# Jacob C. Kimmel

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# **Profile**

Computational biologist with extensive experience in data analysis, machine learning, and image analysis. Deep biological domain knowledge, from molecules to mouse trials. Passionate about finding value in overlooked places.

# Experience

# PhD Candidate, University of California San Francisco, San Francisco, CA, 2015 - Present Dept. of Biochemistry and Biophysics

Thesis: Inferring stem cell state from cell behavior by quantitative image analysis

- Developed *Heteromotility* biological motion analysis package, including feature extraction, unsupervised clustering, and time-series analysis tools to quantify dynamic state transitions in cellular systems
- Quantified rates of muscle stem cell differentiation with single cell resolution for the first time using Heteromotility
- Developed *Lanternfish* deep learning package to enable the analysis of biological motion using convolutional neural networks for the first time
- Demonstrated classification of stem cell differentiation states and cancerous transformation detection using *Lanternfish*

# Summer Intern, National Institute on Aging, Baltimore, MD, 2012 Summer Image Informatics and Computational Biology Unit

- Identified that tissues age at different rates across the anatomy of C. elegans with machine learning image classifiers
- Investigated the relationship between cognitive and age related morphologies in the primate hippocampus with machine learning image classifiers

### Burnett Research Scholar, University of Central Florida, Orlando, FL, 2011 - 2015

- Utilized machine-learning image classifiers to characterize the timing of induced pluripotent reprogramming the process of generating new embryonic-like stem cells from normal adult cells
- Developed a data processing pipeline for mouse behavioral testing data
- Examined the use of polarized optical torques to guide cell motility in collaboration with U. of Central Florida's optics research center

# Amgen Scholar (Summer), University of California San Francisco, San Francisco, CA, 2014

- Implemented quantitative PCR normalization algorithms to enable robust analysis of gene expression
- Investigated the role of microRNA-29 in the regulation of cytokine production in T-cells

### Skills

Data Analysis: Experimental design, hypothesis testing, time series analysis, multivariate statistics Machine Learning: Supervised, unsupervised, feature engineering, distributed computing (MPI, SGE) Deep Learning: Convolutional neural networks, Tensorflow, Theano, Keras

Image Analysis: Image segmentation, classification, object detection, particle tracking **Programming:** Python (scipy stack, pandas, scikit-learn), Matlab, R, git, shell scipt, LATEX

# Education

#### Ph.D. — Dept. of Biochemistry & Biophysics, UC San Francisco, 2015 - Present

Funding: NSF Graduate Fellowship, PhRMA Informatics Fellowship, NIH T32 Awardee

GPA: 4.0 / 4.0

### B.S. — Biotechnology, University of Central Florida, 2012 - 2015

Distinctions: Summa Cum Laude, Top Honors, University Honors

GPA: 4.0 / 4.0

#### Selected Publications & Presentations

- 1. Deep convolutional neural networks allow analysis of cell motility during stem cell differentiation and neoplastic transformation. **Jacob C. Kimmel**, Andrew S. Brack, and Wallace F. Marshall. 2017. bioRxiv. doi: https://doi.org/10.1101/159202. (Submitted for publication).
- 2. Inferring cell state by quantitative motility analysis reveals a dynamic state system and broken detailed balance. **Jacob C. Kimmel**, Amy Y. Chang, Andrew S. Brack, Wallace F. Marshall. 2017. bioRxiv 168534; doi: https://doi.org/10.1101/168534. (Submitted for publication).
- 3. Inferring cell state from cell motility behavior. **Jacob C. Kimmel**, Amy Y. Chang, Andrew S. Brack, Wallace F. Marshall. Selected oral presentation to the NSF Quantitative Cell Biology Network Workshop. Allen Institute for Cell Science, Seattle, WA. 2016.
- 4. Characterization of induced pluripotent reprogramming in mesenchymal stem cells by quantitative image analysis. **Jacob C. Kimmel**, B. Yuan, C. Constant, D. Dogariu, H. Foroosh, K. Sugaya. U. of Central Florida, Showcase of Research Excellence. 2014.
- 5. Do different tissues age at different rates? Machine-learning based characterization of aging rates in C. elegans and rhesus macques. **Jacob C. Kimmel**, D. Mark Eckley, Chris Colleta, John Long, Nikita Orlov, Peter Rapp, Ilya G. Goldberg. National Institute on Aging, Baltimore, MD. 2012.

#### Honors and Awards

- National Science Foundation Graduate Research Fellowship, 2017
- PhRMA Informatics Fellowship, PhRMA Foundation, 2017
- Amgen Scholar, University of California San Francisco, 2014
- Burnett Research Scholar, U. of Central Florida, 2013
- Florida Academic Scholar, State of Florida, 2012-2015
- National Merit Scholar, 2012

# Outreach & Volunteering

#### Science Education Partnership, UCSF/SFUSD, 2016 - 2017

• Partnered with San Francisco public school teachers to design and implement hands-on lessons in cell biology and microbiology for 1st grade students

#### Bay Area Maker Faire Exposition, 2016 - Present

- Bay Area festival to present creative science and technology to the public, 100,000+ attendees
- Presented an exhibit demonstrating adaptive cellular responses to engage the public with life science research