Lorin Crawford

1 Memorial Dr, Cambridge, MA 02142, USA lcrawford@microsoft.com • +1 (857) 453-6156 • www.lorincrawford.com

EDUCATION Duke University, Durham, North Carolina, USA

Aug 2013 – May 2017

Ph.D. in Statistical Science

Co-Advisors: Sayan Mukherjee, Ph.D. and Kris C. Wood, Ph.D.

Thesis: Bayesian Kernel Models for Statistical Genetics and Cancer Genomics

Clark Atlanta University, Atlanta, Georgia, USA

Aug 2009 – May 2013

B.S. in Mathematics

Advisor: Fisseha Abebe, Ph.D.

Valedictorian/Summa Cum Laude (Cumulative GPA: 4.0/4.0)

PROFESSIONAL EXPERIENCE

Microsoft Research New England, Cambridge, Massachusetts, USA

Principal Researcher Jun 2022 – Present Senior Researcher Sep 2020 – Jun 2022

Brown University, School of Public Health, Providence, Rhode Island, USA

Associate Professor of Biostatistics

RGSS Assistant Professor of Biostatistics

Jul 2022 – Present

Jul 2019 – Jul 2019

Assistant Professor of Biostatistics

Jul 2017 – Jul 2019

PUBLICATIONS

REFEREED PAPERS (*CO-FIRST AUTHORS; †CO-SENIOR AUTHORS; #CORRESPONDING AUTHOR(S); ADVISEE)

- [1] **L. Crawford**, V. Ponomarenko#, J. Steinberg, and M. Williams (2014). Accepted elasticity in local arithmetic congruence monoids. *Results in Mathematics*. **66**: 227-245.
- [2] G.R. Anderson, S.E. Wardell, M. Cakir, L. Crawford, J.C. Leeds, D.P. Nussbaum, P.S. Shankar, R.S. Soderquist, E.M. Stein, J.P. Tingley, P.S. Winter, E.K. Zeiser-Misenheimer, H.M. Alley, A. Yllanes, V. Haney, K.L. Blackwell, S.J. McCall, D.P. McDonnell, and K.C. Wood# (2016). PIK3CA mutations enable selective targeting of a breast tumor lineage survival dependency through MTOR-mediated control of MCL-1 translation. *Science Translational Medicine*. 8: 369ra175.
- [3] G.R. Anderson*, P.S. Winter*, K.H. Lin, D.P. Nussbaum, M. Cakir, E.M. Stein, R. Soderquist, L. Crawford, J.C. Leeds, R. Newcomb, P. Stepp, C. Yip, S.E. Wardell, J.P. Tingley, M. Ali, M. Xu, M. Ryan, S.J. McCall, A. McRee, C.M. Counter, C.J. Der, and K.C Wood# (2017). A landscape of therapeutic cooperativity in KRAS mutant cancers reveals principles for controlling tumor evolution. *Cell Reports*. 20(4): 999-1015.
- [4] **L. Crawford**#, P. Zeng, S. Mukherjee, and X. Zhou# (2017). Detecting epistasis with the marginal epistasis test in genetic mapping studies of quantitative traits. *PLOS Genetics*. **13**(7): e1006869.
- [5] K.R. Singleton*, L. Crawford*, E. Tsui, H.E. Manchester, O. Maertens, X. Liu, M.V. Liberti, A.N. Magpusao, E.M. Stein, J.P. Tingley, D.T. Frederick, G.M. Boland, K.T. Flaherty, S.J. McCall, C. Krepler, K. Sproesser, M. Herlyn, D.J. Adams, J.W. Locasale, K. Cichowski, S. Mukherjee, and K.C. Wood# (2017). Melanoma therapeutic strategies that select against resistance by exploiting MYC-driven evolutionary convergence. *Cell Reports.* 21(10): 2796-2812.
- [6] R. Soderquist, **L. Crawford**, E. Liu, M. Lu, A. Agarwal, G.R. Anderson, K.H. Lin, P.S. Winter, M. Cakir, and K.C. Wood# (2018). Systematic mapping of BCL-2 gene dependencies in cancer reveals molecular determinants of BH3 mimetic sensitivity. *Nature Communications*. **9**(1): 3513.
- [7] **L. Crawford**#, K.C. Wood, X. Zhou#, and S. Mukherjee# (2018). Bayesian approximate kernel regression with variable selection. *Journal of the American Statistical Association*. **113**(524): 1710-1721.
- [8] D.E. Runcie# and **L. Crawford** (2019). Fast and general-purpose linear mixed models for genome-wide genetics. *PLOS Genetics*. **15**(2): e1007978.

- [9] A. Monod#, S. Kališnik, J.Á. Patiño-Galindo, and **L. Crawford** (2019). Tropical sufficient statistics for persistent homology with a parametric application to infectious viral disease. *SIAM Journal on Applied Algebra and Geometry*. **3**(2): 337-371.
- [10] **L. Crawford**#, S.R. Flaxman, D.E. Runcie, and M. West (2019). Variable prioritization in nonlinear black box methods: a genetic association case study. *Annals of Applied Statistics*. **13**(2): 958-989.
- [11] T. Borgovan#, **L. Crawford**, <u>C. Nwizu</u>, and P. Quesenberry (2019). Stem cells and extracellular vesicles: biological regulators of physiology and disease. *American Journal of Physiology-Cell Physiology*. **317**(2): C155-C166.
- [12] K.H. Lin, J.C. Rutter, A. Xie, <u>E.T. Winn</u>, B. Pardieu, R. Dal Bello, Y.R. Ahn, Z. Dai, R.T. Sobhan, G.R. Anderson, K.R. Singleton, A.E. Decker, P.S. Winter, J.W. Locasale, **L. Crawford**, A. Puissant#, and K.C. Wood# (2020). Using antagonistic pleiotropy to design a chemotherapy-induced evolutionary trap. *Nature Genetics*. **52**: 408-417.
- [13] W. Cheng, S. Ramachandran#, and **L. Crawford**# (2020). Estimation of non-null SNP effect size distributions enables the detection of enriched genes underlying complex traits. *PLOS Genetics*. **16**(6): e1008855.
- [14] J.S. Sadick, **L. Crawford**, H.C. Cramer, C. Franck, S.A. Liddelow, and E.M. Darling# (2020). Generating cell type-specific protein signatures from non-symptomatic and diseased tissues. *Annals of Biomedical Engineering*. **48**: 2218-2232.
- [15] **L. Crawford**#, A. Monod#, A.X. Chen, S. Mukherjee, and R. Rabadán (2020). Predicting clinical outcomes in glioblastoma: an application of topological and functional data analysis. *Journal of the American Statistical Association*. **115**(531): 1139-1150.
- [16] B.A. Borden, Y. Baca, J. Xiu, F. Tavora, I. Winer, B.A. Weinberg, A.M. VanderWalde, S. Darabi, W.M. Korn, A.P. Mazar, F.J. Giles, L. Crawford, H. Safran, W.S. El-Deiry, and B.A. Carneiro# (2021). The landscape of glycogen synthase kinase-3 beta (GSK-3b) genomic alterations in cancer. *Molecular Cancer Therapeutics*. 20(1): 183-190.
- [17] A.N. Spierer#, J.A. Mossman, S.P. Smith, L. Crawford, S. Ramachandran, and D.M. Rand# (2021). Natural variation in the regulation of neurodevelopmental genes modifies flight performance in *Drosophila. PLOS Genetics.* 17(3): e1008887.
- [18] B. Wang*, T. Sudijono*, H. Kirveslahti*, T. Gao, D.M. Boyer, S. Mukherjee†, and **L. Crawford**†# (2021). A statistical pipeline for identifying physical features that differentiate classes of 3D shapes. *Annals of Applied Statistics*. **15**(2): 638-661.
- [19] D.E. Runcie#, J. Qu, H. Cheng, and **L. Crawford** (2021). Mega-scale linear mixed models for genomic predictions with thousands of traits. *Genome Biology*. **22**: 213.
- [20] P. Demetci*, W. Cheng*, G. Darnell, X. Zhou, S. Ramachandran, and L. Crawford# (2021). Multi-scale inference of genetic architecture using biologically annotated neural networks. *PLOS Genetics*. 17(8): e1009754.
- [21] M. Kamariza#, **L. Crawford**#, D. Jones#, and H.K. Finucane# (2021). Misuse of the term "trans-ethnic" in genomics research. *Nature Genetics*. **50**: 1520-1521.
- [22] S. Raghavan*, P.S. Winter*#, A.W. Navia*, H.L. Williams*, <u>A. DenAdel</u>, R.L. Kalekar, J. Galvez-Reyes, K.E. Lowder, J. Galvez-Reyes, R.L. Kalekar, N. Mulugeta, K.S. Kapner, M.S. Raghavan, A.A. Borah, N. Liu, S.A. Väyrynen, A. Dias Costa, R.W.S. Ng, J. Wang, E.K. Hill, D.Y. Ragon, L.K. Brais, A.M. Jaeger, L.F. Spurr, Y.Y. Li, A.D. Cherniack, M.A. Booker, E.F. Cohen, M.Y. Tolstorukov, I. Wakiro, A. Rotem, B.E. Johnson, J.M. McFarland, E.T. Sicinska, T.E. Jacks, R.J. Sullivan, T.E. Clancy, K. Perez, D.A. Rubinson, K. Ng, J.M. Cleary, **L. Crawford**, S.R. Manalis, J.A. Nowak, B.R. Wolpin†, W.C. Hahn†, A.J. Aguirre†#, and A.K. Shalek†# (2021). Microenvironment drives cell state, plasticity, and drug response in pancreatic cancer. *Cell.* **184**(25): 6119-6137.
- [23] S.P. Smith, S. Shahamatdar, W. Cheng, S. Zhang, J. Paik, M. Graff, C. Haiman, T.C. Matise, K.E. North, U. Peters, E. Kenny, C. Gignoux, G. Wojcik, L. Crawford†, and S. Ramachandran†# (2022). Enrichment analyses identify shared associations for 25 quantitative traits in over 600,000 individuals from seven diverse ancestries. *American Journal of Human Genetics*. 109: 871-884.

- [24] W.S. Tang*, G.M. da Silva*, H. Kirveslahti, E. Skeens, B. Feng, T. Sudijono, K.K. Yang, S. Mukherjee, B. Rubenstein†, and L. Crawford†# (2022). A topological data analytic approach for discovering biophysical signatures in protein dynamics. *PLOS Computational Biology.* **18**(5): e1010045.
- [25] W. Cheng#, S. Ramachandran, and **L. Crawford**# (2022). Uncertainty quantification in variable selection for genetic fine-mapping using Bayesian neural networks. *iScience*. **25**(7): 104553. (Spotlight Talk at the 10th RECOMB Satellite on Computational Methods in Genetics).
- [26] B.L. Trippe#, B. Huang, E.A. DeBenedictis, B. Coventry, N. Bhattacharya, K.K. Yang, D. Baker, and L. Crawford# (2022). Randomized gates eliminate bias in sort-seq assays. *Protein Science*. **31**(9): e4401.
- [27] <u>A. Conard*</u>, <u>A. DenAdel</u>*, and **L. Crawford**# (2023). A spectrum of explainable and interpretable machine learning approaches for genomic studies. *WIREs Computational Statistics*. **15**(5): e1617.
- [28] C. Rios-Martinez, N. Bhattacharya, A.P. Amini, L. Crawford, and K.K. Yang# (2023). Deep self-supervised learning for biosynthetic gene cluster detection and product classification. *PLOS Computational Biology*. 19(5): e1011162.
- [29] J. Stamp#, A. DenAdel, D. Weinreich, and L. Crawford# (2023). Leveraging the genetic correlation between traits improves the detection of epistasis in genome-wide association studies. *G3: Genes, Genomes, Genetics.* **13**(8): jkad118.
- [30] H. Adam#, F. Yin, M. Hu, N. Tenenholtz, **L. Crawford**, L. Mackey, and A. Koenecke (2023). Should I stop or should I go: early stopping with heterogeneous populations. *Advances in Neural Processing Systems (NeurIPS)*.
- [31] <u>E.T. Winn-Nuñez</u>#, M. Griffin, and **L. Crawford**# (2024). A simple approach for local and global variable importance in nonlinear regression models. *Computational Statistics & Data Analysis*. **194**: 107914.

PREPRINTS (*CO-FIRST AUTHORS; †CO-SENIOR AUTHORS; #CORRESPONDING AUTHOR(S); ADVISEE)

- [1] **L. Crawford**# and X. Zhou#. Genome-wide marginal epistatic association mapping in case-control studies. *bioRxiv*. 374983.
- [2] J. Ish-Horowicz*, <u>D. Udwin</u>*, <u>K. Scharfstein</u>, S.R. Flaxman, S.L. Filippi#, and **L. Crawford**#. Interpreting deep neural networks through variable importance. *arXiv*. 1901.09839.
- [3] K.E. Ware, S. Gupta, J. Eng, G. Kemeny, B.J. Puviindran, W.C. Foo, **L. Crawford**, R.G. Almquist, D. Runyambo, B.C. Thomas, M.U. Sheth, A. Agarwal, M. Pierobon, E.F. Petricoin, D.L. Corcoran, J. Freedman, S.R. Patierno, T. Zhang, S. Gregory, Z. Sychev, J.M. Drake, A.J. Armstrong#, J.A. Somarelli#. Convergent evolution of p38/MAPK activation in hormone resistant prostate cancer mediates pro-survival, immune evasive, and metastatic phenotypes. *bioRxiv*. 2020.04.22.050385.
- [4] W. Cheng, G. Darnell, S. Ramachandran, and L. Crawford#. Generalizing variational autoencoders with hierarchical empirical Bayes. *arXiv*. 2007.10389.
- [5] M.C. Turchin#, <u>G. Darnell</u>, **L. Crawford**#, and S. Ramachandran#. Pathway analysis within multiple human ancestries reveals novel signals for epistasis in complex traits. *bioRxiv*. 2020.09.24.312421.
- [6] K. Meng#, J. Wang, **L. Crawford**, and A. Eloyan. Randomness and statistical inference of shapes via the smooth Euler characteristic transform. *arXiv*. 2204.12699.
- [7] S.P. Smith*, <u>G. Darnell</u>*, <u>D. Udwin</u>, A. Harpak, S. Ramachandran†, and **L. Crawford**†#. Accounting for statistical non-additive interactions enables the recovery of missing heritability from GWAS summary statistics. *bioRxiv*. 2022.07.21.501001.
- [8] K. Meng#, M. Ji, J. Wang, K. Ding, H. Kirveslahti, A. Eloyan, and **L. Crawford**. Statistical inference on grayscale images via the Euler-Radon transform. *arXiv*. 2308.14249.
- [9] <u>H. Xie</u>, **L. Crawford**†#, and <u>A. Conard</u>†#. Multioviz: an interactive platform for in silico perturbation and interrogation of gene regulatory networks. *bioRxiv*. 2023.10.10.561790.

- [10] <u>K.Z. Kedzierska</u>#, **L. Crawford**†, A.P. Amini†, and A.X. Lu†#. Assessing the limits of zero-shot foundation models in single-cell biology. *bioRxiv*. 2023.10.16.561085.
- [11] <u>E.T. Winn-Nuñez</u>#, H. Witt, D. Bhaskar, R.Y. Huang, J.S. Reichner, I.Y. Wong, and **L. Crawford**#. Generative modeling of biological shapes and images using a probabilistic α -shape sampler. *bioRxiv*. 2024.01.09.574919.
- [12] <u>C. Nwizu</u>, M. Hughes, M.L. Ramseier, A. Navia, A.K. Shalek, N. Fusi, S. Raghavan†, P.S. Winter†, A.P. Amini†#, and **L. Crawford**†#. Scalable nonparametric clustering with unified marker gene selection for single-cell RNA-seq data. *bioRxiv*. 2024.02.11.579839.
- [13] A. DenAdel, M.L. Ramseier, A. Navia, A.K. Shalek, S. Raghavan, P.S. Winter, A.P. Amini, and L. Crawford#. A knockoff calibration method to avoid over-clustering in single-cell RNA-sequencing. bioRxiv. 2024.03.08.584180.

NON-REFEREED PAPERS

[1] At the intersection of machine learning, biology, and health: an interview with Lorin Crawford (2021). *Communications Biology.* **4**(1): 32. [Link]

SOFTWARE

- [1] α -Shape Sampler: Pipeline for Generating 2D and 3D Biological Shapes and Images
- [2] BAKR: Bayesian Approximate Kernel Regression
- [3] BANNs: Biologically Annotated Neural Networks
- [4] callback: Calibrated Clustering via Knockoffs
- [5] ESNN: Ensemble of Single-Effect Neural Networks
- [6] **gene**- ε : A Recalibrated Hypothesis Test for Sets of SNP-Level Summary Statistics
- [7] **GOALS**: The GlObal And Local Score
- [8] Grid-LMM: Fast and Flexible Linear Mixed Models for Genetic Association Studies
- [9] **HEBAE**: Hierarchical Empirical Bayes Autoencoder
- [10] i-LDSC: Interaction-LD Score Regression
- [11] **MAPIT**: The MArginal ePIstasis Test
- [12] MAPIT-R: The MArginal ePIstasis Test for Regions and SNP-Sets
- [13] MegaLMM: Mega-scale Linear Mixed Models for Multivariate Genomic Prediction
- [14] Multioviz: An Interactive Platform for Analyzing Gene Regulatory Networks
- [15] **mvMAPIT**: The Multivariate MArginal ePIstasis Test
- [16] NCLUSION: Nonparametric Clustering of Single-cell Populations
- [17] RATE: RelATive cEntrality Measures for Variable Prioritization
- [18] **SECT**: The Smooth Euler Characteristic Transform
- [19] **SINATRA**: Pipeline for Sub-Image Analysis and Feature Selection on 3D Shapes
- [20] SINATRA Pro: Protein Structure and Conformation Analysis using Topological Summary Statistics
- [21] **Tropix:** Tropical Sufficient Statistics for Persistent Homology

AWARDS & FELLOWSHIPS

Committee of Presidents of Statistical Societies (COPSS) Emerging Leader Award	2023
International Association for Intelligent Biology and Medicine (IAIBM) Eminent Scholar Award	2023
Mathematical Association of America & National Association of Mathematicians Lecturer	2022
Kavli Fellow of the National Academy of Sciences, Frontiers of Science Symposium	2021
Cell Press: 1000 Inspiring Black Scientists in America	2021
David & Lucille Packard Foundation Fellowship for Science and Engineering	2020

Mathematically Gifted & Black: Black History Month Honoree The Root: 100 Most Influential African Americans in 2019 Endowed Named Assistant Professorship Alfred P. Sloan Research Fellowship Forbes 30 Under 30 Class of 2019: Science Leonard J. Savage Dissertation Award in Applied Methodology National Science Foundation (NSF) Graduate Research Fellowship Duke University Dean Graduate Fellowship Isabella T. Jenkins Outstanding Academic Achievement Award J.J. Dennis Endowed Undergraduate Fellowship Clark Atlanta University Provost Scholarship	2020 2019 2019 2019 2019 2018 2015 2013 2013 2012 2009
P20GM103645 (PI Sanes) NIH/NIGMS Title: COBRE Center for Central Nervous System Function Role: Core B Co-Investigator	09/01/18 - 09/08/20
2U10CA180794 (PIs Gray and Gatsonis) NIH/NCI/Dana Farber Cancer Institute Title: ECOG-ACRIN Network Group Statistics and Data Management Center Role: Biostatistician	03/01/19 – 09/08/20
W81XWH-18-1-018 (PI Somarelli) DoD/PCRP Title: Targeting the p38/Snail/PD-L1 axis in Hormone-therapy Resistance and Me Role: Co-Investigator	09/01/18 – 08/31/21 etastasis
P20GM109035 (PI Rand) NIH/NIGMS Title: COBRE Center for Computational Biology of Human Disease Project Title: Deep learning Methods for Fine Mapping and Discovery in Genomi Role: Principal Investigator of Project	03/01/19 – 02/28/21 c Association Studies
FG-2019-11622 (PI Crawford) Alfred P. Sloan Foundation Research Fellowship Title: Interpretable Machine Learning Methods for Genome-wide Association MagRole: Principal Investigator	09/15/19 – 09/14/21 pping
2020-71387 (PI Crawford) David & Lucille Packard Foundation Fellowship for Science and Engineering Role: Principal Investigator	10/15/20 – 10/14/25
1R35GM138032-01 (PI Crawford) NIH/NIGMS Title: Interpretable Machine Learning for Characterizing Broad-sense Heritabilit Rare Diseases Role: Principal Investigator	Deferred ty in Complex Traits and
AS ASSOCIATE PROFESSOR / PRINCIPAL RESEARCHER	
Joint Statistical Meetings, Invited Paper Session, Portland, OR Biology of Genomes, Computational and Statistical Genomics Session, Cold Sprin	2024 ng Harbor, NY 2024

INVITED TALKS

SPONSORED RESEARCH

Joint Statistical Meetings, Invited Paper Session, Portland, OR

Biology of Genomes, Computational and Statistical Genomics Session, Cold Spring Harbor, NY
University of Tennessee Health Science Center, Biostatistics Seminar Series, Virtual
Joint Mathematical Meetings, AMS Special Session, San Francisco, CA

2024

 $Johns\ Hopkins\ University, Department\ of\ Applied\ Mathematics\ and\ Statistics\ Seminar,\ Baltimore,\ MD\ 2023$

University of Southern California, QCB Seminar Series, Los Angeles, CA Grace Hopper Celebration, Data Science Lecture, Orlando, FL Babson College, MAST Seminar Series, Babson Park, MA International Conference on Intelligent Biology and Medicine, Eminent Scholar, Tampa Bay, FL Mount Sinai, Charles Bronfman Institute for Personalized Medicine (CBIPM) Seminar, NYC, NY ICLR, Machine Learning & Global Health Network Workshop, Hybrid, Kigali, Rwanda Applied Algebraic Topology Research Network, Online Seminar Series, Virtual UMass Chan Medical School, Department of Systems Biology Seminar, Worcester, MA IMSI, Randomness in Topology and its Applications Workshop, Chicago, IL Cedars-Sinai, Computational Biomedicine Grand Rounds Seminar, Los Angeles, CA NCI, Division of Cancer Epidemiology and Genetics, Biostatistics Branch Seminar Series, Virtual	2023 2023 2023 2023 2023 2023 2023 2023
Johns Hopkins University, Statistical Genetics Working Group Seminar, Virtual Stanford University, Department of Biomedical Data Science, Biostatistics Workshop, Palo Alto, CA Georgia Institute of Technology, School of Biological Sciences Seminar, Atlanta, GA University of Rhode Island, Department of Computer Science and Statistics Seminar, Kingston, RI University of Oxford, Applied Topology Seminar, Oxford, England, UK University of Cambridge, MRC Biostatistics Unit Seminar, Cambridge, England, UK University of California Santa Cruz, Department of Statistics Seminar, Virtual SIAM Conference on Mathematics of Data Science, San Diego, CA Joint Statistical Meetings, Biometrics Invited Session, Washington, D.C. SIAM Annual Meeting, Plenary Speaker, Pittsburgh, PA	2022 2022 2022 2022 2022 2022 2022 202
AS ASSISTANT PROFESSOR / SENIOR RESEARCHER	
Bayesian Young Statisticians Meeting (BAYSM), Plenary Speaker, Montréal, Québec, CA WNAR/IMS/JR Meeting, IMS Invited Session, Virtual Population, Evolutionary, and Quantitative Genetics (PEQG), Pacific Grove, CA NES/MAA Spring Meeting, Plenary Talk, Haddam, CT FASEB, The Cell Signaling in Cancer Conference: From Mechanisms to Therapy, New Orleans, LA Yale University, CBDS Distinguished Speaker Seminar Series, Virtual Duke University, Department of Statistical Sciences Seminar Series, Durham, NC KS/MAA Fall Meeting, Plenary Talk, Atchison, KS Conference on Health, Inference, and Learning (CHIL), Keynote Speaker, Virtual Brown University, Provost's By Faculty For Faculty Lecture Series, Providence, RI University of Chicago, Department of Human Genetics Seminar Series, Virtual University of Pennsylvania, Bioinformatics Seminar Series, Virtual Toronto Bioinformatics Users Group (TorBUG) Seminar Series, Virtual Joint Mathematical Meetings, AMS Special Session, Seattle, WA	2022 2022 2022 2022 2022 2022 2022 202
Winston-Salem State University, Math and Stats Club, Virtual University of Chicago, Department of Statistics Colloquium, Virtual Penn State, ICDS Symposium, Keynote Speaker, Virtual Meeting Cornell University, Plant Breeding and Genetic Seminar Series, Virtual Atlanta University Center Data Science Symposium, Keynote Speaker, Virtual Meeting Joint Statistical Meetings, Biometrics Invited Session, Virtual Meeting SMB, Data-Driven Modeling and Analysis in Mathematical Biology, Virtual Meeting Great Lakes Bioinformatics Conference, Keynote Speaker, Virtual Meeting ICLR, Geometrical and Topological Representation Learning Workshop, Virtual Meeting IMSI, Topological Data Analysis Workshop, Virtual New York University and ETH Zürich, Math and Data (MAD+) Seminar, Virtual Meeting Columbia University, DSI Distinguished Speaker Series, Virtual ProbGen, Quantitative Genetics and Association Mapping Session, Virtual Meeting University of Michigan, Dept. of Biostatistics Seminar, Virtual Princeton University, Quantitative and Computational Biology Seminar Series, Virtual Johns Hopkins University, Dept. of Biostatistics Seminar Series, Virtual	2021 2021 2021 2021 2021 2021 2021 2021
NeurIPS, Topological Data Analysis and Beyond Workshop, Virtual Meeting University of Colorado, Biostatistics Seminar Series, Virtual	2020 2020

University of Wisconsin-Madison, Dept. of Statistics Seminar, Virtual	2020
University of North Carolina, Dept. of Biostatistics Seminar, Virtual	2020
University of Pennsylvania, The Wharton School Statistics Seminar, Virtual	2020
Rochester Institute of Technology, Mathematical Modeling Seminar, Virtual	2020
The Black Women in Computational Biology Network, Seminar Series, Virtual Meeting	2020
Joint Statistical Meetings, Biometrics Invited Session, Virtual	2020
Stanford University, Statistics Department Seminar, Virtual	2020
Brown University, Rhode Island IDeA Symposium (Invited Science Talk), Virtual	2020
University of Arkansas for Medical Sciences, Biomedical Informatics Seminar, Virtual	2020
Microsoft Research New England, Seminar Series, Cambridge, MA	2020
ENAR Spring Meeting, Invited Session, Nashville, TN	2020
Brown University and Lifespan, Populations Science Group Meeting, Providence, RI	2020
Duke University, Sloan Research Summit (Keynote Speaker), Durham, NC	2020
University of Massachusetts Amherst, Statistics and Probability Seminar, Amherst, MA	2020
NES/MAA Fall Meeting, Plenary Talk, Wellesley, MA	2019
Broad Institute of MIT and Harvard, Models, Inference & Algorithms Seminar, Cambridge, MA	2019
Duke University, Computational Biology Seminar, Durham, NC	2019
Joint Statistical Meetings, IMS Invited Session, Denver, CO	2019
WNAR/IMS/JR Meeting, IMS Invited Session, Portland, OR	2019
33rd New England Statistics Symposium (NESS), Hartford, CT	2019
UC Irvine, Dept. of Statistics Seminar, Irvine, CA	2019
Brown University, Center for Computational Biology of Human Disease Seminar, Providence, RI	2019
ENAR Spring Meeting, IMS Invited Session, Philadelphia, PA	2019
University of Connecticut, Dept. of Statistics Seminar, Storrs, CT	2018
University of Michigan, Dept. of Biostatistics Seminar, Ann Arbor, MI	2018
Harvard University, Data Science Initiative Conference, Cambridge, MA	2018
Brown University, Pattern Theory Seminar Series, Providence, RI	2018
ISBA World Meeting, Edinburgh, Scotland, UK	2018
College of the Holy Cross, Pi Mu Epsilon (PME) Colloquium, Worcester, MA	2018
ENAR Spring Meeting, Geometry and Topology in Statistical Inference Workshop, Atlanta, GA	2018
42nd SIAM-SEAS, Statistical Topological Data Analysis Mini Symposium, Chapel Hill, NC	2018
ICERM, NSF TRIPODS: Geometry and Topology of Data Workshop, Providence, RI	2017
NeurIPS, Synergies in Geometric Data Analysis Workshop, Long Beach, CA	2017
Brown University, Data Science Initiative Seminar, Providence, RI	2017
Brown University, Center for Computational Molecular Biology Seminar, Providence, RI	2017
Brown University, Applied Topology and Geometry Seminar, Providence, RI	2017
SIAM AG'17, Statistics and Applied Algebraic Topology Workshop, Atlanta, GA	2017

PROFESSIONAL AFFILIATIONS

American Statistical Association (ASA); Genetics Society of America (GSA); International Biometric Society Eastern North American Region (IBS ENAR); Institute of Mathematical Statistics (IMS); International Society for Bayesian Analysis (ISBA)

SERVICE ACTIVITIES

EDITORIAL SERVICE

Biostatistics (Associate Editor)	2021 – Present
Journal of the American Statistical Association (AE of Reproducibility)	2018 – Present

EXTERNAL SERVICE

ENAR Regional Committee	2024 – Present
IMS Outreach Committee	2023 – Present
ENAR Fostering Diversity in Biostatistics Workshop (Planning Committee)	2022 – Present
ISBA Section on Biostats and Pharma (Treasurer)	2021 - 2023
ENAR Regional Advisory Board	2020 - 2022

PROFESSIONAL SERVICE

Banff International Research Station (BIRS), Alberta, Canada Statistical, Computational, Translational, and Ethical Challenges in Biobank Data Analysis Role: Co-Organizer	2024
Biology of Genomes, Cold Spring Harbor, NY Session: Computational and Statistical Genomics Role: Co-Chair	2024
ENAR Spring Meeting, Baltimore, MD Role: Program Committee	2024
Advances in the Genetic Architecture of Complex Human Traits Workshop, Washington, D.C. National Human Genome Research Institute Role: Scientific Organizing Committee	2023
New England Statistical Society (NESS) Conference, Boston, MA Role: Keynote Panel Organizing Committee	2023
The Junior Section of the International Society of Bayesian Analysis Blackwell-Rosenbluth Award Scientific Committee Role: Chair	2022
Conference on Neural Information Processing Systems (NeurIPS), New Orleans, LA Workshop: Learning Meaningful Representations of Life Role: Organizing Committee	2022
Conference on Neural Information Processing Systems (NeurIPS), New Orleans, LA Role: Workshop Proposal Reviewer	2022
35th New England Statistics Symposium (NESS), Storrs, CT MassMutual Student Paper Award Role: Review Committee	2022
Joint Statistical Meetings (JSM), Washington, D.C. Session: Imaging and Clinical Biomarkers in Neurodegenerative Disease Role: Co-Organizer	2022
Population, Evolutionary, and Quantitative Genetics (PEQG), Pacific Grove, CA Role: Session Chair	2022
Intelligent Systems for Molecular Biology (ISMB), Virtual Meeting Role: Proceedings Program Committee	2022
Conference on Neural Information Processing Systems (NeurIPS), Virtual Meeting Workshop: Learning Meaningful Representations of Life Role: Organizing Committee	2021
The Junior Section of the International Society of Bayesian Analysis Blackwell-Rosenbluth Award Scientific Committee Role: Vice Chair	2021
Probabilistic Modeling in Genomics (ProbGen), Virtual Meeting Session: Quantitative Genetics and Association Mapping Role: Co-Chair	2021

2022

Intelligent Systems for Molecular Biology (ISMB), Virtual Meeting European Conference on Computational Biology (ECCB), Virtual Meeting

Role: Proceedings Program Committee

REVIEWER SERVICE

American Journal of Human Genetics; Annals of Applied Statistics; Bioinformatics; Biostatistics; BMC Bioinformatics; Conference on Neural Information Processing Systems (NeurIPS); Genes; Intelligent Systems for Molecular Biology (ISMB); International Conference on Artificial Intelligence and Statistics (AISTATS); International Conference on Machine Learning (ICML); Journal of the American Statistical Association; Journal of Animal Science; Journal of Computational and Graphical Statistics; Journal of Machine Learning Research; Journal of Multivariate Analysis; Nature Communications; Nature Genetics; Nature Protocols; New England Journal of Medicine; PLOS Genetics; Research in Computational Molecular Biology (RECOMB); Transactions of Mathematics and Its Applications

INSTITUTIONAL SERVICE (BROWN UNIVERSITY)

Center for Computational Molecular Biology: Executive Steering Committee	2022 – Present
Data Science Initiative (DSI) Campus Advisory Board	2021 – Present
Center for Computational Molecular Biology: PhD Admissions Committee	2017 – Present
Goldwater Scholarship Nomination Committee	2019 - 2020
School of Public Health: Operational Planning Committee	2019 - 2020
Department of Biostatistics: PhD Admissions Committee	2018 - 2020
Department of Biostatistics: Seminar Series Organizer	2018 - 2020
Department of Biostatistics: Communications Committee	2017 - 2020
Department of Biostatistics: Academic Programs Committee	2017 - 2020

MENTORSHIP ACTIVITIES

POSTDOCTORAL FELLOWS

Greg Darnell (Co-advisee) 2019 – 2020

DOCTORAL STUDENTS

Kun Meng (Biostatistics)

Alan DenAdel (Computational Biology)	Present
Cecile Meier-Scherling (Computational Biology; Co-advisee)	Present
Chibuikem Nwizu (Computational Biology)	Present
Whitney Sloneker (Computational Biology; Co-advisee)	Present
Julian Stamp (Computational Biology)	Present
Ria Vinod (Computational Biology)	Present
Alexandra Wong (Computational Biology)	Present
Emily Winn-Nuñez (Applied Mathematics)	2024
Dana Udwin (Biostatistics)	2023
Ashley Conard (Computational Biology)	2022
Wai Shing Tang (Physics)	2022
Wei Cheng (Computational Biology; Co-advisee)	2022

DOCTORAL DISSERTATION COMMITTEES

Anushka Narayanan (Earth, Environmental, and Planetary Sciences)	Present
Rob Zielinski (Biostatistics)	Present
Alysandra Zhang (Cognitive Linguistic & Psychological Sciences)	Present
Rachel Gaither (Epidemiology)	Present
Isaac Kim (Computational Biology)	Present
Dilum Aluthge (Computational Biology)	Present
Haobo Yang (Chemistry)	Present
Vivek Ramanan (Computational Biology)	Present
Jiaqi Zhang (Computer Science)	Present
Qing Wu (Computational Biology)	2023
Pinar Demetci (Computational Biology)	2023

Topi Paananen (Aalto University - Finland) Samuel Smith (Computational Biology) Haobo Yang (Chemistry) Adrienne Parsons (Molecular Pharmacology, Physiology, and Biotechnology) Dhananjay Bhaskar (Biomedical Engineering) Sahar Shahamatdar (Computational Biology)	2022 2022 2022 2021 2021 2021
MASTERS THESIS ADVISING	
Tara Amruther (Biostatistics)	Present
Colin Small (Biotechnology) Alexander Li (Biostatistics) Isaac Zhao (Biostatistics) Bruce Wang (Data Science Institute)	2023 2021 2019 2018
UNDERGRADUATE HONORS THESIS ADVISING	
Ryan Huang (Computer Science) Helen Xie (Computer Science)	Present Present
Lee Ding (Applied Math) Erin Bugbee (Statistics) Gabrielle Ferra (Applied Math-Biology) Kayla Scharfstein (Applied Math-Computer Science) Zachary Kaplan (Applied Math) Timothy Sudijono (Applied Math)	2022 2020 2020 2020 2019 2019
MICROSOFT RESEARCH INTERNS	
Kasia Kedzierska (Oxford) Zeinab Navidi (University of Toronto) Giovanni Palla (Helmholtz Munich) Avish Vijayaraghavan (Imperial College London) Fatemeh Afrasiabi (UMass Boston) Dan Yuan (U Washington) Brian Trippe (MIT) Yaniv Yacoby (Harvard)	2023 2023 2023 2023 2022 2022 2022 2021 2021
PHP0100 - First Year Seminar: Statistics is Everywhere PHP2601 - Linear Models PHP2605 - Generalized Linear Models PHP2950 - Doctoral Seminar in Public Health	

[CV compiled on 2024-03-28]

COURSES TAUGHT