#### Postdoctoral Research Associate @ Yale University

### Haikuo Li

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#### **EDUCATION**

**Doctor of Philosophy, Biological and Biomedical Sciences (Molecular Genetics and Genomics)** 8/2019 – 12/2023 Washington University in St. Louis, MO, United States

- Thesis mentor: Benjamin D. Humphreys, M.D., Ph.D.
- Thesis committee: Ting Wang, Samantha Morris, Jeffrey Millman, Allegra Petti, Michael Meers

#### Bachelor of Science, Bioscience (Zhiyuan Honors Program)

9/2015 - 6/2019

Shanghai Jiao Tong University, Shanghai, China

- 2019 Top 1% Bachelor Thesis: Rank #1 in the program
- 2019 Outstanding Graduate in Bachelor Degree, Shanghai

#### **WORK EXPERIENCE**

# Postdoctoral Research Associate Yale University | Department of Biomedical Engineering | Rong Fan Lab Visiting Student 6/2018 – 4/2019

Yale University | Department of Immunobiology | Aaron Ring Lab

#### **PUBLICATIONS**

#### First-author research articles

- <u>Li, H.</u>, Li, D., Ledru, N., Xuanyuan, Q., Wu, H., Asthana, A., Byers, L.N., Tullius, S.G., Orlando, G., Waikar, S.S. and Humphreys, B.D., 2024. Transcriptomic, epigenomic, and spatial metabolomic cell profiling redefines regional human kidney anatomy. *Cell Metabolism*, 36(5), pp.1105-1125. (URL)
- Featured Content of the *Cell Metabolism* journal website
- <u>Li, H.</u>, Dixon, E.E., Wu, H. and Humphreys, B.D., 2022. Comprehensive single-cell transcriptional profiling defines shared and unique epithelial injury responses during kidney fibrosis. <u>Cell Metabolism</u>, 34(12), pp.1977-1998. (URL)
- Research featured by NIH NIDDK annual report (URL)
- Research Highlights by Nature Review Nephrology (URL)
- Research Highlights by *Kidney International (URL)*
- 3. <u>Li, H.\*</u>, Li, D.\* (co-first author) and Humphreys, B.D., 2024. Chromatin conformation and histone modification profiling across human kidney anatomic regions. *Scientific Data*, 11(1), p.797. (URL)
- Highlighted in the collection of *Data for epigenetics research* (URL)
- 4. <u>Li, H.</u> and Humphreys, B.D., 2024. Spatially resolved metabolomic dataset of distinct human kidney anatomic regions. *Data in Brief*, 54, p.110431. (URL)

- 5. <u>Li, H.</u> and Humphreys, B.D., 2024. Protocol for multimodal profiling of human kidneys with simultaneous high-throughput ATAC and RNA expression with sequencing. *STAR Protocols*, 5(3), p.103049. (URL)
- 6. <u>Li, H.</u> and Humphreys, B.D., 2022. Mouse kidney nuclear isolation and library preparation for single-cell combinatorial indexing RNA sequencing. *STAR Protocols*, 3(4), p.101904. (URL)
- Ku, X.\*, Wang, J.\*, <u>Li, H.\*</u> (co-first author), Meng, C., Yu, F., Yu, W., Li, Z., Zhou, Z., Zhang, C., Hua, Y. and Yan, W., 2023. Proteomic portrait of human lymphoma reveals protein molecular fingerprint of disease specific subtypes and progression. <u>Phenomics</u>, 3(2), pp.148-166. (URL)
- 8. Wang, J.\*, Ku, X.\*, Ma, Q.\*, <u>Li, H.\*</u> (co-first author), Huang, S.\*, Mao, L., Yu, F., Jin, J. and Yan, W., 2024. Hsa\_circ\_0007099 and PIP4K2A coexpressed in diffuse large B-cell lymphoma with clinical significance. *Genes & Diseases*, 11(4), p.101056. (URL)

#### First-author review/commentary articles

- 9. <u>Li, H.</u> and Humphreys, B.D., 2024. Multimodal characterization of sexual dimorphism in the mammalian kidney. *Kidney International*, 105(4), pp.653-655. (URL)
- 10. <u>Li, H.</u> and Humphreys, B.D., 2024. Targeting de novo lipogenesis to mitigate kidney disease. <u>The Journal of Clinical Investigation</u>, 134(4). (URL)
- 11. <u>Li, H.</u> and Humphreys, B.D., 2022. New functions for basophils identified in kidney fibrosis. <u>Nature Immunology</u>, 23(6), pp.824-825. (URL)
- 12. <u>Li, H.</u> and Humphreys, B.D., 2021. Single cell technologies: Beyond microfluidics. <u>Kidney360</u>, 2(7), pp.1196-1204. (URL)
- 13. <u>Li, H.</u> and Humphreys, B.D., 2020. Surveying the human single-cell landscape. <u>Kidney International</u>, 98(6), pp.1385-1387. (URL)

#### First-author book chapter

- 14. Muto, Y.\*, <u>Li, H.\*</u> (co-first author) and Humphreys, B.D., 2022. Single Cell Transcriptomics.

  <u>Innovations in Nephrology: Breakthrough Technologies in Kidney Disease Care</u> (pp. 87-102). Cham: Springer International Publishing. (URL)
- Chapter has been translated into German (DOI)

#### Other publications

- 15. Bai, Z., Zhang, D., Gao, Y., Tao, B., Zhang, D., Bao, S., Enninful, A., Wang, Y., <u>Li, H.</u>, Su, G. and Tian, X., Zhang, N., Xiao, Y., Liu, Y., Gerstein, M., Li, M., Xing, Y., Lu, J., Xu, ML. and Fan, R. 2024. Spatially exploring RNA biology in archival formalin-fixed paraffin-embedded tissues. <u>Cell</u> (URL)
- 16. Zhao, C., Rong, K., Liu, P., Kong, K., <u>Li, H.</u>, Zhang, P., Chen, X., Fu, Q. and Wang, X., 2024. Preventing periprosthetic osteolysis in aging populations through lymphatic activation and stem cell-associated secretory phenotype inhibition. *Communications Biology*, 7(1), p.962. (URL)
- 17. Wilson, P.C., Verma, A., Yoshimura, Y., Muto, Y., <u>Li, H.</u>, Malvin, N.P., Dixon, E.E. and Humphreys, B.D., 2024. Mosaic loss of Y chromosome is associated with aging and epithelial injury in chronic kidney disease. <u>Genome biology</u>, 25(1), p.36. (URL)
- 18. Cheng, Z. et al. (special journal issue; author list alphabetically ordered) (2023). RNA-seq: questions and answers. *STAR Protocols*. (URL)

#### RESEARCH EXPERIENCE

## Postdoctoral Research Associate, Rong Fan Lab Department of Biomedical Engineering, School of Engineering & Applied Science, Yale University

• Single-cell and spatial genomics technology development

• Investigating mechanisms of gene expression regulation

#### Research Associate, Benjamin Humphreys Lab

12/2023 - 1/2024

2/2024 - present

Division of Nephrology, Washington University in St. Louis

#### PhD Student, Benjamin Humphreys Lab

4/2020 - 12/2023

Division of Nephrology, Washington University in St. Louis

- Multiomics characterization of kidney physiology: developing an anatomically stratified human kidney atlas with single-cell combinatorial indexing multiomics sequencing
- Multiomics characterization of kidney pathology: studying mouse models of kidney fibrosis with single-cell combinatorial indexing RNA-seq and spatially resolved transcriptomics
- Studying metabolic mechanisms that drive kidney fibrosis
- Studying cell fate determination in kidney injury & regeneration and regional differentiation
- Method development: single-cell combinatorial indexing (split-pool barcoding) library generation and unique data analysis; computational pipeline of analyzing spatially resolved metabolomics data

#### PhD Rotation Student, Tim Peterson, Sidharth Puram, Benjamin Humphreys Labs

8/2019 - 4/2020

Washington University in St. Louis

- Peterson Lab: Understanding the intracellular effects of Cationic Amphipathic Drugs on organelles
- Puram Lab: Studying head and neck cancer by CITE-seq
- Humphreys Lab: Characterizing kidney injury and repair markers by RNAscope

#### Visiting Student, Aaron Ring Lab

6/2018 - 4/2019

#### Department of Immunobiology, Yale University

- Modulating immune cytokines by protein engineering
- Bench supervisor: Ting Zhou, Ph.D.

#### Undergraduate Researcher, Wei Yan Lab

9/2016 - 7/2018

#### Shanghai Center for Systems Biomedicine, Shanghai Jiao Tong University

- Identification of biomarkers of lymphoma with mass spectrometry; clinic proteomics
- Bench supervisor: Xin Ku, Ph.D.

#### Summer Intern, Manyuan Long Lab

6/2017 - 8/2017

#### Department of Ecology and Evolution, The University of Chicago

• Identification of mammalian positively selected genes by polygenetic analysis

#### Science Olympiad (Mathematics), Shandong Province Team, China

7/2014 - 2/2015

• Top10 students selected to participate in the Chinese Mathematics Olympics

2/2010 4/2

#### **SOFTWARE DEVELOPMENT**

MALDIpy: Single-cell analysis of MALDI-MS imaging mass spectrometry data (first contributor)

• pip install MALDIpy (https://pypi.org/project/MALDIpy/)

#### **SKILLS**

#### Wet lab experiment

- Extensive experience in spatial genomics library generation including spatial-RNA-seq (DBiT-seq), spatial-ATAC-seq, spatial-CUT&Tag and spatial co-profiling
- Extensive experience in single-cell and single-nucleus library generation from diverse technologies, including 10X Genomics, sci-RNA-seq, SHARE-seq and INTACT, as well as multimodal profiling including RNA-seq, ATAC-seq, Hi-C, CUT&RUN, CARLIN and CITE-seq
- Extensive experience in molecular biology technologies such as cloning, vector construction, qPCR, immunohistology, and in-situ hybridization
- Extensive experience in tissue culture including primary cell isolation, immunocytochemistry and Seahorse metabolic measurement
- Strong experience in animal work such as mouse kidney disease surgery (UUO/IRI) and tumor implantation
- Strong experience in clinical sample management and processing such as human kidney dissection
- Strong background in protein chemistry including mass spectrometry sample preparation & recombinant protein preparation and protein liquid chromatography

#### Computational workflow

- Extensive experience in using Python, R, Shell and Jupyter
- Extensive experience in single-cell sequencing data preprocessing and analysis including UMAP visualization, data integration, sample demultiplexing, cell trajectory interference, fate mapping, gene activity prediction and multimodal analysis at the million-cell level.
- Extensive experience in spatial transcriptomics and epigenomics data preprocessing and analysis
- Extensive experience in spatially resolved transcriptomics and metabolomics analysis
- Strong experience in analysis of bulk RNA-seq, proteomics and metabolomics data
- Extensive experience in data mining and discovering biological insights
- Strong training background in mathematics

#### TEACHING EXPERIENCE

#### 1. Trainee Supervisory Experience

#### Rotation Project Bench Mentor, Washington University in St. Louis

9/2023-11/2023

Rachel W. (Ph.D. student in CSB/DBBS): Functional analysis of human kidney single-cell multiomics

#### Peer Study Mentor, Washington University in St. Louis

3/2023-6/2023

Jenna U. (Ph.D. student in MGG/DBBS): Genomics (Bio5488) and Python-based coding tutoring

#### Rotation Project Bench Mentor, Washington University in St. Louis

12/2022-1/2023

• Qiao X. (Ph.D. student in BIDS/DBBS): Spatially resolved metabolomics analysis and package development

#### Bench Mentor, Washington University in St. Louis

9/2022-7/2023

• Dian L. (Ph.D. student in CSB/DBBS): Single-cell multimodal integration on human kidney physiology

#### Peer Study Mentor, Washington University in St. Louis

1/2022 - 6/2022

Julie C. (Ph.D. student in HSG/DBBS): Genomics (Bio5488) and Python-based coding tutoring

#### 2. Course Instructor Assistant

#### Assistant Instructor, Washington University in St. Louis

8/2022 - 6/2023

• Structural bioinformatics of proteins (Bio4525)

#### Assistant Instructor, Washington University in St. Louis

1/2021 - 6/2021

2023

• Genomics (Bio5488)

#### Undergraduate Teaching Assistant, Shanghai Jiao Tong University

- College Genetics Course (2/2019–6/2019)
- College Macrobiology Course (2/2018–6/2018)
- College Biochemistry Course (9/2017–1/2018)

#### PRESENTATIONS & POSTERS

Oral Talk Speaker, American Society of Nephrology Annual Meeting (ASN Kidney Week)		
Retreat Talk (MGG/CSB/HSG/IMSD programs), Washington University in St. Louis	9/2023	
POSTER (PDF)   PhD program retreat, Washington University in St. Louis		
• Transcriptomic, epigenomic and spatial metabolomic cell profiling redefines regional human kidney anatomy		
Project Talk, CZI Single-Cell Biology Annual Meeting	11/2022	
Speaker, Nephrology Division Research Seminars, Washington University in St. Louis	9/2022	
Retreat Talk (MGG/CSB/HSG/IMSD programs), Washington University in St. Louis	9/2022	
POSTER (PDF)   PhD program retreat, Washington University in St. Louis	9/2022	
Cell profiling defines metabolic dysregulation in kidney fibrosis		
DBBS Friday Talks (MGG/CSB/HSG/IMSD programs), Washington University in St. Louis		
REBUILDING A KIDNEY Spring Meeting   lighting talk	4/2022	
PhD Program Thesis Committee Meeting, Washington University in St. Louis 8/2021, 4/2022, 4/2023, 12/20		
REBUILDING A KIDNEY Work in Progress   small group meeting	12/2021	
PhD Program Qualifying Examination Committee Meeting	9/2020	
POSTER (PDF)   SJTU Academic Festival (Best Poster Award)	2/2019	
<ul> <li>Modulating the Tumor-Targeting Specificity of "Decoy-Resistant" Interleukin-18 by Protein Engineering</li> </ul>		
POSTER (PDF)   Human Proteome Organization World Congress	10/2018	
• Clinical Proteomics Analysis using Data Independent Acquisition-Mass Spectrometry (DIA-MS) Identified Classifiers		
for Molecular Characterization of Lymphoma		
POSTER (PDF)   SJTU Academic Festival (Best Poster Award)	12/2017	

#### **HONORS & ACTIVITIES**

Award for Outstanding Students Abroad [highest award granted by the Chinese government to students overseas (LINK)]

Detecting Positively Selected Genes among Mammalian Species Using Phylogenetic Analysis of Maximum Likelihood

Member, NCFDD (National Center for Faculty Development & Diversity)	2023 - present
Member, ASN (American Society of Nephrology)	2020 - present
Top 0.2% Bachelor Thesis of Shanghai Jiao Tong University (URL) (Thesis PDF)	2019
Outstanding Graduate in Bachelor Degree, Shanghai	2019
Academic Excellence Scholarship (First-class), Shanghai Jiao Tong University	2016, 2017, 2018
Rank #1 Student Presentation, National Biology Education Conference of Chinese Ministry of Educa	tion 2018
Vice President, Students' Union of Zhiyuan Honors Program, Shanghai Jiao Tong University	2017 - 2018
Top 0.1% in Chinese University Entrance Examination (681 points)	2015
Bronze medal, Chinese Mathematical Olympiad (CMO)	2014