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Chong Wu

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☆: https://wuchong.org

RESEARCH INTEREST

Statistical genomics (TWAS, GWAS, multi-omics, integrative analysis, polygenic risk score) Machine learning (unsupervised learning, supervised learning)
Causal inference (Mendelian randomization, instrumental variables regression)

EMPLOYMENT

Assistant Professor

Aug. 2018-Present

Department of Statistics, Florida State University

EDUCATION

University of Minnesota

Minneapolis, MN

Jun. 2018

Ph.D., Biostatistics

Advisors: Drs. Weihua Guan & Wei Pan

Thesis: Statistical methods for high-dimensional genetic and genomic data

Huazhong University of Science & Technology

Wuhan, China Jun. 2013

B.S., Applied Mathematics Advisor: Dr. Xiaoyang Zhou

PEER-REVIEWED ARTICLES

-Methodology

- 1. He, Y., Xu, G., **Wu, C.**, & Pan, W. (2021). Asymptotically independent U-statistics in high-dimensional testing. *Annals of Statistics*, 49(1), 154–181.
- 2. **Wu, C.***, Bradley, J., Li, Y., Wu, L., & Deng, H. (2021). A gene-level methylome-wide association analysis identifies novel Alzheimer's disease genes. *Bioinformatics*, 37(14), 1933–1940.
- 3. Bae Y.[‡], Wu, L., & **Wu, C.*** (2021). InTACT: An adaptive and powerful framework for joint-tissue transcriptome-wide association studies. *Genetic Epidemiology*,1–12.
- Wu, C.*, Xu, G., Shen, X., & Pan, W.* (2020). A regularization-based adaptive test for high-dimensional generalized linear models. *Journal of Machine Learning Research*, 21, 1–67. (This paper won the 2019 ENAR Distinguished Student Paper Award.)
- 5. Xue, H., **Wu, C.**, & Pan, W. (2020). Leveraging existing GWAS summary data to improve power for a new GWAS. *Genetic Epidemiology*, 44(7), 717–732.
- Wu, C. (2020). Multi-trait genome-wide analyses of the brain imaging phenotypes in UK Biobank. Genetics, 215(4), 947–958.
 (This paper won a <u>platform presentation</u> at the American Society of Human Genetics (ASHG) 2019 Annual Meeting; highlighted in *Genetics* August issue)
- Yang, T., Wu, C., Wei, P., & Pan, W. (2020). Integrating DNA sequencing and transcriptomic data for association analyses of low-frequency variants and lipid traits. *Human Molecular Genetics*, 29(3), 515–526.

^{*} Corresponding author; †Co-first author; ‡Students

- 8. Yang, T., Kim, J., **Wu, C.**, Ma, Y., Wei, P., & Pan, W. (2020). An adaptive test for meta-analysis of rare variant association studies. *Genetic Epidemiology*, 44(1), 104–116.
- 9. **Wu, C.***, & Pan, W.* (2020). A powerful fine-mapping method for transcriptome-wide association studies. *Human Genetics*, 139, 199–213.
- Wu, C.*, & Pan, W.* (2019) Integration of methylation QTL and enhancer-target gene maps with schizophrenia GWAS summary results identifies novel genes. *Bioinformatics*, 35(19), 3576–3583.
- 11. **Wu, C.***, Xu, G., & Pan, W.* (2019). An adaptive test on high dimensional parameters in generalized linear models. *Statistica Sinica*, 29, 2163–2186.
- 12. **Wu, C.**, & Pan, W. (2018). Integration of enhancer-promoter interactions with GWAS summary results identifies novel schizophrenia-associated genes and pathways. *Genetics*, 209(3), 699–709.
- 13. Wu, C., & Pan, W. (2018). Integrating eQTL data with GWAS summary statistics in pathway-based analysis. Genetic Epidemiology, 42(3), 303–316. (Highlighted by International Genetic Epidemiology Society (IGES). This paper won a poster talk at the ASHG 2017 Annual Meeting.)
- 14. Park, J.Y., **Wu, C.**, Basu, S., McGue, M., & Pan, W. (2018). Adaptive SNP set association testing in generalized linear mixed models with application to family studies. *Behavior Genetics*, 48(1), 55–66.
- 15. Xu, Z., **Wu, C.**, Wei, P., & Pan, W. (2017). A powerful framework for integrating eQTL and GWAS summary data. *Genetics*, 207(3), 893–902.
- 16. Liu, B., **Wu**, **C.**, Shen, X., & Pan, W. (2017). A novel and efficient algorithm for de novo discovery of mutated driver pathways. *Annals of Applied Statistics*, 11(3), 1481–1512.
- 17. Xu, Z., **Wu, C.**, Pan, W., & Alzheimer's Disease Neuroimaging Initiative (ADNI). (2017). Imaging-wide association study: integrating imaging endophenotypes in GWAS. *NeuroImage*, 159, 159–169.
 - (This paper won a platform presentation at the ASHG 2017 Annual Meeting.)
- 18. **Wu, C.**[†], Kwon, S.[†], Shen, X., & Pan, W. (2016). A new algorithm and theory for penalized regression-based clustering. *Journal of Machine Learning Research*, 17(188), 1–25.
- Wu, C., Chen, J., Kim, J., & Pan, W. (2016). An adaptive association test for microbiome data. Genome Medicine, 8(1), 1–12.
 (This paper won the 2016 Joint Statistical Meetings (JSM) Distinguished Student Paper Award on Statistics in Genomics and Genetics Section.)
- 20. **Wu, C.**, Demerath, E. W., Pankow, J. S., Bressler, J., Fornage, M., Grove, M. L., Chen, W., & Guan, W. (2016). Imputation of missing covariate values in epigenome-wide analysis of DNA methylation data. *Epigenetics*, 11(2), 132–139.

—Applications & Collaborations

- 21. Xie, T., Xu, C., Shi, X., **Wu, C.**, Meng, R., Meng, X., Yu, G., Wang, K., Xiao, H., & Deng, H. (2021). Accurate recognition of colorectal cancer with semi-supervised deep learning on pathological images. *Nature Communications*. Accepted.
- 22. **Wu, C.***, Zhu, J., King, A.[‡], Tong, X., Lu, Q., Park, J. Y., ... & Wu, L.* (2021). Novel strategy for disease risk prediction incorporating predicted gene expression and DNA methylation data: a multi-phased study of prostate cancer. *Cancer Communications*, 1–11.

- 23. Wu, L.*, Zhu, J., Liu, D., Sun, Y., & **Wu, C.*** (2021). An integrative multiomics analysis identifies putative causal genes for COVID-19 severity. *Genetics in Medicine*, 1–11.
- 24. **Wu, C.***, Wu, L., Wang, J., Lin, L., Li, Y., Lu, Q., & Deng, H. (2021). Systematic identification of modifiable risk factors and drug repurposing options for Alzheimer's disease: Mendelian randomization analyses. *Alzheimer's Dement*, 7:e12148.
- 25. Sun, Y., Zhou, D., Rahman, R., **Wu, C.**, Zhu, J., Cox NJ., Beach TG., Wu, C., Gamazon, ER., & Wu, L. (2021). A transcriptome-wide association study identifies novel blood-based gene biomarker candidates for Alzheimer's disease risk. *Human Molecular Genetics*, ddab229.
- Sun, Y., Zhu, J., Zhou, D., Canchi, S., Wu, C., Cox NJ., Rissman, RA. Gamazon, ER., & Wu, L. (2021). A transcriptome-wide association study of Alzheimer's disease using prediction models of related tissues identifies novel candidate susceptibility genes. *Genome Medicine*, 13(1), 1–11.
- 27. Wang, K. S., Yu, G., Xu, C., ..., **Wu, C.**, ... & Deng, H. W. (2021). Accurate diagnosis of colorectal cancer based on histopathology images using artificial intelligence. *BMC Medicine*, 19(1), 1–12.
- 28. Liu, D., Zhu, J., Zhou, D., ..., **Wu, C.**, ..., & Wu, L. (2021). A transcriptome-wide association study identifies novel candidate susceptibility genes for prostate cancer risk. *International Journal of Cancer*, 1–11.
- 29. Zhu, J., **Wu, C.**, & Wu, L. (2021). Associations between genetically predicted protein levels and COVID-19 severity. *The Journal of Infectious Diseases*, 223(1), 19–22.
- 30. Liu, D., Zhou, D., Sun, Y., Zhu, J., Ghoneim, D., **Wu, C.**, Yao, Q., Gamazon, E.R., Cox, N.J., & Wu, L. (2020). A transcriptome-wide association study identifies candidate susceptibility genes for pancreatic cancer risk. *Cancer Research*, 80(20), 4346–4354.
- 31. Wu L., Yang Y., Guo X., ..., **Wu C.**, ..., & Long, J., The Practical Consortium (2020). An integrative multi-omics analysis to identify candidate DNA methylation biomarkers related to prostate cancer risk. *Nature Communications*, 11(1), 1–11.
- 32. Song, M., Greenbaum, J., Luttrell IV, J., Zhou, W., **Wu, C.**, Shen, H., Gong P., Zhang C., & Deng, H. W. (2020). A review of integrative imputation for multi-omics datasets. *Frontiers in Genetics*, 11.
- 33. Zhu, J., Shu, X., Guo, X., ..., **Wu, C.**, ..., & Wu L. (2020). Associations between genetically predicted blood protein biomarkers and pancreatic cancer risk. *Cancer Epidemiology, Biomarkers & Prevention*, 29(7), 1501–1508.
- 34. Zhu, L., Li, Y., Chen, Y., Carrera, C., **Wu, C.**, & Fork, A. (2018). Comparison between two post-dentin bond strength measurement methods. *Scientific Reports*, 8(1), 2350.
- 35. Bose, M., **Wu, C.**, Pankow, J. S., Demerath, E. W., Bressler, J., Fornage, M., Grove, M. L., Mosley, T. H., Hicks, C., North, K., Kao, W. H., Zhang, Y., Boerwinkle, E., & Guan, W. (2014). Evaluation of microarray-based DNA methylation measurement using technical replicates: the Atherosclerosis Risk In Communities (ARIC) Study. *BMC Bioinformatics*, 15(1), 1–10.

-Peer-reviewed Proceedings

- 36. Park, J. Y., **Wu, C.**, & Pan, W. (2018). An adaptive gene-level association test for pedigree data. *BMC Genetics*, 19(1), 68.
- 37. **Wu, C.**, Park, J.Y., Guan, W., & Pan, W. (2018). A powerful gene-based test for methylation data. *BMC Proceedings*, 12(9), 60.

—Book Chapter

38. **Wu, C.** (2021). Using R for cell-type composition imputation in epigenome-wide association studies. Epigenome-Wide Association Studies, Springer, in production.

SUBMITTED MANUSCRIPTS

- * Corresponding author; †Co-first author; ‡Students
- 39. Ma, X., Wang, J.*, & **Wu, C.*** (2021+). Breaking the winner's curse in Mendelian randomization: Rerandomized inverse variance weighted estimator. Alphabetical order; Submitted to *Annals of Statistics*.
- 40. Zhang, Z.[‡], Bae, Y.[‡], Bradley, J., Wu, L, & **Wu, C.*** (2021+). SUMMIT: An integrative approach for better transcriptomic data imputation improves causal gene identification. Submitted. (An earlier version won a <u>poster talk</u> (top 10% of all posters; Reviewers' Choice) at ASHG 2021).
- 41. King, A.[‡], Wu, L., Deng, HW., & **Wu, C.*** (2021+). Polygenic risk score improves the accuracy of a clinical risk score for coronary artery disease. Submitted.
- 42. Guo, X., Wei, W., Liu, M., Cai, T., **Wu, C.**, & Wang, J. (2021+). Assessing heterogeneous risk of type 2 diabetes associated with statins usage: Evidence from electronic health record data. To be submitted. An earlier version is on arXiv.
- 43. Sun, Y., Zhu, J., Yang, Y., Nowakowski, R., Long, J., **Wu, C.**, & Wu, L. (2021+). Identification of candidate DNA methylation biomarkers related to Alzheimer's disease risk by integrative multi-omics analysis. Submitted.
- 44. Sun, Y.[†], Bae, YE.^{†‡}, Zhu, J., **Wu, C.***, & Wu, L.* (2021+) A splicing transcriptome-wide association study identifies novel altered splicing in Alzheimer's disease susceptibility. Submitted.
- 45. Li, Y., Kang, J., **Wu, C.**, Dinov, I., Chalise, P., & Mahnken, J. (2021). A machine-learning approach for detection of local brain networks and marginally weak signals identifies novel AD/MCI differentiating connectomic neuroimaging biomarkers. Submitted. An earlier version is on bioRxiv.
- 46. Hidalgo, B., Lent, S., ..., **Wu, C.**, ..., Hivert, M.F., & Pankow J.S. for the CHARGE consortium (2021+). DNA methylation associated with glycemic traits and type 2 diabetes in multi-ethnic meta-analyses: CHARGE Consortium. *Nature Communications*. Under revision.

GRANTS

—Undergoing

•Novel statistical methods for multi-omics data integration in Alzheimer's disease (Wu, C and Bradley, J)

National Institutes of Health (NIH)

01/01/21–12/31/22 Total cost: \$295,903

NIA 1R03AG070669 Total cost: \$295,903
Role: Contact Pl 20% efforts

-Pending

 Uncovering causal protein markers to improve prostate cancer etiology understanding and risk prediction in Africans and Europeans (Wu, L and Wu, C)

R01

NIH, NCI 04/01/2022–03/31/2027

Role: **PI** 15% efforts in Years 1–2, 20% efforts in Years 3–5

Status: Pending Council Review

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•Discovering causal DNA methylation and protein markers for Alzheimer's disease (Wu, C and Wu, L)

NIH, NIA 04/01/2022–03/31/2027 Role: **Contact PI** 25% efforts

Status: Pending IRG review

•Trans-omics integration of multi-omics studies for osteoporosis: Administrative supplement for COVID-19 studies (Deng, HW)

U19

NIH 09/10/2021–03/31/2022

Role: Subcontract PI 5% efforts

Status: Waiting for the final approval from Tulane University

Subaward cost: \$12,422

•A comprehensive Bridge2Al data generation project (ACB-DGP) for health care and genomics

(Deng, HW)

NIH, Bridge2AI 02/01/2022–01/31/2026

Role: Co-Investigator 5% efforts

Status: Pending IRG review

—Completed

Novel statistical methods for transcriptome-wide association studies

First Year Assistant Professor Grant, Florida State University

05/08/19–08/06/19

Role: contact PI

Total cost: \$20,000

Novel machine learning methods for Alzheimer's disease

The Committee on Faculty Research Support, Florida State University 05/07/20–06/30/20

Role: contact PI Total cost: \$14,000

SOFTWARE

•Owner and maintainer for the following R packages:

-prclust: Penalized Regression-Based Clustering Methods;
21K downloads

-MiSPU: Microbiome Based Sum of Powered Score (MiSPU) Tests;
21K downloads

-aSPU2: A New Version of Adaptive Sum of Powered Score (aSPU) Test;

-GLMaSPU: Adaptive Tests on High Dimensional Parameters in GLMs; 19K downloads

-qlmtlp: Truncated Lasso Regularized Generalized Linear Models; 14K downloads

•Online software tutorial: http://wuchong.org/software.html

•Lab GitHub Repo: https://github.com/ChongWuLab

TEACHING EXPERIENCE

•STA 4321/5323 Introduction to Mathematical Statistics

Instructor, Department of Statistics, Florida State University
 Instructor, Department of Statistics, Florida State University

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•STA 5934 Statistical Genetics (Topic course, redesigned by Wu, C.)

—Instructor, Department of Statistics, Florida State University

Aug. 2020–Dec. 2020

 Instructor, Department of Statistics, Florida State University Aug. 2021-Dec. 2021 •PUBH 6451 Biostatistics (Instructor: Dr. William Thomas) -Lab Instructor, Division of Biostatistics, University of Minnesota Jan. 2014-May 2014 •PUBH 6414 Biostatistical Methods (Instructor: Dr. Ann M. Brearley) -Teaching Assistant, Division of Biostatistics, University of Minnesota Sept. 2013-Dec. 2013 STUDENTS •Ph.D. students who passed the qualifying exam: Shengjie Jiang, Ye Eun Bae, Zhuo Meng, Austin King, Hunter Melton •Other Ph.D. students: Zichen Zhang, Madison Layfield PRESENTATIONS Most of my presentation slides can be found at wuchong.org/talks.html. •"An adaptive association test for microbiome data" -Presentation at Eastern North American Region (ENAR) 2016 Spring Meeting Austin, TX Mar. 2016 –Presentation at 2016 Joint Statistical Meetings (JSM) Chicago, IL Aug. 2016 •"Iterative PCA in epigenome-wide association studies" -Poster presented at American Society of Human Genetics (ASHG) 2016 Annual Meeting Vancouver, BC, Canada Oct. 2016 •"A gene-level adaptive association test for methylation data" -Presentation at Genetic Analysis Workshop (GAW) 20 San Diego, CA Mar. 2017 •"An adaptive test on high dimensional parameters in GLMs" -Presentation at ENAR 2017 Spring Meeting Washington, DC Mar. 2017 -Poster presented at MSI Research Exhibition Minneapolis, MN Apr. 2017 -Presentation at 2017 JSM Baltimore, MD Aug. 2017 •"Imaging-wide association study: integrating imaging endophenotypes in GWAS" -Invited presentation at 2017 JSM (On behalf of Dr. Wei Pan) Baltimore, MD Aug. 2017 -Invited presentation at Third Annual Kliakhandler Conference (On behalf of Dr. Wei Pan) Houghton, MI Aug. 2017 "Integrating eQTL data with GWAS summary statistics in pathway-based analysis" -Poster talk at ASHG 2017 Annual Meeting

Orlando, FL Oct. 2017

•"Integration of methylation QTL and enhancer-target gene maps with schizophrenia GWAS summary results identifies novel genes"

-Oral presentation at IGES 27th Annual Meeting San Diego, CA

Oct. 2018

• "An adaptive test for high-dimensional generalized linear models with application to detect gene- environment interactions"	
Presentation at ENAR 2019 Spring MeetingPhiladelphia, PA	Mar. 2019
 "Complex disease risk prediction via a deep learning method" Presentation at JSM 2019 Denver, CO 	Jul. 2019
• "Multi-trait genome-wide analyses of the brain imaging phenotypes in UK Biobank" —Platform presentation at ASHG 2019	
Huston, TX"A powerful fine-mapping method for transcriptome-wide association studies"	Oct. 2019
-Invited Presentation at JSM 2020 Virtual	Aug. 2020
 "Novel strategy for disease risk prediction incorporating predicted gene expressio methylation: a multi-phased study of prostate cancer" 	n and DNA
 Department seminar at Florida State University School of Medicine Virtual 	Sep. 2020
-Invited presentation at International Indian Statistical Association (IISA) 2021. Virtual	May 2021
•"A gene-level methylome-wide association analysis identifies novel Alzheimer's disease genes" —Poster presented at ASHG 2020	
Virtual	Oct. 2020
 "A regularization-based adaptive test for high-dimensional generalized linear models" Department seminar at Shanghai University of Finance and Economics Statistics 	
Virtual	Dec. 2020
 Department seminar at Washington University Biostatistics Virtual 	Dec. 2020
• "SUMMIT: An integrative approach for better transcriptomic data imputation improves causal gene identification"	
-Invited presentation at New Investigator in AD and AFAR Grantee Conference Virtual	Oct. 2021
HONORS & AWARDS	
•ENAR Distinguished Student Paper Award, —Eastern North American Region International Biometric Society	Mar. 2019
 James R. Boen Student Achievement Award Division of Biostatistics, University of Minnesota 	May 2018
 Pre-Doctoral Trainee Award Association of Chinese Geneticists in America (ACGA) 	Oct. 2017
Poster Award –University of Minnesota Chapter of Sigma Xi	May 2017
•Elected to Delta Omega (Public Health Honorary Society) –University of Minnesota	May 2017
•Elected to Sigma Xi (The Scientific Research Society)	May 2017

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PROFESSIONAL ACTIVITIES

Manuscript Reviewer

- •Statistics/Biostatistics: Biometrics, Journal of the American Statistical Association, Journal of Machine Learning Research, Statistics in Medicine, Electronic Journal of Statistics, Statistical Applications in Genetics and Molecular Biology, Statistics in Biosciences, Journal of Theoretical Biology, Biometrical Journal, Computational Statistics and Data Analysis
- •Genetics/Genomics/Medicine/Bioinformatics: Genetic Epidemiology, Bioinformatics, Genome Biology, Frontiers in Genetics, Alzheimer's & Dementia, Briefings in Bioinformatics, PLOS Computational Biology
- Epidemiology: American Journal of Epidemiology
- •Multidisciplinary: Nature Communications, PLOS One, Scientific Reports

Guest Editor, Frontiers in Genetics

Education Officer, mental health statistics section, American Statistical Association (ASA)

Invited Session Chair & Organizer (with Dr. Jingshen Wang), Recent Advances of Causal Inferences in Human Genetics, JSM 2021

Grant Reviewer

- •NIA Special Emphasis panel, ZAG ZIJ-P J2, Functional genomics, 2020
- •NCI ITCR study section, 2020