Conference Agenda

Tuesday 10th September

Pembroke College, Oxford

16.00 Registration

17.00 Opening Remarks

Professor Matthew Wood, Director, Oxford-Harrington Rare Disease Centre

17.15 Keynote Lectures

The Harrington Discovery Institute

Jonathan Stamler, President, Harrington Discovery Institute

Gene therapy for Retinitis Pigmentosa

Robert MacLaren, Professor of Ophthalmology, Nuffield Department of Clinical Neurosciences, University of Oxford

Gene therapy for Cystic Fibrosis

Eric Alton, Chair in Gene Therapy, Faculty of Medicine, National Heart & Lung Institute, Imperial College London

19.00 Drinks and canapés reception

Wednesday 11th September

9.00 Session I

Title tbc

Rita Balice-Gordon Global Head, Rare and Neurological Diseases Therapeutic Area, Sanofi

Title tbc

Thomas Voit, Director of the NIHR Great Ormond Street Hospital (GOSH) BRC, Professor and Honorary Consultant of Paediatrics at GOSH

Title tbc- new drug targets in Friedreich's Ataxia

Richard Wade-Martins, Professor of Molecular Neuroscience, Department of Physiology, Anatomy and Genetics, University of Oxford

Title tbc

Kevin Talbot (tbc), Professor of Motor Neuron Biology, Nuffield Department of Clinical Neurosciences, University of Oxford

Title tbo

Francesco Muntoni, Chair of Paediatric Neurology, UCL

Title tbc

Esther Becker, Associate Professor of Neurobiology, Departement of Physiology, Anatomy & Genetics, University of Oxford

11.00 Coffee

Keynote Lecture - NIH National Center for Advancing Translational Sciences

Chris Austin, Director, NIH NCATS

12.30 Lunch

13.15 **Session 2**

Title tbc In born errors of metabolism

Tim Cox, Department of Medicine, University of Cambridge

Title tbc

Wyatt Yue, Structural Genomics Consortium, University of Oxford

Title tbc

Atul Chopra, Investigator, Harrington Discovery Institute, University Hospitals, Cleveland

Title tbo

Bob Lightowlers, Professor of Molecular Neuroscience, University of Newcastle

14.35 **Session 3**

Title tbc Gene therapy in Dermatology

John McGrath, King's College London

Gene therapy in Duchenne Muscular Dystrophy and Spinal Muscular Atrophy

Laurent Servais, Professor of Paediatric Neuromuscular Diseases, Department of Paediatrics, University of Oxford

LifeArc Philanthropic Fund for Rare Disease Translational Research

Catriona Crombie, Philanthropic Fund Manager, LifeArc

15.35 Coffee break

15.55 Session 4 Genomics and Genetic Approaches

100k Genomes Project

Richard Scott, Clinical Lead, Rare Diseases, Genomics England

Title the

Lucy Raymond, Professor of Medical Genetics and Neurodevelopment, Department of Medical Genetics, University of Cambridge

Title tbc

Jenny Taylor, Wellcome Trust Centre for Human Genetics, University of Oxford

Title tbc

Henry Houlden, Consultant Neurologist, UCL and Head of Neurological GeCIP Domain, Genome England

Title the

Hugh Watkins, Radcliffe Professor of Medicine, University of Oxford

Title tbc

Tim Barrett, Leonard Parsons Professor of Paediatrics and Child Health, University of Birmingham

Title tbc

Holm Uhlig, Professor of Paediatric Gastroenterology, University of Oxford

Conference residential dinner

Thursday 12th September

Pembroke College, Oxford

8.30 Registration and coffee

9.00 Session 5 Industry

Title tbc

Steve Murray, VP Strategy and Innovation, Mallinckrodt Pharmaceuticals

Translational Research for Repositioning TAK063 in Fragile X Syndrome – A Potential New Treatment

Christine Charman, Senior Director, External Asset Lead, Centre for External Innovation, Takeda Pharmaceuticals

Title tbc

Per Lundin, Chief Operating Officer, Evox Therapeutics

Title tbc

Michael Panzara, Chief Medical Officer, Wave Life Sciences

Title tbc

Madhurima Benekareddy, Project Lead and Lab Head, Roche

Title tho

Stuart Hughes, Vertex Pharmaceuticals

11.20 Session 6 Haematology and Immunology

Emerging therapies for hemophilia and sickle cell anemia

Seng Cheng, CSO Rare Diseases, Pfizer

Title tbc

Bobby Gaspar, CSO, Orchard Therapeutics

Rare blood disorders

Mike Laffan, Imperial College

Title the

Georg Holländer, Hoffmann and Action Medical Research Professor of Developmental Medicine

Title tbc

Willem Owehand, Professor of Experimental Haematology, University of Cambridge

Birmingham Rare Disease Network

David Adams, Director, NIHR Birmingham Biomedical Research Centre, University of Birmingham

13:20 Closing Remarks