954 Fremont St., Apt 1, Menlo Park, CA, 94025, USA

□ (+1) 909-569-4530 | ■ benchu99@hotmail.com | # https://biona001.github.io/ | • https://github.com/biona001

"Expertise in Mathematics, Statistics, Software Engineering, and Quantitative Biology"

Research Interests

Math and Statistics Knockoffs, Penalized regression methods, Generalized linear models, Variance component models

Computation Parallel and High-Performance Computing, Big Data Analysis, Package Development

Biology Polygenic risk scores, Genome-Wide Association Studies, Phasing and imputation, admixture estimation

Education

University of California, Los Angeles

Los Angeles, California

Ph.D BIOMATHEMATICS

09/2016 - 08/2021

- Advisors: Kenneth Lange and Janet Sinsheimer
- Thesis: Scalable Algorithms for Genetic Association Studies, Genotype Imputation, and Ancestry Inference

University of California, Berkeley

Berkeley, California

B.A., APPLIED MATHEMATICS 09/2012 - 05/2016

Positions

Postdoctoral Researcher, Stanford University

Stanford, California

DEPARTMENT OF BIOMEDICAL DATA SCIENCE (ADVISOR: CHIARA SABATTI)

09/2021 - present

Experiences _____

Google Summer of Code

NumFOCUS - Julia Cohort

STUDENT SOFTWARE DEVELOPER

Summer 2018

- An unique summer program where students work on self-proposed open source projects
- · Added 3 additional features to IHT.jl to integrate it with the Open Mendel umbrella program

Academia Sinica Institute of Biomedical Sciences

RESEARCH ASSISTANT WITH PROF CARMAY LIM

Summer 2014, 2015, 2016

- One publication: An Efficient Protocol for Computing the pKa of Zn-Bound Water
- Using computational approaches to model drug-protein and drug-environment interactions

Publications

Knockoffs.jl: Fast, robust, and easy variable selection for statistical and genetic analysis

work in progress

BB CHU, T MORRISON, C SABATTI

Fast Score Tests for Multivariate and Longitudinal GWAS with Non-Gaussian Bases

work in progress

BB Chu, Ji SS, Zhou H, Sinsheimer JS, Sabatti C, Lange K

work in progress

Controlling the False Discovery Rate Improves Polygenic Risk Prediction with sufficient power

BB Chu, C Sabatti

Unsupervised Discovery of Ancestry Informative Markers and Genetic Admixture preprint **Proportions in Biobank-Scale Data Sets** S Ko, BB Chu, D Peterson, C Okenwa, JC Papp, DH Alexander, EM Sobel, H Zhou, K Lange Oct. 2022 A Flexible Quasi-Copula Distribution for Statistical Modeling preprint SM JI, **BB CHU**, JS SINSHEIMER, H ZHOU, K LANGE May, 2022 Multivariate Genomewide Association Analysis by Iterative Hard Thresholding preprint BB CHU, S Ko, JJ Zhou, A Jensen, H Zhou, JS Sinsheimer, K Lange April, 2021 Optimized Replacement T4 & T3 Dosing in Male & Female Hypothyroid Patients with Different BMIs using a Personalized Mechanistic Model of Thyroid Hormone Frontiers in Endocrinology **Regulation Dynamics** M CRUZ-LOYA*, BB CHU*, J JONKLAAS, DF SCHNEIDER, J DISTEFANO III (* DENOTES CO-FIRST AUTHORSHIP) A Fast Data-Driven Method for Genotype Imputation, Phasing, and Local Ancestry **Bioinformatics** Inference: MendelImpute.jl BB CHU, EM SOBEL, R WASIOLEK, JS SINSHEIMER, H ZHOU, K LANGE July, 2021 Iterative Hard Thresholding in GWAS: Generalized Linear Models, Prior Weights, and GigaScience **Double Sparsity** BB CHU, KL KEYS, CA GERMAN, H ZHOU, JIN J. ZHOU, JS SINSHEIMER, K LANGE June, 2020 **OpenMendel: A Cooperative Programming Project for Statistical Genetics Human Genetics** H ZHOU, JS SINSHEIMER, D BATES, BB CHU, CA GERMAN, S JI, KL KEYS, G MOSHER, J PAPP, EM SOBEL, J ZHAI, JJ January, 2020 ZHOU, K LANGE An Efficient Protocol for Computing the pKa of Zn-Bound Water Physical Chemistry Chemical Physics C GRAUFFEL, BB CHU, C LIM November, 2018 **Public Talks** False discovery rate control for polygenic risk prediction University of California, Los Angeles CENTER FOR ADMIXED POPULATIONS AND HEALTH EQUITY (ONLINE) April 21, 2022 Scalable algorithms for genetic association studies, genotype imputation, and University of Southern California ancestry inference University of Southern California, Biostatistics Seminar (online) January 28, 2021 **University of Michigan** Scalable algorithms for GWAS, genotype imputation, and ancestry inference University of Michigan, Biostatistics Seminar (online) December 14, 2020 Julia Meets Mendel: Algorithms and Software for Modern Genomic Data Analysis Online AMERICAN SOCIETY OF HUMAN GENETICS VIRTUAL MEETING 2020 October 27-30, 2020 **OpenMendel Programming Workshop** University of California, Los Angeles INAUGURAL LANGE SYMPOSIUM February 22, 2020

A Multiple Regression Approach for GWAS and High Dimensional Inference

QUANTITATIVE AND COMPUTATIONAL BIOSCIENCES RESEARCH LUNCH

MendelIHT.jl: Generalized Linear Models for High Dimensional Genetics (GWAS) Data JuliaCon 2019, available at: www.youtube.com/watch?v=UPIKafShwFw

University of Maryland Baltimore

July 25, 20

University of California, Los Angeles

February 8, 2020

Teachings_

Bruins-In-Genomics (B.I.G) summer research program

UCLA

GRADUATE STUDENT MENTOR Summer 2018, 2019, 2020, 2021

- Mentors 2 undergraduates for 8 weeks each summer.
- Proposed student projects. Reviewed student code. Guided them through technical aspects of genetics research.

Math 98 and 198 course **UC Berkeley**

Undergraduate student instructor

2013 - 2016

- Taught 7 semesters of beginner/advanced Rubik's cube course (2 units), with about 15 students per semester.
 Average instructor rating 4.8/5.0

Honors & Awards

FOLLOWSHIPS AND SCHOLARSHIPS

2019	JuliaCon Financial Assistance for Travel,	Baltimore, MD
2019	Doctoral Student Travel Grant,	Pisa, Italy
2018	JuliaCon Financial Assistance for Travel,	London, UK
2018	Genomics Analysis Training Grant (T32),	NIH/UCLA
2016	University Fellowship,	UCLA
2016	Registration Fee grant,	UCLA

MISCELLANEOUS

Amer. Red Cross	Certified in lifeguarding, first aid, AED, and CPR.,	2017
Cal Cube Club	International inter-univeristy cube relay, 10th place	2015
UCB Go Club	UC Berkeley semester Go (Weiqi) Tounament, 1st place	2014
WCA	3 by 3 rubik's cube speedsolve (10.52 seconds), 144th place in U.S, 910th in world	2013

Notable Softwares

Knockoffs.jl	Fast and robust variable selection with Knockoffs
MendelIHT.jl	Efficient ℓ_0 penalized regression tool for model selection, optimized for gene-mapping (GWAS) analysis
MendelImpute.jl	Fast data-driven genotype imputation, phasing, and admixture estimation
VCFTools.jl	Julia utilities for handling VCF (variant call format) files
Thyrosim.jl	Individualized mechanistic Thyroid hormone simulator for optimizing LT4 and LT4+LT3 dosing
MendelKinship.jl	Computation of theoretical/empirical kinship and other identity coefficients based on pedigree/genotype data
SnpArrays.jl	Package for handling binary PLINK 1 formatted files

OCTOBER 26, 2022 BENJAMIN CHU · CURRICULUM VITAE