

View Now: 12th Annual Patient-Reported Outcome Consortium Workshop – Held Virtually

April 14–15, 2021

On April 14–15, 2021, the 12th Annual Patient-Reported Outcome Consortium Workshop was held in a virtual format.

The following Workshop Agenda provides an overview of the two-day meeting as well as links to the session recordings and slide decks. Links to posters summarizing the status of the PRO Consortium's working groups, the Rare Disease Subcommittee activities, and the ePRO Consortium are located after the agenda.

Agenda – Day 1

11:00–11:20 am	Welcome and Patient-Reported Outcome Consortium UpdateOverview: Provides a high-level summary of the recent accomplishments and ongoing activities within the Patient-Reported Outcome (PRO) Consortium	Welcome to the 12 th Annual Patient-Reported Outcome Consortium Workshop Livert will begin at 11:01 am US ET April 24 - 15, 2021
	Presenter: Sonya Eremenco, MA – Director, Patient-Reported Outcome (PRO) Consortium, Critical Path Institute (C- Path)	

11:20–12:30 pm	Session 1: 2021 FDA Update	U.S. (FOOD & DRUG
	Overview: Provides an update on FDA's Clinical Outcome Assessment (COA) Qualification Program and other initiatives	2021 FDA Update 12 ¹⁰ Annual PRO Consortium Workshop April 14, 2021
	Moderator:	
	<i>Michelle Campbell, PhD</i> – Senior Clinical Analyst for Stakeholder Engagement and Clinical Outcomes, Office of Neuroscience (ON), Office of New Drugs (OND), Center for Drug Evaluation and Research (CDER), U.S. Food and Drug Administration (FDA)	
	Presenters:	
	<i>Robyn Bent, RN, MS</i> – Director, Patient Focused Drug Development Program, CDER, FDA	
	<i>Laura Lee Johnson, PhD</i> – Director, Division of Biometrics III, Office of Biostatistics, Office of Translational Sciences, CDER, FDA	
	<i>Elektra Papadopoulos, MD, MPH</i> – Acting Deputy Director, Division of Clinical Outcome Assessment, Office of Drug Evaluation Sciences (ODES), OND, CDER, FDA	
	<i>David S. Reasner, PhD</i> – Division Director, Division of Clinical Outcome Assessment, ODES, OND, CDER, FDA	
	Q & A	
12:30–12:50 pm	Break – 20 min	

12:50–2:20 pm	Session 2: Migraine: A Tale of Two Paths to PRO- Based Product Labeling	Migraine: A Tale of Two Paths to PRO-Based Product Labeling Session will begin at 12:51 pm US ET
	Overview: Discusses the experience of two pharmaceutical firms' different paths to obtaining FDA- approved PRO-based label claims for novel migraine drugs	22 th Annual Patient-Reported Outcome Consortium Workshop April 14-15, 2021 CRITICAL PATH
	Moderator:	
	Stephen Joel Coons, PhD – Executive Director, PRO Consortium and Senior Vice President, Clinical Outcome Assessment Program, C-Path	
	Opening Remarks:	
	Billy Dunn, MD – Director, ON, CDER, FDA	
	Presenters:	
	<i>Pooja Desai, PhD</i> – Director, US Health Economics, Therapeutic Area Lead – Inflammation, Nephrology and Bone, Amgen	
	<i>Elizabeth (Nicki) Bush, MHS</i> – Senior Advisor and Head, Patient-Focused Outcomes Center of Expertise, Eli Lilly and Company	
	Panelists:	
	<i>Eric Bastings, MD</i> – Acting Director, Division of Neurology I; Deputy Director, ON, CDER, FDA	
	<i>Nick Kozauer, MD</i> – Director, Division of Neurology II, ON, CDER, FDA	
	<i>Elektra Papadopoulos, MD, MPH</i> – Acting Deputy Director, Division of Clinical Outcome Assessment, ODES, OND, CDER, FDA	
	Q & A	
2:20–2:30 pm	Day 1 Wrap Up	

Agenda – Day 2

11:00–11:20 am	eCOA: Getting Better Together Initiative Update	Welcome to the
	Overview: Provides a brief update on the eCOA: Getting Better Together Initiative, an ongoing collaboration between the PRO Consortium and ePRO Consortium	12th Annual Patient-Reported Outcome Consortium Workshop Event will begin at 11:01 am US ET Agril 14 - 15, 2021 CELTICAL PATH
	Presenter:	
	Sonya Eremenco, MA –Director, PRO Consortium and Acting Director, Electronic Patient-Reported Outcome (ePRO) Consortium, C-Path	
11:20–12:50 pm	Session 3: Where Are We Headed with Activity Monitors in Clinical Trials?	Where Are We Headed with Activity Monitors in Clinical Trials?
	Overview: Discusses practical considerations associated with the use of activity monitors to measure efficacy endpoints in clinical trials	12 th Annual Patient-Reported Outcome Consortium Workshop April 14-15, 2021 CRITICAL PATH
	Moderator:	
	<i>Maria Mattera, MPH</i> – Assistant Director, PRO Consortium, C-Path	
	Presenters:	
	Jennifer Goldsack, MChem, MA, MBA – Executive Director, Digital Medicine Society	
	<i>Bill Byrom, PhD</i> – Vice President, Product Intelligence and Positioning, Signant Health	
	<i>Jiat Ling Poon, PhD</i> – Principal Research Scientist, Eli Lilly and Company	
	<i>Milena Anatchkova, PhD</i> – Senior Research Leader, Evidera, Inc.	
	Panelists:	
	Andrew Potter, PhD – Mathematical Statistician, Division of Biometrics I, CDER, FDA	
	Steven Blum, MBA, MA – Asset and Indication Lead, Patient Reported Outcomes Assessment, WWHEOR, Bristol Myers Squibb	
	Q & A	
12:50–1:10 pm	Break – 20 min	

1:10–2:40 pm	Session 4: Identifying COAs for Use in Rare Disease Treatment TrialsOverview: Provides an update regarding the establishment and goals of the Rare Disease COA Consortium and presents results of work accomplished by the PRO Consortium's Rare Disease SubcommitteeModerator:Lindsey Murray, PhD, MPH – Associate Director, PRO Consortium, C-PathPresenters:Tori Brooks, MPH – Research Associate II, Mapi Research TrustLindsey Murray, PhD, MPH – Associate Director, PRO Consortium, C-PathPresenters:Tori Brooks, MPH – Research Associate II, Mapi Research TrustLindsey Murray, PhD, MPH – Associate Director, PRO Consortium, C-PathKiera Berggren, MA/CCC-SLP, MS – Research Speech- Language Pathologist, Department of Neurology, Virginia Commonwealth UniversityPanelists:Naomi Knoble, PhD – Reviewer, Division of Clinical Outcome Assessment, OND, CDER, FDADawn Phillips, PT, MS, PhD – Director, Clinical Scientist, Outcomes Research, REGENXBIO Inc.Adam Shaywitz, MD, PhD – Chief Medical Officer, BridgeBio Gene TherapyAllison Seebald – Senior Research Program Manager, National Organization for Rare DisordersQ & A	<text><text><text></text></text></text>
2:40–2:45 pm	Workshop Wrap Up – Sonya Eremenco	

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

Posters		
Chronic Heart Failure Working Group	Pediatric Asthma Working Group	
Cognition Working Group	Rheumatoid Arthritis Working Group	
Depression Working Group 2.0	Small Cell Lung Cancer Working Group	
Functional Dyspepsia Working Group	Rare Disease Subcommittee	
Irritable Bowel Syndrome Working Group	ePRO Consortium	
Multiple Sclerosis Working Group		