

**FDA-AACR Workshop on
How Much is Enough? Trial Designs for Treatment Regimens with Multiple Phases**

Bethesda Marriott Pooks Hill

May 9 | 9:00 AM – 4:00 PM

Workshop Co-chairs:

Harpreet Singh, MD, Director, Division of Oncology 2, U.S. Food and Drug Administration

Elizabeth Jaffee, MD, FAACR, FCP, Deputy Director, Sidney Kimmel Comprehensive Cancer Center

AGENDA

9:00 AM INTRODUCTION

9:00 AM **Welcome & Introduction:**
Elizabeth Jaffee, MD, Sidney Kimmel Comprehensive Cancer Center

9:10 AM **Overview of Workshop:**
Harpreet Singh, MD, U.S. Food and Drug Administration

9:20 AM SESSION 1: CURRENT LANDSCAPE FOR PERIOPERATIVE TRIAL DESIGNS

9:20 AM **Moderator Introduction:**
Erin Larkins, MD, U.S. Food and Drug Administration

9:25 AM **Current Therapeutic Landscape for Early-Stage Solid Tumors:**
Oladimeji Akinboro, MD, U.S. Food and Drug Administration

9:35 AM **Biomarker-Guided Perioperative Clinical Trials:**
Valsamo Anagnostou, MD, PhD, Sidney Kimmel Comprehensive Cancer Center

9:45 AM **Implementing Sequential, Multiple, Randomized (SMART) Trial Designs:**
Kelley Kidwell, PhD, University of Michigan School of Public Health

9:50 AM PANEL DISCUSSION

- **Anup Amatya, PhD**, U.S. Food and Drug Administration
- **Paz Vellanki, MD**, U.S. Food and Drug Administration
- **Thelma Brown**, Translational Breast Cancer Research Consortium
- **Roy Herbst, MD, PhD**, Yale University
- **Mark Kris, MD**, Memorial Sloan Kettering Cancer Center
- **Craig Tendler, MD**, Janssen Research & Development, LLC

10:45 AM BREAK

10:55 AM SESSION 2A: OPTIMIZING PERIOPERATIVE TREATMENT REGIMENS

- 10:55 AM **Moderator Introduction:**
Elizabeth Jaffee, MD, Sidney Kimmel Comprehensive Cancer Center
- 11:00 AM **Optimizing the Regimen: Cooperative Group Perspective:**
Jhanelle Gray, MD, Moffitt Cancer Center
- 11:10 AM **Cumulative and Long-Term Toxicity with Immunotherapy:**
Mark Yarchoan, MD, Sidney Kimmel Comprehensive Cancer Center

11:20 AM PANEL DISCUSSION

- **Vishal Bhatnagar, MD**, U.S. Food and Drug Administration
- **Tatiana Prowell, MD**, U.S. Food and Drug Administration
- **Michael Axelson, MD**, Loxo@Lilly
- **Fred Hirsch, MD, PhD**, The Tisch Cancer Institute at Mount Sinai
- **Jane Perlmutter, PhD, MBA**, Gemini Group
- **Sara Tolaney, MD, MPH**, Dana-Farber Cancer Institute

12:20 PM LUNCH BREAK

- 12:50 PM **Fireside Chat with FDA Leaders**

1:20 PM SESSION 2B: THE FUTURE OF REGISTRATIONAL TRIALS WITH MULTIPLE ARMS

- 1:20 PM **Moderator Introduction:**
Bernardo Haddock Lobo Goulart, MD, U.S. Food and Drug Administration
- 1:25 PM **Statistical Considerations for Future Perioperative Trials:**
Chi Song, PhD, U.S. Food and Drug Administration
- 1:35 PM **Industry Perspective on Future Perioperative Trials:**
Minghua Shan, PhD, Bayer Pharmaceuticals

1:45 PM PANEL DISCUSSION

- **Nicole Gormley, MD**, U.S. Food and Drug Administration
- **Pallavi Mishra-Kalyani, PhD**, U.S. Food and Drug Administration
- **Patrick Forde, MD**, Johns Hopkins Medicine
- **Giuseppe Giaccone, MD, PhD**, Weill Cornell Medical College
- **Aaron Sosa Mejia, MD**, European Medicines Agency
- **Joshua Reuss, MD**, MedStar Georgetown
- **Kathleen Winson, MS**, Genentech, Inc.

2:45 PM BREAK

2:55 PM Session 3: Considerations in Other Therapeutic Areas

2:55 PM **Moderator Introduction:**

Mirat Shah, MD, U.S. Food and Drug Administration

3:00 PM **Where Do We Go from Here? Considerations for NSCLC & Other Therapeutic Areas:**

Harpreet Singh, MD, U.S. Food and Drug Administration

3:10 PM Additional Panelists

- **Stephanie Wethington, MD**, U.S. Food and Drug Administration
- **Naomi Horiba, MD, MPH**, U.S. Food and Drug Administration
- **Christine Gause, PhD**, Merck
- **Manju George, PhD, MVSc**, Colontown
- **Cristina Migali, MD, PhD**, European Medicines Agency
- **Thomas Powles, MD**, Barts-Cancer Institute, London

3:55PM **Concluding Remarks**

Harpreet Singh, MD, U.S. Food and Drug Administration

4:00 PM ADJOURN