**Tyler Grimes**

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# Education

**Ph.D. Biostatistics** 2016 – 2020

University of Florida, Gainesville FL

Dissertation Advisor: Somnath Datta

Thesis: “Some contributions to the differential network analysis of –omics data”

**M.S. Mathematics**, concentration in Statistics 2014 – 2016

University of North Florida, Jacksonville FL

Thesis: “A saddlepoint approximation to hypothesis test of variance for non-normal populations” (Thesis Advisor: Ping Sa)

**B.S. Mathematics**, minor in Computer Science 2010 – 2014

University of Central Florida, Orlando FL

# Employment

**Senior Bioinformatics Analyst** 2021 – present

The Emmes Company

* Provide statistical guidance for developing clinical study protocols with -omics endpoints, ensuring the design and methods are scientifically valid and feasible.
* Author statistical analysis plans (SAPs) to outline analysis strategies and algorithms for -omics data analysis.
* Develop workflows for the processing, analysis, and management of -omics data.
* Author study reports and manuscripts for research utilizing -omics endpoints.

**Assistant Professor of Statistics** 2020 – 2023

University of North Florida, Department of Mathematics and Statistics

* Conducted collaborative and independent research focused on applying probabilistic graphical models to high-dimensional genomics data.
* Taught graduate and undergraduate courses on statistics methods, design of experiments, quality control, and probability.
* Directed graduate and undergraduate student projects. Titles included: “Probabilistic graphical models and their applications”, “Survival prediction using autoencoders and AFT models”, and “Unsupervised learning: principal component analysis.”

**Statistician** – VA IPA 2018 – 2020

U.S. Department of Veterans Affairs, Brain Rehabilitation Research Center

* Assisted in the design and analysis of experiments; coded simulations for sample size calculations; wrote the statistical analysis plan for grant applications; analyzed experimental data; reported and discussed results with collaborators.

**Graduate Research Assistant** 2016 – 2020

University of Florida, Department of Biostatistics

* Worked on various research projects involving high-dimensional data analysis, dimension reduction, high-throughput RNA-sequencing, metabolomics data, predictive modeling, data visualization, and differential network analysis.

**Graduate Teaching Assistant**  2014 – 2016

University of North Florida, Department of Mathematics and Statistics

# Teaching Experience

Assistant Professor, University of North Florida 2020 – present

* Courses taught include:
* Sampling (Graduate – STA 6226 and Undergraduate – STA 4664)
* Statistical Quality Control (Graduate/Undergraduate – STA 6666/4664)
* Probability and Statistics (Undergraduate – STA 4321)
* Statistical Methods I & II (Undergraduate – STA 3163/4, Graduate – STA 6166)
* Elementary Statistics for Health and Social Sciences (STA 2014)
* Directed student projects through Capstone and Independent Study. Titles include:
* “Probabilistic graphical models and their applications”
* “Survival prediction using autoencoders and AFT models”
* “Unsupervised learning: principal component analysis”

Temporary Instructor – Large Sample Theory, University of Florida Spring 2019, 2020

* Lectured topics included probability, modes of convergence, U-statistics, and MLE.

Instructor, University of North Florida Spring 2016

* Courses include: MAC 2233 (Calculus for Business).

Teaching Assistant, University of North Florida 2014 – 2016

* Assisted in elementary statistics courses by leading discussions in weekly breakout sessions, managing computer lab sessions, and holding weekly office hours.

# Publications

1. **Grimes, T.**, Walker, A., Datta, S., Datta, S., (2018) Predicting survival times for neuroblastoma patients using RNA-Seq expression profiles. *Biology Direct*. 13(1), 11
2. Walker, A., **Grimes, T.**, Datta, S., Datta, S., (2018) Unraveling bacterial fingerprints of city subways from microbiome 16S gene profiles. *Biology Direct*. 13(1), 10
3. **Grimes, T.**, Potter, S., Datta, S., (2019) Integrating gene regulatory pathways into differential network analysis of gene expression data. *Scientific Reports.* 9(1), 5479
4. Boissoneault, C., **Grimes, T.**, Rose, D., Waters, M.F., Khanna, A., Datta, S. and Daly, J.J., (2020) Innovative long-dose neurorehabilitation for balance and mobility in chronic stroke: a preliminary case series. *Brain Sciences*. 10(8), 555
5. **Grimes, T.**, Datta, S., (2021) SeqNet: an R package for generating gene-gene networks and simulating RNA-seq data. *Journal of Statistical Software.* 98(12), 10.18637/jss.v098.i12
6. Boissoneault, C., Rose, D. K., **Grimes, T**., Waters, M. F., Khanna, A., Datta, S., & Daly, J. J. (2021). Trajectories of stroke recovery of impairment, function, and quality of life in response to 12-month mobility and fitness intervention. *NeuroRehabilitation*. 49(4), 573-584
7. Ahn, S., **Grimes, T.**, Datta, S. (2021) The analysis of gene expression data incorporating tumor purity information. *Frontiers in Genetics.* 12(1), 642759
8. **Grimes, T.**, Datta, S. (2021) A novel probabilistic generator for large-scale gene association networks. *PLoS One*. 16(11), e0259193
9. Ahn, S., **Grimes, T.**, & Datta, S. (2023). A pseudo-value regression approach for differential network analysis of co-expression data. *BMC Bioinformatics*. 24(1), 8
10. Goll, J. B., Bosinger, S. E., Jensen, T. L., Walum, H., **Grimes, T.**, Tharp, G. K., ... & Hoft, D. F. (2023). The Vacc-SeqQC project: Benchmarking RNA-Seq for clinical vaccine studies. *Frontiers in Immunology*. 13(1), 1093242
11. Waid-Ebbs, J. K., Wen, P. S., **Grimes, T.**, Datta, S., Perlstein, W. M., Hammond, C. S., & Daly, J. J. (2023). Executive function improvement in response to meta-cognitive training in chronic mTBI/PTSD. *Frontiers in Rehabilitation Sciences*. 4(1), 1189292
12. DeZern, A. E., Goll, J. B., Lindsley, R. C., Bejar, R., Wilson, S. H., Hebert, D., ..., **Grimes, T.**, … & Walter, M. J. (2023). Utility of targeted gene sequencing to differentiate myeloid malignancies from other cytopenic conditions. *Blood Advances*. 7(14), 3749-3759

## Conference Proceedings:

1. **Grimes, T.**, Datta, S., (2019) A random graph generation model for transcription networks and nonparametric simulator for RNA-seq expression data. *Computer Data Analysis and Modeling*. Proceedings of the XII International Conference, Minsk. September 18-22, 2019, Eds: P. Filzmoser and Y. Kharin, pp. 37-42. ISBN 978-985-566-811-5.

## In Progress:

1. **Grimes, T.**, Datta, S., dnapath: an R package for pathway-based differential network analysis*.*
2. **Grimes, T.**, Datta, S., (tentative) Inferring dynamic networks from scRNA-seq data
3. **Grimes, T.**, Datta, S., (tentative) Identifying cancer driver genes from differential co-expression networks.
4. **Grimes, T.**, Ping, S., A saddlepoint approximation to hypothesis tests of variance for non-normal populations.

# Presentations

## Talks:

1. University of Maryland, statistics colloquium invited talk, virtual seminar 2022

“Predictive modeling with differential co-expression of RNA-seq data”

1. UNF Biomedical Sciences Research in Progress seminar, Jacksonville, FL. 2021

“Identifying cancer driver genes from differential co-expression networks”

1. CMStatistics International Conference, virtual conference 2020

“Identifying cancer driver genes from differential co-expression networks”

1. ISMB/CAMDA International Conference, Chicago, IL 2018 “An exploratory approach for identifying novel biomarkers in high-risk cancer patients from RNA-seq data”
2. ISMB/ECCB International Conference, Prague, Czech Republic 2017 “Predicting survival times for neuroblastoma patients using RNA-Seq expression profiles”

## Posters:

1. JSM, Denver, CO. 2019 “Integrating gene regulatory pathways into differential network analysis of gene expression data”
2. UF College of PHHP Research Day, Gainesville, FL. 2019 *“*A pathway-based approach to differential network analysis of gene expression data”
3. IISA International Conference on Statistics, Gainesville, FL. 2018 “Predicting Survival Times for Neuroblastoma Patients Using RNA-Seq Expression Profiles”
4. UF College of PHHP Research Day, Gainesville, FL. 2018 *“*Predicting Survival Times for Neuroblastoma Patients Using RNA-Seq Expression Profiles”
5. UF College of Medicine Celebration of Research, Gainesville, FL. 2018 *“*Differential network analysis based on next-generation sequencing data”
6. UF Graduate Student Research Day, Gainesville, FL. 2018 *“*Predicting Survival Times for Neuroblastoma Patients Using RNA-Seq Expression Profiles”
7. UF Dept. of Statistics Winter Workshop, Gainesville, FL. 2018 *“*Differential network analysis based on next-generation sequencing”
8. JSM, Baltimore, MD. 2017 *“*Differential network analysis based on next-generation sequencing”
9. FaceBase Scientific Meeting, Boston, MA. 2017 *“*A differential network analysis of palatal development”

# Service

**Committee** **Chair** – Undergraduate Committee 2021 – 2023

Department of Mathematics and Statistics, University of North Florida

**Founding** **President** - Biostatistics Student Organization 2017 – 2019

Department of Biostatistics, University of Florida

**Executive Board Member** - Doctoral Student Council 2017 – 2019

College of Public Health and Health Professionals, University of Florida

**Student Representative - Department of Biostatistics 2019**

**College of PHHP Development Board Meeting, University of Florida**

**Student Representative - Department of Biostatistics 2017**

**College of PHHP Advisory Board Member Meeting, University of Florida**

# Projects

My [github](https://github.com/tgrimes) account hosts various projects including course projects such as a [stochastic approximation algorithm for minimum graph bisection](https://github.com/tgrimes/UF-PHC6068/blob/master/Project/SAA_and_SAMC_for_minimum_graph_bisection.pdf); conference challenges including the [CAMDA 2017 Neuroblastoma challenge](https://github.com/tgrimes/CAMDA-2017-Neuroblastoma); and other personal projects.

My methodological research is implemented in various R packages. These include [SeqNet](https://cran.r-project.org/web/packages/SeqNet/index.html), an R package for simulating RNA-sequencing data from any underlying association network; and [dnapath](https://cran.rstudio.com/web/packages/dnapath/index.html), an R package that performs differential network analysis based on known pathway information, which includes methods for easily summarizing and visualizing the results.

# Professional Memberships

American Statistical Association (ASA)

International Society for Computational Biology (ISCB)

# Journal Referee

BioData Mining

Bioinformatics

BMC Bioinformatics

Briefings in Bioinformatics

Communications in Statistics - Simulation and Computation

Scientific Reports

GigaScience

# Honors and Awards

Outstanding Dissertation Award 2020

Department of Biostatistics, University of Florida

Student Travel Award(multiple awards) 2017, 2019

Department of Biostatistics, University of Florida

and by College of Public Health and Health Professionals, University of Florida

Travel Fellowship 2017, 2018

Granted by CAMDA committee to present at ISMB/CAMDA 2018, Chicago, Illinois

and at ISMB/ECCB 2017, Prague, Czech Republic

Department of Biostatistics, University of Florida

Outstanding Graduate Student in Statistics 2016

University of North Florida

UCF Scholars Award(scholarship and summer grant) 2012 – 2013

University of Central Florida

Florida Academic Scholars Award(four-year scholarship) 2010 – 2014

# Funding

**U.S. Department of Veterans Affairs IPA**, “Analysis of Traumatic Brain Injury and other Brain Rehabilitation Data”, September 2018 – August 2019; VA PIs: Julia Waid-Ebbs and Janis Daly

**National Institutes of Health**, “Exploratory Statistical Analysis of Differential Network Behaviors based on Gene Expression Atlas of Palate Development”, August 2016 - July 2019; PI: Somnath Datta