

# Lawrence Du

Machine Learning • Data Science • Molecular Biology

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## Skills

### Techniques

Neural networks (CNNs, GANs, GraphNN), decision trees, k-fold cross-validation, SVMs, PCA, Louvain/Leiden, k-means clustering

### Tools

Tensorflow, Linux, NumPy, SciPy, h5py, Matplotlib, Flask, Amazon EC2, Conda, LaTeX, Jenkins, Jupyter

### Programming

Python, C#, SQL, Java, Dart, Bash, MATLAB/Octave, HTML/CSS, C/C++, Perl

### Languages

Mandarin Chinese and some Spanish

## Experience

### Software Engineer II - Machine Learning Engineering • 23andMe

Apr 2020 - Present (Sunnyvale, CA)

- Developed systems for automated ML model validation and report generation for ancestry based products.
- Major contributor to new AWS Athena backed service used by researchers and therapeutics for pulling single nucleotide polymorphisms used by machine learning models and GWAS pipelines.

### Data Scientist - Ancestry Product • 23andMe

Nov 2018 - Apr 2020 (Sunnyvale, CA)

- Developed (and now maintain) current version of Recent Ancestor Locations machine learning based country matching algorithm which reaches >10 million of customers worldwide.
- Adapted new Recent Ancestor Locations to run as a microservice on AWS backed hardware using MLflow for model artifact tracking.
- Improved graph-based techniques for unsupervised identification of populations by genetic relationships.

### Bioinformatician IV • Scripps Research

May 2018 - Oct 2018 (San Diego, CA)

- Developed robust automated sequencing pipelines for Oxford Nanopore data using Common Workflow Language (CWL) for realtime microbial diagnostics, *de novo* genome assembly, and variant calling.

### Independent Consultant • Juno Diagnostics/Mindwerks (startup)

Sept 2017 - Feb 2018 (San Diego, CA)

- Developed Tensorflow deep neural networks for inferring genetic abnormalities from sequencing of cell-free DNA.
- Constructed a simulator derived from statistical analysis of human high throughput sequencing data for producing.
- Authored patent description concerning deep learning applications for diagnosis of genetic conditions.

### Data Science Fellow • Insight

Jan 2017 - Apr 2017 (Remote Session - San Diego, CA)

- Wrote [DeepPixelMonster](#) and created an interactive Python Flask web application hosted on Amazon AWS integrated with a Tensorflow backend for GAN based art generation.

### PhD Student Biology • UC San Diego • Scott A. Rifkin Lab

Aug 2010 - May 2017 (La Jolla, CA)

- Performed research on RNA expression noise during animal development by imaging single molecule RNA expression data >5,000 embryos and analyzing data using self-written MATLAB tools for image segmentation, fluorescence quantification, and image deconvolution.
- Wrote [DeepNuc](#) - a CNN model for classifying over 500,000 transcriptional start site (TSS) flanking sequences from humans, mice, fruit flies, and nematodes and over 60,000 microRNA target sequences (from publically available CLEAR-CLIP data).
- Transgenically modified over 70 nematode and Drosophila strains using techniques such as GIBSON assembly, CRISPR/Cas9, MosSCI, and *PhiC31* integrase.

## Education

Ph.D Biology UC San Diego, 2010 - 2017

B.A. Biological Sciences *Genetics and Development, Magna Cum Laude* Cornell University, 2006 - 2010

## Activities and interests

### Hobbies

I enjoy running, painting, and during the pandemic, I worked on making a VR space shooter simulator.

### Extracurricular Activities

23andMe Spitballers Ultimate Frisbee • Machine Learning Society of San Diego • UCSD GSA Lobby Corps • [BioEASI Art and Science Board](#) • Genetics Training Program Grant Hughes Scholar (2009) • [Cornell Undergraduate Research Board \(Vice President\)](#) • Triple Helix (writer) • Friends of Farmworkers (tutor)