

TENTATIVE PROGRAM

2019



10th Annual Congress on

RARE DISEASES AND ORPHAN DRUGS

October 16-17, 2019 Tokyo | Japan

10th Annual Congress on**RARE DISEASES AND ORPHAN DRUGS**

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Program @ Glance

Day 1 October 16, 2019	
Opening Ceremony	
Plenary & Keynote Speeches (09:00-11:30)	
Networking and Refreshments Break	
Group Photo	
Speaker Session I (11:30-13:00)	Different types of Rare Diseases Clinical Research and Public Awareness Mystery Diagnosis of Rare Diseases
	Panel Discussion
Lunch Break	
Speaker Session II (13:00-16:00)	Challenges in Rare Diseases Treatment Rare Infectious Diseases and Immune Deficiencies Rare Diseases in Cancer
	Panel Discussion
Networking and Refreshments Break	
Speaker Session III (16:00-18:00)	Rare Diseases in Aging Orphan Drugs- development trends and strategies Orphan Drugs and Ethical Issues
	Panel Discussion
Day 1 concludes...	
Day 2 October 17, 2019	
Opening Ceremony	
Plenary & Keynote Speeches (09:00-11:30)	
Networking and Refreshments Break	
Speaker Session I (11:30-13:00)	Clinical Research on Orphan Drugs Orphan Drugs and Ethical Issues Future Hereditary of Rare Diseases and Orphan Drugs Entrepreneurs Investment Meet
	Panel Discussion
Lunch Break	
Speaker Session II (13:00-16:00)	Different types of Rare Diseases Mystery Diagnosis of Rare Diseases Patient organizations and their role in drug development or clinical research
	Panel Discussion
Networking and Refreshments Break	
Speaker Session III (16:00-18:00)	Poster Presentations
	Best Poster Award Distribution
Day 2 concludes...	



A patient at the table: How partnership with patients improves research and enhances service delivery

Alastair Kent, Genetic Alliance, UK



Measuring the impact of diagnosis and treatment of rare diseases

Alba Ancochea, EURORDIS, Spain



Targeting familial Alzheimer's disease

Michael S. Wolfe, University of Kansas, USA



Consumer Directed Precision Health - Convergence of Multi- omics, Environment, Life style and Behaviors

Praduman Jain, Vibrent Health Inc., USA



Speeding up access to medicines for patients with unmet medical need: Integrating evidence and regulatory pathways

Stella Blackburn, QuintilesIMS, UK

PAST KEYNOTE SPEAKERS



Pediatric Rare Disease Enrollment Case Study in Latin America

Sara Tylosky, Farmacon, USA



Lifting the burden of Rare Disease by providing access to next generation sequencing

Romina Ortiz, Rare Genomics Institute, USA

PAST WORKSHOP'S



Helping patients cope with, and recover from the effects of living with a rare disease: A patient's perspective

Mark Landiak, Corporate Dynamics, Inc. USA



Overview of rare disease funding at NIH

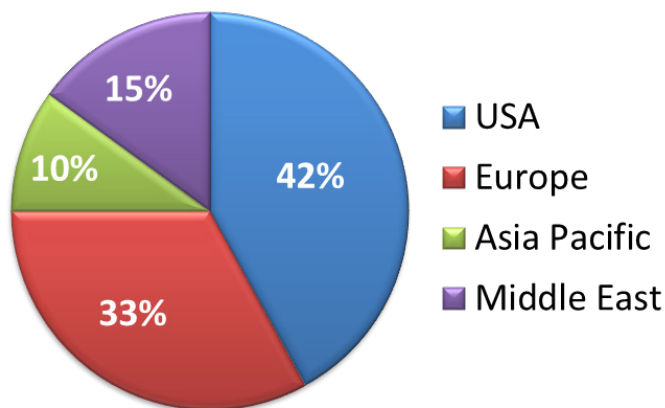
Jason Wan, National Institutes of Health, USA



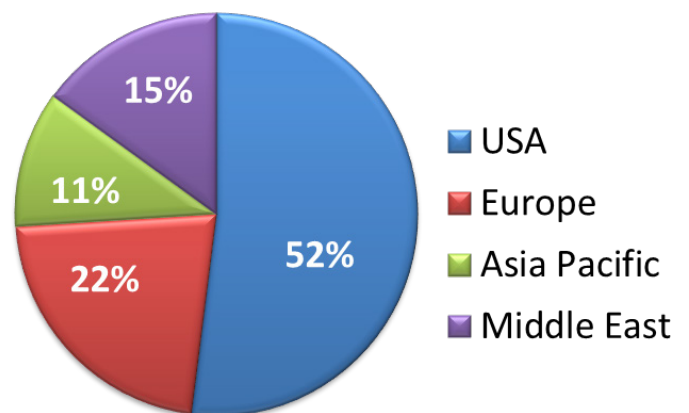
How to educate others about your rare disease

Dana Mauro, National Ataxia Foundation, USA

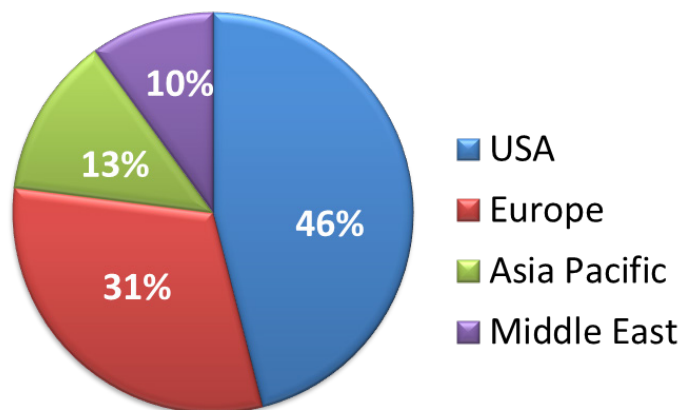
Demographics of Speakers



Demographics of Delegates



Continent Demographics



Our conference is surrounded with Doctors, Deans, Professors, Students, Young researchers, Health Practitioners, Relevant Association and Societies, Business professionals all over the globe, from which most of them are decision makers.

Illustrated in Figure 1, 2 and 3 are the recent Demographics for Speakers, Delegates and attendees from the Continents.

Rare Diseases and Orphan Drugs, past Speakers & Delegates

Abdulaziz Aldawood, King Saud Bin Abdulaziz University,
Saudi Arabia

Alan B. Moy, The John Paul II Medical Research Institute and
Cellular Engineering Technologies, USA

Alan Gilstrap, Akcea Therapeutics, USA

Alastair Kent, Genetic Alliance, UK

Alba Ancochea, EURORDIS, Spain

Alice Abdel Aleem, Weill Cornell Medical College, Qatar

Amrik Sahota, Rutgers University, USA

Anne Marie Finley, Biotech Policy Group LLC, USA

Annemieke Aartsma-Rus, Leiden University Medical Center,
The Netherlands

April Weir Hauptman, InClin, USA

Ashwani K Singal, University of Alabama at Birmingham, USA

Aya Narita, Tottori University Hospital, Japan

Charles Du Mond, InClin, USA

Courtney Smith, Colpitts Clinical, USA

Dana Mauro, National Ataxia Foundation, USA

Daniel Cohen, Pharnext , France

Danilo A Tagle, National Institutes of Health, USA

David Dongliang Ge, Apostle Inc. , USA

Deanna Laurain, InClin, USA

Diego-Abelardo Alvarez-Hernandez, Anahuac University,
Mexico

Dung-Fang Lee, The University of Texas Health Science Center
at Houston, USA

Duygu Kuyuncu Irmak, INC Research, Turkey

E. Dewsnap, Epilepsy Association of Utah, USA

Fahd Al Mulla, Kuwait University, Kuwait

Fernando Ferrer, Multinational Partnerships LLC, USA

Gail Adinamis, GlobalCare Clinical Trials, LLC, USA

Gayathri Balasubramanian, Focus Scientific Research
CenterPhamax, India

George Faurot, InClin, USA

Gianluca Colella, Rizzoli Orthopaedic Institute, Italy

Godwin Oligbu, St Georges Hospital London, UK

Harsha K Rajasimha, George Mason University, USA

Irene Tan, Temple School of Medicine, USA

Isabelle Windheuser, University Hospital Bonn, Germany

Jacques P Tremblay, Universite Laval, Canada

Jacques P. Tremblay, Department of Molecular Medecine,
Université Laval, Québec

Janel Long-Boyle, University of California San Francisco, USA

Jason Wan, National Institutes of Health, USA

Jianmeng Chen, Food and Drug Administration, USA

John A Mauro, National Ataxia Foundation, USA

John Leaman, Selecta Biosciences, USA

Jong Wook Chang, Samsung Medical Center, South Korea

Karen S. Ho, Lineagen, Inc, USA

Karren Williams, Akcea Therapeutics, USA

Katarína Šimeková, University Hospital in Martin, Slovak
Republic

Kei Kishimoto, Selecta Biosciences, USA

Kim Frieze, InClin, USA

Larissa Wester, University Hospital Bonn, Germany

Laura Sunderlin, Beazley, USA

Laxminarayan Bhat, Reviva Pharmaceuticals Inc, USA

Laxminarayan Bhat, Reviva Pharmaceuticals, Inc, USA

Lei He, Food and Drug Administration, USA

Leonardo Cano, Universidad Católica Santiago de Guayaquil,
Ecuador

Lisa Baumbach- Reardon, Translational Genomics Research
Institute, USA

Maite A. Castro, Universidad Austral de Chile, CHILE

Manoj Krishnan, Duke-NUS, Singapore

Maria Shkrob, Elsevier, USA

Marise Abdou, Abo El-Rish Children's Hospital, Cairo, Egypt

Mark Landiak, Patient Advocate, Foundation for Sarcoidosis
Research, USA

Megan O'Boyle, Phelan-McDermid Syndrome Foundation, USA

Michael S Wolfe, University of Kansas, USA	Shipra Agrawal, The Research Institute at Nationwide Childrens Hospital, USA
Mingli Jiao, Harbin Medical University, China	Shmuel Prints, Clalit Health Service, Israel
Moamen Al Zoubi, Advocate Illinois Masonic Medical Center, USA	Stefano Giacomini, Rizzoli Orthopedic Institute, Italy
Moji C. Adeyeye, Roosevelt University, USA	Stella Blackburn, QuintilesIMS, UK
Nadia Ameen, Yale University School of Medicine, USA	Stephen P. Arneric, Critical Path Institute, USA
Nasir Ali Afsar, Alfaisal University College of Medicine, KSA	Stephen Shrewsbury, Fortuna Fix, USA
Ndiaye Mady, Faculty of Medicine of Thiès, Nigeria	Stephen Smolinski, Selecta Biosciences, USA
Patrick J Tighe, University of Nottingham, U.K	Sujatha Kannan, Johns Hopkins University School of Medicine, USA
Philippe Jouvett, Sainte-Justine Hospital University of Montreal, Canada	Tatjana Michel, University Hospital Tuebingen, Germany
Prasanth Puthanveetil, Roosevelt University College of Pharmacy, USA	Thomas C Chen, Keck School of Medicine of USC, USA
Ramune Sepetiene, Lithuanian University of Health Sciences, Lithuania	Timothy Coté, Coté Orphan, USA
Rashmi Gopal-Srivastava, NCATS, NIH, USA	Tony Zbeidy, Orphan-Europe, France
Rob W J Collin, Radboud University Medical Center, Netherlands	Wei Zheng, National Institutes of Health, USA
Róbert Rosol'anka, University Hospital in Martin, Slovak Republic	Xavier Paoli, Pharnext, France
Romina Ortiz, Rare Genomics Institute, USA	Xiaolan Zhang, Sarepta Therapeutics, USA
Ryan Clift, InClin, USA	Yingjun Xie, The Third Affiliated Hospital of Guangzhou Medical University, China
Samuel Ayoola Abati, Iagos university, Nigeria	Yoko Sato, National Defense Medical College, Japan
Sara Tylosky, Farmacon, USA	Yolande van Bever, Erasmus Medical Centre, The Netherlands
Serge Braun, Scientific Director, AFMTelethon, France	Yong Moon Choi, Bio-Pharm Solutions Co., Ltd, South Korea
Shin'ichi TAKEDA, National Institute of Neuroscience, Japan	Yunzhao Ren, Food and Drug Administration, USA
	Yusuf Hovsep Eken, Elkerliek ziekenhuis, Netherland



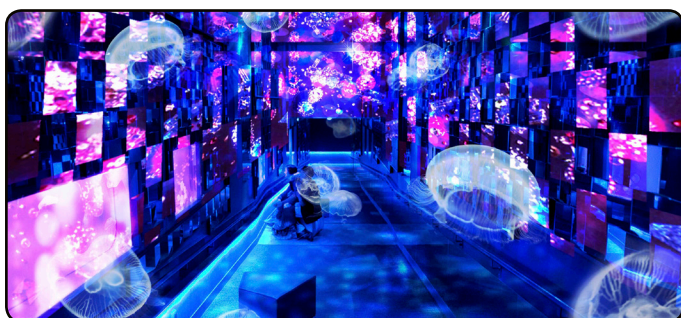
WHAT YOU CAN EXPECT



GLIMPSES OF RARE DISEASES CONFERENCE



PLAN YOUR TRIP @ TOKYO



CONTACT US

For Queries

Natalia Jones | Program Manager
Email: rarediseases@conferenceseries.net
Toll Free: +44-800-014-8923

Hosted by Infectious Diseases conferences
47 Churchfield Road, London, UK, W3 6AY
Toll Free: +44-800-014-8923