Yuehua Cui

CONTACT INFORMATION

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EDUCATION

M. Stat. Statistics (2002) Department of Statistics, University of Florida, Gainesville, FL

M.S. Plant Biology (1998) College of Biological Science, China Agricultural University

B.S. Agronomy (1995) Department of Agronomy, Shanxi Agricultural University

ACADEMIC POSITIONS

Graduate Director	2019 – pres, Department of Statistics & Probability, Michigan State University
Professor	2015 – pres, Department of Statistics & Probability, Michigan State University
Associate Professor	2010 – 2015, Department of Statistics & Probability, Michigan State University
Assistant Professor	2005 – 2010, Department of Statistics & Probability, Michigan State University
Research Scientist	1998 – 2000, Institute of Botany, Chinese Academy of Sciences

Other MSU Affiliations:

Professor (2018 – 2020), College of Nursing, Michigan State University

Faculty Member of MSU Genetics and Genome Sciences Program

Faculty Member of MSU Quantitative Biology Initiative (QBI) Program

Faculty Member of MSU Ecology, Evolution and Behavior (EEB) Program

Faculty Member of MSU Center for Research on Autism, Intellectual and other Neurodevelopmental Disabilities (C-RAIND) Program

RESEARCH INTERESTS

- Statistical genetics/genomics: gene-gene and gene-environment interactions, multi-omics data integration, causal mediation analysis, Mendelian randomization, scRNA-seq and spatial transcriptomics
- Applied functional/longitudinal data analysis, semi- and non-parametric models, high-dimensional data analysis, statistical machine learning
- Statistical applications in public health and biological sciences

ACADEMIC AWARDS AND HONORS

- 2022 Elected Fellow, American Statistical Association (ASA)
- 2010 Elected member, International Statistics Institute (ISI)
- 2008 Junior Faculty Meritorious Research Award, Sigma Xi Scientific Research Society, Michigan State University Chapter
- 2008- Sigma Xi full membership, the Sigma Xi Scientific Research Society
- 2007 William L. Harkness Instructional Innovation Award, Department of Statistics & Probability, Michigan State University
- 2003 Mu Sigma Rho, Statistics Honor Society
- 2002-2004 Superfund Graduate Fellow, Superfund Basic Research Program, Center for Environmental & Human Toxicology, University of Florida

EDITORIAL SERVICES

Senior Editorial Board: BMC Genomic Data (2021-)

Associate Editor:

- Computational and Structural Biotechnology Journal (2023-)
- Frontiers in Systems Biology Integrative Genetics and Genomics (2021-)
- Statistics and Probability Letters (2015-)
- BMC Genetics (2010-2016)

Deputy Section Editor: BMC Genetics, Section on Statistical and Computational Genetics (2016-2020)

Editorial Board Member:

- Mathematics (section board member) (2023-)
- Journal of Computational Systems Biology (2014-)
- Computational and Structural Biotechnology Journal (2021-2023)
- The Scientific World Journal, Computational Biology Domain (2011-2017)
- Recent Patents on Biotechnology (2010-2014)

Review Editor:

- Frontiers in Genetics Statistical Genetics and Methodology (2011-)
- Frontiers in Bioinformatics Network Bioinformatics (2020-)

Guest Editor: Recent Patents on Biotechnology (2011)

Guest Associate Editor

- Special issue on "Design and analysis of genome-wide association and next generation sequencing data in cancer pharmacogenomics", *Frontier in Genetics - Statistical Genetics and Methodology* (2014-2015)

- Special issue on "The Development and Application of Multi-Omics Integration Approaches to Dissecting Complex Traits in Plants", *Frontier in Genetics – Plant Genomics* (2021)

- Special Issue on "Network-Based Statistical Methods and Applications in Transcriptome-Wide Association Studies (TWAS)", *Frontier in Genetics – Computational Genomics* (2021)

-Special Issue on "Algorithms for Single Cell Data Analyses", *Frontiers in Systems Biology – Integrative Genetics and Genomics* (2022)

FUNDED RESEARCH GRANTS

- 2025-2029 (Co-I, with D. Vendramini (PI)) "Endoplasmic Reticulum Stress in the activation and pro-tumor functions of fibroblasts in pancreatic cancer", NIH/NCI R01CA295509 Total Award Amount: \$2,709,577 (5%) Total Award Period Covered: 01/01/2025-12/31/2029
- 2024-2027 (Co-I, with L. Wang (PI) and C. Hong). "Genetic polymorphisms as predisposed risk for apparent treatment-resistant hypertension in CKD", AHA 24TPA1288424 Total Award Amount: \$300,000 (summer effort + student RA support) Total Award Period Covered: 07/01/2024-06/30/2027
- 2024-2025 (Co-PI, with Knickmeyer R and Sadler R), "Understanding gene-environment interactions in neurodevelopment using a global, multicohort study", MSU Tetrad Initiative Award Total Award Amount: \$30,000 (student RA support) Total Award Period Covered: 09/01/2024-12/31/2025
- 2023-2025 (PI) "Statistical analysis of pancreatic cancer spatial transcriptomics data", Henry Ford Health System Total Award Amount: \$76,616 (student RA support) Total Award Period Covered: 05/15/2023-05/15/2025

2023-2025	 (Co-I, with B. Chen (PI)) "AI center for drug discovery (AID)", MSU Strategic Partnership Grants, MSU Research Foundation Total Award Amount: \$240,000 (student RA support) Total Award Period Covered: 09/01/2023-08/31/2025
2024-2024	(Co-I, with B. Chen (PI)) "Interrogating the phenotype and function of γδT cells in pancreatic cancer liver metastasis", MSU+Henry Ford Health Integrative Grant Total Award Amount: \$21,846 (student RA support) Total Award Period Covered: 01/01/2024-12/31/2024
2018-2024	(Co-PI, With J. Wang (PI) and E. Andrechek) "Statistical modeling of long-range chromatin interactions on gene regulation and underlying molecular mechanisms", NIH R01GM131398. Total Award Amount: \$1,300,004 Total Award Period Covered: 09/01/2018-05/31/2024
2019-2022	(Co-I, with H-S Wu), "Effects of bright light on co-occurring cancer-related symptoms in breast cancer survivors: A personalized intervention". National Inst. of Nursing Research, NR016828. Total Award Amount: \$323,837.55 (10% effort) Total Award Period covered: 09/01/2019-08/31/2022
2019-2022	(PI, with P. Zhong, A. Burt and K. Klump) "Novel methods for longitudinal study of synergistic gene-environment interactions in complex diseases". NIH R21HG010073. Total Award Amount: \$422,646.00 Total Award Period Covered: 05/01/2019-04/30/2022
2012-2016	(PI, with P-S Zhong), "Statistical inference for gene regulation with genetical genomic data", NSF (<u>DMS-1209112</u>), \$208,688
2012-2017	(Co-PI, with CR Buell, J Jiang, D Douches and RE Veilleux) "Unraveling the Heterozygosity, Allelic Composition, and Copy Number Variation of Potato", NSF (<u>IOS-1237969</u>). Total Award Amount: \$5,774,862.00 (My portion: \$567,256.00) Total Award Period Covered: 10/01/2012-09/30/2017
2011-2017	(Co-PI, with N Jiang, CR Buell, J Jiang), "The Impact of Pack-MULEs on Plant Genome Evolution and Mechanisms of Sequence Acquisition", NSF (<u>MCB-1121650</u>), \$1,490,000 (my portion: \$144,011) Total Award Period Covered: 09/01/2011-08/31/2017
2009-2017	(Major participant), "The dynamic response of plants to a changing environment", International Research Training Grant, Deutsche Forschungs Gemeinschaft (DFG) of Germany, Partnership between the University of Dusseldorf and Michigan State University.
2013-2015	(Co-PI, with R Freed and D Douches) "SNP Marker Identification for Pre-harvest Sprouting Resistance in Wheat". Michigan GREEN project, \$69,482 (no personal effort, student support).
2010-2011	(Statistician, with D. DellaPenna), "Advancing Drug Development from Medicinal Plants using Transcriptomics and Metabolomics", NIH, \$1,451,717 (10% effort and one ½ time RA)
2007-2011	(sole PI), "Statistical methods for mapping imprinted genes underlying complex traits", National Science Foundation, (<u>DMS-0707031</u>), \$116,817.
2010-2010	(sole PI), "Statistical methods to identify genetic conflicts associated with pregnancy complications", NIH/NICHD subcontract, \$55,932.
2006-2007	(Co-PI, with Wenjiang Fu), "Genetic association study of complications of pregnant women", NIH subcontract, \$492,968 (40% effort).

2006-2008 (sole PI), "Dissecting genomic imprinting and maternal effect underlying triploid endosperm development through genetic mapping", MSU Intramural Research Grants Program (06-IRGP-789), \$38,062.

PUBLICATIONS

Research Highlight: Our work on Gene-centric genetic association study (Cui et al. 2008, *Genetics*) was highlighted in *Nature Reviews Genetics* Vol. 9 No. 6 (2008).

Peer-referred Journal Articles (*senior corresponding author)

- 1. Su, H., Y. Wu, B. Chen, **Y.H. Cui***. (2025) STANCE: a unified statistical model to detect cell-type-specific spatially variable genes in spatial transcriptomics. *Nature Communications* (accepted)
- 2. Zhang, J., X. Liu, H. Wang and **Y.H. Cui***. (2025) Functional varying-index coefficient model for dynamic gene-environment interactions. *Statistics in Biosciences* https://doi.org/10.1007/s12561-024-09472-3.
- 3. Guan, S., X. Liu and **Y.H. Cui***. (2025) Variable Selection for Generalized Single-index Varying-coefficient Models with Applications to Synergistic Gx E Interactions. *Mathematics* 13 (3): 1-23.
- 4. Zhang, J., H. Wang and **Y.H. Cui***. (2025) Generalized functional varying-index coefficient model for dynamic synergistic gene-environment interactions. *PloS One* 20 (1), e0318103.
- Loveless, I., S. Kemp, K. Hartway, J. Mitchell, Y. Wu, S. Zwernik, D. Salas-Escabillas, S. Brender, M. George, Y. Makinwa, T. Stockdale, K. Gartrelle, R. Reddy, D. Long, A. Wombwell, J. Clark, A. Levin, D. Kwon, L. Huang, R. Francescone, D. Barbosa Vendramini-Costa, B. Stanger, A. Alessio, A. Waters, Y.H. Cui, E. Fertig, L. Kagohara, B. Theisen, H. Crawford, N. Steele. (2025) Human pancreatic cancer single cell atlas reveals association of CXCL10+ fibroblasts and basal subtype tumor cells. *Clinical Cancer Research* (published online) doi: 10.1158/1078-0432.CCR-24-2183.
- Liyanage, J. S. S., J.H. Estepp, K. Srivastava, S. R. Rashkin, J. S. Hankins, C. M. Takemoto, Y. Li, Y.H. Cui, M. Mori, M. J. Weiss, and G. Kang. (2024) A Novel Mendelian Randomization Approach for Count Outcome Robust to Correlated and Uncorrelated Pleiotropic Effects. *Genetic Epidemiology* 49: e22602.
- Jia, C., T. Wang, G. Liu, Y. Tian, D. Cui, Z. Xu, R. Fang, H. Yu, Y. Zhang, Y.H. Cui* and H. Cao*. (2024) Multi-omics data integration for subtype identification of renal cell carcinoma: a metagene-based similarity network fusion approach. *Briefings in Bioinformatics* 25: bbae606.
- Liu, Y., B. Thyreau, Y.H. Cui, Y. Zhang, Y. Tatewaki and Y. Taki. (2024) Influence of Intergenerational Social Mobility On Brain Structure and Global Cognition: Findings From the Whitehall II Study Across 20 Years. Age and Aging 53(10): afae221.
- 9. Tan, X., X. Zhang, **Y.H. Cui**, X. Liu. (2024) Uncertainty quantification in high-dimensional linear models incorporating graphical structures with applications to gene set analysis. *Bioinformatics*, 40(9): btae541.
- 10. Yang, H., X. Wang, Z. Zhang, F. Chen, H. Cao, L. Yan, X. Gao, H. Dong, **Y.H. Cui***. (2024) A highdimensional omnibus test for set-based association analysis. *Briefings in Bioinformatics* 25: bbae456.
- 11. Cao, D.Q., S.C. Lei, H. Liu, Y. Jin, Y.F. Wu, **Y.H. Cui**, R. Wu. (2024) Reverse Solute Diffusion Enhances Sludge Dewatering in Dead-End Forward Osmosis. *Membranes* 14(9): 196.
- 12. Shi, R., L. Wang, S. Burgess and **Y.H. Cui***. (2024) MR-SPLIT: a novel method to address selection and weak instrument bias in one-sample Mendelian randomization studies. *PloS Genetics* 20: e1011391.
- 13. Das Adhikari, S., J. Yang, **Y.H. Cui**^{*}, and J. Wang^{*}. (2024) BayesKAT: Bayesian optimal kernel-based test for genetic association studies reveals joint genetic effects in complex diseases. *Briefings in Bioinformatics* 25 (3): bbae182
- Cao, H, Z. Li, H. Yang, R. Fang, Y. Zhang, B. Wang and Y.H. Cui*. (2024) Multi-omics data integration identified novel cancer subtypes with weighted multi-kernel learning. *British Journal of Cancer* 130: 1001-1012.
- 15. He, T., P-S. Zhong, **Y.H. Cui**, and V. Mandrekar. (2023) Unified tests for nonparametric functions in RKHS with kernel selection and regularization. *Statistica Sinica* 33: 919-944.
- 16. Guan, S.[#], M. Zhao[#] and **Y.H. Cui**^{*}. (2023) Variable selection in varying multi-index coefficient models for synergistic gene-environment interactions. *Electronic Journal of Statistics* 17(1): 823-857.

- 17. Wei, Y-F., X. Zhao, L. Li, H. Yang, H. Cao* and **Y.H. Cui***. (2023) Cancer subtyping with multi-omics data via multi-kernel learning. *Briefings in Bioinformatics* 24(1), bbac488.
- Liu, Y., Liu, Y., Cui, Y., Thyreau, B., Zhang, Y., Taki, Y., Chen, H. (2023), Intergenerational Social Mobility, Brain Structure, and Cognitive Trajectory: Findings from the Whitehall II Study. *Alzheimer's Dement.*, 19: e071978.
- Liyanage, J.S.S., J.H. Estepp, K. Srivastava, S.R. Rashkin, V.A. Sheehan, J.S. Hankins, C.M. Takemoto, Y. Li, Y.H. Cui, M. Mori, S. Burgess, M.R. DeBaun and G. Kang. (2022) A Versatile and Efficient Novel Approach for Mendelian Randomization Analysis with Application to Assess the Causal Effect of Fetal Hemoglobin on Anemia in Sickle Cell Anemia. *Mathematics*, 10(20): 3743.
- Wang, J[†], Y. Miao[†], L. Li, Y. Wu, Y. Ren^{*}, Y.H. Cui^{*} and H Cao^{*}. (2022) Multi-omics data integration for hepatocellular carcinoma subtype identification with multi-kernel learning. *Frontiers in Genetics-Computational Genomics* 13: 962870. ([†]contributed equally to this work)
- Li, L-M., G. Shi, Y. Wei, H. Yang, Z. Li, R. Fang, H. Cao* and Y.H. Cui*. (2022) Multi-omics data integration for subtype identification of Chinese lower-grade gliomas: A joint similarity network fusion approach. *Computational and Structural Biotechnology Journal* 20: 3482-3492.
- Wang, H.L., J.Y. Zhang, K.L. Klump, S.A. Burt and Y.H. Cui*. (2022) Multivariate partial linear varyingcoefficients model for gene-environment interactions with multiple longitudinal traits. *Statistics in Medicine* 41: 3643-3660.
- 23. Qu, J. and **Y.H. Cui***. (2022) Gene set association analysis with graph embedded kernel association test. *Bioinformatics* 38(6): s1560-1567.
- 24. Zheng, X., Y.H. Cui, Y. Xue, L. Shi, Y. G, F. Dong, C. Zhang. (2022) Adverse childhood experiences in depression and the mediating role of multimorbidity in mid-late life: A nationwide longitudinal study. *Journal of Affective Disorders* 301:217-224.
- 25. Shen, X., Y. Wen, **Y.H. Cui** and Q. Lu. (2022) A conditional autoregressive model for genetic association analysis accounting for genetic heterogeneity. *Statistics in Medicine* 41: 517-542.
- Zhao, M., Y. Gao and Y.H. Cui*. (2022) Variable selection for longitudinal varying coefficient errors-invariables models. *Communications in Statistics: Theory and Method* 51(11): 3713-3738.
- Liu, Y., Y. Gao, R. Fang, H. Cao, J. Sa, J. Wang, H. Liu, T. Wang* and Y.H. Cui*. (2021) Identifying complex gene-gene interactions: a mixed kernel omnibus testing approach. *Briefings in Bioinformatics* 22(6): bbab305.
- 28. Fang, R., H. Yang, Y. Gao, H. Cao, E.L. Goode and **Y.H. Cui***. (2021) Gene-based mediation analysis in epigenetic studies. *Briefings in Bioinformatics* 22(3): bbaa113.
- 29. Yang, H., X. Li, H. Cao, **Y.H. Cui**, Y. Luo, J. Liu and Y. Zhang. (2021) Using machine learning methods to predict hepatic encephalopathy in cirrhotic patients with unbalanced data. *Computer Methods and Programs in Biomedicine* 211: 106420.
- 30. Zhou, L., Z. Guo, B. Wang, Y. Wu, Z. Li, H. Yao, R. Fang, H. Yang, H. Cao* and **Y.H. Cui***. (2021) Risk prediction in Heart Failure with Preserved Ejection Fraction patients with gene expression data using machine learning. *Frontier in Genetics: Statistical Genetics and Methodology* 12: 412.
- Wu, Y., H. Wang, Z. Li, J. Cheng, R. Fang, H. Cao* and Y.H. Cui*. (2021) Subtypes identification on heart failure with preserved ejection fraction via network enhancement fusion using multi-omics data. *Computational and Structural Biotechnology Journal* 19: 1567-1578.
- 32. Liu, X., P-S. Zhong and **Y.H. Cui***. (2020) Joint test of parametric and nonparametric effects in partial linear models for gene-environment interaction. *Statistica Sinica* 30: 325-346.
- 33. Sun, H., Y.H. Cui, H, Wang, H. Liu and T. Wang. (2020) Comparison of methods for the detection of outliers and associated biomarkers in mislabeled omics data. *BMC Bioinformatics* 21: 1-23.
- 34. Wang, H., Y. Wu, R. Fang, J. Sa, Z. Li, H. Cao and **Y.H. Cui***. (2020) Time-varying gene network analysis of human prefrontal cortex development. *Frontiers in Genetics: Computational Genomics* 11: 1409.
- 35. Deng, Y., T. He, R. Fang, S. Li, H. Cao, and **Y.H. Cui***. (2020) Genome-wide gene-based multi-trait analysis. *Frontiers in Genetics: Statistical Genetics and Methodology* 11: 437.

- Kao, T-S. A., J. Ling, M. Dalaly, L.B., Robbins, and Y.H. Cui. (2020) Parent–Child Dyad's Collective Family Efficacy and Risky Adolescent Health Behaviors. *Nursing Research* 69(6): 455-465.
- Zhang, H., W. Bi, Y.H. Cui, H. Chen, J. Chen, Y. Zhao, and G. Kang. (2020) Extreme-value sampling design is cost-beneficial only with a valid statistical approach for exposure–secondary outcome association analyses. *Statistical Methods in Medical Research* 29: 466-480.
- Yang, H., H. Cao, T. He, T. Wang and Y.H. Cui*. (2020) Multi-level heterogeneous omics data integration with kernel fusion. *Briefings in Bioinformatics* 21: 156-170.
- 39. Sun, H.W., Y.H. Cui, Q. Gao, and T. Wang. (2020) Trimmed LASSO Regression Estimator for Binary Response. *Statistics and Probability Letters* 159: 108679.
- 40. Jing, L.W., **Y.H. Cui**, Q. Lu, and H. Yu. (2020) Multiplier method estimates of the population of men who have sex with men: the effect of privacy protection. *Journal of Public Health* 42: 429-434.
- 41. Gao, B., X. Liu, H. Li and **Y.H. Cui***. (2019) Integrative analysis of genetical genomics data incorporating network structures. *Biometrics* 75(4): 1063-1075.
- 42. Gao, Y., H. Yang, R. Fang, Y. Zhang, E. L. Goode and **Y.H. Cui***. (2019) Testing mediation effects in highdimensional epigenetic studies. *Frontiers in Genetics: Epigenomics and Epigenetics* 10: 1195.
- 43. He, T., S. Li, P-S. Zhong* and Y.H. Cui*. (2019) An optimal kernel-based method for gene set association analysis. *Genetic Epidemiology* 43: 137-149.
- Kang, Y., I. Torres-Jerez, Z. An, V. Greve, D. Huhman, N. Krom, Y.H. Cui and M. Udvardi. (2019) Genome-wide association analysis of salinity responsive traits in Medicago truncatula. *Plant, Cell & Environment*.13508 42: 1513–1531.
- 45. Wu, C., P-S. Zhong and **Y.H. Cui***. (2018) Additive varying-coefficient model for nonlinear geneenvironment interactions. *Statistical Applications in Genetics and Molecular Biology* 17(2).
- 46. Wang, H.L., P-S. Zhong and Y.H. Cui. (2018) Empirical likelihood ratio tests for coefficients in high dimensional heteroscedastic linear models. *Statistica Sinica* 28: 2409-2433.
- 47. Wu, C., Y. Jiang, J. Ren, Y.H. Cui and S. Ma. (2018) Dissecting gene-environment interactions: a penalized robust approach accounting for hierarchical structures. *Statistics in Medicine* 37(3): 437-456.
- 48. Wang, H.L., P-S. Zhong, Y.H. Cui and Y. Li. (2018) Unified empirical likelihood ratio tests for functional concurrent linear models and the phase transition from sparse to dense functional data. *Journal of the Royal Statistical Society B*. 80(2): 343-364.
- 49. Jing, L., Q. Lu, **Y.H. Cui**, H. Yu and T. Wang. (2018) Combining the randomized response technique and the network scale-up method to estimate the female sex worker population size: an exploratory study. *Public Health* 160:81-86.
- Zhao, D., J.P. Hamilton, B. Vaillancourt, W. Zhang, G.C. Eizenga, Y.H. Cui, J. Jiang, C.R. Buell, and N. Jiang. (2018) The unique epigenetic features of Pack-MULEs and their impact on chromosomal base composition and expression spectrum. *Nucleic Acids Research* 46: 2380-2397.
- Wu, D., H. Yang, S.J. Winham, Y. Natanzon, D.C. Koestler, T. Luo, B.L. Fridley, E.L. Goode, Y.B. Zhang and Y.H. Cui*. (2018) Mediation analysis of alcohol consumption, DNA methylation, and epithelial ovarian cancer. *Journal of Human Genetics* 63: 339-348.
- 52. Yang, H., S. Li, H. Cao, C. Zhang and **Y.H. Cui***. (2017) Predicting disease trait with genomic data: A composite kernel approach. *Briefings in Bioinformatics* 18(4):591–601.
- Cao, H.Y., Z. Li, H. Yang, Y.H. Cui* and Y. Zhang*. (2017) Longitudinal next-generation sequencing data analysis for rare variants detection with penalized quadratic inference function. *Scientific Reports* 7(1): 650.
- 54. Zhao, J., S. Li, L. Wang, L. Jiang, R. Yang* and Y.H. Cui*. (2017) Genome-wide random regression analysis for parent-of-origin effects of body composition allometries in mouse. *Scientific Reports* 7: 45191.
- 55. Liu, X., B. Gao and **Y.H. Cui***. (2017) Generalized partial linear varying-index coefficient model for geneenvironment interactions. *Statistical Applications in Genetics and Molecular Biology* 16(1): 59-74.
- 56. Cao, HY, P. Zeng, Z. Li, **Y.H. Cui** and Y.B. Zhang. (2017) The application of penalized generalized estimating equations in genetic association with longitudinal data. *Chinese Journal of Health Statistics* 34(4): 534-537.

- 57. Ren, Yan, J. Jia, J. Sa, L-X. Qiu, Y. Cui, Y-A. Zhang, H. Yang, G-F. Liu. (2017) Association between Nterminal proB-type natriuretic peptide and depressive symptoms in patients with acute myocardial infarction. *Chinese Medical Journal* 130(5): 542-548.
- Zhao, D., J. Hamilton, M. Hardigan, D. Yin, T. He, B. Vaillancourt, M. Reynoso, G.G. Pauluzzi, S. Funkhouser, Y. Cui, J. Bailey-Serres, J. Jiang, C.R. Buell and N. Jiang. (2017) Analysis of ribosome-associated mRNAs in rice reveals the importance of transcript size and GC content in translation. G3 7(1): 203-219.
- 59. Li, G.X. and **Y.H. Cui***. (2016) Assessing statistical significance in variance components linkage analysis: a theoretical justification. *Journal of Statistical Planning and Inference* 178: 70-83.
- 60. Liu, X., H. Wang and **Y.H. Cui***. (2016) Statistical identification of gene-gene interactions triggered by nonlinear environmental modulation. *Current Genomics* 17(5): 388-395.
- 61. Liu, X., Y.H. Cui* and R. Li. (2016) Partial linear varying multi-index coefficient model for integrative geneenvironment interactions. *Statistica Sinica* 26: 1037-1060.
- Li, Z., H.Y. Cao, Y.H. Cui*, Y.B. Zhang*. (2016) Extracting DNA Words Based on the Sequence Features: Non-uniform Distribution and Integrity. *Theoretical Biology and Medical Modelling* 13:2 (DOI 10.1186/s12976-016-0028-3).
- 63. Luo, T., X. Liu and **Y. Cui***. (2016) A genome-wide association analysis in four populations reveals strong genetic heterogeneity on birth weight. *Current Genomics* 17(5): 416-426.
- 64. Sa, J., X. Liu, T. He, G. Liu* and **Y.H. Cui***. (2016) A nonlinear model for gene-based gene-environment interaction. *International Journal of Molecular Sciences* 17(6): 882; doi:10.3390/ijms17060882.
- 65. Wu, C., X. Shi, **Y.H. Cui** and S.G. Ma. (2015) A penalized robust semi-parametric approach for geneenvironment interactions. *Statistics in Medicine* 34(30):4016-30.
- 66. Gao, B. and **Y.H. Cui**^{*}. (2015) Learning directed acyclic graphical structures with genetical genomics data. *Bioinformatics* 31(24): 3953-60.
- 67. Li, G., **Y.H. Cui** and H.Y. Zhao (2015) An Empirical Bayes risk prediction model using multiple traits for sequencing data. *Statistical Applications in Genetics and Molecular Biology* 14(6): 551-73.
- Bi, W., G. Kang, Y. Zhao, Y.H. Cui, S. Yan, Y. Li, C. Cheng, S.B. Pounds, M.J. Borowitz, M.V. Relling, J.J. Yang, C-H. Pui, S.P. Hunger, C.M. Hartford, W. Leung and J-F. Zhang. (2015) SVSI: fast and powerful set-valued system identification approach to identifying rare variants in sequencing studies for ordered categorical traits. *Annals of Human Genetics* 79(4): 294-309.
- 69. Cao, H.Y., X. Wei, X. Guo, C. Song, Y. Luo, **Y. Cui**, X. Hu, and Y. Zhang. (2015) Screening high-risk clusters for developing birth defects in mothers in Shanxi Province, China: application of latent class cluster analysis. *BMC Pregnancy and Childbirth* 15: 343.
- 70. Habing, G.G., S. Manning, C. Bolin, **Y.H. Cui**, J. Rudrik, S. Dietrich and J.B. Kaneene. (2015) Withinfarm changes in dairy farm associated Salmonella subtypes and comparison to human clinical isolates in Michigan, USA, 2000-2001 and 2009. *Applied and Environmental Microbiology* 81(17): 5724-35.
- 71. Manrique-Carpintero, N. C., J.J. Coombs, **Y.H. Cui**, R.E. Veilleux, C. R. Buell, and D. Douches. (2015) Genetic map and QTL analysis of agronomic traits in a diploid potato population using SNP markers. *Crop Science* 55(16): 2566-2579.
- 72. Ren Y, **Cui Y.H.**, Li X, Wang B, Na L, Shi J, Wang L, Qiu L, Zhang K, Liu G, Xu Y. (2015) A co-expression network analysis reveals IncRNA abnormalities in peripheral blood in early-onset schizophrenia. *Progress in Neuro-psychopharmacology & Biological Psychiatry* 63:1-5.
- 73. Wu, C., **Y.H. Cui** and S.G. Ma. (2014) Integrative analysis of gene-environment interactions under a multi-response partially linear varying coefficient model. *Statistics in Medicine* 33: 4988-4998.
- 74. He, T., P-S. Zhong and **Y.H. Cui***. (2014) Gene set association test identifies sex-specific pathways associated with type 2 diabetes. *Frontiers in Genetics: Statistical Genetics and Methodology* 5: 395.
- 75. Li*, S., **Y.H. Cui*** and R. Romero. (2014) Entropy-based selection for maternal-fetal genotype incompatibility with application to preterm pre-labor rupture of membranes. *BMC Genetics* 15: 66.
- 76. He, T., J. Sa, P-S. Zhong and **Y.H. Cui***. (2014) Statistical dissection of cyto-nuclear epistasis subject to genomic imprinting in line crosses. *PloS One* 9(3): e91702.
- 77. Wu, C., and **Y.H. Cui***. (2014) Boosting signals in gene-based association studies via efficient SNP selection. *Briefings in Bioinformatics* 15(2):279-291.

- 78. Dai, H.Y., T. Srivastava and **Y.H. Cui**. (2014) A modified generalized fisher method for combining probabilities from dependent tests. *Frontiers in Genetics: Evolution and Population Genetics* 5:23.
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- 123. Cui, Y.H. and R.L. Wu. (2005) Mapping genome-genome epistasis: a high-dimensional model. *Bioinformatics* 21(10): 2447-2455.
- 124. Sun, K.L., Y.H. Cui and B. Hauser. (2005) Changes in gene expression during ovule abortion in Arabidopsis. *Planta* 222: 632-642.
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- 126. Lu, Q., **Y.H. Cui** and R.L. Wu. (2004) A multilocus likelihood approach to joint modeling of linkage, parental diplotype and gene order in a full-sib family. *BMC Genetics* 5: 20.
- 127. Sun, K.L., M. Wang and **Y.H. Cui**. (2000) Study on the primary vascular system of the seedling of *Nelumbo Nucifera* Gaertn. *Acta Botanica Yunnanica*, 22(3): 286-290
- 128. **Cui, Y.H.**, M. Wang and K.L. Sun. (1999) Study on the heterogeneity of endosperm cell development in *Eucommia ulmoides* Oliv. *Journal of Agricultural Biotechnology* 7(1): 89-93.
- 129. Cui, Y.H., M. Wang and K.L. Sun. (1999) Morphology of gutta-containing cells in *Eucommia ulmoides* Oliv. *Chinese Bulletin of Botany* 16(4): 439-443.
- 130. Wang, M., Y.H. Cui and K.L. Sun. (1999) Study on the programmed cell death in the process of endosperm degeneration in *Eucommia ulmoides* Oliv. *Bulletin of Botanical Research* 19(4): 401-406.
- 131. Zhao, Y.J., W. Zhang, G. C. Li, T. Wang and **Y.H. Cui**. (1998) Genetic Studies on several morphological index of columnar apple trees. *Journal of China Agricultural University* Vol. 3, supplement: 96-98.

Review and Comment Papers

- 132. Cui, Y.H. (2025) GLMY homology theory meets idopNetwork: Dissecting soil microbiota resilience under forest thinning and climate change. Comment on "Topological change of soil microbiota networks for forest resilience under global warming" by Gong et al. *Physics of Life Reviews* 52: 44-45.
- 133. Yang, J., SD. Adhikari, H. Wang, B. Huang, W. Qi, **Y.H. Cui** and J. Wang. (2024) De novo prediction of functional effects of genetic variants from DNA sequences based on context-specific molecular information. *Front. Syst. Biol. Integrative Genetics and Genomics* 4: 1402664.
- 134. Das Adhikari, S., J. Yang, J. Wang*, and Y.H. Cui*. (2024) Recent advances in detecting spatially variable genes for spatial transcriptomics. *Computational and Structural Biotechnology Journal* 23: 883-891.
- 135. **Cui**, **Y.H.*** and H. Yang. (2017) Dissecting genomic imprinting and genetic conflict from a game theory perspective: Comment on: "Epigenetic game theory: How to compute the epigenetic control of maternal-to-zygotic transition" by Qian Wang et al. *Physics of Life Reviews* 20:161-163.
- 136. Wu, C., S. Li and **Y.H. Cui***. (2012) Genetic association studies: an information content perspective. *Current Genomics* 13(7): 566-573.
- 137. Wu, R.L., W. Hou, Y.H. Cui, H.Y.Li, T. Liu, S.Wu, C-X. Ma, and Y. Zeng. (2007) Modeling the genetic architecture of complex traits with molecular markers. *Recent Patents on Nanotechnology* 1: 41-49.

Conference Proceedings

- 138. Ma D., W. Bu, **Y.H. Cui**, Y. Xie, and X. Wu. (2018) Learning Collaborative Model for Visual Tracking. 24th International Conference on Pattern Recognition (ICPR), 2582-2587.
- 139. Ma D., W. Bu, Y. Xie, Y.H. Cui, and X. Wu. (2018) Segmentation-Guided Tracking with Prior Map Decision. 2018 24th International Conference on Pattern Recognition (ICPR), 2014-2019.

Invited Book Chapters

- 140. Liu, T., C. Tong, J.S. Wu, B. Zhang, Y.H. Cui, Y. Li, Y. Zeng and R.L. Wu. (2011) Haplotype Analysis of Complex Traits in Outcrossing Tree Species: Allele Discovery of Quantitative Trait Loci. In: *Genetics, Genomics and Breeding of Poplar* (C.P. Joshi, S.P. DiFazio, and C. Kole eds.) Science Publishers, Northants, UK.
- 141. Cui, Y.H.*, G.X. Li, S.Y. Li and R.L. Wu. (2010) Designs for Linkage Analysis and Association Studies of

Complex Diseases. In: *Statistical Methods in Molecular Biology* (Eds.: H. Bang, X.K. Zhou, H.L. Van Epps and M. Mazumdar) Humana Press.

Book Review

- 142. **Cui, Y.H.** (2022) "Multivariate Data Integration Using R: Methods and Applications with the mixOmics Package" by Kim-Anh LeCao, Zoe Marie Welham. *Biometrics*.
- 143. **Cui, Y.H.** (2015) "Statistical methods for QTL mapping" by Z. Chen. *Journal of the American Statistical Association*.
- 144. **Cui, Y.H.** (2009) "Multivariate Statistics: Exercises and Solutions" by W. Hardle and Z. Hlavka. *Journal of the American Statistical Association* 104(488): 1722.

Papers under review/revision

- 1. Miao, R, D. Fan, H. Wu, J. Du, H. Wang, **Y. Cui**, T. Cheng, Q. Zhang and L Sun. Identification of DEADbox RNA Helicase Genes in Prunus mume Reveals the Role of PmDDX39 in Cold Stress Tolerance. *Horticultural Plant Journal* (under review)
- 2. Das Adhikari, S., N.G. Steele, B. Theisen, J. Wang, **Y.H. Cui***. SPACE: Spatially variable gene clustering adjusting for cell type effect for improved spatial domain detection. *Nucleic Acids Research* (under revision)
- 3. Liu, X., Y. Liu, X. Zhang, J. Huang, **Y.H. Cui** and X. Shi. Multi-omics data integration with multi-view learning via composed tensors. *Bioinformatics* (under review)
- Cao, H.*, T. Wang, Z. Xu, X. Zhao G. Liu, X. Yang, R. Fang, Y. Luo, P. Zeng, H. Yu, Y. Zhang, Y.H. Cui*. Interactive multi-kernel learning for integrating multi-omics data in cancer subtyping. *Briefings in Bioinformatics* (under review)
- 5. Liu, Y., Y. Tatewaki, C-J. Boraxbekk, B. Thyreau, **Y. Cui**, Y. Zhang, Y. Hidaka, N. Karalija, S. Yuki, M. Takebayashi, T. Ninomiya, Y. Taki and N. Karalija. Women at Risk: Women at risk: a comparative study on socioeconomic status, lifestyle, and brain cognition among older females in Japan and Sweden. (under review)

PROFESSIONAL ACTIVITIES

Panelist and Ad hoc Reviewer for Grant Proposals

- University of Sharjah Research Funding Department, 2022
- NSF Panelist, 2016, 2022
- Ad hoc NSF Reviewer, 2009, 2011, 2020
- Netherlands Genomics Initiative Reviewer, 2009
- National Natural Science Foundation of China Reviewer, 2018, 2019
- Intramural Funding Program Reviewer, the Office of the Vice President for Research and Graduate Studies, Michigan State University, 2009
- Michigan AgBio Research Program Reviewer, 2012
- UK Medical Research Council Reviewer, 2017
- Hong Kong Research Grants Council Reviewer, 2019

Book proposal reviewer for Prentice-Hall and Springer

Reviewer for Mathematical Reviews (2010-)

Journal referee services for the following journals (>300 papers)

- Statistics related journals
- 1. Annals of Applied Statistics
- 2. Biometrics
- 3. Biometrical Journal
- 4. Biostatistics

- 5. Communications in Statistics T&M
- 6. Computational Statistics and Data Analysis
- 7. Journal of American Statistical Association
- 8. Journal of Biopharmaceutical Statistics

- 9. Journal of Business & Economic Statistics
- 10. Journal of Royal Statistical Society B
- 11. Journal of Statistical Planning and Inference
- 12. Journal of Statistical Software
- 13. Scandinavian Journal of Statistics
- 14. Science China Mathematics
- 15. Stat

Statistical Genetics and Bioinformatics journals

- 21. American Journal of Human Genetics
- 22. Annals of Human Genetics
- 23. Bioinformatics
- 24. BMC Bioinformatics
- 25. BMC Genetics
- 26. BMC Genomics
- 27. BMC Medicine
- 28. Briefings in Bioinformatics
- 29. Communications in Information and Systems
- 30. Computational and Structural Biotechnology Journal
- 31. Current Genomics
- 32. Frontiers in Genetics, Statistical Genetics and Methodology
- 33. G3: Genes, genomics, Genetics
- 34. Genetica
- 35. Genetics
- 36. Genetic Epidemiology

Natural Science and Public Health journals

- 54. African Journal of Biotechnology
- 55. BioMed Research International
- 56. Biomolecules
- 57. BMC Plant Biology
- 58. British Journal of Psychiatric
- 59. Cellular and Molecular Life Sciences
- 60. Drug Discovery
- 61. Environmental Science and Pollution Research
- 62. Environmental International
- 63. Euphytica
- 64. European Journal of Public Health
- 65. Food Science and Technology
- 66. Frontiers in Cardiovascular Medicine
- 67. Frontiers in Publish Health
- 68. Gene
- 69. Health Expectations
- 70. Hypertension
- 71. International Journal of Obesity
- 72. Journal of Affective Disorder
- 73. Journal of Hazardous Materials

Organizing Committee in Conferences

74. Journal of Internal Medicine

- 16. Statistica Sinica
- 17. Statistical Applications in Genetics and Molecular Biology
- 18. Statistics and Its Interface
- 19. Statistics and Probability Letters
- 20. Statistics in Medicine
- 37. Genetics Research
- 38. Genomics
- 39. Heredity
- 40. Human Genetics
- 41. Human Genetics and Genomics Advances
- 42. IEEE/ACM Transactions on Computational Biology and Bioinformatics
- 43. Journal of Chemical Information and Modeling
- 44. Journal of Neuroscience methods
- 45. Journal of Theoretical Biology
- 46. Mathematical Biosciences and Engineering
- 47. Methods in Ecology and Evolution
- 48. Nature Computational Science
- 49. Nucleic Acids Research
- 50. PloS Computational Biology
- 51. PloS Genetics
- 52. Theoretical and Applied Genetics
- 53. Theoretical Biology and Medical Modeling
- 75. Journal of Public Health
- 76. Journal of Nutrition Education and Behavior
- 77. Journal of Sleep Research
- 78. Mammalian Genome
- 79. Marine Biotechnology
- 80. Molecular Genetics and Genomics
- 81. Molecular Plant
- 82. Nature Communications
- 83. Nutrition, Metabolism and Cardiovascular Diseases
- 84. Plant Cell
- 85. Plant Journal
- 86. Plant Physiology
- 87. Plants
- 88. PloS One
- 89. Proceedings of National Academy of Sciences
- 90. Public Health Nutrition
- 91. Science in China
- 92. Scientific Reports
- 93. Scientific World Journal
- 94. Translational Psychiatry
- DahShu 2024 Data Science Symposium, East Lansing, MI (May 16-18, 2024) (as co-chair)
- International Symposium in Graduate Researches on Data Sciences and Technology by Dokuz Eylul University, Türkiye (Dec. 2, 2023)

- 9th International Forum on Statistical Genetics and Genomics, Jinan, China (July 2-July 4, 2023) (as co-chair)
- The Data Science Section of the First International Symposium on Graduate Researches by Dokuz Eylul University, Türkiye (Dec. 2-3, 2022) (as a member of both the Organizing Committee and Scientific Committee)
- 5th International Workshop on Statistical Genetics and Genomics, Jinan, China (June 30-July 2, 2018) (as co-chair)
- 4th International Workshop on Statistical Genetics and Genomics, Taiyuan, China (June 21-23, 2017) (as chair)
- Program Committee member, 2017 ICSA Applied Statistics Symposium, Chicago, IL
- 3rd International Workshop on Statistical Genetics and Genomics, Taiyuan, China (June 20-23, 2016) (as chair)
- 2nd International Workshop on Statistical Genetics and Genomics, Taiyuan, China (June 23-24, 2015) (as chair)
- 1st International Workshop on Statistical Genetics and Genomics, Taiyuan, China (June 24-25, 2014) (as chair)
- 2nd International Conference on Epidemiology & Evolutionary Genetics, Beijing, China (Aug 18-19, 2014)
- International Year of Statistics: Statistics in Applications Forum, MSU (October 1, 2013)
- Statistical Genetics and Genomics Workshop, East Lansing, MI (May, 2008) (as chair)

Conference Invited Session Organizer/Chair:

- "Recent Statistical Advances in Complex Genetic and Genomic Data Analysis", 2025 ICSA-China Conference, Zhuhai, China (June 28-30, 2025)
- "Recent Methodological Advances in Single-Cell and Spatial Multi-omics", STATGEN 2025: Conference on Statistics in Genomics and Genetics, Minneapolis, MI (May 21-23, 2025)
- "Advancing Statistical and Machine Learning Models for Single-Cell and Spatial Omics", *MCBIOS* 2025: Data-Driven Discovery: Harnessing the power of AI to transform health, University of Utah, Salt Lake City (March 27-29, 2025)
- "Statistical and computational advances for complex biological data analysis", *2024 ICSA-China Conference*, Wuhan, China (June 28-30, 2024)
- "Recent Advances in Mendelian Randomization for Causal Inference", STATGEN 2024: Conference on Statistics in Genomics and Genetics, Pittsburg, PA (May 1-3, 2024)
- "Unraveling Intricate Molecular Pathways in Complex Diseases: Insights from Recent Statistical and Computational Advances", *MCBIOS 2024: From Codes to Cells to Care, Transforming Health Care with AI*, University of Emory, Atlanta (March 22-24, 2024)
- "Statistical advances in Mendelian randomization for causal inference", The 16th International Conference of the ERCIM WG on Computational and Methodological Statistics (CMStatistics 2023), Berlin, Germany (December 16-18, 2023)
- "Statistical innovations in scRNA-seq and spatial transcriptomics analysis", The 16th International Conference of the ERCIM WG on Computational and Methodological Statistics (CMStatistics 2023), Berlin, Germany (December 16-18, 2023)
- "Recent statistical advances for complex multi-omics data analysis", *The 2023 ICSA China Conference*, Chengdu, China (June 30-July 3, 2023)
- "Statistical advances for biomedical data with complex structures", *The 2023 ICSA Applied Symposium*, Ann Arbor, MI (June 14-16, 2023)
- "Recent advances in statistical methods for complex genetic/genomic data", 15th International Conference of the ERCIM WG on Computational and Methodological Statistics (CMStatistics 2022), London, UK (December 17-19, 2022)
- "Statistical advances for genetic data analysis", 2021 *ICSA Applied Statistics Symposium*, Washington DC (Sept 12-15, 2021)

- "Statistical methods for complex human genetic data", 2020 *ICSA Applied Statistics Symposium*, Houston, TX (Dec 17-20, 2020)
- "Recent advances in high-dimensional genomic data analysis", *The 3rd International Conference on Econometrics and Statistics (EcoSta 2019)*, Taichung, Taiwan (June 25-27, 2019)
- "Recent advances in statistical analysis of genetic/genomic data", 2018 ICSA China Statistics Conference, Qingdao, China (July 2-5, 2018)
- "Recent advances in statistical analysis of genetic/genomic data", 2017 ICSA Applied Statistics Symposium, Chicago, IL (June 25-28, 2017)
- "Novel statistical methods for genetic data analysis", 2017 *ICSA Applied Statistics Symposium*, Chicago, IL (June 25-28, 2017)
- "Recent advances in statistical analysis of genetic data", *2016 ICSA China Statistics Conference*, Qingdao, China (June 24-25, 2016)
- "Statistical advances for the detection of gene-environment interactions", *Joint Statistical Meeting*, Seattle, WA (August 8-13, 2015)
- "Statistical advances for genetic data analysis", *Joint 24th ICSA Applied Statistics Symposium and 13th Graybill Conference*, Fort Collins, CO (June 14-17, 2015)
- "Recent advances in statistical genetics: methods and applications", *The 3rd Biostatistics Symposium*, Chengdu, China (June 27-28, 2014)
- "Recent advances in statistical analysis of high-dimensional genetic data", *The 4th IMS-China International Conference on Statistics and Probability*, Chengdu, China (June 30-July 4, 2013)
- "Public Health Genomics", *BIT's 3rd World DNA and Genome Day-2012*, Xi'an, China (April 25-28, 2012)
- "New statistical challenges and advancements in eQTL mapping studies", XXVth International Biometric Conference, Florianópolis-SC, Brazil (December 5-10, 2010)
- "Statistical genetics: challenges, advances and prospects", *First Joint Biostatistics Symposium*, Beijing, China (July 1-3, 2010)
- "Recent advances and statistical challenges in genetical genomics analysis", *ICSA Applied Statistics Symposium*, Indianapolis, IN (June 20-23, 2010)

CONFERENCE AND DEPARTMENTAL SEMINAR PRESENTATIONS

Invited Plenary Talks

- 1. "From Association to Causation: Causal Inference with Mendelian Randomization in Biomedical Studies", 2024 International Congress of Basic Science, Beijing, China (July 14-26, 2024)
- 2. "Dealing with weak instruments in Mendelian randomization: an adaptive data splitting approach", *The 9th International Statistical Genetics and Genomics (ISGG) Forum*, Jinan, China (July 3, 2023)
- 3. "Genome-wide gene-based multi-trait analysis", *The 6th International Symposium on Statistical Genetics/Genomics*, Nanjing, China (June 29-July 1, 2019)
- 4. "Network-constrained genetical genomics analysis with nonlinear environmental modification", *The Third Annual Kliakhandler Conference on Bayesian Inference in Statistics and Statistical Genetics*, Michigan Technological University, Houghton, MI (Aug 16–20, 2017)
- 5. "Statistics in post genomic era: challenges and opportunities", *Marrakesh International conference on Probability and Statistics* 2016, Marrakech, Morocco (April 25-28, 2016)

Invited Conference and Workshop Presentations (since 2010, V=virtual)

- 1. "TBD", 2025 Joint Statistical Meeting, Nashville, TN (August 2-7, 2025)
- "TBD", The 11th International Forum on Statistical Genetics and Genomics, Xi'an, China (July 4-July 5, 2025)
- 3. "Spatial deconvolution and cell type-specific spatially variable gene detection in spatial transcriptomics", *2025 ICSA-China Conference*, Zhuhai, China (June 28-30, 2025)

- 4. "Cell-type-specefic spatially variable gene detection in spatial transcriptomics", 2025 WNAR/IMS Annual Meeting, Whistler, BC, Canada (June 15-18, 2025)
- "Spatial deconvolution and cell type-specific spatially variable gene detection in spatial transcriptomics", STATGEN 2025: Conference on Statistics in Genomics and Genetics, Minneapolis, MI (May 21-23, 2025)
- "A unified model for cell-type-specific spatially variable gene detection in spatial transcriptomics", MCBIOS 2025: Data-Driven Discovery: Harnessing the power of AI to transform health, University of Utah, Salt Lake City (March 27-29, 2025)
- 7. "Spatially informed trajectory-based differential expression analysis for spatial transcriptomics", 2024 *Joint Statistical Meeting*, Portland, OR (August 3-8, 2024)
- 8. "A unified statistical model to detect cell-type-specific spatially variable genes in spatial transcriptomics", *The 10th International Forum on Statistical Genetics and Genomics*, Wuhan, China (June 30-July 2, 2024)
- 9. "MR-SPLIT: a novel method to deal with weak instruments in one-sample Mendelian randomization studies", *ICSA China 2024*, Wuhan, China (June 28-30, 2024)
- 10. "Mendelian randomization for causal inference with MR-SPLIT", STATGEN 2024: Conference on Statistics in Genomics and Genetics, Pittsburg, PA (May 1-3, 2024)
- 11. "From association to causation: causal inference with Mendelian randomization in biomedical studies", *WDD 2024*, Nagoya, Japan (April 22-24, 2024).
- 12. "Causal inference with MR-SPLIT", *MICOBIO*, University of Emory, Atlanta (March 22-24, 2024)
- 13. "Addressing weak instruments in one sample MR analysis with MR-SPLIT", 16th International Conference of the ERCIM WG on Computational and Methodological Statistics (CMStatistics 2023), Berlin, GA (Dec. 18, 2023)
- 14. "Dealing with weak instruments in Mendelian randomization: an adaptive data splitting approach", *The 9th International Statistical Genetics/Genomics Forum*, Jinan, China (July 3-4, 2023)
- 15. "Gene set analysis with graph embedded kernel association test", 2023 ICSA applied statistics symposium, Ann Arbor, MI (June 11-14, 2023)
- 16. "FDR control in high-dimensional mediation analysis", 2023 New England Statistics Symposium, Boston, MA (June 3-6, 2023)
- "Causal inference with Mendelian randomization for longitudinal traits", Advances in Statistical and Computational Methods for Analysis of Biomedical, Genetic, and Omics Data (in celebrating Prof. Shili Lin's 60th birthday), The University of Texas at Dallas, Richardson, TX (March 17-19, 2023)
- (V)"Mendelian randomization for causal inference on longitudinal traits", Frontier in Mathematical Science at Tsinghua-Sanya International Mathematics Forum (TSIMF), Sanya, Hainan Province, China (Dec 22 to Dec 29, 2022)
- 19. (V)"Causal inference with Mendelian randomization for longitudinal traits", *15th International Conference of the ERCIM WG on Computational and Methodological Statistics* (CMStatistics 2022), London, UK (December 17-19, 2022)
- 20. (V)"Recent developments in causal inference", *International Symposium on Statistical Theory and Methods*, Anhui University of Finance and Economics, China (December 11, 2022)
- 21. (V)"Causal inference with Mendelian randomization for longitudinal traits", *The 8th International Symposium on Statistical Genetics/Genomics*, Taiyuan, China (Nov 26-27, 2022)
- 22. (V)"Variable selection in varying multi-index coefficient model for synergistic gene-environment interactions", *2022 ENAR Spring Meeting*, Houston, TX (March 27-30, 2022)
- 23. (V)"Multi-omics data integration with kernel learning", 2020 ICSA applied statistics symposium, Houston, TX (Dec 17-20, 2020)
- 24. (V)"Multivariate partial linear varying-coefficient model for G×E studies with longitudinal traits", *2020 JSM Meeting*, Philadelphia, PA (Aug. 1-6, 2020)
- 25. (V)"Nonlinear moderated mediation analysis with genetical genomics data", 2020 ENAR Meeting, Nashville, TN (March 22-25, 2020)

- 26. "High-dimensional mediation analysis in (epi)genomics studies", 2019 CoCoH Symposium on Computational Systems Biology, Grand Rapids, MI (Aug. 27, 2019)
- 27. "Nonlinear moderated mediation analysis with genetical genomics data", 2019 ICSA China Statistics Conference, Tianjin, China (July 1-4, 2019)
- "Functional partial linear varying-index coefficients model for gene-environment interaction", *The 3rd International Conference on Econometrics and Statistics (EcoSta 2019)*, Taichung, Taiwan (June 25-27, 2019)
- 29. "Omics data integration with kernel fusion", *San Francisco Bay area chapter of ASA*, San Francisco, CA (Oct. 26, 2018)
- 30. "Statistics in post-genomic era: challenges and opportunities", *Western Meeting of AMS*, San Francisco, CA (Oct. 27-28, 2018)
- 31. "Integrative genetical genomics analysis incorporating network structures", 2018 ICSA China Statistics Conference, Qingdao, China (July 2-5, 2018)
- 32. "Gene selection in big data era: biologically guided statistical treatment", *The 5th International Symposium on Statistical Genetics/Genomics*, Jinan, China (June 30-July 2, 2018)
- 33. "Embracing the complexity of genomic data: some statistical challenges and solutions", *Symposium on Big Data and the Future of Statistics*, Shanxi University of Finance and Economics, Taiyuan, China (Dec. 16-18, 2017)
- 34. "An optimal kernel-based method for gene set association analysis", 2017 *IMS-China International Conference on Statistics and probability*, Nanning, China (June 28-July 2, 2017)
- 35. "Gene selection with nonlinear instrumental regression incorporating network structures", *The 10th R Conference*, Taiyuan University of Technology, Taiyuan, China (June 24-25, 2017)
- 36. "An optimal kernel-based method for gene set association analysis", 4th International Workshop on *Statistical Genetics/Genomics*, Taiyuan, China (June 22-23, 2017)
- 37. "Integrative genetical genomics analysis incorporating network structures", Biostatistics workshop, *Statistical Inference for Biomedical Big Data*, Gainesville, FL (April 7-8, 2017)
- 38. "Integrative analysis of genetical genomics data incorporating network structures", *Joint Statistical Meeting*, Chicago, IL (July 30-August 4, 2016)
- 39. "Integrative genetical genomics analysis incorporating network structures", *The 4th International Symposium on Biostatistics*, Shanghai, China (July 2-3, 2016)
- 40. "Integrative genetical genomics analysis incorporating network structures", 2016 ICSA China Statistics Conference, Qingdao, China (June 24-25, 2016)
- 41. "Integrative analysis of gene-environment interactions", 3rd International Workshop on Statistical Genetics/Genomics, Taiyuan, China (June 21-22, 2016)
- 42. "Integrative genetical genomics analysis incorporating network structures", HDDA-VI: The 6th International Workshop on the Perspectives on High-Dimensional Data Analysis, Toronto, Canada (May 25-27, 2016)
- 43. "Integrative analysis of gene-environment interactions". *Joint Statistical Meeting*, Seattle, WA (August 7-13, 2015)
- 44. "Integrative genetical genomics analysis incorporating network structures", *ICSA-China Statistics Conference*, Shanghai, China (July 6-7, 2015)
- 45. "Integrative genetical genomics analysis incorporating network structures", *IMS-China International Conference on Statistics and Probability*, Kunming, Yunnan, China (July 1-4, 2015)
- 46. "Integrative genetical genomics analysis incorporating network structures", 2nd International Workshop on Statistical Genetics/Genomics, Taiyuan, China (June 22-24, 2015)
- 47. "Econometrics in the age of big data: the opportunities, challenges and prospects", *The 2nd Innovation and Global Industrial Development Forum: 2015 International Symposium on Innovation Creating New Competitive Advantage*, Wuhan, China (June 3, 2015)
- 48. "Statistical genetics: current status and perspectives", *Statistics in Applications Symposium* East Lansing, MI (October 2, 2014)
- 49. "Partial linear models for integrative gene-environment interactions", *3rd International Biostatistics Symposium*, Chengdu, China (June 27-28, 2014)
- *50.* "Challenges and strategies in gene-environment interactions", 1st International Workshop on Statistical Genetics/Genomics, Taiyuan, China (June 24-25, 2014)

- 51. "Novel strategies in high-dimensional genetic data analysis", *The 4th IMS-China International Conference on Statistics and Probability*, Chengdu, China (June 30-July 4, 2013)
- 52. "Linear or Non-linear: how genes respond to environmental stimuli", International *Symposium on Mapping the Genetic Regulation of Forest Ecosystems*, Beijing, China, (Oct. 6-13, 2012)
- 53. "Varying coefficient model for nonlinear gene-environment interaction", *The 8th Purdue Symposium on Statistics*, Lafayette, IN (June 20-24, 2012)
- 54. "Statistical identification of gene-gene and gene-environment interactions for complex disease", Epigenetics and the Elucidation of Complex Genetic Traits and Diseases section, *BIT's 3rd World DNA and Genome Day-2012*, Xi'an, China (April 25-29, 2012).
- 55. "Gene-centric gene-gene interaction", Statistical Genomics Workshop, *Plant and Animal Genome XX Conference*, San Diego, CA (January 15-17, 2012)
- 56. "eQTL mapping: the added value in complex trait dissection", *iGRAD-Plant Symposium/Retreat*, Henrich Heine University, Düsseldorf, Germany (September 12-17, 2011)
- 57. "Gene-centric gene-gene interaction in genome-wide association studies", *International workshop on statistical and computational genetics*, Beijing Forestry University, China, (August 4-5, 2011)
- 58. "Statistical dissection of genetic conflicts associated with pregnancy complications", (Session title 'Design and Analysis Issues for Assessing Complex Joint Maternal-Fetal Genetic Effects"), *Joint Statistical Meeting* (JSM), Vancouver, Canada (August 1-5, 2010)
- 59. "Gene-based genome-wide association studies", *Summer Workshop on Statistical & Computational Genetics*, Beijing Forest University, Beijing, China (July 19, 2010)
- 60. "eQTL mapping: the added value in complex trait dissection", *First Joint Biostatistics Symposium*, Beijing, China, (July 15-18, 2010).
- 61. "Enriching our knowledge in network regulation in eQTL mapping: a combined p-value approach", Statistical Genomics Workshop, *Plant and Animal Genome XVIII Conference*, San Diego, CA, (January 9-13, 2010).

Invited Department Colloquium/Seminar Talks (since 2010)

- 1. University of Texas at San Antonio, Department of Management Science and Statistics (4/18/2025)
- 2. University of Texas at Arlington, Department of Mathematics (4/4/2025)
- 3. (V)Texas Tech University, Department of Mathematics & Statistics (3/2025)
- 4. Zhejiang University, College of Agriculture and Biotechnology, China (7/4/2024)
- 5. Zhengzhou University, School of Public Health, China (7/7/2023)
- 6. Tsinghua University, Yau Mathematical Sciences Center, China (6/26/2023)
- 7. Shanxi Medical University, School of Public Health, China (6/20/2023)
- 8. (V)Indiana University and Purdue University at Indianapolis (IUPUI), Department of Mathematical Science (4/4/2023)
- 9. (V)Anhui University of Finance and Economics, School of Statistics, China (12/10/2022)
- 10. (V)Shanghai University of Finance and Economics, School of Statistics, China (12/7/2022)
- 11. (V)University of Illinois at Chicago, Department of Math, Stat, & Comp Sci (10/26/2022)
- 12. (V)University of Georgia, Department of Statistics (10/29/2020)
- 13. University of Florida, Department of Biostatistics (03/14/2019)
- 14. Michigan State University, Department of Statistics and Probability (1/22/2019)
- 15. University of Notre Dame, Department of Applied and Computational Mathematics and Statistics (11/16/2018)
- 16. Zhengzhou University (China), School of Public Health, China (05/28/2018)
- 17. Anhui University of Finance and Economics, School of Statistics, China (06/05/2018)
- 18. University of Minnesota, Division of Biostatistics (04/07/2017)
- 19. University of Notre Name, Department of Applied and Computational Mathematics and Statistics (9/26/2016)
- 20. Beijing Academy of Agriculture and Forestry Sciences (China) (07/13/2016)
- 21. Taiyuan University of Technology (China), School of Mathematics (06/03/2016)

- 22. Western Michigan University, Department of Statistics (02/12/2016)
- 23. Stony Brook University, Department of Applied Mathematics and Statistics (11/20/2015)
- 24. University of Alabama at Birmingham, Section on Statistical Genetics, Department of Biostatistics (09/25/2014)
- 25. Wuhan University (China), School of Mathematics and Statistics (05/26/2014)
- 26. Wright State University, Department of Mathematics and Statistics (04/11/2014)
- 27. Zhengzhou University China), School of Public Health (06/24/2013)
- 28. St. Jude Children's Research Hospital, Department of Biostatistics (01/08/2013)
- 29. Chinese Academy of Sciences, Beijing Institute of Botany (05/21/2012)
- 30. Wuhan University of Science and Technology (China), College of Science (05/07/2012).
- 31. Xi'an Jiaotong University (China), College of Mathematics and Statistics (04/28/2012)
- 32. Beijing Forestry University (China), College of Biological Sciences and Biotechnology (04/24/2012)
- 33. Yale University, Division of Biostatistics (11/8/2011)
- 34. Michigan State University, Department of Statistics and Probability (10/17/2011)
- 35. Shanxi Medical University (China), School of Public Health (6/15/2011)
- 36. Haerbin Medical University (China), College of Bioinformatics and Technology (6/10/2011)
- 37. Virginia Tech, Department of Statistics (09/30/2010)
- 38. University of Connecticut, Department of Statistics (04/19/2010)
- 39. UT Southwestern Medical Center, Department of Clinical Science (04/15/2010)
- 40. University of South Carolina, Department of Epidemiology and Biostatistics (04/07/2010)
- 41. Georgia State University, Department of Mathematics and Statistics (03/31/2010)
- 42. Saint Louis University, Division of Biostatistics (03/19/2010)
- 43. University of Nebraska Medical Center, Department of Biostatistics (03/17/2010)
- 44. Indiana University, Department of Medical and Molecular Genetics (03/09/2010)
- 45. Emory University, Department of Biostatistics and Bioinformatics (02/18/2010)
- 46. Marquette University, Department of Mathematics and Statistics, and Computer Science (02/14/2010)
- 47. MSU Computational Biology Group kick-off seminar (02/10/2010)
- 48. Michigan State University, Department of Statistics & Probability (02/02/2010)
- 49. Ohio State University, Department of Statistics (01/28/2010)
- 50. Indiana University and Purdue University at Indianapolis, Department of Mathematical Science (01/25/2010)

STUDENTS ADVISING

Current Ph.D. and MS students in statistics (as advisor/co-advisor)

1.	Ruxin Shi	Ph.D.	Ongoing
2.	Aoqi Xie	Ph.D.	Ongoing
3.	Yuesong Wu	Ph.D. (co-advisor, joint with Yuying Xie)	Ongoing
4.	Phuong Vo	Ph.D.	Ongoing
5.	Haohao Su	Ph.D.	Ongoing
6.	Juhee Lee	Ph.D. (co-advisor, joint with Chenxi Li from Epi/Biost)	Ongoing
7.	Zun Wang	Ph.D. (co-advisor, joint with Hao Zhang)	Ongoing

Past PhD students, their thesis title, and job placement

As advisor

1. Sikta Das Adhikari (joint with Jianrong Wang from CMSE), Dual Ph.D. in Statistics and Computational Mathematics, degree conferred in 08/2024

Thesis title: "Advanced Statistical and Computational Techniques for Genomic Data Analysis" 1st job: Statistician, FDA, Washington DC

 Jialin Qu, Ph.D. in Statistics, degree conferred in 08/2022
 Thesis title: "Causal Inference with Mendelian Randomization for Longitudinal Data" 1st job: Statistician, Moderna, Inc. Boston

- 3. Shunjie Guan, Ph.D. in Statistics, degree conferred in 08/2017
 - Thesis title: "Variable selection in varying multi-index coefficient models with applications to geneenvironmental interactions"
 - 1st job: Senior Research Statistician, AbbVie; currently manager at Pfizer
- 4. Jingyi Zhang, Ph.D. in Statistics, degree conferred in 08/2017
 - Thesis title: "Functional varying index coefficient model for gene-environment interactions with longitudinal data"
 - 1st job: Statistician, Wells Fargo; currently statistician at Amazon
- 5. Tao He, (joint with Ping-Shou Zhong) Dual Ph.D. in Statistics and Quantitative Biology, degree conferred in 08/2015
 - Thesis title: "Kernel-based Nonparametric Testing in High-dimensional Data with Applications to Gene Set Analysis".
 - 1st job: Assistant Professor, Department of Mathematics, San Francisco State University
- Honglang Wang, (joint with Ping-Shou Zhong) Dual Ph.D. in Statistics and Quantitative Biology, degree conferred in 08/2015

Thesis title: "Empirical likelihood based functional data analysis and high dimensional inference with applications in biology".

- 1st job: Assistant Professor, Department of Mathematical Sciences, IUPUI
- 7. Bin Gao, Ph.D. in Statistics, degree conferred in 08/2015
- Thesis title: "Graph estimation and network constrained regularization with application in genetical genomics"

1st job: Senior Statistician, Johnson & Johnson

- Cen Wu, Dual Ph.D. in Statistics and Quantitative Biology, degree conferred in 08/2013 Thesis title: "High-dimensional statistical methods for gene-environment interactions". 1st job: Postdoc, Department of Biostatistics, Yale University. Currently Associate Professor, Department of Statistics, Kansas State University
- Shaoyu Li, Dual Ph.D. in Statistics and Quantitative Biology, degree conferred in 08/2011 Thesis title: "Statistical issues and novel strategies for eQTL mapping".
 1st job: Assistant member, Department of Biostatistics, St. Jude Children's Research Hospital. Currently Associate Professor, Department of Mathematics and Statistics, University of North Carolina at Charlotte
- Gengxin Li, Dual Ph.D. in Statistics and Quantitative Biology, degree conferred in 08/2010 Thesis title: "Variance component models in mapping imprinted genes: statistical theory and applications".
 - 1st job: Postdoc, Department of Biostatistics, Yale University.
 - Currently Associate Professor, Dept. of Mathematics and Statistics, University of Michigan at Dearborn.

As co-chair

- Shan Zhang, Ph.D. in Statistics (joint with Qing Lu from Biostat/UF), degree conferred in 2021 Thesis title: "Neural networks models with applications to genetic studies" 1st job: Statistician, FDA.
- Jinghang Lin, Ph.D. in Statistics (joint with Qing Lu from Biostat/UF), degree conferred in 2021 Thesis title: "A neural network-based method with genetic data analysis of complex diseases".
 1st job: Postdoc, Department of Biostatistics, Yale University. Currently, Quantitative analytics specialist, Wells Fargo
- Xiaoxi Shen, Ph.D. in Statistics (joint with Q. Lu from Epi/Biostat), degree conferred in 08/2019 Thesis title: "Statistical Analysis for Network-Based Models with Application to Genetic Association Tests and Predictions".

1st job: Postdoc, Department of Biostatistics, University of Florida.

Currently Assistant Professor, Dept of Mathematics, Texas State University

 Daewoo Pak, Ph.D. in Statistics (joint with C. Li from Epi/Biostat), degree conferred in 08/2018 Thesis title: "Analysis of complex life-history data and variable selection in survival analysis under interval censoring".

1st job: Postdoc, MD Anderson Cancer Center.

Currently Associate Professor, Yonsei University, South Korea

15. Wei-Wen Hsu, Ph.D. in Statistics (joint with D. Todem from Epi/Biostat), degree conferred in 08/2011 Thesis title: "Tests of homogeneity in two-component mixture models".

1st job: Assistant Professor, Department of Statistics, Kansas State University

Currently Associate Professor, Division of Biostatistics and Bioinformatics in the Department of Environmental and Public Health Sciences, University of Cincinnati

16. Yanwei Zhang, Ph.D. in Statistics (joint with D. Todem from Epi/Biostat), degree conferred in 05/2008

Thesis title: "A hierarchical Bayesian approach to model spatially correlated binary data with applications to dental research".

1st job: Statistician, Pfizer; Currently Senior Director, Statistical & Quantitative Sciences, Takeda

Past postdocs and their current position

- 17. Hongyan Cao, Postdoctoral Fellow, 2019-2021 Currently Associate Professor, School of Public Health, Shanxi Medical University, China
- Haitao Yang, Postdoctoral Fellow, 2016-2017
 Currently Professor, School of Public Health, Hebei Medical University, China
- Xu Liu, Postdoctoral Fellow, 2013-2016
 Currently Associate Professor with tenure, School of Statistics and Management, Shanghai University of Finance and Economics, China.
- 20. Guolian Kang, Postdoctoral Fellow, 2006-2007 Currently Full Member, Department of Biostatistics, St. Jude Children's Research Hospital.

Past MS students and their 1st position

- 1. Hanyu Yang, M.Sc., degree conferenced in 5/2024, PhD student in Biostatistics, MSU
- 2. Di Zhang, M.Sc., degree conferred in 5/2020, PhD student at National University of Singapore
- 3. Ze Meng, M.Sc., degree conferred in 5/2020, Statistician, Henry Ford Health, Detroit
- 4. Yimin Wu, M.Sc., degree conferred in 12/2017, Statistician, DTE Energy, Ann Arbor
- 5. Ran Cao, M.Sc., degree conferred in 05/2013, Statistician, i360 at Washington DC
- 6. Chenyang Gu, M.Sc., degree conferred in 05/2012, Ph.D. student, Department of Biostatistics, Brown University
- 7. Chunyu Chen, M.Sc., degree conferred in 05/2012, Ph.D. student, Department of Animal Science, Michigan State University
- 8. Wenzhao Yang, M.Sc., degree conferred in 08/2008. She obtained her Ph.D. degree at Michigan State University in 2014.
- 9. Guanghui Liu, M.Sc., degree conferred in 12/2006, currently Biostatistician, Mayo Clinic

Other MSU Ph.D. Students (as a member of the Guidance Committee)

Name		Department	Time thesis defended
1.	Wenzhuo Tang	Dept. of Statistics and Probability	ongoing
2.	Kenia Seguraaba	Genetics and Genome Sciences Program	ongoing
3.	Jiachen Liu	Dept. of Counseling, Educational	ongoing
		Psychology and Special Education	
4.	Harold Wu	Dept. of Epidemiology and Biostatistics	ongoing
5.	Dali Liu	Dept. of Statistics and Probability	ongoing
6.	Jianrui Zhang	Dept. of Statistics and Probability	ongoing

7. Meiqi Liu	Dept. of Statistics and Probability	ongoing
8. Arash Yunesi	Dept. of Statistics and Probability	ongoing
9. Fei Zhang	Dept. of Animal Science	12/2024
10. Liang Wang	Dept. of Epidemiology and Biostatistics	07/2024
11. Sang Kyu Lee	Dept. of Statistics and Probability	07/2024
12. Jiaxin Yang	Dept. of Computational Mathematics, Science and Engineering	05/2024
13. Zichun Cao	Dept. of Epidemiology and Biostatistics	07/2023
	i Dept. of Statistics & Probability	07/2023
15. Wenjie Qi	Dept. of Computational Mathematics, Science and Engineering	08/2022
16. Sanket Jantre	Dept. of Statistics and Probability	08/2022
17. Nilanjan Chakraborty	Dept. of Statistics and Probability	07/2022
18. Di Wu	Dept. of Epidemiology and Biostatistics	06/2022
19. Runze Su	Dept. of Statistics and Probability	06/2022
20. Hao Wang	Dept. of Computational Mathematics, Science and Engineering	05/2022
21. Binbin Huang	Dept. of Computational Mathematics, Science and Engineering	12/2021
22. Ningyu Sha	Dept. of Computational Mathematics, Science and Engineering	07/2021
23. Bixi Zhang	Dept. of Counseling, Educational	06/2021
-	Psychology and Special Education	
24. Xiaoran Tong	Dept. of Epidemiology and Biostatistics	04/2020
25. Yuning Hao	Dept. of Statistics and Probability	12/2019
26. Bing Tong	Dept. of Counseling, Educational	08/2019
	Psychology and Special Education	
27. Shawn Santo	Dept. of Statistics and Probability	12/2018
28. Yaohui Ding	Dept. of Computer Science and Engineering	01/2018
29. Sneha Jadhav	Dept. of Statistics and Probability	07/2017
30. Chunyu Chen	Dept. of Animal Science	08/2017
31. Ashwini Maurya	Dept. of Statistics and Probability	05/2016
32. Keyin Wang	Dept. of Counseling, Educational	08/2016
	Psychology and Special Education	
33. Jikai Lei	Dept. of Computer Science	07/2015
34. Aritro Nath	The Genetics Program	05/2015
35. Changshuai Wei	Dept. of Epidemiology and Biostatistics	05/2014
36. Wenzhao Yang	Dept. of Animal Science	04/2014
37. Yvonne Badke	Dept. of Animal Science	08/2013
38. Greg Habing	Center for Comparative Epidemiology	08/2012
39. Xuechun Zhou	Dept. of Counseling, Educational Psychology and Special Education	08/2012
40. Ming Wu	Dept. of Computer Sciences and Engineering	08/2012
41. Shannon Bell	Dept. of Biochemistry & Molecular Biology	04/2012
42. Ming Li	Division of Biostatistics, Dept. of Epidemiology	05/2011
43. Shujie Ma	Dept. of Statistics and Probability	07/2011
44. Qiongxia Song	Dept. of Statistics and Probability	07/2010
45. Nora Bello	Dept. of Animal Science	07/2010
46. Menghan Liu	Dept. of Crop & Soil Science	03/2010
47. Young Gui Kim	Dept. of Economics	07/2009
48. Tianshu Pan	Dept. of Counseling, Educational	10/2008
49. Lan Xiao	Psychology and Special Education Dept. of Animal Science	08/2007
TJ. LAH AIdU	בפיני אין אווווזמן סטופוונפ	00/2007

Lab rotation student

• Shubham Koirala (fall 2023), BioMolecular Science Program, MSU

Undergraduate Students Advised:

- Ning Jiang (2019-2020), exchange student from Xi'an Jiaotong University
- Ruobing Jia (2017-2018), exchange student from Jilin University
- Joseph Xiao (1/2016-5/2016), exchange student from Nankai University
- Yuchen Guo (2015-2016), Regression analysis of DNA methylation data
- Supervised undergraduate Sebastian Raschka from biochemistry for a project titled "Statistical Identification of Potential CLAVATA2 Interactors by Fluorescence Resonance Energy Transfer Analysis" to participate in the online OpenIntro Student Project Competition. He was one of the students in my STT421 class in fall 2011. The work was published in http://www.openintro.org/stat/.

TEACHING EXPERIENCES

Graduate Level

- STT843 Multivariate Analysis
- STT847 Survival Data Analysis
- STT855 Statistical Genetics
- STT863 Statistics Methods I (Linear Regression)
- STT864 Statistics Methods II (Generalized Linear Model)
- STT890 Statistical Problems
- HRT892/GEN800 Quantitative Genetics Seminar

Undergraduate Level

- STT481 Issues in Statistical Practice
- STT442 Probability & Statistics II (Statistics)
- STT421 Statistics I
- STT351 Probability and Statistics for Engineering
- STT315 Intro Prob & Stat for Business
- STT231 Statistics for Scientists
 - STT200 Statistical Methods

COMMITTEE SERVICES

Professional Societies

International Congress of Basic Science (ICBS)

- 2025 Manager for the Bioinformatics and AI for Life Sciences section, Frontier of Science Award (FSA)
- 2024 Chair of the Nomination Committee and Selection Committee for the Bioinformatics and AI for Life Sciences section, Frontier of Science Award (FSA)

American Statistical Association (ASA) Committee

2021-2023 ASA Archives and Historical Materials Committee

2020-2022 Membership Engagement Committee, ASA Section on Statistics in Genomics and Genetics

Department Committee

- 2024-2025 Faculty Search Committee (as chair)
- 2023-pres Major Curriculum Committee (as chair)
- 2023-2024 Dept Bylaws Committee
- 2022-pres Reappoint, Promotion and Tenure (RPT) Committee (as chair)
- 2022-2023 Junior Faculty Search Committee (as chair)
- 2021-2023 Faculty Advisory Committee (as chair)
- 2020-2021 Mentoring Policy Committee (as chair)
- 2020-2021 Prelim Policy Committee (as chair)

- 2019-pres Graduate Support Committee (as chair)
- 2019-pres Master Support Committee
- 2019-2022 Major Curriculum Committee (as chair from 2019-2021)
- 2018-2019 Committee of the Whole (as President)
- 2018-2019 Department Tenure & Promotion Committee
- 2017-2018 Faculty Search Committee (as chair)
- 2016-2017 Data Science Master Program Committee
- 2014-2015 Faculty Search Committee for the joint CMSE-STT faculty search (as chair)
- 2014-2016 Department Advisory Committee
- 2014-2015 Committee of the Whole (as President)
- 2014-2015 Graduate Support Committee
- 2014-2015 Master Admissions Committee (as chair)
- 2014-2017 Committee on Responsible Conduct for Research training
- 2012-2013 Colloquium Committee (as chair)
- 2005-pres Committee of the Whole
- 2010-2011 Internal Review Committee
- 2009-2011 Major Curriculum Committee (as chair)
- 2010-2011 Service Course Committee
- 2007-2008 Faculty Search Committee
- 2007-2008 Graduate Support Committee
- 2006-2009 Colloquium Committee
- 2005-2007 Secretary of the Advisory Committee
- 2005-2006 Comupter Committee

College and University Committee

- 2023-pres University Honors College Committee
- 2019-pres NatSci College Graduate Committee
- 2016-2019 CMSE Undergraduate Data Science Major Committee
- 2016-2018 Committee on University Student-Faculty Judiciaries
- 2016-2017 Faculty Search Committee for Quantitative/Statistical/Computational Genomics faculty hire
- 2015-2017 Faculty Search Committee for CMSE data science faculty hire
- 2014-2015 Umbrella Committee (committee for the new CMSE department)
- 2013-2015 Executive Committee, Quantitative Biology Program
- 2009-2011 Faculty Advisory Council, College of Natural Science
- 2009 All-University Awards Committee, College of Natural Science
- 2009 Awards Committee, MSU chapter of Sigma Xi Scentific Research Society
- 2009-2011 Genetics Program Executive Committee
- 2009 Dean's Representative, College of Social Science
- 2007-2008 Faculty Search Committee in Human Statistical Genetics, Genetics Program
- 2006-2007 Faculty Search Committee in quantitative genetics/genomics, Departments of Animal Science, and Fishery and Wild Life

Other Committee Service

- 2020-2021 President of MSU Chinese Faculty Club
- 2018-2022 Executive Committee, MSU Chinese Faculty Club

PROFESSIONAL MEMBERSHIPS

- American Association for the Advancement of Science (AAAS) (2018-)
- Institute of Mathematical Society (IMS) (2024-)
- American Statistical Association (ASA) (lifetime member)
- International Chinese Statistical Association (ICSA) (lifetime member)
- Sigma Xi full membership, The Scientific Research Society (2008-)