

TAKING STEPS TOWARDS ending the diagnostic odyssey

APRIL 19-20, 2023

FINAL AGENDA

DAY ONE	Wednesday, April 19, 2023	8:00 am to 4:00 pm PDT	
MORNING	MORNING FOCUS: rapid Whole Genome Sequencing™		
8:00 am	Registration/Exhibit Hall/Breakfast		
9:00 am	Welcome	Charlotte Hobbs, MD, PhD VP and Chief Research & Clinical Management Officer, Rady Children's Institute for Genomic Medicine (RCIGM) Stephen Kingsmore, MD, DSc President & CEO, RCIGM	
9:15 am	Genomic Medicine for III Neonates and Infants: Lessons Learned from the GEMINI Study	Kristen Wigby, MD Assistant Professor of Pediatrics, University of California, San Diego / Physician Investigator, RCIGM	
9:45 am	Streamlining Clinical Genetic Testing: The Promise of Long-Read Sequencing	Danny Miller, MD, PhD Assistant Professor of Pediatrics & Genome Sciences, University of Washington School of Medicine	
10:15 am	BREAK – Exhibit Hall Open		
10:45 am	Infant Mortality Associated with Monogenic Disorders	Liana Protopsaltis, MS, CGC Research & Project Manager, RCIGM	
11:15 am	Family Panel	Moderator: Jerica Lenberg, MS, CGC Licensed Clinical Genetic Counselor, RCIGM Patient families	
Noon	Lunch & Networking		
	Presentation by lunch sponsor Fabric Genomics: Rapid Diagnosis in the NICU and PICU: Using AI to Overcome the Challenges of Genomic Variant Interpretation.	Mark Yandell, PhD Professor of Human Genetics, Co-Director Utah Center for Genetic Discovery School of Medicine, University of Utah	
AFTERNOON FOCUS: Newborn Genomic Screening			
1:00 pm	Introduction to Newborn Genomic Screening	Stephen Kingsmore, MD, DSc President & CEO, RCIGM	
1:05 pm	Genomics England Newborn Genomes Programme	David Bick, MD, PhD Principal Clinician, Genomics England	
1:40 pm	Genome Sequencing in Newborns: A Public Health Perspective	Laurie Smith, MD, PhD RCIGM Consultant	

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		Gunter Scharer, MD Chief Medical Officer, RPRD Diagnostics
		Jennifer Schleit, PhD, FACMG Laboratory Director, RCIGM
		Moderator: Mei Baker, MD, FACMG Professor of Pediatrics, University of Wisconsin-Madison / Newborn Screening Laboratory Director, Wisconsin State Laboratory of Hygiene
2:15 pm	BREAK – Exhibit Hall Open	
2:45 pm	The Path to Universal Newborn Sequencing	Robert Green, MD, MPH Professor of Medicine (Genetics), Harvard Medical School and Mass General Brigham
3:15 pm	Challenges and Opportunities in Screening Newborns and Young Children for Complex Conditions using Genetic Risk Scores	Holly Peay, PhD, MS, CGC Early Check / Senior Research Scientist, RTI International
3:45 pm	Closing Remarks	Drs. Stephen Kingsmore and Charlotte Hobbs, RCIGM
4:30 – 7:00 pm	Reception/Networking Heavy hors d'oeuvres and cocktails Brief sponsor presentation: Illuming Opening	Chric Kunard, Sr. Director, Open Innovation Customer
	Brief sponsor presentation: Illumina Opening Remarks and Role in Whole Genome Sequencing	Chris Kunard, Sr. Director, Open Innovation Customer Collaboration, Illumina

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DAY TWO	Thursday, April 20, 2023	8:00 am to 4:30 pm PDT
MORNING	FOCUS: Newborn Genomic Screening	
7:30 am	Registration/Exhibit Hall/Breakfast	
8:00 am	Welcome	Charlotte Hobbs, MD, PhD VP and Chief Research & Clinical Management Officer, RCIGM
8:05 am	Screen4Care EU-IMI Project: Accelerating Rare Disease Diagnosis by Genetic Newborn Screening and Digital Technology	Alessandra Ferlini, MD, PhD Professor & Head of the Medical Genetics Unit, University o Ferrara / Project Scientific Coordinator, Screen4Care Nicolas Garnier, PhD EFPIA Lead, Screen4Care / Head of Patient Advocacy, Pfizer Global Product Development
8:40 am	Newborn Genomic Screening in Greece	Petros Tsipouras, MD Scientific Director, First Steps Greece / CEO, PlumCare
9:15 am	The BeginNGS Perspective	Tom DeFay, PhD Deputy Chair, BeginNGS Consortium / Deputy Head of Diagnostics Strategy & Development, Alexion
9:50 am	Panel: Newborn Genomic Screening Programs Across the Globe	David Bick, MD Principal Clinician, Genomics England Alessandra Ferlini, MD, PhD Professor & Head of the Medical Genetics Unit, University o Ferrara / Project Scientific Coordinator, Screen4Care Nicolas Garnier, PhD EFPIA Lead, Screen4Care / Head of Patient Advocacy, Pfize Global Product Development Holly Peay, PhD, MS, CGC Senior Research Scientist, RTI International

	and Mass General Brigham
	Petros Tsipouras, MD Scientific Director, First Steps Greece / CEO, PlumCare
	Moderator: Tom DeFay, PhD Deputy Chair, BeginNGS Consortium / Deputy Head of Diagnostics Strategy & Development, Alexion
BREAK – Exhibit Hall Open	
Unraveling the Diagnosis of ENPP1 Deficiency	Catherine Nester, RN, BSN VP, Physician & Patient Strategies, Inozyme Pharma
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Robert Green, MD, MPH Professor of Medicine (Genetics), Harvard Medical School

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11:30 am	Panel: Addressing the Diagnostic Odyssey Through Pharma Participation in NBS-by-WGS programs	Catherine Nester, RN, BSN VP, Physician & Patient Strategies, Inozyme Pharma Charlotte Chanson, MSc Senior Director, Global Diagnostics/Newborn Screening, Orchard Therapeutics Steve Rodems, PhD VP, Research, Travere Therapeutics Tom DeFay, PhD Deputy Chair, BeginNGS Consortium / Deputy Head Diagnostics Strategy & Development, Alexion Nicole Miller, PhD VP, Molecular Diagnostics, Ultragenyx Moderator: Walter Kowtoniuk, PhD Venture Partner, Third Rock Ventures
12:10pm	Lunch & Networking	
AFTERNOON FOCUS: Diagnostic rWGS [®] , Effectiveness & Reimbursement		
1:10 pm	Welcome Back	Charlotte Hobbs, MD, PhD VP and Chief Research & Clinical Management Officer, RCIGM
1:15 pm	Lightning Talks	
	DNA Methylation Detection Using PacBio Sequencing	Matthew Bainbridge, PhD Assistant Director of Translational Research, RCIGM
	Improving Outcomes of Critical Congenital Heart Disease Through Genome-Informed Management	Nathaly Sweeney, MD, MPh, MS Assistant Clinical Professor of Pediatrics, University of California, San Diego / Physician Investigator, RCIGM
	Pediatric Genomic Psychiatry: Precision Medicine for Youth Mental Health	Aaron Besterman, MD Assistant Clinical Professor of Psychiatry, University of California, San Diego / Physician Investigator, RCIGM
1:45 pm	Next-Generation Advocacy Driving Forward Collaborative Progress in Rare Disease	Charlene Son Rigby, MBA CEO, Global Genes
2:15 pm	Bears, Manatees, and Deer, Oh My! – Michigan's Implementation of rWGS Affirms Strong Clinical and Economic Utility	Caleb Bupp, MD FACMG Assistant Clinical Professor of Pediatrics and Human Development, Michigan State University / Geneticist, Division Chief, Corewell Health & Helen DeVos Children's Hospital
2:45 pm	Economic Utility of First Line rWGS [®] in Commercial Payor Model Empowers Coverage and Implementation	Christy Moore, MS, CGC Genetics Program Manager, Blue Shield of California
3:15 pm	BREAK – Exhibit Hall Open	

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3:45 pm	Panel: rWGS — Michigan's Journey to Coverage and Implementation	Andrea Scheurer-Monaghan, MD Clinical Assistant Professor of Pediatrics, Western Michigan University / Neonatologist, Southwestern Michigan Neonatology, P.C.
		Laura Appel Executive VP, Michigan Health & Hospital Association
		David Dimmock, MD Staff Physician, Valley Children's Hospital
		Moderator: Caleb Bupp, MD FACMG Assistant Clinical Professor of Pediatrics and Human Development, Michigan State University / Geneticist, Division Chief, Corewell Health & Helen DeVos Children's Hospital
4:25 – 4:40 pm	Closing Discussion	Drs. Stephen Kingsmore and Charlotte Hobbs, RCIGM