JIACHEN AI

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EDUCATION

University of California, Los Angeles, Fielding School of Public Health, CA

| GPA: 4.00/4.00

Advisor: Dr. Matteo Pellegrini

Master of Science in Biostatistics

Southwest University, School of Life Sciences, China

Sept. 2018 – June 2022

Sept. 2023 – Current

Bachelor of Science in Biological Sciences | GPA: 3.84/4.00 (Ranked 2/65)

Outstanding Graduate

Thesis: Analytic Hierarchy Process for Invasive Plant Management

HONORS

Warren Alpert Computational Biology and AI Network Fellow	2024-2025
Fielding Dean's Leadership Grant	2023-2025
Steve Wallace Policy Fellowship	2024
APEx Funding, UCLA Human Genetics	2024
International Research Award, Southwest University	2021
Second- and Third-Class Academic Excellence Scholarships, Southwest University	2019, 2020, 2021

RESEARCH

Influenza Vaccination: DNA Methylation and Immune Response Mechanisms

Los Angeles, CA

Supervised by Dr. Matteo Pellegrini

Mar. 2024 – Current

- Established a multivariate prediction model with LOOCV to link targeted bisulfite sequencing DNA methylation profiles to sero-protection (day 0, day 28) and sero-conversion (day 28) post-influenza injection, identifying key CpG sites and RIG-I signaling pathway as the key regulator of vaccine-induced immunity.
- Integrated DNA methylation, bulk **RNA-seq**, and **cytokine** data through **eQTL** and co-expression analyses to uncover **regulatory mechanisms** and potential therapeutic targets.

Nonlinear Aging Dynamics in Co-methylation Network

Los Angeles, CA

Graduate Student Researcher | Supervised by Dr. Feng Gao

Sept. 2024 – Current

Analyzed non-linear **DNA methylation** patterns in **aging** mice colon tissue, identifying **graph-based aDMR clusters** linked to gene expression and pathways, and uncovering age-related functional changes through gene set enrichment analysis.

Inferring Immune Histories from SARS-CoV-2 RNA-seq Data

Los Angeles, CA

Supervised by Dr. Vanessa D. Jonsson

Sept. 2024 – Current

Designed a **pipeline** integrating immune cell deconvolution, **metagenomic annotation**, TCR/BCR repertoire analysis, and HLA-epitope mapping from SARS-CoV-2 RNA-seq data to infer immune histories, validated against publicly available antibody datasets.

Single-Cell RNA-Seq for Cell Type Annotation

Los Angeles, CA

Supervised by Dr. Jingui Jessica Li

May 2024 – June 2024

Analyzed **single-cell RNA-seq** data from human bone marrow and pancreas to identify cell types through **Seurat**based preprocessing, dimensional reduction, clustering, and **biomarker** identification.

Statistical Analysis for Biology and Biomedicine Research

Chongging, China

Undergraduate Research Assistant

Jun. 2021 – Jan. 2023

- Modelled random walks using **MCMC** to analyze time-homogeneous, recurrent, and transient properties in biology.
- Identified significant stage-specific effects of irrigation on rice yield across five growth stages by **multivariate linear regression**.
- Applied **logistic regression** to identify dietary habits, BMI, and stress as significant factors influencing semen quality.

Lymphocyte Regeneration after Thymocyte Damage in Zebrafish

Chongging, China

Supervised by Dr. Li Li

Oct. 2019 – June 2020

- Established a zebrafish thymocyte damage model with 2 mM Metronidazole, revealing the CXCL12/CXCR4 signaling pathway as critical for T-lymphocyte regeneration through WISH, confocal microscopy, and RT-qPCR.
- Demonstrated that pathway inhibition with WZ811 significantly impaired lymphocyte recovery.

PUBLICATION

- Ai, J. (2021) Lifestyle does affect semen quality: factor determination in a statistical way. Lecture Notes in *Medicine and Life Sciences*, 4, 163–171. DOI: 10.23977/behdp.2021021.
- Ai, J., Pickering, H., Pellegrini, M. et al. Influenza Vaccination: DNA Methylation and Immune Response Mechanisms. Genome Biology. (forthcoming).
- **Ai, J.**, Fan, Y., Gao, F. et al. Nonlinear aging dynamics in co-methylation network. *Cell Reports*. (forthcoming).

TEACHING

MCD BIO 187AL: Research Immersion Laboratory in Genomic Biology (5 units)

UCLA, Los Angeles, CA

June 2024 – Aug. 2024

Served as *Graduate Teaching Assistant*, leading lab sessions and office hours, demonstrating bioinformatics tools (e.g., NCBI Blast, InterProScan, JPRED4, PSIPRED), and mentoring student final projects.

CONFERENCES

2025 Computational Genomics Summer Institute (CGSI)	UCLA, Los Angeles, CA Aug. 2025
2025 Lange Symposium on Computational Statistics	UCLA, Los Angeles, CA Jan. 2025
10th Annual QCBio Retreat	UCLA, Los Angeles, CA Sept. 2024
2024 Computational Genomics Summer Institute (CGSI)	UCLA, Los Angeles, CA Aug. 2024

POSTERS & PRESENTATIONS

PROFESSIONAL EXPERIENCE

Intern at Western Biomedical Technology Inc.

Mar. 2021 – June 2021

• Designed Python and R pipelines to improve **western blot** and **ELISA** accuracy and managed experiments and biweekly reports across **immunology**, **histopathology**, and **molecular biology**.

Office Assistant at SWU Life Science Administrative Office

Oct. 2018 - Jun. 2021

• Mentored 65 students, facilitating academic advising sessions, and coordinating research opportunities.

Volunteer at SWU Central Library

Chongqing, China | Oct. 2019 – Jan. 2020

Organized a **charity event** for 200+ attendees and supported library operations.

EXPERTISE

Computational & Statistical

- Languages: R (tidyverse, ggplot2, glmnet, RShiny), Python (NumPy, Pandas, Biopython, Scikit-learn, Pybedtools, Matplotlib), MATLAB, SAS, STATA, SQL, HTML/CSS/JavaScript, LaTeX
- Platforms: Linux scripting, Git/GitHub, Hoffman2 cluster, Docker, Cloud Platforms (GCP, AWS, Azure, OCI)
- Data Analysis & Modeling: ML, Bayesian inference, deep learning (Keras, TensorFlow, PyTorch), simulation, linear models, optimization, graph theory
- High-Performance Computing (HPC): Workflow orchestration, parallelized analysis, cluster-based computations

Bioinformatics

- Analysis: GWAS, EWAS, eQTL analysis, gene enrichment analysis, polygenic risk scores, Mendelian
 randomization, Gene Ontology analysis, multi-omics integration, genome annotation, single-cell data
 analysis (scRNA-seq, scDNA-seq)
- Tools: BLAST, InterProScan, PSIPRED, AlphaFold, ChimeraX, UCSC Genome Browser, Ensembl, Galaxy, Bioconductor, BEDTools, STAR, DESeq2, Seurat, edgeR, Enrichr

Mathematical

• *Core Mathematics*: Linear Algebra, Real Analysis, Probability and Statistics, Stochastic Process, Calculus, Partial Differential Equations, Discrete Mathematics, Infinite Theory

Laboratory

- Molecular Biology: PCR, qPCR, DNA/RNA extraction, CRISPR/Cas9, cloning, electrophoresis, next-generation sequencing (NGS) library preparation
- Genomics: RNA-seq, ChIP-seq, single-cell RNA-seq, bisulfite sequencing
- Proteomics: Western blot, ELISA, mass spectrometry, chromatin immunoprecipitation (ChIP)
- Cell Biology: Cell culture, transfection, confocal and fluorescence microscopy, immunofluorescence, flow cytometry

[&]quot;Influenza Vaccination: DNA Methylation and Immune Response Mechanisms", Matteo Lab, UCLA, Nov. 2024

[&]quot;Higher Education in Public Health", Fielding School of Public Health, UCLA, Apr. 2024

[&]quot;Key Factors Linked to Depression Before Adjuvant Therapy in Breast Cancer", UCLA. Mar. 2024

[&]quot;Lymphocyte Regeneration After Thymocyte Damage in Zebrafish", Li Lab, Southwest University, June 2020