Innovations in Regulatory Science Summit
Sunday, January 12, 2020 | 8 am - 5 pm

Keynote Speakers
Janet Woodcock, MD
Center for Drug Evaluation and Research, FDA

Peter Marks, MD, PhD
Center for Biologics Evaluation and Research, FDA

Panel Discussions
• Accelerating Clinical Trials in the Development and Approval of Innovative Medical Products
• Academia, Government and Industry in Regulatory Science: Cross-Sector Collaboration
• Real-World Evidence, Artificial Intelligence and Novel Medical Devices
• Advancing Discovery to First-In-Human Clinical Trials for New Medical Products

Amy Abernethy (FDA)
Russ Altman (Stanford)
Hal Barron (GlaxoSmithKline)
Howard Bauchner (JAMA)
Jay Bradner (Novartis)
Robert Califf (Verily)
Edward Chang (UCSF)
Mildred Cho (Stanford)
Arthur Ciociola (Novartis)
Laura Esserman (UCSF)
Malvina Eydelman (FDA)
Adam Gazzaley (UCSF)
Kathy Giacomini (UCSF)
Steven Goodman (Stanford)
Sam Hawgood (UCSF)
Sandra Horning (Genentech)
Daphne Koller (Insitro)
Carol Linden (FDA)
Mathai Mammen (Janssen)
Carl Peck (NDA Partners)
Alfred Sandrock (Biogen)
George Scangos (Vir)
Kuldev Singh (Stanford)
Marc Tessier-Lavigne (Stanford)
Andrew Weber (GlaxoSmithKline)
Anne Wojcicki (23andMe)
Joseph Wu (Stanford)

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Information and Registration
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