# Ginny Xiaohe LI

# Bioinformatician

+1 773-383-2781| ginnyli056@gmail.com| https://github.com/ginnyintifa

### **EXPERIENCE**

# Department of Pathology, University of Michigan

MI, USA

Research Fellow

07/2021 – present

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

- Collaborate with pathologists in research hypothesis generation towards comprehensive molecular characterization of heterogenous non-ccRCC.
- Design data analysis workflow integrating genomic, transcriptomic, proteomic and metabolomic data.
- Supervise and conduct data analysis leading to discovery of diagnostic markers distinguishing 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.

# Perform statistical analysis in multiple collaborative projects

- Conduct genomics and proteomics data quality assessment and preprocessing in various cancer cohorts including lung cancer and acute myeloid leukemia.
- Build statistical models on multi-omic datasets to test hypotheses of biological relevance.
- 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

**Singapore** 

Research Fellow

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 **Singapore** 08/2018 - 08/2019 Research Assistant

## 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".

### **EDUCATION**

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

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

# School of Mathematics, Shandong University

Jinan, China

Bachelor of Science

09/2010 - 06/2014

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

### **DOCTORAL PROJECTS**

# 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

01/2015 - 05/2018

- 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**

- 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.
- **MS based proteomics pipeline:** experienced in performing proteomics identification and quantitation with FragPipe.
- Downstream data analysis: proficient in handling high dimensional -omics data such as RNA-seq (bulk and single cell level), proteomics, phosphoproteomics and metabolomics data, conducting differential expression, functional enrichment, network analysis with statistical modelling and machine learning algorithms.
- Large datasets exposure: familiar with TCGA, COSMIC, CPTAC and many other public data resources.

# **Programming and Data Visualization**

- **R**: proficient in scientific programming including and statistical modelling and machine learning; Proficient with *dplyr*, *ggplot2*, *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/Conda**: experienced in data wrangling with *pandas* and machine learning tools in *scikit-learn*; Proficient in calling transcriptomics tools in conda environments.
- **Cytoscape and Adobe Illustrator**: proficient in publication-quality data visualization.
- Other: experienced in working on Linux servers and collaborating through Git.

### **PUBLICATIONS**

- **Li GX\***, Lijun Chen\*, Hsiao Yi\*, Saravana Mohan Dhanasekaran, Alexey Nesvizhskii, CPTAC consortium. Comprehensive proteogenomics characterization of rare kidney tumors. *Submitted*.
- **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. https://github.com/ginnyintifa/PTMscape
- **GPD:** protein-centric exome variants aggregation for cancer biomarker discovery. https://github.com/ginnvintifa/GPD
- **KSA2D**: differential analysis for discovery of co-varied kinase-substrate pairs. https://github.com/ginnyintifa/KSA2D
- **KOPI**: tracking and visualizing protein abundance changes in various experiment settings. https://ginnyintifa.shinyapps.io/ProteomeNodesShiny/

# **CONFERENCE PRESENTATIONS**

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