

# Feifei Xiao, PhD

## CONTACT INFORMATION

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Department of Biostatistics  
College of Public Health and Health Professions & College of Medicine  
University of Florida  
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## EDUCATION

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06/2013	<b>Ph.D.</b> in Biomathematics and Biostatistics <b>The University of Texas Health Science Center MD Anderson Cancer Center</b> , Graduate School of Biomedical Sciences, Houston, TX, USA Mentor: Christopher I. Amos, PhD Dissertation: <i>Natural and orthogonal interaction framework for modeling gene-gene/gene-environmental interactions and imprinting effects.</i>
06/2009	<b>M.S.</b> in Microbiology <b>Wuhan University</b> , Wuhan, China State Key Laboratory of Virology College of Life Sciences
06/2006	<b>B.S.</b> in Biology <b>Wuhan University</b> , Wuhan, China College of Life Sciences

## POST-GRADUATE TRAINING

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06/2013 – 07/2015	<b>Postdoctoral Research Associate</b> <b>Yale School of Public Health</b> , New Haven, CT, USA Department of Biostatistics, School of Public Health Mentor: Heping Zhang, PhD Projects: <i>Algorithms and methods for copy number variation detection; women and children health.</i>
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**EMPLOYMENT**

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01/2022 – present	<b>Associate Professor</b> Department of Biostatistics, College of Public Health and Health Professions & College of Medicine, <b>University of Florida</b> , Gainesville, FL, USA
07/2021 – 01/2022	<b>Associate Professor</b> Department of Epidemiology and Biostatistics, Arnold School of Public Health, <b>University of South Carolina</b> , Columbia, SC, USA
05/2017 – 07/2017	<b>Visiting Assistant Professor</b> Department of Biomedical Data Science, Geisel School of Medicine at <b>Dartmouth College</b> , Lebanon, NH, USA
08/2015 – 06/2021	<b>Assistant Professor</b> Department of Epidemiology and Biostatistics, Arnold School of Public Health, <b>University of South Carolina</b> , Columbia, SC, USA

**MAJOR RESEARCH INTERESTS**

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Methodology and Application in:

Statistical Genetics	Multi-Omics Integration
Single Cell Sequencing	Machine Learning
Cancer Omics	Bioinformatics

**RESEARCH AWARDS AND GRANTS**

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**Ongoing*****The evolution of copy number variations in the AT-rich Plasmodium genome***

R01AI150856                      Guler(PI)                      04/01/2021 – 03/31/2026

NIH/NIAID

Role: Co-Investigator (15% of effort in first three years, plus 1 RA each year)

Total Subaward: \$219,494

***Dysfunctional Myelopoiesis and Myeloid-Derived Suppressor Cells in Sepsis Pathobiology***

RM1GM139690-03                      Moldawer (PI)                      05/01/2021 - 4/30/2026

NIH/NIGMS

Role: Co-Investigator (25% of effort)

Total Award: \$1,689,645

**Completed**

***Copy Number Variation and Lung Cancer: Disease Risk and Mechanisms***

R21HG010925

Xiao (PI)

09/01/2020 - 08/30/2023

NIH/NHGRI

Role: Principal Investigator (25% of effort, plus 1 RA each year)

Total Award: \$426,122

***Assessing safety and mobility benefits***

Classen (PI) 10/01/2022 – 09/30/2023

US Department of Veteran Affairs Gainesville Medical Center

Role: Co-Investigator (5% of effort)

Total Subaward: \$9,687

***Molecular Basis of Exercise-induced Changes in HDL Function***

R01 HL146462

Sarzynski (PI)

04/01/2019 - 03/31/2024

NIH/NHLBI

Role: Co-Investigator (year 1-4: 5%, 5%, 10%, 15% of 9-month, year 5: 25% of 12-month)

Total Award: \$3,466,346

*\*Early termination in 01/15/2022 after relocation to UF****Aging Brain Cohort at the University of South Carolina (ABC@USC)***

USC Excellence Initiative Award Fridriksson (PI)

07/01/2019 - 06/30/2023

Role: Co-Investigator (18% of effort)

Total Award: \$1,750,000

*\* Early termination 01/15/2022 after relocation to UF****A Novel Statistical Algorithm for Copy Number Variation Detection with Application to Lung Cancer***

USC Office of the Vice President for Research Xiao (PI)

07/01/2020 - 9/30/2021

ASPIRE-I Innovation grant: Advanced Support for Innovative Research Excellence

Role: Principal Investigator

Total Direct Cost: \$14,964

***Mechanisms of Early Onset Colorectal Cancer (EOCRC)***

USC Office of the Vice President for Research Hofseth (PI)

07/01/2020 - 9/30/2021

ASPIRE-II Integration grant: Advanced Support for Integration Research Excellence

Role: Co-Investigator

Total Direct Cost: \$84,105

***Medicare Shared Savings Cost Determinants Analysis, 2015-2017***

ASPH Dean's office

Rudisill (PI)

07/01/2020 - 06/30/2021

Role: Co-Investigator (8% of effort)

Total Direct Cost: \$14,315



2. Qin F#, Cai G, Amos CI, **Xiao F\***. A statistical learning method for simultaneous copy number estimation and subclone clustering with single-cell sequencing data. *Genome Research*. 2024. 34(1):85-93.
3. Sun Z, Neelon B, Ethier SP, **Xiao F**, Wallace K, Chung D. A Bayesian framework for pathway-guided identification of cancer subgroups by integrating multiple types of genomic data. *Statistics in Medicine*. 2023;42(28):5266-5284.
4. Islam F, Thrasher JF, **Xiao F**, Moran RR, Hardin JW. Data Management and Techniques for Best-Worst Discrete Choice Experiments. *The Stata Journal Promoting communications on statistics and Stata*. 2023; 23(4):1020-1044.
5. Luo X#, Qin F#, **Xiao F**, Cai G. BISC: accurate inference of transcriptional bursting kinetics from single-cell transcriptomic data. *Briefing in Bioinformatics*. 2022;23(6).
6. Luo X#, Cai G, McLain AC, Amos CI, Cai B, **Xiao F\***. BMI-CNV: a Bayesian framework for multiple genotyping platforms detection of copy number variants. *Genetics*. 2022; 222(4).
7. Cai G, Yu X#, Youn C, Zhou J, **Xiao F\***. SCANNER: A web server for annotation, visualization and sharing of single cell RNA-seq data. *Database* (2022). Online.
8. Qin F#, Luo X#, **Xiao F**, Cai G. SCRIP: an accurate simulator for single-cell RNA sequencing data. *Bioinformatics*. 2022; 38(5):1304-1311
9. Cui X, Qin F#, Yu X, **Xiao F**, Cai G. SCISSOR™: a single-cell inferred site-specific omics resource for tumor microenvironment association study. *NAR Cancer*. 2021; 3(3): zcab037.
10. Cai G, Du M, Bossé Y, Albrecht H, Qin F#, Luo X#, Androulakis M, Yi C, Cheng C, Nagarkatti M, Nagarkatti P, Christiani DC, Whitfield ML, Amos CI, **Xiao F\***. SARS-CoV-2 Impair Dendritic Cells and Regulate DC-SIGN Expression in Tissues. *International Journal of Molecular Sciences*. 2021; 22(17): 9228.
11. Qin F#, Luo X#, Cai G, **Xiao F\***. Shall genomic correlation structure be considered in copy number variation detection? *Briefings in Bioinformatics* (2021). Online.
12. Liang Q, Tan C, Xiao F, Yin F, Liu M, Lei L, Wu L, Yang Y, Tan HJ, Liu S, Zeng X. Integrated profiling identifies ITGB3BP as prognostic biomarker for hepatocellular carcinoma. *Bosnian Journal of Basic Medical Sciences*. 2021, 21(6):712-723.

13. Cai G, Zhu X, Charvet L, **Xiao F**, Datta A, Androulakis XM. A Systematic review and meta-analysis on the efficacy of transcranial direct current stimulation for migraine. *Journal of Pain Research*. 2021. 14:1171-1183.
14. Luo X#, Qin F#, Cai G, **Xiao F\***. Integrating genomic correlation structure improves copy number variations detection. *Bioinformatics*. 2021. 37(3):312-317.
15. Zhou J, Cui X, **Xiao F**, Cai G. A cluster-based approach for identifying prognostic microRNA signatures in digestive system cancers. *International Journal of Molecular Sciences*. 2021. 22(4):1529.
16. Kristinsson S, Zhang W, Rorden C, Newman-Norlund R, Basilakos A, Bonilha L, Yourganov G, **Xiao F\***, Hillis A, Fridriksson J. Machine learning-based multimodal prediction of language outcomes in chronic aphasia. *Human Brain Mapping*. 2021. 42(6):1682-1698.
17. Hao N, Niu YS, **Xiao F**, Zhang H. A super scalable algorithm for short segment detection. *Statistics in Biosciences*. 2021. 13(1):18-33.
18. Cai G, Bossé Y, **Xiao F**, Kheradmand F, Amos CI. Tobacco smoking increases the lung gene expression of ACE2, the receptor of SARS-CoV-2. *American Journal of Respiratory and Critical Care Medicine*. 2020. 201(12):1557-1559.
19. Wang L#, Luo X#, Cheng C, Amos CI, Cai G, **Xiao F\***. A gene expression based immune signature for lung adenocarcinoma prognosis. *Cancer Immunology and Immunotherapy*. 2020. 69(9):1881-1890.
20. Deng S, Hardin J, Amos CI, **Xiao F\***. Joint modeling of eQTLs and parent-of-origin effects using an orthogonal framework with RNA-seq data. *Human Genetics*. 2020.139(8):1107-1117.
21. Liu M, Liu X, Liu S, **Xiao F**, Guo E, Qin X, Wu L, Liang Q, Liang Z, Li K, Zhang D, Yang Y, Luo X, Lei L, Tan HJ, Yin F, Zeng X. Big data-based identification of multi-gene prognostic signatures in liver cancer. *Frontiers in Oncology*. 2020. 10:847.
22. Kristinsson S, **Xiao F**, Yourganov G, Bonilha L, Stark BC, Rorden C, Basilakos A, Fridriksson J. BDNF Genotype Specific Differences in Cortical Activation in Chronic Aphasia. *Journal of Speech, Language, and Hearing Research*. 2019. 62(11):3923-3936.
23. **Xiao F\***, Luo X#, Hao N, Niu YS, Xiao X, Cai G, Amos CI, Zhang H. An accurate and powerful method for copy number variation detection. *Bioinformatics*. 2019. 35(17):2891-2898.
24. Zhao Y, Varn F, Cai G, **Xiao F**, Amos CI, and Cheng C. A P53-deficiency gene signature predicts recurrence risk of patients with early stage lung adenocarcinoma. *Cancer Epidemiology, Biomarkers & Prevention*. 2018; 27(1):86-95.

25. Bukowski R, Sadovsky Y, H. Goodarzi, Zhang H, Biggio JR, Varner M, Parry S, **Xiao F**, and et al. Onset of human preterm and term birth is related to unique inflammatory transcriptome profiles at the maternal fetal interface. *Peer J*. 2017; 5:e3685.
26. Cai G, Zheng X, Liang S, **Xiao F\***. Local sequence and sequencing depth dependent accuracy of RNA-seq reads. *BMC Bioinformatics*. 2017; 8(1): 364.
27. **Xiao F**, Niu Y, Hao N, Xu Y, Jin Z, Zhang H. modSaRa: a computationally efficient R package for CNV identification. *Bioinformatics*. 2017; 33(15):2384-2385.
28. Cai G, **Xiao F**, Cheng C, Li Y, Amos CI, Whitfield ML. Population effect model identifies gene expression predictors of survival outcomes in lung adenocarcinoma for both Caucasian and Asian patients. *PLoS One*. 2017; 12(4):e0175850.
29. **Xiao F**, Cai G, Zhang H. Segregation analysis suggests that a genetic reason may contribute to "the dress" colour perception. *PLoS One*. 2016; 11(10):e0165095.
30. **Xiao F**, Min X, Zhang H. Modified screening and ranking algorithm for copy number variant detection. *Bioinformatics*. 2015; 31(9):1341-8.
31. Zhang H, Baldwin DA, Bukowski RK, Parry S, Xu Y, Song C, Andrews WW, Saade GR, Esplin MS, Sadovsky Y, Reddy UM, Ileakis J, Varner M, Biggio JR Jr; Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD) Genomic and Proteomic Network for Preterm Birth Research (GPN-PBR) cohort (including **Xiao F**). A genome-wide association study of early spontaneous preterm delivery. *Genet Epidemiol*. 2015; 39(3):217-26.
32. Manuck TA, Esplin MS, Biggio J, Bukowski R, Parry S, Zhang H, Huang H, Varner MW, Andrews W, Saade G, Sadovsky Y, Reddy UM, Ileakis J; Eunice Kennedy Shriver NICHD GPN-PBR cohort (including **Xiao F**). The phenotype of spontaneous preterm birth: application of a clinical phenotyping tool. *Am J Obstet Gynecol*. 2015; 212(4):487.
33. Parry S, Zhang H, Biggio J, Bukowski R, Varner M, Xu Y, Andrews WW, Saade GR, Esplin MS, Leite R, Ileakis J, Reddy UM, Sadovsky Y, Blair IA; Eunice Kennedy Shriver NICHD GPN-PBR cohort (including **Xiao F**). Maternal serum serpin B7 is associated with early spontaneous preterm birth. *Am J Obstet Gynecol*. 2014; 211(6):678.e1-12.
34. **Xiao F**, Ma J, Cai G, Fang S, Lee JE, Wei Q, Amos CI. Natural and orthogonal interaction model for estimating gene-gene interactions: applied to cutaneous melanoma. *Human Genetics*. 2014; 133(5):559-74.
35. **Xiao F**, Ma J, Amos CI. A unified framework integrating parent-of-origin effects for association study. *PLoS ONE*. 2013; 8(8):e72208.

36. Ma J, **Xiao F**, Xiong M, Andrew AS, Brenner H, Duell EJ, Haugen A, Hoggart C, Hung RJ, Lazarus P, Liu C, Matsuo K, Mayordomo JI, Schwartz AG, Staratschek-Jox A, Wichmann E, Yang P, Amos CI. Natural and orthogonal interaction framework for modeling gene-environment interactions with application to lung cancer. *Human Heredity*. 2012; 73(4):185-94.
37. Zhou S, Xiao W, Wan Q, Yi C, **Xiao F**, Liu Y, Qi Y. Nogo-B mediates HeLa cell adhesion and motility through binding of Fibulin-5. *Biochemical and biophysical research communications*. 2010; 398(2):247-53.
38. **Xiao F**, Zuo Z, Cai G, Kang S, Gao X, Li T. miRecords: an integrated resource for microRNA-target interactions. *Nucleic acids research*. 2009; 37:D105-10.
39. Ren Y, Gong W, Zhou H, Wang Y, **Xiao F**, Li T. siRecords: a database of mammalian RNAi experiments and efficacies. *Nucleic acids research*. 2009; 37: D146-9.
40. Gong W, Zhou D, Ren Y, Wang Y, Zuo Z, Shen Y, **Xiao F**, Zhu Q, Hong A, Zhou X, Gao X, Li T. PepCyber: P~PEP: a database of human protein-protein interactions mediated by phosphoprotein-binding domains. *Nucleic acids research*. 2008; 36: D679-83.

**Under review or submitted** (\* indicates corresponding author; # indicates advisee):

41. Qin F#, Luo X#, Cai B, **Xiao F**, Cai G. Spatial pattern and differential expression analysis with spatial transcriptomic data. *Nucleic Acid Research*. Under Revision.
42. Rudisill A., Qin F#, Chapman C, **Xiao F**. Data-led policy design using Medicare Shared Savings Program (MSSP) health care cost trajectories. In preparation.

## HONORS AND AWARDS

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- North Coast Conference on Precision Medicine Travel Award, 2018
- March of Dimes Award for Best Research in Prematurity, SMFM's Meeting, 2015
- NIGMS Short Course on Statistical Genetics & Genomics Travel Award, 2014
- City Federation of Women's Clubs Endowed Scholarship in Biomedical Sciences, 2013
- Summer Institute of Statistical Genetics (SISG) Travel Award, 2011
- Scholarship for Excellent Graduate Students, Wuhan University, 2006-2009
- Outstanding Undergraduate Student Scholarship, Wuhan University, 2005 & 2006
- Superior Academic Activities Award, Wuhan University, 2004
- Outstanding Freshman Scholarship, Wuhan University, 2002

## SHORT COURSES/EDUCATIONAL WORKSHOPS

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- Arnold School of Public Health NIH grant writing bootcamp. University of South Carolina. 2020-2021.



- USC Center for Teaching Excellence Workshop: *Active Learning Made Easy*. Columbia, SC, November 2018.
- Educational Workshop on Genetic Epidemiology: *Fundamentals for Precision Medicine*, IGES meeting, Cambridge, UK, September 2017
- USC Center for Teaching Excellence New Faculty Academy Workshop: *Stress Reduction and Resilience: Strategies for Busy Faculty Member*. Columbia, SC, December 2016
- Educational Workshop: *Big data phenotyping: opportunities, analytic challenges and solutions*, IGES meeting, Toronto, ON, Canada, October 2016
- Statistical Genetics and Genomics, University of Alabama, Birmingham, AL, July 2014
- Summer Institute in Statistical Genetics, University of Washington, Seattle, WA, May 2011

## TEACHING EXPERIENCE

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### University of Florida

#### *Instructor*

PHC 6937    Causal Inference    *Spring 2023-2024*

### University of South Carolina

#### *Instructor*

BIOS 759	Theory and Methods of Discrete Data Analysis	<i>Fall 2021</i>
BIOS 894	Topics: Statistical Methods in Bioinformatics	<i>Spring 2021</i>
BIOS 805	Categorical Data Analysis	<i>Fall 2020</i>
STAT 770	Categorical Data Analysis	<i>Fall 2020</i>
BIOS 794	Topics: Biostatistical Modeling in Genetic Data Analysis	<i>Fall 2019</i>
BIOS 757	Intermediate Biostatistics	<i>Fall 2017, 2018, 2019</i>
BIOS J757	Intermediate Biostatistics (Distance)	<i>Spring 2018, 2019</i>
BIOS 890	Independent Study of Teaching Practicum	<i>Fall 2017, Fall 2019</i>
BIOS 890	Independent Study of Research Practicum	<i>Summer 2018, Spring 2021</i>
BIOS 794	Special topics in Biostatistics-Statistical Genetics	<i>Spring 2017</i>
BIOS 700	Introduction to Biostatistics	<i>Spring 2016, Fall 2016</i>

#### *Guest Lectures*

EPID 777	Fundamentals of Genetic Epidemiology	<i>Spring 2021</i>
EPID 800	Epidemiology Methods II	<i>Fall 2015, 2016, 2017, 2019</i>
BIOS 745	Seminar in Biostatistics	<i>Spring 2018</i>
ENHS 793	Bioinformatics and Computational Biology	<i>Fall 2018, Fall 2019</i>

## MENTORING EXPERIENCE (University of Florida)

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**PhD Dissertation Advisee***Ongoing*

- Dayuan Wang, PhD candidate in Biostatistics 08/2022 – now

**Doctoral Dissertation Committee***Ongoing*

- Chen Bai, PhD in Biomedical Informatics and Health Outcomes (advisor: Dr. Mamoun Mardini) 06/2023 – now  
Title: *Predicting frailty of older adults and investigating the association of predicted frailty with their health outcomes*

**Academic Advisor (for advice on courses and other academic activities)***Ongoing*

- Vinai Modem, MS candidate in Biostatistics
- Zhongping He, MS candidate in Biostatistics
- Dayuan Wang, PhD candidate in Biostatistics

**MENTORING EXPERIENCE (University of South Carolina)**

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**PhD Dissertation Advisee***Completed*

- Xizhi Luo, PhD in Biostatistics Defense: 07/2021  
Topic: *Accurate and integrative detection of copy number variants with high-throughput data*
- Fei Qin, PhD candidate in Biostatistics Defense: 06/2023  
Topic: *Statistical Methods for Single Cell Sequencing Data Analysis*

*Ongoing*

- Xuanxuan Yu, PhD candidate in Biostatistics 08/2019 – now

**Doctoral Dissertation Committee***Completed*

- Rahimi Gelareh, PhD in Biostatistics (advisor: James Hardin) Defense: 08/2018  
Title: *Adjusting for mis-reporting in count data*
- Xiangyang Cao, PhD in Statistics (U of SC, advisor: Dr. Karl Gregory), Defense: 08/2018  
Title: *High-dimensional inference based on the Leave-One-Covariate-Out LASSO path.*
- Yuan Hong, PhD candidate in Biostatistics (advisor: Dr. Alexander McLain), Defense: 11/2020  
Title: *Incorporation and Measurement of Uncertainty in Longitudinal and*

*Spatial Data*

- Zequn Sun, PhD candidate in Biostatistics (MUSC, advisor: Dr. Dongjun Chung),  
Defense: 07/2021  
Title: *Bayesian latent factor model by integrating multiple types of genomic data*
- Farahnaz Islam, PhD candidate in Biostatistics (advisor: Dr. James Hardin)  
Defense: 09/2022 Title: *Extensions of discrete choice experiment theory for public health*

**Master's Thesis Committee***Completed*

- Andrew Fogner, MPH in Epidemiology (advisor: Dr. Myriam Torres), Defense: 05/2016  
Thesis title: *Prenatal Depression in South Carolina Latinos*
- Danielle Sill, MSPH in Epidemiology (advisor: Dr. Myriam Torres), Defense: 03/2017  
Thesis title: *The association between sexual risk behaviors of Latinos and HIV Knowledge in South Carolina*
- Amanda Collins, MSPH in Epidemiology (advisor: Dr. Anwar Merchant), Defense: 07/2018  
Title: *Effect Modification of Stroke in The Relationship Between Tooth Loss and Cognitive Functioning*
- Diana Diaz, MPH in Epidemiology (advisor: Dr. Susan Steck), Defense: 07/2019  
Title: *Association between SNPs in the vitamin D binding protein, vitamin D status, and aggressive prostate cancer by race*
- Brittany Crawford, MSPH in Epidemiology (advisor: Dr. Susan Steck), Defense: 03/2020  
Title: *Serum Calcium, Phosphorus, and PTH and Prostate Cancer Aggressiveness*
- Haley Davis-Martin, MSPH in Epidemiology (advisor: Dr. Matthew Lohman),  
Defense: 04/2020  
Title: *The interaction between caffeine consumption, alcohol use, and amount of sleep on bone health*
- Marie Knoll, MSPH in Epidemiology (advisor: Dr. Susan Steck) Defense: 06/2020  
Title: *Association between urinary enterolignans as a marker for gut microbiome diversity and depression in NHANES*
- Huizhong Yang, MPH in Biostatistics (advisor: Dr. Bo Cai) Defense: 06/2020  
Title: *Bayesian Zero Inflation Model for Ordinal Data*
- Wanfang Zhang, MPH candidate in Biostatistics (advisor: Dr. Bo Cai)  
Title: *Multiple frailty model for spatially correlated interval-censored data*  
Defense: 09/2021
- Avery Ulrich, MSPH candidate in Epidemiology (advisor: Dr. Alyssa Clay-Gilmour),  
Title: *Associations of the FTO gene and risk of Acute Myeloid Leukemia.*  
Defense: 03/2022

**Academic Advisor (for advice on courses and other academic activities)***Completed*

- Huizhong Yang, MS in Biostatistics, 2020
- Fei Qin, PhD candidate in Biostatistics, 2023

*Ongoing*

- Xuanxuan Yu, PhD candidate in Biostatistics

### Undergraduate Thesis Advisee (Visiting scholars from Nanjing Medical University, China)

- Zilong Bian, Title: *DNA methylation-based signature prediction model for cancer* 02/2020-06/2020
- Jingting Lu, Title: *CNV based signature prediction model for cancer*. 02/2020-06/2020
- Jiani Xu, Title: *Comprehensive Analysis of Somatic Copy Number Aberrations and Gene Expression in Lung Adenocarcinoma* 01/2018-06/2018
- Yaqian Liu, Title: *Integrative analysis of Copy number aberrations in Esophageal Cancer* 01/2018-06/2018
- Lijuan Wang, Title: *Differential expression of immunology pathway related genes in Lung carcinoma* 01/2017-06/2017
- Lu Xu, Title: *Implications of immunotherapy in breast cancer treatment*. 01/2017-06/2017

### CONFERENCE PRESENTATIONS

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1. A statistical learning method for simultaneous copy number estimation and subclone clustering with single cell sequencing data. International Chinese Statistical Association. *Invited Talk*. Chengdu, China, July 2023.
2. A statistical learning method for simultaneous copy number estimation and subclone clustering with single cell sequencing data. American Association of Cancer Research, *Poster Presentation*. Orlando, FL, April 2023.
3. Integrating Genomic Correlation Structure Improves Copy Number Variations Detection. JSM Virtual Meeting, *Student's contributed talk*. August 2021.
4. SCRIP: An Accurate Simulator for Single-Cell RNA Sequencing Data. JSM Virtual Meeting, *Student's contributed talk*. August 2021.
5. Data-led policy design using Medicare Shared Savings Program (MSSP) health care cost trajectories. *Abstract*. Academy Health, 2021.
6. Multimodal neuroimaging prediction models carry complementary information that can be harnessed to enhance prediction of language function in aphasia. *Abstract*. Society for the Neurobiology of Language, 2020.
7. Joint modeling of eQTLs and parent-of-origin effects with RNA-seq data. *Contributed*. JSM Virtual Meeting, August 2020.
8. Adjusting for misreporting in count data. *Abstract*. ASA Women in Statistics and Data Science Conference, Bellevue, Washington, October 2019.
9. An accurate and powerful method for copy number variation detection. *Invited Poster Presentation*. Houston, TX, October 2019.

10. Assessment of statistical batch effect correction methods for different RNA sequencing data types. *Abstract*. Houston, TX, October 2019.
11. BDNF Genotype Specific Differences in Cortical Activation in Chronic Aphasia. *Abstract*. Society for the Neurobiology of Language, Helsinki, Finland, August 2019.
12. LDSaRa: A Powerful Method for High Resolution Copy Number Variation Detection. *Invited talk*. International Chinese Statistical Association (ICSA), June 2019.
13. Common genetic variations in the calcium sensing receptor (CaSR) gene, plasma 25-hydroxyvitamin D and aggressive prostate cancer in the North Carolina-Louisiana Prostate Cancer (PCaP) study. *Abstract*. Annual Meeting-American Society of Preventive Oncology, March 2019.
14. LDSaRa: a powerful tool for high resolution copy number variation detection. *Abstract*. South Carolina Chapter American Statistical Association Annual Meeting, March 2019.
15. Copy number variation detection with complex genetic data. *Invited talk*, American Statistics Association South Carolina Chapter, Clemson, SC, October 2018.
16. Improving copy number estimation by incorporating BAF using modSaRa. *Contributed talk*, Joint Statistical Meeting (JSM), Vancouver, Canada, August 2018.
17. Integrating genetic source improves power for copy number variation detection. *Invited talk*, ICSA, Qingdao, China, July 2018.
18. Integrating genetic source boosts power for copy number variation detection. *Invited talk*, *International Statistics Forum*, Beijing, China, July 2018.
19. A super scalable algorithm for short segment detection. *Abstract*. International Conference on Econometrics and Statistics, Hongkong, China. June 2018.
20. Analyses of germline copy number variation and gene expression in cutaneous melanoma. *Invited talk*, International Genetic and Epidemiology Society (IGES), Cambridge, UK, Sep 2017.
21. Efficient epigenetic effect identification in eQTL mapping with RNA-seq data. *Invited talk*, International Conference on Computational Methods (ICCM), Guilin, Guangxi, China, July 2017.
22. Modeling parent-of-origin effect in eQTL mapping using RNA-seq data. *Invited talk*, ICSA, Chicago, IL, June 2017.
23. Testing of parent-of-origin effect in eQTL mapping using RNA-seq data. *Invited talk*, IGES, Toronto, ON, Canada, Oct 2016.
24. A change-point based method for copy number variation detection with application to melanoma. *Invited talk*, ICSA, Atlanta, GA, June 2016.
25. SNPs in vitamin D-related genes are associated with prostate cancer aggressiveness in the North Carolina-Louisiana Prostate Cancer Project (PCaP). *Abstract*, American Association for Cancer Research, New Orleans, LA, April 2016.
26. A new change-point model based method for copy number variation detection. *Invited Poster presentation*, American Society of Human Genetics (ASHG), Baltimore, MD, USA, Oct 2015.

27. Modified screening and ranking algorithm for copy number variation detection. *Invited Poster presentation*, IGES, Baltimore, MD, USA, Oct 2015.
28. Using expression data to define patient specific predictors for survival outcomes in lung adenocarcinoma. *Abstract*, ASHG, Baltimore, MD, USA, Oct 2015.
29. MicroRNA as biomarkers of spontaneous preterm birth. *Abstract*, Society for Maternal-Fetal Medicine (SMFM). March 2015.
30. Neonatal, not Maternal, Copy Number Variants are Associated with Spontaneous Preterm Birth. *Abstract*, SMFM. March 2015.
31. Natural and orthogonal model for gene-gene interactions applied to cutaneous melanoma. *Poster presentation*, IGES, Chicago, IL, USA, Sep 2013.
32. Natural and orthogonal association framework to detect parent-of-origin effects. *Poster presentation*, IGES, Stevenson, WA, USA, Oct 2012.
33. Natural and orthogonal interaction framework for modeling GxG and GxE interactions. *Poster presentation*, IGES, Stevenson, WA, USA, Oct 2012.

## INVITED TALKS AND SEMINARS

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34. Genome wide association study of copy number variation in lung cancer. University of Florida Cancer AI workshop, November 2021.
35. Integrating genomic correlation structure improves copy number variants detection. University of Florida, June 2021.
36. Identification and characterization of genomic features from high throughput data. University of California Log Angeles, March 2021.
37. Copy number variation, epigenetics and cancer. School of Medicine, Wuhan University, China, July 2019.
38. Genetics and Environment Factors in Complex Diseases. School of Public Health, Guangxi Medical University, China, June 2019.
39. An accurate and powerful method for copy number variation detection. Department of Biology, Fudan University, China, June 2018.
40. Integrating genetic source boosts power for copy number variation detection. Department of Statistics, University of South Carolina, Columbia, SC, April 2018.
41. A powerful statistical framework for copy number variation detection with application to melanoma. Medical University of South Carolina, Charleston, SC, April 2018.
42. Copy number variations, parent-of-origin effects and cancer. Nanjing Medical University, Nanjing, Jiangsu, China, Jul 2017.
43. Imprinting effect detection and copy number variation in cancer. Wuhan University, Wuhan, Hubei, China, Jul 2017.
44. Copy number variations, epigenetics and cancer. Dartmouth College, Hanover, NH, USA, Feb 2017.

45. Testing of parent-of-origin effect in eQTL mapping using RNA-seq data. University of South Carolina, Columbia, SC, USA, Nov 2016.
46. A statistical framework for copy number variation detection with application to melanoma study. Wuhan University, Wuhan, China, June 2016.
47. A framework of testing interactions and parent-of-origin effect with application to cancer studies. Moffitt Cancer Center, Tampa, FL, USA, Dec 2015.
48. Testing of interactions and parent-of-origin effect using orthogonal models. University of South Carolina, Columbia, SC, USA, Nov 2015.
49. Modified Screening and Ranking Algorithm for Copy Number Variation Detection. City University of New York, New York, NY, USA, April 2015.
50. A new change-point model based method for copy number variation detection. University of Massachusetts Amherst, Amherst, MA, USA, March 2015.
51. Modified Screening and Ranking Algorithm for Copy Number Variation Detection. University of South Carolina, Columbia, SC, USA, February 2015.
52. Modified Screening and Ranking Algorithm for Copy Number Variation Detection. *Seminar*, University of Colorado Boulder, Boulder, CO, USA, January 2015.
53. Natural and Orthogonal Model for Gene-Gene interactions Applied to Cutaneous Melanoma. Geisel School of Medicine at Dartmouth College, Lebanon, NH, USA, July 2013.

## **INSTITUTIONAL SERVICES IN UFL**

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### *University*

- Member on the UF Artificial Intelligence Initiative Decision Support Committee (2022-)
- Member of Faculty Senate (2023-)

### *College*

- Poster judge for College of Medicine Research Day (2022, 2023)
- AI Seminar Committee Member (2023-)

### *Department*

- Member of Graduate Students Admission Committee (2022-)

## **INSTITUTIONAL SERVICES IN USC**

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### *University*

- University of South Carolina Faculty Senate (2019 - 2021)
- Member of AI Institute and Center of Excellence in Autism & Neurodevelopment Disorder joint position search committee (2020-2021)
- USC Office of the Vice President for Research ASPIRE II Grant Reviewer (2021)
- USC Office of the Vice President for Research ASPIRE I Grant Reviewer (2019)

- Member of the Book Store Committee for the University of South Carolina (2017 – 2020)

#### *School*

- Member of Department of Communication Sciences and Disorders Faculty Search Committee (2017 – 2018)

#### *Department*

- Division of Biostatistics Admissions Committee (2016 - 2021)
- Member of Department of Epidemiology and Biostatistics Students Handbook Committee (2020)
- Chair of the Biostatistics Forum for the Department of Epidemiology and Biostatistics (2017-2019)
- Chair of the Graduate Exam Committee for the Department of Epidemiology and Biostatistics (2017 - 2018)
- Member of the Graduate Exam Committee for the Department of Epidemiology and Biostatistics (2016 - 2017)
- Grader of the Graduate Students Qualifying and Comprehensive Exams (2016 – 2020)

### **EXTERNAL SERVICES and MEMBERSHIP**

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- Reviewer for the Student Paper Award for International Chinese Statistical Association (ICSA) Applied Statistics Symposium, 2022
- Early Career Reviewer Panel for NIH Cancer Genetic Study Section, June 2020
- Chair and Organizer, *Statistical learning advancement for inference with complex biomedical data*. ICSA Applied Statistics Symposium, 2020
- Chair and Organizer, *New developments in High Dimensional Data Analysis*. ICSA 2019 Applied Statistics Symposium, Raleigh, NC, 2019
- Platform session chair of International Conference on Computational Methods (ICCM), Guilin, Guangxi, China, 2017
- Platform session chair ICSA, Atlanta, GA, 2016
- Program committee: European Conference on Computational Biology (ECCB), 2016
- *Ad hoc* reviewer for *American Journal of Human Genetics*, *Biomedical Sciences*, *BioData Mining*, *Bioinformatics*, *BMC Cancer*, *BMC Medical Genomics*, *Briefings in Bioinformatics*, *Clinical Genetics*, *European Conference on Computational Biology*, *Environmental Science and Pollution Research*, *Frontier in Genetics*, *Genetic Epidemiology*, *Genomics*, *Human Genetics*, *Human Heredity*, *IEEE Signal Processing Letters*, *Journal of Nervous and Mental Disease*, *Journal of Theoretical Biology*, *Molecular Genetics and Genomics*, *Nature Genetics*, *Neoplasia*, *Nutrition & Diabetes*, *Nutrition & Metabolism*, *Nucleic Acid Research*, *PLoS ONE*, *PLoS Computational Biology*, *PLoS Genetics*, *Peer J*, *Scientific Report*, *Statistics & Its Interface*, *Environmental Science and Pollution Research*.
- Invited reviewer for *Annals of Public Health and Research* and *Human Genetics*
- Member of the International Genetic Epidemiology Society (IGES)



- Member of the American Society of Human Genetics (ASHG)
- Member of the American Association of Cancer Research (AACR)
- Member of the International Chinese Statistical Association (ICSA)

## **SOFTWARE**

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### **CORRseq and SARaseq**

- R packages for copy number variation detection with whole exome sequencing data.  
<https://github.com/FeifeiXiaoUSC/CORRseq-and-SARaseq>.

### **LDcnv**

- A R package for whole genome correlation struction integrated copy number variation detection with SNP array data. <https://github.com/FeifeiXiaoUSC/LDcnv>.

### **modSaRa2**

- An R package for accurate and powerful method for copy number variation detection with SNP array data by integrating relative allele frequencies.  
<https://github.com/FeifeiXiaoUSC/modSaR2>.

### **modSaRa**

- An R package for a change-point model based method for copy number variation detection with SNP array data. <https://github.com/FeifeiXiaoUSC/modSaRa>.

### **miRecords**

- An integrated resource for animal microRNA-target interactions providing both computational predicted and experimental validated miRNA-target interactions.  
<http://c1.accurascience.com/miRecords/>.

## **COMPUTER SKILLS**

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### Computational Programming:

- R/S-plus
- Unix Shell Script
- SAS
- Stata
- JavaScript
- C