

ericdotau@gmail.com | https://edotau.github.io/ (408)-786-4934 | San Francisco, CA

### **EXPERIENCE**

# Mammoth Biosciences, Inc.

Software Engineer

Mar 2022 - Present

- Develops full stack applications and infrastructure to expand diagnostics CRISPR gene-editing platform
- Improves cloud infrastructure for submitting cloud batch jobs by optimizing cost and resource allocation
- Computational support with scientific analysis for experiments to enable cross functional team efforts

# Berkeley Lights, Inc.

Software Engineer

Jun 2021 – Mar 2022

- Build scalable automation testing infrastructure for continuous integration and deployment using cloud compute engines, storage buckets, and data warehouses to enhance cutting-edge bioinformatics software
- Developed computing pipelines for scRNA-SEQ to advance T-Cell-Receptor discovery analysis toolbox
- Engineer high quality genomic software to optimize data wrangling features on R&D Life Sciences products

# **Invitae Corporation**

Research Associate

Jan 2021 – Apr 2021

- Designed and established validation strategies for library prep and sequencing of SARS-CoV-2 samples as public health surveillance that identified the novel strain variant CAL.20C as well as other variants
- Supported clinical sample management and developed scaling assays with laboratory automation, data analysis, software and leaded end-to-end pipeline development for exploratory projects

# **Duke University**

Research Analyst, Molecular Genetics and Microbiology

Jul 2018 – Dec 2020

- Programed a framework which computed Illumina, Pacific Biosciences, Oxford Nanopore sequencing that produced a full 22 chromosome genome assembly that identified candidate genes and genomic changes
- Developed graph data structure that incorporated genome diversity within populations which captured detailed and accurate variant analyses of haplotype-resolved genes and/or genomic regions
- Identified allele specific enhancers from regions of open chromatin that overlap differentially expressed genes to pinpoint biological discovery of functional enhancers in the developing brain

### **Stanford University**

Life Science Research Professional, Genetics

Oct 2013 - Jul 2018

- Managed RNA-Seq analysis on responses to physical activity from a cohort and produced a comprehensive molecular map that accelerated the development of therapeutics and exercise recommendations
- Designed DNA/RNA isolation assays at production scale which optimized workflow and reduced cost

### COMPUTATIONAL SKILLS

- Programing Languages: Go(Lang), Bash/Linux, Python, HTML/CSS/JS/TS, C/C++, Mysql/Postgres, R
- Cluster & Cloud Computing: AWS, Google Cloud, Dask, Kubernetes, Slurm, Sun Grid Engine (SGE)
- APIs & Frameworks: Apache, Docker, Django, Prefect, REST API, React, Terraform, TensorFlow

### **MOLECULAR BIOLOGY**

- NGS Library Preparation & High-throughput Sequencing: Illumina, Oxford Nanopore Technologies, Pacific Bioscience, RNA-Seq, ATAC-Seq, Whole Genome, Hi-C, Target Enrichment
- Liquid Handling and Automation: Beckman Coulter Biomek, Agilent Bravo
- Quality Control & Quantification: Nanodrop, Qubit, Tape Station, qPCR, DNA/RNA extraction

# San Jose State University

**EDUCATION** 

### Bachelor of Science in Applied & Computational Mathematics

 Relevant Coursework: Bioinformatics Numerical Analysis, Scientific Computing, Mathematical Modeling, Statistics, Data Structures and Algorithms, Differential Equations and Dynamical Systems