Xiang Ji, Ph.D.

Last Updated on 08/11/2025

Assistant Professor, Department of Mathematics Tulane University, New Orleans, LA, 70118

xji4@tulane.edu • GitHub @ xji3

Personal Website: http://xiang-ji-ncsu.github.io/

Google Scholar: https://scholar.google.com/citations?user=YH4BbIMAAAAJ&hl=en



EDUCATION

| Doctor of Philosophy in Bioinformatics and Statistics (Co-Major) | December |
|--|-----------|
| North Carolina State University, Raleigh, NC | 2017 |
| Dissertation: Phylogenetic Approaches for Quantifying Interlocus Gene Conversion | |
| Committee: Jeffrey Thorne, Eric Stone, Nadia Singh and Sujit Ghosh | |
| Master of Science in Material Science and Engineering | September |
| North Carolina State University, Raleigh, NC | 2013 |
| Thesis: Laser Interference Lithography for Fabrication of Gas Sensors | |
| Committee: John Muth, Lew Reynolds, Michael Kudenov and Jon-Paul Maria | |
| Bachelor of Science in Economics (Double Major) | July 2011 |
| Peking University, Beijing, China | |
| Bachelor of Science in Physics | July 2011 |
| Peking University, Beijing, China | |
| Advisor: Jia-Sen Zhang | |

APPOINTMENTS

| Faculty Investigator | November 2024 – Present |
|--|--------------------------|
| Tulane Biostatistics and Bioinformatics Core, Tulane University, New C | Orleans, LA |
| Faculty Member | October 2021 – Present |
| Cancer Center, Tulane University, New Orleans, LA | |
| Assistant Professor | July 2020 – Present |
| Department of Mathematics, Tulane University, New Orleans, LA | |
| Postdoctoral Fellow | January 2018 – July 2020 |
| University of California, Los Angeles, CA | |

RESEARCH INTERESTS

Multigene family evolution; cancer systems biology; phylogenetics with large data sets; development of statistical techniques and software

PUBLICATIONS

- 51. Fourment, M., Macaulay, M., Swanepoel, C.J., **Ji, X.**, Suchard, M.A. and Matsen IV, F.A. (2025) torchtree: flexible phylogenetic model development and inference using PyTorch. *Systematic Biology*, p.syaf047.
- 50. Qi, K., Chen, J., Ma, X., Li, D., Baele, G., Li, X., Qin, Y., **Ji, X.**, Zhu, M., Li, X.X. and Guo, D. (2025) Novel Coronaviruses Identified in Livestock: The Urgent Need to Enhance Coronavirus Surveillance to Mitigate Zoonotic Risks. *Journal of Infection*.
- Cong, F., Bao, H., Wang, X., Tang, Y., Bao, Y., Poulton, J., Liu, X., Wong, A., Ji, X. and Deng,
 W. (2025) Translocation of gut bacteria promotes tumor-associated mortality by inducing immune-activated renal damage. *The EMBO Journal*
- 48. Niu, Q., Jiang, Z., Wang, L., **Ji, X.**, Baele, G., Qin, Y., Lin, L., Lai, A., Chen, Y., Veit, M. and Su, S. (2025) Prevention and control of avian influenza virus: Recent advances in diagnostic technologies and surveillance strategies. *Nature Communications*
- 47. Baele, G., **Ji, X.**, Hassler, G. W., McCrone, J. T., Shao, Y., Holbrook, A. J., Lemey, P., Drummond, A., Rambaut, A., Suchard, M. A. (2025) BEAST X for Bayesian phylogenetic, phylodynamic and phylogeographic inference. *Nature Methods*
- 46. Pekar, J., et al., (2025) The recency and geographical origins of the bat viruses ancestral to SARS-CoV and SARS-CoV-2. *Cell*
- 45. Jiang, Z., Yan, Z., Hou, Y., Tang, J., Zheng, M., Lu, M., **Ji, X.**, Gangavarapu, G., Li, X., and Su, S., (2025). The RodentGPOmics Atlas: a comprehensive database of rodent biology for genomes and pathogens. *Nucleic Acids Research*.
- 44. Kang, M., Wang, L.F., Sun, B.W., Wan, W.B., **Ji, X.**, Baele, G., Bi, Y.H., Suchard, M.A., Lai, A., Zhang, M., ... Su, S., (2024). Zoonotic infections by avian influenza virus: changing global epidemiology, investigation, and control. *The Lancet Infectious Diseases*.
- 43. Magee, A. F., Holbrook, A. J., Pekar, J. E., Caviedes-Solis, I. W., Matsen IV, F. A., Baele, G., Wertheim, J. O., **Ji, X.**, Lemey, P., and Suchard, M. A., (2024). Random-effects substitution models for phylogenetics via scalable gradient approximations. *Systemic Biology*.
- 42. Yang, J., Schuemie, M.J., **Ji, X.** and Suchard, M.A., (2024). Massive parallelization of massive sample-size survival analysis. *Journal of Computational and Graphical Statistics*, *33*(1), pp.289-302.
- 41. Gangavarapu, K., **Ji, X.**, Baele, G., Fourment, M., Lemey, P., Matsen IV, F.A. and Suchard, M.A., (2024). Many-core algorithms for high-dimensional gradients on phylogenetic trees. *Bioinformatics*, 40(2), p.btae030.
- 40. Huang, S., He, J., Yu, L., Guo, J., Jiang, S., Sun, Z., Cheng, L., Chen, X., **Ji, X.** and Zhang, Y., (2024). ASTK: A Machine Learning-Based Integrative Software for Alternative Splicing Analysis. *Advanced Intelligent Systems*, p.2300594.
- 39. Fisher, A., **Ji, X.**, Nishimura, A., & Suchard, M. A. (2023) Shrinkage-based random local clocks with scalable inference. *Molecular Biology and Evolution*, 40(11), msad242.
- 38. **Ji, X.,** Fisher, A. A., Su, S., Thorne, J. L., Potter, B., Lemey, P., Baele, G., & Suchard, M. A., (2023) Scalable Bayesian divergence time estimation with ratio transformations. *Systematic Biology*, syad039, arXiv preprint arXiv:2110.13298.

- 37. Yang, Y., Xu, T., Conant, G., Kishino, H., Thorne, J.L. and **Ji, X.**, (2023) Interlocus Gene Conversion, Natural Selection, and Paralog Homogenization. *Molecular Biology and Evolution*, msad198.
- 36. Matteson, N. L., et al., (2023) Genomic surveillance reveals dynamic shifts in the connectivity of COVID-19 epidemics. *Cell*, 186(26), pp.5690-5704.
- 35. He, W.T., Li, D., Baele, G., Zhao, J., Jiang, Z., **Ji, X.**, Veit, M., Suchard, M.A., Holmes, E.C., Lemey, P., Boni, M.F. and Su, S., (2023) Newly identified lineages of porcine hemagglutinating encephalomyelitis virus exhibit respiratory phenotype. *Virus Evolution*, *9*(2), p.vead051.
- 34. Zhang, Z., Nishimura, A., Trovão, N.S., Cherry, J.L., Holbrook, A.J., **Ji, X.**, Lemey, P. and Suchard, M.A., (2023) Accelerating Bayesian inference of dependency between mixed-type biological traits. *PLOS Computational Biology*, *19*(8), p.e1011419.
- 33. Zhao, J., Kang, M., Wu, H., Sun, B., Baele, G., He, W.T., Lu, M., Suchard, M.A., **Ji, X.**, He, N. and Su, S., (2023) Risk assessment of SARS-CoV-2 replicating and evolving in animals. *Trends in Microbiology*.
- 32. Zhao, J., Dellicour, S., Yan, Z., Veit, M., Gill, M.S., He, WT, Zhai, X., **Ji, X.**, Suchard, M.A., Lemey, P. and Su, S., (2023) Early Genomic Surveillance and Phylogeographic Analysis of Getah Virus, a Reemerging Arbovirus, in Livestock in China. *Journal of Virology*, 97(1), pp.e01091-22.
- 31. Tsui, L.H., et al., (2023) Genomic assessment of invasion dynamics of SARS-CoV-2 Omicron BA. 1. *Science*, 381(6655), pp.336-343.
- 30. Hassler, G.W., Magee, A.F., Zhang, Z., Baele, G., Lemey, P., **Ji, X.**, Fourment, M. and Suchard, M.A., (2023) Data Integration in Bayesian Phylogenetics. *Annual Review of Statistics and Its Application*, 10, pp.353-377.
- 29. Fisher, A. A., Hassler, G. W., **Ji, X.**, Baele, G., Suchard, M. A., & Lemey, P. (2022) Scalable Bayesian phylogenetics. *Philosophical Transactions of the Royal Society B*, 377.1861(2022): 20210242.
- 28. Zhao, J., Dellicour, S., Yan, Z., Veit, M., Gill, M.S., He, WT, Zhai, X., **Ji, X.**, Suchard, M.A., Lemey, P. and Su, S., (2022) Early Genomic Surveillance and Phylogeographic Analysis of Getah Virus, a Reemerging Arbovirus, in Livestock in China. *Journal of Virology*, pp.e01091-22.
- 27. Holbrook, A. J., **Ji, X.**, & Suchard, M. A. (2022) From viral evolution to spatial contagion: a biologically modulated Hawkes model. *Bioinformatics*, 38(7), pp.1846-1856. arXiv:2103.03348 [stat.ME]
- 26. McCrone, J.T., Hill, V., Bajaj, S. et al. (2022) Context-specific emergence and growth of the SARS-CoV-2 Delta variant. *Nature*, 610(7930), pp.154-160.
- 25. He, W., et al., (2022). Virome characterization of game animals in China reveals a spectrum of emerging pathogens. *Cell*, 185(7), 1117-1129.
- 24. Tian, J., Sun, J., Li, D., Wang, N., Wang, L., Zhang, C., Meng, X., Ji, X., Suchard, M. A., Zhang, X., Lai, A., Su, S., & Veit, M. (2022) Emerging viruses: Cross-species transmission of Coronaviruses, Filoviruses, Henipaviruses and Rotaviruses from bats. *Cell Reports*
- 23. Zhao, J., Sun, J., Li, X., Xing, G., Zhang, Y., Lai, A., Baele, G.*, **Ji, X.*,** Su, S.* (2022). Divergent Viruses Discovered in Swine Alter the Understanding of Evolutionary History

- and Genetic Diversity of the Respirovirus Genus and Related Porcine Parainfluenza Viruses. *Microbiology Spectrum*, 10(3), pp.e00242-22. *Senior author
- 22. He, W., et al., (2022) Phylogeography reveals association between swine trade and the spread of porcine epidemic diarrhea virus in China and across the world. *Molecular Biology and Evolution*, 39(2), msab364
- 21. Fan, Y., et al. (2022). Systematic analysis of inflammation and pain pathways in a mouse model of gout. *Molecular Pain*, 18, 17448069221097760.
- 20. Xie, S., et al. (2022). Disrupted myelination network in the cingulate cortex of Parkinson's disease. *IET Systems Biology*, 16(3-4), pp.98-119.
- 19. Holbrook, A. J., **Ji, X.**, & Suchard, M. A. (2022) Bayesian mitigation of spatial coarsening for a fairly flexible spatiotemporal Hawkes model. *Annals of Applied Statistics*, 16 (1), 573-595.
- 18. Holbrook, A. J., Nishimura, A., **Ji, X.**, & Suchard, M. A. (2021). Computational Statistics and Data Science in the Twenty-first Century. *Wiley StatsRef: Statistics Reference Online*. arXiv preprint arXiv:2204.05530.
- 17. Lemey, P., et al., (2021) Untangling introductions and persistence in COVID-19 resurgence in Europe. *Nature*, 595 (7869), 713-717.
- 16. Landeros, A., **Ji, X.**, Lange, K., Stutz, T. C., Xu, J., Sehl, M. E., & Sinsheimer, J. S. (2021) An examination of school reopening strategies during the SARS-CoV-2 pandemic. *PloS One*, 16(5), e0251242.
- 15. Zhang, Z., Nishimura, A., Bastide, P., **Ji, X.**, Payne, R. P., Goulder, P., ... & Suchard, M. A. (2021). Large-scale inference of correlation among mixed-type biological traits with phylogenetic multivariate probit models. *The Annals of Applied Statistics*, 15(1), 230-251.
- 14. Lindelof, K., Lindo, J. A., Zhou, W., **Ji, X.**, & Xiang, Q. Y. J. (2020) Phylogenomics, biogeography, and evolution of the blue- or white-fruited dogwoods (Cornus) insights into morphological and ecological niche divergence following intercontinental geographic isolation. *Journal of Systematics and Evolution*, 58(5), pp.604-645.
- 13. Sun, J., He, W. T., Wang, L., Lai, A., **Ji, X.**, Zhai, X., ... & Veit, M. (2020). COVID-19: epidemiology, evolution, and cross-disciplinary perspectives. *Trends in Molecular Medicine*, 26(5), pp.483-495.
- 12. He, W.*, **Ji, X.***, He, W. *, Dellicour, S. *, ..., & Su, S. (2020) Genomic epidemiology, evolutionary dynamics, and transmission patterns of porcine deltacoronavirus. *Molecular Biology and Evolution*, 37(9), pp.2641-2654. *equal contribution
- Ji, X., Zhang, Z., Holbrook, A., Nishimura, A., Baele, G., Rambaut, A., Lemey, P., & Suchard, M. A. (2020) Gradients do grow on trees: a linear-time O(N)-dimensional gradient for statistical phylogenetics. *Molecular Biology and Evolution*, 37(10), pp.3047-3060. arXiv:1905.12146 [stat.CO]
- 10. Fisher, A., **Ji, X.**, Zhang, Z., Lemey, P., & Suchard, M. A. (2020) Relaxed random walks at scale. *Systematic Biology*, 70(2), pp.258-267. arXiv:1906.04834[q-bio.PE]
- 9. Blestsa, M., Suchard, M. A., **Ji, X.**, Gryseels, S., Vrancken, B., Baele, G., Worobey, M., & Lemey, P. (2019) Divergence dating using mixed effects clock modelling: an application to HIV-1. *Virus Evolution*, 5(2), vez036.

- 8. Li, G., Zhang, W., Wang, R., Xing, G., Wang, S., **Ji, X.**, ... & Zhou, J. (2019). Genetic Analysis and Evolutionary Changes of the Torque teno sus Virus. *International journal of molecular sciences*, 20(12), 2881.
- 7. Zhou, W., **Ji, X.**, Obata, S., Pais, A., Dong, Y., Peet, R., & Xiang, Q., (2018) Resolving relationships and phylogeographic history of the Nyssa sylvatica complex using data from RAD-seq and species distribution modeling. *Molecular Phylogenetic and Evolution*, 126, 1-16.
- 6. **Ji, X.** (2017). Phylogenetic approaches for quantifying interlocus gene conversion. Doctoral Dissertation
- 5. **Ji, X.**, Griffing, A., & Thorne, J. L. (2016). A phylogenetic approach finds abundant interlocus gene conversion in yeast. *Molecular Biology and Evolution*, 33(9), 2469-2476.
- 4. Wang, K., Yu, S., **Ji, X.**, Lakner, C., Griffing, A., & Thorne, J. L. (2015). Roles of Solvent Accessibility and Gene Expression in Modeling Protein Sequence Evolution. *Evolutionary Bioinformatics*, 11, 85.
- 3. Ji, X. (2013). Laser Interference Lithography for Fabrication of Gas Sensors. Master Thesis
- 2. Han, X., **Ji, X.**, Wen, H., & Zhang, J. (2012). H-shaped resonant optical antennas with slot coupling. *Plasmonics*, 7(1), 7-11.
- 1. Xiao, G., **Ji, X.**, Gao, L., Wang, X., & Zhou, Z. (2012). Effect of dipole location on profile properties of symmetric surface plasmon polariton mode in Au/Al2O3/Au waveguide. *Frontiers of Optoelectronics*, 5(1), 63-67.

Articles in Submission and Preparation

- 6. Gangavarapu, K., **Ji, X.**, Shao, Y., Rambaut, A., Lemey, P., Baele, G. and Suchard, M. A. (2025) Tensor cores unlock efficient and lower-energy massive parallelization on phylogenetic trees. *In revision with Systematic Biology*
- 5. Baele, G., Carvalho, L. M., Brusselmans, M., Dudas, G., **Ji, X.**, McCrone, J. T., Lemey, P., Suchard, M. A. and Rambaut, A. (2025) HIPSTR: highest independent posterior subtree reconstruction in TreeAnnotator X. *In revision with Bioinformatics*
- 4. Bao, Y., Deng, W., Su, S., Lemey, P., Suchard, M. A., Glatt-Holtz, N.* and **Ji, X.*** (2025) Reflection Hamiltonian Monte Carlo for divergence time estimations. *Senior author. *In preparation*
- 3. Zhang, M. N., Bao, H. and **Ji, X.** (2025) Identification of a novel palmitoylation-related prognostic signature in head and neck squamous cell carcinoma. *In review with Translational Cancer Research*
- 2. **Ji, X.**, Redelings, B., Su, S., Bao, H., Deng, W., Hong, S. L., Baele, G., Lemey, P. and Suchard, M. A. (2025) Detecting Shifts in Natural Selection with Branch-Specific Substitution Models and Shrinkage Priors. *In review with Systematic Biology*, arXiv:2507.08386 [q-bio.PE][stat.CO]
- 1. **Ji, X.**, Thorne, J. L. (2024) A phylogenetic approach disentangles the tract length and initiation rate of interlocus gene conversions. *In wait/delay for submission*, arXiv:1908.08608 [q-bio.PE]

TEACHING EXPERIENCE

Department of Mathematics, Tulane University, New Orleans, LA

Summary: I regularly teach graduate-level Math 7260 (Linear Models) and Math 7360 (Data Analyses, a Tulane version of statistical learning), and undergraduate-level Math 3070 (Intro to Probability). Periodically, I teach Math 1230 Stats for Scientists, which is designed for students with non-math majors.

• Fall 2025

o Math 3070/6070 Intro to Probability (3 credits)

GitHub site: TBD

Level: Undergraduate, Graduate (Master)

Topics: Axioms of probability, discrete and continuous random variables, multivariate distributions, expectation, limit theorem

Enrollment: TBD

Spring 2025

o Math 6040/7260 Linear Models (3 credits)

GitHub site: https://tulane-math-7260-2025.github.io/

Level: Graduate (Master + Ph.D.)

Topics: Simple and multiple linear regressions, hypothesis testing, analysis of variance, bootstrap, logistic regression

Enrollment: 7

• Fall 2024

o Math 1230 Stats for Scientists (3 credits)

Level: Undergraduate

Topics: Probability, random variable, discrete and continuous distributions, sampling distributions, one and two-sample estimations, hypothesis testing, simple linear regression

Enrollment: 62

o Math 3070/6070 Intro to Probability (3 credits)

GitHub site: https://tulane-math-3070-2024.github.io/

Level: Undergraduate, Graduate (Master)

Topics: Axioms of probability, discrete and continuous random variables, multivariate distributions, expectation, limit theorem

Enrollment: 33

Spring 2024

Teaching Relief

• Fall 2023

o Math 7360 Data Analysis (3 credits)

GitHub site: https://tulane-math-7360-2023.github.io/

Level: Graduate (Master + Ph.D.)

Topics: R, RMarkdown, Rcpp, git, data visualization using ggplot2, web scraping, linear models, generalized linear models, neural network, classification

Enrollment: 22

Math 3070/6070 Intro to Probability (3 credits)

GitHub site: https://tulane-math-3070-2023.github.io/

Level: Undergraduate, Graduate (Master)

Topics: Axioms of probability, discrete and continuous random variables, multivariate distributions, expectation, limit theorem

Enrollment: 48

Spring 2023

o Math 6040/7260 Linear Models (3 credits)

GitHub site: https://tulane-math-7260-2023.github.io/

Level: Graduate (Master + Ph.D.)

Topics: Simple and multiple linear regressions, hypothesis testing, analysis of variance, bootstrap, logistic regression

Enrollment: 13

Math 7980 Independent Study (3 credits)

Level: Graduate (Master)

Topics: Item response theory models

Enrollment: 1

• Fall 2022

Math 3070/6070 Intro to Probability (3 credits)

GitHub site: https://tulane-math-3070-2022.github.io/

Level: Undergraduate, Graduate (Master)

Topics: Axioms of probability, discrete and continuous random variables, multivariate distributions, expectation, limit theorem

Enrollment: 37

Math 1230 Stats for Scientists (3 credits)

Level: Undergraduate

Topics: Probability, random variable, discrete and continuous distributions, sampling distributions, one and two-sample estimations, hypothesis testing, simple linear regression

Enrollment: 66

Math 7980 Independent Study (3 credits)

Level: Graduate (Ph.D.)

Topics: Bayesian Phylogenetics

Enrollment: 1

Spring 2022

o Math 6040/7260 Linear Models (3 credits)

GitHub site: https://tulane-math-7260-2022.github.io/

Level: Graduate (Master + Ph.D.)

Topics: Simple and multiple linear regressions, hypothesis testing, analysis of variance, bootstrap, logistic regression

Enrollment: 16

Math 7980 Independent Study (3 credits)

Level: Graduate (Master)

Topics: Review of optimization methods

Enrollment: 1

• Fall 2021

o Math 7360 Data Analysis (3 credits)

GitHub sites: https://tulane-math-7360-2021.github.io/

Level: Graduate (Master + Ph.D.)

Topics: R, RMarkdown, Rcpp, git, data visualization using ggplot2, web scraping, linear models, generalized linear models, neural network, classification

Enrollment: 19

Spring 2021

o Math 6040/7260 Linear Models (3 credits)

GitHub site: https://tulane-math-7260-2021.github.io/

Level: Graduate (Master + Ph.D.)

Topics: Simple and multiple linear regressions, hypothesis testing, analysis of variance, bootstrap, logistic regression

Enrollment: 17

• Fall 2020

o Math 7360 Data Analysis (3 credits)

GitHub site: https://tulane-math7360.github.io/

Level: Graduate (Master + Ph.D.)

Topics: R, RMarkdown, Rcpp, git, data visualization using ggplot2, web scraping, linear models, generalized linear models, neural network, classification

Enrollment: 12

STUDENT ADVISING

Co-Supervisor (Doctoral) 2024 -Present Department of Microbiology and Immunology, The University of Melbourne Melbourne, Australia Student name: John Tay Supervisor: Sebastian Duchene • Research Title: TBD **Committee Chair (Doctoral)** 2023 -Present Department of Mathematics, Tulane University, New Orleans, LA Student name: Yufei Zou • Research Title: Evolutionary inference with interlocus gene conversion 2022 -**Committee Chair (Doctoral)** Present Department of Mathematics, Tulane University, New Orleans, LA Student name: Yuwei Bao • Research Title: Reflective Hamiltonian Monte Carlo method

Prospectus Exam Committee Member (Doctoral)

2025

Department of Mathematics, Tulane University, New Orleans, LA

• Student name: John Argentino

• Chair: Michelle Lacey

| Research title: Multivariate Modeling Techniques for Estimating Assistance Impacts | |
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| from Daily Food Security Surveys | |
| Dissertation Defense Committee Member (Doctoral) | 2025 |
| Department of Mathematics, Tulane University, New Orleans, LA | |
| Student name: Linh Do | |
| Chair: Scott McKinley | |
| Research title: Statistical techniques for comparing populations of continuous piecewise- | |
| linear stochastic processes | |
| Prospectus Exam Committee Chair (Doctoral) | 2024 |
| Department of Mathematics, Tulane University, New Orleans, LA | |
| Student name: Yuwei Bao | |
| Chair: Xiang Ji | |
| Research title: Scalable Bayesian statistical phylogenetics models | |
| Dissertation Defense Committee Member (Doctoral) | 2024 |
| Department of Mathematics, Tulane University, New Orleans, LA | |
| Student name: Daniela A. Florez Pineda | |
| Chair: Ricardo Cortez | |
| Oral Exam Committee Member (Doctoral) | 2024 |
| Department of Mathematics, Tulane University, New Orleans, LA | |
| Student name: John V. Argentino | |
| Chair: Michelle Lacey | |
| Prospectus Exam Committee Member (Doctoral) | 2024 |
| Department of Mathematics, Tulane University, New Orleans, LA | |
| Student name: Daniela A. Florez Pineda | |
| Chair: Ricardo Cortez | |
| • Research title: Mathematical models for transmission and control of mosquito-borne | |
| diseases | |
| Prospectus Exam Committee Member (Doctoral) | 2023 |
| Department of Mathematics, Tulane University, New Orleans, LA | |
| Student name: Linh Do | |
| Chair: Scott McKinley | |
| Research title: Statistical techniques for comparing populations of continuous | |
| piecewise-linear stochastic processes | |
| Oral Exam Committee Member (Doctoral) | 2023 |
| Department of Mathematics, Tulane University, New Orleans, LA | |
| Student name: Moslem Uddin | |
| Chair: Lisa Fauci | |
| Oral Exam Committee Chair (Doctoral) | 2023 |
| Department of Mathematics, Tulane University, New Orleans, LA | |
| Student name: Yuwei Bao | |
| Chair: Xiang Ji | |

| Oral Exam Committee Member (Doctoral) | 2023 |
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| Department of Mathematics, Tulane University, New Orleans, LA | |
| Student name: Sinchita Lahiri | |
| Chair: Kyle Kun Zhao | |
| Oral Exam Committee Member (Doctoral) | 2022 |
| Department of Mathematics, Tulane University, New Orleans, LA | |
| • Student name: Linh Do | |
| Chair: Scott McKinley | |
| Oral Exam Committee Member (Doctoral) | 2022 |
| Department of Mathematics, Tulane University, New Orleans, LA | |
| Student name: Daniela A. Florez Pineda | |
| Chair: Ricardo Cortez | |
| Independent Study Advisor (Master) | 2022 |
| Department of Mathematics, Tulane University, New Orleans, LA | |
| Student name: Paul Pluscht | |
| Project topic: Review of optimization methods | |
| Undergraduate Research Advisor (Undergraduate) | 2020 – |
| Department of Statistics, North Carolina State University, Raleigh, NC | 2022 |
| Student name: Yixuan Yang | |
| Co-advisor: Jeffrey L. Thorne | |
| • Research topic: Quantifying interlocus gene conversion and natural selection | |
| Undergraduate Research Advisor (Undergraduate) | 2021 |
| Department of Mathematics, Tulane University, New Orleans, LA | |
| Student name: Jiayun Ling | |
| Co-advisor: Xin Jiang | |
| Research topic: Item response theory model, Add Health data, social science | |
| | |
| MENTEE ACHIEVEMENTS | |
| Megan (Jiayun) Ling (undergraduate research advisor) | 2021 |
| Oral presentation at ASA Louisiana Chapter meeting in 2021 | |
| Graduated from master's program in Data Analytics of Duke University | |
| Now working as Data Analyst at Blizzard Entertainment | |
| Yuwei Bao (PhD thesis advisor) | 2022 – |
| Oral presentation at ASA Louisiana Chapter meeting in 2022 | Present |
| Oral presentation at Tulane Math Graduate Student Colloquium in 2022 | |
| Oral presentation at Evolution Meeting 2023 | |
| Oral presentation at Scientific Computing Around Louisiana Meeting 2023 | |
| Oral presentation at Math for All in Nola meeting 2023 | |
| Poster presentation at New England Statistical Symposium 2024 | |
| Poster presentation at Pharmaceutical Data Science Conference online 2024 | |
| F | |

- Poster presentation at ASA Conference on Statistical Practice 2024
- Tuition Scholarship & Travel Award at Summer Institute in Statistical Genetics 2023
- Travel Award from Society for Study of Evolution at Evolution Meeting 2023
- Summer Research Fund from Math Department at Tulane University 2023
- Travel Award from Graduate Studies Student Association at Tulane University 2023 & 2024
- Student Poster Award at New England Statistical Symposium 2024
- FDA-OCE-ASA Oncology Educational Fellow 2024
- Internship at Moderna Spring 2025

Yufei Zou (undergraduate & graduate research advisor)

2022 -

- Tuition Scholarship & Travel Award at Summer Institute in Statistical Genetics 2023 Present
- Oral presentation at ASA Louisiana Chapter Meeting on May 19, 2025
- Oral presentation at Evolution Meeting 2025
- SSE Caregiver Awards for attending Evolution Meeting 2025
- Attending PhD program in Data Science at UNC Charlotte

AWARDS

| HPC Fund Research Cloud Allocation Award ¹ | AMD Corporation | 2023 |
|---|---------------------------------|------|
| NVIDIA Academic Hardware Grant ² | NVIDIA Corporation | 2022 |
| Tuition Scholarship | $SISMID^3$ | 2018 |
| NIEHS Fellowship ⁴ | North Carolina State University | 2015 |
| SAMSI Graduate Fellow | SAMSI ⁵ | 2014 |
| Tuition Scholarship | ${ m SISG^6}$ | 2013 |
| University Graduate Fellowship | North Carolina State University | 2011 |

FUNDED PROJECTS

External

Louisiana Board of Regents Research Competitiveness Subprogram

07/01/23 -

• Title: Molecular epidemiology through scalable statistical phylogenetic Modelling

06/30/26

• Principle Investigator: Xiang Ji

• Total Amount: \$159,000

• Direct Cost: \$159,000

National Science Foundation

05/01/23 -

• Title: Evolutionary inference with interlocus gene conversion

07/31/24

¹1,080 node hours (targeting 4 MI100 GPUs/node)

² Donation of one A100 80Gb PCIE GPU card

³ SISMID: Summer Institute in Statistics and Modeling in Infectious Diseases at the University of Washington at Seattle

⁴ The funds were matched through North Carolina State University

⁵ SAMSI: The Statistical and Applied Mathematical Sciences Institute

⁶ SISG: Summer Institute in Statistical Genetics at the University of Washington at Seattle

| Principle Investigator: Jeffrey L. Thorne | |
|--|------------|
| Co-Investigator: Xiang Ji | |
| • Award number: DEB1754142 | |
| • Total Amount: \$564,338 | |
| • Shared Amount: \$94,351 | |
| National Institute of Health | 07/01/22 - |
| • Title: Statistical innovation to integrate sequences and phenotypes for scalable | 03/31/25 |
| phylodynamic inference | |
| Principle Investigator: Marc A. Suchard | |
| Subaward Investigator: Xiang Ji | |
| • Award number: R01AI153044 | |
| • Total Amount: \$2,321,335 | |
| • Shared Amount: \$117,259 | |
| National Institute of Health | 04/01/22 - |
| • Title: Notch signaling and germline-soma interactions in <i>Drosophila</i> ovarian model | 03/31/27 |
| Principle Investigator: Wu-Min Deng | |
| Co-Investigator: Xiang Ji | |
| • Award number: R01GM072562 | |
| • Total Amount: \$ 2,432,208 | |
| • Shared Amount: \$86,460 | |
| National Institute of Health | 02/01/20 - |
| • Title: Consortium for viral systems biology (CViSB) | 01/31/22 |
| Principle Investigator: Robert F. Garry | |
| Co-Investigator: Xiang Ji | |
| Award number: U19AI135995 | |
| • Total Amount: \$1,143,554 | |
| • Shared Amount: \$51,774 | |
| Internal | |
| Tulane University CoR Research Fellowship | 05/01/23 - |
| Title: Scalable molecular epidemiology models | 05/31/24 |
| | 03/31/21 |
| Principle Investigator: Xiang Ji Total Amount: \$5,500 | |
| Total Amount. \$5,500 | |
| SUBMITTED PROPOSALS | |
| National Institute of Health | 2025 |
| • Title: Statistical innovation to integrate sequences and phenotypes for | |
| scalable phylodynamic inference | |
| Principle Investigator: Marc A. Suchard | |
| Subaward Investigators: Xiang Ji | |

| • Total Amount: \$1,833,000 | |
|---|------|
| • Shared Amount: \$204,805 | |
| Status: Pending | |
| National Institute of Health | 2025 |
| • Title: COBRE for Basic and Translational Research in Cancer (BATRIC) | |
| Principle Investigator: Wu-Min Deng | |
| Core Co-Director: Xiang Ji | |
| • Total Amount: \$11,420,264 | |
| • Shared Amount: \$86,460 | |
| • Status: Pending | |
| National Science Foundation | 2025 |
| • Title: FEC: Identifying factors that mitigate the effect of poverty on child | |
| brain development using computational neural fingerprinting techniques | |
| Principle Investigator: Yu-ping Wang | |
| Co-Investigators: Xiang Ji | |
| • Total Amount: \$4,000,000 | |
| • Shared Amount: \$20,000 | |
| Status: Pending | |
| National Institute of Health | 2025 |
| Title: Scalable statistical advances to quantify interlocus gene conversion | |
| in multigene family evolution | |
| Principle Investigator: Xiang Ji | |
| Co-Investigators: None | |
| • Total Amount: \$1,833,000 | |
| Status: Pending | |
| National Science Foundation | 2024 |
| Title: Scalable phylogenomic inference through massively parallelized action-based BEAGLE library | |
| Principle Investigator: Xiang Ji | |
| Co-Investigators: Robert F. Garry and Marc A. Suchard | |
| • Total Amount: \$1,600,775 | |
| Status: Not Funded | |
| National Institute of Health | 2024 |
| Title: Tracking and controlling viral outbreaks through scalable | |
| biologically realistic phylodynamic integrations | |
| Principle Investigator: Xiang Ji | |
| • Co-Investigators: Robert F. Garry and Marc A. Suchard | |
| • Total Amount: \$2,693,909 | |
| Status: Not Funded | |
| National Institute of Health | 2024 |

 Title: Characterizing intra- and inter-host Nov evolution using human challenge study samples

Principle Investigator: Pengbo LiuSubaward Investigators: Xiang Ji

• Shared Amount: \$120,018

• Status: Not Funded

CONFERENCE PRESENTATIONS

Upcoming Conference Presentations

Ji, X. (2025, October) Scalable phylogenetic inference with Hamiltonian Monte Carlo method. Statistics and AI in Precision Health and Biology, AMS Sectional Meeting, Tulane University, New Orleans, LA

Oral Presentations

- 11 **Ji, X.** (2025, August) *Quantitative traits on massive trees with many incomplete measurements.* JSM 2025, Nashville, TN
- 10 **Ji, X.** (2025, June) *Action-based phylogenetic likelihood calculations for large state-space models.* Evolution 2025 Meeting, University of Georgia, Athens, GA
- 9 **Ji, X.** (2025, June) Scalable divergence time estimations with Hamiltonian Monte Carlo method. 2025 International Indian Statistical Association Conference, University of Nebraska-Lincoln, Lincoln, NE
- 8 **Ji, X.** (2025, May) Scalable evolutionary analysis with Hamiltonian Monte Carlo methods. STATGEN 2025: Conference on Statistics in Genomics and Genetics, ASA Section on Statistics in Genomics and Genetics, University of Minnesota, Minneapolis, MN
- Ji, X. (2025, May) Action-based phylogenetic likelihood calculations for large state-space models. 2025 Spring ASA Louisiana Chapter Meeting, LSU Health Science Center, New Orleans, LA
- 6 **Ji, X.** (2023, June) Scalable phylogenetic inference with Hamiltonian Monte Carlo method. Evolution 2023 meeting, Albuquerque, NM
- 5 **Ji, X.** (2023, March) *Scalable phylogenetic inference*. SCALA 2023: Scientific Computing Around Louisiana, Tulane University, New Orleans, LA
- 4 **Ji, X.** (2021, August) Divergence time estimation with Hamiltonian Monte Carlo sampling and ratio transform. Invited speaker at 2nd AsiaEvo Conference, Online
- 3 **Ji, X.** (2021, July) Divergence time estimation with Hamiltonian Monte Carlo sampling and ratio transform. Society of Molecular Biology and Evolution Meeting, Online
- 2 Ji, X. (2018, September) Large-scale molecular epidemiology for viruses: efficient algorithms and new models. Oral session presented at the Taishan Forum on Viral Infectious Diseases, Taishan Medical University, Tai'an, Shandong Province, China
- 1 **Ji, X.** (2017, August) *Phylogenetic approaches for quantifying interlocus gene conversion.* Quest for Ortholog 5 Meeting, University of Southern California, Los Angeles, CA

Poster Presentations

- 2 **Ji, X.** (2017, July). *Phylogenetic approaches for quantifying interlocus gene conversion.* Society of Molecular Biology and Evolution Meeting, Austin, TX
- 1 **Ji, X.** (2014, June). *A Phylogenetic approach for quantifying interlocus gene conversion*. Evolution Meeting, Raleigh, NC

INVITED TALKS

| 16 | Scalable phylogenetic Hamiltonian Monte Carlo method and its application in | 03/05/2025 |
|----|--|------------|
| | molecular epidemiology | |
| | Iowa State University, Ames, IA | |
| 15 | Scalable phylogenetic Hamiltonian Monte Carlo method and its application in | 08/29/2024 |
| | divergence time estimations | |
| | Southeastern Louisiana University, Hammond, LA | |
| 14 | Phylogenetic approach for estimating amounts of interlocus gene conversion in duplications | 12/06/2023 |
| | AMS/AWM at Tulane University, New Orleans, LA | |
| 13 | Scalable phylogenetic inference via Hamiltonian Monte Carlo method | 02/03/2023 |
| | Indiana University, Bloomington, IN | |
| 12 | Scalable phylogenetic algorithm, modeling, and inference | 10/10/2022 |
| | North Carolina State University, Raleigh, NC | |
| 11 | Smooth non-parametric coalescent priors for scalable divergence time estimations | 10/05/2022 |
| | AMS/AWM at Tulane University, New Orleans, LA | |
| 10 | Divergence time estimation with Hamiltonian Monte Carlo sampling and ratio | 08/05/2020 |
| | transform | |
| | KU Leuven, Leuven, Belgium | |
| 9 | Scalable modeling and inference for phylogenetics – from interlocus gene | 12/04/2019 |
| | conversion to evolving pathogens | |
| | Tulane University, New Orleans, LA | |
| 8 | Scalable modeling and inference for phylogenetics – from interlocus gene | 11/04/2019 |
| | conversion to evolving pathogens | |
| | Louisiana State University, Baton Rouge, LA | |
| 7 | Large-scale molecular epidemiology for viruses | 11/08/2018 |
| | University of California, Los Angeles, CA | |
| 6 | Phylogenetic approaches for quantifying interlocus gene conversion | 09/18/2018 |
| | Zhejiang University, Hangzhou, China | |
| 5 | Phylogenetic approaches for quantifying interlocus gene conversion | 09/04/2018 |
| | Chinese Academy of Sciences, Beijing, China | |
| 4 | Phylogenetic approaches for quantifying interlocus gene conversion | 05/30/2018 |
| | CGM Online | |
| 3 | Phylogenetic approaches for quantifying interlocus gene conversion | 02/15/2018 |
| | Duke University, Durham, NC | |
| 2 | Phylogenetic approaches for quantifying interlocus gene conversion | 04/25/2017 |
| | Temple University, Philadelphia, PA | |
| | | |

Phylogenetic approaches for quantifying interlocus gene conversion *North Carolina State University, Raleigh, NC*

SOFTWARE

IGCexpansion

- My software for studying interlocus gene conversion.
- Available at https://github.com/xji3/IGCexpansion.

IGC BEAST Tutorial

- A tutorial for the Bayesian method (under development) of quantifying interlocus gene conversion in BEAST.
- Available at https://github.com/xji3/IGC BEAST Tutorial.

BEAST X

- I am a <u>developer</u> of the BEAST X software package.
- Available at https://github.com/beast-dev/beast-mcmc.

BEAGLE

- I am a <u>developer</u> of the BEAGLE library.
- Available at https://github.com/beagle-dev/beagle-lib.

Bito

- I collaborate with a group of statisticians and programmers to help them use my linear-time gradient algorithm implementations for their variational Bayesian phylogenetics development.
- Available at https://github.com/phylovi/bito.

SYNERGISTIC ACTIVITIES

Grant Panelist

• I served as a guest panel member for the UK MRC Better Methods, Better Research (BMBR) Panel in 2023.

Professional service

- I serve as an Editorial Board member for *Discover Viruses* by *Springer Nature*.
- I serve as a Review Editor for Frontiers in Statistical Genetics and Methodology.
- I have reviewed manuscripts for
 - o Nature Medicine (3)
 - o Proceedings of National Academy of Sciences (2)
 - o Molecular Biology and Evolution (9)
 - Systematic Biology (1)
 - Annals of Applied Statistics (2)
 - o Journal of Computational and Graphical Statistics (1)
 - o Theoretical Population Biology (1)
 - The American Journal of Human Genetics (1)
 - o *Plos Pathogens* (1)
 - o Microbiology Spectrum (1)

- o IEEE/ACM Transactions on Computational Biology and Bioinformatics (1)
- o Frontiers in Public Health (2)
- o Frontiers in Virology (2)
- o BMC Ecology and Evolution (1)
- o Evolutionary Bioinformatics (2)
- o Scientific Reports (1)
- o International Journal of Data Mining and Bioinformatics (1)
- o Rapid Reviews: COVID-19 (1)
- Open Veterinary Journal (1)

Departmental service

- I led the development and establishment of a new Master of Science in Data Science program jointly hosted by the Math and Computer Science Departments at Tulane University.
- I served as the Director of Master Programs of the Math Department at Tulane University from Fall 2021 to Spring 2024.
- I served on the Graduate Study Committee of the Math Department at Tulane University from Fall 2021 to Spring 2024.
- I served on the Search Committee for a Professor of Practice in Statistics position from Fall 2021 to Spring 2023.
- I organize the Prob & Stats research seminar series of the Math Department at Tulane University since Fall 2023.
- I ran a tutorial workshop on building personal website for graduate students in the Math Department on 04/05/2022 (tutorial link: xji3.github.io/Tutorial).
- I serve on the School of Science & Engineering Cypress HPC committee as the representative for the Math Department since Spring 2025.

Outreach

- I served as an ASN student poster award judge for the Evolution 2025 meeting.
- I participated in the LA FIRST Lego League State Championship as a volunteer judge in January 2023.
- I participated in Boys at Tulane in STEM (BATS) and Girls in STEM at Tulane (GIST), organized by Tulane Center for K-12 System Education
 - o Volunteer, Boys at Tulane in STEM (BATS), 03/18/2023
 - O Volunteer, Boys at Tulane in STEM (BATS), 09/16/2023
 - o Volunteer, Girls in STEM at Tulane (GiST), 11/04/2023
 - o Volunteer, Girls in STEM at Tulane (GiST), 03/09/2024
 - o Volunteer, Boys at Tulane in STEM (BATS), 04/13/2024
 - o Volunteer, Boys at Tulane in STEM (BATS), 09/21/2024
 - o Volunteer, Girls in STEM at Tulane (GiST), 10/26/2024
 - o Volunteer, Boys at Tulane in STEM (BATS), 02/01/2025
 - o Volunteer, Girls in STEM at Tulane (GiST), 03/22/2025
- I served as treasurer of the ASSIST⁷ Student Leadership Council in 2012 and 2013. I participated in the Magnet Fair at South Raleigh Magnet High School as an ASSIST center graduate representative in 2012.

⁷ ASSIST: The NSF Center for Advanced Self-Powered Systems of Integrated Sensors and Technologies at NC State University