WENJING MA, PhD

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Beijing, China

Postdoctoral Research Fellow, Department of Biostatistics, University of Michigan 1415 Washington Heights, Ann Arbor, MI 48109-2029

mawenjing1993@gmail.com | 917-496-1226 | https://marvinquiet.github.io/

Research Interests

- Develop deep learning methods integrated with statistical modeling to analyze single-cell genomics data, including spatial transcriptomics, epigenomics, and multi-omics
- Investigate temporal dynamics of cellular processes using time-resolved single-cell data to reveal biological mechanisms such as spatial domain transitions, cell state changes, and gene regulatory network dynamics
- Apply developed methodologies to study dynamics in epigenetic disease progression, with a focus on neurodegenerative disorders and cancer

Education and Training

University of MichiganAnn Arbor, MIPostdoctoral Research Fellow, Department of Biostatistics07/2023 - current

Advisor: Dr. Xiang Zhou

Emory University Atlanta, GA

Ph.D., in Computer Science and Informatics (Biomedical Informatics Track) 08/2019 - 05/2023

Dissertation Title: Cell type identification in single-cell genomics and its application

Advisor: Dr. Hao Wu

University of Virginia Charlottesville, VA

Visiting Scholar in Bioinformatics, Center for Public Health Genomics 08/2018 - 06/2019

Advisor: Dr. Chongzhi Zang

Beijing University of Posts and Telecommunications (BUPT)

M.S. and B.S. in Computer Science and Technology 09/2010 - 06/2017

Publications (Reverse chronological order)

Note: * denotes equal-contribution;

Preprints

- 1. **Wenjing Ma**, Siyu Hou, Lulu Shang, Jiaying Lu, Xiang Zhou. Optimal transport modeling uncovers spatial domain dynamics in spatiotemporal transcriptomics. *Under Review* at *Nature Communications* (2025).
- 2. Siyu Hou, **Wenjing Ma**, Xiang Zhou. FastCCC: A permutation-free framework for scalable, robust, and reference-based cell-cell communication analysis in single cell transcriptomics studies. *Under Revision* at *Nature Communications*, (2025). DOI
- 3. Jie Xu*, **Wenjing Ma***, Shaojun Yu*, Yujing Li, Jidong Goo, Christina Michalski, Mengjun Zhang, Jianjun Wang, Yunhee Kang, Xiangxue Guo, Weibo Niu, Ying Zhou, Zhexing Wen, Peng Jin. A dynamic

- temporal regulatory role of human FMRP underlying altered developmental trajectories in fragile X syndrome-derived hippocampal organoids. *Under Review at Nature Communications*, (2025).
- 4. Yulin Jin*, Yiqu Cao*, **Wenjing Ma***, Ronghua Li, Yujing Li, Yunhee Kang, Xiangxue Guo, Jing Huang, Michael Epstein, Natalia Rivera, Junghwa Lim, Emily Allen, Peng Jin. Integrative transcriptome-wide association reveals PRKCG-linked GABAergic dysfunction in Fragile X-associated Tremor/Ataxia Syndrome. *Under Revision at Nature Communications*, (2025).
- 5. Sumeet Sharma*, **Wenjing Ma***, Kerry J. Ressler, Thea Anderson, Dan. C. Li, Peng Jin, Shannon L. Gourley, Zhaohui Qin. Dysregulation of prefrontal oligodendrocyte lineage cells across mouse models of adversity and human major depressive disorder. *bioRxiv*, (2022). DOI

Peer-Reviewed Journals

- 6. Xin Wei, **Wenjing Ma**, Zhijin Wu, Hao Wu. TORC: Target-oriented Reference Construction for supervised cell-type identification in scRNA-seq. *In press at Genome Biology*, (2025). (*A conference version has been accepted by* IEEE BIBM 2024)
- 7. Hejie Cui, Jiaying Lu, Ran Xu, Shiyu Wang, **Wenjing Ma**, Sue Ye, Shaojun Yu, Xuan Kan, Chen Ling, Liang Zhao, Zhaohui S. Qin, Joyce Ho, Tianfan Fu, Jing Ma, Mengdi Huai, Fei Wang, Carl Yang. A survey on knowledge graphs for healthcare: Resources, applications, and promises. *In press at Journal of Biomedical Informatics*, (2025). (*A conference version has been accepted by ICML 2023 Workshop on Interpretable Machine Learning in Healthcare*)
- 8. Yulin Jin, Karen N Conneely, **Wenjing Ma**, Robert K. Naviaux, Teepu Siddique, Emily G. Allen, Sandra Gingrich, Robert M. Pascuzzi, Peng Jin. Whole-genome bisulfite sequencing of cell-free DNA unveils age-dependent and ALS-associated methylation alterations. *Cell & Bioscience*, (2025). <u>DOI</u>
- 9. Shilin Yu, Guanquan Meng, Wen Tang, **Wenjing Ma**, Rui Wang, Xiongwei Zhu, Xiaobo Sun, Hao Feng. cypress: an R/Bioconductor package for cell-type-specific differential expression analysis power assessment. *Bioinformatics*, (2024). DOI [R package]
- 10. **Wenjing Ma**, Jiaying Lu, Hao Wu. Cellcano: supervised cell type identification for single cell ATAC-seq data. *Nature Communications*, (2023). <u>DOI [Python package] [Documentation]</u>
- 11. **Wenjing Ma**, Sumeet Sharma, Peng Jin, Shannon L. Gourley, Zhaohui S. Qin. *LRcell*: detecting the source of differential expression at the sub–cell-type level from bulk RNA-seq data. *Briefings in Bioinformatics*, (2022). DOI [R package]
- 12. Yulin Jin, Kenong Su, Ha Eun Kong, **Wenjing Ma**, Zhiqin Wang, Yujing Li, Ronghua Li, Emily Allen, Hao Wu, Peng Jin. Cell-type specific DNA methylome signatures reveal epigenetic mechanisms for neuronal diversity and neurodevelopment disorder. *Human Molecular Genetics*, (2022). DOI
- 13. Shengen S. Hu, Lin Liu, Qi Li, **Wenjing Ma**, Michael J. Guertin, Clifford A. Meyer, Ke Deng, Tingting Zhang, Chongzhi Zhang. Intrinsic bias estimation for improved analysis of bulk and single-cell chromatin accessibility profiles using SELMA. *Nature Communications*, (2022). DOI, [package]
- 14. Wenjing Ma, Kenong Su, Hao Wu. Evaluation of some aspects in supervised cell type identification for single-cell RNA-seq: classifier, feature selection, and reference construction. *Genome Biology*, (2021).
 DOI [project link]

15. **Wenjing Ma***, Zhenjia Wang*, Yifan Zhang, Neal E. Magee, Yayi Feng, Ruoyao Shi, Yang Chen, Chongzhi Zang. BARTweb: a web server for transcriptional regulator association analysis. *NAR Genomics and Bioinformatics*, (2021). DOI [website] [RECOMB 2019 poster]

Book Chapters

16. Ronnie Y. Li, **Wenjing Ma**, Zhaohui S. Qin. Approaches to Marker Gene Identification from Single-Cell RNA-Sequencing Data. In *Handbook of Statistical Bioinformatics* (pp. 71-84). Springer, Berlin, Heidelberg. (2022). DOI

Teaching Experience

• BIOS 555: High-throughput data analysis using R and Bioconductor

Emory University

<u>Teaching assistant and guest lecturer</u> for Dr. Hao Wu (Course for *PhD students*)

Fall 2022

Lecture about advanced single-cell genomics sequencing technologies along with analysis techniques

• QTM 100: Intro to Statistical Inference

Emory University

<u>Instructor</u> (Course for *undergraduates*)

Fall 2021

Teach basic R coding and apply to perform statistical tests on real-world applications

• CS 584: Biomedical Image Analysis

Emory University

<u>Teaching assistant</u> for Dr. Imon Banerjee (Course for *PhD students*)

Spring 2021

• CS 253: Data Structures and Algorithms

Emory University

Teaching assistant for Dr. Jinho Choi (Course for undergraduates)

Fall 2020

• CS 224: Foundations of Computer Science

Emory University

Teaching assistant for Dr. Timothy Sun (Course for *undergraduates*)

Spring 2020

Mentoring Experience

• Xin Wei Emory University

Master in Biostatistics (Now PhD student at Brown University)

Summer 2022

Co-mentored with Dr. Hao Wu.

Project: Identify reference datasets combination strategy for accurate cell type identification for single cell genomics data

Yielded a paper accepted by BIBM 2024 and under revision at Genome Biology.

· Rohan Dalal and Francis Wu

High school students (Now undergraduates at Georgia Institute of Technology)

Summer 2022

Project: Evaluate how different deep neural network structures affect cell type prediction performance Yielded a poster presentation.

Fellowships, Awards and Honors

Fellowships

• Women in Natural Sciences Fellowship

Emory University 2019 - 2023

Awards

• Student Paper Award [News] [Interview, Page 9] American Statistical Association 2023 ASA Section: Section on Statistics in Genomics and Genetics (SGG)

• 1st place Young Scientist Excellence Award (YSEA) Category: Graduate Student **MCBIOS 2023**

1st place Poster Award

MCBIOS 2023

• 3rd place Student Poster Presentation award (out of 47) [News]

Georgia Statistics Day 2022

• Honorable Mention Poster

Georgia Statistics Day 2021

• Best 5-minute presentation award

2nd Annual Workshop Emerging Data Science Methods for

Complex Biomedical and Cyber Data 2021

Travel Grants

 2023 Advances in Statistical and Computational Methods for Analysis of Biomedical, Genetic, and Omics Data (ABGOD)
 Dallas, TX

• 2022 Grad Cohort for Women

New Orleans, LA

Talks and Posters

<u>Talks</u>

 2025 Tools and Technology seminar series, University of Michigan [video] Remote SpaDOT: Optimal transport analysis of spatial genomics data reveals functional domain changes along time trajectories

• 2024 32nd International Biometric Conference

Atlanta, GA

SpaDOT: Optimal transport analysis of spatial genomics data reveals functional domain changes along time trajectories

Invited session: New advances in statistical methods and applications for spatial transcriptomics and cellular heterogeneity [link]

• 2023 Eastern North American Region

Nashville, TN

Cellcano: Supervised cell type identification for single cell ATAC-seq data

Contributed papers session: Machine learning - Methods and Applications

• 2023 MCBIOS: Big data and artificial intelligence for genomics and therapeutics Cellcano: Supervised cell type identification for single cell ATAC-seq data

Dallas, TX

• 2023 The Chinese University of Hong Kong, Shenzhen, School of Data Science forum

Remote

Supervised cell type identification for single-cell genomics and its applications

• 2022 Atlanta Workshop on Single-cell Omits (AWSOM)

Atlanta, GA

LRcell: detecting the source of differential expression at the sub-cell-type level from bulk RNA-seq data

• 2022 Computational Genomics Summer Institute (CGSI)

Los Angeles, CA

Supervised cell type identification for single-cell omics and its application

• 2021 Emerging Data Science Methods for Complex Biomedical and Cyber Data

Remote

Evaluation of some aspects in supervised cell type identification for single-cell RNA-seq

Posters

• 2022 Georgia Statistics Day

Arthens, GA

Cellcano: supervised cell type identification for single cell ATAC-seq data

Evaluation of some aspects in supervised cell type identification for single-cell RNA-seq

Professional Activities and Services

- Invited reviewer for Journal of Applied Statistics.
- Invited reviewer for American Medical Informatics Association (AMIA) 2025.
- Invited reviewer for IEEE Transactions on Computational Biology and Bioinformatics.
- Invited reviewer for BMC Bioinformatics.
- Invited reviewer for Bioinformatics Advances.
- Invited reviewer for Bioinformatics.
- Invited Program Committee (PC) member for IEEE International Conference on Bioinformatics and Biomedicine (BIBM) 2024, IEEE BIBM 2023; Invited reviewer for IEEE BIBM 2022.
- Invited reviewer for International Journal of Computational Biology and Drug Design (IJCBDD).
- Invited sub-reviewer for ISMB/ECCB 2023.
- Invited reviewer for Journal of Agricultural, Biological, and Environmental Statistics (JABE).

List of References

Xiang Zhou

Professor

Department of Biostatistics

University of Michigan

Address: 1415 Washington Heights, Ann Arbor, MI 48109

Email: xzhousph@umich.edu

Tel: (734) 764-5722

Hao Wu

Distinguished Professor

Computer Science and Control Engineering

Shenzhen University of Advanced Technology

Address: No.1 Gongchang Road, Guangming District, Shenzhen, P.R.China

Email: wuhao@suat-sz.edu.cn

Peng Jin

Robert W. Woodruff Professor of Human Genetics

Chair, Department of Human Genetics, School of Medicine

Director, Emory Stephen T. Warren National Fragile X Research Center

Emory University

Address: 615 Michael Street, Suite 305E, Atlanta, GA 30322

Email: peng.jin@emory.edu

Tel: (404) 727-3729