LIANG (ADAM) DING

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# EXPERIENCE

**Spatial Genomics, Inc.** March 2024 - Present

*Senior Bioinformatics Scientist Pasadena, CA*

* Spearheaded collaboration with assay scientists to leverage spatial transcriptomics for improving cell- type identification accuracy in human solid tumors and mouse MASH liver, identifying disease-associated spatial niches and immune-tumor cell communications.
* Led development of a novel barcode optimization algorithm using bulk and/or scRNA-seq references to resolve optical crowding in raw image decoding; documented implementation protocols for software

engineers, enhancing decoding efficiency.

* Designed and delivered customer-facing [tutorials and analysis workflows](https://tutorials.spatialgenomics.com/intro.html) (e.g., clustering, annotation, integration, imputation) to support adoption of [GenePS](https://spatialgenomics.com/product/), Spatial Genomics’ imaging platform.
* Translated technical insights into business collateral (brochures, posters) showcasing GenePS applica- tions for key stakeholders.

**Spatial Genomics, Inc.** September 2022 - February 2024

*Bioinformatics Scientist Pasadena, CA*

* Pioneered a deep generative model-based algorithm (autoencoders and Bayesian inference) for spatial niche alignment, correcting transcript misassignments by integrating imaging, spatial coordinates, and expression data.
* Developed spatial data showcases (e.g., [mouse kidney Minerva story](http://kidneyviewer.spatialgenomics.com/)) to demonstrate GenePS capabil- ities at launch.
* Conducted comparative benchmarking of commercial in situ platforms to inform second-generation platform and assay development.
* Designed standardized/custom gene panels and probes for internal/external use.

**St. Jude Children’s Research Hospital** December 2018 - September 2022

*Bioinformatics Research Scientist Memphis, TN*

* Directed development of systems biology tools for multiomics analysis (scRNA-seq, spatial transcrip- tomics, gene networks), including:

[scMINER](https://jyyulab.github.io/scMINER/) (Python/R): Preprocessing, QC, clustering, gene network reconstruction, hidden driver in- ference, and visualization.

[NetBID2](https://jyyulab.github.io/NetBID/) (R): Data-driven network-based Bayesian inference of drivers. [SJARACNe](https://github.com/jyyulab/SJARACNe) (Python/C++): Scalable gene network reverse engineering.

* Applied computational approaches to identify cancer biomarkers/therapeutic targets with direct clinical

relevance.

* Contributed to research funding through co-authorship of peer-reviewed publications and successful grant proposals, demonstrating translational impact.

**St. Jude Children’s Research Hospital** July 2016 - December 2018

*Senior Software Engineer Memphis, TN*

* Developed bioinformatics workflows for clinical/research genomics (WES, WGS), improving patient data analysis pipelines.
* Collaborated with scientists to develop algorithms for multiomics data analysis (e.g., [RNAIndel](https://github.com/stjude/RNAIndel), [Epi-](https://github.com/stjude/Episomizer) [somizer](https://github.com/stjude/Episomizer)).
* Analyzed clinical sequencing data to inform treatment decisions and enhance patient care.

**University of Georgia, Athens, Georgia** September 2011 - June 2016

*Graduate Instructor/Lab Instructor Athens, Georgia*

* Designed/lectured system programming courses (C, Java); developed labs, projects, and assessments for 30+ students.

# EDUCATION

## University of Georgia, Athens, Georgia *July 2016*

Ph.D., Computer Science

## University of Texas - Pan America, Edinburg, Texas *August 2011*

M.S., Computer Science

**Zhengzhou University, Henan, China** *December 2008*

M.S., Applied Mathematics B.S., Applied Mathematics

# TECHNICAL STRENGTHS

**Computer Languages** Python, Rust, R, Bash, C++/C, SQL, Latex, Javascript

**Databases** PostgreSQL, SQLite

**Tools** PyTorch, PyG, Git, Nvidia RAPIDS, Nextflow, Snakemake, Docker

**Single-cell/Spatial analysis** Scanpy, Seurat, scvi-tools, Squidpy, SpatialData, Minerva

**Image analysis** ImageJ/Fiji, OpenCV, Baysor, Cellpose

**Cloud computing** AWS, Microsoft Azure

# PUBLICATIONS

\* First authors contributed equally. *#* Corresponding authors. Sort by year in descending order.

## Peer Reviewed Journals

1. Pan Q.\*, **Ding L.**\*, Hladyshau S., Yao X., et al., Chi H., Yu J.*#* [scMINER](https://www.nature.com/articles/s41467-025-59620-6): a mutual information- based framework for clustering and hidden driver inference from single-cell transcriptomics data. Nature Communications 16, 4305, 2025.
2. McCastlain K.\*, Welsh C.\*, Ni Y.\*, **Ding L.**\*, et al., Yu J., Pounds S., Kundu M.*#* Somatic mitochondrial DNA mutations are a source of heterogeneity among primary leukemic cells. Science Advanced, 2025.
3. Huang X.\*, Li Y.\*, Zhang J.\*, Yan L., Zhao H., **Ding L.**, et al., Yu J.*#*, Yang J.*#* Single-cell systems pharmacology identifies development-driven drug response and combination therapy in B cell acute lymphoblastic leukemia. Cancer cell, 42(4), 2024.
4. Yan K.\*, Condori J., Ma Z., **Ding L.**, Dhungana Y., et al., Gottschalk S.*#*, Yu J.*#* Integrome signatures of lentiviral gene therapy for SCID-X1 patients. Science Advances, 9(40), 2023.
5. Zhang Y.\*, Pool A.H., Wang T., Liu L., Kang E., Zhang B., **Ding L.**, Frieda K., Palmiter R., Oka Y.*#* Parallel neural pathways control sodium consumption and taste valence. Cell, 186(26), 2023.
6. Dong X.\*, **Ding L.**, et al., Chi H., Zhang J., Yu J.*#* [NetBID2](https://www.nature.com/articles/s41467-023-38335-6) provides comprehensive hidden driver analysis. Cell, 14(1), 2023.
7. Chang D.\*, **Ding L.**\*, Malmberg R., Robinson D, Wicker M., Yan H., Martinez A., Cai L.*#* [Optimal learning of Markov *k*-tree topology](https://www.sciencedirect.com/science/article/pii/S277241582200013X). Journal of Computational Mathematics and Data Science, 4(100046), 2022.
8. Shi H.\*, Yan K.\*, **Ding L.**, Qian C., Chi H., Yu J.*#* Network Approaches for Dissecting the Immune System. iScience 23(8), 2020.
9. Silveira A.B., Kasper L.H., Fan Y., et al., **Ding L.**, Zhang J., Finkelstein D., et al., Baker S.J.*#* H3.3 K27M depletion increases differentiation and extends latency of diffuse intrinsic pontine glioma growth in vivo. Acta Neuropathol. 137(4): 637-655, 2019.
10. Kohei K.\*, **Ding L.**, Michael N.E., Stephen V.R., Scott N., John E., Juncheng D., Soheil M., Rhonda E.R., Michael R., Zhang, J.*#* [RNAIndel](https://academic.oup.com/bioinformatics/article/36/5/1382/5583731): discovering somatic coding indels from tumor RNA-Seq data. Bioinformatics, btz753, 2019.
11. Silveira A.B.\*, Kasper L.H., Fan Y., Jin H., Wu G., Shaw T., Zhu X., Larson J.D., Easton J., Shao Y., Yergeau D.A., Rosencrance C., Boggs K., Rusch M.C., **Ding L.**, Zhang J., et al., Zhang J., Baker S.J.*#* H3.3 K27M depletion increases differentiation and extends latency of diffuse intrinsic pontine glioma growth in vivo. Acta Neuropathologica. 137(4):637-655, 2019.
12. Xu K.\*, **Ding L.**, Chang T.C., et al., Baker S.J., Wu G.*#* Structure and evolution of [double](https://link.springer.com/article/10.1007/s00401-018-1912-1) [minutes](https://link.springer.com/article/10.1007/s00401-018-1912-1) in diagnosis and relapse brain tumors. Acta Neuropathologica. 137(1):123-137, 2019.
13. Alexander T.B.\*, Gu Z.\*, Iacobucci I.\*, Dickerson K., Choi J.K., Xu B., et al., **Ding L**, Liu Y., Zhang J., et al., Inaba H.*#*, Mullighan C.G.*#* The genetic basis and cell of origin of mixed phenotype acute leukaemia. Nature, 562(7727):373-379, 2018.
14. Mohebbi M.\*, **Ding L.**, Malmberg R.L., Momany C., Rasheed K., Cai L.*#* Accurate prediction of human miRNA targets via graph modeling of miRNA-target duplex. Journal of Bioinformatics and Computational Biology, 7:1850013, 2018.
15. **Ding L.**\*, Xue X., LaMarca S., Mohebbi M., Samad A., Malmberg R.L.*#*, Cai L.*#* Accurate Prediction of [RNA Nucleotide Interactions](https://academic.oup.com/bioinformatics/article/31/16/2660/321332) with Backbone *k*-Tree Model. Bioinformatics, 31(16): 2660-2667, 2015.

## Conference Proceedings

1. **Ding L.**\*, Xue X., LaMarca S., Mohebbi M., Samad A., Malmberg R.*#*, and Cai L.*#* (2014) Abinitio prediction of RNA nucleotide interactions with backbone k-tree model, Proceedings of ECCB’14 Workshop on Computational Methods for Structural RNAs, Strasbourg France, 25-42.
2. **Ding L.**\*, Samad A., Xue X., Huang X., Malmberg R.*#*, and Cai L.*#* (2014) Stochastic k- tree grammar and its application in bimolecular structure modeling, International Conference on Language and Automata Theory and Applications (LATA 2014) Vol 8370, 308-322.
3. **Ding L.**\*, Robertson J., Malmberg R.*#*, and Cai L.*#* (2013) Protein closed loop prediction from contact probabilities, International Symposium on Bioinformatics Research and Applications (IS- BRA 2013), 199-210.
4. **Ding L.**, Fu B., and Zhu B. (2011) Minimum Interval Cover and Its Application to Genome Sequencing. International Conference on Combinatorial Optimization and Applications (COCOA 2011), 287- 298.
5. **Ding L.**, Fu B., and Fu Y. (2010) Improved Sublinear Time Algorithm for Width- bounded Separators. International Workshop on Frontiers in Algorithmics, 101-112.
6. **Ding L.**, Fu B., Fu Y., Lu Z., and Zhao Z. (2010) *O*((*logn*)*2*) Time Online Approximation Schemes for Bin Packing and Subset Sum Problem. International Workshop on Frontiers in Algorithmics, 250-261.

## Book Chapters

1. Cai L., **Ding L.**, Huang X., Malmberg R.L., and Xue X. (2014) Stochastic grammar systems for biomolecular structure modeling, Integrative Bioinformatics for Biomedical Research: A No Boundary Thinking Approach, Huang and Moore ed. Cambridge Press, to appear.
2. **Ding L.**, Fu B. (2013) Algebrization and Randomization Methods. Handbook of Combinatorial Optimization, Springer, pp 171-220.
3. **Ding L.**, Fu B., Fu Y., Wan Y. (2012) Application of Width-Bounded Separators to Protein Side- Chain Packing Problem, Sequence and Genome Analysis: Methods and Applications, iConcep Press.

## Preprints

1. Michael R., **Ding L.**, Sasi A., Andrew T., Hongjian J., Michael M., Lawryn K., Andre S., Michael A.D., Suzanne J.B., Zhang J.. XenoCP: Cloud-based BAM cleansing tool for RNA and DNA from Xenograft. bioRxiv 843250.
2. **Ding L.**, Samad, A., Xue, X., Huang, X., Cai, L. (2013) Polynomial kernels collapse the W- hierarchy. arXiv preprint arXiv:1308.3613.

# HONORS AND AWARDS

* + Ovation Silver Award, St. Jude Children’s Research Hospital, 2018
  + Dissertation Completion Award, Graduate School, University of Georgia, Athens, Georgia, 2015
  + Outstanding Graduate Teaching Assistant Award, Graduate School, University of Georgia, Athens, Georgia, 2015
  + CUDA and GPU Programming Certificate CUDA, Department of Computer Science, University of Georgia, 2013
  + Outstanding Student Scholarship, Zhengzhou University, Henan, China, 2005