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Information	Carnegie Mellon University	email: ftownes@andrew.cmu.edu
	Pittsburgh, PA USA	

#### Summary

I am an applied statistician working primarily with biomedical data. I am involved with infectious disease as part of the Delphi Group and have developed methods for spatial and single-cell genomics. I am also interested in wearable devices, ecology, and economics. I strive to produce open source software implementations to help scientists achieve their research goals more effectively.

Professional	2022 - present	Assistant Professor, Department of Statistics and Data Science
EXPERIENCE		Carnegie Mellon University, Pittsburgh, PA
	2019 - 2022	Postdoctoral Researcher, Department of Computer Science
		Princeton University, Princeton, NJ (advisor: Barbara Engelhardt)
	2011 - 2014	Software Test Engineer, SRA International, Arlington, VA
	2008 - 2010	Software Analyst, Perrin Quarles Associates, Charlottesville, VA
	2007 - 2008	Fulbright Scholar, University of the Philippines
		Conducted independent forest ecology research while living with an
		indigenous Aeta community in Bataan Province.

## EDUCATION

2019 Ph.D. Biostatistics, Harvard University, Cambridge, MA USA

Dissertation Committee: Rafael Irizarry (Dana-Farber Cancer Institute, Data Science)

Jeff Miller (Harvard University, Biostatistics)

Martin Aryee (Massachusetts General Hospital, Pathology)

2013 M.S. Mathematics & Statistics, Georgetown University, Washington, DC USA

2007 B.S. Biology, Washington & Lee University, Lexington, VA USA

## Research In Review

- [1] Jones A, **Townes FW**, Li D, Engelhardt BE. Alignment of spatial genomics and histology data using deep Gaussian processes (2022). Preprint: doi:10.1101/2022.01.10.475692.
- [2] Verpeut JL, Bergeler S, Kislin M, **Townes FW**, Klibaite U, Dhanerawala ZM, Hoag A, Jung C, Lee J, Pisano TJ, Seagraves KM, Shaevitz JW, Wang SSH. Cerebellar contributions to a brainwide network for flexible behavior (2021). Preprint: doi:10.1101/2021.12.07.471685.

# In Press

[1] **Townes FW**, Engelhardt BE. Nonnegative spatial factorization. *Nature Methods* (2022). Preprint: https://arxiv.org/abs/2110.06122.

# Peer-reviewed Journal Articles

- [1] Gewirtz ADH, **Townes FW**, and Engelhardt BE. Telescoping bimodal latent Dirichlet allocation to identify expression QTLs across tissues. *Life Science Alliance* (2022). doi:10.26508/lsa.202101297.
- [2] Jones A, Townes FW, Li D, Engelhardt BE. Contrastive latent variable modeling with application to case-control sequencing experiments. Annals of Applied Statistics (2022). doi:10.1214/21-AOAS1534.
- [3] Hecker J, Townes FW, Kachroo P, Lasky-Su J, Ziniti J, Cho MH, Weiss ST, Laird NM, Lange C. A unifying framework for rare variant association testing in family-based designs, including higher criticism approaches, SKATs, and burden tests. *Bioinformatics* (2020). doi:10.1093/bioinformatics/btaa1055.
- [4] Townes FW, Carr K, Miller JW. Identifying longevity associated genes by integrating gene expression and curated annotations. *PLOS Computational Biology* (2020). doi:10.1371/journal.pcbi.1008429.

- [5] **Townes FW**, Irizarry RA. Quantile normalization of single-cell RNA-seq read counts without unique molecular identifiers. *Genome Biology* (2020). doi:10.1186/s13059-020-02078-0.
- [6] Townes FW, Hicks SC, Aryee MJ, Irizarry RA. Feature selection and dimension reduction for single-cell RNA-Seq based on a multinomial model. *Genome Biology* (2019). doi:10.1186/s13059-019-1861-6.
- [7] Marsh DM, **Townes FW**, Cotter K, Farroni K, McCreary K, Petry R, Tilghman J. Thermal Preference and Species Range in Mountaintop Salamanders and Their Widespread Competitors. *Journal of Herpetology* (2019). doi:10.1670/18-110.
- [8] Hicks SC, **Townes FW**, Teng M, Irizarry RA. Missing Data and Technical Variability in Single-Cell RNA-Sequencing Experiments. *Biostatistics* (2018). doi:10.1101/025528.
- [9] Hecker, J, Xu X, **Townes FW**, Fier HL, Corcoran C, Laird N, Lange C. Family-Based Tests for Associating Haplotypes With General Phenotype Data: Application to Asthma Genetics. *Genetic Epidemiology* (2017). doi:10.1002/gepi.22094.
- [10] Valeri L, Patterson-Lomba O, Gurmu Y, Ablorh A, Bobb J, Townes FW, Harling G. Predicting Subnational Ebola Virus Disease Epidemic Dynamics from Sociodemographic Indicators. *PLOS One* (2016). doi:10.1371/journal.pone.0163544.
- [11] **Townes W**. Seed dispersal of the genus Leea in forest patches of Bataan, Philippines. *Ecotropica* (2010).

# Technical Reports

- [1] **Townes FW**. Review of Probability Distributions for Modeling Count Data. arXiv (2020). https://arxiv.org/abs/2001.04343.
- [2] **Townes FW**. Generalized Principal Component Analysis. *arXiv* (2019). https://arxiv.org/abs/1907.02647.

#### Talks Invited

- [1] Biomedical Informatics Department, University of Colorado Anschutz Medical Center. October 2022.
- [2] Advanced Biomedical Computation Seminar, Computational Pathology Department, Brigham and Women's Hospital. March 2022.
- [3] Statistics and Data Science Department, Carnegie Mellon University. March 2022.
- [4] Biomedical Data Science Department, Stanford University. February 2022.
- [5] Biostatistics Department, University of Michigan. January 2022.
- [6] Population and Public Health Sciences Department, University of Southern California. January 2022.
- [7] Emerging Concepts in Microbial Informatics, Molecular Medicine and Biotechnology (panelist). Trinity University of Asia. November 2021.
- [8] Don't Normalize: the GLM-PCA approach to normalization. *Normjam*, New York Genome Center. November 2019.
- [9] Biostatistics and Data Science as a Career Path. Biology Department, Washington & Lee University. October 2019.

# Contributed Talks

[1] Latent factorization methods for genomics. New England Statistical Symposium (virtual presenter). May 2022.

- [2] Nonnegative process factorization for multivariate spatial count data. *Joint Statistical Meetings* (virtual conference). August 2021.
- [3] Quantile normalization of single-cell RNA-seq read counts without unique molecular identifiers. Joint Statistical Meetings (virtual conference). August 2020.
- [4] Dimension reduction for massive single-cell datasets. *Bioconductor Annual Meeting* (virtual conference). July 2020. slides doi:10.7490/f1000research.1118084.1
- [5] Dimension Reduction for Single Cell RNA-Seq based on a Multinomial Model. *Joint Statistical Meetings* (Denver, CO). August 2019.
- [6] Varying-Censoring Aware Matrix Factorization for Single Cell RNA-Sequencing. *Bioconductor Annual Meeting* (Boston, MA). July 2017.

#### Seminars

- [1] Human tumor atlas network steering committee meeting. April 2022.
- [2] Nonnegative spatial factorization. Hansen and Hicks labs, Department of Biostatistics, Johns Hopkins Bloomberg School of Public Health. November 2021.
- [3] Primer: Generalized linear models and latent factor models. *Broad Institute Models, Inference, and Algorithms*. October 2020.
- [4] Feature Selection and Dimension Reduction for Single Cell RNA-Seq. Greene Lab, Department of Systems Pharmacology and Translational Therapeutics, Perelman School of Medicine, University of Pennsylvania. February 2019.
- [5] Unsupervised Learning for Single Cell Gene Expression. *Harvard Biostatistics Cancer Working Group*. September 2018.
- [6] Single Cell Housekeeping Genes and Normalization. Dana-Farber Cancer Institute cBio Seminar. April 2018.
- [7] Informative Missing Data in Single Cell RNA-Seq. Dana-Farber Cancer Institute Genomics Seminar. October 2016.
- [8] Family Based Association Tests for Rare Variants. Brigham & Women's Hospital Channing Network Medicine Seminar. September 2015.

# Interviews

[1] The Bioinformatics Chat. March 2020. https://bioinformatics.chat/glm-pca

# Posters

- [1] Townes FW, Shukla C. Gene Expression Autoencoders. Harvard Biomedical Informatics 707 (Deep Learning in Healthcare). April 2018.
- [2] Townes FW, Marquez-Luna C. Mixture of Experts Analysis of Infectious Disease Outbreak Characteristics. Harvard Computer Science 281 (Advanced Machine Learning). December 2015.
- [3] Townes FW, Karaayvaz M, Gillespie S, Bernstein B, Ellisen L, Aryee M. Single Cell RNA-Seq Technical and Biological Confounders. Program in Quantitative Genomics Conference, Harvard Medical School. November 2015.

PEER REVIEW **Journals**: American Journal of Human Genetics, Bioinformatics, Biostatistics, Genome Biology, Journal of Machine Learning Research, NAR Genomics and Bioinformatics, Nature Communications

Professional American Statistical Association, International Society for Computational Biology, Sigma Xi Societies

FUNDING	2017 2016 - 2018 2016	NIH T32 Training Grant: Cancer Chan-Zuckerberg Foundation Travel Grant Human Cell Atlas Jamboree. European Bioinformatics Institute, Hinxton, UK NIH T32 Training Grant: Big Data to Knowledge NSF-CBMS Travel Award: Topology, Geometry, and Statistics NIH T32 Training Grant: Clinical Epidemiology of Lung Diseases
TEACHING EXPERIENCE	2021 Fall 2020 Fall 2018 Fall 2018 Summer	Guest lecture, Advanced Computational Genomics (Princeton - COS 597D) Guest lecture, Computational Biology of Single Cells (Princeton - COS 597F) Teaching Assistant, Applied Regression (Harvard - BST 210) Grade homeworks, hold office hours, and teach a weekly lab section. Co-Instructor, StatStart (Harvard) Statistics program for under-represented high school students.
	2017 Fall	Gave two interactive lectures on graphing data and regression. <b>Teaching Assistant</b> , Applied Bayesian Analysis (Harvard - BST 228)  Grade homeworks, provide solutions, and hold office hours.
	2017 Summer	Lead Instructor, Introduction to Data Science (PARSE Ltd.) Nonprofit, week-long statistics program for high school students. Developed course material and gave interactive lectures using R.
	2017 Spring	Teaching Assistant, Applied Longitudinal Analysis (Harvard - BST 226) Grade homeworks and hold office hours.
	2016 Fall	<b>Teaching Assistant</b> , Intro to Statistical Methods (Harvard - BST 201) Grade homeworks, provide solutions, hold office hours, and teach a weekly lab section.
	2016 Summer	Project Mentor, Pipelines into Biostatistics (Harvard) Program for under-represented undergraduates. Guided three students on a pharmacogenomics data analysis.
	2016 Spring	<b>Teaching Assistant</b> , Rates and Proportions (Harvard - BST 210) Grade homeworks, provide solutions, hold office hours, and teach a weekly lab section.

# SOFTWARE PACKAGES

- [1] **glmpca**: dimension reduction for non-normally distributed data. R package: https://cran.r-project.org/package=glmpca, python package: https://pypi.org/project/glmpca/
- [2] scry: Small-Count Analysis Methods for High-Dimensional Data. Bioconductor R package: https://bioconductor.org/packages/release/bioc/html/scry.html
- [3] **quminorm**: Quantile normalization of non-UMI single cell read counts. R package: https://github.com/willtownes/quminorm
- [4] **spatial-factorization**: Spatially-aware probabilistic factor models. Python package: https://github.com/willtownes/spatial-factorization-py

TECHNICAL **Programming**: R, Python, SQL, Git, IATEX, Shell SKILLS **Operating Systems**: MacOS, Linux, Windows

Machine Learning Frameworks: Caret, Scikit-learn, Keras, Tensorflow, Stan