# MATTHEW N. BERNSTEIN

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### **EDUCATION**

N.D. C 1. C.	D 2015 A 2010
Ph.D., Computer Sciences University of Wisconsin – Madison	Dec. 2015 – Aug. 2019 Madison, WI
M.S., Computer Sciences University of Wisconsin	Aug. 2013 – Dec. 2015 Madison, WI
<b>B.S.</b> <i>Magna Cum Laude</i> , <b>Computer Science</b> University of Notre Dame	Aug. 2009 – May 2013 South Bend, IN
Work Experience	
Postdoctoral Fellow  Morgridge Institute for Research  • Advised by Christina Kendziorski, Ph.D. and Ron Stewart, Ph.D.	2019 – Present Madison, WI
Research Assistant University of Wisconsin – Madison • Advised by Colin Dewey, Ph.D.	2014 - 2019 Madison, WI
Software Development Engineering Intern Amazon	Summer 2014 Seattle, WA
<b>Teaching Assistant (Lecturer)</b> University of Wisconsin – Madison	2013 - 2014 Madison, WI
Software Development Engineering Intern Amazon	Summer 2013 New York, NY
Research Intern Space and Naval Warfare Systems Command (SPAWAR)	Summer 2012 San Diego, CA
Awards and Fellowships	
NIH/BD2K Young Investigator Travel Scholarship International Conference on Intelligent Systems for Molecular Biology	July 6-10, 2018 Chicago, IL
<b>Best Plenary Talk</b> National Library of Medicine Informatics Training Conference	June 6, 2017 San Diego, CA
CIBM Predoctoral Fellowship University of Wisconsin – Madison	2015-2018 Madison, WI
University Housing Honored Instructor Award University of Wisconsin – Madison	Fall 2013 Madison, WI
PROFESSIONAL MEMBERSHIPS	

## **International Society for Computational Biology**

Tau Beta Pi

Upsilon Pi Epsilon

### † corresponding author

- 1. **Bernstein, M.N.**, Ni, Z., Collins, M., Burkard, M.E., Kendziorski, C., Stewart, R. (2021). CHARTS: A web application for characterizing and comparing tumor subpopulations in publicly available single-cell RNA-seq datasets. *BMC Bioinformatics*. 22(83).
- 2. **Bernstein, M.N.**, Ma, J., Gleicher, M., Dewey, C.N. (2021). CellO: Comprehensive and hierarchical cell type classification of human cells with the Cell Ontology. *iScience*, 24(1), 101913.
- 3. Overmyer, K.A., Shishkovaa, E., Miller, I.J., Balnisa, J., **Bernstein, M.N.**, Peters-Clarke, T.M., Meyer, J.G., Quan, Q., Muehlbauer, L.K., Trujillo1, E.A., Hei, Y., Chopra, A., Chieng, H.C., Anupama Tiwari, A., Judson, M.A., Paulson, B., Brademan, D.R., Zhu, Y., Serrano, L.R., Linke, V., Drake, L.A., Adam, A.P., Schwartz, B.S., Singer, H.A., Swanson, S., Mosher, D.F., Stewart, R., Coon, J.J., Jaitovich, A. (2021). Large-scale Multi-omic Analysis of COVID-19 Severity. *Cell Systems*, 12(1), 23-40.
- 4. **Bernstein, M.N.** †, Gladstein, A., Latt, K.Z., Clough, E., Busby, B., Dillman, A. (2020). Jupyter notebook-based tools for building structured datasets from the Sequence Read Archive. *F1000Research*, 9(376).
- 5. **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.

#### SUBMITTED MANUSCRIPTS

 Simonett, S.P., Shin, S., Bacher, R., Dong, C., Smith, L., Rabaglia, M., Stapleton, D., Schueler, K., Choi, J., Bernstein, M.N., Perez-Cerventes, C., Herring, J., Spaeth, J., Stein, R., Tessem, J., Kendziorski, C., Keles, S., Moskowitz, I., Attie, A., Keller, M. (2020). Identification of direct transcriptional targets of Nfatc2 that promote β-cell proliferation in human islets.

#### TALKS AND SEMINARS

Workshop on Computational Advances for Single-Cell Omics Data Analysis CellO: Comprehensive and hierarchical cell type classification of human cells with the Cell Ontology	Dec. 12, 2020 Virtual
International Conference on Intelligent Systems for Molecular Biology (ISMB)  MetaSRA: Normalized sample-specific metadata for the Sequence Read Archive	July 7, 2018 Chicago, IL
Center for Predictive Computational Phenotyping Annual Retreat  Cellular phenotyping with mass, heterogeneous transcriptomic data	May 31, 2018 Madison, WI
Computation and Informatics in Biology and Medicine Seminar Towards cell type prediction with public RNA-seq data	January 30, 2018 Madison, WI
Center for Predictive Computational Phenotyping Annual Retreat MetaSRA: Normalized sample-specific metadata for the Sequence Read Archive	June 1, 2017 Madison, WI

#### POSTER PRESENTATIONS

Cold Spring Harbor Laboratory conference on Biological Data Science	ence Virtual, Nov. 4-6, 2020
International Conference on Intelligent Systems for Molecular Bio	<b>ology</b> Virtual, July 13-16, 2020
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 27-28 2016

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Aparna Maddala (Georgia Institute of Technology)	Summer 2019
<ul> <li>Integrated Biological Sciences Summer Research Program</li> <li>Project: Hierarchical deconvolution of bulk RNA-seq data in the MetaSRA</li> </ul>	
<ul> <li>Brett VanTassel (Ramapo College of New Jersey)</li> <li>Integrated Biological Sciences Summer Research Program</li> <li>Project: Compression of RNA-seq alignments</li> </ul>	Summer 2017
<ul> <li>Kevin Liao (University of North Carolina at Chapel Hill)</li> <li>Integrated Biological Sciences Summer Research Program</li> <li>Project: Comparing similarity measures over RNA-seq derived gene expression properties.</li> </ul>	Summer 2016 profiles
Teaching .	
Guest lecture, STAT 877 - Statistical Methods for Molecular Biology Visualizing high dimensional data with dimensionality reduction	Sept. 22, 2020 University of Wisconsin – Madison
Guest lecture, STAT 877 - Statistical Methods for Molecular Biology Cellular phenotyping with public, heterogeneous RNA-seq data	Feb. 7, 2019 University of Wisconsin – Madison
Primary lecturer, CS 302 - Introduction to Programming	Aug. 2013 - May 2014 University of Wisconsin - Madison
<ul> <li>Primary lecturer to approx. 30 out 700 students enrolled in CS 302</li> <li>Designed programming assignments for all approx. 700 students enrolled</li> </ul>	in CS 302
SERVICE	
Journal Reviewer	Ongoing
<ul> <li>Bioinformatics (1 review)</li> <li>BMC Bioinformatics (3 reviews)</li> <li>Journal of Computational Biology (1 review)</li> <li>Nature Biotechnology (1 review)</li> <li>Nature Communications (1 review)</li> <li>Nucleic Acids Research (1 review)</li> </ul>	
Review Editor for Frontiers in Genetics	2021-Presen
Committee Member, Prospective Student Welcome Weekend  Department of Computer Sciences, University of Wisconsin – Madison  Organized the department's graduate student recruitment weekend  Coordinated lodging, transportation, and social events	Spring 2010
Committee Chair, Prospective Student Welcome Weekend  Department of Computer Sciences, University of Wisconsin – Madison  • Led the organization of the department's graduate student recruitment we	Spring 2015 eekend
<ul> <li>Club Leader, After School Computer Programming Club</li> <li>Stephen's Point Elementary School, Madison, WI</li> <li>Led an after school computer programming club for 4th and 5th graders</li> <li>Taught the Scratch computer programming language</li> </ul>	Spring 2015
Committee Member, Prospective Student Welcome Weekend Department of Computer Sciences, University of Wisconsin – Madison	Spring 2014
Professional Activities	
NCBI Single-cell in the Cloud Codeathon  Team Lead  • Led a team at a bioinformatics codeathon held at the New York Genome Codeathon	January 15-17, 2020 New York, NY
• Led a team at a bioinformatics codeathon neighbor the New York Genome C  NCBI RNA-seq in the Cloud Codeathon	Lenter March 11-13, 2019

Team Lead • Led a team at a bioinformatics codeathon held at the University of North Carolina, Chapel Hill