# **Jared Croyle**

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### Industry-Focused Resume for jaredcroyle.com

#### Education

University of California, Berkeley, B.S. in Molecular Environmental Biology & Minor in Data Science

May 2025

Relevant Coursework: Machine Learning, Data Analytics, Computational Structures, Practical Genomics, General Genetics

## **Technical Skills**

Languages & Tools: Python, R, SQL, BASH, HTML5/CSS3, Git, GitHub, Linux, MacOS, VSCode, Excel Data Engineering: Apache Spark, Databricks, Delta Lake, Azure, Jupyter Notebooks, Pandas, NumPy ML & Data Analysis: Scikit-learn, SciPy, Matplotlib, Seaborn, OpenCV, Feature Engineering, Clustering

#### **Experience**

Genomic Data Engineering Intern, Evolab at UC Berkeley – Berkeley, CA

February 2024 - May 2025

- Developed and productionized scalable Python/BASH pipelines to ingest and transform 10K+ semi-structured genomic records, supporting biodiversity modeling and adaptive trait analysis.
- Automated ETL workflows for data integration using open-source tools and custom scripts, improving throughput by 30%.
- Built modular components for statistical validation of predictive models, visualized results using Matplotlib to support reporting for scientific and engineering stakeholders.

**Genomic Data Intern,** Evolutionary Genomics Lab at UC Berkeley – Berkeley, CA September 2024 – December 2024

- Designed structured metadata systems using Excel and SQL for 500+ genomic samples to ensure data integrity, reproducibility, and scalability in downstream analytics pipelines.
- Streamlined QC and cataloging pipelines, supporting long-term genomic research access.

Bioinformatics Fellow (NSF-REU), Stajich Lab at UC Riverside – Riverside, CA

June 2023 - August 2023

- Developed reproducible pipelines (AMPtk, DADA2, Python) to process 7M+ metabarcode reads, generating 10K+ high-confidence ASVs and contributing to the discovery of 3 novel fungal species and 2 peer-reviewed publications.
- Applied clustering algorithms and phylogenetic analysis to microbial communities, uncovering statistical patterns to predict ecological interactions.

Veterinary Assistant and X-ray Technician, West Riverside Veterinary Hospital – Riverside, CA June 2021 – May 2023

Supported diagnostics and medical data analysis to optimize patient care and identify trends in clinical outcomes.

#### **Featured Project**

Trunkline: Full-Stack ML Pipeline for Predictive Synthetic Biology – github.com/jaredcroyle/Trunkline-SynBioML

- Built a modular, YAML-configurable ML pipeline with decoupled components for data ingestion, preprocessing, model training, and ensemble prediction. Enabled CLI-based execution for reproducibility and scalability.
- Developed an automated reporting engine using SHAP, Matplotlib, and Jinja2 to generate interpretable diagnostics (e.g., learning curves, residuals, feature importances) for rapid model evaluation and iteration.

#### **Leadership & Service**

General Member, Epsilon Eta, University of California, Berkeley

January 2024 - May 2025

• Mentored undergraduate students in biology and contributed to community outreach focused on environmental justice.

Science Education Ambassador, The Aquarium of the Pacific – Long Beach, CA

May 2018 - August 2019

• Supported international biodiversity monitoring and translated research into educational content for 10K+ visitors.

### **Awards and Publications**

- Kelly, K., Liu, X., Croyle, J., and Stajich, J.E. (2025). Climate impact on the biocrust mycobiome.
- Croyle, J., Kelly, K., Liu, X., and Stajich, J. E.. Draft genome sequence of endophytic fungus Coniochaeta sp. (In Prep)
- Nona & Innovation Award Most impactful and innovative bioinformatics software, Nonaworks Hackathon (2024)