BORAN GAO

● 4331 SPH II, 1415 Washington Heights, Ann Arbor, MI ■ (734)-546-2396

EDUCATION

Ph.D. in Biostatistics

Expected Apr 2024

University of Michigan, Ann Arbor, MI

Advisor: Dr. Xiang Zhou

Dissertation Title: Statistical Methods and Computational Tools for Genetics and Genomics Data

M.S. in Biostatistics, MPH in Epidemiology

Sep 2014 - Apr 2017

University of Michigan, Ann Arbor, MI

Bachelor of Medicine, Bachelor of Surgery

Sep 2006 - Jul 2011

Soochow University, Suzhou, China

RESEARCH INTEREST

Methods: Bayesian variable selection, mixed effects models, data integration, high-dimensional data analysis, and machine learning

Applications: Heritability and genetic correlation estimation, fine-mapping, colocalization, polygenic risk score prediction, and multi-omics integration

Data types: Genome-wide association study (GWAS), bulk RNA-seq, spatially resolved transcriptomics (SRT), perturb-seq, proteomic, and electronic health records (EHR) data

PUBLICATIONS

Peer-reviewed Publications

- 1. **Boran Gao** and Xiang Zhou (2023). MESuSiE enables scalable and powerful multi-ancestry fine-mapping of causal variants in genome-wide association studies. *Nature Genetics, in press.*
- 2. **Boran Gao**, Can Yang, Jin Liu, and Xiang Zhou (2021). Accurate genetic and environmental covariance estimation with composite likelihood in genome-wide association studies. *PLOS Genetics*, 17(1), e1009293.
- 3. Lu Wang, **Boran Gao**, Yue Fan, Fuzhong Xue, and Xiang Zhou (2021). Mendelian randomization under the omnigenic architecture. *Briefings in Bioinformatics*, 22(6), bbab322.

Manuscript under Review

1. Daiwei Zhang, Boran Gao, Qidi Feng, Ani Manichaikul, Gina M. Peloso, Russell P. Tracy, Peter Durda, Kent D. Taylor, Yongmei Liu, W. Craig Johnson, Stacey Gabriel, Namrata Gupta, Joshua D. Smith, Francois Aguet, Kristin G. Ardlie, Thomas W. Blackwell, Robert E. Gerszten, Stephen S. Rich, Jerome I. Rotter, Laura J. Scott**, Xiang Zhou**, and Seunggeun Lee** (2023+). Proteome-wide association studies for blood lipids and comparison with transcriptome-wide association studies. Under revision in PLOS Genetics. ** co-corresponding authorship.

Manuscripts in Preparation

- 1. Qidi Feng*, **Boran Gao***, Laura J. Scott**, Seunggeun Lee**, and Xiang Zhou**. Transcriptomewide association studies in lipid traits in diverse populations. * Co-first authorship, ** co-corresponding authorship.
- 2. **Boran Gao**, Zheng Li, and Xiang Zhou. Logica: A powerful approach for estimating local genetic correlations across ancestries via GWAS summary statistics.

- 3. **Boran Gao**, Peijun Wu, Lulu Shang, and Xiang Zhou. MECO enables powerful multi-ancestry colocalization analysis.
- 4. Robert Langefeld, **Boran Gao**, and Xiang Zhou. pyGEMMA: A user-friendly Python implementation of linear mixed model for genome-wide association studies.

RESEARCH EXPERIENCE

Graduate Student Research Assistant

Jan 2018 - Present

Department of Biostatistics, University of Michigan

Advisor: Dr. Xiang Zhou

Genetic and Environmental Correlation Estimation in GWAS

Jan 2018 - Oct 2020

- Statistical Modeling
 - Developed a scalable method for genetic and environmental correlation estimation, GECKO
 - Utilized a bivariate linear mixed model to capture both genetic and environmental correlations
 - Derived and implemented a composite likelihood based optimization algorithm
- Application
 - Identified genetic and environmental correlated traits and enhanced understanding of trait co-evolution
 - Experience with five large scale GWASs and gene set enrichment analysis

Multi-Ancestry Fine-Mapping

Jan 2021 - Oct 2022

- Statistical Modeling
 - Developed a scalable and powerful method for multi-ancestry fine-mapping, MESuSiE
 - Used a multivariate sum of single effects model for causal variant detection and employed a mixture of multivariate normal to model both shared and ancestry-specific effects
 - Derived and implemented the multivariate iterative bayesian stepwise selection algorithm
- Application
 - Identified shared and ancestry-specific causal SNPs, facilitated understanding of disease etiology across diverse ancestries, realized the benefits of GWAS for underrepresented populations
 - Experience with multi-ancestry large-scale GWASs and function enrichment analysis

Multi-ancestry Colocalization

Jun 2022 - Present

- Statistical Modeling
 - Developed a scalable and powerful method for multi-ancestry colocalization, MECO
 - Employed multivariate normal mixture assumption to model colocalization signals, extended the MESuSiE algorithm towards multi-ancestry colocalization
- Application
 - Identified colocalization signals across ancestries, provided insights into shared biological pathways between molecular phenotype and complex traits across ancestries
 - Bridged GWAS with bulk RNA-seq studies

Local Genetic Correlation Estimation across Ancestries

Nov 2022 - Present

• Statistical Modeling

- Developed a scalable and powerful method for local genetic correlation estimation, Logica
- Utilized a bivariate linear mixed model for genetic correlation estimation and testing
- Derived and implemented the parameter expansion expectation—maximization algorithm for scalable estimation

• Application

- Estimated and detected local genetic correlations, uncovered shared genetic architecture across ancestries
- Experience with multi-ancestry large-scale GWASs

Graduate Student Research Assistant

Sep 2019 - Apr 2021

Department of Biostatistics, University of Michigan

Advisor: Dr. Xiang Zhou, Dr. Seunggeun Lee, Dr. Laura Scott

eQTL Mapping and TWAS in the Multi-Ethnic Study of Atherosclerosis

- Conducted multiple ancestry expression quantitative trait locus (eQTL) mapping study, and compared genetic architecture underlying expression variation
- Utilized PrediXcan to train gene expression predictors and performed transcriptome-wide association study (TWAS) across multiple ancestries
- Developed highly efficient and scalable eQTL detection and TWAS analysis pipeline

SELECTED PRESENTATIONS

- Platform talk, American Society of Human Genetics (ASHG), Los Angeles, CA
 Oct 2022
 MESuSiE: a novel method for discovering shared and unique putative causal variants by finemapping across diverse ancestries
- Center for Statistical Genetics, Department of Biostatistics, University of Michigan 2019-2021
 MESuSiE: a novel method for discovering shared and unique putative causal variants by fine mapping across diverse ancestries
 GECKO: Accurate genetic and environmental covariance estimation with composite likelihood in
 genome-wide association studies

PROFESSIONAL EXPERIENCE

Intern Statistical Genetics, 23andMe, CA

Jun 2022 - Aug 2022

- Developed an R package for multi-ancestry fine-mapping of 23andMe data
- Accounted for varying number of SNPs across ancestries, assumed causal SNPs are shared

Medical Consultant, Gongren's Hospital, Tangshan, China

Mar 2012 - Aug 2013

- Assisted with various administrative tasks, arranged rotations for resident physicians
- Managed the resolution of medical disputes

TEACHING AND MENTORING EXPERIENCE

Graduate Student Instructor (GSI)

• EPID 503: Strategies and Uses of Epidemiology

Jan 2016 - Apr 2016

Advisor: Dr. Aubree Gordon, Department of Epidemiology, University of Michigan.

• BIOSTAT 523: Statistical Methods for Epidemiology Sep 2016 - Dec 2016 Advisor: Dr. Susan Murray, Department of Biostatistics, University of Michigan

Research Mentor for:

• Peijun Wu, B	Biostatistics Ph.D. student, University of Michigan	Jun 2023 - Present
• Robert Lange	efeld, Biostatistics M.S. student, University of Michigan	Mar 2023 - Present
• Zheng Li, Bio	ostatistics Ph.D. student, University of Michigan	Nov 2022 - Present

SELECTED HONORS AND AWARDS

• Excellence in Research Award, Department of Biostatistics, University of Michigan	2023
• Charles J. Epstein Trainee Award, Predoctoral semifinalist, ASHG	2022
• Rackham Travel Award, University of Michigan	2022
• Outstanding GSI, Department of Biostatistics, University of Michigan	2017

LEADERSHIP AND SERVICES

Diversity, Equity and Inclusion Co	ommittee of Biostatistics	Sep 2019 -	July 2021
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SOFTWARE

- MESuSiE: Multiple ancEstry fine-mapping via Sum of the Single Effect model https://github.com/borangao/MESuSiE
- GECKO: Genetic and Environmental covariance estimation via Composite liKelihood Optimization https://github.com/borangao/GECKO

TECHNICAL SKILLS

- Programming: R, Python, C++, Shell, Git, LATEX, SAS
- Operating Systems: Mac OS X, Linux, Windows.

PROFESSIONAL SERVICE

Journal Reviewer: American Journal of Human Genetics, Journal of Orthopaedic Surgery and Research, Frontiers in Immunology, BMC Bioinformatics

ACADEMIC MEMBERSHIP

American Statistical Association, American Society of Human Genetics

REFERENCE LIST

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2. Seunggeun Lee

Professor, Graduate School of Data Science, Seoul National University, Seoul, Republic of Korea Adjunct Professor, Department of Biostatistics, University of Michigan, Ann Arbor, MI

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3. Zhongshang Yuan

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