# Matthew N. Bernstein

Morgridge Institute for Research Regenerative Biology Group Madison, WI 53715  $\begin{tabular}{ll} mbernstein@morgridge.org\\ mbernste.github.io \end{tabular}$ 

Madison, WI

Madison, WI

Madison, WI 2014 - 2019

2019 - Present

Aug. 2019

Education University of Wisconsin - Madison

Ph.D., Computer Sciences

Advisor: Colin Dewey, Ph.D.

University of Wisconsin - Madison Madison, WI M.S., Computer Sciences Dec. 2015

University of Notre Dame

B.S., Computer Science

South Bend, IN

May 2013

Experience Morgridge Institue for Research

Postdoctoral Fellow

Advisors: Christina Kendziorski, Ph.D. Ron Stewart, Ph.D.

University of Wisconsin - Madison

Research Assistant Advisor: Colin Dewey, Ph.D.

AmazonSeattle, WASoftware Development Engineering InternSummer 2014

University of Wisconsin - Madison Madison, WI

Teaching Assistant (Lecturer) Fall 2013, Spring 2014

Course taught: CS302 - Introduction to Programming

Amazon New York City, NY

Software Development Engineering Intern Summer 2013

Space and Naval Warfare Systems CommandSan Diego, CAResearch InternSummer 2012

Awards and NIH/BD2K Young Investigator Travel Scholarship

Fellowships International Conference on Intelligent Systems for Molecular Biology Chicago, IL, July 6-10, 2018

#### Best Plenary Talk

Tau Beta Pi

National Library of Medicine Informatics Training Conference San Diego, CA, June 6, 2017

#### **CIBM Training Program**

University of Wisconsin - Madison Feb. 2015 - Feb. 2018

#### University Housing Honored Instructor Award

University of Wisconsin – Madison Fall 2013

## Professional Memberships

International Society for Computational Biology Upsilon Pi Epsilon

## Peer-reviewed Publications

Bernstein, M.N., Doan, A., Dewey, C.N. (2017). MetaSRA: normalized human sample-specific metadata for the Sequence Read Archive. *Bioinformatics*, 33(18), 2914–2923.

## Pre-prints

Bernstein, M.N., Gladstein A., Latt K.Z. *et al.* (2020). Jupyter notebook-based tools for building structured datasets from the Sequence Read Archive [version 1; peer review: awaiting peer review]. *F1000Research*, 9:376

Bernstein, M.N. and Dewey, C.N. (2019). Hierarchical cell type classification using mass, heterogeneous RNA-seq data from human primary cells. *bioRxiv*.

## Talks and Seminars

Guest lecture for STAT 877 - Statistical Methods for Molecular Biology Cellular phenotyping with public, heterogeneous RNA-seq data University of Wisconsin - Madison, Feb. 7, 2019

International Conference on Intelligent Systems for Molecular Biology MetaSRA: Normalized sample-specific metadata for the Sequence Read Archive Chicago, IL, July 7, 2018

Center for Predictive Computational Phenotyping Annual Retreat Cellular phenotyping with mass, heterogeneous transcriptomic data Madison, WI, May 31, 2018

#### Computation and Informatics in Biology and Medicine Seminar

Towards cell type prediction with public RNA-seq data University of Wisconsin - Madison, January 30, 2018

#### National Library of Medicine Informatics Training Conference

MetaSRA: Normalized sample-specific metadata for the Sequence Read Archive San Diego, CA, June 6, 2017

Center for Predictive Computational Phenotyping Annual Retreat

MetaSRA: Normalized sample-specific metadata for the Sequence Read Archive

Madison, WI, June 1, 2017

Poster Presentations Great Lakes Bioinformatics Conference

Madison, WI, May 19-22, 2019

RNA-Seq Summit

San Francisco, CA, April 25-27, 2017

National Library of Medicine Informatics Training Conference

Columbus, OH, June 27-28m 2016

Undergraduate Mentorship

Aparna Maddala (Georgia Institute of Technology)

Integrated Biological Sciences Summer Research Program, Summer 2019 Project: Hierarchical deconvolution of bulk RNA-seq data in the MetaSRA

Brett Van Tassel (Ramapo College of New Jersey)

Integrated Biological Sciences Summer Research Program, Summer 2017

Project: Compression of RNA-seq alignments

Kevin Liao (University of North Carolina at Chapel Hill)

Integrated Biological Sciences Summer Research Program, Summer 2016

Project: Comparing similarity measures over RNA-seq derived gene

expression profiles

Teaching

CS 302 - Introduction to Programming

University of Wisconsin - Madison, Fall 2013 & Spring 2014

Teaching Assistant (primary lecturer to class of approximately 30 students)

Service

Journal Reviewer: Nature Biotechnology, Bioinformatics

Committee Member, Prospective Student Welcome Weekend

Department of Computer Sciences, University of Wisconsin - Madison,

Spring 2016 & Spring 2014

Committee Chair, Prospective Student Welcome Weekend

Department of Computer Sciences, University of Wisconsin - Madison,

Spring 2015

Club Leader, After School Computer Programming Club

Stephen's Point Elementary School, Madison, WI, Spring 2015

Professional Activities Team Lead, NCBI's Single-cell in the Cloud Codeathon

New York Genome Center, NY, January 15-17, 2020

Team Lead, NCBI's RNA-Seq in the Cloud Codeathon

University of North Carolina at Chapel Hill, NC, March 11-13, 2019