

Justin Cha

Education

Cornell University

August 2021 – Present

PhD student

Dr. Frank Pugh Laboratory; Computational Biology

The Pugh Lab is a leader in developing protocols for high-resolution chromatin immunoprecipitation (ChIP-exo) sequencing. I am currently involved in several projects investigating the fundamental characteristics of chromatin. I have investigated the role of the intrinsically disordered region in the LGE1/BRE1 ubiquitin-depositing complex in yeast, as well as the roles of specific components in the SWR1 H2A.Z-exchange complex. Currently I am working on methods for identifying and characterizing TF-TF interactions.

Georgia Institute of Technology

May 2018

BS with Highest Honors

Major in Biomedical Engineering; Minor in Physics

Experience

Broad Institute of Harvard and MIT – Cambridge, MA

Associate Computational Biologist II

September 2018 – July 2021

At the Broad Institute, I am a member of the Getz Lab, one of the world's leading labs for cancer genomics. I have worked on several exciting projects pushing at the forefront of the field. One was an analysis of genomic progression in head and neck squamous cell carcinoma (HNSCC). For this project, I made use of a novel set of computational methods to reconstruct the trajectory of genomic events from exome sequencing data. This allowed us to see which mutations and other variants tend to occur early on in the progression of cancer, which will be useful in treatment development and prognosis.

Integrative Systems Biology Lab – Atlanta, GA

Undergraduate Researcher

August 2016 – May 2018

In the Integrative Systems Biology Lab at Georgia Tech, I worked with Dr. Denis Tsygankov on a project to simulate the biomechanics of solid tumors. For this project, I developed a tool to simulate densely packed cell populations under high pressure using time-evolving Voronoi diagrams, which is useful for analyzing tumor growth under specified conditions.

Publications

*These authors contributed equally

- Burr, R. *et al.* (2024, in press) "Germline mutations and developmental mosaicism underlying EGFR-mutant lung cancer," *Nature Cancer*.
- Khalsa, J.*, Cha, J.*, Utro, F.*, Naeem, A.*, Murali, I.*, *et al.* (2023) "Genetic events associated with venetoclax resistance in CLL identified by whole exome sequencing of patient samples," *Blood*. Available at: <https://doi.org/10.1182/blood.2022016600>.
- Leshchiner, I.*, Mroz, E.*, Cha, J.*, *et al.* (2023) "Inferring early genetic progression in cancers with unobtainable premalignant disease," *Nature Cancer*. Available at: <https://doi.org/10.1038/s43018-023-00533-y>.

- Bustoros, M. *et al.* (2020) “Genomic profiling of smoldering multiple myeloma identifies patients at a high risk of disease progression,” *Journal of Clinical Oncology*, 38(21), pp. 2380–2389. Available at: <https://doi.org/10.1200/jco.20.00437>.

Presentations

- “Mutating the intrinsically disordered region of LGE1 in yeast impacts H2B ubiquitination,” Cornell Computational Biology student seminar, 2022
- “Mechanisms of Primary and Acquired Resistance to Venetoclax in Chronic Lymphocytic Leukemia (CLL),” American Association for Cancer Research, 2020
- “Genomic landscape of metastatic breast cancer (MBC): comprehensive cell-free DNA analysis from over 10,000 patients and comparison with primary breast cancer,” San Antonio Breast Cancer Symposium, 2020

Posters

- “High-resolution characterization of transcription factor binding in *S. cerevisiae*,” Great Lakes Bioinformatics, 2023
- “Inferring early genetic progression in cancers with unobtainable premalignant disease,” Massachusetts General Hospital Center for Cancer Research, 2019

Training

- Open Science Grid (OSG) User School; Madison, WI; all expenses paid

Skills

Programming Languages: Python, JavaScript, Julia, SQL, Matlab

Quantitative: Data analysis, Statistics, Genomics

Communication: Technical presentation/writing, Data visualization, Web development, Teaching

Links

- jcha40.github.io
Personal website
- <https://github.com/broadinstitute/PhylogicNDT>
PhylogicNDT: A tool for clustering mutations, inferring tumor phylogeny, and inferring the order of mutations in cancer
- <https://github.com/broadinstitute/getzlab-SignatureAnalyzer>
SignatureAnalyzer: A tool for identifying mutational signatures in a cohort of tumor samples
- <https://github.com/jcha40/DPMCMC>
DPMCMC: A library for a tensorflow implementation of the Dirichlet Process Monte Carlo Markov Chain algorithm