



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

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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

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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

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