

JAMES SACCO

Laboratory for Genomics Research, University of California, San Francisco,
San Francisco, CA 94158, USA

Self-motivated computational biologist with seven years of both academic and industry experience in genetic disease, functional genomics, and biomedical software development.

An analytical thinker and quick learner, with broad experience in AI/ML, gene regulatory network inference, and multi-omic single cell data analysis. Published manuscript on association of curated genotype, phenotype, and rare variant data to sub-type genetic disorders. Advocate for diversity and inclusion in STEM.



View this CV online with links at
<https://jsacco1.github.io/cv/>

SELECTED EXPERIENCE

Present
|
2022


Bioinformatics Programmer III


Laboratory for Genomics Research, University of California,
San Francisco

 San Francisco, CA

- Led investigation of computational rules for **CRISPR interference (CRISPRi) guide RNA design** optimization, using **Python machine learning modules**.
- Improved whole-genome targeting of promoter regions in a **genome-wide CRISPRi dual-guide library** in K562 cancer cell lines.
- **Identified over 100 putative therapeutic targets**, as computational lead on a CRISPRi screen focused on cellular dysfunction in Niemann-Pick Type C Disease.
- Integrated **single-cell epigenetic and transcriptomic data** to computationally reconstruct **gene regulatory networks** for seventy-six transcription factors in iPSC-derived cells.
- Mentored two junior programmers on the computational biology team.
- Co-authored a manuscript officially intended for publication, on in-house experimental findings.


CONTACT INFO

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 jsacco1.github.io

 github.com

 [linkedin.com](https://www.linkedin.com)

 +1 (786)-473-9993

For more information, please
contact me via email.

Work authorization: US Citizen

KEY SKILLS

Expert programming in **Python, R,**
and **Linux** shell scripting

Proficiency with **single cell gene expression, gene regulation, and epigenetic data analyses**, genomic data repositories, **machine learning**, experimental design, and CRISPR screening tools.

High scientific rigor & willingness
to teach and learn about new
computational methods and
biology

2022
|
2021

Bioinformatics Scientist



ASC Therapeutics

📍 Milpitas, CA

- Quantified and characterized **CRISPR gene modifications** in a sickle-cell disease model, with both in-house and open-source tools for genomic data analysis.
- Queried, retrieved, and integrated data from public genomics databases, to enhance on-target gene editing with **sequence alignment** protocols.
- Examined concordance and sensitivity of **five computational CRISPR off-target error detection methods**.
- Streamlined high-performance analyses for bench scientists via **custom data pipelines** and **user interfaces**, enabling round-the-clock analyses.

2020
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2019

Biomarker Data Analyst II



Genentech (Contract)

📍 South San Francisco, CA

- Established department-first **machine learning pipeline** to study effects of biomarker operations on quality of cancer immunotherapy assays.
- Upgraded **data operations** for six data streams, including flow cytometry, biomarker operations, and oncology data warehouses.
- Monitored and solved sample data quality issues for two cancer immunotherapy trial arms.

2019
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2018

Data Curator



Genentech (Contract)

📍 South San Francisco, CA

- Collaborated with AI engineers to create an ETL pipeline for **multi-modal survival prediction and patient stratification**, by using Python ML and R to integrate **gene panel** and **RNA-Seq data**.
- Designed ETL data pipeline for integration of **real-world EHRs** into **deep learning** module, using Python, SQL, and Apache Spark.
- Organized and managed close coordination of Artificial Intelligence, data management, and DevOps functions, to bring **clinical machine learning** from prototype to production.
- Provided AI and genomic analytic insights in collaboration with Roche-wide data curators and integrators network, to establish F.A.I.R. data management strategies for legacy, current, and next-generation datasets
- Tested data querying capabilities of newly-acquired AWS-based tech stack.
- Scoped design of in-house proteomics assay data ingestion and storage

PROGRAMMING SKILLS

🐍 Python ★★★★★

📊 R ★★★★★☆

🐧 Linux ★★★★★☆

🔑 Git ★★★★★

📄 Markdown ★★★★★

🔧 C++ ★★★★★☆

📐 MATLAB ★★★★★☆

2018
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2017

Data Curator, Bioinformatics Analyst

BIOMARIN

BioMarin Pharmaceutical (Contract)

📍 San Rafael, CA

- Established an integrated genetic and curated literature workflow to predict incidence and prevalence rates of **over sixty rare, genetic disorders**, with **R statistical and data visualization packages** and **MATLAB**.
- Upgraded and tested **RNA-Seq data pipeline (STAR2, SAMtools, Bioconductor)** to discern disease contribution of **rare variants** to neural disease.
- Spearheaded development of three relational databases derived from **Hail population genomics platform**, in collaboration with software engineer.
- Co-authored manuscript submitted to peer-reviewed publication, on **prediction of disease severity** in metachromatic leukodystrophy.
- Developed **allele-specific association models and GWAS analysis software** for two later published manuscripts.

2017
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2016

Research Associate



Pearse Lab: The Miami Project to Cure Paralysis, The Miller School of Medicine, University of Miami

📍 Miami, FL

- Mentored research associates in developing bioinformatic skills, specifically in **multiple sequence alignment** and **standard molecular biological protocols** (restriction enzyme digestion, gel electrophoresis, spectrophotometry).
- Identified over forty putative **conserved vertebrate phosphodiesterase proteins**, by using multiple sequence alignment and genomic evidence.



EDUCATION

Rowan University, Graduate School of Biomedical Sciences

Master of Biomedical Science (Distinction)

📍 Stratford, NJ

"Mechanisms of synaptic signaling dysfunction in hippocampal long-term potentiation"

Related coursework:

- Biochemistry
- Molecular Biology
- Cell Biology
- Mechanisms of Disease
- Pharmacology

Active member of Biostars



Biostars
BIOINFORMATICS EXPLAINED

I have participated in bioinformatics and software workshops and *Meetup.com* groups, in both Seattle, WA and San Francisco, CA, including:

SF/Bay AI Developers Group

San Jose Data Science and AI/ML Meetup

Hacker Dojo

Seattle Sequencing

LANGUAGES

English: Native Proficiency

Spanish: Native Proficiency

Dutch: Intermediate Proficiency

- Florida International University, The Honors College
BSc, Biological Sciences  Miami, FL



PROFESSIONAL DEVELOPMENT

- 2022 ● Python for Data Science: Fundamentals Part I Course
Dataquest.io  Online
- 2021 ● Introduction to Genomic Technologies
Johns Hopkins University  Online
- 2021 ● Statistics for Genomic Data Science
Johns Hopkins University  Online
- 2016 ● Software Carpentry for R and UNIX
University of Miami  Miami, FL







PUBLICATIONS

- 2023 ● Predicting disease severity in metachromatic leukodystrophy using protein activity and a patient phenotype matrix.
Genome Biology. 2023 Jul 24:172.
Trinidad M., Hong X., Froelich S., Daiker J., **Sacco J.**, Phuc Nguyen H., Campagna M., Suhr D., Suhr T., LeBowitz J.H., Gelb M.H., Clark W.T.
- 2016 ● Regulating Axonal Responses to Injury: The Intersection between Signaling Pathways Involved in Axon Myelination and The Inhibition of Axon Regeneration.
Frontiers in Molecular Neuroscience. 2016 Jun 8;9:33.
Rao S.N., Pearse D.D. **Manuscript Editor**



CONFERENCES

- 2023 ● Single Cell Analyses
 *Cold Spring Harbor Laboratory Single Cell Analyses Meeting.*
 Cold Spring Harbor, New York
- 2020 ● Prediction of disease severity in metachromatic leukodystrophy using measures of protein activity and a novel phenotype matrix.
 *American Society of Human Genetics Virtual Meeting.*
[Poster]  Online
Trinidad M., Hong X., **Sacco J.**, Nguyen H.P., Clark W.T., Froelich S., LeBowitz J.H., Gelb M.H.

- 2018 • Association of HGMD and gnomAD variants of unknown significance with prediction of disease incidence and prevalence.
- BIOMARIN *BioMarin Internal Research and Development Asilomar Conference*. [Poster]
- 📍 Pacific Grove, CA
- Sacco J., Clark W.T., Yu K., Wu K., LeBowitz J.H.

RESEARCH PROJECTS

- 2019 • Characterization of glycan substrates accumulating in GM1 gangliosidosis
- Molecular Genetics and Metabolism Reports*. 2019. 21, p.100524.
- Lawrence R., et al.
- 2018 • Utilizing ExAC to assess the hidden contribution of variants of unknown significance to Sanfilippo Type B incidence
- PLoS One*. 2018. 13(7):e0200008.
- Clark W.T., et al.
- 2017 • Phosphodiesterase Inhibitors as a Therapeutic Approach to Neuroprotection and Repair
- International Journal of Molecular Sciences*. 2017. 18(4): 696.
- Knott E.P., et al.

ORAL COMMUNICATIONS

- 2022 • Data Visualization of RNA-Seq Differential Expression Analysis with DESeq2
- hhmi** Doudna Lab: Howard Hughes Medical Institute, Department of Molecular and Cell Biology, University of California, Berkeley
- 📍 Berkeley, CA
- Topics: Introduction to RNA-Seq, Standard Workflows with DESeq2, Reference-Based Assembly, Exploratory Data Analysis, Statistical Analysis, Data Visualization, Gene Set Enrichment, Variant Annotation
- Presented publication-quality RNA-Seq data summaries and representations to post-doctoral associates
- 2022 • Changes in chromatin accessibility are not concordant with transcriptional changes for single-factor perturbations
- UCSF** Laboratory for Genomics Research & Department Biochemistry & Biophysics, University of California, San Francisco
- 📍 San Francisco, CA
- Presented now-published preprint in contrast with established research at internal Journal Club to inform future experimental design in epigenetic editing.

HOBBIES

Stonewall Sports kickball. San Francisco, CA. 2023 – Present.

Ultimate frisbee player. San Francisco, CA. 2018 – Present.

Rugby at *Seattle Quake RFC*. Seattle, WA. 2016 – 2017.



TEACHING

2009



Peer-Led Team Learning

Florida International University

📍 Miami, FL

General Biology II facilitator for two concurrent PLTL courses

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Last update: 2023-11-14. The most recent version of this resume is [available here](#).