Matthew G. Jones

378 Stanley Hall Email: matthew.jones@ucsf.edu
University of California, Berkeley Home: mattjones315.github.io

Berkeley, CA ORCID iD: orcid.org/0000-0002-0363-4493

Education

† Indicates expected

2017–2021 † Ph.D., Bioinformatics, University of California, San Francisco

Thesis Title: Computational tools for single-cell lineage tracing & applica-

tions to cancer biology

Research Advisors: Jonathan Weissman and Nir Yosef

2013-2017 B.A., Computer Science, University of California, Berkeley

Research Advisors: Rasmus Nielsen, Russ Corbett-Detig, & Nir Yosef

Publications

Preprints

(1) Ouardini K, Lopez R, **Jones MG**, Prillo S, Zhang R, Jordan MI, Yosef N. Reconstructing unobserved cellular states from paired single-cell lineage tracing and transcriptomics data. *BioRxiv*. 2021.

Journal Articles

- (1) Gong W*, Granados A*, Hu J*, **Jones MG***, Raz O*, Salvador-Martinez I*, Zhang H*, ..., Meyer P. Benchmarked approaches for cell lineage reconstructions of in vitro dividing cells and in silico models of Caenorhabditis elegans and Mus musculus developmental trees. *Cell Systems* (In press). 2021.
- (2) Quinn JJQ*, **Jones MG***, Okimoto RA, Nanjo S, Chan MM, Yosef N, Bivona TG, Weissman JS. Single-cell lineages reveal the rates, routes, and drivers of metastasis in cancer xenografts. *Science*. 2021
- (3) **Jones MG***, Khodaverdian A*, Quinn JJ*, Chan MM, Hussmann JA, Wang R, Xu C, Weissman JS, Yosef N. Inference of Single Cell Phylogenies from Lineage Tracing Data with Cassiopeia. *Genome Biology*. 2020
- (4) Newberry RW, ..., **Jones MG**, ..., DeGrado WF, Kampmann M. Robust Sequence Determinants of α -Synuclein Toxicity in Yeast Implicate Membrane Binding. *ACS Chem. Biol.*. 2020
- (5) DeTomaso D*, **Jones MG***, Subramaniam M, Ashuach T, Ye JC, Yosef N. Functional Interpretation of Single-Cell Similarity Maps. *Nature Communications*. 2019

- (6) Chan MM*, Smith ZD*, Grosswendt S, Kretzmer H, Norman T, Adamson B, Jost M, Quinn JJ, Yang D, **Jones MG**, Khodaverdian A, Yosef N, Meissner A, Weissman JS. <u>Molecular recording of mammalian embryogenesis</u>. *Nature*. 2019
- (7) Corbett-Detig, R. and **Jones, M**. SELAM: Simulation of Epistasis and Local adaptation during Admixture with Mate choice. *Bioinformatics*. 2016

Teaching Assistant

2020	CS176: Algorithms for Computational Biology	University of California, Berkeley
2020	BP205B: Dynamical Systems Modeling	University of California, San Francisco
2020	BP205B: Dynamical Systems Modeling	University of California, San Francisco

Funding, Awards & Honors

2019-2021	UCSF Discovery Fellowship
2020	Allen Institute Cell Lineage Reconstruction DREAM Challenge Best Performer
2019	Quantitative Biology Consortium Retreat Best Poster
2018	Quantitative Biology Consortium Retreat Best Poster

Presentations

Invited Talks

2021	Weinberg Lab @ MIT, journal club
2021	Royer Group @ Biohub & Wagner Group @ UCSF, group meeting
2021	AI/ML @ Genentech, seminar
2021	NIH Single Cell Users Group, seminar
2021	Yale Center for Biomedical Data Science, seminar
2020	UC Berkeley Computational Biology Skills Seminar, seminar
2019	UC Berkeley Computational Biology Skills Seminar, seminar

^{*} denotes equal contribution

Contributed Talks

2021	The Cancer Target Discovery and Development Alliance NIH Site Visit
2021	CZI Biohub Seed Networks Computational Biology Meeting
2020	Hindsight 2020 - The Allen Institute Developmental Recording Symposium
2020	CZI Seed Network 2020 Annual Meeting
2020	Center for Genomic Editing and Recording (CGER) Retreat
2019	UC Berkeley Computational and Genomic Biology Retreat
2019	Quantitative Biology Consortium Retreat

Posters

2019	Quantitative Biology Consortium Retreat
2019	Chan-Zuckerberg Biohub Confab
2019	Next Generation Genomics
2018	Quantitative Biology Consortium Retreat

Industry Experience

2020	Google Health Genomics, research intern
2016	United States Medical Affairs @ Genentech, intern

Appointments

2020 Weizmann Institute - Rehovot, Israel (Hosts: Ido Amit & Nir Yosef)

Professional Activities

2020-2021	UC Berkeley Computational Biology Skills Seminar, coordinator
2019	UC Berkeley Computational Biology Skills Seminar, participant
2018-2019	Northern California Computational Biology Student Symposium, coordinator
2019	UCSF Integrative Program in Quantitative Biology (IPQB) Bootcamp, coordinator
2018	UCSF Integrative Program in Quantitative Biology (IPQB) Bootcamp, Bioinformatics Module Leader

Supervision

2019-	Richard Zhang, Research Engineer	UC Berkeley
2020-2021	Khalil Ouardini, MSc Student	ENS Cachan, MVA
2019-2021	Yanay Rosen, Undergraduate	UC Berkeley
2019-2021	Robert Wang, Undergraduate	UC Berkeley

Software

Cassiopeia	Analysis tools for single-cell lineage tracing data.
VISION	Functional annotation of scRNA-seq data.
SELAM	Simulation framework for large-scale population admixture.

References

Available on request.