

Coalition Against Major Diseases and FDA 2015 Annual Scientific Workshop

October 15, 2015

FDA White Oak Campus

CO-SPONSORED BY:

Critical Path Institute

U.S. Food and Drug Administration

Overview and Objectives

The Coalition Against Major Diseases (CAMD) is a public-private-partnership aimed at creating new tools and methods that can be applied to increase the efficiency of the development process of new treatments for Alzheimer's disease (AD) and Parkinson's disease (PD). The annual meeting brings together members from the pharmaceutical industry, academic key opinion leaders, NIA, FDA, EMA and advocacy groups. The objectives of the meeting are: understand accomplishments of CAMD scientific projects, discuss how these tools are currently or will be applied in drug development, obtain commitment for sharing information/data to begin quantifying benefits of these tools, and facilitate robust and open discussion among all parties of drug development in Alzheimer's and Parkinson's diseases. Experts in the fields of Alzheimer's disease and Parkinson's disease, and leaders of the patient stakeholder community will deliver keynote presentations and regulatory science will be prominently featured throughout the meeting.

Click here for the 2015 Annual Meeting Minutes.

Agenda

7:30-8:30 am	Continental Breakfast
8:30-9:00 am	Welcoming Remarks Martha Brumfield, CEO (Critical Path Institute) Janet Woodcock, Director (Center for Drug Evaluation and Research, FDA) Diane Stephenson & Stephen Arneric, Executive Co-Directors (CAMD)
9:00-9:20 am	Keynote Address Manuel Haas (EMA) Coalition Against Major Diseases and FDA 2015 Annual Scientific Workshop

9:20-9:40 am	Regulatory Perspectives
	ShaAvrée Buckman-Garner (FDA)
	FDA Biomarker Learnings and the Future
9:40-9:55 am	BREAK
a Gordon, Chair	elopments in CAMD Working Groups Meeting the Needs of the Parkinson's Community
9:55-10:10 am	Steve Ford (Parkinson's UK)
10.10.10.27	Computational Modeling for AD
10:10-10:25 am	Julie Stone (Merck) & Klaus Romero (C-Path)
	Where has CAMD come and where do we need to go?
	How can we achieve better understanding of disease progression and efficient clinical populations?
	AD Hippocampal Volume Team
10:25-10:40 am	AD Hippocampal Volume Team Derek Hill (Ixico)
10:25-10:40 am	
10:25-10:40 am 10:40-10:55 am	Derek Hill (Ixico)
	Derek Hill (Ixico) AD CSF Biomarkers Team

SESSION II:

Strategies for Successful Implementation of Biomarkers in Clinical Trials CAMD Data Sharing and Integration...Looking to the Future

11:40-12:25 pm **LUNCH & AWARDS**

Peter Loupos, Chair (Sanofi)

12:25-12:40 pm	Data SharingWhat Can Be Learned from ALS? Melanie Leitner (Biogen)
12:40-12:55 pm	PPMI Paving the Way for Defining Prodromal PD Ken Marek (MNI)
12:55-1:10 pm	Data sharingSuccess Story from Multiple Sclerosis (MSOAC) Jesse Cedarbaum (Biogen)
1:10-1:25 pm	Panel Discussion on Prospective Directions for CAMDFocus on Data Melanie Leitner, Jesse Cedarbaum, Ken Marek, Paul Maruff (Cogstate, AIBL)
1:25-1:35 pm	BREAK
1:35-1:50 pm	Transforming Alzheimer's Disease Therapies Through Collaboration James Hendrix (Alzheimer's Association)

Integrated Focus Sessions		
1:50-2:30 pm	Session III: Modeling	
	How can we achieve better understanding of disease progression and efficient clinical populations?	
	Vikram Sinha (FDA Co-chair) & Klaus Romero (CAMD Co-chair)	
2:30-3:10 pm	Session IV: Biomarkers	
	Chris Leptak (FDA Co-chair) & Richard Meibach (CAMD Co-chair)	
3:10-3:50 pm	Session V: Digital Biomarker Technologies	
	Medical Device Regulatory Decision Points	
	Defining Context of Use and Challenges to Deploying Wearables and Digitial Techno Peter Como (FDA Co-chair) & Jesse Cedarbaum (CAMD Co-chair)	
3:50-4:05 pm	BREAK	
4:05-4:35 pm	KEY RECOMMENDATIONS: SESSIONS III-V	
4:35-4:45 pm	Wrap-up and Looking Ahead Diane Stephenson & Stephen Arneric	