FDA-CRCG Workshop on Navigating the Transition to Low Global Warming Potential Propellants

Public Workshop December 4-5, 2024 Agenda

Day 1	December 4	
8:30 AM – 8:40 AM	Welcome and Opening Remarks	
6.30 AIVI - 6.40 AIVI	James Polli, PhD	Co-Director, CRCG
	Anna Schwendeman, PhD	Co-Director, CRCG
8:40 AM – 8:50 AM	FDA Opening Remarks	
0.40 AIVI 0.30 AIVI	Markham Luke, MD, PhD	Director, DTP I, ORS, OGD, FDA
8:50 AM – 8:55 AM	Workshop Day 1 Overview	
	Markham Luke, MD, PhD	Director, DTP I, ORS, OGD, FDA

Session 1: LGWP Propellant Transition for New Drug Product Metered Dose Inhalers (MDIs)

In this session, regulators will discuss the current propellant transition including lessons learned from previous transitions and scientific considerations for the LGWP propellants. FDA presenters will cover the scientific and regulatory considerations for new drug LGWP propellant MDI development programs and expectations for the pharmacology/toxicology, quality, pharmacokinetics, pharmacodynamics, efficacy, and safety data to support a new drug application.

8:55 AM – 9:00 AM	Speaker Introductions Sally Seymour, MD	Director, DPACC, OII, OND, FDA
9:00 AM – 9:20 AM	The Global Phasedown of HFC: A U.S Luke Hall-Jordan, MPH	. Regulatory Perspective Supervisory Environmental Policy Analyst, Phasedown Implement Branch, EPA
9:20 AM – 9:30 AM	Lessons Learned from CFC Transition Sally Seymour, MD	Director, DPACC, OII, OND, FDA
9:30 AM – 9:55 AM	Propellant Properties and the Impact Stephen Stein, MS	t on Product Development Scientific Director, Inhalation Product Development, Kindeva Drug Delivery
9:55 AM – 10:10 AM	Coffee Break	
10:10 AM – 10:20 AM	Introduction to OND Framework for LGWP Transition Stacy Chin, MD Clinical Team Leader, DPACC, OII, OND, FDA	
10:20 AM – 10:30 AM	Nonclinical Considerations for LGWP Luqi Pei, PhD	Transition Master Pharmacologist, DPT II, OII, OND, FDA
10:30 AM – 10:40 AM	Quality Considerations for LGWP Pro Craig Bertha, PhD	pellant Transitions CMC Reviewer, DPQA VII, OPQA II, OPQ, FDA
10:40 AM – 10:55 AM	Clinical Pharmacology Consideration Sneha Dhapare, PhD	s for LGWP Transition Senior Clinical Pharmacologist, DIIP, OTS, FDA
10:55 AM – 11:10 AM	Clinical Considerations for LGWP Translated Alishah Ali, MD	nsition Senior Physician, DPACC, OII, OND, FDA
11:10 AM – 12:00 PM Moderator: Panelists:	Q&A Session with Panel Sally Seymour, MD Aishah Ali, MD Craig Bertha, PhD Stacy Chin, MD Sneha Dhapare, PhD	Director, DPACC, OII, OND, FDA Senior Physician, DPACC, OII, OND, FDA CMC Reviewer, DPQA VII, OPQA II, OPQ, FDA Clinical Team Leader, DPACC, OII, OND, FDA Senior Clinical Pharmacologist, DIIP, OTS, FDA

Luke Hall-Jordan, MPH Supervisory Environmental Policy Analyst, Phasedown Implement Branch, EPA

Luqi Pei, PhD Master Pharmacologist, DPT II, OII, OND, FDA

Stephen Stein, MS Scientific Director, Inhalation Product Development, Kindeva Drug Delivery

Hailing Zhang, PhD Division Director, DPQA XII, OPQA II, OPQ, FDA

12:00 PM - 12:50 PM **Lunch Break**

Session 2: Current Industry Experience with New Drug LGWP MDI Development

This session will include industry presentations on their experiences with new drug LGWP propellant MDI development, including development strategy and challenges encountered thus far.

12:50 PM – 12:55 PM **Speaker Introductions**

Christy Gilbert, BS, RAC Associate Director, CMC Regulatory Affairs, AstraZeneca

12:55 PM - 1:10 PM An Introduction to the IPAC-RS Scenarios - Transition to LGWP Propellants in MDIs: Proposed Pathways to U.S. FDA

Approval

Ann Purrington, BS Regulatory Affairs Director, Kindeva Drug Delivery

1:10 PM - 1:30 PM Statistical Considerations for the In Vitro Comparisons of Critical Product Performance Attributes for an Approved

MDI Product Transitioning to LGWP Propellant

Richard (Rik) Lostritto, PhD Consultant, Lostritto Consulting, LLC

Helen Strickland, MS Sr. Statistical Consultant, Manufacturing Science & Technology, GSK

1:30 PM - 1:50 PM Case Study: Development Considerations for Transitioning MDI Products to LGWP Propellant

Laura Clow, MChem Medicine Development Leader, GSK

Poonam Gulati, PhD, MBA, PGCert Senior Director & Team Lead, Global Regulatory Affairs, GSK

1:50 PM - 2:10 PM Engineered Excipient Particles Facilitate Transition to LGWP Propellants in Combination MDI Products

David Lechuga-Ballesteros, PhD Head of Innovation, IPD, AstraZeneca

2:10 PM - 2:30 PM Innovative Drug Development Approach to Address the Transition to LGWP Propellant Using HFA 152a, for

a Triple Combination MDI Product Targeting Small Airways

Angelo Benedetto Matturo, MS R&D Global Technical Leader, CHIESI Farmaceutici

2:30 PM – 3:00 PM **Q&A Session with Panel**

Moderators: Christy Gilbert, BS, RAC Associate Director, CMC Regulatory Affairs, AstraZeneca

Sue Holmes, MS CMC Regulatory Consultant, Sue Holmes CMC Consulting LLC

Panelists: Laura Clow, MChem Medicine Development Leader, GSK

Poonam Gulati, PhD, MBA, PGCert Senior Director & Team Lead, Global Regulatory Affairs, GSK

David Lechuga-Ballesteros, PhDHead of Innovation, IPD, AstraZenecaRichard (Rik) Lostritto, PhDConsultant, Lostritto Consulting, LLCMarkham Luke, MD, PhDDirector, DTP I, ORS, OGD, FDA

Angelo Benedetto Matturo, MS R&D Global Technical Leader, CHIESI Farmaceutici Ann Purrington, BS Regulatory Affairs Director, Kindeva Drug Delivery

3:00 PM – 3:05 PM Closing Remarks for the Virtual Session (End of Virtual Session for Day 1)

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

3:05 PM – 3:15 PM *Coffee Break*

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

3:15 PM -4:45 PM

Lead: Elizabeth Bielski, PhD Senior Pharmacologist, DTP I, ORS, OGD, FDA

Moderators: Richard (Rik) Lostritto, PhD Consultant, Lostritto Consulting, LLC

Christy Gilbert, BS, RAC Associate Director, CMC Regulatory Affairs, AstraZeneca

Shyamala Ivatury, MS Senior Director, IPD, PT&D, AstraZeneca

Uwe Niesner, PhD VP, Head Respiratory & Biologics Regulatory Strategy, Viatris

Bryan Newman, PhD
Lead Pharmacologist, DTP I, ORS, OGD, FDA
Ross Walenga, PhD
Senior Chemical Engineer, DQMM, ORS, OGD, FDA
Anubhav Kaviratna, PhD
Biomedical Engineer, DTP I, ORS, OGD, FDA
Hailing Zhang, PhD
Division Director, DPQA XII, OPQA II, OPQ, FDA

4:45 PM – 5:00 PM	Closing Remarks (Day 1) James Polli, PhD Anna Schwendeman, PhD	Co-Director, CRCG Co-Director, CRCG

Day 2 December 5

8:30 AM – 8:35 AM Workshop Day 2 Overview

Bryan Newman, PhD Lead Pharmacologist, DTP I, ORS, OGD, FDA

Session 1: Generic LGWP MDI Development and the Generic Industry Experience

This session will include presentations from FDA on the scientific and regulatory considerations for generic LGWP propellant MDI development programs from the quality and bioequivalence perspectives. Industry presentations will then discuss the current experiences and challenges encountered by generic developers for an LGWP propellant MDI.

8:35 AM – 8:40 AM	Speaker Introductions Bryan Newman, PhD	Lead Pharmacologist, DTP I, ORS, OGD, FDA
8:40 AM – 9:00 AM	Summary of Small Group Working Se Bryan Newman, PhD	Lead Pharmacologist, DTP I, ORS, OGD, FDA
9:00 AM – 9:20 AM	Policy Considerations for Generic MD Rachael Dippold, PhD, JD	Pls Transitioning to an LGWP Propellant Regulatory Counsel, DPD, OGDP, OGD, FDA
9:20 AM – 9:40 AM	Generic MDI LGWP Propellant Transi Elizabeth Bielski, PhD	tion: OGD Framework and Data Submission Recommendations Senior Pharmacologist, DTP I, ORS, OGD, FDA
9:40 AM – 10:00 AM	Coffee Break	
10:00 AM – 10:20 AM	Alternative In Vitro Bioequivalence A Lucas W. S. Silva, BSc	pproaches for the Low GWP Propellant Transition Senior Specialist, Analytical Development, Nanopharm, An Aptar Pharma Co.
10:20 AM – 10:40 AM	No Time to Lose: Adopting a Science- Rupi Pannu, PhD	Based Approach to Ensure Continued Access to Generic pMDI Products Senior Director, Respiratory R&D Project Leader, Respiratory R&D, Viatris
10:40 AM – 11:00 AM	Perspective on Generic LGWP MDI De Siva Vaithiyalingam, PhD	evelopment Senior Vice President/Head of US Regulatory Affairs, Cipla, Ltd
11:00 AM – 11:50 AM Moderator: Panelists:	Q&A Session with Panel Bryan Newman, PhD Pradeep Bhadauria, MPharm Andrew Clerman, MD, PhD William Feldman, MD, DPhil, MPH Dhaval Gaglani, MS Bing Li, PhD Rupi Pannu, PhD Lucas W. S. Silva, BSc Siva Vaithiyalingam, PhD Ross Walenga, PhD	Lead Pharmacologist, DTP I, ORS, OGD, FDA President and Global Chief Scientific Officer, Cipla Ltd Acting Lead Physician, DTP I, ORS, OGD, FDA Assoc. Physician, Pulmonary & Critical Care Med, Faculty, Regulation Program Therapeutics & Law, Pharmacoepidemiology & Pharmacoeconomics, Assoc. Dir., Ethics Service, Brigham & Women's Hosp., Asst. Prof, Harvard Med School Supervisor, DPQQ V, OPQA I, OPQ, FDA Associate Director for Science, OB, OGD, FDA Senior Director, Respiratory R&D Project Leader, Respiratory R&D, Viatris Senior Specialist, Analytical Development, Nanopharm, An Aptar Pharma Co. Senior Vice President/Head of US Regulatory Affairs, Cipla, Ltd Senior Chemical Engineer, DQMM, ORS, OGD, FDA

Session 2: The Global LGWP Propellant Transition

Lunch Break

11:50 AM - 12:50 PM

This session will cover the global perspective on the LGWP propellant transition. Presentations will include discussion on how similarities and differences in regulatory thinking may impact LGWP propellant MDI development strategies and areas for study harmonization that could minimize potential challenges for drug developers seeking approval in different markets.

12:50 PM – 12:55 PM	Speaker Introductions Sarah Ibrahim, PhD	Associate Director, Stakeholder and Global Engagement, OGD, FDA
12:55 PM – 1:10 PM	Considerations from the Global Pro Karolina Törneke, DVM	opellant Transition Assistant Prof and Senior Clinical Assessor, Swedish Medical Products Agency
1:10 PM – 1:25 PM	Current MHRA Approach on Data Requirements for the Transition to Low GWP Propellants in pMDIs Nithyanandan Nagercoil, MBBS, MD Senior Medical Assessor, MHRA	

Orla Ní Ógáin, PhD, PGDip Stat QI Senior Pharmaceutical Assessor, Medicines & Healthcare Products, MHRA

1:25 PM - 1:40 PM Navigating the Regulatory Landscape: Sustaining Patient Care with Next-Gen LGWP MDI Propellants

> Mark Boelens, PhD Global Senior Director, Product Stewardship & Toxicology, Honeywell

1:40 PM - 1:55 PM Key Considerations in the Business Decision to Reformulate a HFA-based pMDI with LGWP

> Geraldine Venthoye, PhD, BPharm, Chief Scientific Officer & EVP Inhaled Technology Platforms, Vectura Group

MRPharmS

1:55 PM - 2:10 PM Considerations and Challenges Facing Generic Manufacturers Transitioning to LGWPs

> Giuseppe Randazzo, MS Senior Vice President, Sciences and Regulatory Affairs Association for

> > Accessible Medicines (AAM)

2:10 PM - 2:55 PM **Q&A Session with Panel**

> Moderator: Sarah Ibrahim, PhD Associate Director, Stakeholder and Global Engagement, OGD, FDA Panelists:

Mark Boelens, PhD Global Senior Director, Product Stewardship & Toxicology, Honeywell

Nithyanandan Nagercoil, MBBS, MD Senior Medical Assessor, MHRA

Orla Ní Ógáin, PhD, PGDip Stat QI Senior Pharmaceutical Assessor, Medicines & Healthcare Products, MHRA

Sally Seymour, MD Director, DPACC, OII, OND, FDA

Karolina Törneke, DVM Assistant Prof and Senior Clinical Assessor, Swedish Medical Products Agency

Geraldine Venthoye, PhD, BPharm, Chief Scientific Officer & EVP Inhaled Technology Platforms, Vectura Group **MRPharmS**

Paul Wielowieyski, MSc

Sr. Drug Evaluator, Division of Biopharmaceutics Evaluation, Health Canada

Deputy Director, ORS, OGD, FDA Lei Zhang, PhD, FAAPS

Giuseppe Randazzo, MS Senior Vice President, Sciences and Regulatory Affairs Association for

Accessible Medicines (AAM)

2:55 PM - 3:05 PM Coffee Break

Session 3: Lessons Learned and Closing Remarks

Following the presentations across both Day 1 and 2, this panel discussion will cover the lessons learned, along with identifying the areas where the FDA can provide additional guidance on areas that are most needed to address the ongoing challenges affecting LGWP propellant MDI development.

3:05 PM - 3:50 PM **Holistic Panel Discussion**

> Moderator: Darby Kozak, PhD Deputy Director, OGD, FDA

Panelists: Scientific Director, Inhalation Product Development, Kindeva Drug Delivery Stephen Stein, MS

> Richard (Rik) Lostritto, PhD Consultant, Lostritto Consulting, LLC

Lucas W. S. Silva, BSc Senior Specialist, Analytical Development, Nanopharm, An Aptar Pharma Co.

Global Senior Director, Product Stewardship & Toxicology, Honeywell Mark Boelens, PhD

Poonam Gulati, PhD, MBA, PGCert Senior Director & Team Lead, Global Regulatory Affairs, GSK

Uwe Niesner, PhD VP, Head Respiratory & Biologics Regulatory Strategy, Viatris Siva Vaithiyalingam, PhD Senior Vice President/Head of US Regulatory Affairs, Cipla, Ltd

3:50 PM - 4:00 PM Closing Remarks (Day 2)

Robert Lionberger, PhD Director, ORS, OGD, FDA

Appendix of Abbreviations

ANDA Abbreviated New Drug Application

Assoc. Associate

BS or BSc Bachelor of Science
BPharm Bachelor of Pharmacy

Chem Chemistry

CMC Chemistry, Manufacturing and Controls

Co. Company

CRCG Center for Research on Complex Generics

Dept Department

DIIP Division of Inflammation and Immune Pharmacology

Dir Director

DPACC Division of Pulmonary Allergy and Critical Care

DPD Division of Policy Development

DPhil Doctor of Philosophy

DPQA VII Division of Product Quality Assessment VII
DPQA XII Division of Product Quality Assessment XII

DPT II Division of Pharmacology/Toxicology for Immunology and Inflammation

Dr Doctor

DTP I Division of Therapeutic Performance I

DQMM Division of Quantitative Methods and Modeling

DVM Doctor of Veterinary Medicine
EPA Environmental Protection Agency

EVP Executive Vice President FDA Food and Drug Administration

FAAPS Fellow of the American Association of Pharmaceutical Scientists

GSK GlaxoSmithKline

IPD Inhalation Product Development
LGWP Low Global Warming Potential
LLC Limited Liability Company

LTD Limited

MBA Master of Business Administration

MBBS Bachelor of Medicine, Bachelor of Surgery in India

MChem Master of Chemistry MD Doctor of Medicine

Med Medicine

MHRA Medicines and Healthcare products Regulatory Agency

MPH Master of Public Health MPharm Master of Pharmacy

MRPharmS Member of the Royal Pharmaceutical Society

MS or MSci Master of Science

NDA New Drug Application

OB Office of Bioequivalence

OGD Office of Generic Drugs

OGDP Office of Generic Drug Policy

OII Office of Immunology and Inflammation
OPQ Office of Pharmaceutical Quality
OPQA I Office of Pharmaceutical Quality I

OPQA II Office of Pharmaceutical Quality Assessment II

ORS Office of Research and Standards
OTS Office of Translational Sciences
PGCert Post Graduate Certification

PGDip Stat QI Post Graduate Diploma in Statistics and Quality Improvement

PhD Doctor of Philosophy

Prof Professor

PT&D Pharmaceutical Technology & Development

RAC Regulatory Affairs Certification
R&D Research and Development

Sr Senior

VP Vice President