Overview

One in every ten Americans has a rare disease, a health condition by definition that afflicts fewer than 200,000 people per year in the United States and collectively affects an estimated 25 million people. Treatment options are few as only 400 FDA-approved drugs are available to treat the diverse spectra of 7,000 rare diseases. In this complimentary live webinar, WuXi brings together key thought leaders and players to explore the unique challenges and opportunities in the search for effective therapies to address this large unmet medical need.

Invited Speakers

- **Philip Reilly, MD, JD**
  Venture Partner
  Third Rock Ventures
- **Alexandra Glucksmann, PhD**
  Chief Operating Officer
  Editas Medicine
- **Phil Vickers, PhD**
  Global Head of Research and Development
  Shire
- **Gerald Cox, MD, PhD**
  Vice President of Clinical Development
  Genzyme, a Sanofi company
- **Richard Soll, PhD**
  SVP and Head of International Discovery Service Unit
  WuXi AppTec

Speaker Biography

**Philip Reilly, MD, JD**
Venture Partner
Third Rock Ventures

Phil joined Third Rock Ventures in 2009 to discover, launch and build companies pursuing breakthrough treatments for rare genetic diseases. He is a highly respected clinical geneticist with an extensive track record of launching and building companies in the rare disease space.

In addition to his role at Third Rock, Phil is currently a trustee of Cornell University and is on the board of overseers of Weill Cornell Medical College. Prior to joining Third Rock, he was the chief executive officer and chairman of Interleukin Genetics, and before that, he was the executive director of the Eunice Kennedy Shriver Center for Mental Retardation. He served twice (2000 and 2003) as president of the American Society of Law, Medicine and Ethics. He is a founding fellow of the American College of Medical Genetics. He has authored or co-authored more than 100 articles in scholarly journals as well as several books about clinical genetics. Phil has held teaching positions at Tufts University School of Medicine and Harvard Medical School. Phil received his BA from Cornell University and holds an MD from Yale University, a JD from Columbia University, is a member of the Massachusetts Bar and is board certified in internal medicine and clinical genetics.

When he was much younger, Phil spent a year bartending in the Caribbean.
Phil is Global Head of Research and Development at Shire. Phil is responsible for overseeing preclinical research and development, clinical research, regulatory affairs, and medical affairs at Shire. He oversees the organization's growing product portfolio and plays a key role in developing and executing Shire's global business strategy. Phil is a member of Shire's Executive and Pipeline Committees.

Phil has over 25 years in the pharmaceutical industry experience. He joined Shire from Resolvyx Pharmaceuticals, where he was Chief Scientific Officer and President. While at Resolvyx, he was a member of the Board of Directors, with accountability for all preclinical and clinical research, as well as partnering with investors, external business development partners, and establishing external collaborations.

Prior to this, he was Senior Vice President and US Head of Research at Boehringer-Ingelheim Pharmaceuticals, where he had responsibility for the Ridgefield CT research site, which focused on therapeutic areas of cardiovascular and immunology & inflammation. Before BI, he spent over 13 years with Pfizer in a number of R&D leadership positions of increasing responsibility in the US and UK. These roles included drug discovery leadership across therapeutic areas, and leading the Pfizer Research and Technology Center in Cambridge, MA. He started his career with Merck Frosst in Canada, where he played a key role in the successful leukotriene and cyclooxygenase programs.

Phil holds a PhD in Biochemistry from the University of Toronto, and a Bachelor of Science degree in Applied Biochemistry from the University of Salford, Manchester. He was also a Visiting Fellow at the National Cancer Institute in Bethesda, Maryland.
Richard Soll, PhD is SVP and Head of International Discovery Service Unit at WuXi AppTec, a Shanghai headquartered and New York Stock Exchange listed (NYSE: WX) premium provider of pharmaceutical R&D services across the pharmaceutical R&D value chain. In this capacity, Rich has advanced more than 40 integrated programs across major target classes and therapeutic indications from hits to lead optimization campaigns, giving rise to clinical candidates including those designated by the FDA as breakthrough therapies. Previously, Rich was CSO and VP of R&D at TargeGen where he led innovative clinical-stage R&D programs including a highly selective JAK2 inhibitor fedratinib (SAR302503; TG101348) for the treatment of myeloproliferative disorders, leading to the acquisition of TargeGen by Sanofi. Before TargeGen, Rich founded the chemistry department at 3-Dimensional Pharmaceuticals as VP of Chemistry and spent 10 years at Wyeth Pharmaceuticals. Rich has been an SAB member to biotechs, advisor to entrepreneurs, and has produced more than 100 patents and papers.

Gerry is Vice President of Clinical Development in the Rare Diseases Group at Genzyme, a Sanofi company. He also is a practicing clinical geneticist at Boston Children’s Hospital and an Instructor in Pediatrics at Harvard Medical School. Since joining Genzyme in 2000, Gerry has been involved in the development of several enzyme replacement therapies (ERT) and substrate reduction therapies (SRT) for lysosomal storage disorders. He played a key role in the worldwide approval of Aldurazyme (larondase) for Mucopolysaccharidosis type I in 2003, Japan approval of Elaprase (idursulfase) for Mucopolysaccharidosis type II in 2007, and US approval of Cerdelga (eliglustat) for Gaucher disease type 1 in August 2014. His group recently completed a successful phase 1b study of ERT for Niemann-Pick disease type B and is planning phase 2 studies of a CNS-accessible SRT for neurodegenerative glycosphingolipidoses.

Gerry graduated magna cum laude with a BA in biology from Harvard College in 1980 and obtained an MD and a PhD in biology from UC San Diego in 1989. Following a pediatrics residency and clinical and research fellowships in genetics at Boston Children’s Hospital, he became a staff physician and directed the Human Genomic Mapping Facility. Gerry is board-certified in Clinical, Biochemical, and Molecular Genetics and formerly in Pediatrics. He served on the scientific advisory boards of the Barth Syndrome Foundation and the Pediatric Cardiomyopathy Registry, and he has authored more than 60 peer-reviewed publications. In addition to lysosomal storage disorders, Gerry’s clinical interests include genetic causes of cardiomyopathy and eye disease.