# Lincoln J. Harris

Curriculum Vitae

Last Updated: December 11, 2019

Aspiring computational biologist with strong interests in big data, disease evolution and genomics, teaching, and open source code/science

## Education

2017 **B.A.**, **Biology**, Swarthmore College, Swarthmore, PA.

# Experience

2017– **Chan Zuckerberg Biohub**, Spyros Darmanis Laboratory, San Francisco, CA, Research Present Associate II.

Investigated large biological datasets with a variety of tools including Docker, Python (pandas, SciPy, Matplotlib, scikit-learn), AWS, and workflow management systems such as Reflow. Worked as part of a collaborative team in a fast-paced, cutting-edge scientific environment.

#### Projects:

- Built a software tool for fast and memory-efficient summarizing of vcf (variant calling format)
   file entries following a sequencing experiment (https://github.com/czbiohub/cerebra)
- Identified mutations in single-cell RNA-seq patient samples as part of a larger effort to better characterize changes in the lung tumor microenvironment across disease progression. In collaboration with the Bivona lab of UCSF, manuscript submitted to *Cell*
- Investigated the gene expression and alternate splicing dynamics of neuroendocrine cells in the developing lung, in collaboration with the Christin Kuo lab of Stanford
- 2016–2017 **Brad Davidson Laboratory**, Swarthmore College Biology Department, Swarthmore, PA, Computational Biology Research Assistant.

Assembled the genome and embryonic transcriptome of *Corella Inflata* (sea squirt). Helped construct an improved phylogenetic tree for the *Tunicate* subphylum, in collaboration with the Joe Ryan Lab of the University of Florida Whitney Marine Station

# Research Training

- 2015–2016 Nick Kaplinsky Laboratory, Swarthmore College, Swarthmore, PA.
  - Developed web-based pipelines for differential gene expression tests in RNA-seq samples
- 2013–2014 **Evan Vosburgh Laboratory**, Cancer Institute of New Jersey, New Brunswick, NJ. Investigated the effects of several cancer drugs on expression of key oncogenes in cell culture.

#### Honors and Awards

2019 NSF Graduate Research Fellowship Program – Applicant

## Publications

#### Articles

- [1] Maynard A, McCoach C, Rotow J\*, **Harris L**\*, Darmanis S, Blakely C, Bivona T, et al. Heterogeneity and targeted therapy-induced adaptations in lung cancer revealed by single-cell RNA-seq. *submitted to Cell*, 2019.

  \*These authors contributed equally to this work.
- [2] DeBiasse M, Colgan W, Harris L, Ryan J, and Davidson B. The genome of Corella inflata and phylogeny of Tunicata. submitted to Molecular Biology and Evolution, 2019.
- [3] Schaum N, **The Tabula Muris Consortium**, et al. The murine transcriptome reveals global aging nodes with organ-specific phase and amplitude. *bioRxiv*, 2019.
- [4] **Harris L**, Vanheusden R, and Darmanis S. Cerebra: a tool for high-throughput variant call processing. *manuscript in preparation*, 2019.
- [5] The Tabula Muris Consortium et al. A single cell transcriptomic atlas characterizes aging tissues in the mouse. bioRxiv, 2019.

#### Conference Posters

- [1] **Harris L**, Maynard A, McCoach C, Rotow J, , Darmanis S, Blakely C, Bivona T, et al. Single-cell characterization of the tumor microenvironment in advanced stage Lung Adenocarcinoma. In *Chan Zuckerberg Biohub Inter-lab Confab*, San Francisco, CA, October 2018.
- [2] **Harris L**, Schug J, Kaplinsky N, and Davidson B. Towards an assembled genome and transcriptome of *Corella Inflata*. In *Sigma Xi National Honor Society Student Research Showcase*, Swarthmore, PA, October 2016.
- [3] Harris L, Vanheusden R, and Darmanis S. Cerebra: a new tool for high-throughput variant call processing. In *Algorithms and models for Single-cell genomics*, Irvine, CA, June 2019.

## Talks

- \* indicates invited talk
- 2019 New methods for high-throughput variant call processing, Chan Zuckerberg Biohub Interlab Confab, San Francisco, CA.
- 2019 New methods for high-throughput variant call processing, Beyond the Cell Atlas Conference, Berkeley, CA\*.
- 2018 Single-cell characterization of cancer hallmarks in advanced stage Lung Adenocarcinoma, Chan Zuckerberg Biohub Internal Seminar Series, San Francisco, CA.
- 2017 The genome and embryonic transcriptome of *Corella Inflata*, *Undergraduate Biology Department Research Symposium*, Swarthmore College, Swarthmore, PA.

## Software

- cerebra Performs high-throughput summarizing of vcf entries following a sequencing experiment. https://github.com/czbiohub/cerebra (19 stars on GitHub)
- motifScan Scans an enhancer region for putative transcription factor binding sites. https://github.com/lincoln-harris/motifScan

# Teaching, Leadership and Interests

- 2017–2019 **Instructor**, Cupcakes and Coding, Chan Zuckerberg Biohub, San Francisco, CA. Volunteered as an instructor at weekly beginner-friendly coding sessions
- 2016–2017 **Co-host**, WSRN Radio Station, Swarthmore, PA. Co-hosted a popular weekly radio program for two semesters
- 2016–2017 **Volunteer Instructor**, *Youth Swim Lessons*, Swarthmore, PA. Taught bi-weekly swim lessons for children ages 5-12
  - 2016 **Student teacher**, Swarthmore Rutledge Elementary School, Swarthmore, PA. Taught science lessons for 5th grade students at a local elementary school
- 2013–2017 **Member**, Swarthmore College Swim Team, Swarthmore, PA. Participated as a four-year varsity member on a Division III swim team
- 2012–2013 **Captain**, South Brunswick High School Swim Team, South Brunswick, NJ. Lead varsity team to a dual-meet record of 6-4

## Mentees

2019 Rohan Vanheusden, Nueva High School, San Mateo, CA, High School Student.
Mentored a local high school student with strong interests in computational biology for 11 weeks.