Department of Biostatistics and Bioinformatics Email: zhicheng.ji@duke.edu

Duke University School of Medicine Homepage: https://zji90.github.io/

### Education

2020 Ph.D. in Biostatistics, Johns Hopkins Bloomberg School of Public Health

2020 M.S.E. in Computer Science, Johns Hopkins Whiting School of Engineering

2013 B.S. in Statistics, Fudan University

## Professional Experience

08/2020-present Tenure-track Assistant Professor

Department of Biostatistics and Bioinformatics , Duke University School of Medicine

### Honors and Awards

2021 Whitehead Scholar, Duke Universit	2021	Whitehead	Scholar.	Duke	Universit
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2018 Margaret Merrell Award, Department of Biostatistics, Johns Hopkins University

2018 June B. Culley Award, Department of Biostatistics, Johns Hopkins University

2017 Runner-up, ENCODE-DREAM in vivo Transcription Factor Binding Site Prediction Challenge

2016 ASA Section on Statistics in Genomics and Genetics Distinguished Student Paper Award

2015 Top Performers, Prostate Cancer DREAM Challenge

2014 Kocherlakota Award, Department of Biostatistics, Johns Hopkins University

2012 First-class Scholarship, Fudan University

### **Publications**

### Google Scholar

equal contributions\*, corresponding author†, mentee

### Journal Articles and Articles under Review

- 1 Wenpin Hou, **Zhicheng Ji**<sup>†</sup>. Palo: Spatially-aware color palette optimization for single-cell and spatial data. Submitted.
- 2 Yang Liu, Zhiyuan Huang, Honghe Liu, **Zhicheng Ji**, Amit Arora, Danfeng Cai, Zhe Liu, Eric A. J. Simko, Yanjun Zhang, Goran Periz, Jiou Wang. C9orf72 Hexanucleotide Repeat DNA Initiates Pathogenic Cascades Through Chromatin and Epigenome Regulator DAXX. Submitted.
- 3 Yuqi Fang, **Zhicheng Ji\***, Weiqiang Zhou\*, Jordi Abante\*, Michael Koldobskiy, Hongkai Ji, Andrew Feinberg. DNA methylation entropy is associated with DNA sequence features and developmental epigenetic divergence. Submitted.
- 4 Jifeng Zhang, Xiao Wang, **Zhicheng Ji**, Weidong Tian. Large-scale prediction of key dynamic interacting proteins in multiple cancers. Submitted. bioRxiv

5 Boyang Zhang, **Zhicheng Ji**, and Hongkai Ji. Tree-based Correlation Screen and Visualization for Exploring Phenotype-Cell Type Association in Multiple Sample Single-Cell RNA-Sequencing Experiments. Genome Biology. Under review. bioRxiv

- Haotian Zhuang, Huimin Wang, **Zhicheng Ji**<sup>†</sup>. findPC: An R package to automatically select number of principal components in single-cell analysis. Bioinformatics. In revision. bioRxiv
- 7 Wenpin Hou, **Zhicheng Ji**, Zeyu Chen, E. John Wherry, Stephanie C. Hicks, Hongkai Ji. A statistical framework for differential pseudotime analysis with multiple single-cell RNA-seq samples. Nature Portfolio. In revision. bioRxiv
- 8 Lex RK, Zhou W, **Ji Z**, Falkenstein KN, Schuler KE, Windsor KE, Kim JD, Ji H, Vokes SA. GLI transcriptional repression is inert prior to Hedgehog pathway activation. Nature Communications. 2022;13(1):808.
- 9 Hou W, Ji Z<sup>†</sup>. Unbiased visualization of single-cell genomic data with SCUBI. Cell Reports Methods. 2022;2(1):100135.
- 10 Stephens KE, Zhou W, Renfro Z, **Ji Z**, Ji H, Guan Y, Taverna SD. Global gene expression and chromatin accessibility of the peripheral nervous system in animal models of persistent pain. Journal of Neuroinflammation. 2021;18(1):185.
- 11 Caushi JX, Zhang J, **Ji Z**, Vaghasia A, Zhang B, Hsiue EH, Mog BJ, Hou W, Justesen S, Blosser R, Tam A, Anagnostou V, Cottrell TR, Guo H, Chan HY, Singh D, Thapa S, Dykema AG, Burman P, Choudhury B, Aparicio L, Cheung LS, Lanis M, Belcaid Z, El Asmar M, Illei PB, Wang R, Meyers J, Schuebel K, Gupta A, Skaist A, Wheelan S, Naidoo J, Marrone KA, Brock M, Ha J, Bush EL, Park BJ, Bott M, Jones DR, Reuss JE, Velculescu VE, Chaft JE, Kinzler KW, Zhou S, Vogelstein B, Taube JM, Hellmann MD, Brahmer JR, Merghoub T, Forde PM, Yegnasubramanian S, Ji H, Pardoll DM, Smith KN. Transcriptional programs of neoantigen-specific TIL in anti-PD-1-treated lung cancers. Nature. 2021;596(7870):126-132.
  - Reviewed in Nature Reviews Clinical Oncology "Tumour antigen-induced T cell exhaustion the archenemy of immune-hot malignancies"
- Ji Z, Ji H. Discussion of "Exponential-family Embedding with Application to Cell Developmental Trajectories for Single-cell RNA-seq Data". Journal of the American Statistical Association. 2021;116(534):471-474.
- 13 Zhang J, Miao G, Hu S, Sun Q, Ding H, **Ji Z**, Guo P, Yan S, Wang C, Kan X, Nie L. Quantification and evolution of mitochondrial genome rearrangement in Amphibians. BMC Ecology and Evolution. 2021;21(1):19.
- 14 Chamling X, Kallman A, Fang W, Berlinicke CA, Mertz JL, Devkota P, Pantoja IEM, Smith MD, **Ji Z**, Chang C, Kaushik A, Chen L, Whartenby KA, Calabresi PA, Mao HQ, Ji H, Wang TH, Zack DJ. Single-cell transcriptomic reveals molecular diversity and developmental heterogeneity of human stem cell-derived oligodendrocyte lineage cells. Nature Communications. 2021;12(1):652.
- Dangi A, Natesh NR, Husain I, **Ji Z**, Barisoni L, Kwun J, Shen X, Thorp EB, Luo X. Single-cell transcriptomics of mouse kidney transplants reveals a myeloid cell pathway for transplant rejection. JCI Insight. 2020; 5(20):e14132.
- 16 Hou W, **Ji Z**, Ji H, Hicks S. A Systematic Evaluation of Single-cell RNA-sequencing Imputation Methods. Genome Biology. 2020; 21(1):218.
- Ji Z, Zhou W, Hou W, Ji H. Single-cell ATAC-seq signal extraction and enhancement with SCATE. Genome Biology. 2020; 21(1):161.
- 18 Zhang J, Yan S, Jiang C, **Ji Z**, Wang C, Tian W. Network Properties of Cancer Prognostic Gene Signatures in the Human Protein Interactome. Genes (Basel). 2020; 11(3):247.
- Zhang J\*, Ji Z\*, Caushi JX\*, El Asmar M\*, Anagnostou V, Cottrell TR, Chan HY, Suri P, Guo H, Merghoub T, Chaft JE, Reuss JE, Tam AJ, Blosser RL, Abu-Akeel M, Sidhom JW, Zhao N, Ha JS, Jones DR, Marrone KA, Naidoo J, Gabrielson E, Taube JM, Velculescu VE, Brahmer JR, Housseau F, Hellmann MD, Forde PM, Pardoll DM, Ji H\*, Smith KN\*. Compartmental Analysis of T-cell Clonal Dynamics as a Function of Pathologic Response to Neoadjuvant PD-1 Blockade in Resectable Non-Small Cell Lung Cancer. Clinical Cancer Research. 2020; 26(6):1327-1337.

20 Lex RK\*, **Ji Z\***, Falkenstein KN\*, Zhou W, Henry JL, Ji H, Vokes SA. GLI transcriptional repression regulates tissue-specific enhancer activity in response to Hedgehog signaling. Elife. 2020; 9:e50670.

- 21 Stephens KE, Zhou W, **Ji Z**, Chen Z, He S, Ji H, Guan Y, Taverna SD. Sex differences in gene regulation in the dorsal root ganglion after nerve injury. BMC Genomics. 2019; 20(1):147.
- Chen Z\*, **Ji Z\***, Ngiow SF, Manne S, Cai Z, Huang AC, Johnson J, Staupe RP, Bengsch B, Xu C, Yu S, Kurachi M, Herati RS, Vella LA, Baxter AE, Wu JE, Khan O, Beltra JC, Giles JR, Stelekati E, McLane LM, Lau CW, Yang X, Berger SL, Vahedi G, Ji H, Wherry EJ. TCF-1-Centered Transcriptional Network Drives an Effector versus Exhausted CD8 T Cell-Fate Decision. Immunity. 2019; 51(5):840-855.e5.
  - Reviewed in Immunity "Back to the future: effector fate during T cell exhaustion"

220+ citations on Google Scholar

- 23 Zhou W, **Ji Z**, Fang W, Ji H. Global prediction of chromatin accessibility using small-cell-number and single-cell RNA-seq. Nucleic Acids Research. 2019; 47(19):e121.
- 24 Kuang Z, **Ji Z**, Boeke JD, Ji H. Dynamic motif occupancy (DynaMO) analysis identifies transcription factors and their binding sites driving dynamic biological processes. Nucleic Acids Research. 2018; 46(1):e2.
- 25 Zhou W, Sherwood B, **Ji Z**, Xue Y, Du F, Bai J, Ying M, Ji H. Genome-wide prediction of DNase I hypersensitivity using gene expression. Nature Communications. 2017; 8(1):1038.
- 26 Han F, Ji H, **Ji Z**, Wang H. A provable smoothing approach for high dimensional generalized regression with applications in genomics. Electronic Journal of Statistics. 2017; 11(2):4347-4403.
- 27 Seyednasrollah F, et al. A DREAM Challenge to Build Prediction Models for Short-Term Discontinuation of Docetaxel in Metastatic Castration-Resistant Prostate Cancer. JCO Clinical Cancer Informatics. 2017;1:1-15. (Listed as consortium coauthor)
- 28 Guinney J, et al. Prediction of overall survival for patients with metastatic castration-resistant prostate cancer: development of a prognostic model through a crowdsourced challenge with open clinical trial data. Lancet Oncology. 2017; 18(1):132-142. (Listed as consortium coauthor)
- 29 Ji Z\*, Zhou W\*, Ji H. Single-cell regulome data analysis by SCRAT. Bioinformatics. 2017; 33(18):2930-2932.
- 30 Li Q, Lex RK, Chung H, Giovanetti SM, Ji Z, Ji H, Person MD, Kim J, Vokes SA. The Pluripotency Factor NANOG Binds to GLI Proteins and Represses Hedgehog-mediated Transcription. Journal of Biological Chemistry. 2016; 291(13):7171-82.
- 31 Norrie JL, Li Q, Co S, Huang BL, Ding D, Uy JC, **Ji Z**, Mackem S, Bedford MT, Galli A, Ji H, Vokes SA. PRMT5 is essential for the maintenance of chondrogenic progenitor cells in the limb bud. Development. 2016; 143(24):4608-4619.
- 32 Ji Z, Ji H. TSCAN: Pseudo-time reconstruction and evaluation in single-cell RNA-seq analysis. Nucleic Acids Research. 2016; 44(13):e117.
  - Winner of ASA Section on Statistics in Genomics and Genetics Distinguished Student Paper Award 390+ citations on Google Scholar
- 33 Hong X, Ladd-Acosta C, Hao K, Sherwood B, Ji H, Keet CA, Kumar R, Caruso D, Liu X, Wang G, Chen Z, Ji Y, Mao G, Walker SO, Bartell TR, **Ji Z**, Sun Y, Tsai HJ, Pongracic JA, Weeks DE, Wang X. Epigenome-wide association study links site-specific DNA methylation changes with cow's milk allergy. Journal of Allergy and Clinical Immunology. 2016; 138(3):908-911.e9.
- Wang G, Hu FB, Mistry KB, Zhang C, Ren F, Huo Y, Paige D, Bartell T, Hong X, Caruso D, **Ji Z**, Chen Z, Ji Y, Pearson C, Ji H, Zuckerman B, Cheng TL, Wang X. Association Between Maternal Prepregnancy Body Mass Index and Plasma Folate Concentrations With Child Metabolic Health. JAMA Pediatrics. 2016; 170(8):e160845.
- 35 Deng D, Du Y, **Ji Z**, Rao K, Wu Z, Zhu Y, Coley RY. Predicting survival time for metastatic castration resistant prostate cancer: An iterative imputation approach. F1000Research. 2016; 5:2672.
- **Ji Z**, Vokes SA, Dang CV, Ji H. Turning publicly available gene expression data into discoveries using gene set context analysis. Nucleic Acids Research. 2016; 44(1):e8.

### **Book Chapters**

37 Ji Z, Ji H. Pseudotime Reconstruction Using TSCAN. Methods in Molecular Biology. 2019; 1935:115-124.

38 Zhang J, **Ji Z**, Smith KN. Analysis of TCR  $\beta$  CDR3 sequencing data for tracking anti-tumor immunity. Methods in Enzymology. 2019; 629:443-464.

#### Thesis

39 Ji Z. Statistical Methods for Decoding Gene Regulation in Single Cells. PhD thesis. 2020. Johns Hopkins University.

## Funding Support

### Current

Targeting Ferroptosis in Lethal RB1 Deficient Prostate Cancer (NIH R01CA269211)

Dates: 04/01/2022-03/31/2027. Principal Investigator: Ming Chen.

Role: Co-Investigator

The Duke Senescent Cell Evaluations in Normal Tissues (SCENT) Mapping Center (NIH U54AG075936)

Dates: 09/30/2021-08/31/2026. Principal Investigator: Patty Lee. Amount: \$12.7M.

Role: Data Analysis Core Co-PI

### Pending

High-resolution Spatial Maps of Gene Expression and Gene Regulatory Programs (NIH)

Principal Investigator: Zhicheng Ji.

Role: PI

Benchmarking deconvolution tools for spatial transcriptomics (Chan Zuckerberg Initiative)

Principal Investigator: Zhicheng Ji.

Role: PI

Tensor methods for multi-sample multi-omics single-cell data (Chan Zuckerberg Initiative)

Principal Investigator: Zhicheng Ji, Anru Zhang.

Role: PI

Interrogating the Molecular Basis of Alzheimer's Disease Susceptibility by Spatiotemporal Analysis (NIH)

Principal Investigator: Simon Gregory, Shih-Hsiu Wang, Zhicheng Ji.

Role: Co-PI

### Completed

Duke School of Medicine Whitehead Scholar award

Dates: 2020. Principal Investigator: Zhicheng Ji, PhD. Amount: \$250K.

Role: PI

Translating Duke Health award

Dates: 2020. Principal Investigator: Zhicheng Ji, PhD. Amount: \$50K.

Role: PI

### Software

Palo: Color palette optimization [Github]

SCUBI: Single-cell unbiased visualization [Github]

SCATE: Single-cell ATAC-seq signal extraction and enhancement [Github]

TSCAN: Pseudo-time reconstruction in single-cell RNA-seq analysis [Bioconductor] [Github] [GUI]

SCRAT: Single-cell regulome analysis tool [Github] [GUI]

STIP: State transition inference prediction [Github]

GSCA: Gene set context analysis [Bioconductor] [Github] [GUI]

RMRCE: Regularized maximum rank correlation estimator [Github]

## **Editorial Activities**

### Editor

Guest editor of Biomedicines Special Issue: Single-Cell Genomics in Biomedicines

Guest editor of Frontiers in Genetics: Statistical Genetics and Methodology

### Journal Referee

Nature Methods, Nucleic Acids Research, Nature Communications, Genome Biology, Genomics, Proteomics & Bioinformatics, PLOS Computational Biology, Computational and Structural Biotechnology Journal, PLOS Genetics, Bioinformatics, Bioinformatics, Bioinformatics, Bioinformatics, BMC Medical Genomics, Cancers, Genes, Viruses, Biology, Life, Statistics in Biosciences, Biochemical Genetics

## Teaching

### Postdoctoral Associate

Huifang Ma, Department of Biostatistics and Bioinformatics, Duke University (co-advise with Dr. Anru Zhang)

### PhD Students

Mengyi (Miko) Liu, Computational Biology and Bioinformatics Program, Duke University (co-advise with Dr. Simon Gregory)

Changxin Wan, Computational Biology and Bioinformatics Program, Duke University

Tzu-Chieh (Jackie) Liao, Department of Immunology, Duke University (rotation student)

### PhD Thesis Committee

Qi Gao, Department of Biostatistics and Bioinformatics, Duke University

Arinze Okafor, Department of Cell Biology, Duke University

### Master's Students

Tianbei Zhang, Department of Biostatistics and Bioinformatics, Duke University

Xiaotan Sun, Department of Biostatistics and Bioinformatics, Duke University (co-advise with Dr. Anru Zhang)

Constantine Stavrianidis, Department of Biostatistics and Bioinformatics, Duke University

Dezhao Fu, Department of Biostatistics and Bioinformatics, Duke University

Caiwei Zhong, Department of Biostatistics and Bioinformatics, Duke University

Haotian Zhuang, Department of Biostatistics and Bioinformatics, Duke University

Huimin Wang, Department of Biostatistics and Bioinformatics, Duke University

### **Undergraduate Students**

Brian Du, Computer Science major, Duke University

Wei Wang, Applied Mathematics major, The College of William & Mary

### Guest Lecturer

BIOSTAT 900: Current Problems in Biostatistics. 2020, Duke University

140.688: Statistics for Genomics. 2018-2020, Johns Hopkins University

## **Professional Activities**

### **Academic Services**

Co-chair, SenNet imaging and mapping working group

Reviewer, Duke University School of Medicine Virtual Research Week Poster Session

Reviewer, ASA Section on Statistics in Genomics and Genetics Student Paper Competition

### **Professional Memberships**

Eastern North American Region (ENAR) (2017-present)

International Society for Computational Biology (ISCB) (2018-2019)

American Statistical Association (ASA) (2016-2017)

International Chinese Statistical Association (ICSA) (2016-2017)

American Society of Human Genetics (ASHG) (2015-2016)

International Genetic Epidemiology Society (IGES) (2015-2016)

### Presentations

### **Invited Talks**

Practical Genomics Workshop: From Biology to Biostatistics. Center for Computational Genomics at Johns Hopkins, October, 2021

Integrative analysis of multi-study single-cell RNA-sequencing datasets identifies conserved immune landscape of COVID-19 patients. *Inaugural Duke Center for Human Systems Immunology Virtual Symposium*, August, 2021

Integrative analysis of multi-study single-cell RNA-sequencing datasets identifies conserved immune landscape of COVID-19 patients. Single-cell and Spatial Colloquium, Duke University, March, 2021

Computational Methods for Decoding Gene Regulation in Single Cells. Computational Biology and Bioinformatics Seminar, Duke University, February, 2021

Single-cell RNA-seq, power to analyze at a single cell level and broader view of statistical approach to employ in single cell analysis. *Duke Transplant Infectious Diseases Research Symposium*, November, 2020

### Contributed Talks

Single-cell Unbiased Visualization with SCUBI. Joint Statistical Meeting, August, 2021

Reconstruction of conserved immune landscape in COVID-19 patients via multi-study integrative analysis of single-cell RNA-sequencing data. Cold Spring Harbor meeting on Systems Immunology, April, 2021

Single-cell ATAC-seq signal extraction and enhancement with SCATE. ENAR, March, 2020

Reproducible interactive data visualization and exploration with iXplore. ENAR, March, 2017

Reproducible interactive data visualization and exploration with iXplore. The 10th International Chinese Statistical Association International Conference, December, 2016

TSCAN: Pseudo-time reconstruction and evaluation in single-cell RNA-seq analysis. *Joint Statistical Meeting*, August, 2016

### Posters

Single-cell ATAC-seq signal extraction and enhancement with SCATE. RECOMB/ISCB Conference on Regulatory & Systems Genomics, December, 2018

Turning publicly available gene expression data into discoveries using gene set context analysis. The American Society of Human Genetics Annual Meeting, October, 2015

Turning publicly available gene expression data into discoveries using gene set context analysis. *International Genetic Epidemiology Society Annual Meeting*, October, 2015