Ginny Xiaohe LI

Computational Biologist

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EXPERIENCE

Department of Pathology, University of Michigan

Research Fellow

Lead CPTAC non-Clear Cell Renal Cell Carcinoma (non-ccRCC) proteogenomics analysis

- Collaborate with pathologists to generate research hypothesis for comprehensive molecular characterization of heterogenous non-ccRCC.
- Design data analysis workflow integrating genomic, transcriptomic, proteomic and metabolomic data.
- Supervise and conduct data analysis to discover diagnostic markers that distinguish RCC subtypes that are histologically similar and novel prognostic markers for rare RCC subtypes.
- Write manuscript and generate data visualization from scratch for high-impact scientific publication. **Conduct statistical analysis in multiple collaborative projects**
- Conduct transcriptomic and proteomic data normalization, imputation and batch effect correction in lung cancer and acute myeloid leukemia.
- Evaluate downstream impact of tumor suppressor gene bi-allelic loss in large lung cancer cohort.
- Build statistical models on multi-omic datasets to test hypotheses of biological relevance.
- Refine algorithms for allele-level quantification of polymorphic HLA proteins.
- Test, benchmark and improve open-source software automating downstream proteomics analysis.

Complete You3: A Leadership and Management Program for Postdocs

• 8-week structured program developing project and team leadership and management skills

Yong Loo Lin School of Medicine, National University of Singapore *Research Fellow*

Singapore 01/2020 – 06/2021

Model phosphorylation signaling in insulin-stimulated human myotubes

- Build an empirical Bayesian statistical framework to detect significant abundance changes in covarying pairs of kinases and substrates based on phosphoproteomics profiling.
- Identify uniquely activated signaling transduction cascades in insulin-stimulated myotubes.
- Develop an R package *KSA2D* to achieve the analysis and draft the full manuscript.

Annotate and visualize proteome-wide impact caused by kinase inhibitor Staurosporine

- Collaborate with molecular biologists from Institute of Molecular and Cellular Biology in research hypothesis generation towards delineating multiple response patterns.
- Design statistical analysis and develop an R package and Shiny App *KOPI* for protein changes annotation and visualization under various experimental settings.
- Draft and revise a manuscript with collaborators resulting in publication in *Scientific Reports*.

Saw Swee Hock School of Public Health, National University of Singapore
Research AssistantSingapore
08/2018 - 08/2019

Multi-omic marker discovery in thyroid and gastric cancer patients

• Aggregate genome variants to protein units and compare the results with respective TCGA cohorts.

• Implement differential expression analysis with transcriptome data and compare with TCGA cohorts. *Teaching Assistant* 01/2015 - 05/2017

- Facilitate "Public Health in Action" modules hosting more than 200 undergraduates for 3 semesters.
- Tutor graduate students with R programing in the module "Quantitative Epidemiology Methods" and "Advanced Biostatistics".

MI, USA 07/2021 – present

EDUCATION

Saw Swee Hock School of Public Health, National University of Singapore Singapore 08/2014 - 07/2019

Doctor of Philosophy

Thesis title: Cancer genome biomarker discovery with integrated molecular data and system biologyoriented approaches.

School of Mathematics, Shandong University

Bachelor of Science

Mathematical training in calculus, linear algebra, numerical analysis, probability, and statistics.

DOCTORAL RESEARCH

Prognostic exome variants discovery in protein-centric approach 01/2018 -12/2019

- Map exome variants across 33 cancer types in TCGA data to domains and PTM sites on proteins.
- Construct a survival analysis model to detect prognostic protein units, resulting in an R package GPD.
- Draft and revise a manuscript resulting in publication in Human Mutation.

Proteome-scale post-translational modification prediction

- Collect a total of 173 sequence and structural features to predict generic PTM sites.
- Develop an R package PTMscape incorporating Liblinear as classifier.
- Draft and revise a manuscript resulting in publication in Molecular Omics.

PROFESSIONAL SKILLS

Computational Biology

- Next generation sequencing (NGS) based genomics, transcriptomics pipeline: experienced in conducting DNA sequence alignment with Bowtie2, post-processing with SAMtools, RNA-seq pseudo-alignment and quantification with Kallisto, single cell RNA-seq quantification with Kallisto bustools.
- Mass spectrometry (MS) based proteomics pipeline: experienced in performing proteomics identification and quantitation with FragPipe.
- Statistical analysis and machine learning: proficient in handling high dimensional -omics data such as genomic variants, DNA methylation, RNA-seq (bulk and single cell), proteomics, phosphoproteomics and metabolomics data, performing data preprocessing including normalization, batch effect correction and imputation:

Proficient in conducting differential expression, functional enrichment, network analysis and other advanced analysis with statistical modelling and machine learning algorithms.

Large datasets exposure: familiar with TCGA, COSMIC, CPTAC and other public data resources. •

Programming and Data Visualization

- **R**: proficient in scientific programming including and statistical modelling and machine learning; Proficient with *dplyr*, *qqplot2*, *limma*, *biomaRt*, and many other packages on CRAN/Bioconductor; Proficient in writing reproducible reports with R Markdown; Proficient in R package and Shiny App development.
- **Python**: experienced in data analysis with *numpy* and *pandas*: • Proficient in calling transcriptomics tools in conda environments.
- Cloud Computing: experienced in working on Linux servers with command-line interfaces and • collaborating through Git.
- Cytoscape and Adobe Illustrator: proficient in publication-quality data visualization.

Jinan, China

09/2010 - 06/2014

01/2015 - 05/2018

PUBLICATIONS

- Li GX*, Lijun Chen*, Hsiao Yi*, Saravana Mohan Dhanasekaran, Nesvizhskii A, CPTAC consortium. Comprehensive proteogenomics characterization of rare kidney tumors. *Submitted*.
- Mumphrey, M. B., Li GX., Hosseini N., Nesvizhskii A., & Cieslik, M. HLAProphet: Personalized allele-level quantification of the HLA proteins. *bioRxiv*, 2023-01.
- Li GX*, Zhao T*, Choi H, Lim YT*, Sobota R. KOPI: Kinase inhibitOr Proteome Impact analysis. *Scientific Reports*. 2022 12(1),1-8.
- Ko S, Li GX, Choi H, Won JH. Computationally scalable regression modeling for large-scale clinical omics data with ParProx. *Briefings in bioinformatics. 2021 22 (6).*
- Han B*, **Li GX***, Chan E*, Liew W, Huang S, Liu MH, Choi H, Kim KP, Tai ES. Proteome and phosphoproteome dynamics in insulin-stimulated myotubes of a multi-ethnic Asian cohort. *In preparation*.
- Li GX, Munro D, Fermin D, Vogel C, Choi H. A protein-centric approach for exome variant aggregation enables sensitive association with clinical outcomes. *Human Mutation. 2020 41(5):* 934-945.
- Li GX, Vogel C, Choi H. PTMscape: an open source tool to predict generic post-translational modifications and map modification crosstalk in protein domains and biological processes. *Molecular Omics. 2018 14.(3)*, 197-209.

SOFTWARE PORTFOLIO

- **PTMscape:** proteome-scale PTM prediction using SVM with a comprehensive set of features. <u>https://github.com/ginnyintifa/PTMscape</u>
- **GPD:** protein-centric exome variants aggregation for cancer biomarker discovery. <u>https://github.com/ginnyintifa/GPD</u>
- **KSA2D**: differential analysis for discovery of co-varied kinase-substrate pairs. <u>https://github.com/ginnyintifa/KSA2D</u>
- **KOPI**: tracking and visualizing protein abundance changes in various experiment settings. <u>https://ginnyintifa.shinyapps.io/ProteomeNodesShiny/</u>

CONFERENCE PRESENTATIONS

•	CPTAC Annual Consortium Meeting 2022	Online
	Proteogenomics characterization of Pan-Renal Cell Carcinoma (talk).	10/2022
•	US Human Proteome Organization Conference 2022	Charleston
	Assessing discordance between peptide and protein level differential expression analysis (poster).	03/2022
•	CPTAC Annual Consortium Meeting 2021	Online
	Proteogenomics characterization of non-Clear Cell Renal Cell Carcinoma (poster).	10/2021
•	8 th Annual Biomedical Scientific Congress	Singapore
	GPD: a segmentation-based approach for exome association analysis (talk).	10/2019
•	25 th Intelligent Systems for Molecular Biology Conference	Prague
	Protein-centric Exome Association (PREXA) analysis for prognostic coding variants discovery (po	oster).07/2017
•	15 th Human Proteome Organization World Congress	Taipei
	PTMtopographer: an interface for proteome-scale PTM prediction (poster).	09/2016

AWARDS

•	Best Visualization Award, Yale-NUS Data1.0 Hackathon.	05/2017
•	Outstanding Graduate of Class 2014, Shandong University.	06/2014
•	First Class Scholarship, Shandong University.	06/2013