San Francisco, California – jck.bio

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## **EDUCATION**

## Ph.D. – University of California, San Francisco

San Francisco, CA

Dept. of Biochemistry & Biophysics; Fellowships: NSF GRFP, PhRMA, NIH T32, Discovery

Sept. 2015 - Apr. 2018

B.S. (Honors) - University of Central Florida

Orlando, FL

Biotechnology; GPA: 4.0/4.0; National Merit Scholar: Full Tuition & Stipend Funding

Aug. 2012 - Apr. 2015

### RECENT EXPERIENCE

# Calico Life Sciences

South San Francisco, CA

Computational Fellow

March 2020 - Present

- Leading a research program focused on programming cell identity to address age-related diseases
- Lead the development of a computational & experimental platform for pooled screening of cell reprogramming strategies, successfully completing a company-wide goal
- Developed scNym semi-supervised adversarial neural networks for classifying single cell genomics profiles. Set new state of the art for performance, used by the Human Cell Atlas scnym.research.calicolabs.com
- Guided biomarker and lead selection decisions for small molecule/mAb programs with functional genomics

#### Calico Life Sciences

South San Francisco, CA

Data Scientist, Computing

May 2018 - February 2020

- Lead an investigation of cell type and tissue environment influences on aging using single cell genomics across three murine tissues mca.research.calicolabs.com
- Lead an investigation of skeletal muscle aging using single cell genomics & dynamical systems modeling that revealed a critical point where old stem cells fail to differentiate myo.research.calicolabs.com
- $\bullet$  Developed an analysis platform for pooled genomics screens, guided target ID & biomarker development
- Developed timelapse image analysis methods for oncology applications, enabling multi-day cell tracking

## University of California, San Francisco

San Francisco, CA

PhD Candidate; Thesis: Inferring cell state from cell behavior; Advisor: Wallace Marshall

Sept. 2015 - Apr. 2018

- Developed Heteromotility, a biological motion analysis package to quantify cell state transitions
- Developed Lanternfish deep learning package to enable discrimination and prediction of cell behaviors
- Discovered age-dampened state transitions in myogenesis with single cell RNA-seq & Heteromotility

IBM Research San Jose, CA

Deep Learning Intern, Cellular Engineering

Sept. 2017 - Dec. 2017

- Developed rapid convolutional neural network segmentation methods for microscopy data
- Implemented a Natural Evolution Strategies (NES) optimization framework to improve multi-cell tracking

### SKILLS

- Data Analysis: Experimental design, linear modeling, time series analysis, multivariate statistics
- Machine Learning: Neural networks, semi-supervised learning, adversarial learning, PyTorch, Tensorflow
- Programming: Python (scipy, scikit-learn, statsmodels), R, Matlab, git, bash, LATEX
- NGS: Single cell and bulk RNA-seq analysis, variant calling, samtools, IGV
- Image Analysis: Image segmentation, classification, particle tracking
- Experimental Biology: Single cell RNA-seq, primary cell culture, molecular biology, automated microscopy

### SELECTED PUBLICATIONS

## Author links for complete listing: Google Scholar, PubMed, bioRxiv

- 1. **Kimmel JC**, Kelley DR. Semi-supervised adversarial neural networks for single-cell classification. 2021. *Genome Research*. doi: 10.1101/gr.268581.120.
- 2. **Kimmel JC**, Hendrickson DG, Kelley DR. Differentiation reveals the plasticity of age-related change in murine muscle progenitors. 2020. *bioRxiv*. doi: 10.1101/2020.03.05.979112. In revision at *Cell Reports*.

- 3. **Kimmel JC**, Hwang AB, Scaramozza A, Marshall WF, Brack AS. Aging induces aberrant state transition kinetics in murine muscle stem cells. 2020. *Development*. doi: 10.1242/dev.183855. **Chosen as a Research Highlight** by *Development*: Muscling in on Stem Cell Aging.
- 4. **Kimmel JC**. Disentangling latent representations of single cell RNA-seq experiments. 2020. bioRxiv. doi: 10.1101/2020.03.04.972166.
- 5. Deuker M, Lewis Hardell KL, Ingaramo M, **Kimmel JC**, Buffenstein R, Settleman J. Unprovoked Stabilization and Nuclear Accumulation of the Naked Mole-Rat p53 Protein. 2020. *Scientific Reports*.
- 6. **Kimmel JC**, Penland L, Rubinstein ND, Hendrickson DG, Kelley DR, Rosenthal AZ. Murine single-cell RNA-seq reveals cell-identity- and tissue-specific trajectories of aging. 2019. *Genome Research*. doi: 10.1101/gr.253880.119. **Featured on the cover of** *Genome Research*.
- 7. **Kimmel JC**, Brack AS, Marshall WF. Deep convolutional and recurrent neural networks for cell motility discrimination and prediction. 2019. *IEEE Transactions on Computational Biology and Bioinformatics*. doi: 10.1109/TCBB.2019.2919307. Preprint featured in *Company of Biologists: the Node*.
- 8. **Kimmel JC**, Chang AY, Brack AS, Marshall WF. Inferring cell state by quantitative motility analysis reveals a dynamic state system and broken detailed balance. 2018. *PLoS Computational Biology* 14(1): e1005927. doi: 10.1371/journal.pcbi.1005927. **Chosen as an Editor's Pick** in *PLoS Editor's Collections*.

# SELECTED PRESENTATIONS

- 1. **Kimmel JC**, Kelley DR. scNym: Semi-supervised adversarial neural networks for single cell classification. Contributed talk at the International Conference on Machine Learning (ICML) 2020, Workshop on Computational Biology. Awarded: Contributor Award for best reviewed papers.
- 2. **Kimmel JC**, Hendrickson DG, Kelley DR. Differentiation reveals latent features of aging and an energy barrier in murine myogenesis. Selected talk at Cold Spring Harbor Mechanisms of Aging 2020.
- 3. **Kimmel JC**, Kelley DR. scNym: Semi-supervised adversarial neural networks for single cell classification. Selected talk at Intelligent Systems for Molecular Biology (ISMB) 2020, Machine Learning in Computational & Systems Biology session.
- 4. **Kimmel JC**, Penland L, Rubinstein ND, Hendrickson DG, Kelley DR, Rosenthal AZ. Cell type and tissue-specific aging trajectories. <u>Invited talk</u> at California QB3 Institute's *Aging and the Single Cell* event. San Francisco, CA. 2019.
- 5. **Kimmel JC**, Penland L, Rubinstein ND, Hendrickson DG, Kelley DR, Rosenthal AZ. Cell type and tissue-specific aging trajectories. <u>Invited talk</u> at Mission Bay Capital Biolabs. San Francisco, CA. 2019.
- 6. **Kimmel JC**, Hwang A, Brack AS, Marshall WF. Inferring cell state dynamics with machine learning. Invited talk at the *Machine Learning in Cell Biology* session at ASCB-EMBO. San Diego, CA. 2018.
- 7. **Kimmel JC**, Brack AS, Marshall WF. Deep neural networks for cell motility analysis. Nvidia Deep Learning in Biomedicine Workshop. San Francisco, CA. 2018. Awarded: Nvidia Most Innovative.

## SCIENTIFIC SERVICE

- Peer Reviewer: Bioinformatics, eLife, IEEE Journal of Biomedical & Health Informatics, Nature Medicine, PLoS Computational Biology, Proc. of the National Academy of Sciences
- Open Source: statsmodels, gseapy, CellBender, scnym<sup>†</sup>, heteromotility<sup>†</sup>, velodyn<sup>†</sup>. <sup>†</sup>Maintainer.

## Selected Honors & Awards

- Contributor Award for top submission, ICML Computational Biology, 2020
- Selected as an *eLife* Early Career Reviewer, 2020
- Nvidia Most Innovative Use of Deep Learning in Biomedicine, Nvidia Deep Learning in Biomedicine, 2018
- National Science Foundation, Graduate Research Fellowship, \$138,000/3 years, 2017
- PhRMA Foundation Informatics Fellowship, \$40,000/2 years, 2017
- UCSF Discovery Fellowship, 2017
- Nvidia GPU Grant, 2017 & 2018
- National Institutes of Health T32 Appointee, \$82,000/2 years, 2015