

# 2024 Clinical Outcome Assessment Program Annual Meeting

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

April 17-18, 2024

**Bethesda North Marriott Hotel & Conference Center**  
**5701 Marinelli Rd**  
**Rockville, MD 20852**





On April 17-18, 2024 the Clinical Outcome Assessment Program Annual Meeting was held in Rockville, MD.

The following agenda provides an overview of the day-and-a-half-long meeting as well as links to the session recordings, slide decks, and posters.

### *Agenda – Day 1*



<b>7:30–8:30 am</b>	<b>Registration and Breakfast – Veranda (Outside Salon E-H)</b>	
<b>8:30–8:45 am</b>	<a href="#"><u>Welcome</u></a> <i>Cheryl D. Coon, PhD</i> – Vice President, Clinical Outcome Assessment Program, Critical Path Institute	
<b>8:45–9:15 am</b>	<a href="#"><u>Rare Disease Clinical Outcome Assessment Consortium Update</u></a> <i>Lindsey Murray, PhD, MPH</i> – Executive Director, Rare Disease Clinical Outcome Assessment Consortium, Critical Path Institute	

<p><b>9:15–10:30 am</b></p>	<p><b><u>FDA Update</u></b>  <b>Moderator:</b>  <i>Michelle Campbell, PhD</i> – Associate Director, Stakeholder Engagement and Clinical Outcomes, Office of Neuroscience, Office of New Drugs, Center for Drug Evaluation and Research, U.S. Food and Drug Administration</p> <p><b>Presenters:</b>  <i>Robyn Bent, RN, MS</i> – Director, Patient-Focused Drug Development Program, Center for Drug Evaluation and Research, U.S. Food and Drug Administration  <i>Selena Daniels, PharmD, PhD</i> – Deputy Division Director, Division of Clinical Outcome Assessment, Office of Drug Evaluation Sciences, Office of New Drugs, Center for Drug Evaluation and Research, U.S. Food and Drug Administration  <i>Lili Garrad, PhD</i> – Master Scientist, Patient-Focused Statistical Scientists (PFSS), Division of Biometrics III, Office of Biostatistics, Office of Translational Sciences, Center for Drug Evaluation and Research, U.S. Food and Drug Administration  <i>Jessica Mavadia-Shukla, PhD</i> – Director, Medical Device Development Tools Program, Center for Devices and Radiological Health, U.S. Food and Drug Administration  <i>David S. Reasner, PhD</i> – Division Director, Division of Clinical Outcome Assessment, Office of Drug Evaluation Sciences, Office of New Drugs, Center for Drug Evaluation and Research, U.S. Food and Drug Administration</p>	
<p><b>10:30–11:00 am</b></p>	<p><b>Break</b></p>	
<p><b>11:00–12:15 pm</b></p>	<p><b><u>Session 1: Leveraging Digital Health Technology and Partnership Opportunities to Measure Clinical Outcomes That Matter</u></b>  <b>Moderator:</b> <i>Maria Mattera, MPH</i> – Scientific Director, Patient-Reported Outcome Consortium, Critical Path Institute</p> <p><b>Presenters:</b>  <i>Katelyn Keyloun, PharmD, MS</i> – Director, R&amp;D Digital and Data Strategy, AbbVie  <i>Chad Gwaltney, PhD</i> – President, Gwaltney Consulting  <i>Christine Guo, PhD</i> – Chief Scientific Officer, ActiGraph</p> <p><b>Panelists:</b>  <i>Jen Horonjeff, PhD</i> – Founder and CEO, Savvy Cooperative  <i>Dilorom (Delia) Sass, PhD, RN, AGPCNP-BC</i> – Clinical Analyst, Division of Clinical Outcome Assessment, Office of Drug Evaluation Sciences, Office of New Drugs, Center for Drug Evaluation Research, U.S. Food and Drug Administration  <i>Michelle Campbell, PhD</i> – Associate Director of Stakeholder Engagement and Clinical Outcomes, Division of Clinical Outcome Assessment, Office of Neuroscience, Office of New Drugs, Center for Drug Evaluation Research, U.S. Food and Drug Administration</p>	
<p><b>12:15–1:30 pm</b></p>	<p><b>Lunch – Veranda (Outside Salon E-H)</b></p>	

<p><b>1:30–2:00 pm</b></p>	<p><b><u><a href="#">Patient-Reported Outcome Consortium Update</a></u></b>  <i>Sonya Eremenco, MA</i> – Executive Director, Patient-Reported Outcome Consortium, Critical Path Institute</p>	
<p><b>2:00–3:15 pm</b></p>	<p><b><u><a href="#">Session 2: eCOA and DHTs in Oncology: Realizing the Digital Dividend</a></u></b>  <b>Moderator:</b>  <i>Scottie Kern</i> – Executive Director, Electronic Clinical Outcome Assessment Consortium, Critical Path Institute  <b>Presenters:</b>  <i>Ari Gnanasakthy, MBA, MSc</i> – Principal Scientist, RTI Health Solutions  <i>Jennifer Lord-Bessen, PhD</i> – Senior Director, Global HEOR Advanced Scientific Capabilities, Patient-Reported Outcomes Assessment (PROA), Hematology-Oncology, Bristol Myers Squibb  <i>Florence Mowlem, PhD</i> – Vice President of Science, ObvioHealth; 2024 eCOA Consortium Industry Co-Director  <i>Elena Izmailova, PhD</i> – Chief Scientific Officer, Koneksa Health  <i>T. J. Sharpe</i> – Patient Engagement Program Manager, Medidata Solutions  <b>Panelist:</b>  <i>Vishal Bhatnagar, MD</i> – Associate Director for Patient Outcomes, Oncology Center of Excellence, Center for Drug Evaluation and Research, U.S. Food and Drug Administration</p>	
<p><b>3:15–3:45 pm</b></p>	<p><b>Break</b></p>	
<p><b>3:45-5:00 pm</b></p>	<p><b><u><a href="#">Session 3: Clinical Outcome Assessments Used in Trial Entry, Stratification, and Endpoints: Evidence and Implications to Ensure They're Fit For Their Purpose</a></u></b>  <b>Moderator:</b>  <i>Cheryl D. Coon, PhD</i> – Vice President, Clinical Outcome Assessment Program, Critical Path Institute  <b>Presenters:</b>  <i>Cole Ayasse, PhD</i> – Clinical Outcome Assessment Scientist, Critical Path Institute  <b>Panelists:</b>  <i>Katherine Bevans, PhD</i> – Director, Patient Reported Outcomes, Johnson &amp; Johnson Innovative Medicine  <i>Onyeka Illoh, OD, MPH</i> – Team Leader, Division of Clinical Outcome Assessment, Office of New Drugs, Center for Drug Evaluation and Research, U.S. Food and Drug Administration  <i>Gahan Pandina, PhD</i> – Senior Director, Compound Development Team Leader, Johnson &amp; Johnson Innovative Medicine  <i>Mitchell Psotha, MD, PhD</i> – Medical Officer, Division of Cardiology and Nephrology, Office of New Drugs, Center for Drug Evaluation and Research, U.S. Food and Drug Administration  <i>RJ Wirth, PhD</i> – CEO &amp; Managing Partner, Vector Psychometric Group, LLC</p>	
<p><b>5:00–5:10 pm</b></p>	<p><b>Day 1 Closing Remarks</b>  <i>Cheryl D. Coon, PhD</i> – Vice President, Clinical Outcome Assessment Program, Critical Path Institute  <b>Adjourn</b></p>	

<b>5:10–5:30 pm</b>	<b>Open</b>
<b>5:30–7:00 pm</b>	<b>Reception – Foyer E-H and Veranda</b>

*Agenda – Day 2*

<b>7:30–8:30 am</b>	<b>Registration and Breakfast – Veranda (Outside Salon E-H)</b>	
<b>8:30-9:00 am</b>	<p><a href="#"><u>Welcome and Electronic Clinical Outcome Assessment Consortium Update</u></a>  <i>Scottie Kern</i> – Executive Director, Electronic Clinical Outcome Assessment Consortium, Critical Path Institute</p>	
<b>9:00–10:15 am</b>	<p><a href="#"><u>Session 4: Mitigating Challenges in Clinical Outcome Assessments With Multiple Respondents: Sharing Lessons Learned From Across the Clinical Outcome Assessment Program</u></a>  <b>Moderator:</b>  <i>Lindsey Murray, PhD, MPH</i> – Executive Director, Rare Disease Clinical Outcome Assessment Consortium, Critical Path Institute  <b>Presenters:</b>  <i>Lindsey Murray, PhD, MPH</i> – Executive Director, Rare Disease Clinical Outcome Assessment Consortium, Critical Path Institute  <i>Adam Scheller, PhD</i> – Senior Scientist, Clinical Outcomes, Denali Therapeutics  <i>Claire Trennery, MSc</i> – Associate Director, Patient-Centered Outcomes, Adelphi Values  <i>Jill V. Platko, PhD</i> – Vice President, Scientific Services, Suvoda  <b>Panelists:</b>  <i>Sarita Edwards, MHA</i> – CEO &amp; President, The E.WE Foundation  <i>Marian M. Strazzeri, MS</i> – Mathematical Statistician, Patient-Focused Statistical Scientists, Office of Biostatistics, Center for Drug Evaluation and Research, U.S. Food and Drug Administration  <i>Nicole Lyn, MPH</i> – Global Health Economics and Value Assessment Business Partner, Sanofi</p>	
<b>10:15–10:45 am</b>	<b>Break</b>	

<p><b>10:45–12:00 pm</b></p>	<p><b><u><a href="#">Session 5: Patient-Reported Outcome Measurement Now and Into the Future: Multi-Stakeholder Perspectives Regarding the Optimization of Patient-Reported Outcome Data in the Decision-Making Process Across the Healthcare Continuum</a></u></b></p> <p><b>Moderator:</b>  <i>Adam Gater, MSc</i> – Senior Director, Adelphi Values</p> <p><b>Presenters:</b>  <i>Sharan Randhawa, MSc</i> – Senior Research Manager, Adelphi Values  <i>Loriana Hernández-Aldama, BA</i> – 2x Survivor, Patient Advocate, ArmorUp for Life  <i>John Powers, III, MD</i> – Professor of Clinical Medicine, George Washington University School of Medicine  <i>Bellinda King-Kallimanis, PhD</i> – Senior Director of Patient-Focused Research, LUNGeivity Foundation  <i>Gary Rice, RPh, MS</i> – Senior Advisor of Specialty &amp; Clinical Strategy, MedImpact Healthcare Systems, Inc.  <i>Eleanor Perfetto, PhD</i> – Professor, University of Maryland  <i>Claire Snyder, PhD</i> – Professor of Medicine, Oncology, and Health Policy &amp; Management, Johns Hopkins University, PROTEUS Consortium (participating as a private consultant)</p> <p><b>Panelist:</b>  <i>Selena Daniels, PharmD, PhD</i> – Deputy Division Director, Division of Clinical Outcome Assessment, Office of Drug Evaluation Sciences, Office of New Drugs, Center for Drug Evaluation and Research, U.S. Food and Drug Administration</p>	
<p><b>12:00–12:15 pm</b></p>	<p><b><u><a href="#">Annual Meeting Wrap Up</a></u></b>  <i>Cheryl D. Coon, PhD</i> – Vice President, Clinical Outcome Assessment Program, Critical Path Institute</p>	
<p><b>12:15–1:30 pm</b></p>	<p><b>Lunch – Veranda (Outside Salon E-H)</b></p>	

**Posters summarizing the status of the PRO Consortium’s working groups, the Rare Disease Subcommittee activities, and the ePRO Consortium are available below:**

- [Chronic Heart Failure Working Group](#)
- [Cognition Working Group](#)
- [Depression Working Group 2.0](#)
- [Irritable Bowel Syndrome Working Group](#)
- [Multiple Sclerosis Working Group](#)
- [Pediatric Asthma Working Group](#)
- [Rheumatoid Arthritis Working Group](#)
- [Small Cell Lung Cancer Working Group](#)
- [Rare Disease COA Consortium](#)
- [eCOA Consortium](#)