Shang-Yang (Sam) Chen

Computational Biologist || Data Scientist

sychen9584@gmail.com || Boston, MA || sychen9584.github.io

Profile

Computational Biology Ph.D. with extensive expertise in analyzing and developing tools for high-throughput genomic data, with a specialization in single-cell data analysis. Demonstrated expertise in innate immunology, leveraging advanced bioinformatics techniques to uncover biological insights and drive impactful research. Skilled in developing innovative solutions to biological problems, including custom pipelines and web-based applications. Passionate about integrating AI/machine learning approaches to advance data analysis, predictive modeling, and biomedical research outcomes.

GPA: 3.89

Education

Northwestern University, Chicago, IL

2016-2022

Ph.D. in Computational Biology

DGP Integrated Biomedical Sciences Program - Biomedical Informatics Track

"Characterizing Macrophage Heterogeneity in Tissues Through High-Throughput Transcriptomics Technologies and Algorithms"

Stony Brook University, Stony Brook, NY

2012-2016

Bachelor of Science in Biochemistry, honors degree GPA: 3.69

Minor in Information Systems

Research and Work Experiences

Scientist II, Computational Science Scientist I, Computational Science

February 2024 - August 2024

June 2022 - January 2024

Dyno Therapeutics

- ❖ Designed and deployed production-grade Python pipelines for automated processing, QC, and analysis of single-nucleus RNA-seq (snRNA-seq) data, enabling scalable analysis of 1.5M+ nuclei per dataset and reducing analysis time by 75%.
- ❖ Served as primary analysis lead for snRNA-seq data across ocular, brain, and liver programs, supporting evaluation and validation of AI-designed AAV gene therapy vectors, contributing to two product launches and a licensing agreement.
- ❖ Performed integrative analysis of bulk NGS datasets from non-human primates, mice, and in vitro assays to guide design and selection of high-performing AAV variants.
- ❖ Collaborated with biologists, engineers, and leadership to optimize experimental design, scale data pipelines, and support IP filings and conference presentations.

Bioinformatics Internship

October 2021 – April 2022

Genentech, Oncology Informatics

- ❖ Developed the R package Epiregulon, published on Bioconductor, enabling inference of transcription factor (TF) activity in single cells via gene regulatory network construction; integrated scATAC-seq, scRNA-seq, and public ChIP-seq data to support therapeutic discovery targeting TFs in oncology.
- Designed and implemented a benchmarking framework to evaluate Epiregulon against reference TF-target gene databases, ensuring rigorous model validation and robust performance assessment.

❖ Analyzed and interpreted scATAC-seq datasets to identify transcriptional regulatory mechanisms in breast cancer subtypes, supporting early oncology discovery efforts.

Ph.D. Candidate Summer 2017-May 2022

Deborah Winter, Northwestern University

Characterizing Macrophage Heterogeneity in Tissues Through High-Throughput Transcriptomics Technologies and Algorithms

- ❖ Led integrative, multi-omics analyses (scRNA-seq, bulk RNA-seq, CITE-seq, ATAC-seq) to characterize immune cell heterogeneity and inflammatory pathways in rheumatoid arthritis and liver cholestasis.
- ❖ Designed and implemented custom R pipelines for scRNA-seq and CITE-seq, enabling high-resolution profiling of immune cell diversity.
- ❖ Developed and deployed MAGNET, a web-based tool for functional enrichment analysis using curated immunology gene sets, adopted by research teams for hypothesis generation and data interpretation.
- Awarded the American Heart Association (AHA) predoctoral fellowship in recognition of research excellence in computational immunology.

Publications

A. Peer-reviewed Original Investigations

MAGNET: A Web-based Application for Improved Functional Enrichment Analysis of Cell-type-specific Data. SY Chen, G. Gadhvi, D. Winter. PLoS One; January 2023; 18(1):e0272166

Tissue-resident, extravascular Ly6c- monocytes are critical for inflammation in the synovium. A. Montgomery, **SY Chen**, Y. Wang, G. Gadhvi, M. Mayr, C. Cuda, S. Dominguez, H-K M. Makinde, M Gurra, A. Misharin, A. Mandelin, E. Ruderman, A. Thakrar, S. Brar, M. Carbs, K. Aren, M. Akbarpour, A. Filer, S. Nayer, A. Teososio, T. Major, A. Bharat, GR. Budinger, D. Winter, H. Perlman. Cell reports; May 2023; 42 (5)

Epiregulon: Single-cell transcription factor activity inference to predict drug response and drivers of cell states. T. Włodarczyk, A. Lun, D. Wu, M. Shi, X. Ye, S. Menon, S. Toneyan, K. Seidel, L. Wang, J. Tan, SY Chen, T. Keyes, A. Chlebowski, A. Waddell, W. Zhou, Y. Wang, Q. Yuan, Y. Guo, LF Chen, B. Daniel, A. Hafner, M. He, A. Chibly, Y. Liang, Z. Duren, C. Metcalfe, M. Hafner, C. Siebel, MR Corces, R. Yauch, S. Xie, X. Yao. Nature Communications 16, 7118 (2025).

Synovial Macrophage Heterogeneity Confers Differential Response to Acute and Chronic Inflammatory Arthritis. SY Chen, A. Montgomery, P. Homan, S. Dominguez, C. Cuda, G. Gadhvi, S. Jung, GR Budinger, A. Misharin, M. Mayr, D. Winter, H. Perlman. (*In preparation*)

Critical role of synovial tissue-resident macrophage niche in joint homeostasis and suppression of chronic inflammation. QQ Huang, R. Doyle, SY Chen, Q. Sheng, AV Misharin, Q. Mao, D. Winter, RM Pope. Science Advances; January 2021; 7(2):eabd0515.

Transcriptional analysis of murine biliary atresia identifies macrophage heterogeneity and subset-specific macrophage functions. Gromer KD, Chen SY, Gadhvi G, Feng L, Shearn C, Antala S, Wechsler JB, Cuda CM, Mack CL, Sokol RJ, Janssen WJ, Green RM, Perlman H, Winter DR, Taylor SA.Front Immunol. 2025 Jan 30;16:1506195

Transcriptional profiling of pediatric cholestatic livers identifies three distinct macrophage populations. S. Taylor, **SY Chen**, G. Gadhvi, L. Feng, K. Gromer, H. Abdala-Valencia, K. Nam, S. Dominguez. A. Montgomery, P. Reyfman, L. Ostilla, J. Wechsler, C. Cuda, R. Green, H. Perlman, D. Winter. PLoS One; January 2021;16(1):e0244743.

Macrophages drive the inflammatory phase in experimental osteoarthritis

A Montgomery, N Fahy, S Hamilton, B Eckman, L De Almeida, S Ishihara, MG Mayr, SY Chen, G Gadhvi, C Cuda, AM Malfait, Y Bastiaansen-Jenniskens, DR. Winter. BioRxiv; June 2020.

Influenza-induced activation of recruited alveolar macrophages during the early inflammatory phase drives lung injury and lethality.

CM. Koch, KR Anakella, YS Hu, M Ciesielski, G Gadhvi, **SY Chen**, M Turner, Y Cheng, BM Coates, H Abdala-Valencia, PA Reyfman, AV Misharin, GR Budinger, DR Winter, KM Ridge. BioRxiv; June 2020

Maintenance DNA methylation is essential for regulatory T cell development and stability of suppressive function. KA. Helmin, L Morales-Nebreda, MA. Torres Acosta, KR. Anekalla, SY Chen, H Abdala-Valencia, Y Politanska, P Cheresh, M Akbarpour, EM. Steinert, SE. Weinberg, BD. Singer; Journal of Clinical Investigation; September 2020

DNA methylation as a regulator of the neonatal CD4+ T cell response to pneumonia McGrath-Morrow SA, Ndeh R, Helmin KA, Chen SY, Anekalla KR, Abdala-Valencia H, Collaco JM, D'Alessio FR, Misharin AV, Singer BD. Journal of Biological Chemistry; July 2018; JBC-RA118

B. Abstracts

Optimizing Intravitreal Delivery to the Non-Human Primate Retina with Machine Guided AAV Capsid Design. S. Abesteh, I. Brenckle, Z. Cargill, I. Chen, S. Chen, D. Deny, J. Fathima, J. Gerold, A. Ghosh, J. Grush, E. Kelsic, H. Kuchwara, J. Kwasnieski, K. Lin, E. Lykken, E. Marrogi, K. Maryak, P. McDonel, H. Levitin, A. Miles, N. Nagabhushana, D. Pignatta, B. Satinsky, B. Saxena, C. Sengel, A. Sheridan, T. Tarbell, M. Thommes, H. Turunen, N. White, S. Wolock 2024 ASGCT Annual Meeting

Applying Artificial Intelligence to Multi Property Optimization of AAV Capsids for Neuronal Gene Delivery. M. Deshpande, D. Bloodgood, B. Diaz-Rohrer, K. Looby, S. Hilton, S. Chen, S. Sinai, K. Lin, J. Kwasnieski, A. Veres, E. Kelsic 2023 ASGCT Annual Meeting

Non-Human Primate Evaluation of an Engineered AAV Capsid for Retinal Cell Specific and Biofactory-Based Ocular Gene Therapies. H.i Turunen, A. Miles, B. Diaz-Rohrer, S. Hilton, S. Chen, K. Maryak, S. Sinai, K. Lin, J. Kwasnieski, A. Veres, E. Kelsic 2023 ASGCT Annual Meeting

Characterizing Heterogeneity of Synovial Macrophages in Rheumatoid Arthritis Patients. SY Chen, Y. Wang, A. Montgomery, S. Dominguez, C. Cuda, G. Gadhvi, H. Perlman, D. Perlman 2020 ACR Convergence 2020

The Dynamics of Macrophage Sub-Populations in the Inflammatory Phase Following Joint Trauma. S. Hamilton, A. Montgomery, N. Fahy, M. Mayr, SY Chen, G. Gadhvi, Y. Bastiaansen-Jenniskens, D. Winter 2020 ACR Convergence 2020

Single Cell RNA-sequencing Reveals Distinct Macrophage Subsets in the Joint with Differing Ontogenies During Steady-state and Arthritis

SY Chen, A Montgomery, P Homan, G Gadhvi, D Winter, H Perlman 2019 ACR/ARHP Annual Meeting

A Novel Subclass of Intravascular Non-classical, Tissue Resident Synovial Monocyte Is Critical for Rheumatoid Arthritis Pathogenesis

A Montgomery, **SY Chen**, D Winter, H Perlman 2019 ACR/ARHP Annual Meeting

The Role of Flip in Differentiation and Survival of Synovial Tissue Resident Macrophages Q Huang, R Doyle, SY Chen, A Misharin, D Winter, R Pope 2019 ACR/ARHP Annual Meeting

Tissue Resident Macrophages Establish a Niche That Limits Monocyte to Macrophage Differentiation in Synovial Tissue during Homeostasis

Q Huang, R Doyle, A Misharin, **SY Chen,** D Winter, R Pope 2018 ACR/ARHP Annual Meeting

Presentations

Poster Presentation, Keystone Symposia: Single Cell Biology, Florence (Cancelled)

May 2020

- Unmasking the heterogeneity of joint macrophages in health and disease using single-cell RNA-sequencing

Oral Presentation, American College of Rheumatology Annual Meeting, Atlanta November 2019 Poster Presentation, Cell Symposia: Transcriptional Regulation, Chicago October 2019

- Single cell RNA-sequencing reveals ontogenically-distinct macrophage subsets in the joint during steady state and arthritis

Poster Presentation, Lewis Landsberg Research Day, Northwestern University April 2019

- MAGNET: Macrophage Annotation of Gene Network Enrichment Tool

Poster Presentation, Lewis Landsberg Research Day, Northwestern University April 2018

- Transcriptional profiling of synovial macrophages in arthritic mice

Poster Presentation, Computational Research Day, Northwestern University April 2017

- Building a Bisulfite Sequencing Data Analysis Workflow

Poster Presentation, URECA, Stony Brook University April 2016

- Seasonal Variation in Lifespan and Starvation Resistance in *D. melanogaster*

Academic Honors & Fellowships

American Heart Association Predoctoral Fellowship

2019 - 2021

Project: Elucidating the Roles of Resident Cardiac Macrophages in Post-Infarction Reperfusion Injury

OSG User School 2017, University of Wisconsin-Madison

July 2017

Workshop on high-performance computing clusters

Selected to represent Northwestern University with full funding support

URECA Summer Research Grant

Summer 2015

Project: Comparison of seasonal variation in starvation resistance between *D. melanogaster* and *D. simulans*.

Tsai Hsing School Foundation Fellowship

2012 - 2016

Provided full tuition and living expenses support for undergrad studies at Stony Brook University

Early Career Experiences

Research Rotations 2016 - 2017

Northwestern University

- ❖ Conducted *in silico* benchmarking of RNA-seq library preparation protocols and collaborated with the sequencing core to evaluate Lab Information Management Systems (LIMS).
- ❖ Developed computational pipelines, including automation scripts for bisulfite sequencing workflows.
- ❖ Built an R-based probabilistic model to simulate and infer cell lineage trees from enhancer profiles.

Undergraduate Researcher

2015 - 2016

Stony Brook University

Comparison of seasonal variation in starvation resistance between D. melanogaster and D. simulans.

❖ Investigated seasonal variation in starvation resistance and sexual polymorphism in *D. melanogaster* and *D. simulans*, leading a project on pupation behavior.

Summer Lab Intern Summer 2013

Taipei Medical University

Assisted in lab mouse care and performed cell culture, PCR, and gel electrophoresis on human cancer cell lines.

Professional Organizations

Member of the International Society for Computational Biology (ISCB)	2017-present
Member of the American Society of Gene & Cell Therapy	2022-2024
Member of the Midwest Taiwanese Biotechnology Association (MTBA)	2019-2023
Member of the American College of Rheumatology (ACR)	2019-2022
Member of the American Heart Association (AHA)	2017-2022

2017 2021

Skills & Relevant Courses:

- <u>Skills:</u> Bioinformatics, Single Cell Omics, Epigenomics, Biostatistics, Data Analytics, Data Visualization, ML/AI, Webapp Development, Version Control, Immunology, Genomics, Cloud Computing
- **Programming Languages/Platforms:** Python, R, UNIX, Java, SQL
- <u>Framework/Packages:</u> Seurat, Scanpy, git, Pandas, Tidyverse, DESeq2, ggplot2, Matplotlib, Seaborn, Scikit-Learn, pyTorch, Django, AWS, Nextflow, BioPython

Certification & Training

Data Structures & Algorithms GTx	2025
Object-Oriented Programming with Java GTx	2025
DeepLearning.AI Deep Learning Specialization	2024
DeepLearning.AI Machine Learning Specialization	2024
Northwestern Kellogg School of Management	
Management for Scientists and Engineers Certificate	2020
Datacamp Coding Best Practices with Python	2020
Datacamp Tidyverse Fundamentals	2020
National Taiwan University Summer Python Bootcamp	2016