CHONG WU Email: cwu3@fsu.edu

# Chong Wu

201C OSB, 117 N. Woodward Ave. P.O. Box 3064330, Tallahassee, FL, 32306-4330

⊠: cwu3@fsu.edu; **☎**: 612-516-2222

http://wuchong.org

## RESEARCH INTEREST

High-dimensional data, big data, machine learning, statistical genetics

#### EDUCATION

University of Minnesota (UMN)

Minneapolis, MN Jun. 2018

**Ph.D.**, **Biostatistics**Advisors: Prof. Weihua Guan & Prof. Wei Pan

GPA: 3.8/4.0

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

Huazhong University of Science & Technology

Wuhan, China

**B.S.**, Applied Mathematics

Jun. 2013

Advisor: Prof. Xiaoyang Zhou

GPA: 3.9/4.0; Ranking: 1/29

## EMPLOYMENT

**Assistant Professor** 

Aug. 2018–Present

Department of Statistics, Florida State University

Biostatistics Intern, Advisor: Dr. Jeff Budd

May 2014–Aug. 2014

Beckman Coulter, Chaska, MN

#### RESEARCH EXPERIENCE

Graduate Research Assistant, Advisor: Prof. Wei Pan

Sept. 2014–Jun. 2018

Research areas: high-dimensional genetic and genomic data, machine learning, and statistical genetics Division of Biostatistics, University of Minnesota

Graduate Research Assistant, Advisor: Prof. Weihua Guan

Sept. 2013-May 2017

Research areas: statistical genetics, and DNA methylation data

Division of Biostatistics, University of Minnesota

Undergraduate Research Assistant, Advisor: Prof. Xiaoyang Zhou

2011–2013

Research areas: data mining on regional economic indicators

School of Mathematics and Statistics, Huazhong University of Science & Technology

#### TEACHING EXPERIENCE

Guest Instructor Jan. 2017–May 2017

Course Name: PUBH 7475/8475 Statistical Learning and Data Mining

Division of Biostatistics, University of Minnesota

**Lab Instructor**, Instructor: Prof. William Thomas Jan. 2014–May 2014

Course Name: PUBH 6451 Biostatistics II Division of Biostatistics, University of Minnesota

**Teaching Assistant**, Instructor: Prof. Ann M. Brearley Sept. 2013–Dec. 2014

Course Name: PUBH 6414 Biostatistical Methods I (Online Section)

Division of Biostatistics, University of Minnesota

### PEER-REVIEWED PUBLICATIONS

- 1. Steven Nguyen, S., Guan, W., **Wu, C.**, Grove, M.L., Xia, R., Roetker, N., Holliday, K., Hibler, E., Zheng, Y., Whitsel, E., Bressler, J., North, K.E., Fornage, M., Boerwinkle, E., Pankow, J.S., Demerath, and E.W. (2018) Epigenome-wide association study of moderate-vigorous physical activity in adult African-Americans identifies loci near HCCA2. *Circulation*, 137 (Suppl 1).
- 2. **Wu, C.** and Pan, W. (2018). Integration of enhancer-promoter interactions with GWAS summary results identifies novel schizophrenia-associated genes and pathways. Accepted by *Genetics*, early online.
- 3. **Wu, C.\***, Xu, G., and Pan, W.\* (2017+). An adaptive test on high dimensional parameters in generalized linear models. Accepted by *Statistica Sinica*. (\* Corresponding author)
- 4. Zhu, L., Li, Y., Chen, Y., Carrera, C., **Wu, C.**, and Fork, A. (2018). Comparison between two post-dentin bond strength measurement methods. *Scientific Reports*, 8(1):2350.
- Wu, C. and Pan, W. (2018). Integrating eQTL data with GWAS summary statistics in pathway-based analysis. Accepted by *Genetic Epidemiology*, early online.
   (Highlight by IGES. This paper won a poster talk (top 24 posters among about 3000 posters) at ASHG 2017 Annual Meeting.)
- 6. Park, J.Y., **Wu, C.**, Basu, S., McGue, M., and Pan, W. (2018). Adaptive SNP set association testing in generalized linear mixed models with application to family studies. *Behavior Genetics*, 48(1):55–66. (IF: 2.4)
- 7. Xu, Z., **Wu, C.**, Wei, P., and Pan, W. (2017+). A powerful framework for integrating eQTL and GWAS summary data. Accepted by *Genetics*, early online.
- 8. Liu, B., **Wu, C.**, Shen, X., and Pan, W. (2017). A novel and efficient algorithm for de novo discovery of mutated driver pathways. *Annals of Applied Statistics*, 17(3):1481–1512.
- 9. Xu, Z., **Wu, C.**, Pan, W., and 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 American Society of Human Genetics (ASHG)

- 2017 Annual Meeting.)
- 10. **Wu, C.\***, Kwon, S.\*, Shen, X., and Pan, W. (2016). A new algorithm and theory for penalized regression-based clustering. *Journal of Machine Learning Research*, 17(188):1–25. (\* Co-first author. A leading journal in machine learning area.)
- Wu, C., Chen, J., Kim, J., and 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.)
- 12. **Wu, C.**, Demerath, E. W., Pankow, J. S., Bressler, J., Fornage, M., Grove, M. L., Chen, W., and Guan, W. (2016). Imputation of missing covariate values in epigenome-wide analysis of DNA methylation data. *Epigenetics*, 11(2):132–139. (The official journal of the DNA Methylation Society.)
- 13. 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., and 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

- 14. Park, J. Y., **Wu, C.**, and Pan, W. (2017+). An adaptive gene-level association test for pedigree data. Accepted by *BMC Proceedings*, a special issue for the Genetic Analysis Workshop (GAW 20).
- 15. **Wu, C.**, Park, J.Y., Guan, W., and Pan, W. (2017+). A powerful gene-based test for methylation data. Accepted by *BMC Proceedings*, a special issue for the Genetic Analysis Workshop (GAW 20).

### SUBMITTED AND IN PREPARATION

- 16. **Wu, C.**, Xu, G., Shen, X., and Pan, W. (2017+). An adaptive test on a high-dimensional parameter in the presence of a high-dimensional nuisance parameter in GLM with application to detect gene-environment interactions. Manuscript.
- 17. Gaeckle, N., Pragman A., Gould T., **Wu, C.**, and Criner, G. (2017+). Current smoking alters the oral microbiome in subjects with COPD. Submitted.
- 18. **Wu, C.**, Xu, G., and Pan, W. An adaptive and powerful test for high dimensional covariance matrices. In Preparation.
- 19. Wu, C. and Pan, W. Network-based support vector machines with a new penalty. In Preparation.
- 20. **Wu, C.**, Kim, J., Wei, P., and Pan, W. Adaptive test for meta-analysis of rare variants in sequencing association studies. In preparation.

# SOFTWARE

- •Owner and maintainer for the following R packages:
  - -prclust: Penalized Regression-Based Clustering Methods;
  - -MiSPU: Microbiome Based Sum of Powered Score (MiSPU) Tests;
  - -aSPU2: A New Version of Adaptive Sum of Powered Score (aSPU) Test;
  - -GLMaSPU: Adaptive Tests on High Dimensional Parameters in Generalized Linear Models;
  - -glmtlp: Truncated Lasso Regularized Generalized Linear Models.
- •Contributor: Theano (Pull request: #6130).

#### PRESENTATIONS

TRESENTATIONS		
<ul> <li>"An adaptive association test for microbiome data."</li> <li>Presentation at Eastern North American Region (ENAR) 2016 Spring Meeting Austin, TX</li> </ul>	Mar.	2016
<ul> <li>Poster presented at School of Public Health Research Day, University of Minnesota Minneapolis, MN</li> </ul>	Apr.	2016
<ul> <li>Poster presented at Minnesota Supercomputing Institute (MSI) Research Exhibition Minneapolis, MN</li> </ul>		2016
–Presentation at 2016 Joint Statistical Meetings (JSM) Chicago, IL	Aug.	2016
<ul> <li>"Iterative PCA in epigenome-wide association studies."</li> <li>Poster presented at American Society of Human Genetics (ASHG) 2016 Annual M</li> <li>Vancouver, BC, Canada</li> </ul>	_	2016
<ul> <li>"A gene-level adaptive association test for methylation data."</li> <li>Presentation at Genetic Analysis Workshop (GAW) 20</li> <li>San Diego, CA</li> </ul>	Mar.	2017
<ul> <li>"An adaptive test on high dimensional parameters in GLMs."</li> <li>Presentation at ENAR 2017 Spring Meeting</li> <li>Washington, DC</li> </ul>	Mar.	2017
–Poster presented at MSI Research Exhibition Minneapolis, MN	Apr.	2017
–Presentation at 2017 JSM Baltimore, MD	Aug.	2017
<ul> <li>"Imaging-wide association study: integrating imaging endophenotypes in GWAS"</li> <li>Invited presentation at 2017 JSM (On Prof. Wei Pan's behalf)</li> <li>Baltimore, MD</li> </ul>	Aug.	2017

-Invited presentation at Third Annual Kliakhandler Conference (On Prof. Wei Pan's behalf)

Houghton, MI	Aug. 2017
<ul> <li>"Integrating eQTL data with GWAS summary statistics in pathway-based analysis"</li> <li>Poster talk at ASHG 2017 Annual Meeting</li> <li>Orlando, FL</li> </ul>	Oct. 2017
HONORS & AWARDS	
	O . 2017
<ul> <li>Pre-Doctoral Trainee Award</li> <li>Association of Chinese Geneticists in America (ACGA)</li> </ul>	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)  –University of Minnesota	May 2017
•Distinguished Student Paper Award, Genomics and Genetics Section –2016 Joint Statistical Meetings	Aug. 2016
•Doctoral Dissertation Fellowship  –University of Minnesota	2016–2017
•Travel Award, Computational Neuroscience Summer School —Statistical and Applied Mathematical Sciences Institute	Jul. 2015
•Dean's Ph.D. Scholarship  —University of Minnesota	Sept. 2013
•Honorable Mention in Mathematical Contest in Modeling  —Consortium for Mathematics and Its Application	Apr. 2012
•National Scholarship —Ministry of Education, China	Sept. 2011
PROFESSIONAL ACTIVITIES	
Manuscript Reviewer  •Journal: Statistics in Biosciences, Journal of Theoretical Biology, Biometrical Journal Statistics and Data Analysis, Genetic Epidemiology, Bioinformatics	rnal, Computa-
•Conference: 2017 Joint Statistical Meetings Student Paper Award Reviewer	
Professional Memberships  •Member, American Statistical Association	2014–Present
•Member, Eastern North American Region	2015–Present
•Member, The American Society of Human Genetics	2016–Present
Programming Skills	

•R, Python, MATLAB, C++