

FOOD AND DRUG ADMINISTRATION (FDA)
Center for Drug Evaluation and Research (CDER)

***Meeting of the Pediatric Oncology Subcommittee of the
Oncologic Drugs Advisory Committee (pedsODAC)***

May 11-12, 2021

DRAFT AGENDA

On May 11, 2021, the subcommittee will discuss the development and successful implementation of the Pediatric Patient-Reported Outcomes Version of the Common Terminology Criteria for Adverse Events (PRO-CTCAE) as a tool for eliciting the patient's voice in oncology clinical trials to more accurately determine tolerability and toxicity of drugs under investigation. The subcommittee will also address the challenges of capturing this type of data across the age spectrum of the pediatric population and possible generalizability of the data. It will consider approaches to address concerns about excluding the patient voice of young children deemed incapable of self-reporting. The subcommittee will also focus on approaches to investigators and commercial sponsors to use the Pediatric PRO-CTCAE in toxicity assessment moving forward.

Day 1: May 11, 2021

10:00 a.m.	Call to Order	Alberto S. Pappo, MD Chairperson, pedsODAC
10:05 a.m.	Introduction of Subcommittee and Conflict of Interest Statement	She-Chia Chen, PharmD Designated Federal Officer, ODAC
10:10 a.m.	Introductory Remarks	Gregory Reaman, MD Associate Director Pediatric Oncology Oncology Center of Excellence (OCE) Office of the Commissioner (OC) Associate Director Pediatric Oncology Office of Oncologic Diseases (OOD) Office of New Drugs (OND), CDER, FDA
10:15 a.m.	FDA PRESENTATIONS	
	FDA Perspective: Patient Self-Reporting in the Evaluation of Cancer Drug Tolerability	Elizabeth S. Duke, MD Medical Officer Division of Oncology 2, OOD, OND, CDER, FDA
	Patient-Reported Outcomes (PROs) in Pediatric Cancer Registration Trials – An FDA Perspective	Meena N. Murugappan, PharmD, MPH Research Fellow OCE, OC, FDA
10:50 a.m.	Clarifying Questions	

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DRAFT AGENDA (cont.)

11:00 a.m. **SPEAKER PRESENTATION**

Patient Self-Reporting and Cancer
Drug Tolerability: Lessons Learned
from the Adult Experience with
PRO-CTCAE

Lori Minasian, MD
Deputy Director
Division of Cancer Prevention
National Cancer Institute
National Institutes of Health

11:15 a.m. **FDA PRESENTATION**

FDA's Project Patient Voice:
Let the Children be Heard

Vishal Bhatnagar, MD
Associate Director for Patient Outcomes
OCE, OC, FDA

11:30 a.m. **GUEST SPEAKER PRESENTATIONS**

Rationale for the Development of
Pediatric PRO-CTCAE

Pamela S. Hinds, PhD, RN, FAAN
Executive Director
Department of Nursing Science,
Professional Practice & Quality
Research Integrity Officer
Children's National Hospital
Professor of Pediatrics
School of Medicine and Health Sciences
George Washington University

Design and Evaluation of the Pediatric
Patient-Reported Outcomes version of
the Common Terminology Criteria for
Adverse Events (Ped-PRO-CTCAE™)
System

Bryce B. Reeve, PhD
Director
Center for Health Measurement
Professor, Population Health Sciences
Professor, Pediatrics
Duke University School of Medicine

12:00 p.m. Clarifying Questions

12:20 p.m. **LUNCH**

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DRAFT AGENDA (cont.)

Pediatric PRO-CTCAE: Future directions beyond tolerability

12:50 p.m.

GUEST SPEAKER PRESENTATIONS

Pediatric PRO-CTCAE: Future
Directions Advancing Supportive Care
Strategies - PROs

Lillian Sung, MD, PhD, FRCPC
Professor, Department of Paediatrics
Senior Scientist, Research Institute
Canada Research Chair in Pediatric
Oncology Supportive Care
The Hospital for Sick Children
Toronto, Canada

Facilitating Survivorship Care and
Research

Tara Henderson, MD, MPH
Professor of Pediatrics
Interim Chief, Pediatric Hematology,
Oncology & Stem Cell Transplantation
Director, Childhood, Adolescent and Young
Adult Survivorship Center, Comer
Children's Hospital, University of Chicago

Incorporating Pediatric PRO-CTCAE
in the National Clinical Trials
Network (NCTN) clinical trials

Doug Hawkins, MD
Professor of Pediatrics
University of Washington
Chair, Children's Oncology Group
Seattle Children's Hospital

1:35 p.m.

Clarifying Questions

1:50 p.m.

BREAK

2:00 p.m.

OPEN PUBLIC HEARING

2:30 p.m.

Questions to the Subcommittee and
Subcommittee Discussion

3:00 p.m.

Wrap-Up

Elizabeth Duke, MD

3:10 p.m.

Closing Remarks

Gregory Reaman, MD

3:15 p.m.

ADJOURNMENT

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DRAFT AGENDA (cont.)

On May 12, 2021, the subcommittee will discuss real-world evidence (RWE) for regulatory use in pediatrics, real-world data (RWD) resources, and RWD and RWE to advance pediatric safety assessments of oncology drug products in children within the context of the FDA framework for RWE. Potential data sources and publicly available platforms, including those made possible through the development and implementation of the National Cancer Institute's Childhood Cancer Data Initiative, will be discussed. The potential for use of data sources to construct external controls to evaluate effectiveness of investigational products will be considered given the frequent dependence on single-arm studies due to extremely small study populations, now exaggerated by molecularly defined subtypes of the rare cancer types that occur in children.

Day 2: May 12, 2021

12:00 p.m. Call to Order

Alberto S. Pappo, MD
Chairperson, pedsODAC

12:05 p.m. Introduction of Subcommittee and Conflict
of Interest Statement

She-Chia Chen, PharmD
Designated Federal Officer, ODAC

12:10 p.m. Introductory Remarks

Gregory Reaman, MD
Associate Director Pediatric Oncology
Oncology Center of Excellence (OCE)
Office of the Commissioner (OC)
Associate Director Pediatric Oncology
Office of Oncologic Diseases (OOD)
Office of New Drugs (OND), CDER, FDA

REAL WORLD EVIDENCE (RWE) FOR REGULATORY USE IN PEDIATRICS

12:15 p.m. **FDA PRESENTATIONS**

The FDA RWE Framework and
Considerations for Possible Use in
Regulatory Decision-Making

Jacqueline Corrigan-Curay, JD, MD
Director
CDER Office of Medical Policy (OMP)
Acting Deputy Center Director for
Operations, CDER, FDA

Designing External Controls Using Real
World Data for Pediatric Cancer Drug
Development

Donna R. Rivera, PharmD, MSc
Associate Director for
Pharmacoepidemiology
OCE, OC, FDA

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DRAFT AGENDA (cont.)

FDA PRESENTATIONS (CONT.)

Statistical Considerations for External
Controls

Pallavi Mishra-Kalyani, PhD
Team Leader
Division of Biometrics V
Office of Biostatistics
Office of Translational Sciences
CDER, FDA

12:45 p.m. Clarifying Questions

PEDIATRIC REAL WORLD DATA (RWD) RESOURCES

1:15 p.m. **SPEAKER PRESENTATIONS**

National Cancer Institute (NCI)
Childhood Cancer Data Initiative

James Doroshow, MD
Deputy Director for Clinical and
Translational Research, NCI
National Institutes of Health (NIH)
Director, Division of Cancer Treatment and
Diagnosis, NCI, NIH

The Childhood Cancer Data Initiative
(CCDI) and RWD Resources for
Pediatric Oncology

Malcolm Smith, MD, PhD
Associate Branch Chief for Pediatrics
Clinical Investigations Branch
Cancer Therapy Evaluation Program
Division of Cancer Treatment and
Diagnosis (DCTD), NCI, NIH

2:15 p.m. Clarifying Questions

**RWE IN EVALUATING PEDIATRIC DRUG SAFETY AND
INFORMING RESEARCH STRATEGIES**

2:30 p.m. **FDA PRESENTATION**

Real World Evidence (RWE) to Assess
Pediatric Medical Product Safety

Ann McMahon, MD, MS, FISPE
Deputy Director of Science
Office of Pediatric Therapeutics, OC, FDA

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DRAFT AGENDA (cont.)

2:40 p.m. **GUEST SPEAKER PRESENTATIONS**

Using RWD and RWE to Evaluate Pediatric
Cancer Drug Safety

**Bruce Carleton, BPharm, PharmD, FCP,
FISPE**

Director, Pharmaceutical Outcomes
Programme, BC Children's Hospital
Professor of Pediatrics and Medical
Genetics, Faculty of Medicine
University of British Columbia
Chair, Division of Translational
Therapeutics, Department of Pediatrics
Senior Clinician Scientist
BC Children's Hospital Research Institute
Vancouver, Canada

Informing pediatric clinical research
strategies and drug development through
RWE

Doug Hawkins, MD

Professor of Pediatrics
University of Washington
Seattle Children's Research Institute
Chair, Children's Oncology Group

3:00 p.m. Clarifying Questions

3:15 p.m. **BREAK**

3:30 p.m. **OPEN PUBLIC HEARING**

4:00 p.m. Questions to the Committee and Committee
Discussion

4:45 p.m. **Closing Remarks**

Gregory Reaman, MD

5:00 p.m. **ADJOURNMENT**