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Xiang(Shawn) Li, Ph.D.

INTERESTS

Computational Solution in Data & NGS

EDUCATION

Ph.D., George Washington University, Biophysics (**Honor: Parke Prize**), 2016 \sim 2021 Thesis Summary Study of Transcriptional and Post-transcriptional Regulation in Immune Systems B.S., Huazhong University of Science & Technology, Applied Physics, (**Honor**) 2011 \sim 2015

SKILLS

Certificate: AWS CCP, DeepLearning.AI, Nextflow Certificate,

Program, Software Skills and Database:

- Advanced: Linux Shell Script, AWS cloud S3 & EC2, Python (data mining→visual→ML), NGS Tools,
- Intermediate: PyPi, Streamlit, Nextflow, WDL, flask, dash, boto3, Fortran, Gitlab, Github, SGE, LATEX
- Basic: Snakemake, Docker, javascript, Web and Server Support(Nginx), MATLAB, C/C++, Java, R
- DB: genomAD, 4DN, NCBI, GEO, ENCODE, HapMap, 1000 Genomes, GIAB, dbSNP

Work Experiences

PSC, Carnegie Mellon University Senior Bioinformatician

Pittsburgh PA, 2024 ~ Present

Support Human BioMolecular Atlas Program(HuBMAP) under Common Fund Data Ecosystem (CFDE)

J&J Scientific Engineer III, RWE

Titusville NJ, 2024

- R&D in Real World Evidence, Patient profiling from medical history (EHR, EMR)
- Dev a toolset from scratch for patient profile, alignment, visualization and analysis(built a fully connected network for 52K patients).

GentiBio, Senior Scientist, Bioinformatics Lead

Boston MA, 2022 \sim 2024

• Translational Research: for GENTI122 IND package (Safety assessment for Gene Editing).

Build SNIP-aware (gnomAD v4.0) insilico Off-target analysis for Cas9 editing On/Off-Target analysis protocol: in-Silico $\Rightarrow GUIDES\ eq \Rightarrow rhAMPS\ eq$.

Build a full-stack web app (aws) for automation. Similar to CRISPResso2 Web

• Early Discovery: Cellular Profiling Platform

scRNAseq & scATACseq integration analysis, Kidney biopsies from Mouse IgA Model

Deploy GentiBio internal chatGPT & Automation pipeline through AWS

Deploy AWS VC-backed startup package and build Bioinformatics Infrastructure.

Write analysis tools (python & R) and pipeline (bash, Nextflow Docker)

Adopt bulk/scRNAseq analysis pipeline and downsteam analysis

CAR design, Implement Alphafold2 & AlphaMissense for IBD, FliC CAR.

Sana Biotechnology Scientist, Computational Biology

Seattle WA, 2021 ~ 2022

- Work as IC in corporate team, focusing on computational pipelines (nextflow) development.
- iPSC Master Cell Bank, Cell Characterization & Safety Assessment
- GUI tools develop for analytical team

Department of Physics, GWU

Washington, DC, 2017 ~ 2021

Ph.D. (Superviser: *Prof. W. Peng*, Collaborator: *Prof. H. Xue*)

- Nature Com: Develop analysis framework (python-base for data representation, visualization and ML)
- Integration of Multi-omics: scRNAseq(10X), RNAseq and DNase_seq, ATAC_seq, ChIPseq, Cut&Run, HiC, MiCroC, HiChIP and etc.
- Chromatin structure and function: Develop a tool (HiCHub, which solved the problem on identifying a mixed range of altered genome structure(from looping to larger than TAD). This tool outperforms similar software in HiC data analysis with a significantly increase in the computational efficiency.
- Nucleic Acids Research: mRNA stability Mathematical Modeling and data analysis.
- Study regulatory roles of Tcf1/Lef1 in CD8⁺ T cells in immunue response. (JEM, 2019, 2018)
- Impact upon deficiency of miRNA-34b/c during somatic reprogramming progress.
- Organize Journal Club and give presentation once per month.
- HPC management. (Linux Admin, data storage &sharing& backup, computation dependencies setup, iupyter notebook & Rstudio server and on-call for tech emergency.
- Build analysis pipeline, dependencies and visualization tools on a cloud platform. (AWS, Linux ssh).

Team Lead, NLP Course Project George Washington University

Washington, DC, 2019

- Natural language processing, use different models for user profiling task in given posts from twitter.
- Write model to implement different language models. Hidden Markov model and bayesian classifier.

Joint Leader of Student Innovation Group (Adviser: B. Yu) Qiming College of Huazhong University of Science & Technology

Boston, 2014

- Built a team for 2014 BIOMOD, Harvard, Cambridge, Massachusetts.
- Invested nano-scale transportation system based with magnetic control device.
- Conducted the final presentation on behalf of our team.
- For more detail, please visit our website: BIOMOD 2014: HUST

Teaching

• Graduate TA. General Physics (with: *X. Qiu / G. Younes / S. Guiriec*)

2016 ~ 2018

• Instructor. Astrophysics (with: *Kalvir Dhuga*)

2019

Membership & Reviewer
• Phenomics, Youth Editorial Board Members

2024 ~ Present

Society of Artificial Intelligence Research SAIR

2023 ~ Present

Machine Learning for Genomics MLGenX

2024

HONORS AND AWARDS

- The Parke Prize, awarded for excellence in theoretical biophysics
- Scholarship in Department of Biostatistics, University of Washington, \$1,950
- Second Prize of Poster in 3rd International Ocean Sciences Summer School. \$800
- Project Awards: Silver; International Biomolecular Design Competition 2014, BIOMOD, Harvard
- Fellowship \$12000 total, Undergraduate Fellowship from College of Life Science & Technology,

- Publications & Patents & Presentations

 HiCHub: A Network-Based Approach to Identify Domains of Differential Interactions from 3D Genome Data Xiang Li[†], ..., Hai-Hui Xue, Weigun Peng., Bioinformatics (BIOINF-2025-0207)
 - Lysine methyltransferase Kmt2d regulates naive CD8+ T cell activation-induced survival Jaekwan Kim, T.N, ..., Xiang Li, ..., Nan-ping Weng., Frontiers in Immunology (2023)
 - Tcf1 CTCF cooperativity shapes genomic architecture to promote CD8+ T cell homeostasis Q. S[†], ..., Xiang Li...., Hai-Hui Xue., Nature Immunology (2022)
 - Tcf1 and Lef1 maintain CD8+ T cell identity by organizing genomic architecture Q. S[†], Xiang Li[†],..., Hai-Hui Xue., Nature Communications (2021)
 - Selected Oral Presentation, 2021 Network Biology, Cold Spring Harbour Laboratory Network analysis reveal the critical role of TCF1/LEF1, Speaker: Xiang Li
 - Sepsis leads to lasting changes in phenotype and function of memory CD8 T cells Isaac J. Jensen[†], Xiang Li[†]..... Weigun Peng and Vladimir P. Badovinac.. 2021 eLife
 - Arid1a-Plagl1-Hh signaling is indispensable for differentiation-associated cell cycle arrest of tooth root progenitors Yang Chai, ..., Li Xiang, Xinguan Jiang. Cell Reports (2021).
 - Transcriptome-wide stability analysis uncovers LARP4-mediated NFB1 mRNA stabilization during T cell activation. Tian Y.[†], Zeng Z.[†], **Li Xiang**[†], ..., Peng W. and Zhu J. Nucleic Acids Research (2020).
 - Integrated analysis of carbon dioxide and oxygen concentrations as a quality control of ocean float data Yingxu Wu, D. Bakker, E. Achterberg, ..., Xiang Li, ... Communications Earth Environment, Accepted. (Mar. 2022)
 - Tcf1 and Lef1 transcription factors are required for the immunosuppressive function of regulatory T cells Xing S., Gai K., Li Xiang,..., Peng W. and Xue H., J Exp Med (2019).
 - Tle corepressors are differentially partitioned to instruct CD8+ T cell lineage choice and identity Xing S., ..., **Xiang Li**,..., Peng W. and Xue H., J Exp Med (2018).

- Work extensively used HiCHub

 Tcf1 CTCF cooperativity shapes genomic architecture to promote CD8+ T cell homeostasis Nature Immunology (2022)
 - CTCF mediates CD8+ effector differentiation through dynamic redistribution and genomic reorganization JEM (2023)