

Seventh Annual Patient-Reported Outcome Consortium Workshop

Partners in Progress: Sharing the Vision, Shaping the Future

April 27 – 28, 2016

Sheraton Silver Spring Hotel 8777 Georgia Avenue Silver Spring, MD 20910

On April 27-28, 2016 the SEVENTH ANNUAL PATIENT-REPORTED OUTCOME CONSORTIUM WORKSHOP was held in Silver Spring, Maryland. The overall Workshop objectives were to:

- Provide updates on FDA's Clinical Outcome Assessment (COA) Qualification Program and COA Compendium
- Discuss progress made within the PRO Consortium, with particular focus on the development of the Symptoms of Major Depressive Disorder Scale (SMDDS) by the Depression Working Group
- Discuss ways of generating evidence to support the content validity of performance outcome (PerfO) measures used to assess efficacy endpoints in clinical trials
- Provide examples of the development and implementation of COA tools in pediatric clinical trials
- Describe NCI's Patient-Reported Outcomes Version of the Common Terminology Criteria for Adverse Events (PRO-CTCAE) and some of the methodological considerations for its implementation in trials
- Build on last year's workshop presentation titled Interpreting Change in Scores on COA Endpoint Measures by discussing emerging approaches to estimating meaningful change on COA (particularly PRO) measures

The following Workshop Agenda provides an overview of the day-and-a-half-long meeting as well as links to the slide sets and posters presented.

Request Session Recordings

Workshop Agenda – Day 1 April 27, 2016

7:30 – 8:30 am	Registration and Continental Breakfast – Cypress Ballroom

	Lunch – Elm I, Elm II and Magnolia Ballroom
	Michelle Campbell, PhD – Reviewer and Scientific Coordinator, COA Qualification Program, COA Staff, OND, CDER, FDA Stephen Joel Coons, PhD – Executive Director, Patient-Reported Outcome Consortium, Critical Path Institute Billy Dunn, MD – Director, Division of Neurology Products, OND, CDER, FDA Q & A
	Panelists:
	Richard S.E. Keefe, PhD – Professor of Psychiatry, Psychology, and Neuroscience, Duke University Medical Center and CEO, NeuroCog Trials, Inc.
	J. Scott Andrews, PharmD – Research Scientist, Global Patient Outcomes and Real World Evidence, Eli Lilly and Company
	Rachel Ballinger, PhD – Lead Outcomes Researcher, Clinical Outcome Assessment, ICON Clinical Research
	Presenters:
10:45 – 12:15 pm	Session 2: Assessing the Content Validity of Performance Outcome (PerfO) Measures Moderator: Elizabeth (Nicki) Bush, MHS – Research Scientist, Global Patient Outcomes and Real World Evidence, Eli Lilly and Company
10:20 – 10:45 am	Break – 25 min
	Q & A
	Hematology Products, Office of Hematology and Oncology Products (OHOP), OND, CDER, FDA
	Elektra Papadopoulos, MD, MPH – Acting Associate Director, COA Staff, OND, CDER, FDA Virginia Kwitkowski, MS, ACNP-BC – Associate Director for Labeling, Division of
	Presenters:
	COA Staff, Office of New Drugs (OND), Center for Drug Evaluation and Research (CDER), U.S. Food and Drug Administration (FDA)
8:50 – 10:20 am	Session 1: Update from FDA Regarding the Clinical Outcome Assessment Qualification Program and the COA Compendium Moderator: Michelle Campbell, PhD – Reviewer and Scientific Coordinator, COA Qualification Program,
8:30 – 8:50 am	Welcome and Patient-Reported Outcome Consortium UpdateStephen Joel Coons, PhD – Executive Director, Patient-Reported (PRO) Outcome Consortium, Critical Path Institute
	Coordinator, Clinical Outcome Assessments (COA) Qualification Program, COA Staff, Office of New Drugs (OND), Center for Drug Evaluation and Research (CDER), U.S. Food and Drug Administration (FDA)
	Day 1 Morning Moderator: Michelle Campbell, PhD – Reviewer and Scientific

	Day 1 Afternoon Moderator: <i>Stephen Joel Coons, PhD</i> – Executive Director, Patient-Reported Outcome Consortium, Critical Path Institute
1:15 – 2:45 pm	Session 3: Addressing Key Challenges in Developing, Testing and Implementing Clinical Outcome Assessments in Pediatric TrialsModerator: Linda Abetz-Webb - Paediatric PRO Expert, CEO/Senior Research Director, Patient-Centered Outcome Assessments, Ltd. (P-COA)
	Presenters:
	Linda Lowes, PT, PhD – Clinical Therapies Research Director, Nationwide Children's Hospital
	Rob Arbuckle – Vice President and UK Managing Director, Patient-Centered
	Outcomes, Adelphi Values Valdo Arnera, MD – Scientific Advisor and General Manager ERT Geneva, ERT
	Panelists:
	Laura Lee Johnson, PhD – Associate Director, Division of Biometrics III, Office of Biostatistics, Office of Translational Sciences (OTS), CDER, FDA
	Andrew E. Mulberg, MD, FAAP, CPI – Division Deputy Director, Gastroenterology and Inborn Errors Products, OND, CDER, FDA
	Josephine Norquist, MS – Patient-Reported Outcome (PRO) & Study Endpoint Group Lead, Center for Observational and Real-world Evidence (CORE), Merck Sharp & Dohme, Corp.
	Anna Rydén, PhD – Director, Patient Reported Outcomes, AstraZeneca
	Q & A
2:45 – 3:10 pm	Break – 25 min

	Session 4: NCI's Patient-Reported Outcomes Version of the Common
3:10 – 4:40 pm	Terminology Criteria for Adverse Events (PRO-CTCAE): Selecting Items and
	Assessment Frequency for Cancer TrialsModerator: Sandra A. Mitchell, PhD,
	CRNP – Research Scientist and Program Director, Outcomes Research Branch,
	National Cancer Institute (NCI)
	Presenters:
	Paul G. Kluetz, MD – Associate Director of Clinical Science, OHOP, OND, CDER, FDA
	Lori Minasian, MD, FACP – Deputy Director, Division of Cancer Prevention, NCI Ethan Basch, MD, MSc – Director, Cancer Outcomes Research Program, University of North Carolina
	Katarina Halling, MSc – Global Head Patient Reported Outcomes, AstraZeneca and Industry Co-Director, PRO Consortium
	Panelist:
	Selena R. Daniels, PharmD, MS – Reviewer and Acting Team Lead, COA Staff, OND, CDER, FDA
	Q & A
4:40 – 4:55 pm	A Review of FDA PRO Labeling (2011 – 2015) Ari Gnanasakthy, MSc, MBA – Head, Patient Reported Outcomes, RTI Health Solutions
4:55 – 5:00 pm	Day 1 Closing Remarks Adjourn
5:30 – 7:00 pm	Reception and Poster Session – Magnolia Ballroom

Workshop Agenda – Day 2 April 28, 2016

7:30 – 8:30 am	Registration and Continental Breakfast – Cypress Ballroom
	Day 2 Moderator : <i>Katarina Halling, MSc</i> – Global Head Patient Reported Outcomes, AstraZeneca and Industry Co-Director, Patient-Reported Outcome Consortium

Session 5: Developing the Symptoms of Major Depressive Disorder Scale 8:30 – 10:00 am (SMDDS): From Patient Input to Final InstrumentModerator: Elizabeth (Nicki) Bush, MHS - Research Scientist, Global Patient Outcomes and Real World Evidence, Eli Lilly and Company **Presenters**: Elizabeth (Nicki) Bush, MHS – Research Scientist, Global Patient Outcomes and Real World Evidence, Eli Lilly and Company Kelly P. McCarrier, PhD, MPH – Senior Research Scientist, Health Research Associates, Inc. Donald Bushnell, MA – Associate Director, Health Research Associates, Inc. Valdo Arnera, MD - Scientific Advisor and General Manager ERT Geneva, ERT Panelists: Stephen Joel Coons, PhD – Executive Director, Patient-Reported Outcome Consortium, Critical Path Institute Tiffany R. Farchione, MD – Deputy Director, Division of Psychiatry Products (DPP), CDER, FDA Q & A Break – 25 min 10:00 - 10:25 am Session 6: Expanding Our Understanding of Meaningful Change from a Patient 10:25 – 11:55 am **PerspectiveModerator**: Cheryl D. Coon, PhD – Principal, Outcometrix **Presenters**: Mona Martin, RN, MPA – Executive Director, Health Research Associates, Inc. Allison Martin Nguyen, MS – Senior Principal Scientist, Patient-Reported Outcomes & Study Endpoints, Merck Research Laboratories, Merck & Co., Inc. Katarina Halling, MSc - Global Head Patient Reported Outcomes, AstraZeneca and Industry Co-Director, PRO Consortium Karon F. Cook, PhD – Research Professor, Department of Medical Social Sciences, Feinberg School of Medicine, Northwestern University

Panelists:

Wen-Hung Chen, PhD – Reviewer, COA Staff, OND, CDER, FDA *Tara Symonds, PhD* – Strategic Lead, Clinical Outcomes Assessments, Clinical Outcomes Solutions

Kathleen (Kathy) Wywrich, PhD – Vice President, Outcomes Research, Evidera

Q & A

11:55 – 12:15 pm | Closing Remarks | Adjourn |