#### **Confirmed Speakers**

#### **Keynote Speakers**

KL01 - Recent Developments in Strategies and Tactics Towards the Synthesis of Complex Secondary Metabolites as Enabling Tools for the Study of Biology and Medicine



Erick M. CARREIRA (ETH ZÜRICH, Zürich, Switzerland)

#### KL02 - Activity-based proteomics - Protein and Ligand Discovery on a Global Scale



Benjamin CRAVATT (THE SCRIPPS RESEARCH INSTITUTE, La Jolla, United States)

#### KL03 - Novel Approaches in the Design of CNS Drug Candidates and PET Ligands



Anabella VILLALOBOS (BIOGEN, Cambridge, United States)

#### Addressing Preclinical Toxicity – Approaches and Lessons Learned

#### PL06 - Reducing Bioactivation Potential of Drug Candidates: Implications for Preclinical Drug Optimization



Andreas BRINK (F. HOFFMANN-LA ROCHE, Basel, Switzerland)

#### PL04 - Mechanism-Based Toxicities Associated With NAMPT Inhibition and Related Mitigation Strategies



Peter DRAGOVICH (GENENTECH INC., San Francisco, United States)

## OC02 - Small Structural Changes Leading to Major Impact on Safety: Developing Safety Strategies in Medicinal Chemistry



Martin PETTERSSON (PFIZER, Cambridge, United States)

## PL05 - Utilizing in Depth Understanding of a Molecules Off-Target Profile to Tailor Clinical and Preclinical Safety Assessments



Douglas THOMSON (CELLZOME GMBH, Heidelberg, Germany)

#### **Advances in Lead Generation**

#### PL08 - A Chemist's Guide to Modern Phenotypic Drug Discovery

#### **Confirmed Speakers**



Monika ERMANN (EVOTEC LTD, Oxfordshire, United Kingdom)

### OC03 - CDK8 Inhibitors with Pre-Engineerd Long Residence Time, Exhibiting Efficacy in Tumor Xenograft Models



Koen HEKKING (MERCACHEM-SYNCOM, Nijmegen, The Netherlands)

#### PL07 - ADAS (Affinity Directed Automated Synthesis): A New Technology to Accelerate Lead Generation



Eva Maria MARTIN (ELI LILLY, Madrid, Spain)

#### PL09 - From Multiple Hit Series to (Pre)Clinical Candidates Using DNA-Encoded Library Technology



Sanne SCHRODER GLAD (NUEVOLUTION A/S, Copenhagen, Denmark)

#### **Advances in Synthetic Methods**

#### **PL01 - Assembly Line Synthesis**



Varinder K. AGGARWAL (UNIVERSITY OF BRISTOL, Bristol, United Kingdom)

#### PL02 - Photochemical Reactions en route to Structurally Complex Molecules



Thorsten BACH (TECHNISCHE UNIVERSITÄT MÜNCHEN, Garching, Germany)

#### PL03 - Expanding the Potential of Organocatalysis with Light



Paolo MELCHIORRE (INSTITUTE OF CHEMICAL RESEARCH OF CATALONIA (ICIQ), Bologna, Italy)

### OC01 - Exploring 3-D Pharmaceutical Space: New CH Functionalisation Reactions of Oxygen and Sulfur Heterocycles



Peter O'BRIEN (UNIVERSITY OF YORK, York, United Kingdom)

#### **Alternative Modalities**

#### OC05 - Proteolysis Targetting Chimera: A New Frontier in Medicinal Chemistry

#### **Confirmed Speakers**



Niall ANDERSON (GLAXOSMITHKLINE, Hertfordshire, United Kingdom)

#### PL13 - Messenger RNA as a Novel Therapeutic Approach



Kerry BENENATO (MODERNA THERAPEUTICS, Cambridge, United States)

#### PL11 - Intracellular Delivery of Macromolecules



David TELLERS (MERCK & CO. INC (MSD), West Point, United States)

### PL12 - New Modalities Probe and Hit Finding for Challenging Targets in Cardiovascular and Metabolic Diseases



Eric VALEUR (ASTRAZENECA, IMED BIOTECH UNIT, Cambridge, United States)

#### **Challenges and Opportunities in Fragment Based Drug Discovery**

#### OC07 - Fragment-Centric Methodologies for the Discovery of DOT1L Inhibitors



Christoph GAUL (NOVARTIS, Basel, Switzerland)

## OC12 - Rational Design of Small-Molecules Inhibitors of Human Cyclophilins with a Pan Viral Activities by Fragment Based Drug Design Using a Linking Strategy



Jean-Francois GUICHOU (CBS INSERM U1054, Montpellier, France)

#### PL17 - Drug Discovery for Challenging Targets by X-ray Crystallographic Fragment Screening



Tom HEIGHTMAN (ASTEX PHARMACEUTICALS, Cambridge, United Kingdom)

#### PL18 - The Impact of Fragments on Drug Discovery



Rod HUBBARD (UNIVERSITY OF YORK & VERNALIS, Cambridge, United Kingdom)

#### **Chemical Biology in Drug and Target Discovery**

#### OC04 - Chemical Physiology of Antibody Conjugates and Natural Products

#### **Confirmed Speakers**



Gonçalo BERNARDES
(INSTITUTO DE MEDICINA MOLECULAR, PORTUGAL & UNIVERSITY OF CAMBRIDGE, Cambridge, United Kingdom)

#### PL10 - Fluorescent and Bioluminescent Sensor Proteins



Kai JOHNSSON (MAX-PLANCK INSTITUTE FOR MEDICAL RESEARCH, Heidelberg, Germany)

#### Late Stage Functionalization

#### PL15 - The Quest for Efficient Ligands in Asymmetric C-H Functionalizations



Nicolai CRAMER (ECOLE POLYTECHNIQUE FÉDÉRALE DE LAUSANNE, Lausanne, Switzerland)

#### PL16 - Catalytic Approaches to Simplifying Synthesis



Darren J. DIXON (UNIVERSITY OF OXFORD, Oxford, United Kingdom)

#### PL14 - New Chemical Tools for the Late Stage Functionalization of Biomolecules



Matthew GAUNT (UNIVERSITY OF CAMBRIDGE, Cambridge, United Kingdom)

#### OC06 - Synthetic Routes to Oxindoles via Metal Catalysis



Mark LAUTENS (UNIVERSITY OF TORONTO, Toronto, ON, Canada)

#### **Drug Discovery Tales**

OC10 - Molecular Accessibility - Measuring and Understanding the Intracellular Free Concentration of Drugs During Lead Optimisation

(EYEDPHARMA)

### OC09 - Discovery of Tak-041: A Potent and Selective Gpr139 Agonist for the Treatment of Negative Symptoms Associated with Schizophrenia



Holger MONENSCHEIN (TAKEDA CALIFORNIA, INC, San Diego, United States)

#### OC08 - Discovery of Allosteric Malt1 Protease Inhibitors with High in Vivo Efficacy

#### **Confirmed Speakers**



Jean QUANCARD (NOVARTIS, Basel, Switzerland)

OC11 - Discovery of a Ketohexokinase Inhibitor for the Treatment of Nafld/Nash: Fragment-to-Candidate via Structure-Based Drug Design and Parallel Chemistry



Brian RAYMER (PFIZER, Cambridge, United States)