ZHENXING GUO

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Google scholar: https://scholar.google.com.hk/citations?user=hmOmP4EAAAAJ&hl=zh-CN

Education

• Ph.D. candidate in Biostatistics and Bioinformatics

2017 - Present

Emory University, Laney Graduate School

— Advisor: Hao Wu

• M.S. in Operational Research

2014 - 2017

Academy of Mathematics and System Science (AMSS) Chinese Academy of Sciences (CAS), Beijing, China.

- Advisor: Shihua Zhang
- Thesis: Models and Algorithms for Sparse deep nonnegative matrix factorization

• B.S. in Applied Mathematics

2010 - 2014

College of Mathematics & Econometrics Hunan University, Changsha, China.

Published Methodological work

- Z. Guo, A. M. Shafik, P. Jin, H. Wu (202x) Differential RNA Methylation Analysis for MeRIP-seq Data under General Experimental Design. Bioinformatics under revision.
- Z. Guo, A. M. Shafik, P. Jin, Z. Wu, H. Wu (2021) Detecting m6A methylation regions from Methylated RNA Immunoprecipitation Sequencing. Bioinformatics, 37(18): 2818–2824. https://doi.org/10.1093/bioinformatics/btab181
- Z Li, **Z. Guo**, Y Cheng, P Jin, H Wu. Robust partial reference-free cell composition estimation from tissue expression. **Bioinformatics**, (2020), 36(11): 3431-3438. https://academic.oup.com/bioinformatics/article/36/11/3431/5804977
- Z. Guo*, Y Cui*, X Shi, JA Birchler, I Albizua, SL Sherman, ZS Qin, T Ji (2020) An empirical bayesian approach for testing gene expression fold change and its application in detecting global dosage effects. NAR Genomics and Bioinformatics, 2(3):lqaa072. https://academic.oup.com/nargab/article/2/3/lqaa072/5908375
- Z. Guo, S Zhang (2020) Sparse deep nonnegative matrix factorization. Big Data Mining and Analytics, 3(1):13-28. https://ieeexplore.ieee.org/document/8935092

Collaborations

- M. C. Runnstrom1, A. Morrison-Porter, M. Ravindran, H. Quehl, R. Ramonell, M. Woodruff, M. Kuruvilla, F. Anam, R. Swerlick, C. Swenson, C. Polito, W. Neveu, R. Patel, C. Kim, I. Hentenaar, Z. Guo, H. Wu, N. S. Haddad, J. L. Daiss, I. Sanz, F. E. Lee* (202x) Reduced SARS-CoV-2 Vaccine Response in Patients Treated with Biologic Therapies for Asthma. In preparation.
- Y. Pan*, Y. Yang, K. Kathleen, G. Khalil, R. Kelly, B. Switzer, **Z. Guo** (202x) Simulation Study for Improving Efficacy Estimates for New HIV Treatments. In preparation.
- A. M. Shafik, F. Zhang, **Z. Guo**, Q. Dai, K. Pajdzik, Y. Li, Y. Kang, B. Yao, H. Wu, C. He, E. G. Allen, R. Duan, P. Jin* (2021) N6-methyladenosine dynamics in neurodevelopment and aging, and its potential role in Alzheimer's disease. **Genome Biology**, 22(1):1-19. https://link.springer.com/article/10.1186/s13059-020-02249-z

Software Developed

- TRESS: Toolbox for mRNA Epigenetics Sequencing analysis. An R/Bioconductor package for analysis of RNA methylation sequencing data. Available at https://bioconductor.org/packages/devel/bioc/html/TRESS.html.
 - Software based on publication Guo 2021 Bioinformatics.
- Foldseq: Fold change detection from RNA-seq data. An R package for testing gene expression fold change. Available on github at https://github.com/cuiyingbeicheng/Foldseq.
 - Software based on publication Guo 2020 NAR Genomics and Bioinformatics

Presentations

- Guo, Z. Differential m⁶A Analysis for MeRIP-seq Data under General Experimental Design. Poster Presentation at Georgia Statistics Day, Atlanta, GA Oct 2021
- Guo Z., Shafik A. M., Jin P., Wu Z., Wu H. (2021) Detecting m6A methylation regions from Methylated RNA Immunoprecipitation Sequencing. JSM 2021, Virtual Aug 2021

Invited talks / lectures at other academic institutes

- Statistical method and tools for RNA methylation sequencing analysis. Department of Population and Quantitative Health Sciences, School of Medicine, Case Western Reserve University, Virtual

 May 2022
- Including Historical Data in Clinical Trial Analysis. Summer school at Shandong University, July 2021

Mentoring Experience

- PhD student mentor for 1^{st} year theory and 2^{nd} year method qualifying exam, 2020.
- \bullet PhD student mentor for 2^{nd} year theory qualifying exam, 2021.

Teaching Experience

- Teaching assistant. Fall 2020: BIOS 731 Advanced statistical computing.
- Teaching assistant. Spring 2020: BIOS 545 R programming.
- Teaching assistant. Fall 2019: BIOS 555 High-throughput data analysis using R/Bioconductor.
- Teaching assistant. Spring 2019: BIOS 591P Lab "Biostatistics Methods II".
- Teaching assistant. Fall 2018: BIOS 500/003: Statistical Methods I Group C.

Work Experience

• June 2020 - August 2020, Technical Intern at Bristol Myers Squibb

Mentors: Rong Liu

Role: Augmentation of current clinical trial with external real world data using Bayesian techniques.

 Oct 2018 – May 2021, Research assistant, Emory University, Department of Biostatistics and Bioinformatics, Atlanta, GA

Advisor: Prof. Hao Wu

Role: Research projects on high-throughput genomic sequencing data

Honors and Award

PhD career:

- Honorable Mention, Emory Biostatistics senior PhD student presentation competition, 2022.
- Excellent Poster Award of Georgia Statistics Day, 2021.
- Michael H. Kutner Doctoral Student Award, Emory University, 2021-2022.
- JSM Distinguished Student Paper Award, 2021.
- SAS Certified Base Programmer for SAS 9, 2017.

Undergraduate career:

- Outstanding Graduate of Hunan University, 2014.
- First Prize of Dissertation, Hunan University, 2014.
- Merit Student of Hunan University (top 6%), Hunan University, 2011-2012, 2012-2013.
- National Scholarship(<1%), Hunan University, 2011-2012.

Professional Membership

- American Statistical Association (ASA)
- The International Biometric Society Eastern North American Region (ENAR)

• International Chinese Statistical Association (ICSA)

Computational Skills

• Programming Languages: R, SAS, Matlab, Python

• Operating Systems: Windows, Unix