ANTICIPATING FUTURE INTERVENTIONS:

DISCOVERING NEW TARGETS, INTEGRATIVE BIOMARKERS, AND BEYOND
Dear Colleagues,

Welcome to Arizona!

On behalf of the American Society of Clinical Psychopharmacology (ASCP) I am pleased to welcome you to our very exciting annual meeting. I want to thank both the Program Committee and the Steering Committee for their wonderful work putting together an extraordinary meeting. Our meeting includes not only a very stimulating Latin American Satellite Symposia but also the 24th iteration of our very successful New Investigators’ Program. Our meeting has something for everyone. There are sessions that discuss innovation across not only syndrome-states and the life cycle but also in terms of public-private partnerships, teaching, as well as public health and dissemination research. And just when you think there could be nothing more to entice you out of the Arizona sun, there are talks on the challenges posed by medical marijuana, to the role of technology in research and clinical practice to a session that describes how you can learn “new tricks” by studying an old medication, Lithium. There is, of course, the very important and unique Regulatory Session that traditionally serves as the closing session of our meeting.

The ASCP is committed to finding and testing new therapies for our patients. We want to advance not only the field of psychopharmacology but treatment research in general. Many advances first presented at our annual meeting over the years have become mainstays not only in our treatment of serious mental disorders but in the way we design and conduct our clinical trials. I am sure that we will see presentations and posters at this meeting that will become important methods for caring for our patients in the future.

Our society has taken on a new responsibility in the past year and is actively becoming one of the places that legislators, federal agencies, and other groups interested in public policy turn to for advice on therapy and research questions about brain diseases. I want to thank Alan Gelenberg who was the initial co-chair of this working group. We have now formed a standing liaison committee to address these issues as they arise. As the experts in the field, we have a responsibility to our patients and society to weigh in on these important issues. Please contact the ASCP Executive Office if you have an interest in this or any other ASCP activities or committees.

I hope you have a wonderful time learning, presenting, and seeing old and new friends at our annual meeting.

Mark Rapaport, M.D.
President
American Society of Clinical Psychopharmacology
Welcome to the ASCP Annual Meeting

On behalf of the ASCP Annual Meeting Steering and Program Committees, we are delighted to welcome you to the ASCP Annual Meeting. The ASCP is committed to continue building on the past success of NCDEU with program innovation while preserving the rich history of this meeting. Below are some of the highlights of the 2016 meeting.

The annual meeting brings together academic investigators, industry scientists, U.S. and international regulators, National Institutes of Health (NIH) and other professionals who work in drug development and clinical trials.

- **2016 Program Highlights**
  - **Monday, May 30th**
    - Latin America Psychopharmacology Update
  - **Tuesday, May 31st**
    - Conference Opening
    - Pharma Pipeline: 8 presentations of Phase 1 and Phase 2 developments
    - Individual Research Reports
  - **Wednesday, June 1st**
    - 17th Annual Fun Run/Walk
    - Regulatory Plenary: *Evolving Views on Pseudospecifcity and Comparing Drug and Device Regulatory Pathways*
    - ASCP Lifetime Awardee Talk
    - Poster Session I
    - ASCP Reception
  - **Thursday, June 2nd**
    - Keynote Plenary Session: *Beyond Single Magic Bullets: True Innovation in Neuropsychiatric Conditions*
    - NIH Institute Directors Plenary
    - Poster Session II
    - Clinical Updates in Pharmacology
    - Workshops
  - **Friday, June 3rd**
    - Regulatory Wrap-Up Plenary with FDA regulators
  - **Throughout the meeting**
    - NIMH, NIDA, and NCCIH panels
  - **The New Investigator Program**
    - A closed workshop for 18 New Investigators and informal breakfast sessions.
  - **Workshops**: 2 hour intensive interactive sessions focused on problems and solutions
    - Wednesday and Thursday Afternoons
  - **“Clinical Track”** – sessions focused on topics of immediate clinical relevance

- **Organization**
  - The meeting is sponsored by the American Society for Clinical Psychopharmacology (ASCP).
  - The Steering Committee organizes the meeting.
  - The Program Committee evaluates submitted proposals and develops program innovations.
  - NIH collaborations:
    - NIMH - National Institute of Mental Health
    - NIDA - National Institute of Drug Abuse
    - NIAAA - National Institute on Alcohol Abuse and Alcoholism
    - NINDS - National Institute of Neurological Disorders and Strokes
    - CSR – Center for Scientific Review
    - NCATS - National Center for Advancing Translational Sciences
  - Regulatory agency collaborations:
    - Food and Drug Administration (FDA)
    - Center for Devices and Radiological Health (CDRH)
  - Parthenon Management Group organizes the ASCP Annual Meeting.

- **And remember**
  - The Fun Run/Walk is Wednesday, June 1st at 6:30 a.m. All are welcome to join!
  - The Opening Reception is Wednesday, June 1st from 6:15 p.m. – 7:15 p.m.

The ASCP Annual Meeting is an opportunity for education and networking. We welcome your suggestions to make the event even better. Seek us out during the meeting or provide your views by completing the evaluation form.

Best Regards,

Husseini Manji, M.D., FRCPC
Steering Committee Co-Chair

Michael E. Thase, M.D.
Steering Committee Co-Chair

Alan Gelenberg, M.D.
Program Committee Co-Chair

Holly A. Swartz, M.D.
Program Committee Co-Chair
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DISCLOSURES FOR ALL ASCP PRESENTERS CAN BE VIEWED AT WWW.ASCPMEETING.ORG
Recipient of the Donald Klein Lifetime Achievement Award

John Davis, M.D.
UIC Psychiatric Institute / University of Il at Chicago

At the time Dr. Davis entered Psychiatry, drugs to treat serious mental illness had been discovered but were not recognized by academic psychiatrists who believed that both treatment and research should be exclusively psychoanalytic, based on individual Freudian case studies. Dr. Davis was one of the early psycho-pharmacologists who transformed psychiatry into a science through introduction of a variety of research methodologies, such as random assignment controlled studies, the development of lithium as an anti-manic agent (Bunney et al., 1968, Am J Psychiatry, 1968; Colburn RW et al., 1967 Nature), the application of pharmacology to understand the mechanisms of how drugs produce improvement, the development of biologically-based theories of mental illness, the study of the clinical pharmacology of psychotropic drugs (such as drug metabolism or drug-drug interactions) and the pooled analysis of clinical trial results.

Dr. Davis performed the first meta-analysis (a statistical technique for combining evidence from different studies) in psychiatry; the second in general medicine, showing that ‘at maintenance’ antipsychotic treatment prevents relapse. The resulting paper (Davis JM, 1975, Am J Psychiatry) was an ISI classic, one of the most cited papers in the year--evidence that the paper had an impact on the field. In 1976, part 2 of this meta-analysis (Davis JM, 1976, Am J Psychiatry) was published showing that maintenance-lithium reduced relapses in bipolar disease and antidepressants did so in depression. Meta-analysis has become a common form of data synthesis particularly after the Cochrane collaboration systematized it, in the 1990s. Drs. Stefan Leucht and Davis have recently updated these analyses in several papers in Lancet (Leucht et al., 2008, 2013), the American Journal of Psychiatry (2009) and other journals, which were highly cited papers and are now called “Hot New Papers” rather than “ISI Science Citation Classics.”
Recipient of the Paul Wender Best Paper in the Journal of Clinical Psychiatry Award

Lee Baer, Ph.D.
Massachusetts General Hospital

Nominated for: *Prevalence and Impact of Obsessive-Compulsive Symptoms in Depression: A STAR*D Report*
Lee Baer, Madhukar H. Trivedi, Ilana Huz, A. John Rush, Stephen R. Wisniewski, and Maurizio Fava

Lee Baer, Ph.D. is an internationally-known clinician and researcher in obsessive-compulsive disorders and their overlap with other Axis I and Axis II disorders. He cofounded the Massachusetts General Hospital OCD Program in 1986, and the McLean OCD Institute in 1997. He co-authored the first comprehensive textbook on OCD, and has written two books on OCD for the general public. Dr. Baer led a team that developed the HANDS depression scale used by National Depression Screening Day for the past quarter century to screen hundreds of thousands of employees, students, and other individuals for major depression. Dr. Baer served for 25 years as Associate Chief of Psychology in the MGH Department of Psychiatry, and is currently senior biostatistician in the MGH Depression Research and Clinical Program, and Professor of Psychiatry, Part-Time in the Harvard Medical School Department of Psychiatry.
New Investigator Awardees

Greta Bushnell, B.S.
University of North Carolina at Chapel Hill

Daniel Eskenazi, M.D., Ph.D.
New York State Psychiatric Institute, Columbia University

Britta Galling, M.D.
The Zucker Hillside Hospital

Carolina Haass-Koffler, Pharm.D.
Brown University

Amy Hilty, M.D.
Otsuka Pharmaceuticals

Yusuke Iwata, M.D.
CAMH Toronto

Manish Jha, M.D.
UT Southwestern

Katelyn Keyloun, Pharm.D.
University of Washington

Luca Lavagnino, M.D.
McGovern Medical School at the University of Texas Health Science Center at Houston

Katarzyna Liwski, B.S., M.P.H.
LECOM

Brian Mickey, M.D., Ph.D.
University of Utah School of Medicine

Dahlia Mukherjee, Ph.D.
Penn State Milton S. Hershey Medical Center

Malik Nassan, M.D.
Mayo Clinic

Primavera Spagnolo, M.D., Ph.D.
National Institute on Alcohol Abuse & Alcoholism, National Institutes of Health

Sunny Tang, M.D.
University of Pennsylvania

Saulo Tractenberg, M.D.
Pontifical Catholic University of Rio Grande do Sul

Hanjing Emily Wu, M.D.
University of Texas Health Science Center at Houston

Xuefeng Zhang, M.D.
Baylor College of Medicine
Regulatory Plenary
Wednesday, June 1st from 8:30 a.m. – 10:00 a.m.

Tiffany Farchione, M.D.
Food and Drug Administration

Dr. Farchione received her medical degree from Wayne State University in Detroit, Michigan, and completed adult residency and child & adolescent fellowship training at the University of Pittsburgh’s Western Psychiatric Institute and Clinic. Dr. Farchione is board certified in both general and child & adolescent psychiatry. Prior to joining FDA in 2010, Dr. Farchione was affiliated with the University of Pittsburgh Medical Center, and was on the faculty of the University of Pittsburgh.

As the Deputy Director in the Division of Psychiatry Products at FDA, Dr. Farchione is involved in the oversight of new drug review for all psychiatric drug development activities conducted under INDs, and the review of all NDAs and supplements for new psychiatric drug claims.

William Heetderks, M.D., Ph.D.
Division of Neurological and Physical Medicine Devices

Dr. Heetderks is the Deputy Director for Clinical Activities in the Division of Neurological and Physical Medicine Devices (DNMPMD) at the Center for Devices and Radiological Health, FDA. This division reviews a wide variety of devices related to neurological disease and rehabilitation including neurosurgical devices, neurostimulation devices, neurodiagnostic devices, and physical medicine and rehabilitation devices. Before coming to the FDA he was the Director of Extramural Science Programs at the National Institute of Biomedical Imaging and Bioengineering, NIH. There he directed broad research programs in medical imaging and bioengineering research and research training.

He received his Ph.D. in bioengineering from the University of Michigan, his M.D. from the University of Miami and is boarded in Internal Medicine. Earlier in his career he served for fifteen years, eventually as Director in the Neural Prosthesis Program, NINDS, a research program focused on development of enabling technologies and devices for rehabilitation of neurological disability including spinal cord injury, deafness, and blindness.
Main Plenary: Beyond Single Magic Bullets: True Innovation in Neuropsychiatric Conditions
Thursday, June 2nd from 8:15 a.m. - 9:45 a.m.

Husseini Manji, M.D.
Johnson & Johnson Pharmaceutical Research & Development

Husseini K. Manji, M.D., FRCPC is the Global Therapeutic Head for Neuroscience at Janssen Research & Development, LLC., a division of Johnson & Johnson. Previously, he was Chief, Laboratory of Molecular Pathophysiology & Experimental Therapeutics, NIH, and Director of the NIH Mood and Anxiety Disorders Program. Dr. Manji received his B.S. and M.D. from the University of British Columbia. He completed fellowship training at the NIMH and completed additional training in cellular and molecular biology. His research has focused on investigation of disease-and treatment-induced changes in gene and protein networks that regulate synaptic and neural plasticity. His work has led to investigation of novel therapeutics for patients with refractory neuropsychiatric illnesses. Dr. Manji has also been involved in medical and postgraduate neuroscience education and has published extensively on the molecular and cellular neurobiology of neuropsychiatric disorders and the development of novel therapeutics. Dr. Manji has received numerous distinguished scientific and academic awards, including the NIMH Director’s Career Award for Significant Scientific Achievement, and was inducted into the U.S. Institute of Medicine of the National Academies in 2008. He has served as Chair of the American College of Neuropsychopharmacology, is a Counselor to the Society of Biological Psychiatry and serves on a variety of editorial boards of scholarly journals. He holds voluntary leadership positions in many organizations devoted to advancement of neuroscience and advocacy for people with neuropsychiatric illnesses. He has been a member of the Howard Hughes Medical Institute and NIH Research Scholars Program Advisory Committee.
Featured Speakers

Vaibhav Narayan, Ph.D.
Johnson & Johnson

Vaibhav is currently Head of Neuroscience Integrated Solutions, Informatics and Disease Interception at Janssen Neuroscience. The Neuroscience Therapeutic Area at Janssen is pioneering a more personalized and holistic approach to therapeutic intervention that goes ‘beyond the pill’, to offer data-driven and science-based ‘integrated solutions’ for preventing, diagnosing, treating, and monitoring CNS diseases. Vaibhav’s work is currently focused on utilizing state-of-the-art informatics methods for developing markers for early diagnosis, disease progression, drug response and treatment monitoring in Alzheimer’s and Mood; and to develop novel ‘point-of-need’ tools and technologies for management of adherence and prediction of relapse in patients with Schizophrenia.

Vaibhav joined Johnson and Johnson from Eli Lilly and Co., where he headed the Discovery and Biomedical Infomatics group to enable Lilly’s translational, biomarker, and tailored therapeutics strategies. Prior to Lilly, Vaibhav held multiple leadership roles in various small to mid-size biotech companies such as Celera Genomics, CuraGen Corporation and Vertex pharmaceuticals during which time he participated in multiple landmark ‘Big Biology’ projects, including sequencing, assembly and analysis of the human genome (Human Genome Project) and elucidation of the first complete protein-protein interaction map of a whole organism.

Vaibhav obtained his Ph.D. from Yale University jointly from the Departments of Chemistry, and Molecular Biophysics & Biochemistry in 1998, and an Executive MBA from Kellogg School of Management, Northwestern University in 2009.

Tanzeem Choudhury, Ph.D.
Cornell University / HealthRhythms

Tanzeem Choudhury is an associate professor in Computing and Information Sciences at Cornell University and a co-founder and CTO of HealthRhythms. At Cornell, she directs the People-Aware Computing group, which works on inventing the future of technology-assisted wellbeing. Tanzeem received her Ph.D. from the Media Laboratory at MIT. Tanzeem was awarded the MIT Technology Review TR35 award, NSF CAREER award and a TED Fellowship. For more information about her research please visit http://pac.cs.cornell.edu. Follow the group’s work on twitter @pac_cornell.
Barbara Sahakian, Ph.D.
University of Cambridge

Barbara J. Sahakian is Professor of Clinical Neuropsychology at the University of Cambridge Department of Psychiatry and Behavioural and Clinical Neuroscience Institute. She is also an Honorary Clinical Psychologist at Addenbrooke’s Hospital, Cambridge. She holds a Ph.D. and a D.Sc. from the University of Cambridge. She is a Fellow of the Academy of Medical Sciences and a Past-President of the International Neuroethics Society and the British Association for Psychopharmacology. In 2016, she was recipient of the Robert Sommer Award. Sahakian is also a Member of the International Expert Jury for the 2017 Else Kröner-Fresenius-Stiftung Prize. She is a member of ACNP, CINP Council and ECNP Review Board. She is co-author of ‘Bad Moves: How Decision Making Goes Wrong and the Ethics of Smart Drugs’ (Oxford University Press, 2013) and co-editor of The Oxford Handbook of Neuroethics (OUP, 2011).

Sahakian has an international reputation in the fields of psychopharmacology, neuropsychology, neuropsychiatry, neuroimaging and neuroethics. She is perhaps best known for her work on ‘hot’ and ‘cold’ cognitive deficits in depression and early detection and early treatment with cholinesterase inhibitors in Alzheimer’s disease. She has over 400 publications in high impact scientific journals. The ISI Web of Science database credits her with a Hirsch (h) index of 102, with some publications having over 300 citations. Sahakian co-invented the neuropsychological CANTAB tests. She serves as a Senior Consultant to Cambridge Cognition, a University of Cambridge spin-out that provides CANTAB (www.cantab.com). She is also a Consultant for Peak (Brainbow) (https://itunes.apple.com/gb/app/peak-brain-training/id806223188?mt=8). There is now the Cambridge University & Peak Advanced Training Plan. Sahakian has contributed to Neuroscience and Mental Health Government Policy and has spoken on resilience, brain health, neuroscience and mental health at the World Economic Forum, Davos, 2014. She was also a finalist for a World Technology Award 2014 under the category of ‘Health and Medicine’. She is a member of the World Economic Forum Global Agenda Council on Brain Research.
Institute Directors Plenary
Thursday, June 2nd from 10:00 a.m. – 11:00 a.m.

Michael E. Thase, M.D.
Perelman School of Medicine at the University of Pennsylvania

Michael E. Thase, M.D., joined the faculty of the University of Pennsylvania School of Medicine in January 2007, as Professor of Psychiatry after more than 27 years at the University of Pittsburgh Medical Center and the Western Psychiatric Institute and Clinic. Dr. Thase’s research focuses on the assessment and treatment of mood disorders, including studies of the differential therapeutics of both depression and bipolar affective disorder. A 1979 graduate of the Ohio State University College of Medicine, Dr. Thase is a Distinguished Fellow of the American Psychiatric Association, a Founding Fellow of the Academy of Cognitive Therapy, a member of the Board of Directors for the American Society of Clinical Psychopharmacology, and Vice Chairman of the Scientific Advisory Board of the National Depression and Bipolar Support Alliance. Dr. Thase has been elected to the membership of the American College of Psychiatrists and the American College of Neuropsychopharmacology. Dr. Thase has authored or co-authored more than 500 scientific articles and book chapters, as well as 15 books.
George Koob, Ph.D.
National Institute on Alcohol Abuse and Alcoholism

George F. Koob, is Director of the National Institute on Alcohol Abuse and Alcoholism (NIAAA) as of January 27, 2014. As NIAAA Director, Dr. Koob oversees a wide range of alcohol-related research, including genetics, neuroscience, epidemiology, prevention, and treatment.

As an authority on alcoholism, drug addiction and stress, he has contributed to our understanding of the neurocircuity associated with the acute reinforcing effects of alcohol and drugs of abuse and the neuroadaptations of the reward and stress circuits associated with the transition to dependence. Dr. Koob has published over 650 peer reviewed papers and several books including the “Neurobiology of Addiction,” a comprehensive treatise on emerging research in the field, and a textbook for upper division undergraduates and graduate students called “Drugs, Addiction and the Brain.” He has mentored 11 Ph.D. students and over 80 post-doctoral fellows.

He received his Ph.D. in Behavioral Physiology from Johns Hopkins University in 1972. He spent much of his early career at the Scripps Research Institute as the Director of the Alcohol Research Center, and as Professor and Chair of the Scripps’ Committee on the Neurobiology of Addictive Disorders. He has also served as a researcher in the Department of Neurophysiology at the Walter Reed Army Institute of Research and the Arthur Vining Davis Center for Behavioral Neurobiology at the Salk Institute for Biological Studies.

Amir Tamiz, Ph.D.
National Institute of Neurological Disorders and Stroke

Dr. Tamiz is a Program Director at the National Institute of Neurological Disorders and Stroke (NINDS), Office of Translational Research (OTR) who oversees NIH Blueprint Neurotherapeutics network (BPN) and Innovation Grants to Nurture Initial Translational Efforts (IGNITE).

Prior to joining NIH in 2012, Dr. Tamiz had held scientific and management positions in research and development of therapeutic programs at Corvas International (acquired by Dendreon), CovX (now part of Pfizer), and Alba Therapeutics. Dr. Tamiz received his Ph.D. at University of Oregon and conducted postdoctoral research at the Department of Neuroscience at Georgetown University Medical Center.
Christopher P. Austin, M.D.
National Center for Advancing Translational Sciences

Christopher P. Austin, M.D., is Director of the National Center for Advancing Translational Sciences (NCATS) at the U.S. National Institutes of Health (NIH). Austin leads the Center’s work to improve the translation of observations in the laboratory, clinic and community into interventions that reach and benefit patients—from diagnostics and therapeutics to medical procedures and behavioral changes. Under his direction, NCATS researchers and collaborators are developing new technologies, resources and collaborative research models; demonstrating their usefulness; and disseminating the data, analysis and methodologies for use by the worldwide research community.

Austin’s career has spanned the spectrum of translational research, in the public and private sectors. Austin joined NIH in 2002 as the senior advisor to the director for translational research at the National Human Genome Research Institute, where he was responsible for conceptualizing and implementing research programs to derive scientific insights and therapeutic benefit from the newly completed Human Genome Project. While at NHGRI, he founded and directed the NIH Chemical Genomics Center, Therapeutics for Rare and Neglected Diseases program, Toxicology in the 21st Century initiative, and NIH Center for Translational Therapeutics. Upon creation of NCATS in 2011, he became the Inaugural Director of the NCATS Division of Pre-Clinical Innovation, and was appointed NCATS director in 2012. Prior to joining NIH, Austin worked at the pharmaceutical company Merck, where he directed programs on genome-based discovery of novel targets and drugs, with a particular focus on schizophrenia and Alzheimer’s disease.

Austin is trained as a clinician and geneticist. He trained in internal medicine and neurology at the Massachusetts General Hospital in Boston, and practiced medicine in academic and community hospital settings as well as in urban primary care and in rural Alaska and Africa. He completed a research fellowship in developmental neurogenetics at Harvard, studying genetic and environmental influences on stem cell fate determination. Austin earned an M.D. from Harvard Medical School and A.B. summa cum laude in biology from Princeton University.
Featured Speakers

Ivan Montoya, M.D., MPH
National Institutes for Neurological Disorders and Stroke / National Institutes of Health

Dr. Montoya is the Deputy Director of the Division of Therapeutics and Medical Consequences (DTMC) of the National Institute on Drug Abuse (NIDA). He received an M.D. from the University of Antioquia (Colombia), a Master’s in Public Health from The Johns Hopkins School of Public Health, and completed residency training in Psychiatry at the University of Antioquia and the University of Maryland Hospital (Baltimore). He has been a Fulbright Fellow at The Johns Hopkins School of Public Health, Visiting Foreign Fellow at the Intramural Research Program of NIDA, Director of the Practice Research Network of the American Psychiatric Association. He has published extensively in the areas of etiology, prevention, treatment (pharmacological and non-pharmacological), and medical consequences of drug abuse.

Richard K. Nakamura, M.D.
Center for Scientific Review

Dr. Richard K. Nakamura is the Director of the Center for Scientific Review. In that capacity, he leads the review of grant applications of the National Institutes of Health. Dr. Nakamura received his Bachelor of Arts in Psychology from Earlham College and his Ph.D. in Psychology from State University of New York (Stony Brook, NY). He was with the National Institute of Mental Health from 1976 to 2011. In 2001, he received the NIH-Asian/Pacific American Organization (APAO) Outstanding Achievement Award for Administrative Work. In 2002, Dr. Nakamura was elected by the American Association for the Advancement of Science (AAAS) to the status of AAAS Fellow. Also in 2002, Dr. Nakamura was awarded the Presidential Rank Award for outstanding leadership. In 2004 and 2005 respectively, he received leadership awards from the Federation of Behavioral Psychological and Cognitive Sciences, and from the International Society for Behavioral Neuroscience. In 2009, he was awarded the NIH Director’s Award for Outstanding Administration.
Sarah Lisanby, M.D.
Division of Translational Research
National Institute of Mental Health

Sarah Hollingsworth Lisanby, M.D. is the director of Division of Translational Research at the National Institute of Mental Health (NIMH). As director for the Division of Translational Research, Dr. Lisanby oversees a research funding portfolio of approximately $400 million and help set a national agenda for research on mental illness. She also works with Dr. Carlos Zarate and colleagues in the Division of Intramural Research Programs, creating an important bridge between the Institute’s extramural and intramural research efforts.

Dr. Lisanby is one of the leading researchers in the area of neuromodulatory interventions for treating major depression, serving as a principal investigator on studies that range from basic research through clinical trials. Additionally, she is a prolific author with approximately 200 scientific articles and book chapters, and she has also received national and international recognition.

Dr. Lisanby’s prodigious research life has been matched by extensive service to NIMH and beyond. She has been a member of the NIMH Board of Scientific Counselors since 2013, and has chaired or been a member of a variety of NIH Study Sections since 2004. Dr. Lisanby also serves on the FDA Neurological Devices Advisory Panel, is on five editorial boards, and has held key leadership positions with numerous professional associations, including Chair of the American Psychiatric Association Task Force to Revise the Practice on Electroconvulsive Therapy (ECT).

Regulatory Wrap-Up
Friday, June 3rd from 10:15 a.m. – 11:45 a.m.

Tiffany Farchione, M.D.
Food and Drug Administration
See previous bio

William Heetderks, M.D., Ph.D.
Division of Neurological and Physical Medicine Devices
See previous bio
Acknowledgements

Steering Committee Chairs

Husseini Manji, M.D., FRCPC
Michael E. Thase, M.D.

Program Committee Chairs

Alan Gelenberg, M.D.
Holly A. Swartz, M.D.

New Investigator Award Program Chairs

Mark H. Rapaport, M.D.
Christopher Sarampote, Ph.D.
Acknowledgements

Steering Committee Members

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Otsuka Pharmaceuticals

★ Lori Davis, M.D.
Tuscaloosa VA Medical Center

Elliot Ehrich, M.D.
Alkermes Pharmaceuticals

★ Tiffany Farchione, M.D.
US Food and Drug Administration

Alan Gelenberg, M.D.
Journal of Clinical Psychiatry

David J. Kupfer, M.D.
University of Pittsburgh School of Medicine

Thomas P. Laughren, M.D.
Food and Drug Administration

Raye Litten, Ph.D.
National Institute of Alcohol Abuse and Alcoholism

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Joseph Palumbo, M.D.
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Luca Pani, M.D.
AIFA The Italian Medicines Agency

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National Institute on Drug Abuse

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University of Pittsburgh School of Medicine

★ New Investigator Alumni  ▲ Representing ASCP CME Committee
Acknowledgements

Steering Committee Members (continued)

★ Madhukar Trivedi, M.D.
University of Texas Southwestern Medical Center

Ben Vitiello, M.D.
National Institute of Mental Health

Program Committee

Scott Aaronson, M.D.
Sheppard Pratt Health Systems

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University of Texas Health Center, San Antonio

Leslie Citrome, M.D., M.P.H.
New York Medical College

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University of Virginia

★ Crystal Clark, M.D.
Northwestern University

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University of North Texas Health Science Center

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Massachusetts General Hospital

★ Christoph U. Correll, M.D.
The Zucker Hillside Hospital

Lynn Crismon, Pharm.D.
The University of Texas at Austin

★ Eden Evins, M.D.
Harvard Medical School

★ Tiffany Farchione, M.D.
Food and Drug Administration

Maurizio Fava, M.D.
Massachusetts General Hospital

▲★ Bradley Gaynes, M.D.
University of North Carolina

★ New Investigator Alumni  ▲ Representing ASCP CME Committee
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Ahmad Hameed, M.D.
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Ebrahim Haroon, M.D.
Emory University School of Medicine

Richard Keefe, Ph.D.
Duke University Medical Center

Terence Ketter, M.D.
Stanford University School of Medicine

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Stephen Marder, M.D.
Semel Institute, UCLA

Tim Mariano, M.D., Ph.D.
Brigham and Women’s Faulkner Hospital

Barbara Milrod, M.D.
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New Investigator Alumni  Representing ASCP CME Committee
Program Committee (continued)

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University of Washington

Benedetto Vitiello, M.D.
National Institute of Mental Health

★ Carlos A. Zarate, M.D.
National Institute of Mental Health

★ Jianping Zhang, M.D., Ph.D.
The Zucker Hillside Hospital

New Investigator Award Program Committee

Mark H. Rapaport, M.D., Co-chair
Emory University School of Medicine (ASCP Board Member)

Christopher Sarampote, Ph.D., Co-chair
National Institute on Mental Health

★ Christoph U. Correll, M.D.
The Zucker Hillside Hospital

★ Lori Davis, M.D.
Tuscaloosa VA Medical Center

★ Tiffany Farchione, M.D.
Food and Drug Administration

Lindsey Grandison, Ph.D.
National Institute on Alcohol Abuse and Alcoholism

Bruce Kinon, M.D.
Lundbeck

Ivan Montoya, M.D., M.P.H.
National Institute on Drug Abuse

★ Katharine Phillips, M.D.
Rhode Island Hospital / Brown University

Nina R. Schooler, Ph.D.
State University of New York, Downstate Medical Center

★ Holly A. Swartz, M.D.
University of Pittsburgh School of Medicine

★ New Investigator Alumni  ★ Representing ASCP CME Committee
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Emory University School of Medicine

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★ Christoph U. Correll, M.D.
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Charles E. Schmidt College of Medicine, Florida Atlantic University

Katharine Phillips, M.D.
Rhode Island Hospital / Brown University

★ Madhukar Trivedi, M.D.
UT Southwestern Medical Center

Sidney Zisook, M.D.
University of California, San Diego

★ New Investigator Alumni ▲ Representing ASCP CME Committee
Meeting Services

Registration Desk Hours:
Monday 10:00 a.m. – 5:00 p.m.
Tuesday 7:30 a.m. – 6:00 p.m.
Wednesday 7:30 a.m. – 6:45 p.m.
Thursday 7:30 a.m. – 6:00 p.m.
Friday 7:30 a.m. – 12:00 p.m.
*The registration/meeting information desk is located in the East Foyer, outside of the Princess Ballroom.

The ASCP Speaker Ready Room is open on the below dates and times for presenters to upload slides. The meeting organizers ask that all speakers upload their slides 24 hours prior to their scheduled presentation time.

The Speaker Ready Room is located in the Bourbon 11 meeting room.
Monday 12:00 p.m. – 5:00 p.m.
Tuesday 8:00 a.m. – 6:00 p.m.
Wednesday - Thursday 7:30 a.m. – 6:30 p.m.
Friday 7:30 a.m. – 12:00 p.m.

Americans with Disabilities Act - It is the policy of ASCP not to discriminate against any person on the basis of disabilities. If you feel you need services or auxiliary aids mentioned in this act in order to fully participate in this continuing education activity, please call the Executive Office at 615-649-3085 or send an email to info@ascpp.org.

Discounts – All ASCP Annual Meeting attendees who booked their room at the Fairmont Scottsdale Princess through the ASCP meeting website will have the following resort amenities complimentary:
- Complimentary internet in the guest rooms
- Fitness Center admittance

ASCP Membership - If you would like to join ASCP, you can log onto our website at www.ascpp.org and register online. Membership applications are available at the ASCP membership booth, located next to the meeting registration desk in the East Princess Foyer. You may also contact the ASCP office at 615-649-3085 for more information.
Disclosures are available for all ASCP Annual Meeting presenters online at www.ASCPMeeting.org.

Continuing Education Credits are available for physicians, pharmacists and psychologists. Applications for credit must be completed online with the meeting evaluation survey. The survey may be completed during or after the conference at www.ASCPMeeting.org. **Surveys for continuing education credit must be submitted no later than June 29, 2016.** There is a $40.00 administrative fee for CME/CE applications. It is the policy of the ASCP to require disclosure of financial relationships from individuals in a position to control the content of a CME activity; to identify and resolve conflicts of interest related to those relationships; and to make disclosure information available to the audience prior to the CME activity. Presenters are required to disclose discussions of unlabeled/unapproved uses of drugs or devices during their presentations.

**Physicians/Nurse Practitioners**

The American Society of Clinical Psychopharmacology designates this live meeting for a maximum of 24.25 **AMA PRA Category 1 Credit(s)**™. Physicians should claim only the credit commensurate with the extent of their participation in the activity.

**Pharmacists**

USF Health is accredited by the Accreditation Council for Pharmacy Education as a provider of continuing pharmacy education. This knowledge-based program has been approved for 24.25 contact hours. Universal program number is as follows: 0230-999-16-008-L01-P.

To receive continuing education credit, a pharmacist must attend the accredited sessions, actively participate in questions and answers and must return the program evaluation instrument. In order to receive full credit, registrants must arrive no later than 10 minutes after the start of the meeting and must attend the entire meeting.
USF Health is approved by the American Psychological Association to sponsor continuing education for psychologists. USF Health maintains responsibility for this program and its content. This activity has been approved for 24.25 CE credits. Full attendance of the live activity is required. Partial credit will not be awarded.

All participants who request continuing education credits by June 29, 2016, should expect to receive their statement of credits via email in July.

The Meeting Evaluation Survey will be available at www.ASCPMeeting.org. We encourage all registrants to complete the evaluation. Attendees requesting CME or CE credits must complete the survey in order to obtain credits. There is a $40.00 administrative fee for CME/CE applications. Your candid input on the 2016 meeting is appreciated as we strive to improve the meeting each year.

ASCP would like to acknowledge the generosity of the following companies whose unrestricted educational grants have contributed to the overall quality of this meeting:

- Alkermes, Inc.
- Otsuka America Pharmaceutical, Inc.

2017 ASCP Annual Meeting – Save the Date

The 2017 ASCP Annual Meeting will take place May 29 – June 2, 2017 at the Loews Miami Beach Hotel in Miami, Florida. Details regarding abstract submission for the 2017 Meeting will be released in September, 2016.
AT-A-GLANCE

Monday, May 30, 2016

8:30 a.m. – 4:15 p.m.  NIA Workshop (Invitation Only)
                        Palomino 8 & 9

10:00 a.m. – 5:00 p.m.  Registration
                        Princess East Foyer

12:00 p.m. – 5:00 p.m.  Speaker Ready Room
                        Bourbon 11

12:00 p.m. – 2:00 p.m.  Latin America Satellite Symposia: An Update on Biomarkers and Clinical Outcomes in Psychiatry
                        Salon I

2:00 p.m. – 4:00 p.m.  Latin America Satellite Symposia: New Therapeutic Approaches for Psychiatric Disorders*
                        Salon I

4:00 p.m. – 5:00 p.m.  ASCP Curriculum Committee Meeting (Invitation Only)
                        Bourbon 9

*of special interest to clinicians
FULL SCHEDULE

Monday, May 30, 2016

8:30 a.m. – 4:15 p.m.  NIA Workshop (Invitation Only)
Palomino 8 & 9

Co-Chairs:  Mark H. Rapaport, M.D., Emory University School of Medicine
Christopher Sarampote, Ph.D., National Institute of Mental Health

The ASCP Annual Meeting offers a special program for New Investigators in an effort to promote the education and training of junior investigators in psychopharmacology. Established investigators were asked to nominate individuals who may be interested in a research career in psychopharmacology for this special program. These nominees submitted an abstract describing their current research or area of research interest, a letter of recommendation from their chair or mentor, a career statement and a curriculum vitae. The selection of awardees was based upon the scientific merit of the abstract, the level of training of the nominee and a committee of internal and external reviewers’ assessment of the relative value of the specialized program to each applicant at this time in his/her career. The awardees will participate in this special educational workshop and present their posters during the scheduled poster sessions and orals during the Individual Research Reports on Tuesday. In addition, they will receive a travel expense award and a certificate acknowledging their participation in the program at an award ceremony on Tuesday evening, May 31, 2016. This year's 18 New Investigator awardees are indicated with a ribbon in the poster section of this program; they will also be notated with a ribbon icon in the program book.

Faculty

Christoph U. Correll, M.D.
The Zucker Hillside Hospital

Lori Davis, M.D.
Tuscaloosa VA Medical Center

Lindsey Grandison, Ph.D.
National Institute on Alcohol Abuse and Alcoholism

Bruce Kinon, M.D.
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<tr>
<th>Time</th>
<th>Event</th>
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</table>
| 10:00 a.m. – 5:00 p.m. | Registration Open  
   *Princess East Foyer*  |
| 12:00 p.m. – 5:00 p.m. | Speaker Ready Room  
   *Bourbon 11*  |
| 12:00 p.m. – 2:00 p.m. | Latin America Satellite Symposia: An Update on Biomarkers and Clinical Outcomes in Psychiatry  
   *Salon I*  |

**Chair:** Mauricio Tohen, University of New Mexico

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<tr>
<th>Time</th>
<th>Event</th>
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| 12:00 p.m. – 12:20 p.m. | Small Non-coding RNA as Predictors and Mediators of Antidepressant Response  
   Gustavo Turecki, McGill University  |
| 12:20 p.m. – 12:40 p.m. | Outcome in First Episode Non-affective Psychosis  
   Mauricio Tohen, University of New Mexico  |
| 12:40 p.m. – 1:00 p.m. | Temporal Lobe Epilepsy: Where Neurology Meets Immunopsychiatry  
   Antonio Teixeira, University of Texas Health Science Center at Houston  |
| 1:00 p.m. – 1:10 p.m. | Discussion  |
| 1:10 p.m. – 1:30 p.m. | Neurobiology of Alcohol Use Disorders  
   Nancy DiazGranados, National Institute on Alcohol Abuse and Alcoholism  |
| 1:30 p.m. – 1:50 p.m. | Paediatric Bipolar Disorder – Brain Mechanisms, Early Detection and Prospects for New Interventions  
   Jair Soares, University of Texas School of Medicine at Houston  |
| 1:50 p.m. – 2:00 p.m. | Discussion/Break  |
2:00 p.m. – 4:00 p.m.  Latin America Satellite Symposia: New Therapeutic Approaches for Psychiatric Disorders*  
Salon I

**Chair:** Rodrigo Machado-Vieira, NIMH, NIH

2:00 p.m. – 2:20 p.m.  *Early Life Stress in Mood Disorders: HPA Axis Response to GR and MR Agonists and Antagonists*  
Mario Jurujena, King’s College London/University of Sao Paulo

2:20 p.m. – 2:40 p.m.  *Translating Neurotrophic and Plasticity Pathways into New Treatments for Mood Disorders*  
Rodrigo Machado-Vieira, NIMH, NIH

2:40 p.m. – 3:00 p.m.  *New Treatments for Alcoholism: a Focus on the Gut-Liver-Brain Axis*  
Lorenzo Leggio, NIAAA, NIDA, NIH

3:00 p.m. – 3:10 p.m.  Discussion

3:10 p.m. – 3:30 p.m.  *Promising Medications to Treat Substance Use Disorders*  
Ivan Montoya, NIDA, NIH

3:30 p.m. – 3:50 p.m.  *New Glutamate Modulators for Mood Disorders*  
Carlos A. Zarate, NIMH, NIH

3:50 p.m. – 4:00 p.m.  Final Remarks

4:00 p.m. – 5:00 p.m.  ASCP Curriculum Committee Meeting (Invitation Only)  
*Bourbon 9*

*of special interest to clinicians*
### AT-A-GLANCE

**Tuesday, May 31, 2016**

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<tr>
<th>Time</th>
<th>Event</th>
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<tbody>
<tr>
<td>6:30 a.m. – 8:30 a.m</td>
<td><strong>ASCP Board Meeting (Invitation Only)</strong>&lt;br&gt;Moor 1</td>
</tr>
<tr>
<td>7:30 a.m. – 6:00 p.m</td>
<td><strong>Registration</strong>&lt;br&gt;Princess East Foyer</td>
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<tr>
<td>7:30 a.m. – 8:30 a.m</td>
<td><strong>NIA Breakfast Roundtable (Invitation Only)</strong>&lt;br&gt;Castile 1</td>
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<tr>
<td>7:30 a.m. – 6:00 p.m</td>
<td><strong>Speaker Ready Room</strong>&lt;br&gt;Bourbon 11</td>
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<tr>
<td>8:30 a.m. – 9:00 a.m</td>
<td><strong>Conference Opening</strong>&lt;br&gt;Princess Ballroom A-E</td>
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<tr>
<td>9:00 a.m. – 10:30 a.m</td>
<td><strong>Panel Sessions</strong></td>
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<tr>
<td>ADHD in Adults with a Focus on Assessment, Comorbidity and Associated Features*</td>
<td>Salon F</td>
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<tr>
<td>Public-private Partnerships to Develop Medications for Alcohol Use Disorder: Recent Successes and New Opportunities for Collaboration with NIAAA</td>
<td>Salon G</td>
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<tr>
<td>Mobile Health: Real-time Monitoring in Bipolar Disorder and Addictions*</td>
<td>Salon H</td>
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<tr>
<td>Pathway to Treatment of Cognitive Impairment in Depression</td>
<td>Salon I</td>
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<tr>
<td>10:30 a.m. – 10:45 a.m</td>
<td><strong>Break</strong>&lt;br&gt;Princess East Foyer</td>
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*of special interest to clinicians
**Panel Sessions**

<table>
<thead>
<tr>
<th>Time</th>
<th>Session</th>
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| 10:45 a.m. – 12:15 p.m. | **State Versus Trait Anhedonia: A Neurobiological Link Between Depression, ADHD and Substance Abuse**
|               | **Unraveling the Complexities of Psychotropic Prescribing during Pregnancy**
|               | **Frontiers of Pharmacotherapy for Autism Spectrum Disorders**
|               | **Biomarker Based Clinical Trials in Drug Development**
|               | Salon F | Salon G | Salon H | Salon I |

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<tr>
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<tr>
<td>12:15 p.m. – 2:00 p.m.</td>
<td><strong>Lunch On Own</strong></td>
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<tr>
<th>Time</th>
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| 2:00 p.m. – 4:00 p.m. | **Pharmaceutical Pipelines**
|               | Princess Ballroom A-E                                                  |

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<tr>
<th>Time</th>
<th>Session</th>
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| 4:00 p.m. – 4:15 p.m. | **Break**
|               | Princess East Foyer                                                    |

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<thead>
<tr>
<th>Time</th>
<th>Session</th>
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</table>
| 4:15 p.m. – 5:30 p.m. | **Individual Research Reports**
| Substance Use Disorders, Treatments for Alzheimer’s Disease, and Speeding Drug Discovery |
| Improving Assessment and Clinical Trial Methodology |
| Mood Disorders: Targets, Approaches, and Outcomes |
| Advances in Schizophrenia* |
| Salon F | Salon G | Salon H | Salon I |

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<tr>
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| 6:15 p.m. – 7:45 p.m. | **New Investigator Awards Ceremony (Invitation Only)**
|               | Arabian A/B                                                            |

*of special interest to clinicians*
FULL SCHEDULE

Tuesday, May 31, 2016

6:30 a.m. – 8:30 a.m.  ASCP Board Meeting (Invitation Only)
  Moor 1

7:30 a.m. – 8:30 a.m.  NIA Breakfast Roundtable (Invitation Only)
  Castile 1

7:30 a.m. – 6:00 p.m.  Registration
  Princess East Foyer

7:30 a.m. – 9:00 a.m.  Morning Break
  Princess East Foyer

7:30 a.m. – 6:00 p.m.  Speaker Ready Room
  Bourbon 11

8:30 a.m. – 9:00 a.m.  Conference Opening
  Princess Ballroom A-E

Panel Sessions

9:00 a.m. – 10:30 a.m.  ADHD in Adults with a Focus on Assessment,
  Comorbidity and Associated Features*
  Salon F

Chair: Frederick Reimherr, University of Utah School of Medicine
Discussant: Mark Stein, University of Washington

9:00 a.m. – 9:10 a.m.  Introduction

9:10 a.m. – 9:30 a.m.  ADHD in Adults with a Focus on Comorbidity and
  Associated Features
  Calvin Sumner, NCS Pearson - Clinical Assessment

9:30 a.m. – 9:50 a.m.  Personality Disorder in Adult ADHD
  Thomas Gift, University of Rochester, Rochester, NY

*of special interest to clinicians
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| 9:00 a.m. – 9:30 a.m. | ADHD in Adults with a Focus on Assessment, Comorbidity and Associated Features*  
(Salon F) |
| 9:50 a.m. – 10:10 a.m. | Violence in Adult ADHD - The Role of ADHD on Type of Aggression  
Florence Philipp-Wiegmann, Institute for Forensic Psychology and Psychiatry, Saarland Universität, Homburg/Saar, Germany |
| 10:10 a.m. – 10:30 a.m. | Discussion |
| 9:00 a.m. – 10:30 a.m. | Public-private Partnerships to Develop Medications for Alcohol Use Disorder: Recent Successes and New Opportunities for Collaboration with NIAAA  
(Salon G)  
Chair: Raye Litten, NIAAA |
| 9:00 a.m. – 9:10 a.m. | Introduction |
| 9:10 a.m. – 9:30 a.m. | Infrastructure to Facilitate Drug Development: Public-Private Partnerships  
Raye Litten, NIAAA |
| 9:30 a.m. – 9:50 a.m. | Moderators of Varenicline Treatment Effects in a Double-blind, Placebo-controlled Trial for Alcohol Dependence: An Exploratory Analysis  
Daniel Falk, NIAAA/NIH |
| 9:50 a.m. – 10:10 a.m. | A Double-blind, Placebo-controlled Trial Assessing the Efficacy of ABT-436 (V1b antagonist) for Alcohol Dependence  
Megan Ryan, NIAAA |
| 10:10 a.m. – 10:30 a.m. | Discussion |

*of special interest to clinicians
9:00 a.m. – 10:30 a.m. Mobile Health: Real-time Monitoring in Bipolar Disorder and Addictions*

Salon H

**Chair:** Erika Saunders, Penn State College of Medicine, Penn State Milton S. Hershey Medical Center

**Discussant:** Terence Ketter, Stanford University School of Medicine

9:00 a.m. – 9:10 a.m. **Introduction**

9:10 a.m. – 9:30 a.m. **Detection of Mood States from Acoustic Features Acquired Using Smartphones**

Melvin McInnis, University of Michigan Medical School

9:30 a.m. – 9:50 a.m. **Daily Mood Monitoring of Symptoms Using Smartphones in Bipolar Disorder: Feasibility of Ecological Momentary Assessment**

Erika Saunders, Penn State College of Medicine, Penn State Milton S. Hershey Medical Center

9:50 a.m. – 10:10 a.m. **Understanding the Relationships Between Affect and Craving in Early Abstinence: Ecological Momentary Assessment of Patients in Treatment for Prescription Opioid Dependence**

Scott Bunce, Penn State Milton S. Hershey Medical Center, Penn State College of Medicine

10:10 a.m. – 10:30 a.m. **Discussion**

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9:00 a.m. – 10:30 a.m. Pathway to Treatment of Cognitive Impairment in Depression

Salon I

**Chair:** Richard Keefe, Duke University Medical Center

**Discussant:** Tiffany Farchione, US Food and Drug Administration

9:00 a.m. – 9:10 a.m. **Introduction**

9:10 a.m. – 9:30 a.m. **Cognitive Impairment and Disability in Major Depression: What is the Nature of the Problem?**

Philip Harvey, Miller School of Medicine, University of Miami

*of special interest to clinicians*
<table>
<thead>
<tr>
<th>Time</th>
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</table>
| **9:00 a.m. – 10:30 a.m. (continued)** | Pathway to Treatment of Cognitive Impairment in Depression  
**Salon I** |
| 9:30 a.m. – 9:50 a.m. | Review of Treatment Studies to Date – What Do We Need?  
Maurizio Fava, Massachusetts General Hospital |
| 9:50 a.m. – 10:10 a.m. | Methods and Design of Trials for Treatment of Cognitive Impairment of MDD  
Richard Keefe, Duke University Medical Center |
| 10:10 a.m. – 10:30 a.m. | Discussion                                                                                       |
| 10:30 a.m. – 10:45 a.m. | Break  
*Princess East Foyer* |
| **Panel Sessions** |                                                                                                  |
| 10:45 a.m. – 12:15 p.m. | State Versus Trait Anhedonia: A Neurobiological Link Between Depression, ADHD and Substance Abuse  
**Salon F** |
| **Chair:** Martin Katzman, START Clinic for Mood and Anxiety Disorders  
**Discussant:** Larry Klassen, Eden Mental Health Centre |                                                                                                  |
| 10:45 a.m. – 10:55 a.m. | Introduction                                                                                     |
| 10:55 a.m. – 11:15 a.m. | Understanding the Biological Basis of Co-occurring Depression and Attention-Deficit/Hyperactivity Disorder  
Irvin Epstein, University of Toronto |
| 11:15 a.m. – 11:35 a.m. | Anhedonia: A Predictor of Cognitive Dysfunction, Attention-Deficit/Hyperactivity Disorder and Treatment Outcome in a Subset of Depressed Patients  
Tia Sternat, START Clinic for Mood & Anxiety Disorders |
10:45 a.m. – 12:15 p.m. (continued)  
**State Versus Trait Anhedonia: A Neurobiological Link Between Depression, ADHD and Substance Abuse**  
*Salon F*

11:35 a.m. – 11:55 a.m.  
**The Neurobiology of Cognitive Dysfunction and Anhedonia: A Common Link Between Psychiatric Disorders and the Effects on Treatment Selection**  
Martin Katzman, START Clinic for Mood and Anxiety Disorders

11:55 a.m. – 12:15 p.m.  
**Discussion**

10:45 a.m. – 12:15 p.m.  
**Unraveling the Complexities of Psychotropic Prescribing During Pregnancy*  
*of special interest to clinicians**  
*Salon G*

**Chair:** Lee Cohen, Massachusetts General Hospital  
**Co-Chair:** Marlene Freeman, Massachusetts General Hospital  
**Discussant:** Alan Gelenberg, Journal of Clinical Psychiatry

10:45 a.m. – 10:55 a.m.  
**Introduction**

10:55 a.m. – 11:15 a.m.  
**Second Generation Antipsychotics in Reproductive Age Women: Current Results from the National Pregnancy Registry for Atypical Antipsychotics**  
Lee Cohen, Massachusetts General Hospital

11:15 a.m. – 11:35 a.m.  
**Baseline BMI and Gestational Weight Gain in Women with Psychiatric Illness: Impact of Psychotropic Use and Clinical Implications**  
Marlene Freeman, Massachusetts General Hospital

11:35 a.m. – 11:55 a.m.  
**Obstetrical, Pregnancy, and Socioeconomic Predictors for Postpartum Psychiatric Disorders**  
Samantha Meltzer-Brody, University of North Carolina at Chapel Hill

11:55 a.m. – 12:15 p.m.  
**Discussion**
10:45 a.m. – 12:15 p.m.  Frontiers of Pharmacotherapy for Autism Spectrum Disorders*

*of special interest to clinicians

Salon H

Chair: Stephen Zukin, Johns Hopkins University School of Medicine
Discussant: Jeremy Veenstra-Vanderweele, Columbia University & New York State Psychiatric Institute

10:45 a.m. – 10:55 a.m.  Introduction

10:55 a.m. – 11:15 a.m.  Endpoint Selection and Stratification Variables for Clinical Trials in Autism: Lessons from the Arbaclofen Program
Paul Wang, Autism Speaks

11:15 a.m. – 11:35 a.m.  Designing Clinical Trials to Enhance Functioning in Developmental Disorders: Moving from Single Dose Studies of Oxytocin to Sustained Treatment Trial in Autism
Linmarie Sikich, Duke University Medical Center

11:35 a.m. – 11:55 a.m.  Opportunities and Challenges in Autism Drug Development: The Industry Perspective
Federico Bolognani, Roche Innovation Center Basel / F. Hoffmann-La Roche, Ltd.

11:55 a.m. – 12:15 p.m.  Discussion

10:45 a.m. – 12:15 p.m.  Biomarker Based Clinical Trials in Drug Development

Salon I

Chair: Madhukar Trivedi, UT Southwestern Medical Center
Discussant: William Potter, National Institute of Mental Health

10:45 a.m. – 10:55 a.m.  Introduction

10:55 a.m. – 11:15 a.m.  Towards Imaging Biomarkers for Treatment Selection in Major Depressive Disorder
Helen Mayberg, Emory University School of Medicine
10:45 a.m. – 12:15 p.m. Biomarker Based Clinical Trials in Drug Development
*Salon I*

11:15 a.m. – 11:35 a.m. Rationale for the Design of the NIMH FAST-MAS Biomarker-based Clinical Trial
Andrew Krystal, Duke Clinical Research Institute and Department of Psychiatry and Behavioral Sciences Duke University School of Medicine

11:35 a.m. – 11:55 a.m. Interleukin-6 Inhibition as a Potential Treatment Strategy for Depression: A Proof of Concept Study with Sirukumab in Depressed Patients with Suboptimal Response to Antidepressants and Elevated Peripheral Inflammatory Markers
Giacomo Salvadore, Janssen Pharmaceuticals

11:55 a.m. – 12:15 p.m. Discussion

12:15 p.m. – 2:00 p.m. Lunch - On Own

2:00 p.m. – 4:00 p.m. Pharmaceutical Pipeline Presentations
*Princess Ballroom A-E*

Chair: Michael E. Thase, Perelman School of Medicine at the University of Pennsylvania

2:00 p.m. – 2:15 p.m. Centanafadine Sr (CTN-SR) Demonstrates Brain Occupancy at Norepinephrine Transporter (NET), Serotonin Transporter (SERT) and Dopamine Transporter (DAT) Using Single Photon Emission Tomography (SPECT) in Healthy Volunteers (HVs)
Anthony McKinney, Bio-Pharma

2:15 p.m. – 2:30 p.m. Dasotraline: A Novel Drug Candidate Being Evaluated for the Treatment of Attention-Deficit/Hyperactivity Disorder and Binge Eating Disorder
Robert Goldman, Sunovion
<table>
<thead>
<tr>
<th>Time</th>
<th>Presentation Title</th>
<th>Speaker(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2:30 p.m. – 2:45 p.m.</td>
<td>A Phase 1 Single- and Multiple-rising Dose Study of the Safety &amp; Pk of EMB-001, a Potential Treatment for Substance Use Disorders, with Exploratory Efficacy Measures in Tobacco Use Disorder</td>
<td>Michael Detke, Indiana University School of Medicine</td>
</tr>
<tr>
<td>2:45 p.m. – 3:00 p.m.</td>
<td>A Randomized Placebo-controlled Multicenter Trial of a Low-dose Bedtime Sublingual Formulation of Cyclobenzaprine (TNX-102 SL*) for the Treatment of Military-related PTSD</td>
<td>Gregory Sullivan, Tonix Pharmaceuticals, Inc.</td>
</tr>
<tr>
<td>3:00 p.m. – 3:15 p.m.</td>
<td>PeRSEVERe: A Study of Esketamine for the Rapid Reduction of the Symptoms of Major Depressive Disorder, including Suicidal Ideation, in Subjects Assessed to be at Imminent Risk for Suicide</td>
<td>Carla Canuso, Janssen Research &amp; Development</td>
</tr>
<tr>
<td>3:15 p.m. – 3:30 p.m.</td>
<td>SRX246: A First-in-class Vasopressin 1a Receptor Antagonist in Phase Ii Trials for Mood and Behavioral Disorders</td>
<td>Michael Brownstein, Azevan Pharmaceuticals, Inc.</td>
</tr>
<tr>
<td>3:30 p.m. – 3:45 p.m.</td>
<td>Drug Development Strategies for Schizophrenia Using a Novel PDE10A Inhibitor: TAK-063</td>
<td>Tom Macek, Takeda Development Center - Americas</td>
</tr>
<tr>
<td>3:45 p.m. – 4:00 p.m.</td>
<td>Phase 2 Study of Bremelanotide in Premenopausal Women with Female Sexual Dysfunctions: Responder Analyses based on Minimum Clinically Important Differences Derived from Receiver Operating Characteristic Curves</td>
<td>Stanley Althof, Case Western Reserve University School of Medicine</td>
</tr>
</tbody>
</table>
4:00 p.m. – 4:15 p.m. Break
Princeess East Foyer

Individual Research Reports

4:15 p.m. – 5:30 p.m. Substance Use Disorders, Treatments for Alzheimer’s Disease, and Speeding Drug Discovery
Salon F

Chair: Timothy Mariano, Brigham and Women’s Faulkner Hospital

4:15 p.m. – 4:30 p.m. Pharmacogenetics of Dopamine Beta Hydroxylase in Cocaine Dependence Therapy with Doxazosin
Xuefeng Zhang, Baylor College of Medicine

4:30 p.m. – 4:45 p.m. Ghrelin as a Novel Possible Target to Treat Alcohol Craving and Role of Endogenous Hormones Serum Levels as a Biomarker
Carolina Haass-Koffler, Brown University

4:45 p.m. – 5:00 p.m. Impact of Concomitant Antidepressant Use on Drinking and Mood Outcomes in Bipolar Alcoholics: Results from a Randomized Controlled Trial of Lamotrigine
Bryan Tolliver, Medical University of South Carolina

5:00 p.m. – 5:15 p.m. The Coding and Noncoding Transcriptional Landscape of Neurons and Glia in Vivo
Adarsh Reddy, Trinitas Regional Medical Center

5:15 p.m. – 5:30 p.m. Effect of Renal and Hepatic Impairment on the Pharmacokinetics of Encenicline
Gordon Loewen, FORUM Pharmaceuticals

= New Investigator Awardee
<table>
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<tr>
<th>Time</th>
<th>Topic</th>
<th>Speaker and Affiliation</th>
</tr>
</thead>
</table>
| 4:15 p.m. – 5:30 p.m. | Improving Assessment and Clinical Trial Methodology | *Salon G*  
Chair: Ahmad Hameed, Penn State College of Medicine  
4:15 p.m. – 4:30 p.m. | How Do Key Co-primary Measures of Functional Capacity Predict Real World Function in Schizophrenia? | Richard Keefe, Duke University Medical Center |
|              |                                                                     |                                                                 |
|              | Do Suicidality Phenomena Follow a Linear or a Non-linear Progression over Time? | David V. Sheehan, University of South Florida College of Medicine |
|              |                                                                     |                                                                 |
|              | Is it Possible to Reduce Biases and Inaccuracies in Large Scale Clinical Trials? | Mark Opler, ProPhase, LLC                                         |
|              |                                                                     |                                                                 |
| 5:00 p.m. – 5:15 p.m. | Nine Problems in the Descriptions of the Psychiatric Inclusion and Exclusion Criteria in Publications of Antidepressant Efficacy Trials | Mark Zimmerman, Brown University |
|              |                                                                     |                                                                 |
| 5:15 p.m. – 5:30 p.m. | Validation of the Tablet-based Brief Assessment of Cognition (BAC App) for Schizophrenia | Brian Saxby, NeuroCog Trials                                    |
| 4:15 p.m. – 5:30 p.m. | Mood Disorders: Targets, Approaches, and Outcomes | *Salon H*  
Chair: Crystal Clark, Northwestern University  
4:15 p.m. – 4:30 p.m. | MoodNetwork: An Innovative Approach to Patient-centered Care | Andrew Nierenberg, Massachusetts General Hospital |
4:15 p.m. – 5:30 p.m. Mood Disorders: Targets, Approaches, and Outcomes

Salon H

4:30 p.m. – 4:45 p.m. A Genome Wide Association Study Implicates Gabaergic Neurotransmission in Early Onset Bipolar Disorder
Malik Nassan, Mayo Clinic

4:45 p.m. – 5:00 p.m. Early and Sustained Work Productivity Improvement Predicts Subsequent Clinical Course in Major Depressive Disorder
Manish Jha, UT Southwestern

5:00 p.m. – 5:15 p.m. Adherence and Persistence across Antidepressant Therapeutic Classes: A Retrospective Claims Analysis among Insured US Patients with Major Depressive Disorder
Katelyn Keyloun, University of Washington

5:15 p.m. – 5:30 p.m. Vilazodone Inhibiting Pro-inflammatory Gene Expression and Immunologic Activation Compared to Paroxetine in Geriatric Depression
Helen Lavretsky, David Geffen School of Medicine at UCLA

4:15 p.m. – 5:30 p.m. Advances in Schizophrenia*

Salon I

Chair: Mohammed Ahmed, University of California, San Diego and Veterans Medical Center, San Diego

4:15 p.m. – 4:30 p.m. Prevalence, Risk Factors and Outcome of Metabolic Syndrome in Veterans with Serious Mental Illness
Stanley Caroff, Perelman School of Medicine University of Pennsylvania

= New Investigator Awardee  *of special interest to clinicians
<table>
<thead>
<tr>
<th>Time</th>
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</table>
| 4:15 p.m. – 5:30 p.m. (continued) | Advances in Schizophrenia*  
Salon I                                                                                     |
| 4:30 p.m. – 4:45 p.m.   | Relative Efficacy and Safety of Individual Second-generation Antipsychotics in Treating First Episode Psychosis: A Systematic Review and Meta-analysis  
Jianping Zhang, The Zucker Hillside Hospital                                                   |
| 4:45 p.m. – 5:00 p.m.   | Worldwide Clozapine Therapeutic Drug Monitoring (TDM) from Capillary Blood, using the Dried Blood Spot (DBS) Technique: A Major Step Forward in Adequate Dosing and Treatment of Therapy-resistant Schizophrenia  
Dan Cohen, Mental Health Organization North-Holland North                                          |
| 5:00 p.m. – 5:15 p.m.   | Successfully Overcoming Clozapine Under prescription: The Dutch Approach as an Evidence-based Model for Scaling-up Innovation and Good Practice Abroad  
Dan Cohen, Mental Health Organization North-Holland North                                          |
| 5:15 p.m. – 5:30 p.m.   | Threshold of Dopamine D2/3 Receptor Occupancy for Hyperprolactinemia in Older Patients with Schizophrenia  
Yusuke Iwata, CAMH Toronto                                                                 |
| 6:15 p.m. – 7:45 p.m.   | New Investigator Awards Ceremony  
Arabian A/B                                                                                   |

= New Investigator Awardeee  *of special interest to clinicians
Wednesday, June 1, 2016

6:30 a.m. – 8:00 a.m.  17th Annual ASCP Fun Run/Walk
Open to ALL Attendees!
Palomino Conference Center North Drive

7:30 a.m. – 6:45 p.m.  Registration
Princess East Foyer

7:30 a.m. – 8:30 a.m.  NIA Breakfast Roundtable (Invitation Only)
Castile 1

7:30 a.m. – 6:30 p.m.  Speaker Ready Room
Bourbon 11

8:30 a.m. – 10:00 a.m.  Regulatory Plenary with FDA: Evolving Views
on Pseudospecificity and Comparing Drug and
Device Regulatory Pathways
Princess Ballroom A-E

10:00 a.m. – 10:15 a.m.  Break
Princess East Foyer

10:15 a.m. – 11:15 a.m.  ASCP Best Paper in Journal of Clinical
Psychopharmacology Award Presentation
and ASCP Lifetime Awardee Talk - Evidence
in Clinical Psychiatry: 60 Years in Psychiatric
Practice and Research
Princess Ballroom A-E

11:15 a.m. – 1:00 p.m.  Poster Session I with Lunch
Palomino 1 - 7
1:00 p.m. – 2:30 p.m. **Panel Sessions**

<table>
<thead>
<tr>
<th>Drinking Change, Consequences, and Biomarkers in Alcohol Clinical Trials: Results from the Alcohol Clinical Trials Initiative (ACTIVE)</th>
<th>Novel Approaches to Treatment-Resistant Depression (TRD)*</th>
<th>Revisiting Experimental and Clinical Therapeutics of Cannabidiol</th>
<th>Lithium: The Old New Wonder Drug*</th>
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<td>Salon F</td>
<td>Salon G</td>
<td>Salon H</td>
<td>Salon I</td>
</tr>
</tbody>
</table>

2:45 p.m. – 3:15 p.m. **ASCP Business Meeting (Members Only)**
Salon I

3:15 p.m. – 3:30 p.m. **Break**
Princess East Foyer

3:30 p.m. – 5:30 p.m. **Workshops**

<table>
<thead>
<tr>
<th>An Integrated Technology Approach for Preventing Relapse in Recently Hospitalized Schizophrenia Patients</th>
<th>Managing the Unique Challenges Associated with Rapid Acting Antidepressant Trials</th>
<th>The Hidden Truth in Psychiatric Trials - Medication Adherence is Highly Variable - Problems, Implications, and Solutions</th>
<th>Is Suicidal Ideation a Symptom or a Syndrome?</th>
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<td>Salon G</td>
<td>Salon H</td>
<td>Salon I</td>
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</table>

5:45 p.m. – 6:15 p.m. **First Timer Reception**
Moor 2

6:15 p.m. – 7:15 p.m. **ASCP Reception**
Princess Ballroom A-E

*of special interest to clinicians*
## FULL SCHEDULE

### Wednesday, June 1, 2016

<table>
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| 6:30 a.m. – 8:00 a.m. | **17th Annual ASCP Fun Run/Walk - Open to ALL Attendees!**  
All participants receive a FREE ASCP T-shirt! |
| 7:30 a.m. – 8:30 a.m. | **NIA Breakfast Roundtable (Invitation Only)**  
*Castile 1* |
| 7:30 a.m. – 9:00 a.m. | **Morning Break**  
*Princess East Foyer* |
| 8:30 a.m. – 10:00 a.m. | **Regulatory Plenary with FDA: Evolving Views on Pseudospecificity and Comparing Drug and Device Regulatory Pathways**  
*Chair & Discussant: Mark H. Rapaport, Emory University School of Medicine* |

- **This year’s plenary session will focus on two topics:**
  1. **The Division of Psychiatry Products’ evolving stance on pseudospecificity.**  
     Historically, the Division has viewed a number of potential labeling claims (e.g., cognitive dysfunction associated with major depressive disorder) to be artificially narrow and, thus, pseudospecific. DPP is now more willing to consider indications that were previously deemed pseudospecific. Dr. Farchione will discuss the Division’s current views, as well as trial design considerations for any development programs designed to evaluate an indication that was once considered pseudospecific.
  2. **A comparison of the regulatory review process for drugs and devices.**  
     Representatives from both the Division of Psychiatry Products in the Center for Drug Evaluation and Research (Dr. Farchione) and the Division of Neurological and Physical Medicine Devices in the Center for Devices and Radiological Health (Dr. Heetderks) will compare and contrast the regulatory requirements and review processes in the two Centers as they related to the treatment of psychiatric illnesses.

### Wednesday, June 1, 2016

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<tr>
<td>8:30 a.m. – 8:40 a.m.</td>
<td><strong>Introduction</strong></td>
</tr>
</tbody>
</table>
| 8:40 a.m. – 9:05 a.m. | **The Division of Psychiatry Products’ Evolving Stance on Pseudospecificity**  
*Tiffany Farchione, US Food and Drug Administration* |
| 9:05 a.m. – 9:30 a.m. | **A Comparison of the Regulatory Review Process for Drugs and Devices**  
*William Heetderks, CDRH, FDA* |
| 9:30 a.m. – 10:00 a.m. | **Discussion and Q&A**                                                                   |
10:00 a.m. – 10:15 a.m.  Break  
*Princess East Foyer*

10:15 a.m. – 11:15 a.m.  ASCP Best Paper in *Journal of Clinical Psychopharmacology* Award Presentation and ASCP Lifetime Awardee Talk - Evidence in Clinical Psychiatry: 60 Years in Psychiatric Practice and Research  
*Princess Ballroom A-E*

**Best Paper Award Presentation to:**  Lee Baer, Massachusetts General Hospital  
Nominated for:  *Prevalence and Impact of Obsessive-Compulsive Symptoms in Depression: A STAR*D Report*

**Lifetime Awardee:**  John Davis, UIC Psychiatric Institute, University of Il at Chicago, Chicago, IL

When I wrote my first scientific papers 60 years ago, academic psychiatry was entirely psychoanalytic. Don Klein and I participated in the development of modern psychopharmacology, writing Diagnosis and the Drug treatment of Mental Disorders, which was both research and clinically based. Over the last 40 years, the clinical side has atrophied, and research is less clinically relevant. I will place modern research in historical prospective, with a focus interpretation of research to make it more clinically relevant. I will start with the application of pharmacology to understand mechanisms of how drugs produce improvement, the biologically-based theories of mental illness, and the clinical pharmacology of psychotropic drugs (such as drug metabolism or drug-drug interactions), updating this with current work in molecular biology. I performed, in 1975 the first meta-analysis in psychiatry; the second in general medicine, showing maintenance treatment prevents relapse, update this with recants network meta-analyses. I will summarize clinical evidence, provide prospective, but the focus is on how to apply clinical studies to individual patients, to tell sense from nonsense, and to the balance of benefit to risk. I will present the efficacy of psychotropic drugs in the perspective of drug used in internal medicine.

11:15 a.m. – 1:00 p.m.  Poster Session I with Lunch  
*Palomino 1 - 7*  
*See pages 73 - 82 for a complete listing of posters*
## Panel Sessions

**1:00 p.m. – 2:30 p.m. Drinking Change, Consequences, and Biomarkers in Alcohol Clinical Trials: Results from the Alcohol Clinical Trials Initiative (ACTIVE)**

*Salon F*

**Chair:** Raymond Anton, Medical University of South Carolina  
**Discussant:** Raye Litten, NIAAA

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<tr>
<th>Time</th>
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<tr>
<td>1:00 p.m. – 1:10 p.m.</td>
<td>Introduction</td>
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</tr>
<tr>
<td>1:10 p.m. – 1:25 p.m.</td>
<td>FDA Guidance on Developing Drugs for the Treatment of Alcoholism</td>
<td>Rachel Skeete, Division of Anesthesia, Analgesia, and Addiction Products, Center for Drug Evaluation and Research, US Food and Drug Administration</td>
</tr>
<tr>
<td>1:25 p.m. – 1:40 p.m.</td>
<td>The Relationship of who Drinking Risk Categories to Alcohol Consequences: Consistency in Population Studies and Clinical Trials</td>
<td>Katie Witkeiwitz, University of New Mexico</td>
</tr>
<tr>
<td>1:40 p.m. – 1:55 p.m.</td>
<td>Novel Efficacy Endpoints Based on Shifts in the World Health Organization (WHO) Risk Levels of Drinking: Treatment Effects in Alcohol Pharmacotherapy Trials</td>
<td>Daniel Falk, NIAAA/NIH</td>
</tr>
<tr>
<td>1:55 p.m. – 2:10 p.m.</td>
<td>Use of Alcohol Consumption Biomarkers in Clinical Trials</td>
<td>Raymond Anton, Medical University of South Carolina</td>
</tr>
<tr>
<td>2:10 p.m. – 2:30 p.m.</td>
<td>Discussion</td>
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</tr>
</tbody>
</table>
### Novel Approaches to Treatment-Resistant Depression (TRD*)

**Salon G**

**Chair:** Maurizio Fava, Massachusetts General Hospital  
**Co-Chair:** George Papakostas, Massachusetts General Hospital  
**Discussant:** Carlos Zarate, National Institute of Mental Health

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<tr>
<td>1:00 p.m. – 1:10 p.m.</td>
<td>Introduction</td>
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</tbody>
</table>
| 1:10 p.m. – 1:25 p.m. | NIMH Approach to Fostering New Rapidly Acting Therapies for TRD  
                        Mi Hillefors, National Institute of Mental Health |
| 1:25 p.m. – 1:40 p.m. | Low-field Magnetic Stimulation in TRD: The Rapid Study in TRD  
                        Maurizio Fava, Massachusetts General Hospital |
| 1:40 p.m. – 1:55 p.m. | The Use of Alternative and Complementary Therapies in TRD  
                        Mark H. Rapaport, Emory University School of Medicine |
| 1:55 p.m. – 2:10 p.m. | Novel Non-monoamine-based Drug Therapies for TRD  
                        George Papakostas, Massachusetts General Hospital |
| 2:10 p.m. – 2:30 p.m. | Discussion                                                                               |

### Revisiting Experimental and Clinical Therapeutics of Cannabidiol

**Salon H**

**Chair:** Antonio Teixeira, University of Texas Health Science Center at Houston  
**Discussant:** Antonio de Oliveira, Federal University of Minas Gerais

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<thead>
<tr>
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<tr>
<td>1:00 p.m. – 1:10 p.m.</td>
<td>Introduction</td>
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</tbody>
</table>
| 1:10 p.m. – 1:30 p.m. | Potential Role of Cannabidiol in Neurodegenerative Diseases  
                        Antonio Teixeira, University of Texas Health Science Center at Houston |

*of special interest to clinicians*
<table>
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<th>Time</th>
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<tr>
<td>1:00 p.m. – 2:30 p.m. (continued)</td>
<td>Revisiting Experimental and Clinical Therapeutics of Cannabidiol</td>
<td>Salon H</td>
</tr>
<tr>
<td>1:30 p.m. – 1:50 p.m.</td>
<td>Potential Role of Cannabidiol (CBD) on the Mood Disorders Therapeutics</td>
<td>João Luciano de Quevedo, The University of Texas Health Science Center at Houston</td>
</tr>
<tr>
<td>1:50 p.m. – 2:10 p.m.</td>
<td>Cannabidiol and Endocannabinoids as Potential Approaches against Cocaine Neurotoxicity</td>
<td>Fabricio Moreira, Federal University of Minas Gerais</td>
</tr>
<tr>
<td>2:10 p.m. – 2:30 p.m.</td>
<td>Discussion</td>
<td></td>
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<tr>
<td>1:00 p.m. – 2:30 p.m.</td>
<td>Lithium: The Old New Wonder Drug*</td>
<td>Salon I</td>
</tr>
<tr>
<td><strong>Chair:</strong> James Kocsis, New York Presbyterian Hospital</td>
<td><strong>Discussant:</strong> Andrew Nierenberg, Massachusetts General Hospital</td>
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<tr>
<td>1:00 p.m. – 1:10 p.m.</td>
<td>Introduction</td>
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<tr>
<td>1:10 p.m. – 1:25 p.m.</td>
<td>Why, When and How Do I Prescribe Lithium Today?</td>
<td>James Kocsis, New York Presbyterian Hospital</td>
</tr>
<tr>
<td>1:25 p.m. – 1:40 p.m.</td>
<td>Can Lithium Extend the Antidepressant Effects of Ketamine? A Randomized Controlled Trial</td>
<td>James Murrough, Icahn School of Medicine at Mount Sinai</td>
</tr>
<tr>
<td>1:40 p.m. – 1:55 p.m.</td>
<td>Results of Recent Pragmatic Studies of Lithium in Bipolar Disorder</td>
<td>Michael E. Thase, Perelman School of Medicine at the University of Pennsylvania</td>
</tr>
<tr>
<td>1:55 p.m. – 2:10 p.m.</td>
<td>Neuronal Hyperexcitability in a Stem Cell Model of Bipolar Disorder is Reversed by Lithium</td>
<td>John Kelsoe, University of California - San Diego</td>
</tr>
<tr>
<td>2:10 p.m. – 2:30 p.m.</td>
<td>Discussion</td>
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*of special interest to clinicians*
**Workshops**

<table>
<thead>
<tr>
<th>Time</th>
<th>Event</th>
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</table>
| 3:30 p.m. – 5:30 p.m. | **An Integrated Technology Approach for Preventing Relapse in Recently Hospitalized Schizophrenia Patients**  
**Salon F** |

**Chair:** John Kane, The Zucker Hillside Hospital  
**Co-Chair:** Nina R. Schooler, SUNY Downstate Medical Center  
**Discussant:** Donald Goff, NYU Langone Medical Center/Nathan Kline Institute

- **3:30 p.m. – 3:40 p.m.** **Introduction**
- **3:40 p.m. – 4:00 p.m.** **Improving Care and Reducing Costs: Background and Rational**  
  John Kane, The Zucker Hillside Hospital
- **4:00 p.m. – 4:20 p.m.** **Components of the Health Technology Program**  
  Delbert Robinson, Hofstra NS-LIJ School of Medicine
- **4:20 p.m. – 4:40 p.m.** **Baseline Characteristics and Initial Usage Data for Recently Hospitalized Schizophrenia Patients using an Integrated Technology Approach to Prevent Relapse vs Controls Receiving Usual Care**  
  Eric Achtyes, Michigan State University College of Human Medicine
- **4:40 p.m. – 5:00 p.m.** **Patient Experience of Technology Enhanced Treatment: What they liked; Barriers to Use and skills they Acquired**  
  Nina R. Schooler, SUNY Downstate Medical Center
- **5:00 p.m. – 5:30 p.m.** **Discussion**
### Managing the Unique Challenges Associated with Rapid Acting Antidepressant Trials

**Salon G**

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<tr>
<th>Time</th>
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<td>3:30 p.m. – 3:45 p.m.</td>
<td>Introduction</td>
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<tr>
<td>3:45 p.m. – 4:00 p.m.</td>
<td>Challenges of Measuring Rapid Changes in Suicidality: Implications for Clinical Trials</td>
<td>James Murrough, Icahn School of Medicine at Mount Sinai</td>
</tr>
<tr>
<td>4:00 p.m. – 4:15 p.m.</td>
<td>Integrating Human Biomarkers in Trials Involving Interventions with Rapid Antidepressant and Antisuicidal Effects</td>
<td>Carlos A. Zarate, National Institute of Mental Health</td>
</tr>
<tr>
<td>4:15 p.m. – 4:30 p.m.</td>
<td>Assessment of Rapid Improvement of Depression: Challenges Across the Age Spectrum</td>
<td>Jaskaran Singh, Neuroscience TA, Janssen R &amp; D, LLC., Janssen Pharmaceutical Companies of JNJ</td>
</tr>
<tr>
<td>4:30 p.m. – 4:45 p.m.</td>
<td>Study Design Challenges in Developing Rapid Onset Antidepressants</td>
<td>Suresh Durgam, Forest Research Institute, A Subsidiary of Actavis, plc</td>
</tr>
<tr>
<td>4:45 p.m. – 5:00 p.m.</td>
<td>Managing Placebo Response and Patient Expectations in Clinical Trials</td>
<td>Ronald Marcus, Cerecor</td>
</tr>
<tr>
<td>5:00 p.m. – 5:30 p.m.</td>
<td>Discussion</td>
<td></td>
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</tbody>
</table>
### 3:30 p.m. – 5:30 p.m.  The Hidden Truth in Psychiatric Trials - Medication Adherence is Highly Variable - Problems, Implications, and Solutions  
*Salon H*

**Chair:** Daniel Burch, PPD  
**Co-Chair:** Atul Mahabaleshwar, Takeda Global Research & Development

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<td>3:30 p.m. – 3:40 p.m.</td>
<td><strong>Introduction</strong></td>
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</tbody>
</table>
| 3:40 p.m. – 3:55 p.m. | **Evidence and Impact of Non-adherence**                                                   
Daniel Burch, PPD |
| 3:55 p.m. – 4:15 p.m. | **Methodological Approaches to Enhancing Medication Adherence in CNS Clinical Trials**   
Maurizio Fava, Massachusetts General Hospital |
| 4:15 p.m. – 4:30 p.m. | **Brief Overview of Technical Solutions**                                                 
Atul Mahabaleshwar, Takeda Global Research & Development |
| 4:30 p.m. – 4:40 p.m. | **Regulatory View**                                                                       
Tiffany Farchione, FDA |
| 4:40 p.m. – 5:30 p.m. | **Discussion**                                                                            |

### 3:30 p.m. – 5:30 p.m.  Is Suicidal Ideation a Symptom or a Syndrome?  
*Salon I*

**Chair:** Steven Targum, Bracket Global

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<td><strong>Introduction</strong></td>
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</table>
| 3:40 p.m. – 4:10 p.m. | **Suicidal Ideation: One Piece of a Larger Puzzle**                                       
Jill Harkavy-Friedman, American Foundation for Suicide Prevention |
| 4:10 p.m. – 4:40 p.m. | **There are Several Suicidality Disorders**                                               
David V. Sheehan, University of South Florida College of Medicine |
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| 3:30 p.m. – 5:30 p.m. | Is Suicidal Ideation a Symptom or a Syndrome?  
(Salon I)            |
| 4:40 p.m. – 5:10 p.m.  | Characterization of Suicide Ideation in Depressed Patients Identified to be at Risk for Suicide and Ketamine Treatment Response  
Larry Alphs, Janssen |
| 5:10 p.m. – 5:30 p.m.  | Discussion                                                           |
| 5:45 p.m. – 6:15 p.m.  | First Timer Reception  
(Moor 2)                |
| 6:15 p.m. – 7:15 p.m.  | ASCP Reception  
(Princess Ballroom A-E)  |
**AT-A-GLANCE**

**Thursday, June 2, 2016**

7:00 a.m. – 8:30 a.m.  **ASCP Steering Committee Meeting**  
(Invitation Only)  
Moor 1

7:30 a.m. – 8:30 a.m.  **NIA Breakfast Roundtable (Invitation Only)**  
Castile 1

7:30 a.m. – 6:00 p.m.  **Registration**  
Princess East Foyer

7:30 a.m. – 6:30 p.m.  **Speaker Ready Room**  
Bourbon 11

8:15 a.m. – 9:45 a.m.  **Keynote: Beyond Single Magic Bullets: True Innovation in Neuropsychiatric Conditions**  
Princess Ballroom A-E

9:45 a.m. – 10:00 a.m.  **Break**  
Princess East Foyer

10:00 a.m. – 12:00 p.m.  **NIH Institute Directors Plenary**  
Princess Ballroom A-E

12:00 p.m. – 2:00 p.m.  **Poster Session II with Lunch**  
Palomino 1-7

2:00 p.m. – 3:30 p.m.  **Psychopharmacology State-of-the-Art Updates**  
Princess Ballroom A-E

3:30 p.m. – 3:45 p.m.  **Break**  
Princess East Foyer

3:45 p.m. – 5:45 p.m.  **Workshops**

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<tr>
<th>Implementation of Universal Depression Screening and Measurement Based Care in Busy Clinical Practices: Lessons Learned from Project VitalSign6</th>
<th>Medical Marijuana: Promise and Peril*</th>
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3:45 p.m. – 5:45 p.m.  **Special Session: How to Use the Model Psychopharmacology Curriculum in Various Teachings**  
Salon G

*of special interest to clinicians*
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| 7:00 a.m. – 8:30 a.m. | ASCP Steering Committee Meeting (Invitation Only)  
                                          *Moor 1* |
| 7:30 a.m. – 8:30 a.m. | NIA Breakfast Roundtable (Invitation Only)  
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| 7:30 a.m. – 9:00 a.m. | Morning Break  
                                          *Princess East Foyer* |
| 8:15 a.m. – 9:45 a.m. | Keynote Session: *Beyond Single Magic Bullets: True Innovation in Neuropsychiatric Conditions*  
                                          *Princess Ballroom A-E* |

**Chair:** Husseini Manji, M.D., Johnson & Johnson

Worldwide, neuropsychiatric diseases are some of the most debilitating and disabling medical conditions. In addition, because of demographic trends and the lack of effective treatments to modify disease course, the economic and societal burden of these disorders are projected to increase in the coming years. These diseases are caused by dysregulations that span genetic, epigenetic, proteomic, and brain circuitry (connectomics) networks, and they affect complex phenotypes, including memory, cognition, emotion, function, and behavior. Effectively addressing such complex, multi-scale, multi-level disorders will necessarily require multi-pronged interventions that go ‘beyond the pill’ and use newly-developed advances in mobile computing, devices, and computer-based therapies to intercept these diseases in their earliest stages, when they still modifiable, and before irreversible damage has occurred.

Psychiatric diseases are chronic diseases of the young, in that they often take hold in late adolescence/early adulthood and have a subsequent dynamic course characterized by relapses and recurrences of increasing frequency. Each episode of illness inflicts considerable trauma to patients and caregivers, as well as considerable cost to the healthcare system. For example, patients with schizophrenia and a recent history of relapse generate direct medical costs that are three- to five-fold higher than other medical disorders; relapses are also disproportionately associated with negative outcomes such as higher prevalence of substance use disorders and worsening functional status. Furthermore, recent neuroimaging studies have revealed that relapses and
reurrences are not just symptomatic exacerbations of the illness, but actually may cause irreversible atrophic changes in the brain that represent progression of underlying neuropathology. As a result, there is a strong imperative to move from reactive treatment of relapses and recurrences to a more proactive regimen of prediction and preemption. Significant technological advances have been made, particularly with regard to methods capable of remotely capturing rich phenotypic information from streamed data drawn from wearable devices over prolonged observation periods. Such advances provide new ways to monitor long-term outcomes and, via real-time analysis of data streams, to remotely detect changes in clinical status. The reliable identification of such changes and biosignatures will fundamentally re-engineer health care systems to provide more precise care and interventions early in the course of relapse/deterioration, or even before clinical changes are apparent. It should be noted that, while predicting and preempting relapse has immense value, intercepting these diseases in their prodromal stages—prior to conversion to clinical diagnosis—should remain the ultimate goal. This is particularly true of diseases like Alzheimer’s disease wherein overt clinical symptoms typically appear almost a decade before the onset of pathology. Interestingly, computer-based cognitive tests have the potential to convert cognition from a subjective and noisy endpoint to a sensitive and quantitative biomarker that will facilitate early detection of Alzheimer’s disease and aid the development of sensitive efficacy endpoints for early AD clinical trials. In the future, more sensitive measures of functional decline based on wearable technology and data from smart homes and devices (‘internet of things’) will help us track disease course and develop customized interventions (cognitive prosthetics) that increase independence and improve quality of life for Alzheimer’s patients.

Finally, increasing our understanding of the role that dysfunction in circuit dynamics and synaptic plasticity play as causal drivers of cognitive impairment and psychiatric disorders will lead to increased use of device- and computer-based interventions. Indeed, the most effective therapeutic combinations may well be those that meld pharmacological and non-pharmacological approaches to simultaneously target molecular pathology and brain circuitry.

In this session, the speaker will explore—via specific examples—how novel technologies can be harnessed to develop integrated solutions that will help us move from a ‘diagnose and treat’ to a ‘predict and preempt’ paradigm, with the ultimate goal of modifying disease course and improving patient outcomes.
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<td>Introduction</td>
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<tr>
<td>8:25 a.m. – 8:45 a.m.</td>
<td>Data and Informatics Challenges in Moving from a ‘Diagnose and Treat’ to a ‘Predict and Preempt’ Paradigm</td>
<td>Vaibhav Narayan, Ph.D., Johnson &amp; Johnson</td>
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<td>8:45 a.m. – 9:05 a.m.</td>
<td>Improving Brain Health and Trajectories and Outcomes for Neuropsychiatric Disorders using Technologies</td>
<td>Barbara Sahakian, Ph.D., D.Sc., FMEDSci, University of Cambridge</td>
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<tr>
<td>9:05 a.m. – 9:25 a.m.</td>
<td>Sensing Behavioral Symptoms of Mental Health and Delivering Personalized Interventions Using Mobile and Wearable Technologies</td>
<td>Tanzeem Choudhury, Ph.D., Cornell University / HealthRhythms</td>
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<tr>
<td>9:25 a.m. – 9:45 a.m.</td>
<td>Discussion</td>
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<td>9:45 a.m. – 10:00 a.m.</td>
<td>Break</td>
<td>Princess East Foyer</td>
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| 10:00 a.m. – 12:00 p.m. | NIH Institute Directors Plenary | }

**Chair:** Michael E. Thase, M.D., Perelman School of Medicine at the University of Pennsylvania

The NIH Institute Directors session will provide updates on NIAAA, NINDS, NCATS, NIDA, CSR, and NIMH.

- George Koob, Ph.D., NIAAA
- Amir Tamiz, Ph.D., NINDS
- Christopher P. Austin, M.D., NCATS
- Ivan Montoya, M.D., MPH, NIDA
- Richard K. Nakamura, Ph.D., CSR
- Sarah Lisanby, M.D., NIMH
2:00 p.m. – 3:30 p.m.  Psychopharmacology State-of-the-Art Updates

*Princess Ballroom A-E*

**Chair:** Holly A. Swartz, University of Pittsburgh School of Medicine

The purpose of this symposium is to provide an overview of the recent advances in clinical psychopharmacology leading to the development of novel treatments for mood disorders. This session will focus on the following topics:

- Terence Ketter will discuss advances in treatments for Bipolar Disorder.
- Barbara Mason will discuss the evidence base for FDA-approved medications to treat alcohol use disorder, including potential differential treatment effects for males and females.
- Mark Rapaport will provide updates on complementary and alternative approaches to treating mood and anxiety disorders.

2:00 p.m. – 2:20 p.m.  Terrence Ketter, M.D., Stanford University School of Medicine

2:20 p.m. – 2:40 p.m.  Barbara Mason, Ph.D., The Scripps Research Institute

2:40 p.m. – 3:00 p.m.  Mark H. Rapaport, M.D., Emory University School of Medicine

3:00 p.m. – 3:30 p.m.  Discussion

3:30 p.m. – 3:45 p.m.  Break

*Princess East Foyer*
Workshops

3:45 p.m. – 5:45 p.m. Implementation of Universal Depression Screening and Measurement Based Care in Busy Clinical Practices: Lessons Learned from Project VitalSign6

Salon F

Chair: Manish Jha, UT Southwestern Medical Center
Co-Chair: Madhukar Trivedi, UT Southwestern Medical Center
Discussant: Tracy Greer, University of Texas Southwestern Medical Center at Dallas

3:45 p.m. – 3:55 p.m. Introduction

3:55 p.m. – 4:15 p.m. Making Evidence-based Treatment of Depression Easily Accessible: Forging Collaborations with Primary Care Clinics
Madhukar Trivedi, UT Southwestern Medical Center

4:15 p.m. – 4:35 p.m. Enhancing Measurement Based Care: Why Does Functional Recovery Matter?
Tracy Greer, University of Texas Southwestern Medical Center at Dallas

4:35 p.m. – 4:55 p.m. Harnessing Health Information Technology to Implement Measurement Based Care: Demonstration of VitalSign6 Software
Manish Jha, UT Southwestern Medical Center

4:55 p.m. – 5:45 p.m. Discussion

= New Investigator Awardee
### 3:45 p.m. – 5:45 p.m.  Medical Marijuana: Promise and Peril*
*Salon H*

**Chair:** Susan Weiss, National Institute of Health/NIDA  
**Co-Chair:** Steven Gust, NIH

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<td><strong>Introduction</strong></td>
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<td>3:55 p.m. – 4:15 p.m.</td>
<td><strong>Medical Marijuana in 2016: What a Clinician Needs to Know</strong></td>
<td>Kevin Hill, McLean Hospital</td>
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<td>4:15 p.m. – 4:35 p.m.</td>
<td><strong>The Endocannabinoid System as a Target for Therapeutic Drugs</strong></td>
<td>Daniele Piomelli, University of California, Irvine</td>
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<td>4:35 p.m. – 4:55 p.m.</td>
<td><strong>Medical Marijuana for Psychiatric Indications: Is the Cart Before the Horse?</strong></td>
<td>Deepak D’Souza, Yale University School of Medicine &amp; VACHS</td>
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<td>4:55 p.m. – 5:15 p.m.</td>
<td><strong>Medical Marijuana in Canada</strong></td>
<td>Didier Jutras-Aswad, Centre hospitalier de l’Université de Montréal</td>
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<td>5:15 p.m. – 5:45 p.m.</td>
<td><strong>Discussion</strong></td>
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#### Special Session

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**Chair:** Ira Glick, Stanford University School of Medicine

This workshop will focus on how to teach cutting edge clinical psychopharmacology for a) psychiatric residents, b) primary care physicians, and c) medical students in the US and globally. The target audience is program directors, chairs, and teachers of psychopharmacology and psychiatry.

*of special interest to clinicians*
Friday, June 3, 2016

7:30 a.m. – 12:00 p.m.  Registration
Princess East Foyer

7:30 a.m. – 12:00 p.m.  Speaker Ready Room
Bourbon 11

8:30 a.m. – 10:00 a.m.  Panel Sessions

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<td>Psychotherapy, Pharmacotherapy and Devices to Treat Bipolar II Depression: New Evidence*</td>
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<td>Applicability of Industry/Regulatory Antidepressant Clinical Trials to Clinical Practice</td>
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<td>Salon I</td>
<td>The Intersection of Pharmacology and Neuromodulation in Treatment Refractory Mood Disorders*</td>
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10:00 a.m. – 10:15 a.m.  Break
Princess East Foyer

10:15 a.m. – 11:45 a.m.  Regulatory Wrap-Up Plenary
Princess Ballroom A-E

12:00 p.m.  Meeting Adjourns

*of special interest to clinicians
FULL SCHEDULE

Friday, June 3, 2016

7:30 a.m. – 9:00 a.m. Morning Break
   East Princess Foyer

7:30 a.m. – 12:00 p.m. Registration
   Princess East Foyer

7:30 a.m. – 12:00 p.m. Speaker Ready Room
   Bourbon 11

Panel Sessions

8:30 a.m. – 10:00 a.m. Shared Pharmacological Targets for Substance Use and Other Psychiatric Disorders
   Salon F

Chair & Discussant: Ivan Montoya, DHHS/National Institute on Drug Abuse

8:30 a.m. – 8:40 a.m. Introduction

8:40 a.m. – 9:00 a.m. Pimavanserin: A New Treatment for Parkinson’s Psychosis that may have Efficacy as a Treatment for Cocaine Use Disorders
   Jane Acri, NIDA/NIH/DTMC

9:00 a.m. – 9:20 a.m. Testing the Efficacy and Safety of a FAAH-Inhibitor in the Treatment of Cannabis Dependence
   Deepak D’Souza, Yale University School of Medicine & VACHS

9:20 a.m. – 9:40 a.m. Characterization of Agonist-Antagonist Opioid Modulation with ALKS 5461 in Major Depression
   Elliot Ehrich, Alkermes, plc

9:40 a.m. – 10:00 a.m. Discussion
8:30 a.m. – 10:00 a.m.  Psychotherapy, Pharmacotherapy and Devices to Treat Bipolar II Depression: New Evidence*

Salon G

Chair: Holly A. Swartz, University of Pittsburgh School of Medicine
Discussant: Michael E. Thase, Perelman School of Medicine at the University of Pennsylvania

8:30 a.m. – 8:40 a.m.  Introduction

8:40 a.m. – 9:00 a.m.  A Recent Treatment Study Addressing if Bipolar II Response to Treatment Really is the Same as Bipolar I Disorder
Trisha Suppes, Stanford University

9:00 a.m. – 9:20 a.m.  Interpersonal and Social Rhythm Therapy and Quetiapine as Treatments for Bipolar II Depression
Holly A. Swartz, University of Pittsburgh School of Medicine

9:20 a.m. – 9:40 a.m.  Safety and Efficacy of Cranial Electrotherapy Stimulation in Treatment of Bipolar II Depression
Igor Galynker, Icahn School of Medicine at Mount Sinai

9:40 a.m. – 10:00 a.m.  Discussion

8:30 a.m. – 10:00 a.m.  Applicability of Industry/Regulatory Antidepressant Clinical Trials to Clinical Practice

Salon H

Chair: Arifulla Khan, Northwest Clinical Research Center
Discussant: Walter Brown, Alpert Medical School, Brown University

8:30 a.m. – 8:40 a.m.  Introduction

8:40 a.m. – 9:00 a.m.  Do Depressed Patients in Registration Clinical Trials Reflect the “Real World” Practice?
Maurizio Fava, Massachusetts General Hospital

*of special interest to clinicians
8:30 a.m. – 10:00 a.m.  Applicability of Industry/Regulatory Antidepressant Clinical Trials to Clinical Practice

Salon H

9:00 a.m. – 9:20 a.m.  Should the Lack of Generalizability of Antidepressant Efficacy Trials Have Implications for Product Labelling?
Mark Zimmerman, Brown University

9:20 a.m. – 9:40 a.m.  Incidence of Suicidal Behavior among Antidepressant Clinical Trial participants: 1991-2013
Arifulla Khan, Northwest Clinical Research Center

9:40 a.m. – 10:00 a.m.  Discussion

8:30 a.m. – 10:00 a.m.  The Intersection of Pharmacology and Neuromodulation in Treatment Refractory Mood Disorders*

Salon I

Chair: Michael Henry, Massachusetts General Hospital
Discussant: William Potter, National Institute of Mental Health

8:30 a.m. – 8:40 a.m.  Introduction

8:40 a.m. – 9:00 a.m.  The Use of Ketamine in Treatment-Resistant Depression: From Research to Clinical
Cristina Cusin, Massachusetts General Hospital

9:00 a.m. – 9:20 a.m.  The Effects of ECT on the Depressed Brain: A Meta Analysis of Imaging and EEG Studies
Michael Henry, Massachusetts General Hospital

9:20 a.m. – 9:40 a.m.  Circuit-level Mechanism of Action of Neuromodulation Therapies
Joan Camprodon, Harvard Medical School/ Massachusetts General Hospital

9:40 a.m. – 10:00 a.m.  Discussion

*of special interest to clinicians
### Friday, June 3, 2016

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**Chair:** Mark H. Rapaport, Emory University School of Medicine

- Tiffany Farchione, US Food and Drug Administration
- William Heetderks, CDRH, FDA

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Wednesday, June 1, 2016
Poster Session I with Lunch
Palomino 1-7
11:15 a.m. - 1:00 p.m.

1. **HLD200, A Novel Delivery System of Methylphenidate, in Children with Attention-Deficit/Hyperactivity Disorder**
   Floyd Sallee*, University of Cincinnati
   Sharon Wigal, Ann Childress, Mary Ann McDonnell, Scott Kollins, Norberto DeSousa

2. **Centanafadine Sr (CTN-SR) Demonstrates Brain Occupancy at Norepinephrine Transporter (NET), Serotonin Transporter (SERT) and Dopamine Transporter (DAT) Using Single Photon Emission Tomography (SPECT) in Healthy Volunteers (HVs)**
   Anthony McKinney*, Bio-Pharma
   Danna Jennings, Olivier Barret, Gary Wisniewski, Kenneth Marek, Catherine O’Brien, Gary Maier, Connie Reining, Gilles Tamagnan, David Alagille, John Seiby

3. **Dasotraline: A Novel Drug Candidate being Evaluated for the Treatment of Attention-Deficit/Hyperactivity Disorder and Binge Eating Disorder**
   Robert Goldman*, Sunovion
   Nga Tong, Tracy Wetter, Kenneth S. Koblan, Seth C. Hopkins, Antony Loebel

4. **A Randomized Placebo-controlled Multicenter Trial of a Low-dose Bedtime Sublingual Formulation of Cyclobenzaprine (TNX-102 SL*) for the Treatment of Military-related PTSD**
   Gregory Sullivan*, Tonix Pharmaceuticals, Inc.
   Judith Gendreau, R. Michael Gendreau, Amy Schaberg, Bruce Daugherty, Heather Jividen, Ashild Peters, Perry Peters, Seth Lederman

5. **An Open Label Pilot Study of Adjunctive Asenapine for the Treatment of Posttraumatic Stress Disorder**
   Lori Davis*, Veterans Affairs Medical Center
   Patricia Pilkinson, Badari Birur, Seth Norrholm, Felicia Moody

© = New Investigator Awardee  ✨ = Pharmaceutical Pipeline Presentation
6. **Lurasidone in the Treatment of Bipolar Depression: Effect of Baseline Depression Severity on Clinical Outcome**
   Andrei Pikalov*, Sunovion Pharmaceuticals, Inc.
   Joyce Tsai, Josephine Cucchiaro, Antony Loebel

7. **A Retrospective Study of Transcranial Magnetic Stimulation (TMS) in the Treatment of Bipolar Depression**
   Scott Aaronson*, Sheppard Pratt Health System
   Kathy Daddario

8. **Discrepancy Between Subjective and Objective Sleep Parameters in Symptomatic and Euthymic Bipolar Disorder Compared to Healthy Controls**
   Venkatesh Krishnamurthy*, Penn State Milton S. Hershey Medical Center

9. **Treating Pediatric Anxiety: The use of SSRIs and other Prescription Medications**
   Greta Bushnell*, University of North Carolina at Chapel Hill
   Stacie Dusetzina, Scott Compton, Bradley Gaynes, Alan Brookhart, Til Stürmer

10. **Virgil Investigative Study Platform: Improving Signal Detection in Psychiatry Clinical Trials**
    Janet Williams*, MedAvante
    Selam Negash, Briana Webber-Lind, Lisa Stein, Christopher Randolph

11. **Access to Information on Schizophrenia Patients’ Medication Adherence Can Change Prescriber’s Treatment Practices**
    Felicia Forma*, Otsuka Pharmaceutical Development & Commercialization, Inc.
    Jason Shafrin, Suepattra May, Anshu Shrestha, Charles Ruetsch, Nicole Gerlanc, Ainslie Hatch, Darius Lakdawalla, Jean-Pierre Lindenmayer

    David Daniel*, Bracket Global, LLC
    Larry Alphs, Dawn Velligan

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= New Investigator Awardee  ★ = Pharmaceutical Pipeline Presentation
13. A Retrospective Analysis of the Effects of Subject Characteristics on Completion Rates in Phase 1 Studies in Subjects with Stable Schizophrenia
David Krefetz*, PRA Health Sciences
Juliet Brown

14. Reduced Inhibitory Control Mediates the Relationship Between Cortical Thickness in the Right Superior Frontal Gyrus and Body Mass Index
Luca Lavagnino*, McGovern Medical School at the University of Texas Health Science Center at Houston
Benson Mwangi, Isabelle Bauer, Bo Cao, Sudhakar Selvaraj, Alan Prossin, Jair C. Soares

15. Attention-Deficit/Hyperactivity Disorder and Depression: Sequential and Concurrent Disorders
Tia Sternat*, START Clinic for Mood & Anxiety Disorders
Munira Mohamed, Melissa Furtado, Alex Canzonieri, Sara Armata, Catherine Cameron, Irvin Epstein, Martin Katzman

16. Improving Alzheimer's Disease Data Quality Through Video Monitoring
Theresa Shackleford*, ePharmasolutions

17. OPEN BOARD

18. Categorical Improvements in Disease Severity in MDD Patients Treated with Vilazodone: Post Hoc Analysis of 4 Randomized, Placebo-controlled Trials
Suresh Durgam*, Forest Research Institute, A Subsidiary of Actavis, plc
Changzheng Chen, John Edwards, Carl Gommoll, Leslie Citrome

19. Adjunctive Brexpiprazole (OPC-34712) in Patients with MDD and Anxiety Symptoms: Results from Post-hoc Analyses of Two Pivotal Studies
Dusan Kostic, Emmanuelle Weiller, Peter Zhang, Anna Eramo, Ross Baker, Catherine Weiss

20. OPEN BOARD

= New Investigator Awardee  ♦ = Pharmaceutical Pipeline Presentation
21. A Post Hoc Subgroup Analysis of the Impact of Vortioxetine on Functional Capacity, as Measured by UPSA, in MDD Patients with Subjective Cognitive Dysfunction
Richard Keefe*, Duke University Medical Center
William Jacobson, George Nomikos, Elizabeth Merikle, Wei Zhong, Christina Kurre Olsen, Michael Cronquist Christensen

22. PeRSEVERe: A Study of Esketamine for the Rapid Reduction of the Symptoms of Major Depressive Disorder, including Suicidal Ideation, in Subjects Assessed to be at Imminent Risk for Suicide
Carla Canuso*, Janssen Research & Development
Jaskaran Singh, Maggie Fedgchin, Larry Alphs, Rosanne Lane, Christine Pinter, Husseini Manji, Wayne C. Drevets

23. Moderating Factors Affect Signal Detection with MSI-195 vs. Placebo in a Major Depressive Disorder Augmentation Trial
Beth Cameron*, MSI Methylation Sciences, Inc.
Steven Targum, Maurizio Fava, David MacDonald, Ludvina Ferreira

24. Lurasidone for Major Depressive Disorder with Mixed Features: Effect of Concurrent Anxiety Severity
Marlene Freeman*, Massachusetts General Hospital
Joyce Tsai, Yongcai Mao, Daisy Ng-Mak, Andrei Pikalov, Antony Loebel

25. Effect of Adjunctive Brexipiprazole and Adjunctive Aripiprazole on Weight: An Analysis of Long-term Trials in Major Depressive Disorder
Emmanuelle Weiller*, H. Lundbeck A/S
Ruth A. Duffy, Keva K. Gwin, Ross Baker, Catherine Weiss

26. Validation and Determination of Minimal Clinically Important Differences and Treatment Response for the UCSD Performance-based Skills Assessment (UPSA) in Major Depressive Disorder
Philip Harvey*, Miller School of Medicine, University of Miami
William Jacobson, Wei Zhong, George Nomikos, Christina Kurre Olsen, Michael Cronquist Christensen, Elizabeth Merikle

27. Symptomatic and Functional Remission as a Therapeutic Objective in Major Depressive Disorder: Vortioxetine Comparative Data in Working Population
George I. Papakostas*, Massachusetts General Hospital
Rebecca Nielsen, Melanie Brignone, Brigitte Tonnoir

= New Investigator Awardee  = Pharmaceutical Pipeline Presentation
28. A Population Dose-response Analysis of Lurasidone in the Treatment of Major Depressive Disorder with Mixed Features
Yu-Yuan Chiu*, Sunovion Pharmaceuticals, Inc.
Jongtae Lee, Sunny Chapel, Felix Agbo, Antony Loebel

29. Vilazodone Efficacy in Subgroups of Patients with MDD: Post Hoc Analysis of 4 Randomized, Double-blind, Placebo-controlled Trials
Rocsanna Namdar*, Allergan
Susan Kornstein, Suresh Durgam, Changzheng Chen, Carl Gommoll, John Edwards

30. Efficacy of Vortioxetine on Cognitive Functioning in Working Subjects with Major Depressive Disorder
Michael Cronquist Christensen*, H. Lundbeck A/S
Roger S. McIntyre, Ioana Florea, Brigitte Tonnoir, Henrik Loft, Raymond Lam

31. Measurement of Adherence to Antidepressants and Outcomes Among Patients with Both Major Depressive Disorder and Type 2 Diabetes
Charles Vega*, University of California, Irvine, College of Medicine
Russell Becker, Lisa Mucha, Betty Lorenz, Michael Eaddy, Augustina Ogbonnaya

32. The DSM-5 Anxious Distress Specifier Interview: Reliability and Validity
Mark Zimmerman*, Brown University
Emily Walsh, Lia Rosenstein, Douglas Gazarian, Heather Clark

33. Ethics of Medical Marijuana: Medicalization of Treatment with Paucity of Proof
Kimberly Kjome*, Ascension, UT-Dell Medical School, Texas A&M Medical School, UT-Southwestern Medical School, UTMB-Galveston Medical School
Leigh Brown

34. A Classification of Suicidality Disorder Phenotypes
David V. Sheehan*, University of South Florida College of Medicine
Jennifer M. Giddens

= New Investigator Awardee  = Pharmaceutical Pipeline Presentation
35. Chime (Childhood Impulsive Aggression and Molindone ER): Double-blind, Placebo-controlled Study of SPN-810 Added to Standard ADHD Therapy in Children with Impulsive Aggression and ADHD  
Scott Brittain*, Supernus Pharmaceuticals, Inc.  
Gianpiera Ceresoli-Borroni, Tesfaye Liranso, Welton O’Neal, Stefan Schwabe, Robert Findling

36. Lost in Translation: Translatability of Psychiatric Terms - The Example of the Mini-international Neuropsychiatric Interview (M.I.N.I.)  
Marie-Pierre Emery*, Mapi Research Trust  
Anne Boudrot, David V. Sheehan, Catherine Acquadro

37. Human Factors Evaluation of a Novel Digital Health Feedback System in Psychiatry  
Timothy Peters-Strickland*, Otsuka Pharmaceutical  
Jane L. Smith, Benjamin Bartfeld, Linda Pestreich, Shashank Rohatagi, Ainslie Hatch, Felicia Forma, Praveen Raja, John Docherty

38. Involving the Caregivers in Digital Health-enhanced Care of Patients with Serious Mental Illness: Recommendations from an Expert Consensus Survey  
Ainslie Hatch*, Otsuka America Pharmaceutical, Inc.  
John Docherty, Julia E. Hoffman, Ruth Ross

39. Factors Affecting the Use of Digital Health Tools by Healthcare Professionals for Patients with Serious Mental Illness: An Expert Consensus Survey  
Ainslie Hatch*, Otsuka America Pharmaceutical, Inc.  
Julia E. Hoffman, Ruth Ross, John Docherty

40. Web-based Curriculums for Teaching Psychopharmacology: Revision of the Resident, Medical Student, and Primary Care Curriculums  
Ira Glick*, Stanford University School of Medicine

41. Predictors of Medication Adherence Evaluated by Urine Drug Monitoring in Patients Prescribed Antipsychotic Agents  
Mancia Ko*, Ingenuity Health, A Service of Ameritox, Ltd.  
Patricia Woster, Stephen Marder, Thomas Smith

42. Prevention of Drug Interactions Involving Psychotropic Drugs  
Marina Ts’oy-Podosenin*, St. John’s Episcopal Hospital  
Subramoniam Madhusoodanan

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43. Psychiatric Comorbidity in Patients with Pseudobulbar Affect
   Andrea Formella*, Avanir Pharmaceuticals, Inc.
   Joao Siffert

44. Intranasal Oxytocin Modulates Neural Responses to Incentive Stimuli in Humans
   Brian Mickey*, University of Utah School of Medicine
   Joseph Heffernan, Curtis Heisel, Marta Pecina, David Hsu, Jon-Kar Zubieta, Tiffany Love

45. Efficacy of Lurasidone in Patients with Schizophrenia with Prominent Positive Symptoms: A Pooled Analysis of Short-term Placebo-controlled Studies
   Steven Potkin*, University of California, Irvine, School of Medicine
   Michael Tocco, Yongcai Mao, Josephine Cucchiaro, Antony Loebel

46. Effects of Aripiprazole Once-monthly and Paliperidone Palmitate on Work Readiness in Patients from the Qualify Study Stratified by Age
   Steven Potkin*, University of California, Irvine, School of Medicine
   Jean-Yves Loze, Carlos Forray, Ross Baker, Christophe Sapin, Timothy Peters-Strickland, Maud Beillat, Anna-Greta Nylander, Peter Hertel, Simon Nitschky Schmidt, Anna Eramo, Karina Hansen, Dieter Naber

47. Preclinical and Initial Clinical Characterization of APN1125, an α7 Nicotinic Acetylcholine Receptor Agonist for the Treatment of Cognitive Disorders
   Michael Detke*, Indiana University School of Medicine
   Gerald Koelsch, John Ng, Raymond Ng, Geoffrey Bilcer, Terence Kelly

48. A Phase 1 Single- and Multiple-rising Dose Study of the Safety & Pk of EMB-001, a Potential Treatment for Substance Use Disorders, with Exploratory Efficacy Measures in Tobacco Use Disorder
   Michael Detke*, Indiana University School of Medicine
   Carol Gloff, Gary Connor, Sherry McKee, Frank Greenway, Mark Leibowitz, Julie Straub, Ann Robbins, Doug Feltner, Nicholas Goeders

49. Symptom Stability in a 52-week Schizophrenia Extension Study of Treatment with Long-acting Injectable Aripiprazole Lauroxil
   Robert Risinger*, Alkermes
   Arielle Stanford, Yangchung Du, Jacqueline Zummo, Hassan Jamal, Chih-Chin Liu, Amy Claxton

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50. Addressing Adherence Challenges among Recently Diagnosed Schizophrenia Patients: Results from an RCT Comparing a Psychoeducation-based vs. a CBT-based Brief Psychosocial Intervention
Katarzyna Liwski*, LECOM
Douglas Turkington, Sara Tai, Peter Weiden

51. The Efficacy of Lurasidone in Improving Cognition
Rachel Vedder*, Affiliated Research Institute, Inc.
Mohammed Ahmed, Walter Litwin, Hunter Hansen, Christina Ellis, Prakash Bhatia

52. Olfactory Deficits in 22q11.2 Deletion Syndrome Are Significant Compared to Non-deleted Individuals with Clinical Risk and Schizophrenia
Sunny Tang*, University of Pennsylvania
James Yi, Erich Dress, Andrew Wiemken, Tyler Moore, Monica Calkins, Paul Moberg, Raquel Gur, Bruce Turetsky

53. Maintenance Electroconvulsive Therapy (ECT) for Clozapine-resistant Schizophrenia
Georgios Petrides*, The Zucker Hillside Hospital
Raphael Braga, Chitra Malur, Samuel Bailine, Nina R. Schooler, Anil Malhotra, Majnu John, Alan Mendelowitz

54. Metabolic Syndrome in Patients with Schizophrenia Receiving Long-term Treatment with Lurasidone, Quetiapine XR, or Risperidone
Michael Tocco*, Sunovion
John Newcomer, Andrei Pikalov, Hanzhe Zheng, Josephine Cucchiaro, Antony Loebel

55. Effects of Paliperidone Palmitate 3-month and 1-month Formulations on Personal and Social Performance Scale Domain Scores in Patients with Schizophrenia
Dong Jing Fu*, Ortho-McNeil Janssen Scientific Affairs, LLC
Adam Savitz, Ibrahim Turkoz

56. Efficacy of Cariprazine in Negative, Cognitive, and Social Function Symptoms in Schizophrenia: A Post Hoc Analysis of a Randomized, Controlled Trial
Andrew Cutler*, Florida Clinical Research Center
Suresh Durgam, Kaifeng Lu, István Laszlovszky, Erzsébet Szalai, Willie Earley

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57. **Randomized, Double-blind, Placebo-controlled, Phase 3 Study of Encenicline as Pro-cognitive Treatment in Patients with Schizophrenia on Chronic Stable Atypical Antipsychotic Therapy**  
Stephen Brannan*, FORUM Pharmaceuticals  
Nancy Dgetluck, Dana Hilt

58. **Long-term Cariprazine Treatment for the Prevention of Relapse in Patients with Schizophrenia: Analysis of Additional Efficacy Outcomes**  
Raffaele Migliore*, Allergan  
Suresh Durgam, Willie Earley, Kaifeng Lu, István Laszlovszky, György Németh, Henry Nasrallah

59. **The Effect of Tropisetron on the Improvement of P50 Deficits and Aspects of Cognitive Performance in Patients with Schizophrenia**  
Hanjing Emily Wu*, University of Texas Health Science Center at Houston  
Xiang Y. Zhang

60. **Psychopharmacological Treatment of Severely Aggressive Inpatients in a State Psychiatric Hospital**  
Elizabeth Sumner*, Duke University Medical Center  
Stephen Oxley

61. **Paliperidone Palmitate 3-monthly vs. 1-monthly Injectable in Schizophrenia Patients with or without prior Exposure to Oral Risperidone or Paliperidone**  
Maju Mathews*, Janssen Research & Development, LLC  
Huiling Pei, Adam Savitz, Isaac Nuamah, Erica Elefant, David Hough, Larry Alphs, Srihari Gopal

62. **Phase 2 Study of Bremelanotide in Premenopausal Women with Female Sexual Dysfunctions: Responder Analyses based on Minimum Clinically Important Differences Derived from Receiver Operating Characteristic Curves**  
Stanley Althof*, Case Western Reserve University School of Medicine  
Johna Lucas, Raymond Rosen, Robert Jordan, Sally Greenberg, Leonard R. DeRogatis

63. **Hemodynamic and Pharmacokinetic Interactions of Intranasal Bremelanotide and Ethanol in a Phase 1, Randomized, Placebo-controlled, Double-blind, Three-period, Three-way Crossover Study**  
Johna Lucas*, Palatin Technologies, Inc.  
Anita Clayton, Robert Jordan, Leonard R. DeRogatis

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64. Effect of Asenapine on Measures of Hostility in Adults with Bipolar I Disorder or Schizophrenia
Leslie Citrome*, New York Medical College
Ronald Landbloom, Xiao Wu, Suresh Durgam

65. Vortioxetine for Major Depressive Disorder: Number Needed to Treat, Number Needed to Harm, and Likelihood to be Helped or Harmed
Leslie Citrome*, New York Medical College

66. Symptomatic Remission Status in Patients with Schizophrenia Treated with Paliperidone Palmitate (1-month and 3-month Formulations)
Arun Singh*, Janssen Pharmaceuticals R&D
Adam Savitz, Srinari Gopal, Haiyan Xu, Isaac Nuamah, David Hough, Maju Mathews

67. Lurasidone in the Treatment of Sleep Disturbance Associated with Bipolar Depression: Post-hoc Analysis of a Placebo-controlled Trial Followed by a Long-term Extension Study
Michael E. Thase*, Perelman School of Medicine at the University of Pennsylvania
Joyce Tsai, Cynthia Siu, Andrei Pikalov, Antony Loebel

68. Direct and Indirect Effects of Levomilnacipran ER on Functional Impairment in Adults with MDD: Post Hoc Path Analyses
Michael E. Thase*, Perelman School of Medicine at the University of Pennsylvania
Pierre Blier, Carl Gommoll, Changzheng Chen, Angelo Sambunaris, Kenneth Kramer

69. Comparative Evaluation of Vortioxetine as a Switch Therapy in Patients with Major Depressive Disorder
Michael E. Thase*, Perelman School of Medicine at the University of Pennsylvania
Natalya Danchenko, Melanie Brignone, Ioana Florea, Francoise Diamand, Paula Jacobsen, Eduard Vieta

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Thursday, June 2, 2016

Poster Session II with Lunch
Palomino 1-7
12:00 p.m. - 2:00 p.m.

1. Correlation of HLD200 Drug Exposure with Permanent Product Measure of Performance (PERMP)-Correct Scores in Children with ADHD
   Floyd Sallee*, University of Cincinnati
   Ann Childress, Norberto DeSousa, Bev Incledon, Angus McLean

2. Algorithm for Adult Attention-Deficit/Hyperactivity Disorder Management from the Psychopharmacology Algorithm Project at the Harvard South Shore Program
   Bushra Awidi*, Harvard South Shore Psychiatry Program
   David Osser

3. Circuit Modulation by Striatal Cholinergic Interneurons
   Daniel Eskenazi*, New York State Psychiatric Institute, Columbia University
   Stephen Rayport

4. Factors Influencing Generalized Anxiety Disorder (GAD) Diagnosis and Management: Perspectives from Practicing Clinicians
   Purvi Smith*, Health and Wellness Partners
   Andrew Goddard, Larry Culpepper, Joseph Lieberman, Katia Zalkind, Anthony Greco, Jani Hegarty, Randi Roberts

5. The Role of Clinical Outcome Assessment (COA) Data in the Drug Approval Process of Products for the Treatment of Autism Spectrum Disorder in the USA and Europe: A Review of Guidance Documents and Authorizations of Medicinal Products
   Marie-Pierre Emery*, Mapi Research Trust
   Caroline Anfray, Catherine Acquadro, Cécile Perret, Patricia Anderson

6. Efficacy of Lurasidone in Bipolar Depression: Pooled Results of Two Adjunctive Studies with Lithium or Valproate
   Joyce Tsai*, Sunovion Pharmaceuticals
   Mauricio Tohen, Andrei Pikalov, Antony Loebel

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| 7. **Safety and Efficacy of Cariprazine in FDA-approved Dose Ranges for Schizophrenia and Bipolar I Disorder: A Pooled Post Hoc Analysis**  
Willie Earley*, Allergan  
Suresh Durgam, Kaifeng Lu, György Németh, István Laszlovszky |
| 8. **Actigraphy Biomarker Data Correlate with Bipolar Disorder Mood Symptoms**  
Terence Ketter*, Stanford University School of Medicine  
Shefali Miller, Nathaniel Stockham, Saloni Shah, Dennis Do, Anshul Sitaram |
| 9. **Potential Biomarkers of Depression and Mania: The Association of Sleep, Kynurenine and Tryptophan in Acute Bipolar Disorder**  
Dahlia Mukherjee*, Penn State Milton S. Hershey Medical Center  
Venkatesh Krishnamurthy, Aubrey Reider, Adem Can, Maureen Groer, Dietmar Fuchs, Teodor Postolache, Erika Saunders |
| 10. **OPEN BOARD** |
| 11. **The Suicide Ideation and Behavior Assessment Tool: Validation of a Novel Measure of Suicidal Ideation and Behavior and Perceived Risk of Suicide**  
Larry Alphs*, Janssen  
Carla Canuso, David Williamson, SIBAT Consortium |
| 12. **Addressing Data Quality Challenges in Rare Disease Clinical Trials**  
Magda Perez*, inVentiv Health  
Chris Brady, Julie Marsh, Kristi Bertzos, Lori Vivian |
| 13. **Marijuana Effect on Differentiating an Opioid from Placebo During the Discrimination Phase of a Human Abuse Potential Study**  
Clark Johnson*, PRA Health Sciences  
Michael Smith, Shawn Searle, Vicky Newton, Lynn Webster |
| 14. **ITI-007 Dose Selection Across Psychiatric and Neurological Therapeutic Indications**  
Jelena Saillard*, Intra-Cellular Therapies, Inc.  
Kimberly E. Vanover, Robert E. Davis, Cedric O’ Gorman, Michal Weingart, Sharon Mates |

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= Pharmaceutical Pipeline Presentation
15. Methodological Considerations in the Assessment of Abuse Potential in Phase 1-3 Clinical Trials of CNS Drugs
Michael Hufford*, Pinney Associates
Jack Henningfield, Reginald Fant, Sidney Schnoll

16. An Evaluation of Urine Drug Monitoring in the Treatment of Patients with Serious Mental Illness
Patricia Woster*, Ingenuity Health
Mancia Ko, Thomas Smith

17. Brain-derived Neurotrophic Factor (BDNF) Val66Met Polymorphism Induces Memory Deficits in Elderly
Saulo Tractenberg*, Pontifical Catholic University of Rio Grande do Sul
Lucas Azeredo, Mateus Levandowski, Tatiana de Nardi, Julia Kommers-Molina, Wieck Andrea, Rodrigo Grassi-Oliveira

18. MIN-117 – A Promising New Antidepressant with a Novel Mechanism of Action Profile
Michael Detke*, Indiana University School of Medicine
Corinne Staner, Nadine Noel, Jay Saoud, Joseph Reilly, Remy Luthringer

19. OPEN BOARD

20. OPEN BOARD

21. Lurasidone for Major Depressive Disorder with Mixed Features: Effect of Irritability
Andrei Pikalov*, Sunovion Pharmaceuticals, Inc.
Alan C. Swann, Joyce Tsai, Yongcai Mao, Antony Loebel

22. OPEN BOARD

23. A Five Year Study of Vagus Nerve Stimulation Compared to Treatment as Usual in Historical ECT Responders and Non-responders
Scott Aaronson*, Sheppard Pratt Health System
Peter Sears, Pradheep Raman, Jennifer Sklar, Mark Bunker

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William Jacobson*, Takeda Pharmaceuticals
Philip D. Harvey, Wei Zhong, George Nomikos, Christina Kurre Olsen, Michael Cronquist Christensen, Elizabeth Merikle

25. SAGE-547 (Allopregnanolone) and SAGE-217: Investigational Neuroactive Steroids Targeting the GABAA Receptors for Treatment of CNS Disorders
Stephen Kanes*, Sage Therapeutics
Helen Colquhoun, Handan Gunduz-Bruce, James Doherty, Shane Raines, Ethan Hoffman, David R. Rubinow, Samantha Meltzer-Brody

26. Evaluation of the Efficacy and Safety of Alks 5461 as Adjunctive Therapy in MDD: Results of FORWARD-3 and FORWARD-4 Studies
Elliot Ehrich*, Alkermes, plc
William Martin, Asli Memisoglu, Sanjeev Pathak, Arielle Stanford, Irena Webster, Ying Jiang, Michael E. Thase, Maurizio Fava

27. Psychiatrist Attitudes about Novel and Emerging Treatments for Depression: Off-label Ketamine
Dawn Roberts*, Bradley University
Aman Singh, Jane Larouche, Laura Jorgenson, Peter Alahi

28. Validation of Patients for a CNS Trial of Major Depressive Disorder
Martina Flynn*, Massachusetts General Hospital
David Mischoulon, Janet Witte, Vincent Pisano, Max Martinson, Marlene Freeman, Mi Hillefors, George Papakostas, Maurizio Fava

29. Susceptibility of Male and Female C57BL/6 Mice to Oxidative Stress in the Hippocampus in an LPS Model of Depression
Caitlin Millett*, Penn State Milton S. Hershey Medical Center
Erika Saunders, Shannon Kelleher

30. Direct and Indirect Effects of Levomilnacipran ER on Functional Impairments in MDD Patients with Cognitive Difficulties: Post Hoc Path Analyses
Ken Kramer*, Allergan
Roger S. McIntyre, Philip Harvey, Carl Gommoll, Changzheng Chen, Keith Wesnes

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31. Effects of Adjunctive Brexpiprazole on Chronobiologic Parameters in Patients with Major Depressive Disorder and Sleep Disturbances
Amy Hilty*, Otsuka Pharmaceuticals
Aurelia Mittoux, Peter Meisels, Ross Baker, Andrew Krystal

32. Treatment Patterns, Healthcare Resource Utilization, and Costs Following First-line Antidepressant Treatment in MDD: A Retrospective US Claims Database Analysis
Vanessa Perez*, Takeda Pharmaceuticals International, Inc.
Genevieve Gauthier, Annie Guerin, Maryia Zhdanava, Clément François, William Jacobson, George Nomikos, Elizabeth Merikle

33. The Digit Symbol Substitution Test (DSST): Psychometric Properties and Clinical Utility in Major Depressive Disorder
Judith Jaeger*, Albert Einstein College of Medicine

34. Into the Island of Addiction: Insights into the Mechanisms of Action of Repetitive Transcranial Magnetic Stimulation
Primavera Spagnolo*, National Institute on Alcohol Abuse & Alcoholism, National Institutes of Health

35. Case Study of Magnesium in the Treatment of Impulse Attack Suicidality Disorder
David V. Sheehan*, University of South Florida College of Medicine
Jennifer M. Giddens

36. Extended-release Molindone for Impulsive Aggression: Phase 2 Double-blind, Placebo-controlled Trial in Children with ADHD Receiving Optimized Stimulant Monotherapy and Behavioral Therapy
Scott Brittain*, Supernus Pharmaceuticals, Inc.
Jennifer D. Stocks, Janet K. Johnson, Tesfaye Liranso, Robert L. Findling

37. Teaching the Teachers of Clinical Psychopharmacology
Ira Glick*, Stanford University School of Medicine
Carl Salzman

38. Assessment of Amphetamine Withdrawal Symptoms of Lisdexamfetamine Dimesylate Treatment for Adults with Binge Eating Disorder
Judith Kando*, Sunovion
Maria Gasior, Barry Herman, Jana Radewonuk, Reginald Fant, Sidney Schnoll, Susan McElroy

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Arun Singh*, Janssen Pharmaceuticals R&D
Srihari Gopal, Haiyan Xu, Kelly McQuarrie, Adam Savitz, Isaac Nuamah, Kimberly Woodruff, Maju Mathews

40. A Case Series on the Effectiveness of Lurasidone in Patients with Stuttering
Janet Charoensook*, UC Riverside
Carlos Fernandez, Shalin Patel, Julia Hoang, Gerald Maguire

41. Psychiatric Stability Maintained in Tardive Dyskinesia Subjects Treated with Valbenazine (NBI-98854)
Jean-Pierre Lindenmayer*, New York University
Richard Josiassen, Joshua Burke, Scott Siegert, Bill Aurora

42. Kinect 3: A Randomized, Double-blind, Placebo-controlled Phase 3 Trial of Valbenazine (NBI-98854) for Tardive Dyskinesia
Mary Ann Knesevich*, University Hills Clinical Research
Stephen Marder, Robert A. Hauser, Grace S. Liang, Christopher O’Brien

43. MoodNetwork.org: A Seminal Online Study
Andrew Nierenberg*, Massachusetts General Hospital
Louisa Sylvia, Casey Hearing, Alexandra K. Gold, Rebecca Montana, Roberta Tovey, Anthony Debeneditis, Allen Doederlein, Susan Edgman-Levitan, Muffy Walker, Donna Holland Barnes, Jonathan Alpert, Jordan Smoller, Marlene Freeman, Thilo Deckersbach

44. Results from an Expert Consensus Survey: Patient-related Factors in the Use of Digital Health Tools in Patients with Serious Mental Illness
Ainslie Hatch*, Otsuka America Pharmaceutical, Inc.
Julia Hoffman, Ruth Ross, John Docherty

45. A Treatment Refinement Study to Optimize the Habit Formation Program for Improving Adherence to Oral Medications in Schizophrenia and Schizoaffective Disorder
Ainslie Hatch*, Otsuka America Pharmaceutical, Inc.
John Docherty, Deborah Profit, Erica Lawson, Anke Adenwala, Dawn Velligan

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46. Randomized, Double-blind, Placebo-controlled, Phase 3 Study of Encenicline as Pro-cognitive Treatment in Patients with Schizophrenia
Steven Potkin*, University of California, Irvine, School of Medicine
Stephen Brannan, Nancy Dgetluck, Dana Hilt

47. Cariprazine for Negative Symptoms of Schizophrenia: A Pooled Post Hoc Analysis of 2 Randomized, Double-blind, Placebo- and Active-controlled Trials
Suresh Durgam*, Forest Research Institute, A Subsidiary of Actavis, plc
Willie Earley, Kaifeng Lu, György Németh, István Laszlovszky, Balázs Szatmári, Henry Nasrallah

48. Understanding Factors Impacting on Cgi-S Vs. Cgi-I Discrepancies: An Exploratory Analysis
David Daniel*, Bracket Global, LLC
Alan Kott

49. Switching Patients with Acute Schizophrenia to Brexpiprazole: Post-hoc Analysis of a Double-blind Randomized Maintenance Treatment Study
Catherine Weiss, John Ouyang, Anna Eramo, Emmanuelle Weiller, Ross Baker

50. Relationship Between Response to Aripiprazole Once-monthly and Paliperidone Palmitate on Work Readiness and Functioning: A Post-hoc Analysis of QUALIFY, A Head-to-head Study in Schizophrenia
Ross Baker*, Otsuka
Steven Potkin, Jean-Yves Loze, Carlos Forray, Christophe Sapin, Timothy Peters-Strickland, Maud Beillat, Anna-Greta Nylander, Peter Hertel, Simon Nitschky Schmidt, Anna Eramo, Karina Hansen, Dieter Naber

51. Long-term Safety and Durability of Effect of Aripiprazole Lauroxil in a One-year Schizophrenia Extension Study
Robert Risinger*, Alkermes
Arielle Stanford, Yangchung Du, Jacqueline Zummo, Hassan Jamal, Chih-Chin Liu, Amy Claxton

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52. A Multicenter, 8-week Study to Assess Usability of a Digital Health Feedback System in Adults with Schizophrenia Treated with Oral Aripiprazole
Timothy Peters-Strickland*, Otsuka Pharmaceutical
Linda Pestreich, Ainslie Hatch, Shashank Rohatagi, Ross Baker, John Docherty, Lada Markovtsova, Praveen Raja, Peter Weiden, David Walling

53. Efficacy and Safety of Intramuscular Ziprasidone in Chinese Schizophrenia Patients with Agitation: A Randomized, Blind, Active Parallel-controlled, Multicenter Clinical Trial
Yifeng Shen*, Shanghai Mental Health Center
Jianfeng Lou, Huafang Li

54. Effect of Brexpiprazole and Aripiprazole on Weight: An Analysis of Long-term Trials in Schizophrenia
Catherine Weiss*, Otsuka Pharmaceutical Development & Commercialization, Inc.
Keva K. Gwin, Ruth A. Duffy, Ross Baker, Emmanuelle Weiller

Marjie Hard*, Alkermes, Inc.
Brian Sadler, Richard Mills, Karen Rowland Yeo, Ryan Turncliff, Leslie Citrome

56. Handwriting Kinematics in the Assessment of Pharmacotherapeutic Outcomes in Psychiatric Populations
Michael Caligiuri*, University of California, San Diego
Hans-Leo Teulings

57. Effects of Aripiprazole Once-monthly and Paliperidone Palmitate in Patients with Schizophrenia and Concomitant Substance use: A Post-hoc Analysis of QUALIFY, A Head-to-head Study
Phyllis Salzman*, Otsuka Pharmaceutical Development & Commercialization, Inc.
Dieter Naber, Ross Baker, Anna Eramo, Carlos Forray, Karina Hansen, Christophe Sapin, Maud Beillat, Timothy Peters-Strickland, Anna-Greta Nylander, Peter Hertel, Jean-Yves Loze, Henrik Steen Andersen, Steven Potkin

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58. Long-term Effectiveness of Aripiprazole Once-monthly is Maintained in the Qualify Extension Study
Anna Eramo*, Lundbeck
Dieter Naber, Ross Baker, Carlos Forray, Karina Hansen, Christophe Sapin, Timothy Peters-Strickland, Anna-Greta Nylander, Peter Hertel, Simon Nitschky Schmidt, Jean-Yves Loze, Steven Potkin

Britta Galling*, The Zucker Hillside Hospital
Alexandra Roldán, Frozan Walyzada, Liz Rietschel, Katsuhiko Hagi, Wei Zheng, Xiao-Lan Cao, Yu-Tao Xiang, Mathias Zink, John M. Kane, Jimmi Nielsen, Stephan Leucht, Christoph Correll

60. A Phase 2, EFFICACY, SAFETY, and Tolerability Study of ALKS 3831 in Schizophrenia with Alcohol Use Disorder
Bernard Silverman*, Alkermes, Inc.
Sanjeev Pathak, David McDonnell, Lauren DiPetrillo, Adam Simmons, Ying Jiang, Jacqueline Zummo, Hassan Jamal

61. Evaluation of Paliperidone Palmitate Long-acting Injectable Therapy by Duration of Illness in Patients with Schizophrenia
Maju Mathews*, Janssen Research & Development, LLC
Brianne Browne, Ibrahim Turkoz, Branislav Mancevski

62. The National Pregnancy Registry for Atypical Antipsychotics: Effects of Fetal Exposure on Risk for Major Malformations
Lee Cohen*, Massachusetts General Hospital
Adele Viguera, Marlene Freeman, Tao Hou, Alexandra Sosinsky, Gina Savella, Danna Moustafa, Sonia Hernández-Díaz

63. Relationship Between Depression and Pregnancy in Rural Guatemala: A Pilot Study
Ahmad Hameed*, Penn State College of Medicine
Myra Qureshi, Pevitr Bansal, Usman Hameed, Scott Bunce

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2016 ASCP Fall Meeting
October 29-30, 2016
Grand Hyatt New York, New York

2017 ASCP Annual Meeting
May 29 – June 2, 2017
Loews Miami Beach, Miami Beach, Florida

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