CDER SMALL BUSINESS AND INDUSTRY ASSISTANCE (SBIA)

# FDA *NanoDay* SYMPOSIUM 2022



Version 2 – Updated September 3, 2022

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# October 11, 2022 AGENDA

All times are Eastern (EDT UTC-4)

View Start Time on World Clock

8:50 - 9:00

#### **Administrative Overview**

# Brenda Stodart, PharmD, MS, BCGP, RAC

CAPT, USPHS

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

9:05 - 9:10

Welcome

Anil Patri, PhD

FDA Nanocore Director

Office of Scientific Coordination (OSC)

National Center for Toxicological Research (NCTR)

9:00 - 9:05

# **Keynote**

#### **Douglas Throckmorton, MD**

Deputy Director for Regulatory Programs

CDER

# Your SBIA Hosts for Day One

Forest "Ray" Ford, PharmD, BCPS

CAPT, USPHS

DDI | OCOMM | CDER

Renu Lal, PharmD LCDR, USPHS DDI | OCOMM | CDER Nora Lim, PharmD, BCPS LT USPHS, Pharmacist SBIA | DDI | OCOMM | CDER 9:10 - 9:40

# CMC Guidance For Development of Products that Contain Nanomaterials

We will discuss the recently finalized Guidance to Industry, "Drug Products, Including Biologicals, that Contain Nanomaterials" and provide feedback for how this can be applied to development of new products that contain nanomaterials

Olen Stephens, PhD

Chemist

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

9:40 - 10:10

# Nonclinical Perspective on Development of Drug Products Containing Nanomaterials

# Wimolnut Manheng, PhD

**Toxicologist** 

Division of Hematology Oncology Toxicology (DHOT) Office of Oncology Drugs (OOD) | CDER

10:10 - 10:30

**Q&A Panel** 

Olen Stephens and Wimolnut Manheng

#### 10:30 - 10:45: BREAK

10:45 - 11:15

# **Development and Characterization of Generic Drug Products Containing Nanomaterials**

We will discuss the recently finalized Guidance to Industry, "Drug Products, Including Biologicals, that Contain Nanomaterials" in context to generic drug development and review. In particular we will discuss FDA funded research into the characterization of generic drug products containing nanomaterials, how this research informs FDA's product-specific guidances for generic drug development, and we will highlight some recently approved generic drug products containing nanomaterials.

Darby Kozak, PhD

Deputy Division Director

Division of Therapeutic Performance 1 (DTPI) Office of Research and Standards (ORS) Office of Generic Drugs (OGD) | CDER

11:15 - 11:45

# Considerations for the Quality, Safety and Efficacy of Prophylactic Lipid Nanoparticle mRNA **Vaccines**

**Keith Peden** 

Microbiologist Division of Viral Products

Office of Vaccines Research and Review (OVRR)

Center for Biologics Evaluation & Research (CBER)

11:45 - 12:05

**Q&A Panel** 

Darby Kozak and Keith Peden

12:05 - 12:50: LUNCH BREAK

12:50 - 1:20

# Safety Evaluation of Food Contact Substances Containing Nanomaterials

In this presentation, we will discuss FDA's food contact notification (FCN) program and how the safety of a food contact substance (FCS) is evaluated. In addition, we will discuss the guidance document on significant manufacturing changes and its relevance to FCSs that contain nanomaterials, as well as provide considerations when assessing the safety of FCSs that contain nanomaterials.

# **Raymond Brinas**

Division of Food Contact Substances Office of Food Safety and Applied Nutrition (OFSAN) Center for Food Safety and Applied Nutrition (CFSAN)

1:20 - 1:50

# Nanomaterial Standards Development at FDA

Anil Patri, PhD

FDA Nanocore Director OSC | NCTR

Jiwen Zheng, PhD

Division of Health Technology 2 C Office of Health Technology 2 Center for Devices and Radiological Health (CDRH)

#### 1:50 - 2:05 PM: BREAK

2:05 - 2:35

# **Future of Continuous Manufacturing of Drug Products Containing Nanomaterials**

The talk will highlight the need and opportunity in advancing manufacturing technology of drug products, with a special focus on those containing nanomaterials such as liposomes and lipid nanoparticles

Xiaoming Xu, PhD

Office of Testing and Research (OTR)

OPQ | CDER

2:35 - 3:05

**Q&A Panel** 

Raymond Brinas, Anil Patri, Jiwen Zheng, and Xiaoming Xu

3:05 - 3:10

## **Symposium Closing**

Olen Stephens, PhD

Chemist

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

3:10 PM: ADJOURM SYMPOSIUM