MATTHEW N. BERNSTEIN

July 5, 2025

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EDUCATION

Ph.D., Computer Sciences University of Wisconsin – Madison	Aug. 2019 Madison, WI
M.S., Computer Sciences University of Wisconsin – Madison	Dec. 2015 Madison, WI
B.S. <i>Magna Cum Laude</i> , Computer Science University of Notre Dame	May 2013 South Bend, IN
Work Experience	
Somite AI Principal Computational Scientist	Boston, MA June 2025 - Present
Stellaromics Staff Scientist, Machine Learning	Boston, MA July 2024 – June 2025
Immunitas Therapeutics Principal Scientist, Computational Biology Senior Scientist, Computational Biology Scientist, Computational Biology	Waltham, MA Oct. 2023 – July 2024 Sept. 2022 – Oct. 2023 March 2022 – Sept. 2022
Morgridge Institute for Research Postdoctoral Fellow	Madison, WI 2019 – 2022
University of Wisconsin – Madison Research Assistant	Madison, WI 2014 - 2019
Amazon Software Development Engineering Intern	Seattle, WA Summer 2014
Amazon Software Development Engineering Intern	New York, NY Summer 2013
Space and Naval Warfare Systems Command (SPAWAR) Research Intern	San Diego, CA Summer 2012

PEER-REVIEWED PUBLICATIONS & PREPRINTS

† corresponding author, * co-first author, ‡ co-second author

- 1. **Bernstein, M.N.***, Scott, D.*, Hession, C.C., Nieuwenhuis, T., Gerritsen, J., Tabrizi, S., Nandivada, V., Huggins, M.A., Duan, M., Malu, S., Tang, M. (2023). Monkeybread: A Python toolkit for the analysis of cellular niches in single-cell resolution spatial transcriptomics data. *bioRxiv*.
- 2. Zhang, J., Webster, S., Duffin, B., **Bernstein, M.N.**, Steill, J., Swanson, S., Forsberg, M.H., Bolin, J., Brown, M.E., Majumder, A., Capitini, C.M., Stewart, R., Thomson, J.A., Slukvin, I.I. (2023). Generation of anti-GD2 CAR macrophages from human pluripotent stem cells for cancer immunotherapies. *Stem Cell Reports*, 2(14), 585-596.
- 3. **Bernstein, M.N.**, Prasad, M., Ni, Z., Brown, J., Mohanty, C., Stewart, R., Newton, M.A., Kendziorski, C. (2022). SpatialCorr: Identifying gene sets with spatially varying correlation structure. *Cell Reports Methods*, 2(12), 100369.

- 4. Simonett, S.P., Shin, S., Herrinng, J.A., Bacher, R., Smith, L.A., Dong, C., Rabaglia, M., Stapleton, D., Schueler, K., Choi, J., Bernstein, M.N., Turkewitz, D.R., Perez-Cerventes, C., Spaeth, J., Stein, R., Tessem, J.S., Kendziorski, C., Keles, S., Moskowitz, I.P., Keller, M.P., Attie, A.D. (2021). Identification of direct transcriptional targets of Nfatc2 that promote β -cell proliferation in human islets. *The Journal of Clinical Investigation*, 131(21), e144833.
- 5. **Bernstein, M.N.**, Dewey, C.N. (2021). Annotating cell types in human single-cell RNA-seq data with CellO. *STAR Protocols*, 2(3), 100705.
- 6. Nimkulrat, S.D., **Bernstein, M.N.**, Ni, Z., Brown, J., Kendziorski, C., Blum, B. (2021). The Anna Karenina model of β -cell maturation in development and their dedifferentiation in type 1 and type 2 diabetes. *Diabetes*, 70(9), 2058-2066.
- 7. **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).
- 8. **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.
- 9. 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.
- 10. **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).
- 11. **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.

TALKS AND SEMINARS

Webinar hosted by Watershed Bio From Data to Discovery: Computational Tools for the Analysis of Spatial Transcriptomics Data	March 26, 2025 Virtual
Panoramics – A Vision, Seminar on dimensionality reduction in single-cell analysis Assessing the utility of data visualizations based on dimensionality reduction	Jan. 20, 2025 Toronto, ON
Workshop on Spatial Data Analysis at the Festival of Genomics Software tools for the analysis of cellular niches in the tissue microenvironment	June 13, 2024 Boston, MA
Spatial Biology US 2024 (Oxford Global) Software tools for the analysis of cellular niches in the tissue microenvironment	June 11, 2024 Boston, MA
Cohere for AI Community Talk A primer on variational autoencoders and their applications in single-cell genomics	April 22, 2024 Virtual
Workshop on AI for Precision Health & Precision Medicine Annotating cell types and cell states in single-cell omics data: A challenging problem	Nov. 16, 2023 Cambridge, MA
Spatial Biology for Drug Development (Hanson Wade) Software tools for the analysis of cellular niches in spatial transcriptomics data	Oct. 12, 2023 Boston, MA
Spatial Biology US 2023 (Oxford Global) Identifying gene sets with spatially varying correlation structure	June 8, 2023 Boston, MA
Discovery Seminar Series (UW-Madison) Unleashing the untapped potential of public genomics data	Oct. 19, 2021 Madison, WI

Computation and Informatics in Biology and Medicine Seminar (UW-Madison) Unleashing the untapped potential of public genomics data		March 23, 2021 Virtual
Workshop on Computational Advances for Single-Cell Omics Data Ar CellO: Comprehensive and hierarchical cell type classification of human cells with the C	•	Dec. 12, 2020 Virtual
International Conference on Intelligent Systems for Molecular Biology MetaSRA: Normalized sample-specific metadata for the Sequence Read Archive	y (ISMB)	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 (UW-Towards cell type prediction with public RNA-seq data	Madison)	Jan. 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		
American Society for Hematology Annual Meeting (ASH)	San Diego,	CA, Dec. 8-12, 2023
Great Lakes Bioinformatics Conference	Virtu	ıal, May 11-13, 2021
Cold Spring Harbor Laboratory conference on Biological Data Science	Vii	ctual, Nov. 4-6, 2020
International Conference on Intelligent Systems for Molecular Biolog	y (ISMB) Virt	ual, July 13-16, 2020
Great Lakes Bioinformatics Conference	Madison, V	WI, May 19-22, 2019
RNA-Seq Summit (Hanson Wade) Sa	ın Francisco, C	CA, Apr. 25-27, 2017
National Library of Medicine Informatics Training Conference	Columbus, (OH, June 27-28 2016
AWARDS AND FELLOWSHIPS NIH/BD2K Young Investigator Travel Scholarship International Conference on Intelligent Systems for Molecular Biology		July 6-10, 2018 Chicago, IL
Best Plenary Talk 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
TEACHING		
Guest lecture - Workshop on RNA-seq Computational methods for the analysis of spatial transcriptomics data		July 26, 2024 Montgomery College
Guest lecture, STAT 877 - Statistical Methods for Molecular Biology Visualizing high dimensional data with dimensionality reduction	University of	Sept. 22, 2020
Guest lecture, STAT 877 - Statistical Methods for Molecular Biology		Wisconsin – Madison
Cellular phenotyping with public, heterogeneous RNA-seq data	University of	Wisconsin – Madison Feb. 7, 2019 Wisconsin – Madison
Primary lecturer, CS 302 - Introduction to Programming	Aı	Feb. 7, 2019

Spatial Biology US 2023 (Oxford Global)

June 8-9, 2023

Panel participant

Boston, MA

• Invited panel member to discuss "Overcoming The Challenges Of Data Processing In Spatial Omics"

Immuno-Oncology Xchange (hubXchange), East Coast 2023

May 23, 2023

Led Round Table Discussion

Woburn, MA

• Led a round table discussion on the applications of machine learning for drug target discovery

NCBI Single-cell in the Cloud Codeathon

Jan. 15-17, 2020

Team Lead

New York, NY

• Led a team at a bioinformatics codeathon held at the New York Genome Center

NCBI RNA-seq in the Cloud Codeathon

Mar. 11-13, 2019

Team Lead

Chapel Hill, NC

· Led a team at a bioinformatics codeathon held at the University of North Carolina, Chapel Hill

SERVICE

Journal Reviewer Ongoing

- Bioinformatics (1 review)
- BMC Bioinformatics (4 reviews)
- Breast Cancer Research and Treatment (1 review)
- Cell Reports Methods (1 review)
- Genes (1 review)
- Journal of Computational Biology (1 review)
- Nature Biotechnology (1 review)
- Nature Communications (2 reviews)
- Nucleic Acids Research (1 review)
- STAR Protocols (1 review)

Review Editor for Frontiers in Genetics

2021-2022

Club Leader, After-school Computer Programming Club

Spring 2015

Stephen's Point Elementary School, Madison, WI

- Led an after-school computer programming club for 4th and 5th graders
- Taught the Scratch computer programming language