Contact	Department of Statistics and Data Science	website: willtownes.github.io
Information	Carnegie Mellon University	email: ftownes@andrew.cmu.edu
	Pittsburgh, PA USA	

SUMMARY

I am an applied statistician working primarily with biomedical and public health data. I am involved with infectious disease projects 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.

Professional Experience	2022 - present	Assistant Professor, Department of Statistics and Data Science
		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)

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

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

Research In Review

2013

- [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] Gewirtz ADH, Townes FW, Engelhardt BE. Expression QTLs in single-cell sequencing data (2022). Preprint: doi:10.1101/2022.08.14.503915.

Peer-reviewed Journal Articles

- [1] Verpeut JL, Bergeler S, Kislin M, **Townes FW**, Klibaite U, Dhanerawala ZM, Hoag A, Janarthanan S, Jung C, Lee J, Pisano TJ, Seagraves KM, Shaevitz JW, Wang SSH. Cerebellar contributions to a brainwide network for flexible behavior in mice. *Communications Biology* (2023). doi:10.1038/s42003-023-04920-0
- [2] **Townes FW**, Engelhardt BE. Nonnegative spatial factorization applied to spatial genomics. *Nature Methods* (2023). doi:10.1038/s41592-022-01687-w
- [3] 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
- [4] 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
- [5] 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

- [6] 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
- [7] **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
- [8] 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.
- [9] 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.
- [10] 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.
- [11] 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.
- [12] 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.
- [13] **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] Biomedical Data Science Department, Stanford University. February 2022.
- [4] Biostatistics Department, University of Michigan. January 2022.
- [5] Population and Public Health Sciences Department, University of Southern California. January 2022.
- [6] Emerging Concepts in Microbial Informatics, Molecular Medicine and Biotechnology (panelist). Trinity University of Asia. November 2021.
- [7] Don't Normalize: the GLM-PCA approach to normalization. *Normjam*, New York Genome Center. November 2019.
- [8] 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] Introduction to single cell genomics. (informal) CMU Machine Learning Department lunch chats. May 2023.
- [2] Wastewater based epidemiology. Auton lab. January 2023.
- [3] Human tumor atlas network steering committee meeting. April 2022.
- [4] Nonnegative spatial factorization. Hansen and Hicks labs, Department of Biostatistics, Johns Hopkins Bloomberg School of Public Health. November 2021.
- [5] Primer: Generalized linear models and latent factor models. *Broad Institute Models, Inference, and Algorithms.* October 2020.
- [6] 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.
- [7] Unsupervised Learning for Single Cell Gene Expression. *Harvard Biostatistics Cancer Working Group*. September 2018.
- [8] Single Cell Housekeeping Genes and Normalization. Dana-Farber Cancer Institute cBio Seminar. April 2018.
- [9] Informative Missing Data in Single Cell RNA-Seq. Dana-Farber Cancer Institute Genomics Seminar. October 2016.
- [10] 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

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

THESIS COMMITTEES

- Aaron Rumack, CMU Machine Learning Department (2023)
- Timothy Barry, CMU Statistics Department (2023)

OTHER SERVICE ACTIVITIES

• Seminar Committee (2022-present)

Professional Societies	American Statistical Association, International Society for Computational Biology, Sigma Xi	
Funding	2018 - 2019	NIH T32 Training Grant: Cancer
		Chan-Zuckerberg Foundation Travel Grant
		Human Cell Atlas Jamboree. European Bioinformatics Institute, Hinxton, UK
	2016 - 2018	NIH T32 Training Grant: Big Data to Knowledge
	2016	NSF-CBMS Travel Award: Topology, Geometry, and Statistics
	2014 - 2016	NIH T32 Training Grant: Clinical Epidemiology of Lung Diseases
Teaching	2022 G	
EXPERIENCE	2023 Spring	Lead Instructor, Introduction to Statistical Computing (CMU - 36-350)
	2021 Fall	Guest lecture, Advanced Computational Genomics (Princeton - COS 597D)
	2020 Fall 2018 Fall	Guest lecture, Computational Biology of Single Cells (Princeton - COS 597F) Too shing: Against Applied Repression (Harvard, RST 210)
	2018 Fall	Teaching Assistant, Applied Regression (Harvard - BST 210)
	2018 Summer	Grade homeworks, hold office hours, and teach a weekly lab section. Co-Instructor, StatStart (Harvard)
	2016 Summer	Statistics program for under-represented high school students.
		Gave two interactive lectures on graphing data and regression.
	2017 Fall	Teaching Assistant, Applied Bayesian Analysis (Harvard - BST 228)
	2011 1 411	Grade homeworks, provide solutions, and hold office hours.
	2017 Summer	, <u>.</u>
	2011 Summer	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)
	1 0	Grade homeworks and hold office hours.
	2016 Fall	Teaching Assistant, Intro to Statistical Methods (Harvard - BST 201)
		Grade homeworks, provide solutions, hold office hours, and teach a weekly lab section.
	2016 Summer	
		Program for under-represented undergraduates.
		Guided three students on a pharmacogenomics data analysis.
	2016 Spring	Teaching Assistant, 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 SKILLS

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

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