panelists:	Megha Barot, PhD	Pharmaceutical Scientist, DPQA IV, OPQA I, OPQ, FDA
	Andrew Clark, PhD	Senior Director of Research and Strategy, digiM Solution LLC
	Young Jhon, PhD	Senior Chemist, DPQA IV, OPQA I, OPQ, FDA
	Coleman Johnson, PhD	Postdoctoral Research Fellow, OPQR, OPQ, FDA
	Kinam Park, PhD	Professor, Purdue University; President, Akina Inc.
	Michael VandenBerg, PhD	Chemical Engineer, DPQR VI, OPQR, OPQ, FDA

## Session 5: Regulatory Science Applied to Drug-Device Combination Products

Lunch Break

11:40 AM - 1:00 PM

This session will address the complexities of ophthalmic drug-device combination products (DDCPs), with a focus on regulatory pathways, performance expectations, and common development challenges. Topics will include comparative threshold analysis for devices, industry perspectives on clinical conduct and quality, lessons learned from ANDA submissions, and the unique challenges faced in developing complex generic ophthalmic drug-device combination products. Presentations from both FDA and industry will highlight strategies to facilitate innovation while ensuring patient safety, therapeutic efficacy, and consistent product quality.

1:00 PM – 1:05 PM	Speaker Introductions		
	Nathan Reed, PhD	Chemist, DPQR II, OPQR, OPQ, FDA	
	Markham Luke, MD, PhD	Director, DTP I, ORS, OGD, FDA	
1:05 PM – 1:25 PM	Challenges with DDCP: Clinical Conduct and Quality from Industry Perspective		
	Tausif Ahmed, PhD	Senior VP & Head of Clinical and Biopharmaceutics, Mankind Pharma	
1:25 PM – 1:45 PM	Ophthalmic Drug-Device Combination Products – Comparative Analyses		
	Shinae Kim, PhD	Pharmacokineticist, DTP I, ORS, OGD, FDA	
1:45 PM – 2:05 PM	Drug-Device Combination Product ANDA Submissions – Lessons Learned		
	CDR Andrew Fine, PharmD, BCPS	Senior Advisor, DCR, OSCE, OGD, FDA	

2:05 PM – 2:25 PM Challenges in the Development of Complex Generic Ophthalmic Products

Ajay Khopade, PhD Vice President R&D, Sun Pharmaceuticals Industries Limited

2:25 PM – 2:55 PM **Q&A Session with Panel** 

Moderators: Nathan Reed, PhD Chemist, DPQR II, OPQR, OPQ, FDA

Markham Luke, MD, PhD Director, DTP I, ORS, OGD, FDA

Panelists: Tausif Ahmed, PhD Senior VP & Head of Clinical and Biopharmaceutics, Mankind Pharma

CDR Andrew Fine, PharmD, BCPS Senior Advisor, DCR, OSCE, OGD, FDA

Ajay Khopade, PhD Vice President R&D, Sun Pharmaceuticals Industries Limited

Shinae Kim, PhD Pharmacokineticist, DTP I, ORS, OGD, FDA

2:55 PM – 3:05 PM Virtual Workshop Closeout

Yan Wang, PhD Deputy Director, DTP I, ORS, OGD, FDA

3:05 PM – 3:20 PM *Coffee Break* 

## Session 6: Small Working Group Sessions (In-Person Only)

This closing in-person session will provide participants with a collaborative forum to discuss and expand on key themes from Day 2. Attendees will engage in focused discussions on regulatory science, technical challenges, and innovation opportunities for ophthalmic drug-device combination products, inserts, and implants. The interactive format will foster direct dialogue among FDA scientists, industry experts, and academic researchers, encouraging the exchange of practical strategies, regulatory insights, and research priorities to advance the development of safe, effective, and high-quality generic ophthalmic products.

3:20 PM – 4:45 PM Session 6: In Person-Only Discussion

4:45 PM – 5:00 PM In Person-Only Workshop Closeout

Andre O'Reilly Beringhs, PhD Pharmacologist, DTP I, ORS, OGD, FDA

## **Appendix of Abbreviations**

ANDA Abbreviated New Drug Application

BCPS Board Certified Pharmacotherapy Specialist

CDR Commander

CEO Chief Executive Officer

CRCG Center for Research on Complex Generics

DB Division of Bioequivalence
DCR Division of Clinical Review

DDCP Drug Device Combination Products

DPQR Division of Pharmaceutical Quality Research
DTP Division of Therapeutic Performance

DQMM Division of Quantitative Methods and Modeling

FDA Food and Drug Administration

Inc Incorporated

IVRT In Vitro Release Testing LLC Limited Liability Company

Ltd Limited

MD Doctor of Medicine
OB Office of Biostatistics
OGD Office of Generic Drugs

OPQ Office of Pharmaceutical Quality

OPQR Office of Pharmaceutical Quality Research

ORS Office of Regulatory Science

PBPK Physiologically Based Pharmacokinetic

PD Pharmacodynamic
PharmD Doctor of Pharmacy
PhD Doctor of Philosophy
PK Pharmacokinetic
Q1 Qualitative 1
Q2 Quantitative 2

R&D Research and Development

VP Vice President