Adam Joseph Gayoso

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Education

Ph.D. Computational Biology 2018 -

> University of California, Berkeley Advisors: Nir Yosef, Aaron Streets

M.S. Computer Science 2016 - 2017

COLUMBIA UNIVERSITY

B.S. Operations Research: Engineering Management Systems, cum laude 2011 - 2015

COLUMBIA UNIVERSITY

Grants, honors & awards

NHGRI Genomics and Computational Biology T₃₂ trainee 2019 - 2020 Ralph W. Haines Scholarship, Columbia Engineering 2012 - 2015 2014

Tau Beta Pi induction, Columbia Engineering (top 1/8 of class)

Publications

MANUSCRIPTS IN SUBMISSION

- 1. Tal Ashuach, Daniel A. Reidenbach, Adam Gayoso, and Nir Yosef. "PeakVI: A Deep Generative Model for Single Cell Chromatin Accessibility Analysis". In: bioRxiv (2021). [PDF].
- 2. Adam Gayoso*, Romain Lopez*, Galen Xing*, Pierre Boyeau, Katherine Wu, Michael Jayasuriya, Edouard Mehlman, Maxime Langevin, Yining Liu, Jules Samaran, Gabriel Misrachi, Achille Nazaret, Oscar Clivio, Chenling Xu, Tal Ashuach, Mohammad Lotfollahi, Valentine Svensson, Eduardo da Veiga Beltrame, Carlos Talavera-Lopez, Lior Pachter, Fabian J Theis, Aaron Streets, Michael I Jordan, Jeffrey Regier, and Nir Yosef. "scvi-tools: a library for deep probabilistic analysis of single-cell omics data". In: bioRxiv (2021). [PDF].

JOURNAL ARTICLES

- 1. Mohammad Lotfollahi, Mohsen Naghipourfar, Malte D. Luecken, Matin Khajavi, Maren Büttner, Marco Wagenstetter, Ziga Avsec, Adam Gayoso, Nir Yosef, Marta Interlandi, Sergei Rybakov, Alexander V. Misharin, and Fabian J. Theis. "Query to reference single-cell integration with transfer learning". In: Nature Biotechnology (In press) (2021). [PDF].
- 2. Adam Gayoso*, Zoë Steier*, Romain Lopez, Jeffrey Regier, Kristopher L Nazor, Aaron Streets, and Nir Yosef. "Joint probabilistic modeling of single-cell multi-omic data with totalVI". In: Nature Methods (2021). [PDF].

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3. Valentine Svensson, Adam Gayoso, Nir Yosef, and Lior Pachter. "Interpretable factor models of single-cell RNA-seq via variational autoencoders". In: *Bioinformatics* (2020). [PDF].

- 4. Manu Setty, Vaidotas Kiseliovas, Jacob Levine, **Adam Gayoso**, Linas Mazutis, and Dana Pe'er. "Characterization of cell fate probabilities in single-cell data with Palantir". In: *Nature Biotechnology* (2019). [PDF].
- 5. Sheila Adams-Sapper, **Adam Gayoso**, and Lee. W. Riley. "Stress-adaptive responses associated with high-level carbapenem resistance in KPC-Producing Klebsiella pneumoniae". In: *Journal of Pathogens* (2018). [PDF].

REVIEW ARTICLES

1. Romain Lopez, **Adam Gayoso**, and Nir Yosef. "Enhancing scientific discoveries in molecular biology with deep generative models". In: *Molecular Systems Biology* (2020). [PDF].

REFEREED WORKSHOP PAPERS

- 1. Pierre Boyeau, Romain Lopez, Jeffrey Regier, Adam Gayoso, Michael I. Jordan, and Nir Yosef. "Deep Generative Models for Detecting Differential Expression in Single Cells". In: *Machine Learning in Computational Biology.* 2019. [PDF].
- 2. Oscar Clivio, Romain Lopez, Jeffrey Regier, Adam Gayoso, Michael I. Jordan, and Nir Yosef. "Detecting Zero-Inflated Genes in Single-Cell Transcriptomics Data". In: *Machine Learning in Computational Biology.* 2019. [PDF].
- 3. Adam Gayoso, Romain Lopez, Zoë Steier, Jeffrey Regier, Aaron Streets, and Nir Yosef. "A Joint Model of RNA Expression and Surface Protein Abundance in Single Cells". In: *Machine Learning in Computational Biology*. 2019. [PDF].

Presentations

INVITED TALKS

Apr 2021 Oct 2019

| May 2021 | Seminar, Celsius Therapeutics |
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| May 2021 | Single cell user group, NIH |
| Apr 2021 | Kundaje Lab Journal Club, Stanford University |
| Feb 2021 | Microsoft Research Health Futures Biomedical Computing team |
| Oct 2020 | Single cell workshop at Pfizer |
| Feb 2020 | Journal club at 10x Genomics |
| | Contributed talks |
| Apr 2021 | CZI Seed Networks Computational Biology Meeting |
| Mar 2021 | UC Berkeley Center for Computational Biology Retreat |
| Nov 2020 | CZI Seed Networks Annual Meeting |
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| | Posters |
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Cold Spring Harbor conference on Probabilistic Modeling in Genomics

UC Berkeley Center for Computational Biology Retreat (Best poster award)

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Other meetings and events

Apr 2021 SingleCellOpenProblems: jamboree for expanding OpenProblems framework

Nov 2020 AstraZeneca single-cell genomics group tutorial Nov 2020 UC Berkeley Computational Biology skills seminar

Nov 2019 Normjam: normalization workshop for scRNA-seq data, CZI/NY Genome Center

Employment

Jan – Jun 2018 Junior Computational Biologist. Sloan Kettering Institute

Advisor: Dana Pe'er. Developed an unsupervised classifier to detect doublets, technical artifacts in single-cell RNA-seq data, and built infrastructure to process and distribute

single-cell datasets to collaborators.

Summer 2017 Research Assistant. SLOAN KETTERING INSTITUTE

Advisor: Dana Pe'er. Designed methods to identify genes predictive of cell fate decisions in single-cell trajectory pseudotime data, as well as clustering of genes based on pseu-

dotemporal patterns.

Spring 2016 Research Assistant. UC BERKELEY SCHOOL OF PUBLIC HEALTH

Advisor: Lee W. Riley. Developed pipeline to process and analyze RNA-seq data with

application to carbapenem-resistant bacteria.

Software

2019 - scvi-tools - A library for deep probabilistic analysis of single-cell omics data.

2017 - DoubletDetection - A package for detecting doublets in scRNA-seq data.

Teaching experience

Fall 2021 Introduction to Computational Molecular and Cell Biology (BIO ENG C131)

Graduate Student Instructor, University of California, Berkeley

Spring 2021 Doctoral Seminar in Computational Biology (COMPBIO 293)

Graduate Student Instructor, University of California, Berkeley

University service

2020 – 2021 Admissions committee for UC Berkeley Computational Biology Graduate Group

UC Berkeley Computational Biology Skills Seminar Coordinator

Journal and conference reviewing

PLOS Computational Biology, 2021

Machine Learning in Computational Biology Workshop (MLCB), 2020

Press

2020 - 2021

"TotalVI: A transformative algorithm". Berkeley Engineering News. Mar 2021.