

Invited Sessions 2015-2018

Invited Sessions 2018:

- [Achieving Genomics Literacy for the Masses: Providing Meaningful Education for Multiple Audiences](#)
- [Advances in Unraveling the Genetic, Molecular, and Clinical Complexity of Autism](#)
- [Chromatin Dysregulation in Neurodevelopmental Disorders](#)
- [Clinical Spotlight: ASHG/ESHG Building Bridges: Prenatal Genetic Testing: Recent Advances and Current Challenges](#)
- [Clinical Spotlight: Realizing the Promise of Common Genomic Variation in Rare and Common Disease: Clinical Implementation of Polygenic Risk Scores](#)
- [Duty to Recontact: Do We Need to Reconsider?](#)
- [Exploring the Medical Phenome: The New Frontier of Genetic Discovery](#)
- [Impact of Natural Selection on the Genetic Architecture of Complex Traits](#)
- [Innovative Strategies to Support the Education of Health Care Providers in Genomic Medicine](#)
- [Large Scale Functional Annotation of Variants of Uncertain Significance](#)
- [New Developments in Mendelian Randomization](#)
- [Novel Insights in Aging: Examining the Interface Between Genetics and the Environment](#)
- [Silent Genomes: Indigenous-led Initiatives on Addressing Equity in Genomics Health Care and Research](#)
- [Technical Approaches and Guidelines for Protecting Privacy of Genetic Data](#)
- [The Genetics of Human Proteomes](#)
- [Uncovering Missing Heritability in Mendelian Diseases: Lessons from Inherited Eye Diseases](#)
- [Understanding Tumor Heterogeneity from Single Cell Sequencing of Genomes, Transcriptomes and Epigenomes](#)
- [What's Sex Got to Do with It: Sexual Dimorphism in Human Disease](#)

Invited Sessions 2017:

- [Analysis of Cancer Genome Variation Using Long-read Sequencing](#)
- [Clinical Spotlight: VUS-busters: Cutting-edge Strategies for Interpreting Variants in Clinical and Research Sequencing](#)
- [Data Sharing, Analysis, and Tools to Catalyze Translation from Genomic to Clinical Knowledge](#)
- [Diversity Matters: Scientific and Ethical Strategies for Achieving Representation in Genomics](#)
- [Emerging Challenges in Complex Traits: One Locus, but Multiple Variants, Genes, and Tissues](#)
- [FACEing the Challenge: Advances in Our Understanding of Facial Development and Disease](#)
- [Novel Insights into Human Brain Evolution from Advanced Genomics](#)
- [Rigor and Reproducibility in Genetic Research](#)
- [Solving the Unsolved: Systems to Facilitate the Discovery of Novel Rare Disease Genes from Genomic Sequencing](#)
- [ASHG/ESHG Building Bridges: Zika: From Virus to Host Response to Vector Control](#)
- [Biology at Single Cell Resolution: Understanding Cell-type-specific Responses in Development and Disease](#)
- [Clinical Spotlight: Translational Genomics: Psychological and Health Behavior Research Outcomes](#)
- [Dosage-sensitive Sex-linked Genes: Role in Aneuploidy and Cancer](#)
- [High-throughput Sequencing of Adaptive Immune Receptors: Insights into Disease Mechanisms and Treatment](#)
- [Illuminating Somatic Mutations in Neurological Development and Disease](#)
- [Layers of Complexity: Dissecting the Etiology of Mendelian Diseases Characterized by Extreme Heterogeneity](#)
- [Metabolomic Perspectives in a Genomic Era: Garrod's Inborn Errors and Warburg's Theory of Cancer](#)
- [Using Controls from External Studies: Issues, Methods, and Successes](#)

Invited Sessions 2016:

- [Augmenting and Interpreting Genomic Data Using Tissue- and Cell-type-specific Networks](#)
- [Celebrating the Centenary of R.A. Fisher's "The Correlation Between Relatives on the Supposition of Mendelian Inheritance"](#)
- [CRISPR: A New Paradigm for Forward Human Genetics](#)
- [Diagnostic Functional Genetics, the Essential Next Step in NGS Translation to the Bedside](#)
- [Education in Action: Meeting the Challenges of 21st Century Genetics](#)
- [From GWAS and Mendelian Genes to Therapeutic Drug Targets](#)
- [Clinical Spotlight: Models and Approaches for Population-based Screening of Cancer Susceptibility Genes](#)
- [The Role of DNA Repair in Genomic Variation, Instability, and Human Disease](#)
- [Unusual Suspects: A Legal Line-Up Beyond GINA](#)
- [ASHG/ESHG Building Bridges Session: Navigating the Myriad Career Paths in Human Genetics](#)
- [Applying Multi-omics to Complications of Solid-organ Transplantation](#)
- [Beyond DNA Fingerprinting: Novel Developments in Forensics](#)
- [Evaluating Effectiveness of Traditional and Novel Methods of Genetic/Genomic Counselling](#)
- [Clinical Spotlight: Gene Discovery, Genetic Counseling, and Clinical Care of Patients with Inherited Retinal Diseases](#)
- [Genome Editing: What Implications and Obligations Does this Emerging Technology Create?](#)
- [How Natural Selection Shapes the Modern Human Genome: Methods and Examples](#)
- [Innovative Approaches to Co-ordinated Genomics Education for all Healthcare Professionals in England](#)
- [Structural Genomics: Integrating Analysis of Chromosome Conformation in Disease and Diagnosis](#)

Invited Sessions 2015:

- [Building the Genetic and Genomic Atlas of Gynecologic Health](#)
- [Cancer Genetics in the Genomics Era](#)
- [Epilepsy Genetics: Exomes, Mechanisms, and Interventions](#)
- [Human Phenotypes for Researchers, Clinicians and Patients](#)
- [Looking Beyond the Genes: Non-coding Mutations and 21st Century Disease Genetics.](#)
- [Maternal Age and Recombination: Risks to Aneuploidy](#)
- [Mendelian Disorders of the Epigenetic Machinery: Genetic Disorders with Epigenetic Consequences.](#)
- [Policy Challenges Affecting Clinical Integration of Next-Generation Sequencing: Advancing Toward Resolution](#)
- [Secure, Efficient, and Scalable Computational Genetics via Summary Statistics](#)
- [When You Know You've Found the One: Fine-Mapping GWAS Hits to a Single Variant](#)
- [Gene Editing/Rewriting the Genome: Moving from Association to Biology and Therapeutics](#)
- [Genetic Control of the Microbiome](#)
- [Integrating Genomes and Transcriptomes to Understand Human Disease](#)
- [Life Beyond Additive Variance](#)
- [Multiplexed and Multimodal Experimental Dissection of Genetic Variants](#)
- [Optimizing Clinical and Molecular Characterisation and Management in Skeletal Dysplasias: An Exemplary Model for Rare Genetic Diseases.](#)
- [Research Partners, Not Subjects: Engaging Indigenous Peoples in Genetics](#)
- [The Landscape of de novo Point Mutations in the Human Genome: How Many, Where, When and Why?](#)
- [Translating Genomic Knowledge into Clinical Practice](#)
- [Understanding Disease Pathogenesis: A Grand Challenge for Model Organisms](#)