CDER SMALL BUSINESS AND INDUSTRY ASSISTANCE

DRUG MASTER FILE (DMF) AND DRUG SUBSTANCE WORKSHOP



Version 5 - Updated February 20, 2021

For files and resources, please visit The Event Page on SBIAevents.com

Add Event to Your Calendar

AGENDA

All times are Eastern (EST UTC-5)

View Start Time on World Clock

DAY ONE: Wednesday, March 3, 2021

7:45 - 8:05

Administrative Overview

Brenda Stodart

CAPT, USPHS

Director, Small Business and Industry Assistance (SBIA) Division of Drug Information (DDI) | Office of Communications (OCOMM) | CDER

8:05 - 8:20

Welcome

Lawrence Yu

Director

Office of New Drug Products (ONDP) Office of Pharmaceutical Quality (OPQ) | CDER

Your SBIA Hosts for Day One

Forest "Ray" Ford, Jr.

CAPT. USPHS. Pharmacist DDI | OCOMM | CDER

Lisa Misevicz

Health Communications Specialist SBIA | DDI | OCOMM | CDER

8:20 - 8:25

Session Introduction

Erin Skoda

Branch Chief (Acting) Division of Lifecycle API ONDP | OPQ | CDER

8:25 - 8:55

Introduction to the DMF Review Process

The DMF review process from a timeline perspective with an emphasis on key takeaways from the workshop will be covered.

Erin Skoda Branch Chief (Acting) Division of Lifecycle API ONDP | OPQ | CDER

8:55 - 9:25

Administrative Aspects of Managing a DMF

We will discuss the administrative timeline of a DMF from requesting a pre-assigned DMF number to progression of status from pending to active and subsequent submissions with advice on administrative aspects of managing a DMF.

Vathsala Selvam Technical Information Specialist

Division of Lifecycle API ONDP | OPQ | CDER

9:25 - 9:50

Q&A Panel

Erin Skoda, Vathsala Selvam, David Skanchy

9:50 - 10:05: BREAK

10:05 - 10:30

Managing Electronic DMF Submissions

Information to manage a DMF in eCTD format including electronic submission requirements, metrics, best practices, frequently asked questions, and where to obtain help will be covered.

Jonathan Resnick

Project Management Officer
Cloud Collaboration Capability Team
Division of Data Management Services and Solutions
Office of Business Informatics (OBI) | CDER

10:30 - 10:45

Drug Master Files from a GDUFA II User Fee Perspective

Information about DMF user fee assessment including fee requirement, payment, best practices, frequently asked questions, and where to obtain help will be covered.

Hanah Pham

Commander, USPHS

Evelyn Hong

Lieutenant Commander, USPHS

Division of User Fee Management and Budget Formulation (DUFMBF)
Office of Management (OM) | CDER

10:45 - 11:10

Timely Consult and Early Information Request (TCIR) Process for Drug Master Files (DMFs)

Background and data on the TCIR process for DMFs, which could have a substantial positive impact on the overall ANDA approval process, is discussed.

Jayani Perera
Chemist
Division of Lifecycle API
ONDP | OPQ | CDER

11:10 - 11:40

Effective Communication Strategies for Drug Master Files (DMF)

Critical pathways, modes, types, and timing of communications during the DMF and application lifecycle along with advice and best practices from an FDA perspective on maximizing the effectiveness of these communications will be covered.

David Skanchy

Commander, USPHS
Director, Division of Lifecycle API
ONDP | OPQ | CDER

Benjamin Danso

Commander, USPHS
Lead DMF Project Manager
Office of Program and Regulatory Operations (OPRO)
| OPQ | CDER

11:40 - 12:00

Q&A Panel

Jonathan Resnick, Hanah Pham, Evelyn Hong, Jayani Perera, David Skanchy, Benjamin Danso

12:00 - 1:00: LUNCH BREAK

1:00 - 1:30

Poster Presentations: Responses to Submitted Questions

NOTE: Poster Presentations will be made available for viewing before the Workshop. Please visit the Workshop Event Page for information on how to view the presentations and submit questions.

Poster Presenters

1:30 - 1:35

Manufacturing Session Introduction

Erin Skoda

Branch Chief (Acting)
Division of Lifecycle API
ONDP | OPQ | CDER

1:35 - 2:00

Drug Substance Facilities – Hidden and Critical Intermediate

This presentation covers critical intermediates and how to avoid DMF hidden facilities in order to prevent delays in referencing application approvals.

Wei Liu

Senior Pharmaceutical Quality Assessor
Division of Lifecycle API
ONDP | OPQ | CDER

Cassandra Abellard

Quality Assessor/ Consumer Safety Officer
Division of Pharmaceutical Manufacturing
Office of Pharmaceutical Manufacturing Assessment
(OPMA) | OPQ | CDER

2:00 - 2:25

ICH Q11 Q&A, a Supporting Document for the Selection and Justification of Starting Materials

This presentation will provide key concepts and clarification for starting materials selection based on ICH Q11 Q&A.

Anita Tiwari Senior Pharmaceutical Quality Assessor Division of Lifecycle API ONDP | OPQ | CDER

2:25 - 2:40

Q&A Panel

Wei Liu, Cassandra Abellard, Anita Tiwari, David Skanchy

2:40 - 2:55: BREAK

2:55 - 3:20

Common Issues Related to LC and GC Methods in Type II DMFs

This presentation will focus on commonly observed issues related to LC and GC analytical procedures and validation.

Xinghua Wu Chemist Division of Lifecycle API ONDP | OPQ | CDER

3:20 - 3:45

Process Validation and ICH Q7

This presentation covers manufacturing validation data from an FDA review perspective.

David Amspacher
Chemist
Division of Lifecycle API
ONDP | OPQ | CDER

3:45 - 4:10

Regulatory Considerations in Demonstrating Complex API Sameness

Regulatory strategies to show API sameness of complex APIs in generic drug product will be discussed.

Bapu R. Gaddam

Chemist

Division of Lifecycle API

ONDP | OPQ | CDER

4:10 - 4:40

Q&A Panel

Xinghua Wu, David Amspacher, Bapu R. Gaddam, David Skanchy

4:40 - 4:45

Day One Closing

Erin Skoda

Branch Chief (Acting)
Division of Lifecycle API
ONDP | OPQ | CDER

4:45: DAY ONE ADJOURN

DAY TWO: Thursday, March 4, 2021

8:05 - 8:15

Administrative Overview

Lisa Misevicz

Health Communications Specialist SBIA | DDI | OCOMM | CDER

8:15 - 8:20

Welcome & Session Introduction

Ramnarayan Randad

Branch Chief Division of Lifecycle API ONDP | OPQ | CDER

Your SBIA Hosts for Day Two

Forest "Ray" Ford, Jr. CAPT, USPHS, Pharmacist DDI | OCOMM | CDER Lisa Misevicz

Health Communications Specialist SBIA | DDI | OCOMM | CDER

8:20 - 8:50

Regulatory Considerations for Impurity Qualification: ICH Q3A/Q3C/Q3D, RLD & MDD

We will present case studies on how to establish clinically relevant impurities specifications.

Hongbiao Liao Chemist Division of Lifecycle API ONDP | OPQ | CDER

8:50 - 9:20

ICH M7(R1) – Chemistry and Manufacturing Control (CMC) Perspective on Hazard Assessment

This presentation outlines the key concepts surrounding hazard assessment and impurity classification per ICH M7.

Barbara O. Scott *Chemist*Division of Lifecycle API

ONDP | OPQ | CDER

9:20 - 9:45

Application of (Q)SAR and Expert Knowledge for ICH M7 Impurity Classification

The basic concepts, technical considerations, and best practices for comprehensive reporting of (Q)SAR results and common deficiencies encountered by FDA in regulatory submissions will be presented.

Naomi L. Kruhlak

Scientific Lead

Computational Toxicology Consultation Service (CTCS)
Division of Applied Regulatory Science (DARS)
Office of Translational Sciences (OTS) | CDER

Office of Translational Sciences (OTS) | ODEN

9:45 - 10:15

Q&A Panel

Hongbiao Liao, Barbara O. Scott, Naomi L. Kruhlak

10:15 - 10:30: BREAK

DAY TWO: Thursday, March 4, 2021

10:30 - 10:55

Safety Evaluation of Drug Substance Impurities in Generics

The OGD-Pharmacology/Toxicology (Pharm/Tox) process for safety evaluation of impurities in drug substances is described and illustrated with case studies, emphasizing critical elements considered in safety evaluations, and commonly occurring deficiencies in DMFs.

Chanchal Gupta

Pharmacology/Toxicology Reviewer
Division of Clinical Review (DCR)
Office of Bioequivalence (OB)
Office of Generic Drugs (OGD) | CDER

10:55 - 11:20

Nitrosamines: Where Are We Now?

We will discuss the Agency's current thinking on nitrosamine risk mitigation.

Deborah F. JohnsonBranch Chief

Division of Lifecycle API

ONDP | OPQ | CDER

11:20 - 11:50

Q&A Panel

Chanchal Gupta, Deborah F. Johnson and Sruthi King

Associate Director of Pharmacology/Toxicology
Division of Clinical Review (DCR)
Office of Bioequivalence (OB) | OGD | CDER

11:50 - 12:50: LUNCH BREAK

12:50 - 1:20

Poster Presentations: Responses to Submitted Questions

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Poster Presenters

1:20 - 1:25

Life Cycle Session Introduction

Ramnarayan Randad Branch Chief

Branch Chief
Division of Lifecycle API
ONDP | OPQ | CDER

1:25 - 1:50

API Facility Inspections

The presentation will cover an overview of FDA's inspection program, approach to various types of inspections, recent compliance trends, and certain API-specific scenarios.

Jay Jariwala

Team Leader, Division of Drug Quality Office of Manufacturing Quality Office of Compliance | CDER

DAY TWO: Thursday, March 4, 2021

1:50 - 2:15

Drug Substance Postapproval Changes Guidance: Determination of Impurity Profile Equivalence

The presentation will provide an overview of the assessment of risk factors with respect to the control of impurities and recommendations for documenting the risk-based determination.

Brian Connell

Senior Pharmaceutical Quality Assessor Division of Lifecycle API ONDP | OPQ | CDER

2:15 - 2:30

Q&A Panel

Jay Jariwala, Brian Connell

2:30 - 2:45: BREAK

2:45 - 3:10

Common CMC Issues in Type II DMFs and How to Avoid Them

This presentation will focus on most common quality issues in DMF submissions and briefly discuss resolution strategies and points to consider to enhance DMF submissions.

Wei Liu

Senior Pharmaceutical Quality Assessor Division of Lifecycle API ONDP | OPQ | CDER

3:10 - 3:35

Modernizing Drug Substance Assessment through KASA

We will discuss the current status of Knowledge-aided Assessment and Structured Application (KASA) for API. Larisa Wu

Associate Director for Science and Communications (Acting) ONDP | OPQ | CDER

3:35 - 4:05

Q&A Panel

Wei Liu, Larisa Wu, David Skanchy

4:05 - 4:15

Day Two Closing

David Skanchy

Commander, USPHS Director, Division of Lifecycle API ONDP | OPQ | CDER

4:15: ADJOURN