# HARRISON WANG R&D DATA SCIENTIST

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https://harrisonized.github.io/

in https://www.linkedin.com/in/harrisonized/

• https://github.com/harrisonized

PROGRAMMING: Python, SQL, Datalog, Matlab

DATABASES & WEB FRAMEWORKS: Postgresql, Datomic, MongoDB, Flask, Django

CLOUD COMPUTING & DASHBOARDING: AWS, EC2, S3, Sagemaker, Redshift, Athena, Quicksight, Tableau, Power BI

TOOLS & LIBRARIES: Pandas, Numpy, Matplotlib, Plotly, Sklearn, Statsmodels, BeautifulSoup, Selenium

MACHINE LEARNING: Regression, Classification, Clustering, Time Series Analysis, Natural Language Processing

## **EXPERIENCE**

#### UNIVERSITY OF PENNSYLVANIA

Doctoral Candidate Aug. 2022 to Current

Ph.D. Student in the NIH-Penn Immunology Partnership Program

- First rotation in Michela Locci's lab, exploring the role Activin A in rheumatoid arthritis.

  Wet lab: tissue culture, mouse IP and tail dermal injections, anesthesia, euthanasia, flow cytometry. Dry lab: Graphpad, Flowjo, Python
- Second rotation in Montserrat Anguera's lab, studying how X-inactivation contributes to female bias for autoimmune diseases. Wet lab: RNA FISH and H3K27me3 immunofluorescence. Dry lab: R, Python

#### THERMO FISHER SCIENTIFIC

**Research and Development Data Scientist** 

South San Francisco Feb. 2021 to Aug. 2022

- Built a scalable, end-to-end ETL process to achieve real-time monitoring of SARS-CoV-2 RT-PCR test data from labs and universities using Amazon AWS and Power BI. Listed as third author on a symposium abstract based on this work
- Lead the development of the data pipeline. Heavily refactored and optimized the existing data pipeline for scalability through clever use of parallelization and batch processing, cutting the time it takes to transform 2000 csv files from 2 hours to 3 minutes. Add new KPIs and support backward compatibility
- Analyzed performance of sample retests using Python and SQL, enabling stakeholders to understand the limit-of-detection of our SARS-CoV-2 RT-PCR tests
- Designed dashboards in Amazon Quicksight and Power BI to monitor data ingestion and filtering, eliminating up-front time in scoping projects for data projects
- Created new Plotly interactive visualization for 96- and 384-well plates, enabling the Data Science team to rapidly identify anomalies in new data
- Saved a legacy Matlab codebase from disrepair through heavy refactoring

INVITAE
Dry Lab Operations Data Scientist

San Francisco Jan. 2020 to Sept. 2020

- Discovered the root cause for a high-visibility. time-sensitive issue on a new assay that resulted in batch failures impacting 500+ samples
- Analyzed trends using Python and SQL to monitor next-generation sequencing (NGS) production-line data for anomalies
- Efficiently diagnosed customer issues by writing a Python script to automate generation of standardized visualizations and statistics
- Wrote maintainable Datalog and SQL queries for Flask app used to migrate data from Datomic to Amazon Redshift for consumption by Tableau
- Built an 8-page **Tableau dashboard** to enable our operations team to carefully monitor a new product launch

#### SANOFI, FORMERLY TRUE NORTH THERAPEUTICS

**Research Associate 2** 

South San Francisco Jan. 2017 to Mar. 2019

- Led **protein-engineering project** on of lead antibody drug (BIVV009). Handled all **cloning and expression** of antibody variants and most of protein purification for the lab. **Collected binding and efficacy (KD and IC50) data** and performed **regression analysis**, revealing a log-linear relationship between KD and IC50. This enabled us to select which variants to use in downstream experiments.
- Led research project to make a protein complex of BIVV009 and its target (C1s of the complement immune system) for crystallography study.
- Wrote Python script to automate design of short DNA oligos, reducing a task that normally took 2 hours/week of manual labor to a 3 minute script.
- · Wet lab: tissue culture, protein expression and purification, ELISAs, affinity and kinetics characterization, crystallography

# GENE YEO LAB, UCSD Staff Research Associate 1

La Jolla May 2013 to Nov. 2016

• Developed standardized protocols to create tagged cell lines for next-generation sequencing (ENCODE)

• Co-authored a Neuron paper that included my experiments on elucidating the mechanism for how a mutation in an RNA-binding protein can cause ALS

- Co-authored a Cell paper that included my experiments on using a new genome-editing technology (CRISPR/Cas9) to track RNA in live cells
- Wet lab: fluorescent cell imaging, next-generation sequencing, high-throughput sequencing, CRISPR-Cas9, qPCR

## **EDUCATION**

Metis Data Science Bootcamp Apr. 2019 to June 2019

University of California, San Diego

2015