## Justin Silverman

Contact jsilve24@gmail.com justin-silverman.com Information statsathome.com Research Bayesian statistics, time series analysis, multivariate analysis, sequence count data, Interests (e.g., microbiome surveys, gene expression studies, single cell RNA-seq), epidemiology, compositional data analysis, decision theory. Pennsylvania State University 2020-EMPLOYMENT Assistant Professor, College of Information Science and Technology (Primary) Assistant Professor, Department of Medicine (Secondary) MD and PhD, Duke University 2012-2020 **EDUCATION** PhD in Computational Biology and Bioinformatics Thesis: Bayesian Multivariate Count Models for the Design and Analysis of Microbiome Studies Mentors: Dr. Lawrence David and Dr. Savan Mukherjee Research Technician, Johns Hopkins University 2011-2012 Mentor: Dr. Nina Markovic Bachelors of Science, Johns Hopkins University 2007-2011 Double Major in Physics and Biophysics, Minor in Mathematics Publications [\* denotes equal contribution]

**Silverman JD\***, Hupert N, Washburne AD\*. Using influenza surveillance networks to estimate state-specific case detection-rates and forecast SARS-CoV-2 spread in the United States. (2020) medRxiv, DOI: 10.1101/2020.04.01.20050542

Morton J\*, Marotz C\*, Washburne A, **Silverman JD**, Edlund A, Zaramela L, Zengler K, Knight R. (2019) Establishing microbial measurement standards with reference frames. *Nature Communications*, 2719; DOI: 0.1038/s41467-019-10656-5

Villa MM\*, Bloom RJ\*, **Silverman JD**, Durand HK, Jiang S, Wu A, Huang S, You L, David LA. (2019) High-throughput isolation and culture of human gut bacteria with droplet microfluidics. *bioRxiv* 630822; DOI:10.1101/630822

**Silverman JD**, Bloom RJ, Jiang S, Durand HK, Mukherjee S, David LA. (2019) Measuring and Mitigating PCR Bias in Microbiome Data. *bioRxiv* 604025; DOI:10.1101/604025

**Silverman JD**, Roche K, Holmes ZC, David LA, Mukherjee S. (2019) Bayesian Multinomial Logistic Normal Models Through Marginally Latent Matrix-T Processes. *arXiv*:1903.11695

Silverman JD, Roche K, Mukherjee S, David LA. (2018) Naught all zeros in sequence count data are the same. bioRxiv 477794; DOI:10.1101/477794.

**Silverman JD**\*, Shenhav L\*, David LA, Halperin E, Mukherjee S. (2018) Statistical Considerations in the Design and Analysis of Longitudinal Microbiome Studies. *bioRxiv* 448332; DOI:10.1101/448332 (*Invited Review at mSystems*)

**Silverman JD**, Durand HK, Bloom RJ, Mukherjee S, David LA. (2018) Dynamic linear models guide design and analysis of microbiota studies within artificial human guts. *Microbiome*. 6:202 DOI:10.1186/s40168-018-0584-3

Shenhav L, Furman O, Bariscoe L, Thompson M, Silverman JD, Mizrahi I, Halperin E. (2018) Modeling the temporal dynamics of the gut microbial community in adults and infants. bioRxiv 212993. DOI:10.1101/212993. (Submitted to PLoS Computational Biology)

Washburne AD, **Silverman JD**, Morton JT, Becker D, Crowley D, Mukherjee S, David LA, Plowright R. (2018) Phylofactorization - a graph partitioning algorithm to identify phylogenetic scales of ecological data. *Ecological Monographs* 00(00):e01353. DOI:10.1002/ecm.1353

Silverman JD, Silverman RK. (2018) The Bayesian Sorting Hat: A Decision-Theoretic Approach to Size-Constrained Clustering. arXiv:1710.06047. (In Revision at Journal of Machine Learning Research)

Midani FS, Weil AA, Chowdhury F, Begum YA, Khan AI, Debela, MD, Durand HK, Reese AT, Nimmagadda SN, **Silverman JD**, Ellis CN, Ryan ET, Calderwood SB, Harris JB, Qadri F, David LA, LaRocque RC. (2018) Human gut microbiota predicts susceptibility to *Vibrio cholerae* infection. *The Journal of Infectious Diseases*. 218:4 DOI:10.1093/infdis/jiy192

Silverman JD, Washburne AD, Mukherjee S, David LA. (2017) A phylogenetic transform enhances analysis of compositional microbiota data. eLife 6:e21887 DOI: 10.7554/eLife.21887

Washburne AD, Silverman JD, Leff JW, Bennett DJ, Darcy JL, Mukherjee S, Fierer N, David LA. (2017) Phylogenetic factorization of compositional data yields lineage-level associations in microbiome datasets. *PeerJ* 5:e2969 DOI:10.7717/peerj.2969

## PATENTS AND SOFTWARE

**Justin D. Silverman** (2019) Multinomial logistic-normal linear models using a marginal Laplace approximation (*stray*)

**Justin D. Silverman** (2019) R/C++ Header Library for Solving Minimum Cost Bipartite Matching Problems (RcppHungarian). R package Version 0.1. https://cran.r-project.org/web/packages/RcppHungarian/index.html

**Justin D. Silverman** (2019) R/C++ Header Library for Compositional Data Analysis (*RcppCoDA*)

**Justin Silverman** (2016) philr: Phylogenetic partitioning based ILR transform for metagenomics data. R package version 1.0.0. http://bioconductor.org/packages/philr/(Downloaded 2951 times; September 6th, 2018).

**Justin David Silverman**, Adam Sean Jermyn, Nina Markovic. (2015) System for Lightweight Image Processing. U.S. Patent 9,097,739.

Funded Research		alth Innovation, Innovation Award k for Predicting Bacteremia and ulture Results"	2020-2021
	and Clinical Sciences Authored (Submitting	tative Approaches to Problems in Basic g PIs: Lawrence David and Sayan Mukherjee) ch to modeling human microbiota dynamics"	2016-2017
Honors and Awards	Mitchell Meritorious I Phi Beta Kappa (JHU Donald E. Kerr Memo H. Keffer Hartline Av JHU Office of Greek I Goldwater Scholar Provost Undergradua Materials Research Sc	cion (Compositional Data Analysis Workshop) Research Travel Award U Highest Honors for Arts and Sciences) orial Award for Excellence in Physics vard for Outstanding Scholarship in Biophysics Life Senior Academic Excellence Award te Research Award cience and Engineering Center, Summer REU sics Honor Society, Invitation Only)	2017 2017 2011 2011 2011 2011 2010 2010
SERVICE	Referee Session Organizer	publons.com/author/1257294/justin-d-silverman Annals of Applied Statistics PLoS Genetics PLoS Computational Biology mSystems Microbiome SORT NAR Genomics and Bioinformatics IMS Invited Session at JSM 2018 "Longitudinal Modeling and Experimental Desig Investigating Host Associated Microbiota"	
Invited Talks	ENAR Compositional Nature of Microbiome Data: Challenges and New Symposium on Advances in Genomics, Epidemiology and Statistics: Phi Biostatistics Seminar Series at University of Massachusetts: Amherst, M. Simons Foundation: New York, NY Penn State University: State College, PA University of Chicago: Chicago, IL Computational Molecular Biology Seminar at Brown: Providence, RI Center of Statistical Genetics Seminar, Univ of Michigan: Ann Arbor, M. IMS Invited Session at JSM: Vancouver, BC Probabilistic Microbial Modeling Symposium: San Diego, CA ENAR Geometry and Topology in Statistical Inference: Atlanta, GA Wharton Statistics Seminar Series: Philadelphia, PA Universitat Politècnica de Catalunya: Barcelona, Spain Seminar in Compositional Data: Girona, Spain Health Analytics Workshop at Duke University: Durham, NC		iladelphia 2019 IA 2019 2019 2019 2019 2019 2018
Conference Talks		n and therapeutic effect in human microbiota, The mpositional Data Analysis, Abbadia San Salvatore,	

RESEARCH MENTORSHIP	Kim Roche (Graduate) Katie Sparks (Undergraduate) Chris Weddel (High School) Adam Jermyn (High School) Roberto Rivera (Undergraduate)	2018-2019 2012 2011-2012 2011-2012 2011
TEACHING EXPERIENCE	DOmath 2018 Research Program: Stochastic Properties of Dynamical Systems: Project Manager	Summer 2018
	Human Microbiome Seminar Guest Lecturer, University of North Carolina, Chapel Hill	Spring 2017
	Statistical Methods in Computational Biology Teaching Assistant and Guest Lecturer, Duke University	Spring 2017
	Spectroscopy (Quantum Mechanics for Biophysics) Teaching Assistant, Johns Hopkins University	Fall 2011

## References

**Sayan Mukherjee**, Professor, Department of Statistical Science, Duke University, (919) 684-4608 sayan@stat.duke.edu

Lawrence David, Assistant Professor, Center for Genomic and Computational Biology, Duke University, (919) 668-5388 lawrence.david@duke.edu

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