Kevin Z. Lin

University of Washington, Department of Biostatistics

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CV last updated: January 1, 2025

EDUCATION

• Princeton University

Princeton, NJ 2010 - 2014

B.S.E. in Operations Research & Financial Engineering

Advisors: Han Liu and Robert Vanderbei

Certificates in "Statistics and Machine Learning" and "Applications of Computing"

• Carnegie Mellon University

Pittsburgh, PA

Ph.D. in Dietrich's Statistics & Data Science, Masters in Machine Learning

2014 - 2020

Thesis advisors: Kathryn Roeder and Jing Lei

EMPLOYMENT

• University of Pennsylvania

Philadelphia, PA

Post-doctoral researcher in Wharton's Statistics & Data Science

2020 - 2023

Advisor: Nancy Zhang

• University of Washington

Seattle, WA

Assistant professor (tenure-track) in School of Public Health's Biostatistics

2023 - Present

Program faculty, UW Computational Molecular Biology

2023 - Present

Honors, Awards, and Scholarships

• Teaching assistant excellence award recipient For Course 36-350: "Statistical Computing" in Fall 2017 Carnegie Mellon University

May 2017

• Winner of Statistical Excellence for Early-Career Writing

Significance Magazine

For article "We, the millennials: The statistical significance of political significance"

June 2017

Competition held jointly with the Young Statisticians Section of Royal Statistical Society

 \bullet Honorable mention in student paper competition

American Statistical Association

For paper "Dependency diagnostic: Visually understanding pairwise variable January 2018 relationships"

For ASA section: Statistical Computing and Statistical Graphics

• PhD TAs of the year

Carnegie Mellon University

For the Spring 2020 semester (1 of 2 total recipients)

May 2020

• Wikimedia Foundation Research Award of the Year

(Remote)

For paper "Controlled analyses of social biases in Wikipedia bios"

2023

• Biometrics Excellent Referee Awards

Atlanta, GA

For the excellent reviews for the journal Biometrics (1 of 3 total recipients)

2024

- Association member for: American Society of Cell Biology (ASCB); American Statistical
 Association (ASA); American Society of Human Genetics (ASHG); Eastern North American
 Region (ENAR); International Chinese Statistical Association (ICSA); Institute of
 Mathematical Statistics (IMS)
- Peer-Reviewer for: Annals of Applied Statistics; Annals of Statistics; Bayesian Analysis;
 Biometrics; Biometrika; Cell Genomics; Electronic Journal of Statistics; IEEE Transactions of
 Network Science and Engineering; Information and Inference: A Journal of the IMA; Journal
 of American Statistical Association; Journal of Molecular Biology; Journal of the Royal
 Statistical Society: Series B; Nature Communications; Nature Methods; Nature Neuroscience;
 Patterns; PLOS Genetics; Statistical Sinica; Statistics and Probability Letters; Statistics in
 Medicine; Technometrics
- Grant Reviewer for:
 - National Science Foundation (Methodology, Measurement, and Statistics): 2024
- Reviewer for Student Paper Award for the American Statistical Association's Section on Statistics in Genomics and Genetics (December in 2023, 2024)
- Area Chair for the conference Machine Learning in Computational Biology (Autumn 2023, 2024).

BIBLIOGRAPHY

Bold denotes PI and his mentored students. Items are ordered in chronological order.

Refereed research articles

- Vanderbei, R., Lin, K. Z., Liu, H., and Wang, L. (2016). Revisiting compressed sensing: Exploiting the efficiency of simplex and sparsification methods. *Mathematical Programming Computation*, 8(3):253–269
 DOI: 10.1007/s12532-016-0105-y
- Lin, K. Z., Sharpnack, J., Rinaldo, A., and Tibshirani, R. J. (2017). A sharp error analysis
 for the fused lasso, with application to approximate changepoint screening. In *Advances in Neural Information Processing Systems*, pages 6884–6893
 DOI: 10.5555/3295222.3295432
- An, J.-Y., Lin, K. Z., Zhu, L., Werling, D. M., Dong, S., Brand, H., Wang, H. Z., Zhao, X., Schwartz, G. B., Collins, R. L., Currall, B. B., Dastmalchi, C., Dea, J., Duhn, C., Gilson, M. C., Klei, L., Liang, L., Markenscoff-Papadimitriou, E., Pochareddy, S., Ahituv, N., Buxbaum, J. D., Coon, H., Daly, M. J., Shin Kim, Y., Marth, G. T., Neale, B. M., Quinlan, A. R., Rubenstein, J. L., Sestan, N., State, M. W., Willsey, A. J., Talkowski, M. E., Devlin, B., Roeder, K., and Sanders, S. J. (2018). Genome-wide de novo risk score implicates promoter variation in autism spectrum disorder. Science, 362(6420)
 DOI: 10.1126/science.aat6576
- 4. Lei, J. and **Lin, K. Z.** (2020). Discussion of 'Network cross-validation by edge sampling'. *Biometrika*, 107(2):285–287
 DOI: 10.1093/biomet/asaa009

- 5. Wang, D., Zhao, Z., **Lin, K. Z.**, and Willett, R. (2021). Statistically and computationally efficient changepoint localization in regression settings. *Journal of Machine Learning Research*, 22:248–1
 - DOI: 10.5555/3546258.3546506
- Hyun, S., Lin, K. Z., G'Sell, M., and Tibshirani, R. J. (2021). Post-selection inference for changepoint detection algorithms with application to copy number variation data. *Biometrics*, 77(3):1037–1049
 - **DOI**: 10.1111/biom.13422
- Lin, K. Z., Liu, H., and Roeder, K. (2021b). Covariance-based sample selection for heterogeneous data: Applications to gene expression and autism risk gene detection. *Journal of the American Statistical Association*, 116(533):54–67
 DOI: 10.1080/01621459.2020.1738234
- 8. **Lin, K. Z.**, Lei, J., and Roeder, K. (2021a). Exponential-family embedding with application to cell developmental trajectories for single-cell RNA-seq data. *Journal of the American Statistical Association*, 116(534):457–470

 DOI: 10.1080/01621459.2021.1886106
- 9. Field, A., Park, C. Y., **Lin, K. Z.**, and Tsvetkov, Y. (2022). Controlled analyses of social biases in Wikipedia bios. In *Proceedings of the ACM Web Conference 2022*, pages 2624–2635 DOI: 10.1145/3485447.3512134
- Lei, J. and Lin, K. Z. (2023). Bias-adjusted spectral clustering in multi-layer stochastic block models. *Journal of the American Statistical Association*, 118(544):2433–2445
 DOI: 10.1080/01621459.2022.2054817
- Guan, P. Y., Lee, J. S., Wang, L., Lin, K. Z., Mei, W., Chen, L., and Jiang, Y. (2023).
 Destin2: Integrative and cross-modality analysis of single-cell chromatin accessibility data.
 Frontiers in Genetics, 14
 DOI: 10.3389/fgene.2023.1089936
- Lin, K. Z. and Zhang, N. R. (2023). Quantifying common and distinct information in single-cell multimodal data with Tilted Canonical Correlation Analysis. *Proceedings of the* National Academy of Sciences, 120(32)
 DOI: 10.1073/pnas.2303647120
- Chen, S., Zhu, B., Huang, S., Hickey, J. W., Lin, K. Z., Snyder, M., Greenleaf, W. J., Nolan, G. P., Zhang, N. R., and Ma, Z. (2024). Integration of spatial and single-cell data across modalities with weakly linked features. *Nature Biotechnology*, 42(7):1096–1106
 DOI: 10.1038/s41587-023-01935-0
- 14. Lin, K. Z., Qiu, Y., and Roeder, K. (2024). eSVD-DE: Cohort-wide differential expression in single-cell rna-seq data using exponential-family embeddings. *BMC Bioinformatics*, 25(1):113 DOI: 10.1186/s12859-024-05724-7
- Prater, K. E. and Lin, K. Z. (2024). All the single cells: Single-cell transcriptomics/epigenomics experimental design and analysis considerations for glial biologists. Glia
 DOI: 10.1002/glia.24633

Other refereed scholarly publications

1. Lin, K. Z. (2017). We, the millennials: The statistical significance of political significance. Significance, 14(5):28–33

DOI: 10.1111/j.1740-9713.2017.01073.x

Other non-refereed scholarly publications

- 1. Lin, K. Z. and Lei, J. (2024). Dynamic clustering for heterophilic stochastic block models with time-varying node memberships. arXiv preprint arXiv:2403.05654

 Arxiv: 2403.05654 (In revision)
- Yang, S. J., Wang, Y., and Lin, K. Z. (2024). LCL: Contrastive learning for lineage barcoded scRNA-seq data. bioRxiv DOI: 10.1101/2024.10.28.620670
- 3. Wu, W., Kennedy, T., Arguello-Miranda, O., and Lin, K. Z. (2024). Measuring regulatory network inheritance in dividing yeast cells using ordinary differential equations. bioRxiv DOI: 10.1101/2024.11.23.624995
- 4. Li, Y., Wei, Z. J., Chen, Y.-C., and Lin, K. Z. (2024). Assessing RNA velocity stability across synthetic replicates using count splitting. *bioRxiv* DOI: 10.1101/2024.11.23.625009

PATENTS AND OTHER INTELLECTUAL PROPERTY

Below are publicly available software made by me and my lab.

- 1. **covarianceSelection**: R package for selecting datasets with similar high-dimensional covariance matrices. (https://github.com/linnykos/covarianceSelection)
- 2. **selectiveModel**: R package for selective inference via binary segmentation for changepoint detection. (https://github.com/linnykos/selectiveModel, developed jointly with Sangwon Hyun)
- 3. **eSVD**: R package for performing dimension reduction for matrices with respect to an exponential-family distribution. (https://github.com/linnykos/esvd)
- 4. **networkSoSD**: R package for performing spectral clustering to estimate node communities based on a collection of networks. (https://github.com/linnykos/networkSoSD)
- 5. **tiltedCCA**: R package for estimating the shared and distinct geometry in paired multimodal data. (https://github.com/linnykos/tiltedCCA)
- 6. **eSVD2**: R package that extends the functionality of the previous eSVD package, and also implements specialized functions to perform subject-level differential expression for single-cell data. (https://github.com/linnykos/eSVD2)
- 7. **Lineage-aware_CL**: Python package to isolate the lineage-specific gene expression signatures from lineage-tracing scRNA-seq data (https://github.com/SZ-yang/Lineage-aware_CL)
- 8. **ODEinherit**: R package to quantify the amount of inheritance of protein regulatory networks during cell division from live imaging microscopy data (https://github.com/WenbinWu2001/ODEinherit)
- 9. **veloUncertainty**: Python package to perform inference on RNA velocity methods based on count-splitting scRNA-seq data (https://github.com/linnykos/veloUncertainty)

"id." denotes identical title as the item above. Items are grouped by talks, and then ordered by chronological order.

Invited departmental seminars

- 1. University of California, Los Angeles: Department of Statistics (12/2021). Exponential-family embedding with application to cell developmental trajectories for single-cell data
- 2. University of California, Irvine: Department of Statistics (12/2022). Tilted-CCA: Quantifying common and distinct information in multiomic single-cell data
- 3. University of Washington: Department of Biostatistics (01/2023). id.
- 4. University of Chicago: Department of Statistics (01/2023). id.
- 5. University of California, Riverside: Department of Statistics (01/2023). id.
- 6. University of Waterloo: Department of Statistics (01/2023). id.
- 7. Pennsylvania State University: Department of Statistics (01/2023). id.
- 8. University of North Carolina: Department of Biostatistics (01/2023). id.
- 9. University of Michigan: Department of Statistics (01/2023). id.
- 10. University of Michigan: Department of Biostatistics (02/2023). id.
- 11. University of Illinois, Urbana-Champaign: Department of Statistics (02/2023). id.
- 12. Emory University: Department of Biostatistics (02/2023). id.
- 13. McGill University: Department of Epidemiology, Biostatistics, and Occupational Health (09/2023). id.
- 14. University of California, Santa Cruz: Department of Statistics (10/2023). id.
- 15. University of Washington: Department of Statistics (05/2024). id.
- 16. Michigan State University: Department of Statistics (02/2024). Single-cell paired RNA & ATAC: Surveying broad multi-modal coordination in development and cancer resistance
- 17. University of Pennsylvania: Biostatistics Division of Department of Biostatistics, Epidemiology, and Informatics (03/2024). id.
- 18. Duke University: Computational Biology and Bioinformatics (04/2024). id.
- 19. Brown University: Center for Computational Molecular Biology (12/2024). eSVD-DE: Cohort-level differential expression in single-cell data via matrix factorization

Conference/Seminar presentations

Note: (*) denotes invited talks.

- 1. Modeling and Optimization: Theory and Applications (08/2014, Bethlehem, PA). Optimization for compressed sensing: New insights and alternatives
- 2. American Society of Human Genetics (10/2015, Baltimore, MD). Gaussian graphical model integrating microarray and sequencing data for autism risk gene detection
- 3. Joint Statistical Meeting (08/2016, Chicago, IL). id.
- 4. Conference on Neural Information Processing Systems (12/2017, Long Beach, CA). A sharp error analysis for the fused lasso, with application to approximate changepoint screening
- 5. Joint Statistical Meeting (08/2017, Baltimore, MD). Hypothesis testing for simultaneous variable clustering and correlation network estimation
- 6. Joint Statistical Meeting (08/2018, Vancouver, Canada). Dependency diagnostic: Visually understanding pairwise variable relationships
- 7. Joint Statistical Meeting (08/2019, Denver, CO). Exponential-family embedding with application to cell developmental trajectories for single-cell data
- 8. Joint Statistical Meeting* (08/2020, Remote). id.
- 9. American Society of Human Genetics (10/2020, Remote). id.
- 10. Joint Statistical Meeting (08/2020, Remote). Time-varying stochastic block models via kernel smoothing, with application to RNA-seq data
- 11. StatScale Seminar* (07/2021, Remote). id.
- 12. Symposium on Data Science and Statistics (06/2022, Pittsburgh, PA). Spectral clustering for multi-layer stochastic block models: Analysis of dynamic heterophilic networks
- 13. International Chinese Statistical Association Applied Statistical Symposium* (06/2023, Ann Arbor, MI). id.
- 14. Joint Statistical Meetings (08/2022, Washington DC). Tilted-CCA: Quantifying common and distinct information in multiomic single-cell data
- 15. Institute of Mathematical Statistics New Researchers Conference 2023 (08/2023, Toronto, Canada). id.
- 16. Scripps Research: Computational Biology and Bioinformatics Seminar* (08/2023, Remote). id.
- 17. Cell Symposia: The conceptual power of single-cell biology (08/2023, San Diego, CA). id.
- 18. American Society of Human Genetics (11/2023, Washington DC). id.
- 19. Keystone Symposia: Regulatory RNA Emerging Mechanisms (12/2023, Banff, Canada). id.
- 20. Joint Statistical Meeting* (08/2024, Portland, OR). id.

- 21. Joint Statistical Meeting (08/2023, Toronto, Canada). eSVD: Cohort-level differential expression in single-cell data via matrix factorization
- 22. UW Genome Science: Combi Seminar* (01/2024, Seattle, WA). Single-cell paired RNA & ATAC: Surveying broad multi-modal coordination in development and cancer resistance
- 23. Fred Hutch Cancer Center: Biostatistics Seminar Series* (01/2024, Seattle, WA). id.
- 24. International Chinese Statistical Association Applied Statistical Symposium* (06/2024, Nashville, KT). eSVD-DE: Cohort-level differential expression in single-cell data via matrix factorization
- 25. American Society of Cell Biology (12/2024, San Diego, CA). Measuring inheritance of regulatory networks in dividing yeast cells using statistical ODE networks

Organized conference sessions

- 1. STATGEN 2024: Conference on Statistics in Genomics and Genetics (05/2024, Pittsburgh, PA). Session title: New mathematical paradigms behind single-cell biology
- 2. Joint Statistical Meeting (08/2024, Portland, OR). Session title: Single-cell Biology: New Frontiers to Understand Gene Expression Through Other Omics

Webinars

1. Institute of Mathematical Statistics – New Researchers Group: Navigating the academic job market (11/2024)

University Service

- UW Department of Biostatistics: Seminar Committee; Member (2024 Present)
- UW Department of Biostatistics: Equity, Diversity, and Inclusion Committee; Member (2023 Present)
 - Lead coordinator for the Wellness working group (Winter 2023, Spring 2024)
- UW School of Public Health: Equity, Diversity, Inclusion Committee; Member, representing the Department of Biostatistics (2023)
- UW Department of Biostatistics: Teaching track assistant professor search; Member (2023)
- UW School of Public Health: Resume Clinic; Reviewer (2023: Autumn, 2024: Spring, Autumn)
- UW School of Public Health: Mock Interview; Interviewer (Winter 2023)

TEACHING HISTORY

At Princeton University

1. Course designer for ORF 350: Analysis of Big Data (Spring 2012, Spring 2013, Spring 2014)

At Carnegie Melon University

- 1. Teaching assistant for 46-921 & 46-923: Financial Data Analysis I and II (Spring 2014)
- 2. Teaching assistant for 36-217: Probability Theory and Random Processes (Spring 2015)
- 3. Teaching assistant for 36-350: Statistical Computing (Fall 2015, Fall 2016, Fall 2017)
- 4. Guest lecturer for 36-750: Statistical Computing (Fall 2016, Fall 2017, Fall 2018, Fall 2019, Fall 2020)
- 5. Assistant instructor for 36-350: Statistical Computing (Spring 2018)
- 6. Instructor for 36-350: Statistical Computing (Summer 2018)
- 7. Data science initiative project fellow for 36-490: Undergraduate Research (Spring 2019)
- 8. Co-instructor for 36-469: Statistical Genomics and High-Dimensional Inference (Spring 2020). With Kathryn Roeder

At University of Washington

- 1. Instructor for BIOST 582: Student Seminar (Autumn 2023)
- 2. Instructor for BIOST 561: Computational Skills for Biostatistics (Spring 2024)

Independent study supervision

- Yuhong Li (2023-24), University of Washington, Department of Biostatistics, Master of Science
 Thesis
- 2. Zhaoheng Li (2023-24), University of Washington, Department of Biostatistics, PhD candidate
- 3. Yimin Zhao (2023-24), University of Washington, Department of Biostatistics, PhD candidate
- 4. Wenjing (Tati) Zhang (2023-24), University of Washington, Department of Biostatistics, Master of Science Thesis
- 5. Amy Watt (2024), University of Washington, Department of Biostatistics, PhD candidate

Teaching and other professional development

Short courses and tutorials taught at conferences:

1. ENAR 2024 (03/2024, Baltimore, MA) Teaching RNA-velocity for single-cell trajectory analysis and discussing its future research directions (Tutorial)

Other professional development:

- Certified by CMU's Eberly Center's Future Faculty Program, which included two observed lectures in two different semesters (Fall 2019 to Summer 2020)
- Certified with Gatekeeper certificate by the QPR's (Question, Persuade, Refer) suicide prevention program (February 2020)
- Certified by Mental Health First Aid USA (Fall 2020)
- Teaching assistant for the Section on Statistical Genomics and Genetics (SSGG)'s online short course (Title: An Introduction to Deep Learning in Omics). Taught by Dr. Wei Sun and Dr. Nancy Zhang (Winter 2022)

MS and PhD committees in non-chair role

- 1. Doctoral Committee Member
 - (a) Connor Finkbeiner (2023-24), University of Washington, Department of Genome Sciences, PhD candidate. (Chair: Manu Setty)
- 2. Graduate School Representative
 - (a) Clifford Rostomily (2024), University of Washington, Department of Genome Sciences, PhD candidate. (Chair: Cole Trapnell)
- 3. Qualifying Exam Member
 - (a) Anna Spiro (2023), University of Washington, Department of Computer Science, PhD candidate. (Chair: Sara Mostafavi)

Other mentoring

Project Supervision (outside of independent study supervision)

- 1. Zeyu (Jerry) Wei (2023), University of Washington, Department of Statistics, PhD (on project jointly with Yen-Chi Chen)
- 2. Wenbin Wu (2023-24), University of Washington, Department of Statistics, Master of Science Advanced Methods and Data Analysis
- 3. Haoye Yang (2023), University of Washington, Department of Statistics, Master of Science Advanced Methods and Data Analysis
- 4. Shizhao (Joshua) Yang (2023-24), University of Washington, Department of Biostatistics, Master of Science Capstone
- 5. Turbo Du (2024), University of Washington, Department of Biostatistics, Master of Science Capstone
- 6. Jingyi Guan (2024), University of Washington, Department of Biostatistics, PhD candidate
- 7. Yifan Lin (2024), University of Washington, Department of Biostatistics, Master of Science Thesis

Department-assigned Academic Advisor

- 1. Alejandro Hernandez (2023), University of Washington, Department of Biostatistics, Master of Science Capstone
- 2. Yutong Jin (2023), University of Washington, Department of Biostatistics, Master of Science Capstone
- 3. Mohamad D. Bairakdar (2024), University of Washington, Department of Biostatistics, PhD candidate