

Jane Liang

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Education

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| Aug. 2017 –
May 2022
(expected) | Doctor of Philosophy in Biostatistics
Harvard T.H. Chan School of Public Health
<i>Boston, MA</i>
Dissertation : <i>Innovative approaches for risk assessment in panel gene testing</i>
Advisor: Giovanni Parmigiani
Committee members: Danielle Braun, Peter Kraft
Cancer training grant fellow (2020 – Present) ↗
Genomics training grant fellow (2017 – 2020) ↗ |
| Aug. 2012 –
Dec. 2015 | Bachelor of Arts in Statistics, with Honors
University of California, Berkeley
<i>Berkeley, CA</i>
Phi Beta Kappa, Distinction in General Scholarship
Honors thesis: <i>Comparing dependence measures using simulation studies</i>
Supervisor: Haiyan Huang |

Experience





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| Aug. 2017 –
Present | Graduate Student Researcher
Department of Biostatistics, Harvard T.H. Chan School of Public Health
Department of Data Sciences, Dana-Farber Cancer Institute
<i>Boston, MA</i>
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 –
July 2017 | Scientific Research Programmer
Division of Biostatistics, Department of Preventive Medicine
University of Tennessee Health Science Center
<i>Memphis, TN</i>
Supervisor: Śaunak Sen
Writing, testing, and documenting software applications based on research requirements; maintaining division's high performance computing systems. |
| Jan. – Sept. 2016 | Programming Analysis Associate
Kaiser Permanente
<i>Pleasanton, CA</i>
Data management, dashboard reporting, and bash scripting to monitor server environment 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 \[stat.AP\]](https://arxiv.org/abs/2010.13011)*, 2020. Submitted to *eLife*.
- [3] **Jane W. Liang** and Saunak Sen. Sparse matrix linear models for structured high-throughput data. *arXiv preprint: [arXiv:1712.05767 \[stat.CO\]](https://arxiv.org/abs/1712.05767)*, 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](https://doi.org/10.1002/sim.8791).
- [5] **Jane W. Liang**, Robert J. Nichols, and Saunak Sen. Matrix linear models for high-throughput chemical genetic screens. *Genetics*, 212(4):1063–1073, 2019. doi: [10.1534/genetics.119.302299](https://doi.org/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](https://doi.org/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](https://doi.org/10.1371/journal.pone.0193496).

* indicates equal contributions

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

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 <i>Houston, TX</i>
June 23, 2017	Matrix linear models for high-throughput data (lightning talk) ↗ JuliaCon ↗ University of California, Berkeley <i>Berkeley, CA</i>

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 2019 – 2020	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, & High-throughput 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 <i>Boston, MA</i> Leading lab sections, designing course materials, holding office hours, monitoring on-line 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 <i>Boston, MA</i> 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 <i>Boston, MA</i> 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