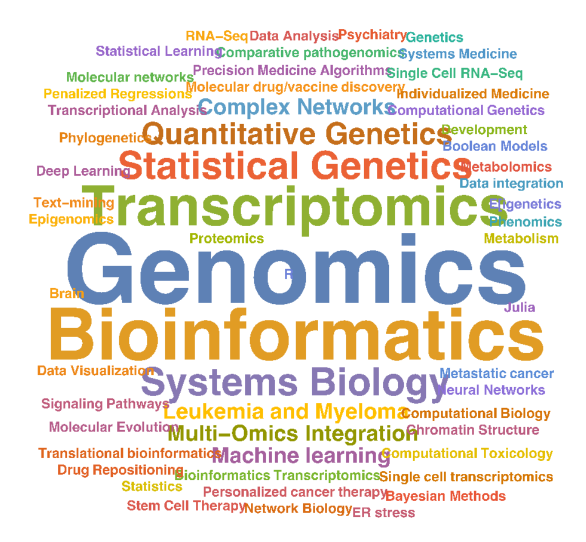
Systems Computation Omics

Michigan State University



Working Draft: 7/1/2019

Mission Statement

Systems approaches are now facilitated with developments in computational statistical methods, and the availability of multiple omics data as indispensable parts of modern biological research. Michigan State University has multiple faculty across colleges and departments involved in numerous ground-breaking research projects in the field ranging from systems biology in basic science to systems medicine. Our common goal is to assemble an interdisciplinary group of experts, based on our research synergies, shared mathematical approaches and collaborative projects that can collaborate and can lead MSU to become a leader in Systems Computational Omics.

Mission

The goal of the Systems Computational Omics working group at Michigan State University is to utilize computational and mathematical methods to:

- Develop state-of-the-art methodology for bioinformatics and statistical omics analyses.
- Employ systems levels analyses to describe biological systems at all scales
- Create a platform for research collaboration not only between computational but also experimental groups
- Develop a robust training program for our students and postdocs

Vision

To utilize the interdisciplinary computational and mathematical expertise across Michigan State University's campuses, to enhance collaboration, development of novel methodology and provide an interactive environment making MSU a leader in computational approaches, both for research as well as training of the next generation of computational scientists.

Faculty Membership

Sudin Bhattacharya, PhD

University of Michigan, Mechanical Engineering, PhD 2006

Assistant Professor in Departments of Biomedical Engineering and Pharmacology & Toxicology http://sbhattac.msu.domains



- Research Interests: The Bhattacharya Lab uses various computational methods to
 understand the signaling and transcriptional regulatory networks that underlie the
 determination of cell fate, and these networks' perturbation by environmental pollutants. We
 are integrating diverse genomic data sets to map and model transcriptional regulatory
 networks and their environmental perturbation in the immune system and the liver.
- Research Methods:
 - o Transcriptome Analysis
 - o Regulatory Network Reconstruction
 - o Single-cell Analysis

Keywords: Genomics; Bioinformatics; Transcriptomics; Systems Biology; Computational Toxicology.



Gustavo de los Campos, PhD

University of Wisconsin-Madison, Quantitative Genetics and Agricultural & Applied Economics, PhD 1997

Associate Professor Departments of Epidemiology & Biostatistics and Statistics & Probability https://quantgen.github.io

- Research Interests: Quantitative, Statistical and Computational Genetics. Prediction of Complex Traits using DNA information and other omics.
- Research Methods:
 - o High-Dimensional Regression
 - o Bayesian and Penalized Methods
 - o Multivariate Analysis
 - o Multi-Omic methods

Keywords: Genomics; Computational Genetics, Statistical Genetics, Quantitative Genetics, Bayesian Methods, Penalized Regressions, R, Julia

Christina Chan, PhD

University of Pennsylvania, Chemical and Biochemical Engineering, PhD 1990

George W. Bissell Professor in Department of Chemical Engineering and Materials Science (and Biochemistry and Molecular Biology) https://www.egr.msu.edu/changroup/

 Research Interests: Her laboratory focuses on understanding the signaling and regulatory mechanisms that are altered in diseases, such as obesity, Alzheimer's disease, and cancer. They develop and



apply metabolic engineering, bioinformatics, and systems biology approaches in combination with biochemical, biophysical, and molecular biology measurements and animal studies to identify targets and disease biomarkers. Her group also is developing polymeric-based drug delivery and tissue engineering platforms to modulate these pathways for treating these diseases.

Research Methods:

- o CRISPR, RT-PCR, Western blot analysis, protein purification
- o Cell culture
- o Network Analysis
- o Regulatory Network Reconstruction

Keywords: ER stress; Transcriptional analysis; Signaling pathways; Stem cell therapy

Bin Chen, PhD

Indiana University at Bloomington, 2012, Informatics
Assistant Professor in Departments of Pediatrics and Human
Development, Pharmacology and Toxicology
http://binchenlab.org

• Research Interests: Rapidly decreasing costs of molecular measurement technologies not only enable profiling of disease sample molecular features (e.g., transcriptome, proteome, metabolome) at different levels (e.g., tissues, single cells), but also enable measuring of molecular

signatures of individual drugs in clinically relevant models. My lab is interested in developing computational methods to relate diseases to potentially efficacious drugs through various molecular features. The lab is also interested in using deep learning and single cell data to discover new therapeutics.

• Research Methods:

- o Large-scale disease and drug molecular data integration and mining
- o RNA-Seq data analysis
- o Deep learning
- o Genomic data analysis

Keywords: RNA-Seq, Single cell RNA-Seq, Drug repositioning, Deep learning, Personalized cancer therapy, Metastatic cancer, Translational bioinformatics

Yuehua Cui, PhD

University of Florida, Statistics, PhD 2005

Professor in Department of Statistics and Probability https://www.stt.msu.edu/~cui/

 Research Interests: Statistical genetics/genomics, statistical machine learning, longitudinal/functional data analysis, statistical applications in applied science.

Research Methods:

- o Gene-gene and gene-environment interactions
- o Casual mediation analysis
- o Multi-level omics data integration
- o Gene network inference



Keywords: Genetics, Genomics; Bioinformatics; Epigenetics; Statistics

Erich Grotewold, PhD

University of Buenos Aires, Argentina, Chemistry & Biochemistry, PhD 1988

Professor in Department of Biochemistry & Molecular Biology https://eglab.osu.edu

• Research Interests: Gene regulatory networks, mechanisms of gene regulation, plant metabolic engineering, trafficking of specialized metabolites

Research Methods:

- o Transcriptome Analysis
- o Chromatin immunoprecipitation High Throughput Sequencing (ChIP-Seq)
- DNA-Affinity Purification High Throughput Sequencing (DAP-Seq)
- o Cap Analysis Gene Expression (CAGE)
- Regulatory Network Reconstruction
- o Metabolic Profiling
- Analysis of Protein-Protein and Protein-DNA Interactions

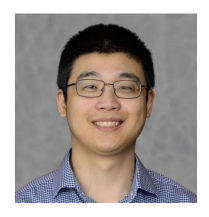
Keywords: Metabolism; Genomics; Bioinformatics; Transcriptomics; Systems Biology

Wen Huang, PhD

University of Wisconsin-Madison, Dairy Science, PhD 2011

Assistant Professor in Department of Animal Science https://msu.edu/~huangw53/index.html

 Research Interests: Genetic basis of complex quantitative traits, context dependent genetic effects, genetic networks



Research Methods:

- o Statistical genetics
- o Bioinformatics
- o Next generation sequencing
- o Molecular genetics

Keywords: Quantitative Genetics; Statistical Genetics; Genomics; Bioinformatics



Arjun Krishnan, PhD

Virginia Tech, Genetics, Bioinformatics, and Computational Biology, PhD 2010

Assistant Professor in the Department of Computational Mathematics, Science and Engineering, and the Department of Biochemistry and Molecular Biology

https://www.thekrishnanlab.org

Research Interests: We develop computational data-driven approaches to build predictive
models that help us link genes and cellular mechanisms to various aspects of health and
disease.

Research Methods:

- Genomics & computational biology
- Applied statistical/machine learning
- Integrative analysis of large-scale genomics and biomedical data
- **o** Genome-wide molecular networks
- Age-specificity and sexual dimorphism in health and disease
- Cross-species models for human traits and diseases
- o Genetic heterogeneity of complex disease and precision medicine

Keywords: Transcriptomics; Text-mining; Machine learning; Molecular networks; Data integration

Qing Lu, PhD

Case Western Reserve University, Statistical Genetics, PhD 2008 Associate Professor in Department of Epidemiology and Biostatistics

• Research Interests: My research interest is primarily in statistical genetics and statistical learning. One area of research is the development of statistical learning methods and software for risk prediction analysis. In parallel with this line of research, I am also interested in developing and applying of new statistical methods for high-dimensional genetic association analysis.



Research Methods:

- o Neural Networks and Deep Learning
- o Tree and Random Forest
- o U-statistics
- o Functional Data Analysis
- o Multi-Omic Methods

Keywords: Statistical learning; Statistical Genetics.

George Mias, PhD

Yale University, Physics, PhD 2007
Assistant Professor of Biochemistry and Molecular Biology
https://georgemias.org

• Research Interests: Using interdisciplinary computational and experimental methods to explore the integration of existing and developing omics technologies and their application in systems precision medicine. We monitor individuals transitioning through various physiological states, from

health to disease (asthma, leukemia, immune disorders, etc), towards universal individualized medicine. A key aspect is the study of longitudinal dynamics from omics data, using complex networks and geometric methods.

• Research Methods:

- o Individualized multi-omics profiling
- o Time series Analysis
- o Combinatorial optimization of drug combinations
- o Algorithms for Data Analysis and Integration
- o Machine Learning

Keywords: Systems Medicine; Systems Biology; Individualized Medicine; Genomics; Bioinformatics; Transcriptomics; Metabolomics; Proteomics; Complex Networks; Leukemia and Myeloma; Multi-Omics Integration.

Carlo Piermarocchi, PhD

Swiss Federal Institute of Technology Lausanne, Switzerland, Physics, PhD 1998

Professor in the Department of Physics and Astronomy

 Research Interests: My research focuses on developing novel mathematical models to describe cellular processes in normal and hematological cancer cells, and predict their response to drugs and drug combinations. Our systems biology models integrate different types of omics with in vitro drug screening data and use concepts from the theory of complex networks and nonlinear dynamics.



Research Methods:

- o Regulatory network reconstruction
- o Neural network models of cell dynamics
- o Time series analysis
- o Combinatorial optimization of drug combinations
- o Algorithms for the analysis of single-cell transcriptomics

Keywords: Genomics; Bioinformatics; Transcriptomics; Precision Medicine Algorithms; Neural Networks, Machine Learning, Complex Networks, Boolean Models; Leukemia and Myeloma; Single cell transcriptomics.



The University of Akron, Integrative Biosciences, PhD 2014

Assistant Professor in Department of Pediatrics and Human Development with second appointment in Pharmacology and Toxicology

 Research Interests: The Prokop lab focuses on integrative biosciences for understanding genes and genetic variants, utilizing computational tools to generate testable hypotheses for genomic variants (coding and noncoding) and protein function that are then used to guide wet lab experiments into disease mechanisms and pharmacological intervention.



- o Transcriptome Analysis
- o Linkage Disequilibrium
- Genome Wide Association Studies

Systems Computational Omics, Michigan State University

- o Whole Genome Sequencing
- o Codon Selection
- o Induced pluripotent stem cells (iPSC)
- o CRISPR
- o CETCh-Seq
- o Molecular Dynamics Simulations
- o Protein Modeling

Keywords: Genotype-phenotype; rare variant analysis; Transcription; GWAS; Clinical Genomics

Janani Ravi, PhD

Virginia Tech, Genetics, Bioinformatics & Computational Biology, PhD 2006

Sr Research Associate, Pathobiology & Diagnostic Investigation, CVM. https://jananiravi.github.io/

Research Interests: My research focuses on developing novel mathematical models to describe cellular processes in normal and hematological cancer cells, and predict their response to drugs and drug combinations. Our systems biology models integrate different types of amino with in vitro drug corresping desired.



- integrate different types of omics with in vitro drug screening data and use concepts from the theory of complex networks and nonlinear dynamics. Unraveling molecular mechanisms common between genetic and infectious diseases, to guide targeted drug repurposing
 - o Identifying genomic signatures, functions, and mechanisms relevant to host-pathogen interactions, to enable understanding and guide diagnosis
 - o Developing tools and resources for seamlessly relating sequence, structure, and function of genes/proteins in poorly characterized pathogens to enable and guide better prevention, diagnostic and treatment regimen
 - o Mathematical/statistical modeling of signaling and regulatory networks: host-pathogen interactions, metabolic pathways, and cellular processes.

Research Methods:

- o Programming: R, Unix/Linux (MATLAB, Python, C/C++)
- o Data science: data analysis & visualization
- o Genomic/Transcriptomic analyses: incl. transcriptional regulatory networks
- o Phylogenetics and molecular evolution: *incl.* sequence-structure-function prediction
- o Dynamical systems and mathematical modeling: *incl.* dynamic Bayesian networks; large quantitative models; deterministic/stochastic modeling with bifurcation analysis

Keywords: Computational Biology; Bioinformatics; Phylogenetics & Molecular Evolution; Comparative pathogenomics; Molecular drug/vaccine discovery; Data Analysis & Visualization; Genomics/Transcriptomics



University of BC, Mathematics, PhD 1987

Associate Professor in Department of BME and in Neuroscience Program https://msu.edu/~reimersm/

o **Research Interests:** Working on i) characterizing functional differences reflected in gene expression differences between psychiatric patients and normals; ii) genetic influences on psychiatric disorders; iii)

Systems Computational Omics, Michigan State University

effects of pre-natal infection on mouse brain transcriptome; iv) single cell transcriptome characterizations of key cell types in mouse and human brain; v) simulation of ways genotypes can give rise to behavioral phenotypes.

• Research Methods:

- o Transcriptome Analysis
- o GWAS accounting for Linkage Disequilibrium
- o Characterizing PPI Networks of GWAS-implicated sets
- o Simulation of regulatory and behavioral systems

Keywords: Genomics; Bioinformatics; Transcriptomics; Brain; Psychiatry; Development.

Shin-Han Shiu, PhD

University of Wisconsin-Madison, Botany, PhD 2010

Professor of Plant Biology, and Computational Mathematics, Science and Engineering

http://shiulab.plantbiology.msu.edu/index.php

Research Interests: The overall goals for our research program are to elucidate: (1) the molecular evolutionary processes that shape eukaryotic genome content, (2) the distinguishing properties of genomic features, particularly those allowing the classification of functional and non-functional sequences, and (3) the mechanisms that underlie the translation of genomic information to molecular, physiological, and morphological phenotypes.

• Research Methods:

- o Evolutionary and population genetics
- o Functional, comparative and phylo-genomics
- o Multi-omics data integration
- o Machine learning
- o Genomic prediction

Keywords: Evolution, Genetics, Bioinformatics, Computational biology, Plant biology.



Associate Professor in Department of ABC https://www.canr.msu.edu/people/dr juan steibel

 Research Interests: Working on methods for Genomic prediction and genome-wide association in structured populations. Modeling of social genetic effects. Methods for phenotyping individuals in groups. E.g. group-housed pigs.

Research Methods:

- o Transcriptome Analysis
- o Linkage Disequilibrium
- o Genomic prediction
- o Social genetic effects
- o Image/video analyses

Keywords: Genomics; Quantitative Genetics; Statistical Genetics; Phenomics.

Ana Vazquez, PhD

University of Wisconsin-Madison, Quantitative Genetics, PhD 2010

Assistant Professor of Epidemiology and Biostatistics https://quantgen.github.io

Research Interests: My research focuses on the development use and implementation of omic-based statistical methods for human health complex traits, including cancer, obesity, exercise and among other health traits. My previous research included studies with generalized linear mixed models (GLMM) with correlated random effects, from which I co-authored an R-package that fits such models.



- o High-Dimensional Regression
- o Generalized Linear Mixed Models
- o Multivariate Analysis
- o Multi-Omic methods

Keywords: Genomics; Computational Genetics, Statistical Genetics, Quantitative Genetics



Jianrong Wang, PhD

Georgia Institute of Technology, Bioinformatics, Ph.D. 2012
Assistant Professor in Department of Computational Mathematics, Science and Engineering

• Research Interests: Develop machine learning and statistical algorithms to construct context-dependent regulatory networks based on large-scale multi-omics dataset. Leverage the constructed regulatory networks to decode the genetic basis of complex human disease mechanisms. Build integrative models to obtain insights on systems biology of gene regulation, especially on

3D chromatin structures, epigenetics and transcription factors.

Research Methods:

- Bayesian statistics
- Machine learning
- Probabilistic graphical models
- Network reconstruction
- Systems genetics
- Next-generation sequencing (NGS) data processing and analysis
- Regulatory grammar prediction

Keywords: Genomics, Network Biology, Bioinformatics, Systems Biology, Epigenomics, Multi-omics integration, Chromatin Structure