Jiebiao Wang, PhD

Department of Biostatistics
University of Pittsburgh

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Education

- 2012-2017 Ph.D., Biostatistics, University of Chicago, Chicago, Illinois
- 2010-2012 Master, Statistics, Renmin University of China, Beijing, China
- 2006–2010 Bachelor, Statistics, Renmin University of China, Beijing, China

Appointments and Positions

- 2020/1-Present Assistant Professor (Tenure Track), Department of Biostatistics, University of Pittsburgh, Pittsburgh, PA
- 2023/7-Present Assistant Professor (Secondary), Clinical and Translational Science Institute (CTSI), University of Pittsburgh, PA
 - 2017–2019 Postdoctoral Researcher, Department of Statistics and Data Science, Carnegie Mellon University, Pittsburgh, PA

Publications/Preprints

Major/methodological work

- +: corresponding author; *: co-first author; : PhD advised
- [1] P. Huang, M. Cai, X. Lu, C. McKennan, and J. Wang⁺, "Accurate estimation of rare cell-type fractions from tissue omics data via hierarchical deconvolution," *The Annals of Applied Statistics*, vol. 18, no. 2, pp. 1178 1194, 2024.
- [2] M. Cai, J. Zhou, C. McKennan, and **J. Wang**⁺, "scMD facilitates cell type deconvolution using single-cell dna methylation references," *Communications Biology*, vol. 7, no. 1, p. 1, 2024.
- [3] P. B. Chandrashekar, S. Alatkar, J. Wang, G. E. Hoffman, C. He, T. Jin, S. Khullar, J. Bendl, J. F. Fullard, P. Roussos, and D. Wang, "DeepGAMI: deep biologically guided auxiliary learning for multimodal integration and imputation to improve genotype–phenotype prediction," *Genome Medicine*, vol. 15, no. 1, p. 88, 2023.
- [4] M. Cai, M. Yue, T. Chen, J. Liu, E. Forno, X. Lu, T. Billiar, J. Celedón, C. McKennan, W. Chen⁺, and **J. Wang**⁺, "Robust and accurate estimation of cellular fraction from tissue omics data via ensemble deconvolution," *Bioinformatics*, vol. 38, no. 11, pp. 3004–3010, 2022.
- [5] **J. Wang**⁺, K. Roeder⁺, and B. Devlin⁺, "Bayesian estimation of cell type–specific gene expression with prior derived from single-cell data," *Genome research*, vol. 31, no. 10, pp. 1807–1818, 2021.
- [6] J. Tian, **J. Wang**, and K. Roeder, "ESCO: single cell expression simulation incorporating gene co-expression," *Bioinformatics*, vol. 37, no. 16, pp. 2374–2381, 2021.
- [7] Y. Qiu, J. Wang, J. Lei, and K. Roeder, "Identification of cell-type-specific marker

- genes from co-expression patterns in tissue samples," *Bioinformatics*, vol. 37, no. 19, pp. 3228–3234, 2021.
- [8] F. Yang, K. Gleason, **J. Wang**, J. Duan, X. He, B. L. Pierce, and L. S. Chen, "CCmed: cross-condition mediation analysis for identifying replicable trans-associations mediated by cis-gene expression," *Bioinformatics*, vol. 37, no. 17, pp. 2513–2520, 2021.
- [9] J. Wang, B. Devlin, and K. Roeder, "Using multiple measurements of tissue to estimate subject-and cell-type-specific gene expression," *Bioinformatics*, vol. 36, no. 3, pp. 782– 788, 2020.
- [10] K. Satterstrom*, J. Kosmicki*, J. Wang*, M. S. Breen, S. De Rubeis, J.-Y. An, M. Peng, R. Collins, J. Grove, L. Klei, C. Stevens, J. Reichert, M. S. Mulhern, M. Artomov, S. Gerges, B. Sheppard, X. Xu, A. Bhaduri, U. Norman, H. Brand, G. Schwartz, R. Nguyen, E. E. Guerrero, C. Dias, Autism Sequencing Consortium, iPSYCH-Broad Consortium, C. Betancur, E. H. Cook, L. Gallagher, M. Gill, J. S. Sutcliffe, A. Thurm, M. E. Zwick, A. D. Børglum, M. W. State, A. E. Cicek, M. E. Talkowski, D. J. Cutler, B. Devlin, S. J. Sanders, K. Roeder, M. J. Daly, and J. D. Buxbaum, "Large-scale exome sequencing study implicates both developmental and functional changes in the neurobiology of autism," Cell, vol. 180, no. 3, pp. 568–584.e23, 2020.
- [11] S. Chen*, **J. Wang***, E. Cicek, K. Roeder, H. Yu, and B. Devlin, "De novo missense variants disrupting protein–protein interactions affect risk for autism through gene coexpression and protein networks in neuronal cell types," *Molecular autism*, vol. 11, no. 1, pp. 1–16, 2020.
- [12] **J. Wang**, P. Wang, D. Hedeker, and L. S. Chen, "Using multivariate mixed-effects selection models for analyzing batch-processed proteomics data with non-ignorable missingness," *Biostatistics*, vol. 20, no. 4, pp. 648–665, 2019.
- [13] R. Gibbons, K. Hur, J. Lavigne, **J. Wang**, and J. J. Mann, "Medications and Suicide: High Dimensional Empirical Bayes Screening (iDEAS)," *Harvard Data Science Review*, vol. 1, no. 2, 2019.
- [14] **J. Wang***, Q. Liu*, B. Pierce, D. Huo, O. Olopade, H. Ahsan, and L. Chen, "A meta-analysis approach with filtering for identifying gene-level gene-environment interactions," *Genetic epidemiology*, vol. 42, no. 5, pp. 434–446, 2018.
- [15] F. Yang, J. Wang, The GTEx Consortium, B. L. Pierce, and L. S. Chen, "Identifying cis-mediators for trans-eqtls across many human tissues using genomic mediation analysis," *Genome research*, vol. 27, no. 11, pp. 1859–1871, 2017.
- [16] L. S. Chen, **J. Wang**, X. Wang, and P. Wang, "A mixed-effects model for incomplete data from labeling-based quantitative proteomics experiments," *The annals of applied statistics*, vol. 11, no. 1, pp. 114–138, 2017.
- [17] J. Wang, E. R. Gamazon, B. L. Pierce, B. E. Stranger, H. K. Im, R. D. Gibbons, N. J. Cox, D. L. Nicolae, and L. S. Chen, "Imputing gene expression in uncollected tissues within and beyond GTEx," *The American Journal of Human Genetics*, vol. 98, no. 4, pp. 697–708, 2016.

Collaborative work

[1] J.-M. Pyun, Y. H. Park, **J. Wang**, D. A. Bennett, P. J. Bice, J. P. Kim, S. Kim, A. J. Saykin, and K. Nho, "Transcriptional risk scores in alzheimer's disease: From pathology to cognition," *Alzheimer's & Dementia*, vol. 20, no. 1, pp. 243–252, 2024.

- [2] J.-M. Pyun, Y. H. Park, J. Wang, P. J. Bice, D. A. Bennett, S. Kim, A. J. Saykin, and K. Nho, "Aberrant GAP43 gene expression is alzheimer disease pathology-specific," *Annals of Neurology*, vol. 93, no. 5, pp. 1047–1048, 2023.
- [3] H.-W. Liang, N. Snyder, J. Wang, X. Xun, Q. Yin, K. LeWinn, K. N. Carroll, N. R. Bush, K. Kannan, E. S. Barrett, R. T. Mitchell, F. Tylavsky, and J. J. Adibi, "A study on the association of placental and maternal urinary phthalate metabolites," *Journal of exposure science & environmental epidemiology*, vol. 33, no. 2, pp. 264–272, 2023.
- [4] W. H. Cuddleston, X. Fan, L. Sloofman, L. Liang, E. Mossotto, K. Moore, S. Zipkowitz, M. Wang, B. Zhang, J. Wang, N. Sestan, B. Devlin, K. Roeder, S. J. Sanders, J. D. Buxbaum, and M. S. Breen, "Spatiotemporal and genetic regulation of A-to-I editing throughout human brain development," *Cell Reports*, vol. 41, no. 5, p. 111585, 2022.
- [5] Z. Xu, X. Wang, L. Fan, F. Wang, B. Lin, J. Wang, G. Trevejo-Nuñez, W. Chen, and K. Chen, "Integrative analysis of spatial transcriptome with single-cell transcriptome and single-cell epigenome in mouse lungs after immunization," iScience, p. 104900, 2022.
- [6] B. C. McKinney, L. L. McClain, C. M. Hensler, Y. Wei, L. Klei, D. A. Lewis, B. Devlin, J. Wang, Y. Ding, and R. A. Sweet, "Schizophrenia-associated differential DNA methylation in brain is distributed across the genome and annotated to MAD1L1, a locus at which DNA methylation and transcription phenotypes share genetic variation with schizophrenia risk," *Translational psychiatry*, vol. 12, no. 1, pp. 1–12, 2022.
- [7] S. C. Piantadosi, L. L. McClain, L. Klei, J. Wang, B. L. Chamberlain, S. A. Springer, D. A. Lewis, B. Devlin, and S. E. Ahmari, "Transcriptome alterations are enriched for synapse-associated genes in the striatum of subjects with obsessive-compulsive disorder," *Translational psychiatry*, vol. 11, no. 1, pp. 1–11, 2021.
- [8] C. A. King, D. Brent, J. Grupp-Phelan, T. C. Casper, J. M. Dean, L. S. Chernick, J. A. Fein, E. M. Mahabee-Gittens, S. J. Patel, R. D. Mistry, S. Duffy, M. Melzer-Lange, A. Rogers, D. M. Cohen, A. Keller, R. Shenoi, R. W. Hickey, M. Rea, M. Cwik, K. Page, T. C. McGuire, J. Wang, R. Gibbons, and Pediatric Emergency Care Applied Research Network, "Prospective development and validation of the computerized adaptive screen for suicidal youth," *JAMA psychiatry*, vol. 78, no. 5, pp. 540–549, 2021.
- [9] M. L. Seney, S.-M. Kim, J. R. Glausier, M. A. Hildebrand, X. Xue, W. Zong, J. Wang, M. A. Shelton, B. N. Phan, C. Srinivasan, A. R. Pfenning, G. C. Tseng, D. A. Lewis, Z. Freyberg, and R. W. Logan, "Transcriptional alterations in dorsolateral prefrontal cortex and nucleus accumbens implicate neuroinflammation and synaptic remodeling in opioid use disorder," *Biological Psychiatry*, vol. 90, no. 8, pp. 550–562, 2021.
- [10] N. F. Fitz, **J. Wang**, M. I. Kamboh, R. Koldamova, and I. Lefterov, "Small nucleolar RNAs in plasma extracellular vesicles and their discriminatory power as diagnostic biomarkers of alzheimer's disease," *Neurobiology of Disease*, vol. 159, p. 105481, 2021.
- [11] S. Chen, R. Fragoza, L. Klei, Y. Liu, **J. Wang**, K. Roeder, B. Devlin, and H. Yu, "An interactome perturbation framework prioritizes damaging missense mutations for developmental disorders," *Nature genetics*, vol. 50, no. 7, pp. 1032–1040, 2018.
- [12] J. Pitt, M. Riester, Y. Zheng, T. Yoshimatsu, A. Sanni, O. Oluwasola, A. Veloso, E. Labrot, S. Wang, A. Odetunde, A. Ademola, B. Okedere, S. Mahan, R. Leary, M. Macomber, M. Ajani, R. Johnson, D. Fitzgerald, J. Grundstad, J. Tuteja, G. Khramtsova, J. Zhang, E. Sveen, B. Hwang, W. Clayton, C. Nkwodimmah, B. Famooto, E. Obasi, V. Aderoju, M. Oludara, F. Omodele, O. Akinyele, A. Adeoye, T. Ogundiran, C. Babalola, K. MacIsaac, A. Popoola, M. Morrissey, L. S. Chen, J. Wang, C. O. Olopade, A. G.

Falusi, W. Winckler, K. Haase, P. V. Loo, J. Obafunwa, D. Papoutsakis, O. Ojengbede, B. Weber, N. Ibrahim, K. P. White, D. Huo, O. I. Olopade, and J. Barretina, "Characterization of nigerian breast cancer reveals prevalent homologous recombination deficiency and aggressive molecular features," *Nature communications*, vol. 9, no. 1, p. 4181, 2018.

Funding

As Principal Investigator

- 2023/2-2027/11 **R01AG080590** (Role: **Contact PI**), NIH, "Statistical methods for population-level cell-type-specific analyses of tissue omics data for Alzheimer's disease", Total: \$1,853,793
- 2022/9-2024/9 **R03OD034501** (Role: **Contact PI**), NIH, "Integration of GTEx and HuBMAP data to gain population-level cell-type-specific insights", \$314,739 (Common Fund initiative through the Office of the NIH Director)
- 2021/7-2023/6 Competitive Medical Research Fund (CMRF) (Role: PI), UPMC, \$25,000
- 2022/1-2023/12 Health Sciences Bridge Funding (Role: PI), University of Pittsburgh Schools of Health Sciences, \$48,720
- 2022/1-2023/12 Brain Institute's Assault on Alzheimer's Seed Grant (Role: PI), University of Pittsburgh Brain Institute, \$25,000
- 2022/6-2023/5 Exploring Existing Data Resources Pilot Award (Role: PI), University of Pittsburgh Clinical and Translational Science Institute (CTSI), \$25,000
- 2022/5-2023/4 Public Health Trans-Disciplinary Collaboration Pilot Award (Pls: Wang and Talisa), Pitt CTSI, "Precision Health Methods for Alzheimer's Disease", \$45,000

As Co-Investigator/Subcontract PI

- 2020/5-2024/2 R01MH123184: Computational methods to integrate and interpret the transcriptome from single cell and tissue level data (PI: Roeder), NIH, 1.2 months support per year
- 2020/7-2025/6 R01HL153058: Mucin sialylation drives epithelial cell senescence and severe asthma (PI: Wenzel), NIH, 1.2 months
- 2020/10-2025/8 U19AG068054: Alzheimer's Biomarker Consortium Down Syndrome (ABC-DS) (PI: Handen), NIH, 1.2 months
- 2022/4-2027/4 P01Al106684: Immune airway-epithelial interactions in steroid-refractory severe asthma (PIs: Ray and Wenzel), NIH, 1.2 months
- 2022/7-2026/4 R01DK133454: Vinyl chloride modifies the risk for nonalcoholic fatty liver disease (PI: Beier), NIH, 0.6 months
- 2023/3-2028/2 R01Al164968: Functional consequences of intergenic autoimmune disease risk variants (Pls: Niewold and Demerci), NIH, 1.8 months and 1/2 GSR

Presentations

Invited Presentations

- 2024 JSM, Portland, OR
- 2024 7th International Conference on Econometrics and Statistics (EcoSta), Beijing, China
- 2024 Senior Vice Chancellor's Research Seminar, University of Pittsburgh
- 2024 STATGEN conference, ASA SSGG
- 2024 NIH Common Fund Data Ecosystem March 2024 All-Hands Meeting, Bethesda, MD
- 2024 Biostatistics and Computational Biology Branch Seminar Series, National Institute of Environmental Health Sciences (NIEHS), NIH

- 2023 The 16th International Conference on Computational and Methodological Statistics (CMStatistics), Berlin, Germany
- 2023 Biostatistics Seminars, University of North Carolina at Chapel Hill
- 2023 Accurate estimation of rare cell type fractions from tissue omics data via hierarchical deconvolution, JSM, Toronto, Canada
- 2023 Cross-Pollination Meeting, NIH Common Fund Data Ecosystem
- 2023 Joint Seminar Series of CMDB (Cell, Molecular, and Developmental Biology) GGB (Genetics, Genomics, and Bioinformatics) MCBL (Microbiology), University of California, Riverside
- 2023 Robust and accurate estimation of cellular fraction from tissue omics data via ensemble deconvolution. ENAR Spring Meeting, Nashville, TN
- 2023 Biostatistics Research Day, University of Pittsburgh
- 2022 Robust and accurate estimation of cellular fraction from tissue omics data via ensemble deconvolution. The 15th International Conference on Computational and Methodological Statistics (CMStatistics 2022), King's College London, UK
- 2021 Robust estimation of cell type fractions from bulk data via ensemble of various deconvolution approaches. ICSA Applied Statistics Symposium
- 2021 Detecting cell-type-specific eQTLs Using Bayesian estimates of cell-type-specific gene expression from tissue samples with single-cell prior. ENAR Spring Meeting
- 2020 Bayesian estimation of cell-type-specific gene expression for each tissue sample with prior derived from single-cell data. Department of Statistics Seminar Series, University of Pittsburgh
- 2020 Gene expression deconvolution implicates cell-type-specific gene expression and coexpression in autism. The 75th Anniversary of the Society of Biological Psychiatry (SOBP) Meeting (Canceled due to COVID-19)
- 2019 High-dimensional mixed-effects models for multi-omics data. Department of Biostatistics Seminar Series, University of Pittsburgh
- 2018 A random-effects model for multi-tissue deconvolution to estimate individual-level celltype-specific gene expression. ICSA Applied Statistics Symposium
- 2017 High-Dimensional Multivariate Selection Models for Proteomics Data with Nonignorable Missingness. Department of Human Genetics, University of Pittsburgh
- 2017 A high-dimensional multivariate selection model for proteomics data with nonignorable missingness. Department of Biomedical Informatics, Arizona State University
- 2017 Statistical methods for genomics data with clustered structure and missingness. Division of Biostatistics, Medical College of Wisconsin
- 2016 A high-dimensional multivariate selection model for proteomics data with batch-level missingness. Workshop on Quantitative Research Methods in Education, Health and Social Sciences, University of Chicago

Contributed Presentations

- 2020 Bayesian Subject-Level Bulk Expression Deconvolution and Application to Cell-Type-Specific Differential Expression Analysis, Joint Statistical Meetings (JSM)
- 2019 An empirical Bayes method for deconvolving multi-measure bulk gene expression, JSM
- 2018 Using multi-tissue gene expression to estimate individual- and cell-type-specific expression via deconvolution, American Society of Human Genetics (ASHG) Annual Meeting (*Platform talk*)

- 2017 A High-Dimensional Multivariate Selection Model for Proteomics Data with Batch-Level Missingness, JSM
- 2016 A multivariate selection model for cluster-level outcome-dependent missing data, JSM
- 2016 Mixed-effects models for multivariate clustered data with nonignorable missing outcomes, ENAR Spring Meeting
- 2015 Imputing the transcriptome in inaccessible tissues in and beyond the GTEx project, **JSM**
- 2015 A mixed-effects model for incomplete data with experiment-level abundance-dependent missing-data mechanism, ENAR Spring Meeting

Honors and Awards

- 2024 Selected to Present at the Senior Vice Chancellor's Research Seminar (\$10,000), University of Pittsburgh
- 2023 Nominated for James L. Craig Endowed Excellence in Education Award, School of Public Health, University of Pittsburgh
- 2018 Outstanding Performance Award in the Field of Public Health Sciences, Division of the Biological Sciences, University of Chicago

Mentoring and Advising

As Ph.D. Advisor

2023-Present Chen Liu

- 2022-Present Penghui Huang
 - Research Poster Award, ICSA Applied Statistics Symposium 2023
 - Student Paper Award Honorable Mention, ICSA Applied Statistics Symposium 2023
 - Student Research Award, New England Statistics Symposium 2023
 - Best PhD Poster Award, Pitt Biostatistics Research Day 2023
 - 2019-2024 Mangi Cai: Statistical Methods for Cellular Deconvolution with Single-Cell Omics
 - Best Research Presentation Award, Graduate Student Research Conference 2022, National Institute of Statistical Sciences
 - Best PhD Poster Award, Pitt Biostatistics Research Day 2023
 - Delta Omega Dissertation Award 2024
 - Delta Omega Membership 2024
 - Dean's Service Award 2024

As MS Thesis Advisor

- 2022-2024 Michelle Sun: Age Related Changes in Human Circadian Rhythms
 - Best Master's Poster Award Honorable Mention, Pitt Biostatistics Research Day 2023
- 2022-2024 Jenny Liang: Pan-Tissue Cell Deconvolution Using Single-Cell RNA-Seg References
 - Outstanding Student Award 2024

As Ph.D. Dissertation Committee Member

2023-Present Huy Le (Statistics)

2023-Present Shuangjia Xue (Environmental and Occupational Health)

2023-Present Rain Katz (Epidemiology): Molecular Biomarkers of Subclinical Cardiovascular Disease in the Tobago Health Study and Long Life Family Study

2023-2024 Xiangning Xue: Statistical Methods for Circadian Analysis on Omics Data

2022-2024 Hai-Wei Liang (Epidemiology): A pathway approach in human pregnancy to quantify the effects of phthalates on fetal health outcomes mediated by placental function 2022-2023 Yang Ou: Estimation of Causal Treatment Effect for Clustered Observational Data with **Unmeasured Confounding** 2022-2023 Jian Zou: Clustering, Biomarker and Cancer Model Selection Using Omics Data 2021-2022 Xinjun Wang: Statistical Learning and Analysis of Single-Cell Multi-Omics Data 2020 Dongjing Liu (Human Genetics): Integrated Genome-Wide Analysis of Human Facial Morphology As MS Thesis Committee Member 2023-2024 Japan Patel: Statistical modeling of Epstein-Barr virus infection using scRNA-Seq host expression Jerry Zhou: Impact of COVID-19 on Adversarial Outcomes for In-Patient Congestive 2023-2024 Heart Failure in the Northeast Mid Atlantic 2023-2024 Nuo Wei: Assessing the Impact of Metal Exposure on Chronic Kidney Disease and Hypertension: A NHANES Data Analysis As Ph.D. Academic Advisor 2021-2023 Gehui Zhang 2020-2022 Lingyi Peng As MS Academic Advisor 2023-Present Nick Heaton Teaching Instructor Fall 2024 BIOST 2154, Statistical Methods for Omics Data, 2 credits Fall 2023 BIOST 2068, Introduction to Causal Inference, 3 credits, 13 enrolled Fall 2022 BIOST 2068, Introduction to Causal Inference, 3 credits, 11 enrolled Fall 2021 BIOST 2068, Introduction to Causal Inference, 3 credits, 35 enrolled **Guest Lecturer** Fall 2023 BIOST 2069: Statistical Methods for Omics Data Fall 2023 BIOST 2065: Analysis of Incomplete Data Spring 2022 HUGEN 2080: Statistical Genetics Spring 2020 HUGEN 2080: Statistical Genetics Spring 2020 EPID 2620: Application of Molecular Biomarkers in Epidemiology Coordinator Spring 2022 BIOST 2025, Biostatistics Seminar, 1 credit, 18 enrolled Fall 2021 BIOST 2025, Biostatistics Seminar, 1 credit, 27 enrolled Spring 2021 BIOST 2025, Biostatistics Seminar, 1 credit, 28 enrolled Fall 2020 BIOST 2025, Biostatistics Seminar, 1 credit, 22 enrolled Spring 2020 BIOST 2025, Biostatistics Seminar, 1 credit, 9 enrolled

Service

Grant Review

- 2024/3 Reviewer, NIH Aging, Injury, Musculoskeletal, Rheumatologic, and Dermatologic disorders (AIMR) Study Section
- Reviewer, NIH BBBT (Bioengineering, Biodata, and Biomodeling Technologies)-02 2023/11 Bioengineering Review Panel
- 2021/6 Reviewer, NIH GCAT (Genomics, Computational Biology and Technology) Study Section
- 2023-2024 Member, Competitive Medical Research Fund (CMRF) Review Committee, University of Pittsburgh
 - 2022 Grant Reviewer, Department of Psychiatry, University of Pittsburgh
 - 2021 Grant Reviewer, CTSI Quantitative Methodologies Pilot Program (QuMP), University of Pittsburgh

Journal Editorship

- 2023-Present Editorial Board Member, BMC Bioinformatics
 - 2020-2022 Guest Associate Editor, Statistical Genetics and Methodology, Frontiers in Genetics
- 2022-Present Review Editor, Frontiers in Genetics & Frontiers in Psychiatry

Manuscript Review

2014-Present Journal of the American Statistical Association, Biometrics, Biostatistics, Annals of Applied Statistics, Nature Communications, Genome Research, Genome Biology, Bioinformatics, American Journal of Human Genetics, PLOS Genetics, PLOS One, PLOS Computational Biology, Psychological Methods, Statistics and Computing, American Journal of Epidemiology, Briefings in Bioinformatics, Genetic Epidemiology, Human Genetics and Genomics Advances, Science Advances, Statistics in Biosciences, Annals of Surgical Oncology, Biometrical Journal, F1000Research, Frontiers in Genetics, Frontiers in Psychiatry, Neurobiology of Aging, PeerJ, Psychological Medicine, Theranostics, Frontiers in Aging Neuroscience, Schizophrenia Bulletin, Communications Biology, Health Services and Outcomes Research Methodology, Journal of Biopharmaceutical Statistics, BMC Bioinformatics, npj Systems Biology and Applications, NAR Genomics and Bioinformatics, Genomics, Proteomics & Bioinformatics, Machine Learning, Canadian Journal of Statistics

International Organizations

- 2023-2024 Co-Chair, Local Organizing Committee, STATGEN 2024: Conference on Statistics in Genomics and Genetics, Section on Statistics in Genomics and Genetics, American Statistical Association
 - 2024 Invited Session Organizer, IBC (International Biometric Society)
 - 2023 Invited Session Organizer, ENAR (Eastern North American Region of The International Biometric Society) Spring Meeting
 - 2023 Invited Session Organizer, WNAR (Western North American Region of The International Biometric Society)
- 2017, 2023 Invited Session Organizer, ICSA Applied Statistics Symposium
- 2021, 2022 Invited Session Organizer, JSM (Joint Statistical Meetings)

Department Committees

- 2022-Present Member, PhD Admissions Committee
- 2023-Present Chair, Master Comprehensive Exam Committee
- 2023-Present Member, Master Student Monitoring Committee

2022	Member, Search Committee for Tenure Stream Faculty
2021-2022	Member, Master Comprehensive Exam Committee
2020-2022	Organizer, Biostatistics Research Day
2020	Member, Biostatistics Teaching Task Force
	Other Services
2021-Present	Member, Scholarly Oversight Committee, Pitt Department of Pediatrics
2023	Judge, ENAR Poster Session
2021	Judge, Public Health Dean's Day Poster Session, University of Pittsburgh
2021	Judge, ASA Pittsburgh Chapter Banquet Poster Session

Last updated on April 19, 2024.