# HAO FENG

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Mailing address 10900 Euclid Ave.

SOM / Robbins E-208

Cleveland, OH 44106

 $\begin{array}{ll} Phone & (216)\ 368\text{-}5510 \\ Email & \text{hxf155@case.edu} \end{array}$ 

Google Scholar https://scholar.google.com/citations?user=YGFvJjwAAAAJ&hl=en

Official Website http://epbiwww.case.edu/haofeng-phd/

Lab Website https://hfenglab.org/

## **Professional Experience**

• Assistant Professor (tenure-track)

2019.08-present

Department of Population & Quantitative Health Sciences

Case Western Reserve University School of Medicine

• Member 2019.08-present

Case Comprehensive Cancer Center Case Western Reserve University

### Education

• Ph.D. in Biostatistics and Bioinformatics

2014-2019

Emory University, Laney Graduate School

- Advisor: Dr. Hao Wu
- Dissertation committee members: Dr. Zhaohui Steve Qin, Dr. Peng Jin, Dr. Karen Conneely
- Dissertation: Statistical Methods for High-throughput Epigenomics Data.
- [\*\* Michael Kutner Distinguished Doctoral Student Award \*\*]
- MSPH in Biostatistics and Bioinformatics

2011-2013

Emory University, Rollins School of Public Health

- Advisors: Hao Wu, Karen Conneely
- Thesis: A Hierarchical Bayesian Approach to Detect Differentially Methylated Loci from Bisulfite Sequencing Data
- B.S. in Biosciences

2007-2011

University of Science and Technology of China (USTC)

## Past Work Experience

- May 2017—August 2017, Statistician/Full-Time Summer Intern Pfizer Inc, Global Research & Development, La Jolla, California
  - Developing Bayesian methods for tumor growth inhibition study
- June 2013—April 2014, Data Analyst, Department of Human Genetics, School of Medicine, Emory University, Atlanta, Georgia.
  - Statistical model development and data analysis for high-throughput data

#### Research Interests

- <u>Statistical Methodology</u>: Biostatistics, Bioinformatics, Computational methods in <u>high-throughput -omics</u> data, Signal deconvolution, Single-cell DNA methylation.
- Applications: Cancer epigenetics/epigenomics, computational biology, cell-free DNA, biomarker discovery, neurodegenerative diseases, PTSD.

### **Publications**

Google Scholar citation statistics (as of June 2024): All citations: 1627. h-index: 14. i10-index: 16.

 $^{\dagger}$  = These authors contributed equally to this work.

\* = Corresponding author.

underline = lab member and trainee.

## Statistical Methodology Publications

- 1. Guanqun Meng, Yue Pan, Wen Tang, Lijun Zhang, Ying Cui, Fredrick R. Schumacher, Ming Wang, Rui Wang, Sijia He, Jeffrey Krischer, Qian Li\*, **Hao Feng**\*. *imply: improving cell-type deconvolution accuracy using personalized reference profiles.* (2024) **Genome Medicine**.
- Zhenxing Guo<sup>†</sup>, Daoyu Duan<sup>†</sup>, Wen Tang, Julia Zhu, William S. Bush, Liangliang Zhang, Xiaofeng Zhu, Fulai Jin, Hao Feng\*. magpie: a power evaluation method for differential RNA methylation analysis in N6-methyladenosine sequencing. (2024) PLOS Computational Biology.
- 3. Hao Feng\*, Guanqun Meng, Tong Lin, Hemang Parikh, Ziyi Li, Jeffery Krischer, Qian Li\*. ISLET: individual-specific reference panel recovery improves cell-type-specific inference. (2023) Genome Biology.
- 4. <u>Daoyu Duan</u>, <u>Wen Tang</u>, Runshu Wang, Zhenxing Guo\*, **Hao Feng**\*. *Evaluation of epitranscriptome-wide N6-methyladenosine differential analysis methods.* (2023) **Briefings in Bioinformatics**.
- 5. Guanqun Meng, Wen Tang, Emina Huang, Ziyi Li, **Hao Feng**\*. A comprehensive assessment of cell type-specific differential expression methods in bulk data. (2023) **Briefings** in Bioinformatics.
- 6. <u>Daoyu Duan, Sijia He</u>, Emina Huang, Ziyi Li\*, **Hao Feng**\*. NeuCA web server: a neural network-based cell annotation tool with web-app and GUI. (2022) **Bioinformatics**.
- 7. Ziyi Li\*, **Hao Feng**\*. A neural network-based method for exhaustive cell label assignment using single cell RNA-seq data. (2022) **Scientific Reports**.
- 8. **Hao Feng**, Hao Wu. Differential methylation analysis for bisulfite sequencing using DSS. (2019) Quantitative Biology.
- 9. **Hao Feng**, Peng Jin, Hao Wu. Disease prediction by cell-free DNA methylation. (2018) **Briefings in Bioinformatics**.
- 10. Weiwei Zhang<sup>†</sup>, **Hao Feng**<sup>†</sup>, Hao Wu, Xiaoqi Zheng. Accounting for tumor purity improves cancer subtype classification from DNA methylation data. (2017) **Bioinformatics**.

11. **Hao Feng**, Karen N. Conneely, Hao Wu. A Bayesian hierarchical model to detect differentially methylated loci from single nucleotide resolution sequencing data. (2014) **Nucleic Acids Research**.

## **Book Chapter**

12. **Hao Feng**, Karen Conneely, Hao Wu. Differential Methylation Analysis for Bisulfite Sequencing (BS-Seq) Data. (2022) **Epigenome-Wide Association Studies**. Methods in Molecular Biology book series. Humana Press, Springer.

#### **Team Science Publications**

- 13. Daxuan Deng, Lijun Zhang, **Hao Feng**, Vern M. Chinchilli, Chixiang Chen, Ming Wang. Robust integration of secondary outcomes information into primary outcome analysis in the presence of missing data. (2024) Statistical Methods in Medical Research.
- 14. Wenzhang Wang, Fanpeng Zhao, Yubing Lu, Sandra L. Siedlak, Hisashi Fujioka, **Hao Feng**, George Perry, Xiongwei Zhu. Damaged Mitochondria Coincide with Presynaptic Vesicle Loss and Abnormalities in Alzheimer's Disease Brain. (2023) Acta Neuropathologica Communications.
- 15. Moeez Rathore, Wei Zhang, Michel'le Wright, Mehrdad Zarei, Ali Vaziri-Gohar, Omid Hajihassani, Ata Abbas, **Hao Feng**, Jonathan Brody, Sandy D. Markowit, Jordan Winter, Rui Wang. Liver Endothelium Microenvironment Promotes HER3-mediated Cell Growth in Pancreatic Ductal Adenocarcinoma. (2022) Journal of Cancer Science and Clinical Therapeutics.
- 16. Grace E Weber, Maria Khrestian, Elizabeth D Tuason, Yvonne Shao, Jagan Pillai, Stephen Rao, **Hao Feng**, Yadi Zhou, Feixiong Cheng, Tara M DeSilva, Shaun Stauffer, James B Leverenz, Lynn M Bekris. *Peripheral sTREM2-Related Inflammatory Activity Alterations in Early-Stage Alzheimer's Disease*. (2022) The Journal of Immunology.
- 17. Grace E Weber, Maria Khrestian, Elizabeth D Tuason, Yvonne Shao, Jagan A Pillai, Stephen M Rao, **Hao Feng**, Tara M DeSilva, Shaun Stauffer, James B Leverenz, Lynn M Bekris. Older adults with Alzheimer's disease exhibit an altered relationship between plasma sTREM2 and inflammatory markers. (2021) Alzheimer's & Dementia.
- 18. Yunhee Kang, Ying Zhou, Yujing Li, Yanfei Han, Jie Xu, Weibo Niu, Ziyi Li, Shiying Liu, **Hao Feng**, Wen Huang, Ranhui Duan, Tianmin Xu, Nisha Raj, Feiran Zhang, Juan Dou, Chongchong Xu, Hao Wu, Gary J Bassell, Stephen T Warren, Emily G Allen, Peng Jin, Zhexing Wen. A human forebrain organoid model of fragile X syndrome exhibits altered neurogenesis and highlights new treatment strategies. (2021) Nature Neuroscience.
- 19. Bin Jiao, Mengli Wang, **Hao Feng**, Han Bao, Feiran Zhang, Hao Wu, Junling Wang, Beisha Tang, Peng Jin, Lu Shen. *Downregulation of TOP2 modulates neurodegeneration caused by GGGGCC expanded repeats.* (2021) Human Molecular Genetics.
- 20. Samaneh K Sarvestani, Steven Signs, Bo Hu, Yunku Yeu, **Hao Feng**, Ying Ni, David R Hill, Robert C Fisher, Sylvain Ferrandon, Reece K DeHaan, Jennifer Stiene, Michael Cruise, Tae Hyun Hwang, Xiling Shen, Jason R Spence, Emina H Huang. *Induced organoids derived from patients with ulcerative colitis recapitulate colitic reactivity.* (2021) Nature Communications.

- 21. E. M. Kennedy, D. R. Powell, Z. Li, J. S. K. Bell, B. G. Barwick, Hao Feng, M. R. McCrary, B. Dwivedi, J. Kowalski, W. S. Dynan, K. N. Conneely, P. M. Vertino. Galactic Cosmic Radiation Induces Persistent Epigenome Alterations Relevant to Human Lung Cancer. (2018) Scientific Reports.
- 22. Yufang Qin, **Hao Feng**, Ming Chen, Hao Wu, Xiaoqi Zheng. InfiniumPurify: An R package for estimating and accounting for tumor purity in cancer methylation research. (2018) Genes & Diseases.
- 23. Lynn M Almli, Adriana Lori, Jacquelyn L Meyers, Jaemin Shin, Negar Fani, Adam X Maihofer, Caroline M Nievergelt, Alicia K Smith, Kristina B Mercer, Kimberly Kerley, Jennifer M Leveille, Hao Feng, Duna Abu-Amara, Janine D Flory, Rachel Yehuda, Charles R Marmar, Dewleen G Baker, Bekh Bradley, Karestan C Koenen, Karen N Conneely, Kerry J Ressler. Problematic alcohol use associates with sodium channel and clathrin linker 1 (SCLT1) in trauma-exposed populations. (2017) Addiction Biology.
- 24. Feiran Zhang, Christy Hammack, Sarah C Ogden, Yichen Cheng, Emily M Lee, Zhexing Wen, Xuyu Qian, Ha Nam Nguyen, Yujing Li, Bing Yao, Miao Xu, Tianlei Xu, Li Chen, Zhiqin Wang, Hao Feng, Wei-Kai Huang, Ki-jun Yoon, Chao Shan, Luoxiu Huang, Zhaohui Qin, Kimberly M Christian, Pei-Yong Shi, Mingjiang Xu, Menghang Xia, Wei Zheng, Hao Wu, Hongjun Song, Hengli Tang, Guo-Li Ming, Peng Jin. Molecular signatures associated with ZIKV exposure in human cortical neural progenitors. (2016) Nucleic Acids Research.
- 25. E Kennedy, DR Powell, Z Li, JSK Bell, Hao Feng, W Dynan, B Dwivedi, J Kowalski, KN Conneely, PM Vertino. Space radiation exposure induces stable epigenome alterations relevant to human lung cancer. (2016) Cancer Research.
- 26. Hao Wu, Tianlei Xu, **Hao Feng**, Li Chen, Ben Li, Bing Yao, Zhaohui Steve Qin, Peng Jin, Karen N. Conneely. Detection of differentially methylated regions from whole-genome bisulfite sequencing data without replicates. (2015) Nucleic Acids Research.
- 27. Lynn M Almli, Richard Duncan, Hao Feng, Debashis Ghosh, Elisabeth B Binder, Bekh Bradley, Kerry J Ressler, Karen N Conneely, Michael P Epstein. Correcting systematic inflation in genetic association tests that consider interaction effects: application to a genome-wide association study of posttraumatic stress disorder. (2014) JAMA Psychiatry.
- 28. Lynn M Almli, Amita Srivastava, Negar Fani, Kimberly Kerley, Kristina B Mercer, Hao Feng, Bekh Bradley, Kerry J Ressler. Follow-up and Extension of a Prior Genome-wide Association Study of Posttraumatic Stress Disorder: Gene × Environment Associations and Structural Magnetic Resonance Imaging in a Highly Traumatized African-American Civilian Population. (2014) Biological Psychiatry.
- 29. Lynn M Almli, Kristina B Mercer, Kimberly Kerley, Hao Feng, Bekh Bradley, Karen N Conneely, Kerry J Ressler. ADCYAP1R1 genotype associates with post-traumatic stress symptoms in highly traumatized African-American females. (2013) American Journal of Medical Genetics Part B: Neuropsychiatric Genetics.

## Funding Support

Current

• 1R01CA264320 (PI: Zhenghe Wang)

08/06/2021 - 07/31/2026

Mechanisms of PIK3CA helical domain mutations driving colorectal tumorigenesis

Role: Biostatistician

• 1R01 AG066526 (PI: Wen-Cheng Xiong)

09/15/2020 - 08/31/2024

NIH/NIA

LRP4 deficiency in Alzheimer's disease development

Role: Biostatistician

• <u>1R01HD105892</u> (PI: Adonis Hijaz)

08/20/2021 - 07/31/2026

NIH/NICHD

Patient-Centered Outcomes of Sacrocolpopexy versus Uterosacral Ligament Suspension for the Treatment of Uterovaginal Prolapse

Role: Biostatistician

09/01/2021 - 08/31/2024

DoD

BG34-200 Triggers Myeloid Recruitment and M1 Re-programming to Enhance Immunotherapy for Pediatric and AYA Osteosarcoma

Role: Biostatistician

 $\bullet \ \underline{\mathrm{P30AG072959}} \ (\mathrm{PI: Jonathan \ Haines})$ 

09/01/2021 - 06/30/2026

NIH/NIA

Cleveland Alzheimers Disease Research Center (CADRC) Data Management and Statistics Core

Role: Biostatistician

• <u>U01NS100610</u> (PI: Jordan Winter)

09/01/2023 - 08/31/2025

NIH/NCI

Targeting pancreatic cancer's metabolic addiction to HuR

Role: Biostatistician

 $\bullet$  <u>U01NS100610</u> (PI: James Leverenz)

09/01/2022 - 08/31/2027

NIH/NINDS

Dementia with Lewy Bodies Consortium

Role: Biostatistician

## Completed

• <u>IRG-16-186-21</u> (PI: Hao Feng)

08/01/2022 - 07/31/2023

American Cancer Society Institutional Research Grants (ACS IRG) pilot award, through Case Comprehensive Cancer Center

Jointly Profiling Microbiome and Transcriptome in Pancreatic Cancer Metastasis

Role: PI

• CASE60007 (PI: Hao Feng)

01/01/2022 - 12/31/2022

Corinne L. Dodero Foundation for the Arts and Sciences

Data Harmonizing and Cell Label Prediction Methods in Autism Single-cell RNA-seq Data

Role: PI

• R01CA237304 (PI: Emina Huang)

03/01/2019 - 02/29/2023

NIH/NCI

The miR-20/c-Myc/E2F Regulatory Axis is Critical for the Tumor Promoting Activity of Inflammatory Fibroblasts in Colitis-Associated Cancer

Role: Biostatistician

• <u>ID: 605146</u> (PI: Alex Huang) 11/01/2018 - 10/30/2021

St. Baldrick's Foundation

High Impact Research for Pediatric Osteosarcoma

Targeting myeloid and lymphoid immune tolerance in metastatic osteosarcoma

Role: Biostatistician

#### Honors and Awards

• UCITE Learning Fellowship Award, 2021

- Senior Ph.D. Student Presentation Award, 2018
- ENAR Distinguished Student Paper Award, 2017
- Michael Kutner Distinguished Doctoral Student Award, 2017
- SAMSI trave award for epigenetics workshop, 2015
- Finalist for the Charles C. Shepard Award, 2013
  - The best master's thesis of the Rollins School of Public Health
- Outstanding Student Scholarship from USTC Overseas Alumni Foundation, 2007

## Software Packages

- **ISLET**: Individual-Specific cell type referencing tool. An R/Bioconductor package deconvolute mixture samples and obtain the individual-specific and cell-type-specific reference panels. Available on Bioconductor: <a href="https://bioconductor.org/packages/ISLET/">https://bioconductor.org/packages/ISLET/</a>.
- magpie: This package aims to perform power analysis for the MeRIP-seq study. It calculates FDR, FDC, power, and precision under various study design parameters. Available on Bioconductor: https://bioconductor.org/packages/magpie/.
- **NeuCA**: Neural network-based cell type annotation tool. Available at https://bioconductor.org/packages/NeuCA/.
- **DSS**: Dispersion Shrinkage for Sequencing. An R/Bioconductor package for differential analysis from high-throughput sequencing data, including differential expression for RNA-seq and differential methylation for bisulfite-sequencing data. Available at <a href="https://bioconductor.org/packages/DSS/">https://bioconductor.org/packages/DSS/</a>.
- InfiniumPurify: A comprehensive R package for estimating and accounting for tumor purity based on DNA methylation Infinium 450k array data. 'InfiniumPurify' provides functionalities for tumor purity estimation. In addition, it can perform differential methylation detection and tumor sample clustering with the consideration of tumor purities. Available at <a href="https://cran.r-project.org/web/packages/InfiniumPurify/index.html">https://cran.r-project.org/web/packages/InfiniumPurify/index.html</a>.
- cfDNAMethy: Reference-free and reference-based methods for disease prediction by cell-free DNA methylation. Available at https://github.com/haoharryfeng/cfDNAmethy.

### Presentations

<u>Invited</u>

- MidSouth Computational Biology and Bioinformatics Society (MCBIOS) 2024, Atlanta GA. March, 2024.
  - Personalized cell-type-specific omics profile deconvolution and inference.
- Departmental Seminar Series, Wayne State University, Center for Urban Responses to Environmental Stressors (CURES), Detroit MI. November, 2023.
  - Personalized cell-type-specific omics profile deconvolution and inference.
- Joint Statistical Meetings (JSM), Toronto, Ontario, Canada. August, 2023.
  - Individual-specific and cell-type-specific gene expression reference panel deconvolution.
- International Chinese Statistical Association (ICSA) International Conference, Hong Kong SAR, China. July, 2023.
  - Individual-specific and cell-type-specific gene expression reference panel deconvolution.
- International Biometric Society (IBS) the Western North American Region (WNAR), Anchorage, AK. June, 2023.
  - Individual-specific and cell-type-specific gene expression reference panel deconvolution.
- Case Comprehensive Cancer Center seminar series. April, 2023.
  - Signal deconvolution methods development in -omics data.
- Departmental Seminar Series, University of Pittsburgh, PA. December, 2022.
  - Individual-specific and cell-type-specific gene expression reference panel deconvolution.
- Departmental Seminar Series, University of South Carolina Statistics Department. Columbia, SC. October, 2022.
  - Individual-specific and cell-type-specific gene expression reference panel deconvolution.
- Bioconductor Asia 2021, Tokyo, Japan. November, 2021.
  - NeuCA: a neural network-based method for exhaustive cell label assignment using single cell RNA-seq data.
- Joint Biostatistics Symposium 2021, Columbus, OH. May, 2021.
  - NeuCA: a neural network-based method for exhaustive cell label assignment using single cell RNA-seq data.
- University Hospitals Seidman Cancer Center, Cleveland, OH. November, 2020.
  - Omics data in cancer research.
- The University of Alabama at Birmingham, Birmingham, AL. March, 2019.
  - Statistical methods for high-throughput epigenomics data.
- Case Western Reserve University, Cleveland, OH. March, 2019.
  - Statistical methods for high-throughput epigenomics data.
- University of Missouri Columbia, Columbia, MO. January, 2019.
  - Statistical methods for high-throughput epigenomics data.

#### Topic-contributed, Contributed, and others

- STATGEN 2024, Pittsburgh, PA. May, 2024.
  - Personalized cell-type-specific omics profile deconvolution and inference.
- JSM 2022, Washington DC, USA. August, 2022
  - NeuCA: a neural network-based method for exhaustive cell label assignment using single cell RNA-seq data.

- ENAR 2022, Houston, TX, USA. March, 2022
  - NeuCA: a neural network-based method for exhaustive cell label assignment using single cell RNA-seq data.
- Affordable Cancer Technology 2021 (ACT 2021), Virtual, USA. October, 2021
  - NeuCA: a neural network-based method for exhaustive cell label assignment using single cell RNA-seq data.
- Senior Ph.D. students presentation competition, Emory University, Atlanta, GA. September, 2018
  - Statistical methods for high-throughput epigenomics data.
- Pfizer Global Research and Development, La Jolla, CA. August, 2017
  - A Bayesian method in tumor growth inhibition study.
- JSM 2017, Chicago, IL. August, 2017
  - Accounting for tumor purity improves cancer subtype classification from DNA methylation data.
- ENAR 2017 Spring meeting, Washington, DC. March, 2017
  - Accounting for tumor purity improves cancer subtype classification from DNA methylation data.
  - [ \*\* ENAR Distinguished Student Paper Award \*\*]
- Statistical and Applied Mathematical Sciences Institute (SAMSI) workshop, Research Triangle Park, NC. March, 2015
  - A Bayesian hierarchical model to detect differentially methylated loci from single nucleotide resolution sequencing data.
- American Society of Human Genetics (ASHG)  $63^{rd}$  annual meeting, Boston, MA. October, 2013
  - A Bayesian hierarchical model to detect differentially methylated loci from single nucleotide resolution sequencing data.

## Teaching

#### Instructor

- PQHS 471: Machine Learning and Data Mining. [3 credits] (2021 current). Department of Population & Quantitative Health Sciences, Case Western Reserve University School of Medicine.
- PQHS 701: Ph.D. Dissertation Research. [varied credits] (2023 current). Department
  of Population & Quantitative Health Sciences, Case Western Reserve University School of
  Medicine.
- PQHS 499: Independent Study on Modern Regression Methods. [6 credits] (2022 2023). Department of Population & Quantitative Health Sciences, Case Western Reserve University School of Medicine.
- STAT 395: **Senior Project in Statistics**. [3 credits] (2022). Department of Mathematics, Applied Mathematics, and Statistics, Case Western Reserve University College of Arts and Sciences.

#### Guest Lecturer

- PQHS 401: Research Seminar for MS program in Biostatistics. (Spring 2024 current). High-throughput Bioinformatics Data Analysis. Department of Population & Quantitative Health Sciences, Case Western Reserve University School of Medicine.
- PQHS 452: Statistical Methods in Human Genetics. (Spring 2021 current). *Multiple Testing and Statistical Power*. Department of Population & Quantitative Health Sciences, Case Western Reserve University School of Medicine.
- PQHS 457: Current Issues in Genetic Epidemiology: Design and Analysis of Sequencing Studies. (Spring 2020 – current). Epigenetics and Its Statistical Methods. Department of Population & Quantitative Health Sciences, Case Western Reserve University School of Medicine.

## Group Discussion Facilitator

• (2024 – current) IBMS 500: **On Being a Professional Scientist**, Case Western Reserve University School of Medicine.

#### Short Courses and Tutorials

• (2023 Spring ENAR Meeting) ENAR Tutorials: Cell-type-aware Differential Analysis for Bulk Transcriptome Data. International Biometric Society (IBS) – ENAR.

#### Teaching Associate

- BIOS 731: Advanced Statistical Computing. (2018). Department of Biostatistics and Bioinformatics, Emory University Rollins School of Public Health.
- BIOS 738: Bayesian and Empirical Bayesian Methods. (2017). Department of Biostatistics and Bioinformatics, Emory University Rollins School of Public Health.
- BIOS 555: **High-throughput data analysis using R and Bioconductor**. (2016). Department of Biostatistics and Bioinformatics, Emory University Rollins School of Public Health.

#### Teaching Assistant

- BIOS 550: **Sampling Applications**. (2016). Department of Biostatistics and Bioinformatics, Emory University Rollins School of Public Health.
- BIOS 500: **Statistical Methods I**. (2015). Department of Biostatistics and Bioinformatics, Emory University Rollins School of Public Health.

## Mentoring

#### Doctoral trainees

- Daoyu Duan, Ph.D. candidate in Epidemiology and Biostatistics
- Guanqun (Leslie) Meng, Ph.D. candidate in Epidemiology and Biostatistics

#### Dissertation committee

- Yanning Wu (2024 ), Ph.D. candidate in Epidemiology and Biostatistics
- Aditi Singh (2023 ), Ph.D. Ph.D. student in Wayne State University

- Noah Lorincz-Comi (2021 2024), Ph.D. candidate in Epidemiology and Biostatistics
- Xueyi Zhang (2021 2024), Ph.D. candidate in Epidemiology and Biostatistics
- Michael Osterman (2020 2023), Ph.D. student in Epidemiology and Biostatistics
- Bowen Jin (2021 2023), Ph.D. student in Systems Biology and Bioinformatics
- Zuxi (Terry) Cui (2020 ), Ph.D. candidate in Epidemiology and Biostatistics
- Ju Zhang (2019 2021), Ph.D. candidate in Epidemiology and Biostatistics

#### Rotation graduate students

- Chenyu Liu (2023 2024), Ph.D. student in Epidemiology and Biostatistics
- Katrina Prantzalos (2020 2021), Ph.D. student in Biomedical and Health Informatics
- Dipak Prasad Upadhyaya (2020 2021), Ph.D. student in Biomedical and Health Informatics
- Jing Zhang (2019 2020), Ph.D. candidate in Epidemiology and Biostatistics
- Shiying Liu (2019 2020), Ph.D. candidate in Epidemiology and Biostatistics

#### Masters students

• Capstone Project Committee for Nicolas Kaplan (2019 - 2021), MPH candidate

#### Visiting students

• Aditi Singh (2022), Graduate Student, Wayne State University, Center for Molecular Medicine and Genetics.

### Undergraduate students and high school students

- Sijia He (2021-2022), CWRU, undergraduate capstone.
- Julia Zhu (2021), Hathaway Brown School, summer research project.

### Service Activities

#### Committee service

- 2023 current, Member, PQHS departmental Committee on Appointments Promotions and Tenure (dCAPT)
- 2023 current, Member, CWRU School of Medicine (SOM) Curriculum Committee
- 2022 2024, Chair, PQHS departmental seminar series committee
- 2021- current, Member, Biomedical & Health Informatics (BHI) admission and curriculum committee
- 2019 2021, Member, Biostatistics Faculty Search Committee
- 2020, Member, Epidemiology and Biostatistics Ph.D. Admission Committee
- 2020, Member, PQHS Student Activity Committee
- 2019, Member, Emory Biostatistics Department Website Committee

#### Conference program

- Organizer and Chair: STATGEN 2024, Topic-contributed Session "Dissecting and testing heterogeneous transcriptome and chromatin signals." STATGEN 2024: Conference on Statistics in Genomics and Genetics, Pittsburgh, PA. May 2024.
- Program Committee: IEEE International Conference on Bioinformatics and Biomedicine (IEEE BIBM). 2022 & 2023.
- Organizer: JSM, Topic-contributed Session "Methodological Advances in Cell Type Deconvolution and Discrimination." JSM 2023, Toronto, Ontario, Canada. August 2023.
- Chair: "WNAR 2023 session Speed Talk 1". WNAR 2023, Anchorage, AK. June 2023.
- Organizer and Chair: ICSA Applied Statistics Symposium, Invited Session "Statistical Methods Development in Single-cell Genomics and Multi-omics Data." ICSA Applied Statistics Symposium, Ann Arbor, MI. June 2023.
- Award Committee: ENAR conference poster award committee. 2023
- Chair: "ENAR session Contributed Papers: Genomics". ENAR 2022, Houston, TX. March 2022.
- Organizer and Chair: Invited Session "Statistical method advancement for analyzing omics data". ICSA 2020 Applied Statistics Symposium, Houston, TX. December 2020.

#### Grant review

- National Science Foundation (NSF), Graduate Research Fellowship Program (GRFP). Panel Reviewer. 2024.
- Department of Defense (DoD) Congressionally Directed Medical Research Programs (CDMRP), Biostatistician Specialist Reviewer. 2022-2024.
- National Institutes of Health (NIH) Genomics, Computational Biology and Technology (GCAT) Study Section, Early Career Reviewer. 2023.
- Invited Reviewer for Sector Innovation Program at Genome British Columbia (GENOME BC), Vancouver, BC, Canada. May 2020.

## **Editorial Activities**

#### Editorial board member

• International Journal of Computational Biology and Drug Design (2020-2023)

#### Journal Reviewer

Aging, American Journal of Tropical Medicine and Hygiene, Annals of Applied Statistics,
Bioinformatics, Bioinformatics Advances, Biomedicine and Pharmacotherapy, Biometrics,
Briefings in Bioinformatics, BMC Bioinformatics, BMC Supplements, Cell Reports Methods,
Cell Systems, Cell Genomics, Cell & Bioscience, Communication Biology, Genes & Diseases,
Genetic Epidemiology, Genome Biology, Journal of Applied Statistics, Journal of Alzheimer's
Disease, Meta Gene, Nucleic Acids Research, NAR Genomics and Bioinformatics, PeerJ,
PLOS Genetics, PLOS ONE, RECOMB, Statistical Analysis and Data Mining, Scientific
Reports, Statistical Methods in Medical Research, Statistics in Biosciences.

# Professional Membership

American Statistical Association (ASA)
The International Biometric Society — Eastern North American Region (ENAR)
International Chinese Statistical Association (ICSA)
International Society for Computational Biology (ISCB)
American Society of Human Genetics (ASHG)