Jane Liang

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janewliang.github.io

Education

Aug. 2017 –

Doctor of Philosophy in Biostatistics

May 2022 (expected)

Harvard T.H. Chan School of Public Health

spected) | Boston, MA

Dissertation: Innovative approaches for risk assessment in panel gene testing

Advisor: Giovanni Parmigiani

Committee members: Danielle Braun, Peter Kraft Cancer training grant fellow (2020 − *Present*)

Genomics training grant fellow (2017 − 2020)

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Aug. 2012 -

Bachelor of Arts in Statistics, with Honors

Dec. 2015 University of California, Berkeley

Berkeley, CA

Phi Beta Kappa, Distinction in General Scholarship

Honors thesis: Comparing dependence measures using simulation studies

Supervisor: Haiyan Huang

Experience

Aug. 2017 –

Graduate Student Researcher

Present | Department of Biostatistics, Harvard T.H. Chan School of Public Health

Department of Data Sciences, Dana-Farber Cancer Institute

Boston, MA

BayesMendel Lab , led by Giovanni Parmigiani & Danielle Braun

Also supervised by: Peter Kraft & Robert C. Green

Statistical methodology and software development for clinical risk assessment in

panel gene testing.

Oct. 2016 -

Scientific Research Programmer

July 2017

Division of Biostatistics, Department of Preventive Medicine

University of Tennessee Health Science Center

Memphis, TN

Supervisor: Śaunak Sen

Writing, testing, and documenting software applications based on research require-

ments; maintaining division's high performance computing systems.

Jan. – Sept. 2016

Programming Analysis Associate

Kaiser Permanente

Pleasanton, CA

Data management, dashboard reporting, and bash scripting to monitor server envi-

ronment activity.

Publications

- [1] **Jane W. Liang**, Gregory Idos, Christine Hong, Stephen Gruber, Giovanni Parmigiani, and Danielle Braun. PanelPRO: a general framework for multi-gene, multi-cancer Mendelian risk prediction models. 2021. In preparation.
- [2] Gavin Lee*, **Jane W. Liang***, Qing Zhang, Theodore Huang, Christine Choirat, Giovanni Parmigiani, and Danielle Braun. PanelPRO: A R package for multi-syndrome, multi-gene risk modeling for individuals with a family history of cancer. *arXiv preprint: arXiv:2010.13011 [stat.AP]*, 2020. Submitted to *eLife*.
- [3] **Jane W. Liang** and Śaunak Sen. Sparse matrix linear models for structured high-throughput data. *arXiv preprint: arXiv:1712.05767 [stat.CO]*, 2020. To appear in *Annals of Applied Statistics*.
- [4] Yunqi Yang, Christine Hong, **Jane W. Liang**, Stephen Gruber, Giovanni Parmigiani, Gregory Idos*, and Danielle Braun*. A likelihood-based approach to assessing frequency of pathogenicity among variants of unknown significance (VUS) in susceptibility genes. *Statistics in Medicine*, 40(3):593–606, 2020. doi: 10.1002/sim.8791.
- [5] **Jane W. Liang**, Robert J. Nichols, and Śaunak Sen. Matrix linear models for high-throughput chemical genetic screens. *Genetics*, 212(4):1063–1073, 2019. doi: 10.1534/genetics.119.302299.
- [6] Alexandra H. Bartlett, **Jane W. Liang**, Jose Vladimir Sandoval-Sierra, Jay H. Fowke, Eleanor M. Simonsick, Karen C. Johnson, and Khyobeni Mozhui. Longitudinal study of leukocyte DNA methylation and biomarkers for cancer risk in older adults. *Biomarker Research*, 7(1):10, 2019. doi: 10.1186/s40364-019-0161-3.
- [7] Hemant Gujar, **Jane W. Liang**, Nicholas C. Wong, and Khyobeni Mozhui. Profiling DNA methylation differences between inbred mouse strains on the Illumina Human Infinium MethylationEPIC microarray. *PloS ONE*, 13(3):e0193496, 2018. doi: 10.1371/journal.pone.0193496.

Presentations

Aug. 12, 2021	Aggregating across genes and cancers in Mendelian risk prediction modeling (speed presentation) Joint Statistical Meetings ♂ Seattle, WA
Jan. 21, 2021	Aggregating across genes and cancers in Mendelian risk prediction modeling (oral presentation) Dana-Farber/Harvard Cancer Center Celebration of Early Career Investigators in Cancer Research (DF/HCC CECI) Boston, MA
March 23, 2020	A general framework for multi-gene, multi-cancer Mendelian risk prediction models (contributed paper) ENAR International Biometric Society Spring Meeting Nashville, TN

^{*} indicates equal contributions

Nov. 11, 2019 | Sparse matrix linear models for structured high-throughput data (contributed poster and lightning talk)

Integrative Biostatistics Research for Imaging, Genomics, & High-throughput Technologies in Precision Medicine (iBRIGHT)

MD Anderson Cancer Center

Houston, TX

June 23, 2017 | Matrix linear models for high-throughput data (lightning talk) 🗹

JuliaCon 🗹

University of California, Berkeley

Berkeley, CA

Honors

Jan. 21, 2021 Best Oral Presentation

 Dana-Farber/Harvard Cancer Center Celebration of Early Career Investigators in Cancer Research (DF/HCC CECI)

 2020 – 2021 Certificate of Distinction in Teaching

 Department of Biostatistics
 Harvard T.H. Chan School of Public Health

Competitive Travel and Conference Support

Nov. 11-13, 2019 Integrative Biostatistics Research for Imaging, Genomics, & Highthroughput Technologies in Precision Medicine (iBRIGHT)
MD Anderson Cancer Center

June 20-24, 2017 | **JuliaCon** ✓ University of California, Berkeley

May 1-3, 2017 | **50th Annual Barrett Memorial Lectures: Mathematical Foundations of Data Science** ☑ University of Tennessee, Knoxville

Feb. 8-10, 2017 | Workshop on the Interface of Statistics and Optimization (WISO)

Statistical and Applied Mathematical Sciences Institute (SAMSI)

Oct. 16-22, 2016 | Short Course on Systems Genetics The Jackson Laboratory

Teaching

Teaching Fellow

Department of Biostatistics

Harvard T.H. Chan School of Public Health

Boston, MA

Leading lab sections, designing course materials, holding office hours, monitoring online discussion boards, grading assignments, and supporting master's, doctoral, and

clinical students from diverse fields.

Spring 2021 | BST 231: Statistical Inference I (remote)

Fall 2020 BST 260: Introduction to Data Science (remote)
Spring 2020 BST 263: Statistical Learning (in-person and remote)
Fall 2019 BST 260: Introduction to Data Science (in-person)
Spring 2019 BST 210: Applied Regression Analysis (in-person)

Teaching Assistant

McGoldrick Professional Development Program in Public Health 🗹

Harvard T.H. Chan School of Public Health, University of KwaZulu-Natal –

Pietermaritzburg, and the ARISE network

Boston, MA

Training researchers from low-and middle-income countries in quantitative methods and building a collaborative network in sub-Saharan Africa.

Feb. 1-12, 2021

Health Data Science (remote)

Curriculum Fellow

Department of Biostatistics

Harvard T.H. Chan School of Public Health

Boston, MA

Designing course materials, including exams, homework, and lab assignments.

Fall 2020 | BST 260: Introduction to Data Science 🛂

Service

Referee: Cancer Epidemiology, Biomarkers & Prevention; PLOS One

Software

R: PanelPRO

Julia: MatrixLMnet, MatrixLM, GeneticScreens

Skills

High proficiency: R, Julia, Python, C++, LaTeX, distributed version control (Git, Mercurial),

high performance cluster computing, Linux environments

Some proficiency: Java, MATLAB, web scraping