Jacob C. Kimmel

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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

RECENT EXPERIENCE

Calico Life Sciences

South San Francisco, CA

March 2020 - Present

 $Computational\ Fellow$

- Leading a research program focused on repurposing developmental programs to address aging pathologies
- Lead the development of a computational & experimental platform for pooled screening of complex perturbations, 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

Calico Life Sciences

South San Francisco, CA

May 2018 - February 2020

Data Scientist, Computing

- 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
- 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

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- 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
- Automated custom microscopy hardware for high-throughput data acquisition

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: Google Scholar, PubMed, bioRxiv

- 1. **Kimmel JC**, Kelley DR. scNym: Semi-supervised adversarial neural networks for single cell classification. 2020. doi: https://doi.org/10.1101/2020.06.04.132324. Accepted at *Genome Research*.
- 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. <u>Contributed talk</u> at the International Conference on Machine Learning (ICML) 2020, Workshop on Computational Biology. <u>Awarded</u>: 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. <u>Selected talk</u> 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. Invited talk 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. <u>Awarded:</u> 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[†], heteromotility[†], velodyn[†]. [†]Maintainer.
- Experimental Oversight: Reviewed all pre-clinical in vivo work at Calico as an Animal Committee member.

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